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Littelfuse Supplier Production Part Approval Process (PPAP) Manual

November 2013, Revision A

Jun.2014, Revision B

April 2016, Revision C

March 2021, Revision D



Littelfuse PPAP General Requirements

Littelfuse uses the Production Part Approval Process to confirm that the supplier understands the design specifications and has a process capable of producing product to meet these requirements, during an actual production run, at the quoted production rate. An industry requirement for all automotive suppliers, PPAP is being expanded to include all of our suppliers.

PPAP requirements vary based on the submission level assigned to a supplier and/or part number. The Littelfuse Supplier Development Engineer and/or Supplier Quality Engineer representative, is responsible for designating the submission level. The submission level is generally determined during the PAF RFQ process and/or PAF. Suppliers are expected to apply these same PPAP requirements to all sub-suppliers.

If the parts and PPAP conform to design record requirements, and the capability data is within specifications, the parts and PPAP will be accepted.

The Part Submission Warrant (PSW) will be signed by Littelfuse Supplier Development engineer and/or Supplier Quality Engineer representative, giving the supplier, the approval to run the parts as submitted. If there is no signed PSW, there is no approval from Littelfuse and parts cannot be shipped, unless otherwise is specified by Littelfuse Supplier Development Engineer and/or Supplier Quality Engineer representative (PPAP with interim approval).

Submission Levels

Level 1	Warrant only (and for designated appearance items, appearance approval report) submitted to Littelfuse.
Level 2	Warrant with product samples and limited supporting data submitted to Littelfuse
Level 3	Default submission level, for all submissions unless otherwise specified. (Refer to PPAP matrix)
Level 4	This level is reserved for special applications only; it cannot be utilized without the consent of Littelfuse Supplier Development Engineer and/or Supplier Quality Engineer, representative.
Level 5	Only applied to on site review as requested by Littelfuse Supplier Development Engineer and/or Supplier Quality Engineer, representative.



Expertise Applied | Answers Delivered Change Management Matrix

2) Does it		tomer's sig			cs? ner affected?	if customer specific requierements exist, the agr obligatory!	reement is	u.	PPAP I FVFI		
		3) is the t	4) type of	- VI	ne custon	iei allecteur	obligatory:		PAF	Ь		
			., ., , , ,		affect cor	ntract docume	nts (e.g., specifications, customer's drawing, data-sets	?		₽dd		
				.,			inction, performance, reliability affected?					
	Υ	Y/N	ALL	Y/N	Y/N	Y/N Change to significant characteritics agreed with the customer for the product, sub-						
				Y	assy., component (electrical/mechanical process,)?							
				1	Y e.g., change to design, tooling, Z Y e.g., change to a dimension not included in the Littelfuse							
					Υ	specification		Z	N	-		
						Change to m		Z	Υ	3		
			*				ternal specification or tolerances outside Littelfuse	Z	.,	١,		
			Σ			specification		2	Υ	2		
			ig.	N		Change to in	ternal specification or tolerances but still within	_	N	2		
			Design MC**		N	Littelfuse spe			<u> </u>			
					IN		lentification of parts/materials but with unchanged	2	Υ	2		
						composition				L		
							arly man'Fing stages (e.g., pre-drilled dimension for a	2	Υ			
						shaft, water	location,)		· ·	L.		
							in process chain, (inc. Supplier, duplicated	Z	N			
				Y	Y/N	production I				L		
					V		s in checks, checking sequence or other reasons.	Z	N	L		
					Υ	e.g., change	in hardening parameters, injection temperature, Change in no. of cavities in tool, progression tools,	Z	N			
							incremental tool.	1	N			
							Duplication of production and checking equipment			t		
							within an existing line	1	N			
						Prod'n-	New of type of machine obtained and installed	I I	N			
						assembly	Change to an existing tool, new equipment, new	_	N			
							Poka Yoke		-	L		
			U				Change in manufacturing stages	-	N			
			Process Mc				Change in setting parameters, production facilities,	-	N			
	N	N	ces				injection temperature Changes in checks, worsened RPN		N			
			Pro	N	N.		Change to checking method, RPN		IN			
					N		Unchanged/improved, same process	I I	N			
						Testing	Extended checks with no change to method (e.g.,			T		
							larget sample size)	-	N			
							Reduction/elimination of check not relevant to the					
							customer (e.g., random sample check)	-	N			
							Tools moved from one line to another, lines are the		N	١,		
							same.	-	IN	Ľ		
						Transfer of		_	N			
						production	no change to the process chain.		Ľ.			
							Location change: equipment, parallel prod'n (not	Z	N			
						Supplier cha	learly mfg stages) nge, new 2nd supplier, supplier has changed sub-			\vdash		
				Y	Y/N	supplier	Ziid supplier, supplier has changed sub-	Z	Υ			
			tics		Y/N	New Carrier	or ESP, SLC	1	N			
			Logistics	7000	Y/N		l/or Littelfuse packing, shipping, invoicing change	Z	Υ			
			2	N			king (e.g., plant to plant, within the plant,) and					
					Y/N	suppliers		-	Υ	4		
				Y		Documents	adjusted to status of approved /released product	Z	2	1		
			ဥ			Documents :	adjusted to status of approved/released product or	_	Υ	1		
			Doc Mc	NI.	Y/N	to correct fo	rmal defects.					
			DO	N		Change to de	ocuments not product-related (e.g., work		NI			
						instructions,)		N			
R	Re-Use of	tools follo	wing 12 or	months or	ut of use			Z	N			
_		qualification						Υ	N	L		
							ar (e.g., turning too, honing tools)	-	N	L		
I R	Replacem	ent of an i	dentical ma	achine or a	machine	with an equal	functionality. Replacement of identical measuring		N			



- y Yes
- n No

np not permitted

- Customer Involvement not essential (Note: PPAP documents must be archived in-house)
- I Customer must me informed as ISO/IATF. Para 4.2.3.1 the customer must have 2 weeks to issue findings.
- Z Customer agreement required, execution of PPA procedure
- Mc Modification
- **ESP** External service provider
- SLC Supplier logistics centre (also applies to warehouse)
- RPN Risk priority number from Process FMEA
- * or other authorized production documents provided to the customer (e.g., quotation drawing, control plan

Remark:

For distributors, it is their responsibility to receive and approve PPAP from the original manufacturer and then submit to Littelfuse along with their own PSW making sure that there is a cross reference among part numbers: manufacturer, distributor and Littelfuse. Any deviations from this requirement need to be approved by Littelfuse in writing prior to first shipment of parts/material

Littelfuse expects that suppliers and distributors, manage and approve their own suppliers base and maintain evidence of compliance.

Regardless of the submission level, PPAP shall be maintained at least for the length of time the part is active plus one calendar year.

PPAP is also applicable for standard catalog purchased parts, components/off the shelf, i.e., electronic, mechanical and/or other component categories.

Supplier must submit PPAP in English, unless otherwise is specified by Littelfuse Supplier Development Engineer and/or Supplier Quality Engineer, representative



Submission Status

The PSW is reviewed by the Supplier Development Engineer and/or Supplier Quality Engineer representative, as follows:

- Approval: Indicates the part meets all specifications and requirements, and the supplier is authorized to ship production quantities.
- Interim Approval: Permits shipment of material for production requirements on a limited time
 or piece quantity basis, when supplier has clearly defined the root cause of the nonconformities preventing production approval and has prepared an interim approval action plan
 agreed upon by Littelfuse.

Note 1: For those parts with disposition "Interim Approval", supplier should issue another PSW once the non- conformities have been corrected.

Note 2: PPAP re-submission is required to obtain a status of approved.

Note 3: Parts with a status of interim approval are not considered fully "Approved"

Rejected: Prevents production quantities from being shipped because the submission, the
production lot from which it was taken and the accompanying documentation does not meet
Littelfuse requirements. In such cases, the submission and/or process as appropriate, shall
be corrected to meet Littelfuse requirements. The re-submission shall be approved before
production quantities may be shipped.

As required, the Supplier Development Engineer and/or Supplier Quality Engineer representative will determine if annual PPAP re-qualification is applicable or not (based on Customer Specific Requirements or other requisites). This requirement must be reviewed and agreed upon during APQP, PAF (Print Acceptance Form) and/or early PPAP stages. When applicable, the annual PPAP re-qualification is separate and in addition to PPAP submissions related to engineering changes.

Annual re-qualification documents may include:

- Full dimensional lavout
- Material testing or certifications (such as flammability compliance)
- Environmental requirements
- · Reliability testing

Note: No additional Supplier costs or fees and charges associated with this requirement are allowed.



P	RODUCTION PART APPROVAL PROCESS		PI	PAP Le	vel	
	(PPAP)	1	2	3	4	5
1	Design Record	R	s	s	*	₩R
2	Engineering Change Documents	R	s	s	*	R
3	Customer Engineering Approval	R	R	s	*	R
4	Design FMEA	R	R	s	*	R
5	Process Flow Diagram	R	R	s	*	R
6	Process FMEA	R	R	s	*	R
7	Control Plan	R	R	s	(*)	R
8	Measurement System Analysis Studies	R	R	s	380	R
9	Dimensional Results	R	s	s	*	R
10	Material, Performance Test Results	R	s	s	*	R
11	Initial Process Studies	R	R	s	*	R
12	Qualified Laboratory Documentation	R	s	s	*	R
13	Appearance Approval Report (AAR)	s	S	s	*	R
14	Sample Product	R	s	s	*	R
15	Master Sample	R	R	R	:*:	R
16	Checking Aids	R	R	R		R
17	Records of Compliance (Substance of Concern & Conflict Minerals)	R	R	R	*	R
18	Part Submission Warrant	S	s	s	s	R

Figure 1 - PPAP Submission Levels from PPAP latest edition by AIAG.

- **S** = The organization shall submit to the customer and retain a copy of records or documentation items at appropriate locations.
- **R =** The organization shall retain at appropriate locations and make available to the
 - customer upon request.
- * = The organization shall retain at appropriate locations and submit to the customer upon request.



Use of the AIAG Bulk Materials Checklist is an acceptable substitute for a regular PPAP if it is applicable.

The requirements of this manual were drafted to be compliant with AIAG PPAP Standard. Littelfuse has specific requirements and additions to this standard that need to successfully submit a PPAP to Littelfuse. The Littelfuse specific requirements are a must for all Level 3 PPAP submission, unless otherwise specified by Littelfuse Supplier Development Engineer and/or Supplier Quality Engineer representative.

Instructions to complete Littelfuse PPAP Submission

Element 1 - Design Record and Ballooned Drawing

Design Record and/or Specification are engineering requirements for judging the acceptability of a part characteristic. For qualification, every feature of the product as identified by engineering in design record/print or specifications must be measured.

Design records are considered as; all Littelfuse and supplier design records.

A ballooned drawing shows the parts or assemblies in a part print with numbered ballooned that point of individual dimensions and requirements of the part. The numbers on the ballooned drawing correlated with the numbers found on the Dimension Result sheet. A ballooned drawing must be submitted as part of PPAP for every submission level when there are dimensional results.

All part requirements on the print must be ballooned and numbered for reference and measurement. These may include:

- 1. Dimensions and tolerances of parts
- 2. Visual features (color, texture, etc.)
- 3. Chemical characteristics (cure time, etc.)
- 4. Physical and mechanical properties (torque, hardness, plating thickness, etc.)
- 5. Any other specified requirement that you have the capability to measure or that is described in print notes or referenced specifications.

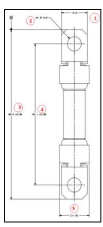


Figure 2 - Example of a ballooned drawing



Element 2 & 3 - Engineering Change

The supplier shall have any authorized engineering change documents not yet recorded in the design record but incorporated in the product, part or tooling.

Suppliers and sub-suppliers of Littelfuse are not authorized to make any change a change to a product and /or process that was previously approved without first receiving written authorization by Littelfuse. Examples of such changes include, but are not limited to: tool moves, changes to manufacturing process / shipping location / plant move, material changes (subsuppliers changes), changes that impact the logistics and deliveries such as ERP and carriers, and others. Any change must be communicated to Littelfuse. In the **Product / Process Change Notification (PCN).**

If apply, PAF (Print/Spec Acceptance Form) will be generated to assure supplier understands requirements regarding changes through the ECR/GCF and PCN/SPCN (e.g. Introduction of new supplier) process.

Is supplier responsibility:

- Confirm their capability to produce the material/component to design record.
- Notify any concerns regarding design records in SECTION 3 before signing off this form.
- Provides any requested quality documents (e.g. PPAP, IMDS, QMP and others as applicable) as established in this form.

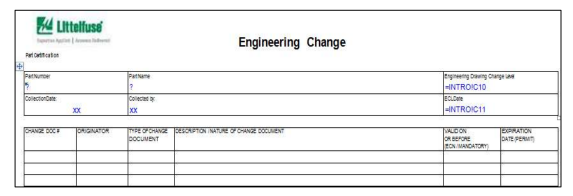


Figure 3 - Form of the Engineering Change



Littelfuse Expertise Applied Answers Delivered	Supplier Prod	duct/Process Cha	ange Notice	Rev B
PCN Information				IVEA D
Supplier Name		PCN#		
Supplier Location(s)		Request Date		
Contact Name		Implement Date		
Phone #		Product		
Email		Identification		
Category of Change				
1. Product Design		2. Assembly / Fa	abrication Process	
3. New Tooling / Mold		4. Manufacturin		
5. Sub Supplier or material s 7. New equipment	source	6. Material type	or component critical equipment pa	aramatara
9. Others		o. Changes on C	critical equipment pa	arameters
Description of Change	& Reasons			
Important Dates ✓ Qualification Samples avalia Final Qualification data avali				
Timal Qualification data avail		1 Last Time orde	i date	25
Method of Distinguishin	ng Changed Pro	oduct		
Product Mark				
Date Code				
Others				
Demonstrated or Anticipoler Qualification F		n Form, Fit, Functior	n or Reliability	
Littelfuse Requirements	ALL Information	on Below is to be filled out by Littelf	fuse	
PPAP Required Y	es ⊛ No ⊜	Level	Due date	
	es □ No□			
	es No			
	es 🗌 No			
Additional Requirements				
Littelfuse Approval Sign	nature	Signature		Date
LF Procurement Represer		eignature		2410
LF Engineering Represent				
LF Supplier Development				-
Additional Requirements/Comm				

Figure 4- Form of the Engineering Change

(excel form available in: https://www.littelfuse.com/about-us/supplier-resources.aspx)



Litti Expertise Applied	telfuse			<u>P</u>	rint/spec	c <u>A</u> cc	ceptance	<u>F</u> orm			
Expertise Applied	Allsweis Deliv		CTION 0	- BASIC	INFORMATION	ı					
Supplier Name							ECO/GCF number				
Vendor Code				1	Select type of chang	je	DGW Type	Select			
Project Description:	Project Description:										
Part Number	Drawing No.	Rev.	Part De	scription	Previous Part Number (If Changed)	Previous Rev.	ECO/GCF Details	Application			
							Select	Select			
							Select	Select			
<u>.</u>							Select Select	Select			
							Select	Select			
							Select	Select			
		SECTIO	ON 1 – L	ITTELFU	ISE REQUIREM	ENT					
1. PPAP with IMDS	Select	Level	3		Due Date		PO#				
Required	(Detailed PPAP re	quirements refe	er to Littelfu	use Supplie	r Production Part App		ess Manual)				
2. Process Audit	Select										
3. LF QMP Updated	Select	(Detailed requ	irements re	efer to Littel	fuse Quality Manager	ment Plan)					
4. DFM Required	Select	Due Date									
5.Technical Review	Select	Due Date									
Meeting Required	(As required based	on information	in append	ix '7.2' for a	guideline of what "co	mplex parts	" refers to)				
6. Section 3: Supplier Review Checklist Required	Select		lates is not	required si	uch as correcting draw		dding tolerance misse				
7. Sample Required 8. Appearance	Select	Due Date	0		E	Se	Specify typ	e of samples			
Approval Required	Select	Such as Sa	afe-Launc	h activities							
9. Additional Requirements											
Purch (Printed Na				Engineerin			Supplier Developmen & Printed Name)				
	,										
	SI	ECTION 2 -	SUPPL	IER RES	PONSE (Updated	by Supplie	r)				
After reviewing SECTIO Accept Print/Spec a Accept Print/Spec v Cannot accept Print	as-is with notes (Specify in	SECTION 3 re		ded informat	tion to be added to D	rawing / Spe	ecification)				
If accept, we (Supplier) of All part Prints/Specification about the disposition and this PAF. All costs were In conclusion, we have the by Littelfuse	declare as below: ions were reviewed i d handling of invento evaluated and are a ne ability to develop,	n detail and are ries of the previ	ious materi	ial/part vers	ion (if change is invol	ved). We wi	Il follow quality requirer to the quality and cost s	nents as stated in standards as defined			
Sales/Proje (Printed Na				neering Ma ed Name 8			Quality Manag (Printed Name &				
,											
will meet all the requirem	nents as documented t to reject componen ponsible for all costs	in the form inc ts from a suppli associated wit	luding thos ier if these	se in the Pri are the res	nt/Specification. ult of a part or proces		ved, Littelfuse understa	303			

Figure 5 - Print/Spec Acceptance Form - Page 1.



Supplier Review Dept.	Review Contents	Yes / No / N/A	Comments & Clarify items with Littelfuse	Due Date	Respon sible
	Are there other similar components in current production?	Select			
	Is all drawing content well understood including notes, specifications, standards and requirements (i.e. EIA J-STD-002D)?	Select			
	Can safety characteristics in drawing be met?	Select			
	Can all dimension in drawing be met with defined Cpk, SPC and critical characteristics requirements (Unless there is an agreement between Littlefluse and Supplier minimum Cpk is 1.67 for critical characteristics).	Select			
	Can current process and production equipment meet design specifications?	Select			
	Is the raw material available and meets the drawing?	Select			
ring	Is apprearance requirement clear and understood in order to create own visual aids?	Select			
inee	Is material free of contaminant agents / sources?	Select			
Technical Engineering Product Project	Is the packaging defined and agreed and validated with Littelfuse (Ground, airfieight, overseas)? If not, there must be an agreement between Littelfuse and supplier.	Select			
Ē.	Are fixtures required to produce this material (i.e welding, machining fixtures and /or gauges) available?	Select		3	
	Are the parting lines, gates, pin position, flash size, sink or protrusions level, deformation level defined with Littlefluse in a measurable method for the components?	Select			
	Is there any agreement to use regrinding or recycle material?	Select		50	
	Is there any warranty, reliability or component life agreement?	Select		59	
	Does the material require special storage/handling method and the shelf life is specified?	Select	ez	38	
	Are there additional characteristics or features requirements for the component?	Select		29	
	Is zero defects target defined within your organization?	Select		54	
	Provide the estimated Internal product yield rate (DPPM or %) per product category.			3	
_	Are you able to measure and test the component based on drawing requirements?	Select		9	
Quality EHS	Are fixtures required to measure / test this material available?	Select		5	
J	Is there a system in place to keep traceability at component level?	Select		9	
	Is material in compliance with RoHS/REACH/Halogen free requirements?	Select		50	
	Are IMDS requirements clear?	Select			
Sourcing urchasing Buyer	Can raw material be supplied according to Littelfuse requirements?	Select			
Sourcing Purchasing Buyer	If outsource process is required (i.e. plating, annealing), Is the sub-supplier manufacturing capability and quality assured?	Select			
동물	Is there any additional equipment to produce this material (Backup equipment)?	Select			
Production Equipment	Is there any additional equipment needed to produce this material (Capacity)?	Select			
Pro	Are there enough operators with required skills to run the process (Certified personnel)?	Select			
, <u>p</u>	Is there a MOQ/MPQ requirement?	Select			
Sales Marketing	Can Littelfuse lead times be met?	Select			
Mar	Was the unit cost quoted considering above items?	Select			I

Figure 6 - Print/Spec Acceptance Form - Page 2.



Element 4-1 - Design FMEA (if supplier is design responsible)

A Design FMEA is an analytical technique utilized by the design responsible to assure that to extent possible, potential Failure Modes and their associated Causes or mechanism of failure have been considered and addressed prior to releasing the part to production.

DFMEA analyzes the functions of a system, subsystem, or component of interest as defined by the boundary shown on the Block/Boundary Diagram.

Severity, Occurrence and Detection ratings are used when performing FMEA activities. These rating scales must be compliant with the AIAG & VDA guidelines for FMEA (latest edition) and definitions are included in the DFMEA worksheet as well as this handbook.

Action Priority (AP) is based on combinations of Severity, Occurrence and Detection ratings to prioritize actions for risk reductions.

- Priority High (H): Highest priority for review and action.
 The team need to either identify and appropriate action to improve Prevention and/or Detection Controls or justify and document why current controls are adequate.
- Priority Medium (M): Medium priority for review and action.
 The team should identify appropriate actions to improve prevention and/or detection controls, or, at the discretion of the company, justify and document why control are adequate.
- Priority Low (L): Low priority for review and action.
 The team could identify actions to improve prevention or detection controls.

Littelfuse requires that any severity ranking of 9 or 10 be addressed with a corrective an action plan. In Addition, Littelfuse recommends suppliers to develop continuous improvement activities for the top Action Priority identified

DFMEA is only required when the part is designed by the supplier. Otherwise, the Design FMEA is the responsibility of Littelfuse.



Figure 7 - Design AIAG &VDA FMEA example

(excel form available in: https://www.littelfuse.com/about-us/supplier-resources.aspx)



Element 5 - Process Flow Diagram

The Process Flow Diagram depicts the flow of materials through the process.

The Process Flow Diagram must follow the process from receipt of raw material and receiving inspection, through any warehousing and shipping steps, and include any "Dock Audits" and Final Inspections. The PFD shall comprehend all potential paths that a part can take in the process, including inspection, containment, rework, scrap, material shipped to sub-contractors and the returning of the material back to the supplier's plant. The Primary process steps must match both the Control plan and the PFMEA.

The Process Flow can be provided in any format used within an organization.

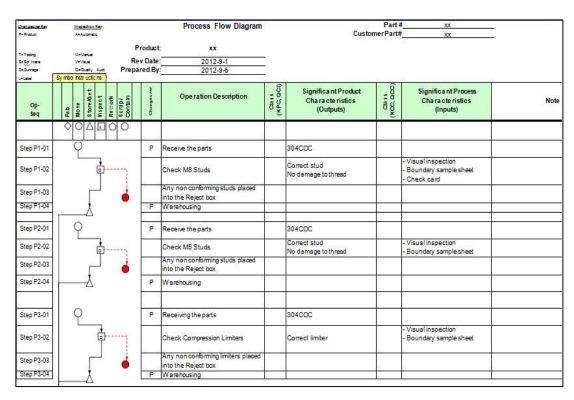


Figure 8 - Example of a Process Flow Diagram (excel form available in: https://www.littelfuse.com/about-us/supplier-resources.aspx)



Element 6 - Process FMEA

Format AIAG for reference while transition to AIAG VDA format is complete

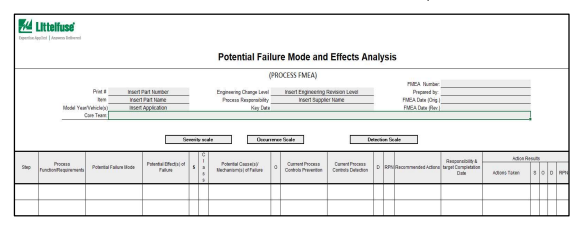


Figure 10 - Example of the Process FMEA form

(excel form available in: https://www.littelfuse.com/about-us/supplier-resources.aspx)



Element 6-1 - Process FMEA AIAG | VDA

Process FMEA analyzes the potential failures of manufacturing, assembly, and logistical processes to produce products which conform to design intent. Process-related failures are different than the failures analyzed in the design FMEA.

The process FMEA analyzes processes by considering the potential failure mode which may result from process variation to stablish priority of actions for prevention and as needed, improve controls, the overall purpose is to analyze processes and take action prior to production start, to avoid unwanted defects related to manufacturing and assembly and the consequences of those defects.

Severity, Occurrence and Detection ratings are used when performing FMEA activities. These rating scales must be compliant with the AIAG & VDA guidelines for FMEA (latest edition) and definitions are included in the DFMEA worksheet as well as this handbook.

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- **Priority Low (L):** Low priority for review and action.

 The team **could** identify actions to improve prevention or detection controls.

Littelfuse requires that any severity ranking of 9 or 10 be addressed with a corrective an action plan. In Addition, Littelfuse recommends suppliers to develop continuous improvement activities for the top Action Priority identified.



											PROCE	ESS FAILURE	MODE AND E	FFECT ANALYSIS	(PROCESS FMEA)																					
LANNING & PRE																																				
	Company Name:				PCP Automotive Pi					MEA ID number:			M4-PRD73-00	01-HEV649			Revi			В																
	Plant Location:			Pi	edras Negras, Coa	ahuila				FMEA Start date:							Process Responsi	oility:																		
	Part Number:			Customer Name:	N/	/A	Model Year(s)/ Program(s):			PFMEA Revision date:							Confidentiality L	evel:																		
Cros	s funtional team																																			
	STRUCTURED A	NALYSIS (Step 2)				FUNCTION A	NALYSIS (Step 3)			ľ	FA	AILURE ANALYSI	5 (Step 4)			RISK ANALYSI	S (Step 5)						OPTIMIZATI	ON (Step 6)												
Process Item System, subsystem, part element or name of process	2.Process Step Station No, and Name of Focus Element	3. Process	Work Element Vi Type	Function of the Process Item Function of System, Subsystem, Part element or Process	2.a Function of the process Step	2.b Product Characteristic as applicable	3.a Function of element and p	of the process work rocess characteristic	3.b Process Characteristics as applicable.	1. Failure Effects (FE)	00	Failure mode (FM) of the process step		use (FC) the work lement	Current Prevention Control (PC) of FC	Current Detection Controls (DC) of FC	Current Detection Controls (DC) of FM	Detection(D) of FC/FM PFMEA AP	Special characteristics	evention action	Detection Action of FC o FM	Responsible Person's Name	Target Completion date	Status	Action taken with pointer to evidence	Completion date										
		Man					Man						Man																							
		Machine	Drill				Machine	Drill body		-8			Machine	incorrect or damage drill bit	NA .	Visual			Imple Equip 909-4 Body Leng	ement Body Drill oment for Body 806 (Poka-Yoke Position, Body	Implement Gauge Hole inspection and position	Gumercindo Galvan	8/27/2020	Open												
Op.10: Body		Material		Able to do the Final	Drill holes on the		Material			Dimensions on	dis	Holes saligment, Not holes, holes	Material				12000		Long																	
drilling	Body drilling	Method	Body drilling process	Fuse Assembly	body to able apply epoxy	according with the drawing 909- 623	Method	Aligment and position according WI: M4- PRD30-1013A- HEV649		the fuse are out 8	the fuse are out 8 diameters	specification spec & damage	cification diameters out spec & damage	diameters out spec & damage	diameters out spec & damage	diameters out spec & damage	diameters out pec & damage	diameters out spec & damage	diameters out spec & damage	diameters out spec & damage	diameters out spec & damage	diameters out spec & damage	Method	Not aligment and incorrect position	NA NA	Visual	Visual	Visual 8 H		ement Body Drill oment for Body 606 (Poka-Yoke Position, Body th)	Implement Gauge Hole inspection and position	Gumercindo Galvan	8/27/2020	Open		
		Measurement					Measurement						Measurement							5600		1														
		M.N (Environment)					M.N (Environment)			-:1			M.N (Environment)																							

Figure 10 - Example of the Process FMEA VDA form (excel form available in: https://www.littelfuse.com/about-us/supplier-resources.aspx



7 - Control Plan

Control plans are written descriptions of the operations, processes, materials, equipment, methodologies, and CTQs for controlling variations in key product or process characteristics integral to the manufacturing process. The supplier shall have a Control Plan that defines all controls used for process control and complies with the Littelfuse specifications. Control plans for "families" of similar parts are acceptable if the new parts have been reviewed for commonality.

The control plan should be developed in stages from proto-type through production. Early planning on the control plan will usually result in a more robust process. Suppliers should develop a pre-launch control plan early in the development of a new product and submit it to Littelfuse for feedback. A pre-Launch would normally show greater inspection size and frequency. Littelfuse may also request that you provide specific documents required at PPAP early in the development phase and the most common ones are the PFMEA and a pre-launch Control Plan.

The process flow diagram and design record/specification provide inputs to the Control Plan. All CTQs identified as Process, First-Piece, or Safety Related by the supplier must be listed on the control plan form. The Process Flow, Process FMEA and Control Plan manufacturing steps should match. The control plan cannot be excessively dependent on visual inspection and should target prevention techniques wherever possible.

NOTE: All critical characteristics / CPK / Safety Characteristic must be identified on the control plan.

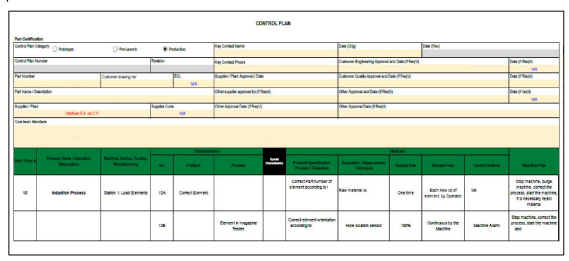


Figure 11 - Example of the Control Plan form (excel form available in: https://www.littelfuse.com/about-us/supplier-resources.aspx)



Figure 12- Example of the Control Plan VDA form - (excel form available in: https://www.littelfuse.com/about-us/supplier-resources.aspx)



Element 8 - MSA,GR&R

Measurement System Analysis (MSA) is an experimental and mathematical method of determining how much the variation within the measurement process contributes to overall process variability.

MSA is a requirement for qualification. Supplier must submit and follow the following:

- All measuring equipment and gauges are calibrated.
- A GR&R must be submitted for devices measuring data on CTQs and for each measurement device mentioned on the control plan on all Level 3 submissions.
- The minimum requirement for Gage R&R is:
 - A Gage R&R study using Total Tolerance samples.
 - % R&R below 10% is acceptable.
 - % R&R between 10% and 30% is marginal acceptable, need an action plan to address and improve the method of measurement.
 - Gages with R&R at 30% or more cannot be used.
 - Number of distinct data categories (ndc) >= 5.
- The ANOVA analysis method is recommended to be used to calculate %R&R.
- For visual devices and Go/ No-Go measuring equipment, the Attribute Gage Study shall be performed. At least must be used one out of the following methods:
 - Attribute gage bias report (Analytical method)
 - Gage repeatability and reproducibility report (Attribute hypothesis test method)
- Any equipment or gauge that is not meeting the %R&R should not be used and must have a plan to fix it or replace it.



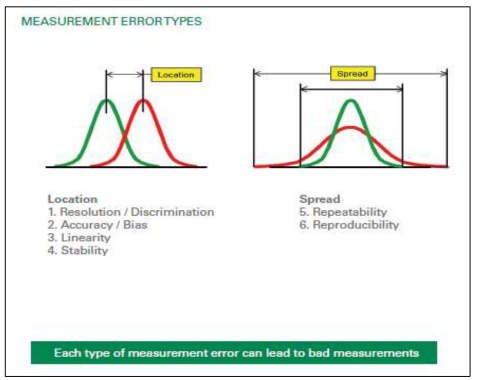


Figure 13 – Measurement Error Types

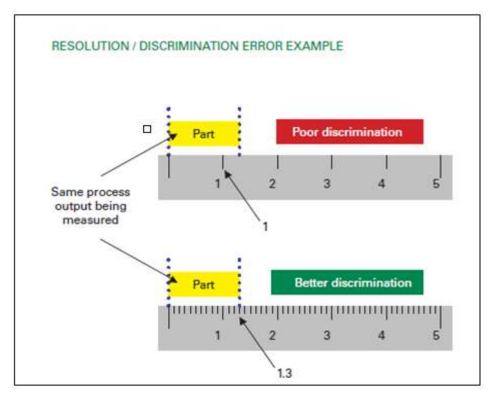


Figure 14 - Example of Resolution/Discrimination Error



					Α	NOV.	AME	THO)				
PartNumb	ë.				Gage Na	me			Appraise	r A			
XX					XX				Chunxiang Yang				
Part Name					Gage Number				Appraise				
XX Characteri	intin		Cnack	Ication	BA025 Gage Ty	-			Tingtin Appraise				
Thicknes			117		PCU-2				Fengiu				
Characteri	stic Cla	esificatio	n		Trials	10.11	Parts			ers.	Date Per	formed	
						3	1	0		3	2012.0	6.25	
APPRAI	SER/		_	_	_	PΔ	RT	_	_	_		Α	VERAGE
TRIAL#	-	1	2	3	4	5	6	7	8	9	10		
1. A	1	138.46	141.89	136,36	138,37	140.68	142.79	139,47	140,96	141.03	135.15		139.516
2.	2	138.45	141.83	136.35	138.32	140.63	142.73	139.35	140,94	141.02	135.21		139.483
3.	3	138,47	141.85	136.34	138.35	140.67	142.75	139.42	140.92	141.13	135.26		139.516
4.	AVE	138.46	141.86	136.35	138.35	140.66	142.76	139,41	140.94	141.06	135.21	Ха≡	139.505
5.	R	0.02	0.06	0.02	0.05	0.05	0.06	0.12	0.04	0.11	0.11	fa=	0.064
6. B	1	138.43	141.86	136.37	138,37	140,68	142.72	139.52	140.95	141.08	135,18		139,516
7.	2	138.47	141.82	136.42	138.45	140.69	142.76	139.42	140.98	141.12	135.51		139,564
8.	3	138.42	141.93	136.45	138.42	140.66	142.82	139,43	140.97	141.12	135.25		139.547
9.	AVE	138.44	141.87	136.41	138.41	140.68	142.77	139,46	140.97	141:11	135.31	Xb≡	139.542
10.	R	0.05	0.11	0.08	0.08	0.03	0.10	0.10	0.03	0.04	0.33	f _b =	0.095
11. C	1	138,43	141.93	136,85	138,34	140,63	142.73	139.46	140.92	141,13	135.12		139,554
12.	2	138,41	141.92	136.32	138,39	140.72	142.72	139.42	140,96	141.52	135.20		139,558
13.	3	138.48	141.88	136.29	138,34	140,69	142.74	139,48	140,93	141.06	135.18		139.507
14.	AVE	138.44	141.91	136,49	138.36	140.68	142.73	139.45	140.94	141.24	135.17	Xe≡	139,540
15.	R	0.07	0.05	0.56	0.05	0.09	0.02	0.06	0.04	0.46	0.08	fe=	0.148
16. PAR	T											X=	139.529
AVER	AGE	138,45	141.88	136.42	138.37	140.67	142.75	139.44	140.95	141.13	135.23	R _p =	7.522

Source	Resis	staResista	9	S	MS		1		Sig
Appraiser		2	0.	03	0.01		1.	72	Ť
Parts		9	472	2.47	52.5	0	694	2.92	
Appraiser-by-Pa	rt	18	0.	11	0.01		0.	79	
Equipment		60	0.	45	0.01				
Total		89	473	3.05					
					•	Signif	icant at	α=0.0	05 level
Anova Rep	ort	Stand Deviation		% To	tal Varia	tion	% C	ontribu	tion
Repeatability (E	V)	0.08	3		3.5%		0.1%		
Reproducibility (AV)	0.0	1		0.6%		0.0%		
GRR		0.09	9		3.6%			0.1%	
Part-to-Part(PV		2.4	1		99.9%			99.9%	
				Gage s	system O.	K			
Note:						90			
Tolerance = 5	8.50				T	otal va	ariation	TV)=	2.42

Figure 15 - Example of a Gage R&R with ANOVA method - form (excel form available in: https://www.littelfuse.com/about-us/supplier-resources.aspx)



THE SECOND P.					MSA Repo	rt					
		Littelfuse Suzho	u Pant		Equipment	59166-503	Auto tester	А	Stati	on 1	
	P/N		59166-503		Equipment ID:	11-	473	В	Station 2		
Character	Hash Test	Spec	OK:1	NG:0	Equipment	N	IA .	С			
Chara	cter Grade				Character: Times	Parts QTY 50	Station 1-3	Date	Station 3 9/30/2015		
					Record	50	1-0				
Number 1	A-FIRST TIME	A-Second Time	A-Inira IIme	B-FIRST TIME	B-Second Time	B-INITA IIME	C-FIRST TIME	C-Second Time	C-Inira Time	Spec	
2	U	U	U	U	U	U	U	U	U	U	
3	0	0	0	0	0	0	0	0	0	0	
5	Ü	Ü	Ü	Ü	Ü	Ü	Ü	Ü	Ü	0	
7	0	0	0	0	0	0	0	0	0	0	
8 9	1	1	1	1	1	1	1	1	1	1	
10	i	i	i	i	i	1	i	i	i	i	
11 12	1	1	1	1	1	1	1	1	1		
13	1	1	1	1	1	1	Ü	1	1		
14 15	1	1 0	1	1	1	1	1	1	 	- 1	
16	1	1	1	1	1	1	1	1	1	1	
17 18	1	1	1	1	1	1	1	1	 	- 1	
19	1	1	1	1	1	1	1	1	1		
20	1	1	1	1	1	1	1	1	1		
22	1	1	1	1	1	1	1	1	1	1	
24	1	1	1	1	1	1	1	1	1		
25 26	1	1	1	1	1	1	1	1	1	1	
26	+	1	1	1	1	1	1	1	1	+	
28 29	1	1	1	1	1	1	1	1	1		
30	- i	1	1	1	1	1	1	1	1	- i	
31	1	1	1	1	1	1	1	1	1	1	
32 33	- i	1	1	1	1	1	- i	1	1	- i	
34 35	1	1	1	1	1	1	1	1	1	1	
36	- i	1	1	1	1	1	1	1	1	- i	
37 38	1	1	0	1	1	1	1	1	1 0	1	
39	- i	- i	1	1	1	+	- i	1	i		
40 41	1	1	1	1	1	1	1	1	1	1	
42	i	Ü	Ü	i	1	1	i	i	1	1	
43 44	1	1	1	1	1	1	1	1	1		
45	1	1	1	1	Ü	1	1	1	1	1	
46 47	1	1	1	1	1	1	1	1	1	- 1	
48	1	1	1	1	1	1	1	1	1	1	
49 50	1	1	1	1	1	1	1	1	1		
			Data summ	ary							
Attendee	Right ajudgement Times for NG parts	Right ajudgementT imes for OK parts	Right ajudgement Times (Total)	Wrong ajudgement Times	Missing ajudgement Times	Times for ajudgement					
A B	124 126	21 21	145 147	5 3	0	150 150					
C	124	21	145	5	0	150					
Count											
		Wrong	Missing								
Attendee	Effectiveness	ajudgement	ajudgement	Bias							
Station 1	96.67%	3.88%	0.00%	#DIV/0!							
Station 2 Station 3	98.00% 96.67%	2.33%	0.00%	#DIV/0! #DIV/0!							
อเลแ งก 3	90.07%	3.88%	0.00%	#DIV/0!							
udgement											
Attendee	Effectiveness	Wrong ajudgement Rate	Missing ajudgement Rate	Bias							
Station 1 Station 2 Station 3	Acceptable Acceptable Acceptable	Acceptable Acceptable Acceptable	Acceptable Acceptable Acceptable	#DIV/0! #DIV/0! #DIV/0!							
Symmary:	Acceptable										
Made by		JH		Checked by:		Zhang					

Figure 16 - Example of an Attribute gage study



Element 9 – Dimensional Results

Dimensional results are the measurement results taken off the at least 5 production parts from production tool, mold or set-up. All dimensions (except reference dimensions), characteristics, specifications as noted on the print and print notes should be measured and listed in a convenient format with the actual results recorded.

The supplier shall provide evidence that dimensional verifications required by the design record print/specification have been completed and results indicate compliance with specified requirements. **Dimensional measurement must be done on all cavities, mold, tools, or dies.** All dimensions, characteristics, and specifications as noted on the print/specification and notes should be measured and listed in a convenient format with the actual results recorded.

For multiple cavity molds, the minimum number of parts to measure for the dimensional element is at least 3 parts from each cavity unless otherwise is specified or agreed by Littelfuse Supplier Development Engineer and/or Supplier Quality Engineer representative. All parts should be the same parts that are submitted as Sample Parts to Littelfuse. All parts should be identified with the corresponding number on the part or the tag.

Any requirement that is non-conforming will result in this requirement being deemed unacceptable. The corrective action is required to be addressed and identified. Any concerns identified in the Dimensional test results should be brought to the attention of Littelfuse Engineering or SDE before submitting PPAP.

The Dimensional Results can be provided in any format used within an organization.

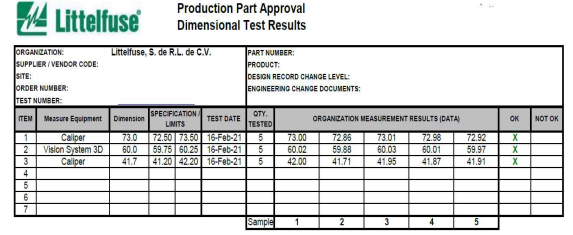


Figure 17 - Example of Dimensional Results

(excel form available in: https://www.littelfuse.com/about-us/supplier-resources.aspx)



Element 10 – Material, Performance Test Results

Material Certifications include any material certifications / material test results relating to the part and the base materials from the supplier's internal lab or outside contracted lab. If there is material specifications noted on the design record/specification, you must provide data that shows conformance to those specifications in the PPAP package.

For products with customer-developed material specifications and a customer-approved subcontractor list, the supplier shall procure materials and/or services (e.g. painting, plating, heat- treating) from subcontractors on that list.

Test Results (Performance, Durability) include any performance or durability tests as prescribed in the design record, including drawings and functional and validation specifications. The required level of detail (i.e. summaries vs. charts vs. raw data) shall be as directed by Littelfuse product engineering.

Supplier need attach the detailed Material/Performance report in the "Material, Performance Test Results" sheets or in a spare sheet.

Littelfuse Expertise Applied Answers Delivered	Product Material		SOLO SELECT MAN CONTRACTOR			
ORGANIZATION: Littelfuse, Inc. SUPPLIER / VENDOR CODE: MATERIAL SUPPLIER: *CUSTOMER SPECIFIED SUPPLIER / VENI	DOR CODE:					
*If source approval is req'd, include the Supplier (Source) &	Customer assigned code.		NAME of LA	ABORATORY:		
MATERIAL SPEC. NO. / REV / DATE	SPECIFICATION / LIMITS	TEST DATE	QTY. TESTED	SUPPLIER TEST RESULTS (DATA)	OK	NOT OK
LF Part Number		Di				
Coil element				See attached	OK	
Solder	2			See attached	OK	
Resin to endbells				See attached	OK	

Figure 18 - Example of Material Performance Test Results

(excel form available in: https://www.littelfuse.com/about-us/supplier-resources.aspx)



Element 11 - Initiall Process Studies (Cpk, Ppk)

Capability or performance shall be determined to be acceptable prior to submission for all special characteristics designated by the customer or supplier. Common requirement for capability study is 25 subgroups containing at least 100 readings and sampled consecutively from a production run that are sampled randomly unless otherwise specified by Littelfuse.

Cpk - The capability index for a stable process. The estimate of sigma is based on within subgroup variation (R-bar/d2 or S-bar/c4). If a supplier is submitting a PPAP but not limited to (a) a new part, (b) a part with revised specifications, (c) a part in which the materials, processes, manufacturing location, or production equipment have significantly changed, or (d) a part in which the material suppliers have changed, then the supplier will be asked to report the Cpk.

Ppk - The performance index. The estimate of sigma is based on total variation (all of individual sample data using the standard deviation [root mean square equation], "s"). If the supplier but not limited to (a) has already been manufacturing the specified part, but is a new supplier to Littelfuse, or (b) is an existing supplier to Littelfuse that has been found to have supplied a large number of nonconforming parts, then the supplier will report Ppk.

Short-term studies. The purpose of the initial process study is to understand the process variation, not just to achieve a specific index value. When historical data is available or enough initial data exist to plot a control chart (at least 100 individual samples), Cpk can be calculated when the process is stable. For chronically unstable processes with output meeting specifications and a predictable pattern, Ppk should be used. When not enough data is available (<100 samples) contact the customer responsible part approval activity to develop a suitable plan.

The minimum required Acceptance Criteria for Initial Study:

- Automotive products: Cpk or Ppk of 1.67
- Non-automotive products: Cpk or Ppk of 1.33

If acceptance criteria cannot be attained by the PPAP submission promise date, the supplier shall submit to the Littelfuse for approval a corrective action plan and a modified Control Plan normally providing for 100% inspection until criteria is achieved or customer full approval is received.

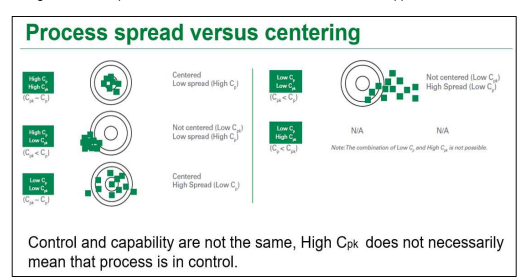


Figure 19 – Process Spread Versus Centering



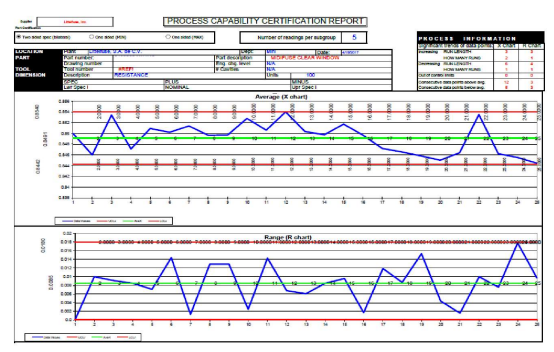


Figure 20 - Example of CPK Chart.

Element 12 – Qualified Laboratory Documentation

If testing is performed in a supplier's internal lab, they must provide a copy of their quality certification. The supplier should also provide documentation of the appropriate laboratory scope. If an external lab is used, send a copy of outside lab certification and the scope of accreditation (Littelfuse prefers that external labs be accredited to known lab accreditation standards such as A2LA and ISO17025 or equivalent).



Figure 21 - Example of a Qualified Laboratory Documentation.



Element 13 - Appearance Approval Report

This requirement is used for more 'print' definition when a specification or design record reference does not exist. Appearance approvals can be used when a specific testing to a known standard or in defining limit samples. This requirement should always be in reference to a specific specification such as color, texture, contrast or paint.

It is not uncommon for projects that have no defined appearance requirements to develop them throughout the course of development. This could be as simple as a paint or color application that has developed into an appearance issue based on Littelfuse feedback or Littelfuse customer feedback. Whenever appearance related issues arise that have no defined specification it is in the best interest of both the supplier and Littelfuse to utilize this element and clearly define what is acceptable and what is not acceptable. When non-conformances arise appearance issues can be readily resolved when there is clear definition of acceptance.

Appearance Approval Report must be provided according to AIAG format.

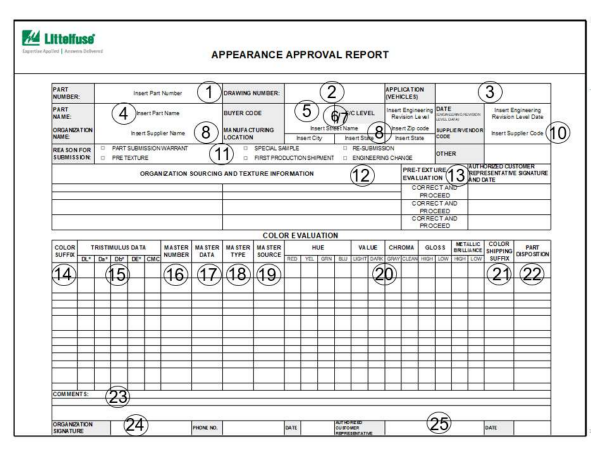


Figure 22 - Example of appearance report

(excel form available in: https://www.littelfuse.com/about-us/supplier-resources.aspx



Completion of the Appearance Approval Report

1	Customer Part Number	Engineering released customer part number
2	Drawing Number	Use the number of the drawing on which the part is shown if different from the part number
3	Application	Enter the model year(s) and vehicle or other program on which the part is used
4	Part Name	Use the finished part name on the part drawing
5	Buyer Code	Enter the code for specific buyer of part
6,7	E/C Level & Date	Engineering change level and E/C date for this submission
8	Organization Name	Organization responsible for submission (include supplier if applicable)
9	Manufacturing Location	Location where part was manufactured or assembled
10	Supplier/Vendor Code	Customer-assigned code for organization location where the part was manufactured or assembled
11	Reason for Submission	Select box(es) explaining the reason for this submission
12	Organization Sourcing & Texture Information	List all first surface tools, graining source(s), grain type(s), and grain and gloss masters used to check part
13	Pre-Texture Evaluation	To be completed by authorized customer representative (not used by GM)
14	Color Suffix	Use alphanumeric or numeric color identification
15	Tristimulus Data	List numerical (colorimeter) data of submission part as compared to the customer-authorized master
16	Master Number	Enter alphanumeric master identification (not used by Ford)
17	Mater Date	Enter the date on which the master was approved
18	Material Type	Identify first surface finish and substrate (e.g., paint/ABS)
19	Material Source	Identify first surface and substrate suppliers, Exampl:Redspot/Dow
20	Color Evaluation, Hue, Value, Chroma, Gloss and Metallic Brilliance	Visual assessment by customer
21	Color Shipping Suffix	Color part number suffix or color number
22	Part Disposition	To be determined by customer (approved or rejected)
23	Comments	General comments by the organization or customer (optional)
24	Organization Signature, Phone No. & Date	Organization certification that the document information is accurate and meets all requirements specified
25	Authorized Customer Representative Signature & Date	Authorized Customer Representative approval signature

THE AREAS INSIDE THE BOLD LINES ARE FOR CUSTOMER USE ONLY Refer to AIAG PPAP Manual.

Element 14 & 15 - Sample product and Master Sample

The Supplier shall provide sample product as requested by the customer and as defined by the submission request.

Supplier shall retain a master sample for the same period as the production part approval records, or

- Until a new master sample is produced for the same customer part number for customer approval,
- Where a master sample is required by the design record. The master sample shall be identified as such, and shall show the customer approval date on the sample.

Supplier shall retain a master sample for each position of a multiple cavity die, mold, tool or pattern.



Element 16 - Checking Aids

Checking aids (fixtures, gages, models, templates etc.) are specific to the part being submitted, used in inspecting or testing. For this item, supplier shall verify that all aspects of the checking aid agree with part dimensional requirements.

Supplier shall document all released engineering design changes that have been incorporated in the checking aid at the time of qualification submission. Suppliers are not required to submit checking aids with qualification unless otherwise is specified; instead they must retain it for future reference. Measurement system analysis studies, e.g. gage R&R, accuracy, bias, linearity, stability studies, shall be conducted in compliance with customer requirements.

Item #	Gage #	Gage ECL	Used for	Location(s)/Process Number(s) on Contro	
1	3D	Α	Measure the size of the material and the product	IQC inspection area	
2	LCR	Α	Measure the capacitor value & inductor value & resistance value of thematerial	IQC inspection area	
3	Paste Gage	В	Measure the paste thickness	SMT SQC Inspection area	

Figure 23 - Example of Checking Aid

Element 17 - Littelfuse Specific Requirements

The Littelfuse specific requirements are must for all Level 3 PPAP submit, unless otherwise specified by Littelfuse Supplier Development Engineer and/or Supplier Quality Engineer representative.

Element 17-1 - Storage conditions

Supplier shall provide in PPAP; the storage conditions and period of time for materials such as but not limited to: plating stamping parts, copper strip, chemical, bulk material, glue, flux, alloy wire, molding part, PCB, PCBA, electronic parts.



Element 17-2 - Certificate of Analysis / Certificate of Compliance

The purpose is to demonstrate that product has passed performance test, quality assurance test, and meets qualification criteria stipulated on drawings or contracts.

The certificate of compliance must minimally include the following information:

- The specific product or type of product certified
- The qualification standard that the product is judged to meet and data that showing the compliance.
- The date of certification (and if applicable, its expiration)

The certificate of analysis can be provided in any format used within an organization.

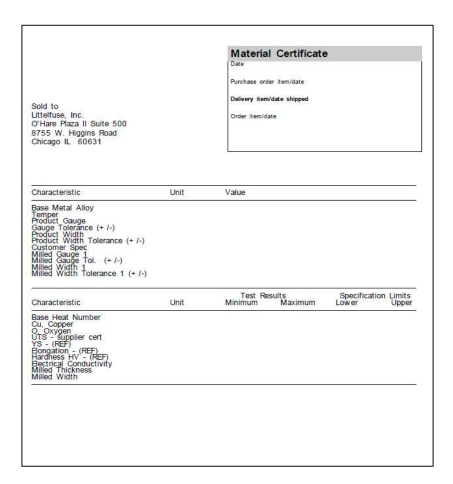


Figure 24 - Example certificate of compliance



Element 17-3 - Preventive Maintenance

Preventive maintenance tends to follow planned guidelines from time-to-time to prevent equipment and machinery breakdown.

The primary goal is to avoid or mitigate the consequences of failure of equipment. This may be by preventing the failure before it actually occurs which Planned Maintenance and Condition Based Maintenance help to achieve. It is designed to preserve and restore equipment reliability by replacing worn components before they actually fail. Preventive maintenance activities include partial or complete overhauls at specified periods, oil changes, lubrication, minor adjustments, and so on. In addition, workers can record equipment deterioration so they know to replace or repair worn parts before they cause system failure. The ideal preventive maintenance program would prevent all equipment failure before it occurs.

The preventive maintenance program can be provided in any format used within an organization.

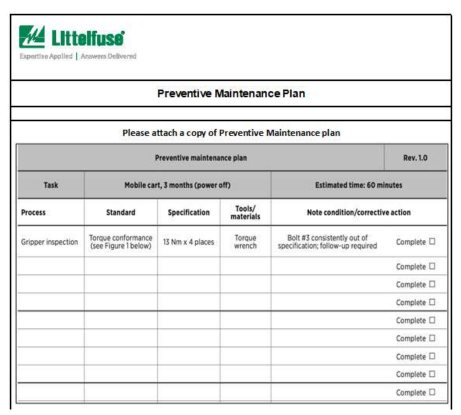


Figure 25 - Example of Preventive Maintenance Plan (excel form available in: https://www.littelfuse.com/about-us/supplier-resources.aspx)



Element 17-4 – Operator Work Instructions

Work instructions should be very detailed on "how" to accomplish a specific job, task or assignment.

Work instructions present a sequence of steps to execute a task or activity. The format is typically text, but a visual depiction of the steps can also constitute work instructions.

A work instruction promotes consistency in execution of work and it is useful when having a frequent turnover of part time helpers.

The work instructions can be provided in any format used within an organization.

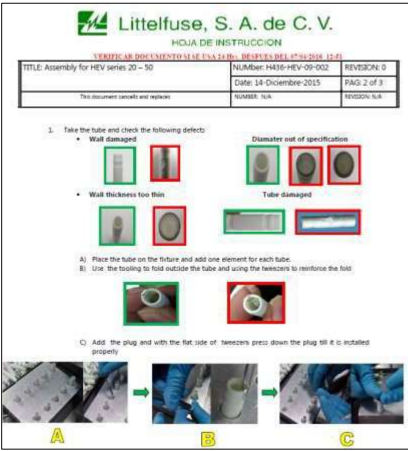


Figure 26 - Example of work instruction.

Element 17-5 - SLC (Safe launch control) Control Plan and S/O

When a new part number has been designed and has been moved into the production phase, Littelfuse requires establishing a pre-launch control plan & procedure in which 100% of the parts manufactured during 1st PO# are contained and inspected.

Please refer to AIAG APQP & control plan manual section 3.7 Pre-launch control plan.



Element 17-6 - IMDS

The IMDS (International Material Data System) is an active tool that aids suppliers in the automotive supply chain to register material data for all components.(the materials they are made from , and the basic substances those materials consist of).

Suppliers must provide the IMDS to Littelfuse thru ID# 2426, and then fill out the PSW format. The module ID#, Version# and date transmitted to Littelfuse must be included.

IMDS report must be also attached on the Littelfuse PPAP format, on section "IMDS".

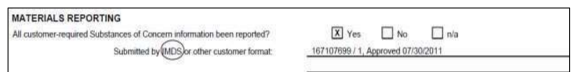


Figure 27 - Example of IMDS.

Element 17-7 - Packing Standard

Purpose is approving the packaging method and material for supplied product. Make sure the package meet all facility related requirements, prevent of shipping and handling defects and addresses any Hazmat related concern. Attach the product package standard / WIs / SOP in this sheet and take the picture at least includes:

- A picture of the part in the packaging position with label A picture of the outside carton with label
- A picture of any dunnage for the container.
- A picture of the final unit load in the shipping configuration.

The Packing Standard can be provided in any format used within an organization

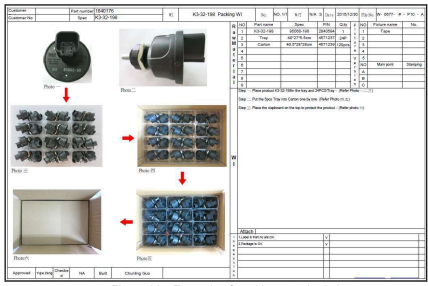


Figure 28 - Example of packing standard sheet.



Element 17-8 - Inspection Plan

The purpose is to approve and record the outgoing inspection requirements, e.g. the inspection items, inspection sample size, inspection gauge etc. Attach the final product outgoing OQC inspection WI / SOP in this sheet.

The Inspection Plan can be provided in any format used within an organization.

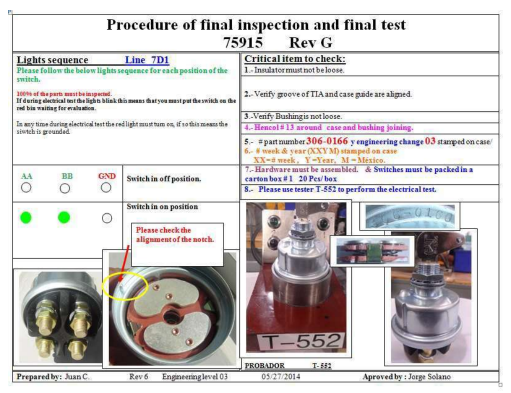


Figure 29 - Example of inspection plan.



Element 17-9 - ISO 9001 / IATF 16949 Certificate

Suppliers must demonstrate compliance with ISO 9001/ IATF 16949 by maintaining a third-party certification issued by a certification body that demonstrates the accreditation mark of an IAF member.

For more details of IAF members visit:

https://www.iaf.nu/articles/IAF Members Signatories/4



Figure 30 - Example of IATF certificate.



Element 17-10 - Sub-contractors PPAP

Sub-contractors PPAP's must be in compliant with the AIAG PPAP standard.

Element 17-11 - Conflict Mineral Report

Littelfuse takes seriously allegations that metals mined in conflict regions throughout the world, including the Democratic Republic of the Congo, may be making their way into the supply chain and that profits from this illegal mining may be fueling human rights atrocities. Littelfuse expects our suppliers to comply with the Electronic Industry Code of Conduct and to only source materials from environmentally and socially responsible suppliers. Littelfuse has procedures in place to help ensure that our suppliers comply with these expectations. In support of this, Littelfuse expects our suppliers to continuously monitor both direct and indirect supply chains to avoid procurement of materials from conflict regions, and to be forthright in sharing compliance information with Littelfuse. Littelfuse suppliers must:

- Comply with all national and other applicable laws and regulations, and require their suppliers do the same (including labor agencies);
- Adopt sound human rights practices and treat workers fairly with dignity and respect.
- Provide a safe and healthy working environment for their workers.
- Conduct business operations in a way that protects and sustains the environment.
- Maintain management systems that measure, improve, and communicate their company's labor, health & safety, environmental performance, and uphold the highest standards of ethics.

For more details of Conflict Mineral Report, contact: pec-cmrt@littelfuse.com



Element 18 – Part Submission Warrant (PSW)

The purpose of the Part Submission Warrant (PSW) is to document the submission and the approval or rejection of purchased parts prior to production. It is critical to make sure the PSW is filled out correctly, and contains accurate and legible information. A sample of the part submission warrant described above can be found below.

Part Name 1	Cust. Part Number 2a
Shown on Drawing No. 3	Org. Part Number
Engineering Change Level	Dated
Additional Engineering Changes5	
Safety and/or Government Regulation Yes 6 No Purchase Ord	(3(1))
Checking Aid No Checking Aid Engineering Change	Level Dated
ORGANIZATION MANUFACTURING INFORMATION	CUSTOMER SUBMITTAL INFORMATION
Supplier Name & Supplier/Vendor Code (12)	Customer Name/Division (14)
Street Address	Buyer/Buyer Code (15)
City Region Postal Code Country	Application
MATERIALS REPORTING	0(6) v
Has customer-required Substances of Concern information been reported	?. Ves No n/a
Submitted by IMDS or other customer format:	
8 -	
Are polymeric parts identified with appropriate ISO marking codes?	Yes No n/a
REASON FOR SUBMISSION (Check at least one) Initial Submission 18 Engineering Change(s) Tooling: Transfer, Replacement, Refurbishment, or additional Correction of Discrepancy Tooling Inactive > than 1 year REQUESTED SUBMISSION LEVEL (Check one) 19 Level 1 - Warrant only (and for designated appearance items, an App Level 2 - Warrant with product samples and limited supporting data states.	
Level 3 – Warrant with product samples and complete supporting dat	ta submitted to customer.
☐ Level 4 – Warrant and other requirements as defined by customer.	
Level 5 – Warrant with product samples and complete supporting dat	ta reviewed at organization's manufacturing location.
SUBMISSION RESULTS (20) The results for dimensional measurements material and function these results meet all design record requirements: Yes NO (If	
Mold / Cavity / Production Process	244.44.14.24
DECLARATION I affirm that the samples represented by this warrent are representative of a Approval Process Manual 4th Edition Requirements. I further affirm that th I also certify that documented evidence of such compliance is on file and avexpLANATION/COMMENTS:	lese samples were produced at the production rate of (23) (24) ours.
25)	
	No 26
Organization Authorized Signature 27	Date
Print Name Phone No	FAX No
Title E-mail	
FOR CUSTOMER USE	ONLY (IF APPLICABLE)
PPAP Warrant Disposition: Approved Rejected Other	*
Customer Signature	Date
Print Name Cus	stomer Tracking Number (optional)

Figure 31 – Part Submission Warrant (PSW).



Completion of the Part Submission Warrant

PART INFORMATION 1 Part Name and 2a. Customer part number: Engineering released finished end item part name and number. 2b Orig. Part Number: Part Number defined by the organization, if any 3 Shown on Drawing Number: The design record that specifies the customer part number beingsubmitted. 4 Engineering Change Level & Date: Show change level and date of the designrecord. 5 Additional Engineering Changes & Date: List all authorized engineering changes not yet incorporated on the drawing but which are incorporated in the part. 6 Safety and/ or Government Regulated: "Yes" if so indicated on part drawing, otherwise "No" 7 Purchase Order Number: Enter this number as found on the purchase order 8 Part Weight: Enter the actual weight inkilograms to four decimal places. 9/10 Checking Aid No. Enter the checking aid number, if one is used for dimensional inspection, and, Its Engineering Change Level and Approval Date ORGANIZATION MANUFACTURING INFORMATION 11 Organization Name & Supplier Code: Show the code assigned to the manufacturing location on the purchase order 12 Supplier Manufacturing Address: Show the complete address of the location where the product was manufactured. CUSTOMER SUBMITTAL INFORMATION 13 Customer Name/Division: Show the corporate name and division or operations group 14 Buyer Name: and Buyer Code: Enter the buyer's name andcode 15 Application: Enter the model year, vehicle name, or engine, transmission, etc MATERIALS REPORTING Substances of Concern: Enter "Yes" "No" or "n/a". IMDS/other customer format: Circle either "IMDS" or "Other customer format" as appropriate 16 If submitted via IMDS include: Module ID#, Version #, and Creation Date. If submitted via other customer format, enter the date customer confirmation was received. 17 Polymeric parts Identification: Enter "Yes" "No" or "n/a". REASON FORSUBMISSION 18 Check the appropriate box. Add explanatory details in the "other" section. REQUESTED SUBMISSION LEVEL 19 Identify the submission level requested by your customer. SUBMISSION RESULTS 20 Check the appropriate boxes for dimensional, material tests, performance tests, appearance evaluation, and statistical data. 21 Check the appropriate box. If "no", enter explanation in "comments" below 22 Molds / Cavity / Production Process: For instruction, see paragraph 2.2.18 DECLARATION 23 Enter the number of pieces manufactured during the significant production run. 24 Enter the time (in hours) taken for the significant production run. Explanation/Comments: Provide any explanatory details on the submission results or any deviations from the Declaration. Additional information may be attached as appropriate. 26 Customer Tool properly tagged and numbered: Are customer-owned tools identified in accord with ISO/IATF 16949 and any customer-specific requirements, answer "Yes" or "No". May not be applicable to OEM internal suppliers. Organization Authorized Signature: A responsible supplier official, after verifying that the results show conformance to all customer requirements and that all required documentation is available shall approve the declaration and provide Title, Phone Number, and Fax Number. FOR CUSTOMER USE ONLY

Element 18 – Part Submission Warrant (PSW) Bulk Materials. (if applicable)

The minimum submission requirement for bulk material is the PSW and the bulk material check list.

For bulk material PPAP only, please refer to AIAG production part approval process manual item F.3 bulk materials requirements check list.

Examples of bulk materials include, but are not limited to: adhesives and sealants (solder, elastomers) chemicals (rinses, polishes, additives, treatments, color/pigments, solvents), coating (top coats, undercoats, primers, phosphates, surface treatments), polymers (rubber, plastics, resins and their precursors).



	Littelfus Expertise Applied Answers De					
Bulk Materials Requirements	s Checklist			Project:		
	Required /		Responsibility	Comments / Conditions	Approve	
Durativet De-	Target Date			Conditions	by / da	
	ign and Developme	iit verilica	ation			
Design Matrix						
Design FMEA						
Special Product Characteristics						
Design Records						
Prototype Control Plan						
Appearance Approval Report						
Master Samples						
Test Results						
Dimensional Results						
Checking Aids						
Engineering Approval						
Process Des	ign and Developme	nt Verific	ation			
Process Flow Diagrams						
Process FMEA						
Special Product Characteristics						
Pre-launch Control Plan						
Production Control Plan						
Measurement System Studies						
Interim Approval						
Prod	uct and Process Val	idation				
Initial Process Studies						
Part Submission Warrant						
Element	s to be Completed a	s Needed	i			
Customer Plant Connection						
Customer Specific Requirements						
Change Documentation						
Supplier Considerations						
Storage conditions and warranty period						
Packaging solutions			+ +			
Operating manual						
-						
n agreed to by: Name / Function			Company / Title / Date			

Figure 32 – Bulk Materials Requirement Checklist