This instruction / checklist is intended for use in upgrading your Quality Management System for the transition from ISO 9001:2015 version to the AS 9100 D revision for Quality management systems used in the aviation, space, and defense industries.

The above Quality Management Systems are compatible with each other and have common requirements.

In the AS 9100 D and ISO 9001:2015 standards the requirements are described in:

- Clause 4 Context of the organization
- Clause 5 Leadership
- Clause 6 Planning
- Clause 7 Support
- Clause 8 Operation
- Clause 9 Performance evaluation
- Clause 10 Improvement

You have the ISO 9001:2015 version in place and now have the objective of upgrading the system to the 2016 Rev D revision. The good news is that since you are familiar with formal management systems, this initiative will be relatively straightforward.

Essentially, the documentation package for the management system will contain:

- One condensed Manual to introduce the documented information required for AS 9100 D.
- A group of procedure/system documents in your QMS with updates to reflect a document numbering system related to the new clause numbers and to incorporate the upgrades for AS 9100 D requirements,
- A group of forms and attachments needed for the documented information and systems.

The documentation will need to be reviewed, upgraded, and implemented. The first step is to assign a person responsible for the QMS, such as with a Management Representative to become familiar with the changes for the 2016 version of the AS 9100 D standard. Visit <u>the9100store.com</u> for training materials, resources, and information on quality management systems requirements.

The following table with detailed instructions focuses on the areas of the documentation required for the AS 9100 D quality management system. As you undertake the task of upgrading your quality management system from the ISO 9001:2015 version to the 2016 version, note that the intent of the main clauses is shown in blue font, and in the first left hand column of the instructions, the clause numbers highlighted in green indicates where specific AS 9100 D additions are made to ISO 9001:2015. Keep in mind that while you need to focus on the new requirements of AS 9100 D, your company now has an opportunity to review the exiting ISO 9001:2015 QMS and improve the system while incorporating the AS 9100 D requirements.

Use a copy of the AS 9100 D standard along with this instruction to pinpoint for your organization the areas that need attention. You may want to make notes and add comments in the space available to the right and the left of the column for reference documentation. Use the upgrade checklist section on the right side of the table to assign the responsibility for the upgrade and to follow up on its completion.

ISO 9001:2015 to AS9100D - QMS Upgrade Instructions / Checklist

			Achieve improvement.		
	This sub-clause focuses on the planning		In P-600, review / upgrade how you plan the actions		
6.1.2	of actions to be taken to address risks	Procedures	to address risks and opportunities, how you integrate		
	and opportunities.		and implement them in the QMS, and how you		
			evaluate the effectiveness of the actions taken.		
			See procedure <u>P-612</u> and related forms <u>F-612-001</u> , \overline{F}_{-612}		
			<u>F-612-002, and F-610-001 & F-810-002</u> for more		
	Quality objectives and the planning to		information on the risk management process. Review / upgrade your system for establishing		
6.2	achieve them applies to both standards		quality objectives at the relevant functions, levels		
0.2	where objectives at relevant functions,		and processes and the planning to achieve them.		
	levels, and processes are included.		Refer to specific requirements in clause 6.2.1 a) thru		
	levels, and processes are included.		g) and clause 6.2.2 a) thru e)		
	Planning for changes applies to both		Include the process for the planning of changes to		
6.3	standards where needed QMS changes		the QMS and for carrying them out in a planned and		
010	are determined and carried out in a		systematic way. Refer to specific requirements in		
	planned manner.		clause 6.3 s a) thru d) dealing with the purpose of		
			change, QMS integrity, resources, and responsibility.		
7	QMS. This section covers the resources th	at support the QI	ide the resources needed to establish, implement, main MS and include people, infrastructure, environment for th	ne operation of pro	ocesses,
7	QMS. This section covers the resources the monitoring and measuring resources, and methods for communication among the per	at support the QI organizational kn sonnel along wit	ide the resources needed to establish, implement, main MS and include people, infrastructure, environment for the owledge. In addition, competence, awareness of the h h systems for documented information need to be deterr	ne operation of pro uman resources /	ocesses,
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7 7.1 7.1.1	QMS. This section covers the resources the monitoring and measuring resources, and methods for communication among the per Clause 7, Support is a requirement in	at support the QI organizational kn rsonnel along wit Documented information	ride the resources needed to establish, implement, main MS and include people, infrastructure, environment for the owledge. In addition, competence, awareness of the h h systems for documented information need to be detern Review your existing system for the management of resources. Review / upgrade the information (in a document P- 710) that outlines the process for managing the resources of infrastructure and environment for the operation of the processes. Review your statement indicating that when determining and providing resources, consideration is given to capabilities, constraints, and external provision. Review the methods required to provide the persons needed for an effective QMS and for the operation and control of the processes. (See also 7.2 competence and 7.3 awareness).	ne operation of pro uman resources /	ocesses,

ISO 9001:2015 to AS9100D - QMS Upgrade Instructions / Checklist

			organizational knowledge (7.1.6) in this document.	
	In AS 9100 D, the periodic review of the		In P-720, consider the periodic review of the	
<mark>7.2</mark>	necessary competence of people is required to be considered.		necessary competence of the people.	
	required to be considered.		Refer to clause 7.3 and in P-720 review the	
7.3				
1.3	In AS 9100 D, new awareness items		awareness requirements a) thru d). In P-720 include the new requirements for:	
<mark>7.3</mark>	e) through h) are required for persons		 relevant QMS documented information, and 	
<mark>7.3</mark>	doing work for the company.			
	doing work for the company.		changes thereto,	
			 their contribution to product or service conformity, 	
			their contribution to product safety,	
			the importance of ethical behavior.	
7.4		Desselves	Review / upgrade the information (in a document P-	
7.4		Procedure	740) that outlines the process for both internal and	
			external communications of quality matters. Refer to	
			7.4 a) thru e) and include the information on what,	
			when, with whom, how, who will be communicating.	
7 4	In AS 9100 D, communication includes		Ensure that your communication system includes	
<mark>7.4</mark>	internal and external feedback		internal and external feedback relevant to the QMS.	
75	In AS 9100 D and ISO 9001:2015,	During	Review /upgrade the information (in a document P-	
7.5	'Documented Information' replaces	Procedure	750) that outlines the process for the control of	
	documented procedure and record.		documented information.	
	Documented procedure is now expressed		Incorporate a document numbering system related to	
	as a requirement to maintain documented		the clause numbers.	
	information. Record is now expressed as		Ensure that documented procedures for Control of Documents and Control of Records are included in	
	a requirement to retain documented information.		P-750, Control of documented information.	
	In AS 9100 D, a note clarifies approval		In P-750, clarify that approval implies authorized	
7.5.2	implications.		persons and approval methods are identified for the	
<mark>1.J.Z</mark>			relevant types of necessary documented information.	
			In P-750 include the method to adequately protect	
7.5.3.1			documented information from loss of confidentiality,	
7.3.3.1			improper use, or loss of integrity.	
			Refer to clause 7.5.3.2 and in P-750 review the	
7.5.3.2			requirements a) thru d).	
	In AS 9100 D, a requirement for control of		In P-750 include the new requirement for a system to	
7.5.3.2	unintended use of obsolete documents is		prevent the unintended use of obsolete documented	
	included at item e).		information by removal or by suitable identification or	
			controls if retained for any purpose.	
	In AS 9100 D, electronically managed		In P-750 and when documented information is	
7.5.3.2	documented information is addressed.		managed electronically, data protection processes	

1.0 Purpose/Scope

1.1 This procedure describes the process used to determine and control the selection of key characteristics and critical items of assemblies, components, materials and processes and the usage of same.

2.0 Responsibilities and Authorities

2.1 The product / process engineer is responsible for the selection of all necessary key characteristics and critical items and determining the appropriate data to be collected and evaluated.

3.0 References and Definitions

3.1 Reference

3.1.1 This document addresses clause 8.1 g of the AS 9100 D standard, covering key characteristics and critical items.

3.2 Definitions

- 3.2.1 Key Characteristic The features of a material, process, or part whose variation has a significant influence on product fit, performance, service life, or producibility.
- 3.2.2 Critical Items Those items such as functions, parts, software, characteristics, processes having significant effect on the provision and use of the products and services. including safety, performance, form, fit, function, producibility, or service life.
- 3.2.3 Risk Analysis To determine the importance of, or value of a situation or condition. Broadly defined to include risk assessment, risk characterization, risk communication, risk management, and policy relating to risk.
- 3.2.4 Pareto Analysis A formal technique for identifying the changes that will provide the greatest benefits. A Pareto analysis is a sorted histogram with two features. First is the cumulative distribution curve. Second, the vital few are identified. The histogram is sorted with the greatest occurrences to the left and descending occurrences moving to the right.
- 3.2.5 Failure Mode Effects Critically Analysis (FMECA) A disciplined review technique that focuses on the development of products, materials and processes based upon prioritized actions and in depth analysis to reduce the risk of product failures, and the associated documentation of those actions and review process.
- 3.2.6 Flowdown Technique for identifying key characteristics and critical items for a product or assembly down to subassemblies, details or processes believed to have a variation effect on upper level key characteristics and critical items.

4.0 Resources

4.1 Is there any special equipment or software required? (i.e.; special analysis

Key characteristics and critical items

INSERT YOUR COMPANY LOGO/NAME HERE

F-810-002 Project Planning Worksheet

Planning Project Type					
Quality Plan	Quality Object	ctive Risk &	Opportunity		
Project Name:		Date:			
Project objective:					
Estimated time frai	me:				
Estimated start dat Project manager:	e:				
Team members:					
Project completion	date:				
Project completion Were objectiv Project status	/es met?	onal Terminat	ed On-hold		
Management appro	oval:				

Every version of the AS 9100 standard has advocated risk avoidance and risk management. The new AS 9100 Rev D standard continues to expect organizations to identify and address risks affecting compliance of products and services, resulting in improved customer satisfaction.

Besides identifying the risks, organizations should address opportunities for improvements and corrective actions based on the risk analysis.

Note that while nonconformity and corrective action are requirements of AS 9100 Rev D, the concept of preventive action can be addressed through a risk-based approach where risks are determined and actions to address risks and opportunities are taken.

This risk analysis exercise is intended to outline several approaches / options for the management of risk at your company.

To prepare for the change, it is time to begin understanding Risk-Based Thinking and begin looking at your processes in terms of risks.

Risk is defined as the combination of the probability of occurrence of harm and the severity of that harm.

When evaluating risk, it is helpful to address it using two (2) metrics or parameters:

- 1. Severity (if harm happens, how serious is the event)
- 2. Likelihood (what is the probability of a harmful event occurring)

Because this topic is so important, it will have an impact on your QMS.

Risk-Based Thinking



INSERT COMPANY NAME/LOGO HERE

AS 9100 Rev D - Quality Management Systems – from ISO 9001:2015 - The Gap Analysis Checklist

This gap analysis checklist is prepared for use in evaluating a Quality Management System (QMS) against the requirements of the new Aerospace standard. The AS 9100 Rev D standard includes the requirements of ISO 9001:2015 and specifies additional aviation, space, and defense (ASD) industry requirements.

In the checklist, each requirement is expressed as a question that the user (auditor / assessor) can use to evaluate your QMS capabilities. You will need to have copies of the AS 9100 D and ISO 9001:2015 standards to use along with this checklist so that, if required, you can refer to the requirements and the clarification sections of Annex A.

While the structure of the AS and ISO standards are the same when comparing the contents, the additional ASD requirements are highlighted in vellow in the relevant sections of the checklist and the intent of the main clauses of the new standard is shown in blue font.

After you have prepared an audit schedule, and assigned responsibility to your auditors for different areas or processes to audit, copy each section of the checklist for the auditors working with that section. As you work through the checklist take notes on what is in place, and what needs to be developed.

In the space for 'currently in place', list or reference the procedures or other documents, or evidence that you have reviewed and that will provide information for the new QMS. Take notes on the status of the documents, that is, will they need to be revised for the new system, or can they be used as is? Also note where processes are in place, but documentation is needed. Focus on what is in place, and what needs to be developed.

While you do want to know if documented information is in place and if procedures and processes are being complied with, compliance is not your main focus for this audit. Remember that the final outcome of this audit should be a list of things that your company needs to do to comply with AS 9100 Rev D.

	QUALITY MANAGEMENT SYSTEMS REQUIREMENTS	Currently in Place	Compliant YES / NO?	If No - % Completed	Items Needed
4	CONTEXT OF THE ORGANIZATION				
Intend of clause	This first clause introduces two sub-clauses relating to the context of the organization, (1) understanding the organization and its context and (2) understanding the needs and expectations of interested parties. Together they require that you determine the issues and requirements that can impact on the planning of the Quality Management System (QMS). In addition, the scope of the QMS and the QMS processes along with their applicability and interactions need to be determined.				

IAQG-Oct-2016 - Audit conducted by:

Date: to