ISO 9001:2000 - CMMI v1.1 Mappings

It is always difficult to determine the appropriate granularity of maps between models. Mapping at a high level may not provide enough insight into similarities and differences. Mapping at a very low level, on the other hand, results in an overwhelming number of connections which also fails to properly illuminate model correspondence.

The mappings presented here address the middle ground. Each ISO 9001 "shall" statement has been mapped to a CMMI practice, using only the most prominent correspondence. If an ISO "shall" statement strongly maps to a CMMI specific practice, we do not indicate mappings to other specific practices that may show some weaker correspondence. The map thus serves as an indicator of correspondence rather than as an implementation guideline.

One should keep in mind that this is a many-to-many mapping, meaning that one ISO statement may correspond to more than one CMMI specific or generic practice, and vice versa.

As with all mappings, it is subjective. Although maps are convenient, they cannot replace an understanding of the frameworks being mapped. Stretching the correspondence may be counterproductive and misleading.

Mapping: ISO 9001:2000 to CMMI

Tables below show the mapping of each ISO 9001:2000 section to the CMMI.

Mapping is done at the "shall-level". Verbatim text from the ISO standard is maintained only in the titles, all other ISO text is replaced with keyword phrases corresponding to the ISO requirements.

"All" in the PA column means that the identified generic practices in every process area correspond to that ISO statement. Similarly, "All" in the Practice column means that all specific practices in the indicated process area correspond to the specific ISO statement or a group of statements. A judgment of the strength of the correspondence is shown as:

S – strong match; M – medium; W – weak

The tables do not indicate a mapping between CMMI generic and specific goals and ISO requirements. Although the goals can be mapped to ISO statements, such a mapping has no meaning in CMMI terms. In the CMMI, specific or generic practices are associated with corresponding goals. In other words, goals aggregate those practices to indicate some unique characteristics of the process area or its institutionalization and do not stand by themselves.

	ISO 9001:2000	CMMI PA	CMMI Practice	Strength	Comments
4.0	Quality Management System				
4.1	General requirements				
	Establish QMS				
	Identify processes	OPD	SP 1.1	S	
		OPF	SP 2.2		
		All	GP 2.1, 2.2,		
			2.3, 2.6, 2.8,		
			2.9		
	Manage using ISO standard	All	GP 2.1	S	
	Control outsourced processes	SAM	SP 2.2	S	
	Outsourced process control in QMS	SAM	SP 1.3	М	CMMI is not as strong
4.2	Documentation requirements				
4.2.1	General				
	Document quality manual	OPD	SP 1.1	М	CMMI satisfies mostly clause (d)
		All	GP 2.1		
4.2.2	Quality Manual				
	Establish quality manual	OPD	SP 1.1	S	
		All	GP 2.2		
4.2.3	Control of documents				
	Control required documents	All	GP 2.6	S	
	Control records				In process improvement "records" are known as "objective evidence"; this evidence is needed to show that a practice was implemented.
	Document control procedure	СМ	All	S	

	ISO 9001:2000	CMMI PA	CMMI Practice	Strength	Comments
4.2.4	Control of records				
	Records provide evidence of conformity				see 4.2.3
	Records identifiable				see 4.2.3
	Record control procedure	All	GP 2.6	S	
		СМ	GP 2.2		

	ISO 9001:2000	СММІ РА	CMMI Practice	Strength	Comments
5.0	Management responsibility				
5.1	Management commitment				
	Provide evidence of commitment	All	GP 2.1	М	CMMI is weaker
5.2	Customer focus				
	Determine customer	All	GP 2.7	М	CMMI not explicitly aimed at
	requirements	RD	SP 1.1-1,		customer satisfaction
			1.1-2, 1.2,		
			SP 2.1		
5.3	Quality policy				
	Top management quality policy responsibility	All	GP 2.1	S	CMMI is more specific on policy
		OPF	SP 1.1		content; GP 2.1 is for each PA and is therefore more detailed (at the PA level)
5.4	Planning				
5.4.1	Quality objectives				
	Objectives established at	OPF	SP 1.1	S	
	appropriate levels	OPP	SP 1.3		
		QPM	SP 1.1, 1.2,		
			1.3		
	Measurable objectives	OPP	SP 1.3	S	
		QPM	SP 1.1		
5.4.2	Quality management system planning				
	Plan to meet quality	OPD	All	S	
	objectives	All	GP 2.2, 3.1		

	ISO 9001:2000	CMMI PA	CMMI Practice	Strength	Comments
5.5	Responsibility, authority and communication				
5.5.1	Responsibility and authority				
	Top management defines responsibility	All	GP 2.4	S	
5.5.2	Management representative				Concept is not explicit in the CMMI
	Appoint member of management				
	Representative responsibility and authority	OPF	GP 2.4	W	
5.5.3	Internal communication				Concept is not explicit in the CMMI
	Establish communication	OPD	GP 2.1	W	CMMI is much weaker. No
	processes	OPF	SP 1.1		effectiveness
5.6	Management review				
5.6.1	General				
	Review QMS	All	GP 2.10	S	
	Assess improvement opportunities	OPF	SP 1.2, 1.3	S	
	Maintain records				Implicit in the CMMI, see 4.2.3 for explanation
5.6.2	Review input				
	Enumerates inputs for	All	GP 2.10	S	
	Teviews	PMC	SP 1.6, 1.7		
			SP 2.1, 2.2,		
			2.3		
5.6.3	Review output				
	Enumerates outputs from	All	GP 2.10	S	
	reviews	PMC	SP 1.6, 1.7		
			SP 2.1, 2.2,		
			2.3		
L	1	4	1	1	

	ISO 9001:2000	CMMI PA	CMMI Practice	Strength	Comments
6.0	Resource Management				
6.1	Provision of resources				
	Determine resource needs	All	GP 2.3	S	
6.2	Human resources				
6.2.1	General				
	Staff has needed skills	All	GP 2.5	S	
6.2.2	Competence, awareness and training				
	Ensure competence, provide training, keep records	OT	SP 1.1, 1.2, 1.3, 1.4, SP 2.1, 2.2, 2.3 SP 1.3	S	
6.3	Infrastructure				
	Provide services and equipment	OEI	SP 1.2	S	This is in the IPPD
6.4	Work environment				
	Maintain environment to meet requirements	PP OEI	SP 2.4 SP 1.2	М	Not explicitly covered in the CMMI. PP address facilities; OEI addresses "integrated work environment"

	ISO 9001:2000	CMMI PA	CMMI Practice	Strength	Comments
7.0	Product realization				
7.1	Planning product realization				
	Develop needed	OPD	SP 1.1, 1.2,	S	
	processes		1.3		
	Planning is consistent with	OPD	SP 1.1	S	
	other processes	All	GP 2.2, 3.1		
	Address objectives and	PP	SP 1.1, 1.2,	S	SP 3.1 – 3.3 are not specifically
	verification		1.3, 1.4		required
			SP 2.1, 2.2,		
			2.3, 2.4, 2.5,		
			2.6, 2.7		
		QPM	1.1		
	Plans in appropriate	PP	SP 2.7	S	
	format	IPM	SP 1.1, 1.3,		
			1.4,		
7.2	Customer—related processes				
7.2.1	Determination of requirements related to the product				
	Determine customer and	RD	SP 1.1-2, 1.2,	S	
	other requirements		SP 2.1, 2.2,		
			2.3,		
			SP 3.1, 3.2		
		REQM	SP 1.1		
		TS	SP 1.2		

	ISO 9001:2000	CMMI PA	CMMI Practice	Strength	Comments
7.2.2	Review of requirements related to the product	RD	SP 3.5	S	ISO is less specific than CMMI
	Organization reviews	RD	GP 2.10	S	
	requirements		SP 1.2, 3.5		
		VER	SP 2.2		
	Review before commitment				
	Requirements are defined	REQM	SP 1.1, 1.2,	S	
			1.5		
	Records are kept				Not specifically required by the CMMI
	Confirm understanding of requirements	RD	SP 2.1, 3.5	S	
	Keep documents current when requirements change	REQM	SP 1.3	S	
7.2.3	Customer communication				
	Communicate effectively	RD	GP 2.7	М	
	with customers	IPM	SP 2.1, 2.2,		
			2.3		
		MA	SP 2.4		CMMI is weaker (less explicit)
		REQM	SP 1.2		

	ISO 9001:2000	CMMI PA	CMMI Practice	Strength	Comments
7.3	Design and development				
7.3.1	Design and development planning				
	Plan development	PP	SP 1.1, 1.2,	S	
		IPM	SP 1.1, 1.3,		
			1.4		
		VER	SP 1.1		
		VAL	SP 1.1		
	Determine development	TS	SP 1.1, 1.2,	S	
	stages and verifications		1.3,		
			SP 2.1, 2.2,		
			2.3, 2.4,		
			SP 3.1, 3.2		
		PI	SP 1.1, 1.2,		
			1.3,		
			SP 2.1, 2.2		
	Manage interfaces	IPM	SP 2.1, 2.2,	S	Note: " 3.x and 4.x are in the IPPD
			2.3,		
			SP 3.1, 3.2,		
			SP 4.1, 4.2,		
			4.3		
	Update plans during	PP	All	S	
	development	IPM	SP 1.3		
7.3.2	Design and development inputs				
	Determine inputs to	RD	SP 1.1, 1.2,	S	
	aevelopment processes		SP 2.1		
			SP 3.2		
	Inputs include product,	RD	SP 1.1, 1.2,	S	
	regulatory, and other requirements		SP 2.1		

	ISO 9001:2000	СММІ РА	CMMI Practice	Strength	Comments
7.3.2	Design and development inputs (continued)				
	Review inputs	RD	SP 3.3, 3.4,	S	
			3.5		
			GP 2.7, 2.10		
	Requirements are	RD	SP 3.3, 3.4,	S	
	consistent and clear		3.5		
7.3.3	Design and development outputs				
	Outputs are verifiable	TS	SP 2.2-3	S	More general
		IPM	SP 1.1		Nore general
	Outputs approved				
	Development requirements are met	TS	All	S	
7.3.4	Design and development review				
	Development reviewed and evaluated	PMC	SP 1.6, 1.7	S	
	Appropriate functions participate in reviews	PMC	GP 2.7	S	
	Records of review are kept				"evidence" in the CMMI
7.3.5	Design and development verification				
	Ensure requirements are	VER	SP 1.1, 1.2,	S	
	met		1.3,		
			SP 2.1, 2.2,		
			2.3,		
			SP 3.1, 3.2		
	Keep verification records				see 4.2.3 for explanation
7.3.6	Design and development validation				
	Validation follows plans	VAL	SP 1.1, 1.2,	S	
			1.3,		
			SP 2.1, 2.2		
	Validate before delivery				Not explicit in the CMMI
	Keep validation records				see 4.2.3 for explanation

	ISO 9001:2000	CMMI PA	CMMI Practice	Strength	Comments
7.3.7	Control of design and				
	Identify changes	TS &	GP 2 6	S	
		PI			
	Review and approve changes	СМ	All	S	
	Evaluate effect of changes	СМ	All	S	
	Keep records of changes	CM	All	S	
7.4	Purchasing				
7.4.1	Purchasing process				
	Purchased product meets	SAM	GP 2.9	S	
	requirements	TS	SP 1.1, 1.2,		
			1.3		
	Control of supplier depends on product	SAM	SP 2.2	М	CMMI is weaker
	Suppliers selected based on ability	SAM	SP 1.2	S	
	Selection criteria established	SAM	SP 1.2	S	
	Records of evaluations kept	SAM	SP 1.2	М	CMMI is weaker
7.4.2	Purchasing information				
	Product requirements	SAM	SP 1.1	S	
	described		SP 2.1		
	Adequate requirements described	SAM	SP 1.3	S	
7.4.3	Verification of purchased product				
	Ensure product meets	SAM	SP 2.2, 2.3	S	
	requirements	VER	SP 3.1		
	Supplier site verification	SAM	SP 2.3	W	CMMI is not explicit

	ISO 9001:2000	CMMI PA	CMMI Practice	Strength	Comments
7.5	Production and service provision				
7.5.1	Control of production and service provision				
	Plan service provision	TS	SP 3.1, .32	М	CMMI is weaker
	Control addresses information, equipment, and activities	TS	GP 2.2, 2.3, 2.6, 2.8	М	CMMI is weaker; "post-delivery" not addressed in the CMMI
7.5.2	Validation of processes for production and service provision				
	Validate production & service processes	VAL	SP 1.1	S	
	Demonstrate ability to meet planned results	VAL	All	S	
	Establish review criteria	VAL	SP 1.2, 1.3,	S	
			SP 2.1, 2.2		
7.5.3	Identification and traceability				
	Identify products	СМ	SP 1.1,	S	
			SP 2.1, 2.2		
	Identify monitoring requirements	Ы	SP 3.1, 3.2,	S	
	Control traceability	PI	SP 3.1. 3.2	S	
	,	REQM	SP 1.4		
		СМ	SP 1.3,		
			SP 2.2,		
			SP 3.1		
7.5.4	Customer property				Not explicitly addressed in CMMI
	Exercise care				
	Identify property				
	Report damage				
7.5.5	Preservation of Product				Not explicitly addressed in CMMI
	Maintain conformity during delivery	PI	SP 3.4	М	CMMI is less explicit
	Preserve identification	PI	SP 3.4	М	CMMI is less explicit
	Preserve product component parts	PI	SP 3.4	М	CMMI is less explicit

	ISO 9001:2000	CMMI PA	CMMI Practice	Strength	Comments
7.6	Control and monitoring of measuring devices				
	Determine monitoring and	VER	GP 2.8	S	
	device needed	VAL	GP 2.8		
	Establish monitoring	MA	GP 2.1, 2.2,	S	
	processes		2.8, 2.9, 2.10		
	Calibrate measuring equipment				Not in the CMMI. CM practices can be used, where applicable
	Assess prior measurement results				Not in the CMMI; CM practices can be used, where applicable
	Take action on equipment				Not in the CMMI; CM practices can be used, where applicable
	Keep calibration records				Not in the CMMI; CM practices can be used, where applicable
	Confirm applicability of				Not in the CMMI; CM practices
	software				can be used, where applicable
	Confirm software before use				Not in the CMMI; CM practices can be used, where applicable

9.0 Maaa	surement, analysis			S	
and i	mprovement				
8.1 Gene	General				
Plan	Plan monitoring and		GP 2.2	S	
meas	surement		SP 1.1, 1.2,		
			1.3, 1.4		
Deter	mine methods and	QPM	SP 2.1, 2.2,	S	
lechn	techniques		2.3, 2.4		
8.2 Moni meas	toring and surement				
8.2.1 Cust	omer satisfaction				
Monit	tor customer	MA	SP 1.1, 1.2	М	CMMI not very strong
peree	perceptions		SP 2.2		Civiliti Hot very strong
		PMC	SP 1.5		
Defin	e methods for suring satisfaction				Not addressed in the CMMI
8.2.2 Interi	nal audit				
Cond	uct planned audits	OPF	SP 1.1, 1.2	S	
		PPQA	All		
Audit	s consider process	OPF	SP 1.1, 1.2	S	
impor	rtance	PPQA	GP 2.2		
Defin	e audit criteria	OPF	SP 1.1, 1.2	S	
Selec	t objective auditors				Not addressed in the CMMI
Don't	audit own output				Not addressed in the CMMI
Docu	mented procedure	OPF	GP 2.4	S	Process Improvement Plan
define	es audits	MA	SP 2.4		CIMIMI IS MORE explicit
		PPQA	GP 2.4		
Action	ns taken promptly	OPF	GP 2.1	S	
			SP 1.3		
		PPQA	GP 2.6		
Verify	/ actions taken	opf Ppqa	SP 2.1, 2.2 GP 2.9	S	CMMI is more detailed (SP 2.3, 2.4 not specifically required by ISO)

	ISO 9001:2000	CMMI PA	CMMI Practice	Strength	Comments
8.2.3	Monitoring and				
	measurement of process	ΔIJ	GP 28	6	
			GF 2.0	3	
	Demonstrate process	MA	GP 2.2	S	
	capability		SP 1.2, 1.3		
	Take corrective actions	QPM	SP 2.2, 2.3	S	
		PMC	SP 2.1, 2.2,		
			2.3		
8.2.4	Monitoring and measurement of product				
	Monitor product	VAL	SP 2.1, 2.2	S	
	characteristics	VER	SP 3.1, 3.2		
	Measure at appropriate	VER	SP 1.1, 1.3	S	
	stages		SP 2.1, 2.2,		
			2.3		
	Maintain conformity record	REQM	SP 1.1	S	
		SAM	SP 1.3		
		PPQA	SP 1.2		
		VAL	SP 1.3		
	Maintain release records				see 4.2.3 for explanation
	Don't release until product realization plans are implemented	СМ	SP 3.2	М	CMMI is weaker
8.3	Control of				I
	Identify and control	СМ			Not explicitly required by CMMI
	nonconforming product				
	Define control of				
	Dealing with	PMC	SP 2.1. 22.		
	nonconforming product		2.3		
	Keep records of				
	Re-verify corrected				
	nonconformance				
	Take action after delivery				

	ISO 9001:2000	CMMI PA	CMMI Practice	Strength	Comments
8.4	Analysis of data				
	Collect data on QMS effectiveness	MA	SP 2.2, 2.3, 2.4	S	
		OPF All	SP 1.3 GP 3.2		
	Include monitoring data				see 4.2.3 for explanation
	Analyze conformance and customer satisfaction	MA RD	SP 2.2 SP 1.1, 1.2,	М	Customer satisfaction is not explicit
			2.1,		
			SP 3.1, 3.2,		
			3.3, 3.4		
		QPM	SP 1.4		
		CAR	SP 1.1, 1.2		
		SAM	SP 2.2		
8.5	Improvement				
8.5.1	Continual improvement				
	Improve QMS	OPF	SP 1.1, 1.3	S	
	effectiveness	OID	SP 1.1		
		MA	SP 1.1, 1.2,		
			1.4,		
			SP 2.1, 2.2		
8.5.2	Corrective action				
	Eliminate causes of nonconformities Take appropriate actions	OPF	SP 2.1, 2.2, 2.3	S	
	Documented procedure defines corrective actions	PMC	SP 2.1, 2.2, 2.3	S	
8.5.3	Preventive action				
	Determine action to prevent nonconformity	OPF	SP 2.4	S	Not in the CMMI
	defines preventive actions	CAR	SP 1.1, 1.2 SP 2.1, 2.2,	IVI	more sophisticated (CAR is Level 5 – staged)
			2.3		

Mapping: CMMI to ISO 9001:2000

The CMMI to ISO 9001 mapping was mechanically constructed from the ISO 9001 to CMMI map rather than from an independent analysis.

The first table shows the per-Process Area mapping from the CMMI to individual ISO sections. The second table shows the mapping from CMMI generic practices to the corresponding ISO sections. A blank in the ISO column indicates that there is no correspondence between the frameworks.

Goal	Specific Practice	Description	ISO 9001:2000
		Organizational Process Focus	
SG 1		Determine Process-Improvement Opportunities	
	SP 1.1-1	Establish Organizational Process Needs	5.3, 5.4.1, 5.5.3,
			5.6.1, 8.2.2, 8.5.1
	SP 1.2-1	Appraise the Organization's Processes	5.6.1, 8.2.2
	SP 1.3-1	Identify the Organization's Process Improvements	8.4, 8.5.1
SG 2		Plan and Implement Process-Improvement Activities	
	SP 2.1-1	Establish Process Action Plans	8.2.2, 8.5.1
	SP 2.2-1	Implement Process Action Plans	4.1, 8.2.2, 8.5.1
	SP 2.3-1	Deploy Organizational Process Assets	8.5.1
	SP 2.4-1	Incorporate Process-Related Experiences into the Organizational Process Assets	8.5.3
		Organizational Process Definition	
SG 1		Establish Organizational Process Assets	
	SP 1.1-1	Establish Standard Processes	4.1, 4.2.1, 4.2.2,
			5.4.2, 7.1
	SP 1.2-1	Establish Life-Cycle Model Descriptions	5.4.2, 7.1
	SP 1.3-1	Establish Tailoring Criteria and Guidelines	5.4.2, 7.1
	SP 1.4-1	Establish the Organization's Measurement Repository	5.4.2
	SP 1.5-1	Establish the Organization's Process Asset Library	5.4.2
		Organizational Training	
SG 1		Establish an Organizational Training Capability	
	SP 1.1-1	Establish the Strategic Training Needs	6.2.2
	SP 1.2-1	Determine Which Training Needs Are the Responsibility of the Organization	6.2.2
	SP 1.3-1	Establish an Organizational Training Tactical Plan	6.2.2
	SP 1.4-1	Establish Training Capability	6.2.2
SG 2		Provide Necessary Training	
	SP 2.1-1	Deliver Training	6.2.2
	SP 2.2-1	Establish Training Records	6.2.2

Goal	Specific Practice	Description	ISO 9001:2000
	SP 2.3-1	Assess Training Effectiveness	6.2.2
		Organizational Process Performance	
SG 1		Establish Performance Baselines and Models	
	SP 1.1-1	Select Processes	
	SP 1.2-1	Establish Process Performance Measures	
	SP 1.3-1	Establish Quality and Process-Performance Objectives	5.4.1
	SP 1.4-1	Establish Process Performance Baselines	
	SP 1.5-1	Establish Process Performance Models	
		Organizational Innovation and Deployment	
SG 1		Select Improvements	
	SP 1.1-1	Collect and Analyze Improvement Proposals	8.5.1
	SP 1.2-1	Identify and Analyze Innovations	
	SP 1.3-1	Pilot Improvements	
	SP 1.4-1	Select Improvements for Deployment	
SG 2		Deploy Improvements	
	SP 2.1-1	Plan the Deployment	
	SP 2.2-1	Manage the Deployment	
	SP 2.3-1	Measure Improvement Effects	
		Project Planning	
SG 1		Establish Estimates	
	SP 1.1-1	Estimate the Scope of the Project	7.1, 7.3.1
	SP 1.2-1	Establish Estimates of Work Product and Task Attributes	7.1, 7.3.1
	SP 1.3-1	Define Project Life Cycle	7.1, 7.3.1
	SP 1.4-1	Determine Estimates of Effort and Cost	7.3.1
SG 2		Develop a Project Plan	
	SP 2.1-1	Establish the Budget and Schedule	7.1, 7.3.1
	SP 2.2-1	Identify Project Risks	7.1, 7.3.1
	SP 2.3-1	Plan for Data Management	7.1, 7.3.1
	SP 2.4-1	Plan for Project Resources	6.4, 7.1, 7.3.1

Goal	Specific Practice	Description	ISO 9001:2000
	SP 2.5-1	Plan for Needed Knowledge and Skills	7.1, 7.3.1
	SP 2.6-1	Plan Stakeholder Involvement	7.1, 7.3.1
	SP 2.7-1	Establish the Project Plan	7.1, 7.3.1
SG 3		Obtain Commitment to the Plan	
	SP 3.1-1	Review Plans that Affect the Project	7.3.1
	SP 3.2-1	Reconcile Work and Resource Levels	7.3.1
	SP 3.3-1	Obtain Plan Commitment	7.3.1
		Project Monitoring and Control	
SG 1		Monitor Project Against Plan	
	SP 1.1-1	Monitor Project Planning Parameters	
	SP 1.2-1	Monitor Commitments	
	SP 1.3-1	Monitor Project Risks	
	SP 1.4-1	Monitor Data Management	
	SP 1.5-1	Monitor Stakeholder Involvement	8.2.1
	SP 1.6-1	Conduct Progress Reviews	5.6.2, 5.6.3, 7.3.4
	SP 1.7-1	Conduct Milestone Reviews	5.6.2, 5.6.3, 7.3.4
SG 2		Manage Corrective Action to Closure	
	SP 2.1-1	Analyze Issues	5.6.2, 5.6.3,
			8.2.3, 8.3, 8.5.2
	SP 2.2-1	Take Corrective Action	5.6.2, 5.6.3,
			8.2.3, 8.3, 8.5.2
	SP 2.3-1	Manage Corrective Action	5.6.2, 5.6.3,
			0.2.3, 0.3, 0.5.2
		Supplier Agreement Management	
SG 1		Establish Supplier Agreements	
	SP 1.1-1	Determine Acquisition Type	7.4.1, 7.4.2
	SP 1.2-1	Select Suppliers	7.4.1
	SP 1.3-1	Establish Supplier Agreements	4.1, 7.4.2, 8.2.4
SG 2		Satisfy Supplier Agreements	
	SP 2.1-1	Review COTS Products	7.4.2, 7.4.3

Goal	Specific Practice	Description	ISO 9001:2000
	SP 2.2-1	Execute the Supplier Agreement	4.1, 7.4.3, 8.4
	SP 2.3-1	Accept the Acquired Product	7.4.3
	SP 2.4-1	Transition Products	
		Integrated Project Management for IPPD	
SG 1		Use the Project's Defined Process	
	SP 1.1-1	Establish the Project's Defined Process	7.1, 7.3.1, 7.3.3
	SP 1.2-1	Use Organizational Process Assets for Planning Project Activities	
	SP 1.3-1	Integrate Plans	7.1, 7.3.1
	SP 1.4-1	Manage the Project Using the Integrated Plans	7.1, 7.3.1
	SP 1.5-1	Contribute to the Organizational Process Assets	
SG 2		Coordinate and Collaborate with Relevant	
		Stakeholders	
	SP 2.1-1	Manage Stakeholder Involvement	7.2.3, 7.3.1
	SP 2.2-1	Manage Dependencies	7.2.3, 7.3.1
	SP 2.3-1	Resolve Coordination Issues	7.3.1
SG 3		Use the Project's Shared Vision for IPPD	
	SP 3.1-1	Define Project's Shared-Vision Context	7.3.1
	SP 3.2-1	Establish the Project's Shared Vision	7.3.1
SG 4		Organize Integrated Teams for IPPD	
	SP 4.1-1	Determine Integrated Team Structure for the Project	7.3.1
	SP 4.2-1	Develop a Preliminary Distribution of Requirements to Integrated Teams	7.3.1
	SP 4.3-1	Establish Integrated Teams	7.3.1
		Risk Management	
SG 1		Prepare for Risk Management	
	SP 1.1-1	Determine Risk Sources and Categories	
	SP 1.2-1	Define Risk Parameters	
	SP 1.3-1	Establish a Risk Management Strategy	
SG 2		Identify and Analyze Risks	
	SP 2.1-1	Identify Risks	
	SP 2.2-1	Evaluate, Categorize, and Prioritize Risks	

Goal	Specific Practice	Description	ISO 9001:2000
SG 3		Mitigate Risks	
	SP 3.1-1	Develop Risk Mitigation Plans	
	SP 3.2-1	Implement Risk Mitigation Plans	-
		Integrated Teaming	
SG 1		Establish Team Composition	
	SP 1.1-1	Identify Team Tasks	
	SP 1.2-1	Identify Needed Knowledge and Skills	
	SP 1.3-1	Assign Appropriate Team Members	-
SG 2		Govern Team Operation	
	SP 2.1-1	Establish a Shared Vision	
	SP 2.2-1	Establish a Team Charter	
	SP 2.3-1	Define Roles and Responsibilities	-
	SP 2.4-1	Establish Operating Procedures	
	SP 2.5-1	Collaborate among Interfacing Teams	
		Quantitative Project Management	
SG 1		Quantitatively Manage the Project	
	SP 1.1-1	Establish the Project's Objectives	5.4.1, 7.1
	SP 1.2-1	Compose the Defined Process	5.4.1
	SP 1.3-1	Select the Subprocesses that Will Be Statistically Managed	5.4.1
	SP 1.4-1	Manage Project Performance	
SG 2		Statistically Manage Subprocess Performance	
	SP 2.1-1	Select Measures and Analytic Techniques	8.1
	SP 2.2-1	Apply Statistical Methods to Understand Variation	8.1, 8.2.3
	SP 2.3-1	Monitor Performance of the Selected Subprocesses	8.1, 8.2.3
	SP 2.4-1	Record Statistical Management Data	8.1
		Requirements Management	
SG 1		Manage Requirements	
	SP 1.1-1	Obtain an Understanding of Requirements	7.2.1, .7.2.2,
			8.2.4
	SP 1.2-2	Obtain Commitment to Requirements	7.2.3
	SP 1.3-1	Manage Requirements Changes	7.2.2

Goal	Specific Practice	Description	ISO 9001:2000
	SP 1.4-2	Maintain Bi-directional Traceability of Requirements	7.5.3
	SP 1.5-1	Identify Inconsistencies between Project Work and Requirements	7.2.2
		Requirements Development	
SG 1		Develop Customer Requirements	
	SP 1.1-1	Collect Stakeholder Needs	5.2, 7.2.1, 7.3.2,
			8.4
	SP 1.1-2	Elicit Needs	5.2, 7.2.1, 7.2.2,
			7.3.2, 8.4
	SP 1.2-1	Develop the Customer Requirements	5.2, 7.2.1, 7.3.2
SG 2		Develop Product Requirements	
	SP 2.1-1	Establish Product and Product-Component Requirements	5.2, 7.2.1, 7.2.2,
			7.3.2, 8.4
	SP 2.2-1	Allocate Product-Component Requirements	7.2.1
	SP 2.3-1	Identify Interface Requirements	7.2.1
SG 3		Analyze and Validate Requirements	
	SP 3.1-1	Establish Operational Concepts and Scenarios	7.2.1, 8.4
	SP 3.2-1	Establish a Definition of Required Functionality	7.2.1, 7.3.2, 8.4
	SP 3.3-1	Analyze Requirements	8.4, 7.3.2
	SP 3.4-3	Analyze Requirements to Achieve Balance	8.4, 7.3.2
	SP 3.5-1	Validate Requirements	7.2.2, 7.3.2
	SP 3.5-2	Validate Requirements with Comprehensive Methods	7.2.2, 7.3.2
		Technical Solution	
SG 1		Select Product-Component Solutions	
	SP 1.1-1	Develop Alternative Solutions and Selection Criteria	7.3.1, 7.3.3, 7.4.1
	SP 1.1-2	Develop Detailed Alternative Solutions and Selection Criteria	7.3.1, 7.3.3, 7.4.1
	SP 1.2-2	Evolve Operational Concepts and Scenarios	7.2.1, 7.3.1,
			7.3.3, 7.4.1
	SP 1.3-1	Select Product-Component Solutions	7.3.1, 7.3.3, 7.4.1
SG 2		Develop the Design	
	SP 2.1-1	Design the Product or Product Component	7.3.1, 7.3.3
	SP 2.2-3	Establish a Technical Data Package	7.3.1, 7.3.3
	SP 2.3-1	Establish Interface Descriptions	7.3.1, 7.3.3

Goal	Specific Practice	Description	ISO 9001:2000
	SP 2.3-3	Design Interfaces Using Criteria	7.3.1, 7.3.3
	SP 2.4-3	Perform Make, Buy, or Reuse Analyses	7.3.1, 7.3.3
SG 3		Implement the Product Design	
	SP 3.1-1	Implement the Design	7.3.1, 7.5.1
	SP 3.2-1	Develop Product Support Documentation	7.3.1, 7.5.1
		Product Integration	
SG 1		Prepare for Product Integration	
	SP 1.1-1	Determine Integration Sequence	7.3.1
	SP 1.2-2	Establish the Product Integration Environment	7.3.1
	SP 1.3-3	Establish Product Integration Procedures and Criteria	7.3.1
SG 2		Ensure Interface Compatibility	
	SP 2.1-1	Review Interface Descriptions for Completeness	7.3.1
	SP 2.2-1	Manage Interfaces	7.3.1
SG 3		Assemble Product Components and Deliver the	
		Product	
	SP 3.1-1	Confirm Readiness of Product Components for Integration	7.5.3
	SP 3.2-1	Assemble Product Components	7.5.3
	SP 3.3-1	Evaluate Assembled Product Components	7.5.3
	SP 3.4-1	Package and Deliver the Product or Product Component	7.5.5
		Verification	
SG 1		Prepare for Verification	
	SP 1.1-1	Select Work Products for Verification	7.3.1, 7.3.5, 8.2.4
	SP 1.2-2	Establish the Verification Environment	7.3.5
	SP 1.3-3	Establish Verification Procedures and Criteria	7.3.5, 8.2.4
SG 2		Perform Peer Reviews	
	SP 2.1-1	Prepare for Peer Reviews	7.3.5, 8.2.4
	SP 2.2-1	Conduct Peer Reviews	7.3.5, 8.2.4
	SP 2.3-2	Analyze Peer Review Data	7.3.5
SG 3		Verify Selected Work Products	
	SP 3.1-1	Perform Verification	7.2.2, 7.3.5,
			7.4.3, 8.2.4

Goal	Specific Practice	Description	ISO 9001:2000
	SP 3.2-2	Analyze Verification Results and Identify Corrective Action	7.3.5, 8.2.4
		Validation	
SG 1		Prepare for Validation	
	SP 1.1-1	Select Products for Validation	7.3.1, 7.3.6, 7.5.2
	SP 1.2-2	Establish the Validation Environment	7.3.6, 7.5.2
	SP 1.3-3	Establish Validation Procedures and Criteria	7.3.6, 7.5.2, 8.2.4
SG 2		Validate Product or Product Components	
	SP 2.1-1	Perform Validation	7.3.6, 7.5.2, 8.2.4
	SP 2.2-1	Analyze Validation Results	7.3.6, 7.5.2, 8.2.4
		Configuration Management	
SG 1		Establish Baselines	
	SP 1.1-1	Identify Configuration Items	4.2.3, 7.3.7,
			7.5.3, 8.3
	SP 1.2-1	Establish a Configuration Management System	4.2.3, 7.3.7, 8.3
	SP 1.3-1	Create or Release Baselines	4.2.3, 7.3.7,
			7.5.3, 8.3
SG 2		Track and Control Changes	
	SP 2.1-1	Track Change Requests	4.2.3, 7.3.7,
			7.5.3, 8.3
	SP 2.2-1	Control Configuration Items	4.2.3, 7.3.7,
			7.5.3, 8.3
SG 3		Establish Integrity	
	SP 3.1-1	Establish Configuration Management Records	4.2.3, 7.3.7,
			7.5.3, 8.3
	SP 3.2-1	Perform Configuration Audits	4.2.3, 7.3.7,
			8.2.4, 8.3
		Process and Product Quality Assurance	
SG 1		Objectively Evaluate Processes and Work Products	8.2.2
	SP 1.1-1	Objectively Evaluate Processes	8.2.2
	SP 1.2-1	Objectively Evaluate Work Products and Services	8.2.2, 8.2.4
SG 2		Provide Objective Insight	8.2.2

Goal	Specific Practice	Description	ISO 9001:2000
	SP 2.1-1	Communicate and Ensure Resolution of Noncompliance	8.2.2
	SP 2.2-1	Establish Records	8.2.2
		Measurement and Analysis	
SG 1		Align Measurement and Analysis Activities	
	SP 1.1-1	Establish Measurement Objectives	8.1, 8.2.1, 8.5.1
	SP 1.2-1	Specify Measures	8.1, 8.2.1, 8.2.3,
			8.5.1
	SP 1.3-1	Specify Data Collection and Storage Procedures	8.1, 8.2.3
	SP 1.4-1	Specify Analysis Procedures	8.1, 8.5.1
SG 2		Provide Measurement Results	
	SP 2.1-1	Collect Measurement Data	8.2.1, 8.4, 8.5.1
	SP 2.2-1	Analyze Measurement Data	8.4, 8.5.1
	SP 2.3-1	Store Data and Results	8.4
	SP 2.4-1	Communicate Results	7.2.3, 8.2.2
		Decision Analysis and Resolution	
SG 1		Evaluate Alternatives	
	SP 1.1-1	Establish Guidelines for Decision Analysis	
	SP 1.2-1	Establish Evaluation Criteria	
	SP 1.3-1	Identify Alternative Solutions	
	SP 1.4-1	Select Evaluation Methods	
	SP 1.5-1	Evaluate Alternatives	
	SP 1.6-1	Select Solutions	
		Organizational Environment for Integration	
SG 1		Provide IPPD Infrastructure	
	SP 1.1-1	Establish the Organization's Shared Vision	
	SP 1.2-1	Establish an Integrated Work Environment	6.3, 6.4
	SP 1.3-1	Identify IPPD-Unique Skill Requirements	6.2.2
SG 2		Manage People for Integration	
	SP 2.1-1	Establish Leadership Mechanisms	
	SP 2.2-1	Establish Incentives for Integration	
	SP 2.3-1	Establish Mechanisms to Balance Team and Home Organization Responsibilities	

Goal	Specific Practice	Description	ISO 9001:2000
		Causal Analysis and Resolution	
SG 1		Determine Causes of Defects	
	SP 1.1-1	Select Defect Data for Analysis	8.4, 8.5.3
	SP 1.2-1	Analyze Causes	8.4, 8.5.3
SG 2		Address Causes of Defects	
	SP 2.1-1	Implement the Action Proposals	8.5.3
	SP 2.2-1	Evaluate the Effect of Changes	8.5.3
	SP 2.3-1	Record Data	8.5.3

Generic Goal	Generic Practices	Description	ΡΑ	ISO 9001:2000
GG 1		Achieve Specific Goals		
	GP 1.1	Perform Base Practices		
GG 2		Institutionalize a Managed Process		
	GP 2.1	Establish an Organizational Policy	All OPD	4.1, 4.2.1, 5.1 5.5.3
			MA OPF	7.6 8.2.2
	GP 2.2	Plan the Process	AII TS MA	4.1, 4.2.2, 5.4.2, 7.1 7.5.1 7.6, 8.1, 8.2.3
	GP 2.3	Provide Resources	All TS	4.1, 6.1 7.5.1
	GP 2.4	Assign Responsibility	All OPF PPQA	5.5.1 8.2.2 8.2.2
	GP 2.5	Train People	All	6.2.1
	GP 2.6	Manage Configurations	All Pl TS	4.1, 4.2.3, 4.2.4 7.3.7 7.3.7, 7.5.1
	GP 2.7	Identify and Involve Relevant Stakeholders	All RD PMC	5.1 7.2.3, 7.3.2 7.3.4
	GP 2.8	Monitor and Control the Process	All TS	4.1, 8.2.3 7.5.1
			VER VAL MA	7.6 7.6 7.6
	GP 2.9	Objectively Evaluate Adherence	All MA PPQA	4.1 7.6 8.2.2

Generic Goal	Generic Practices	Description	РА	ISO 9001:2000
	GP 2.10	Review Status with Higher Level Management	All	5.6.1, 5.6.2, 5.6.3
			RD	7.2.2, 7.3.2
			MA	7.6
GG 3		Institutionalize a Defined Process		
	GP 3.1	Establish a Defined Process	All	5.4.2, 7.1
	GP 3.2	Collect Improvement Information	All	8.4
GG 4		Institutionalize a Quantitatively		
		Managed Process		
	GP 4.1	Establish Quantitative Objectives for the		
		Process		
	GP 4.2	Stabilize Subprocess Performance		
GG 5		Institutionalize an Optimizing		
		Process		
	GP 5.1	Ensure Continuous Process Improvement		
	GP 5.2	Correct Root Causes of Problems		

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