

Supplier Quality System Survey

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Please have your Quality Department or designated representative compete this form and return with the requested documentation within five (5) working days for our review and approval.

(Comp	pany Name			Date	
5	Stree	t Address				
(City		State	Zip Code	Phone	
(Comp	oleted By:				
N	lame	:	Title: _			
E	mail	l :	Phone:	·		
Γ	ate:					
1.	A. B. C. D. E. Em 1. 2. 3. 4. 5. 6.	nployee Breakdown: Total Personnel Quality Assurance: Manufacturing: Engineering: Purchasing: Sales:	Email Email Email Email Emai		Phone#:	
2.	Ye	USINESS IDENTIFICATIO car Business established: AGE Code:	N DUNS Number:	NAICS (Code	
3.		-9 FORM Complete and attach				
4.	BU	JSINESS CLASSIFICATIO	N			
		Large Small Non-Propertification Number Woman Owned Disadve Service Disable Veteran Of Alaskan Native Corporation 8(a) Certified Historica	Certification Da antaged (including Minority wned Ability One Non- ns and Indian Tribes HU	y Owned) 🗌 Veter profit Agency JB Zone		

U]	PPLIER TYPE			
7	Manufacturar/OEM Di	stributor 🗆 Br	okor/Indon	endent Service Other/Please Specify
_	Manufacturer/OEM Di		okei/maep	endent
ΕI	RVICES SUPPLIED			
	Service	YES (X)	NA	Specification/Standard Used
	Brazing			Specification Standard Cott
	Chemical Processing			
	Coating			
	Heat Treating			
	Material Testing			
	Other; Specify			
	Painting			
	Radiation Testing			
3	Radiation resting			
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PPF	Welding RODUCTS SUPPLIED Tope of product currently besome the second s	Certified by an tion Date:	d, are you	d registrar? Yes No

• IF YOUR COMPANY IS NOT REGISTERED TO AS9100 OR ISO9001, COMPLETE ALL PAGES 1-6, AND RETURN WITH

Page **2** of **6**

A COMPLETED W-9 FORM

No.	Supplier Quality Management System Requirements	Yes	No	N/A
1	Does the organization have an established and documented quality management system that			
	includes a Quality Manual and Quality Policy?			
2	Does the organizations' management review, monitor and measure the quality management			
	system yearly as a minimum?			
3	Has documented procedures been established for the quality management system or			
	referenced to them?			
4	Are documents & records controlled?			
5	Is there a procedure that defines the controls needed to approve documents for adequacy			
	prior to issue and to review & update as necessary?			
6	Are records established and maintained to provide evidence of conformity and			
	effectiveness of the quality management system?			
7	How long are records maintained and available for review by customers? Please specify in		Ш	
	years (minimum of 10 years).			
8	Has management appointed a member of management having independent responsibility &	Ш	Ш	
	authority for ensuring the quality management system processes are established,			
	implemented and maintained?			
9	Are appropriate training records of education, skills and experience maintained?	Щ	Щ.	
10	During product realization, does the organization establish processes, documents and	Ш	Ш	ΙШ
4.4	provide resources specific to the product?			
11	Are records available to validate evidence that processes and product meet requirements?	Н	Щ.	
12	Does the organization determine the requirements specified by the customer including	Ш	Ш	Ш
10	those not stated by the customer but necessary for specified or intended use where known?			
13	Does the organization review all requirements related to the product prior to the	Ш	Ш	$ \; \sqcup \; $
1.4	commitment to supply a product to the customer?			
14 15	Are all requirements defined and differences resolved prior to processing?		- -	
15	Where product requirements are changed, does the organization ensure the relevant	Ш	Ш	
	documents are amended, that the change is properly documented and authorized, and that personnel are aware of the change requirements?			
16	Are inputs relating to product requirements determined and records maintained?			
10	Design & Development			
17	Is verification performed to ensure the design and development outputs have met the design			
1 /	and development input requirements?	Ш		
18	Where tests are required for verification and validation, are these controlled, reviewed and	П		П
10	documented to ensure the acceptance criteria are met?	Ш		
19	Are design and development changes identified and records maintained?		П	
20	Does the design and development change procedure provide for customer approval when	Ħ	Ħ	
	required by contract?]	
	Purchasing			
21	Does the organization ensure purchased products conform to the specific purchase			
	requirements?			
22	Are suppliers evaluated and selected based on their ability to supply products in accordance			
	with documented requirements?			
23	Are supplier criteria for selection, evaluation and re-evaluation established and			
	documented?			
24	Are records of the evaluations and any actions as a result of the evaluations maintained?			
25	Is a register of approved suppliers maintained, including the scope of approval?			
26	Are suppliers reviewed periodically and the reviews used to establish levels of controls to			
	be implemented?		1	
27	Does the organization define the necessary actions to take when dealing with suppliers that			
	do not meet requirements?			
28	Do both the organization and all suppliers use customer-approved special process sources			
	where required?			<u> </u>

29	Does the function having the responsibility for approving supplier quality systems have the					
	authority to disapprove the use of sources?					
Supplier Quality Management System Requirements						
30	Does the purchasing authority describe the product to be purchased including the	Ш	Ш	Ш		
	requirements for approval, procedures, process, personnel qualifications requirements &					
2.1	quality management system requirements?					
31	Does the purchasing authority flow down all customer and quality system requirements	Ш	Ш	Ш		
	including right of access by the organization, their customers and authorities to all facilities					
22	involved in the order?					
32	Are the requirements relative to notification of nonconforming product and arrangements for approval of supplier nonconforming material flowed down to the supplier?	ш	ш	ш		
33	Are customer designated special requirements including key characteristics flowed down to					
33	the supplier?	ш	ш			
34	Does the supplier establish and implement inspection necessary to ensure the purchased			П		
51	product meets all specific purchase requirements including objective evidence of the					
	quality of the product and does this include certificates of conformance, test reports,					
	supplier audits, inspection upon receipt and delegation of verification to the supplier of					
	supplier certification?					
35	Is the purchased product held until it has been verified as conforming to requirements					
	unless released under a positive recall procedure?					
36	Where test reports are used to verify purchased product, is the data acceptable per					
	applicable specification?					
	Planning					
37	Does planning consider the establishment of process controls, in-process verification points	Ш	Ш			
	and use of tooling so that variable measurements can be taken?		_			
38	Is the production planned and carried out under controlled conditions?	Щ	<u> </u>	Щ		
39	Does the controlled conditions include characteristics of the product, work instructions,	Ш	Ш	Ш		
40	used of suitable equipment and availability & use of measuring & measuring devices?					
40	Do the controlled conditions also consider release, delivery and post-delivery activities?	H	井	<u> </u>		
41	Do the controlled conditions include accountability for parts quantities, split orders and	ш	ш	ш		
42	nonconforming material and product? Do the control conditions show evidence that all manufacturing & inspection operations		$\overline{}$			
42	have been completed as planned?	ш	ш	Ш		
43	Do provisions exist for the prevention, detection and removal of foreign objects (FOD)?			П		
44	Do provisions exist for workmanship criteria such as written standards, samples or	H	岩	H		
77	illustrations?	ш	ш	Ш		
45						
			П	П		
	Are production operations carried out with approved data such as drawings, process flow					
	Are production operations carried out with approved data such as drawings, process flow charts, mfg. plans, travelers, routers, process cards and inspection documents?					
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54	Does the quality management system provide the level of traceability required by contract			
	with batch/lot traceability maintained, assembly and components traceability maintained			
	and a sequential record of its production maintained and retrievable?			
55	Has customer property been verified, identified, protected and safeguarded for use? (This			
	can include intellectual property including customer furnished data used for design,			
	production or inspection.)			
	Supplier Quality Management System Requirements			
56	Have methods been defined to identify and record customer products lost, damaged or			
	otherwise made unusable and to report such occurrences to the customer?			
57	Does the organization account for preservation of the product and all constituent parts of			
	the product?	-		
58	Is there special handling for sensitive products such as electro-static discharge prevention			
	for electronic components?			
59	Does preservation include shelf life control and special handling of hazardous materials?			
	Monitoring & Measurement			
60	Does the organization maintain a register of monitoring and measuring devices and define			
	process for calibration including equipment type, unique identifier, location, frequency of	-		
	checks, method and acceptance criteria?			
61	Is personally owned equipment subject to the same procedures as company owned			
	equipment?	-		
62	To ensure valid results, is the measuring equipment calibrated or verified at specified			
	intervals, or prior to use, against traceable recognized measurement standards?			
63	Is the measurement equipment identified as to calibration status and safeguarded from			
	adjustments that would invalidate measurement results?			
64	Are there provisions in place to assess and record the validity of previous measuring results		П	П
	when the equipment has been found not to conform to calibration requirements?			
65	Is there a defined method or course of action prescribed to take on equipment and any		П	
	product affected?	_		
66	Are records of the results of calibration and verification maintained?		П	П
67	Are statistical process controls in place and are they being used?		Ħ	
68	Does the organization determine tracking methods and measure customer perception to		Ħ	
00	whether customer requirements have been met?			
	Internal Audits			I
69	Are internal audits conducted at planned intervals to measure and determine whether the		П	
	quality management system conforms to the required standards and how well it is			
	effectively implemented and maintained?			
70	Does the audit program address auditor selection and training to ensure objectivity and		П	
	impartiality of the audit process?			
71	Does the responsible management for those areas being audited ensure actions are taken		П	
	without delay to eliminate identified nonconformities and their causes?			
72	Do follow-up audits include verification of the actions taken and reporting of these		П	П
	verification results?	-		
73	Are suitable methods for monitoring and measuring the quality management system in		П	П
	place?	_		
74	When planned results are not achieved, is correction and corrective action taken where		П	
, .	required to ensure product conformity?			
75	Where process nonconformity has been identified, does the organization take appropriate	П	П	П
, 5	action to correct the process, evaluate whether the process nonconformity has resulted in			
	product non conformity and identify & control the nonconforming product?			
	Product Measurement			
76	Is the product monitored and measured to verify product requirements?		П	
77	Is this performed at appropriate stages during the product realization process in accordance	H	H	H
	with planned arrangements?			
78	Where key characteristics have been identified, are they monitored and controlled?			$\vdash \sqcap$

79	Where sampling inspection is used as a means of product acceptance, is it statistically valid, appropriate for use, and preclude acceptance of lots with known nonconformities? What			
	plan do you use?			
80	Is product held until inspected and verified as conforming to requirements, except when released under positive recall procedures pending completion of required measurement and			
	monitoring activities?			
81	Is the evidence of conformity with the acceptance criteria maintained and do the records indicate the person(s) authorizing release of product?			
	Supplier Quality Management System Requirements			
82				
	Are measurement requirements for product acceptance documented?	$\vdash \vdash \vdash$	$\vdash \vdash$	$\vdash \vdash \vdash$
83	Do test records show actual test result data when required by the acceptance test plan?	片片	H	H
84	Does the system provide for a process to inspect, verify and document a representative item from the first production run of a new part or following any change that invalidates the first article result?			
85	Is product not conforming to requirements identified and controlled to preclude its unintended use of delivery?			
86	Are controls, responsibilities and authorities defined in a documented procedure for dealing with nonconforming product?			
87	Does the system require taking action to eliminated detected non-conformances?			
88	Is the product dispositioned for scrap conspicuously marked and positively controlled until physically rendered unusable?			
89	Are records maintained as to the nature of nonconformities, subsequent actions taken, including concessions obtained?			
90	When corrected, are nonconformities subject to re-verification to demonstrate conformity to the requirements?			
91	Is appropriate action taken to address nonconformities detected after delivery or use?			
92	Is action taken to eliminate the cause of nonconformities to prevent recurrences?			
93	Is there documented procedure defining the requirements to review, determine the causes, evaluate, determine & implementing action needed, recording results of actions taken and reviewing corrective action taken in regards to nonconformities?			
94	Is there a documented procedure to define the requirements for flow down of corrective action to a supplier where it has been determined the supplier is responsible for the root cause and specific actions where timely and effective corrective actions are not achieved?			
UN	NITED DYNAMICS, INC. APPROVAL	1	ı	
Qu	ality Manager: Approval:			
Pro	ocurement Manager: Approval:			
	ΓΕD DYNAMICS, INC. Assessment completed and in file			