HORIZONTAL RECOMMENDATION FOR USE SHEETS (RfUs) OF THE EUROPEAN COORDINATION OF NOTIFIED BODIES IN THE FIELD OF PERSONAL PROTECTIVE EQUIPMENT (PPE)

REGULATION (EU) 2016/425

Number of RfU PPE-R/	Version	Keywords	Approved by Horizontal Committee	Endorsed by PPE Working
00.001	02	Product checks, time interval, random	30/05/18	Group 22/04/19
00.001	02	Module C2, change of EC / EU type-examination certificate	30/05/18	22/04/19
00.002	03	Sample selection	30/05/18	22/04/19
00.003	03	Module C2 testing	30/03/17	23/01/18
00.004	01	Category III product	30/03/17	23/01/18
00.005	01	Retention of samples	30/05/18	22/04/19
00.008	02	Standard template for report content covering annual	30/05/18	22/04/19
00.008	03	assessment process	30/03/10	22/04/19
00.009	03	Failure of C2 samples	30/05/18	22/04/19
00.009	03	Sample selection / production address(es)	30/03/17	23/01/18
00.010	01	C2 samples and process / production dormant	30/05/18	22/04/19
00.011	03	C2 samples and process / production domain	30/05/18	22/04/19
00.012	03	Sample selection / production address(es)	30/03/17	23/01/18
00.013	01	Module D: Suspension, withdrawal or restriction of	30/03/17	23/01/18
		certificates		
00.015	01	Quality assurance system	30/03/17	23/01/18
00.016	03	Re-assessment of approved quality systems	30/05/18	22/04/19
00.017	03	Module D minimum requirements	30/05/18	22/04/19
00.018	02	Module D minimum requirements	30/05/18	22/04/19
00.019	02	sub-contracting, accreditation, acceptance of test results, competence of laboratories	30/05/18	22/04/19
00.020	03	user information, conformity assessment	30/05/18	22/04/19
00.023	03	EU type-examination procedure, harmonised standards	30/05/18	22/04/19
00.024	03	standards, deficiencies	30/05/18	22/04/19
00.025	01	Testing of materials	12/06/17	23/01/18
00.026	03	identification of test samples	30/05/18	22/04/19
00.027	01	Components from different manufacturers	12/06/17	23/01/18
00.028	03	Marking, standard reference, testing according to prEN	30/05/18	22/04/19
00.029	01	Use of pictograms	12/06/17	23/01/18
00.030	01	Test reports, designation of materials	12/06/17	23/01/18
00.031	01	Slip resistance, type examination certificate	12/06/17	23/01/18
00.032	03	type-examination for category I PPE	30/05/18	22/04/19
00.034	02	information to users	30/05/18	22/04/19
00.035	03	interchangeable components, EU type-examination	30/05/18	22/04/19
00.036	01	Harmonised standards, essential requirements, EU type- examination	12/06/17	23/01/18
00.037	01	Technical documentation	12/06/17	23/01/18
00.038	01	Innocuousness of PPE	12/06/17	23/01/18
00.039	01	Conformity to standard	12/06/17	23/01/18
00.040	01	CE marking, separate items of PPE, technical documentation	12/06/17	23/01/18
00.041	02	information supplied by the manufacturer; sensitising or allergenic substances	30/05/18	22/04/19
00.043	02	uncertainty of measurement	30/05/18	22/04/19
00.043	02	dedicated test method standards	30/05/18	22/04/19
00.045	03	Article 3(1)(b); interchangeable components for equipment referred to in point (a) which are essential for its protective function	16/05/23	31/01/24

Number of RfU PPE-R/	Version	Keywords	Approved by Horizontal Committee	Endorsed by PPE Working Group
00.046	03	interchangeable components	30/05/18	22/04/19
00.047	01	Own brand certificates	12/06/17	23/01/18
00.048	01	Sizing	12/06/17	23/01/18
00.049	01	Traceability of technical documentation	12/06/17	23/01/18
00.050	05	Module C2 or D assessment, EU type-examination certificate	22/11/23	26/05/24
00.052	01	Product marking, reference to standards	12/06/17	23/01/18
00.053	02	instructions for use	30/05/18	22/04/19
00.054	01	Modules C2 and D, non-conform product, unsafe design	12/06/17	23/01/18
00.055	01	Control systems	12/06/17	23/01/18
00.056	03	date of manufacture / obsolescence on the product marking	30/05/18	22/04/19
00.058	03	EU type-examination certificate / re-certification / transitional period	30/05/18	22/04/19
00.061	02	Risk assessment	30/05/18	22/04/19
00.069	04	Issues relating to negative conformity assessment results / refused / withdrawn / suspended / restricted	22/11/23	26/05/24
00.075	01	Renewal date, renewal, review, simplified review	22/11/23	26/05/24

(1) : PPE-R/xx.xxx = Coordination of Notified Bodies/PPE-Regulation/Numbering of the RfUs

* PPE * * * *	CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425			PPE-R/00.001 Version 2
$\rightarrow \star \uparrow$	RECOMMENDA	FION FOR	RUSE	
Number of pages: 1			Approval stage :	Approved on :
Origin : Horizontal Comn	nittee		 Vertical Group Horizontal Committee EU PPE Working Group 	n/a 30/05/2018 22/04/2019
Question related to	☑ PPE Regulation	EN/prEl	N:	Other:
Article:	Annex: VII Module C2	Clause: 4.2		
Key words:				
Product checks, time inte	erval, random			
Question:				
What does "random" me	an in module C2, 4.2?			
0.1.1				
Solution:			. I fan in staat in de staat de staat de st	
manufacturer's advance	and sampling, the interval between visits to knowledge, where possible. Where samples its directly with the people concerned.			

* * * * * PPE * * *	CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425			PPE-R/00.002 Version 3		
* * *	RECOMMEN	DATION FO	R USE			
Number of pages: 1			Approval stage :	Approved on :		
Origin : Horizontal Comm	ittee		 □ Vertical Group ☑ Horizontal Committee ☑ EU PPE Working Group 	n/a 30/05/2018 22/04/2019		
Question related to	PPE Regulation	EN/prE	N:	Other:		
Article:	Annex: VII Module C2	Clause:				
Key words:						
Module C2, change of EC	C / EU type-examination certificate					
Question:						
	t in accordance with module C2 give per ification, should the original EC / EU typ			n those stated in the EC /		
Solution:						
Yes; either the original ce	rtificate is amended or a new certificate	issued, and the	e model reference is changed.			
The procedure set out in	4.4 and 4.6 should be followed.					

* PPE * * * * *	CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425			PPE-R/00.003 Version 3
*	RECOMMENDA	TION FO	RUSE	
Number of pages: 1			Approval stage :	Approved on :
Origin : Horizontal Comm	ittee		 Vertical Group Horizontal Committee EU PPE Working Group 	n/a 30/05/2018 22/04/2019
Question related to	☑ PPE Regulation	EN/prE	N:	Other:
Article:	Annex: VII	Clause:		
	Module C2			
Key words:				
Sample selection				
Question:				
What is the minimum req	uirement to be applied to the method of obt	aining samp	es for testing under module C2	2?
Solution:				
	d body or an independent representative of	the notified	hody shall visit a location agre	ed with the manufacturer
	orter, distributor, retail outlet), and shall rand			
		-		

* * * * PPE * * * *	CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425		PPE-R/00.004 Version 1	
Number of pages: 1	RECOMMENDAT			Approved on :
Number of pages: 1		A	pproval stage :	Approved on :
Origin : Horizontal Comm	littee		 ❑ Vertical Group ☑ Horizontal Committee ☑ EU PPE Working Group 	30/03/2017 23/01/2018
Question related to	☑ PPE Regulation	EN/prEN:	· · · ·	Other:
Article:	Annex: VII 0 Module C2	Clause:		
Key words:				
Module C2 testing				
Question:				
When an EU Type Exami the current version?	ination is based upon a withdrawn standard,	should the C2	2 testing be conducted again	st the withdrawn standard or
Solution:				
Whilst the type examinati demonstrate conformity v	on certificate remains valid, the C2 testing sh vith the Regulation.	ould be again	nst the edition of the standar	d used as a basis to

* * * * PPE * * * *	CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425		PPE-R/00.005 Version 1	
×	RECOMMENDATI	Г		
Number of pages: 1			Approval stage :	Approved on :
Origin : Horizontal Comm	ittee		 Vertical Group Horizontal Committee EU PPE Working Group 	30/03/2017 23/01/2018
Question related to	✓ PPE Regulation] EN/prEN	٧:	Other:
Article:	Annex: VII C Module C2	lause: 4.3		
Key words:				
Category III product				
Question:				
	tegory III because the manufacturer claims one 2 be limited to performance against this / these			r category III, can the tests
Solution:				
No. Once a PPE is claime	ed to meet performance requirements that qua	lify for cat	egory III, for whatever reason,	the entire PPE item is
•••	nd not just single performance requirements.			
There should be no differ be tested on C2 samples	rence in approach between all category III PPE	with resp	ect to deciding which performa	ance requirements should

* PPE * * * * *	CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425			PPE-R/00.006 Version 2
	RECOMMEND	ATION FO		
Number of pages: 1			Approval stage :	Approved on :
Origin : Horizontal Comm	nittee		 Vertical Group Horizontal Committee EU PPE Working Group 	n/a 30/05/2018 22/04/2019
Question related to	☑ PPE Regulation	EN/prE	N:	Other:
Article:	Annex: V and VII	Clause:		
	Modules B and C2			
Key words:				
Retention of samples				
Question:				
Is there any requirement (Module B) or tested duri	in the PPE Regulation for notified bodies to ing the annual control of the final product (N	o retain samp Aodule C2)?	ples of the equipment that they	have type-examined
Solution:				
No.				

* * * * * * * * *			PPE-R/00.008 Version 3	
Number of pages: 3	RECOMMENDA	ATION FOR USE Approval stage :	Approved on :	
Origin : Horizontal Comn	ittee	✓ Vertical Group ✓ Horizontal Committee ✓ EU PPE Working Group	n/a 30/05/2018 22/04/2019	
Question related to	PPE Regulation	EN/prEN:	Other:	
Article:	Annex: VII Module C2	Clause:		
Key words:				
Standard template for re	port content covering annual assessment pr	rocess		
Question: What are the minimum requirements for the report content when implementing Recommendation for Use sheet 00.007? NOTE: RfU 00.007 clearly specifies that 2 separate activities are required when assessing module C2, namely: - 1) Annual selection of samples to confirm continued compliance with the reference standard / specification and the type-examined AND 2) Annual assessment of the production control to determine any evidence of non-homogeneity. Solution: See attached pages 2 and 3				

Confidential

Report number and date:

Module C2 Annual Surveillance Report

Notified Body - name / address / number:

Certificate holder:	Period covered by report
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General Reference Documents:

Recommendation for use sheet, 00.007. PPE Regulation 2016/425/EU, Module C2

EU type-examination certificate numbers covered by the surveillance:

Harmonised standards / technical specifications within the scope of the surveillance:

A. Annual assessment of product compliance with standard / specification and type-examined, reference 2A of RfU 00.007

1. Location(s) visited and dates:

- 2a. Selection carried out by..... Relationship to notified body.....
- 2b. Company representative, name and position.....

2c. Relationship of company visited to type-examination certificate holder

Certificate Holder	Production site
Distributor	Retail Outlet
Other (please specify)	

Importer Secondary production site European office of same company

List of PPE - available

- not available
- not selected
- selected plus lot / batch numbers

3. Attached reference documents

Visit report, number xxxxxxx Test report, number yyyyyy

4. Sample selection was positive / negative. Product testing was positive / negative

5. Sample selection and testing demonstrated compliance with the reference specification / standard and type-examined, yes / no.

B. Annual assessment of production not being homogeneous, reference 2B of RfU 00.007

- 1. Method employed to perform assessment, please specify:
 - 2B(i) On-site review of production and test records.
 2B(ii) On-site audit of production control.
 2B(iii) Production non-homogeneity assessed by selection of a single, large sample.
 2B(iv) Production non-homogeneity assessed by assessment of samples throughout the year.
- 2a. Assessment(s) carried by Relationship to notified body.
- 2b. Company representative, name and position.....

Confidential

Report number and date:

Module C2 Annual Surveillance Report

3. Attached reference documents.

Visit report(s), number xxxxxx Test report(s), number yyyyyyy

4. According to our judgement, the assessment concluded that production was not homogeneous, yes / no.

Justification of nonconformities

Conclusion of notified body:

Overall conclusion of the annual surveillance, positive / negative.

Signature...... Date Date

* * * * PPE * * * *	CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425			PPE-R/00.009 Version 3	
	RECOMMENDA	ATION FO	R USE		
Number of pages: 1			Approval stage :	Approved on :	
Origin : Horizontal Comm	ittee		 Vertical Group Horizontal Committee EU PPE Working Group 	n/a 30/05/2018 22/04/2019	
Question related to	☑ PPE Regulation	EN/prE	N:	⊠ Other:	
Article:	Annex: VII Module C2	Clause: 2B(iv)	RfU	sheet 00.007, 2B(iii) and	
Key words: Failure of C2 samples					
Question: What are the necessary actions following failures when samples are taken as required by recommendation for use sheet 00.007, sections 2B(iii) and 2B(iv), assessment of non-homogeneity?					
Solution:					
The following steps shoul	ld be taken:				
1. Manufacturer asked to	investigate the failure(s) and advise the no	otified body o	f their findings.		
2. The manufacturer mus is to be modified, and how	t inform the notified body whether or not th w.	ey consider t	the product acceptable without	modification or if the product	
3. Notified body to then d	etermine what level of additional testing is	required			
4. Additional samples req	uested from the manufacturer and tested ι	under the aut	hority of the notified body		
5. If additional samples pa	ass the required testing, C2 considered con	mpleted.			
6. If additional samples fa	ail, steps 1 to 4 repeated.				
7. If second set of additio NOTE:	nal samples fail, C2 certification to be with	drawn / not re	e-issued.		
1. If the module C2 body	is not the module B body, module B body t	o be kept inf	ormed throughout the process.		
2. Notifying authorities to	2. Notifying authorities to be informed of failures.				

* * * * * * * * * * * *	PPE	ATION OF NOTIFIE Regulation 2016/4 MMENDATION FOR	425	PPE-R/00.010 Version 1	
Number of pages: 1			Approval stage :	Approved on :	
Origin : Horizontal Comm	iittee		 Vertical Group Horizontal Committee EU PPE Working Group 	30/03/2017 23/01/2018	
Question related to	☑ PPE Regulation	EN/prE	N:	Other:	
Article:	Annex: VII Module C2	Clause: 4			
Key words:					
Sample selection / produ	ction address(es)				
Question:					
Certain tests cannot be p	performed on samples of finished	PPE, but require that n	naterials or components are tes	sted.	
In such cases, how shall appropriate tests carried	samples of materials/component out?	ts be obtained in order t	to satisfy the requirement for sa	amples to be selected and	
Solution:					
Where samples are selected from the production plant, the required material/component samples are to be selected at the same time as the finished PPE, either from the company warehouse or production line.					
	Where samples are selected from the importer or similar, advance notice shall be given that materials and components will have to be made available for selection, and size and quantity requirements specified in advance of the C2 visit.				
	d testing (referring to the PPE pro ponent samples with the material			nfirm the identity of the	

* PPE * * * * *	CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425		PPE-R/00.011 Version 3	
$\sim \sim \times \sim$	RECO	MMENDATION FOR	RUSE	
Number of pages: 1			Approval stage :	Approved on :
Origin : Horizontal Comr	nittee		 Vertical Group Horizontal Committee EU PPE Working Group 	n/a 30/05/2018 22/04/2019
Question related to	☑ PPE Regulation	EN/prEl	N:	Other:
Article:	Annex: VII Module C2	Clause:	RfU she	et 00.007, 2B(iii) and 2B(iv)
Key words:	mound of			
C2 samples and process	/ production dormant			
Question: What are the necessary actions where a company follows module C2 and production is dormant for a period, resulting in module C2 not being able to be carried out?				
Solution:				
1. C2 approval is covered by a separate report or certificate with a 1 year validity.				
2. Where the C2 approval does not have a validity period.				
	g cannot be carried out due to na acturer to inform NB. C2 process			
	g cannot be carried out due to no acturer to inform NB. Product is a			

* * * * * * * *	CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425		PPE-R/00.012 Version 3	
$\sim \sim \times \sim$	RECOMMENDA	ATION FOR	USE	
Number of pages: 1			Approval stage :	Approved on :
Origin : Horizontal Comr	littee		 Vertical Group Horizontal Committee EU PPE Working Group 	n/a 30/05/2018 22/04/2019
Question related to	☑ PPE Regulation	EN/prEN	l:	Other:
Article:	Annex: VII Module C2	Clause:		
Key words:				
C2 samples / frequency	of specific tests			
Question:	of the required C2 tests to be carried out o			
Solution:				
Yes, provided that the pr to have been specified by	inciple has been discussed and agreed by t	the applicable	vertical group, and the tests t	hat this principle could apply

* * * * PPE * * * * *	CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425		PPE-R/00.013 Version 1	
×	RECOMMEND			
Number of pages: 1			Approval stage :	Approved on :
Origin : Horizontal Comr	nittee		 Vertical Group Horizontal Committee EU PPE Working Group 	30/03/2017 23/01/2018
Question related to	☑ PPE Regulation	EN/prEN	l:	Other:
Article:	Annex: VII	Clause:		
	Module C2			
Key words:				
Sample selection / produ	uction address(es)			
Question:				
Module C2.3.a requires	only the manufacturer's address to be spea s procedure cover the actual production sit	cified. In many tes?	cases the production site(s) w	ill not be the manufacturers
	under C2.3 shall also include the producti all apply to all production sites.	on site(s) if the	y are different from the manuf	acturer's address. The

* * * * * * * *	CO-ORDINATION C PPE Regula RECOMMEND	ation 2016/4	25	PPE-R/00.014 Version 1
Number of pages: 1	RECOMMEND	1	Approval stage :	Approved on :
Origin : Horizontal Comm	ittee		 ❑ Vertical Group △ Horizontal Committee △ EU PPE Working Group 	30/03/2017 23/01/2018
Question related to	PPE Regulation	EN/prEN	:	Other:
Article:	Annex: VIII Module D	Clause:		
Key words:				
Module D: Suspension, w	vithdrawal or restriction of certificates			
Question:				
what procedure should b	e followed in the event of failures during n	nodule D asses	isments ?	
Solution:	ring module D accessments, the petified k		t has to deside in each individ	ual case, taking into
	rring module D assessments, the notified b lead to the failure and the risks involved.	body concerned		uai case, taking into
	ajor nonconformities issued against either raw their module D approval; in that case			
NOTE: The failures can c notified body shall be info	concern both quality system failures and promed.	roduct performa	ance failures. In the case of pr	oduct failures, the module B

* * * * PPE * * *	CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425		PPE-R/00.015 Version 1	
· · × × ·	RECOMMENDA	TION FOR	RUSE	
Number of pages: 1			Approval stage :	Approved on :
Origin : Horizontal Comn	nittee		 Vertical Group Horizontal Committee EU PPE Working Group 	30/03/2017 23/01/2018
Question related to	☑ PPE Regulation	EN/prEN	٧:	Other:
Article:	Annex: VIII Module D	Clause:		
Key words:				
Quality assurance system	n			
Question:				
Must existing certificates	relating to QA-Systems (ISO 9001) be acce	epted by a no	tified body?	
Solution:				
	is able to take into account existing certifica cation body (accreditation, mutual recognitions) ts.			

CO-ORDINATION OF NOTIN PPE Regulation 201	PPE-R/00.016 Version 3	
RECOMMENDATION F		
Number of pages: 1 Origin : C2D Ad hoc group	Approval stage : ☐ Vertical Group ⊠ Horizontal Committee ⊠ EU PPE Working Group	Approved on : n/a 30/05/2018 22/04/2019
Question related to PPE Regulation	prEN:	Other:
Article: Annex: VIII Clause: Module D		
Key words: Re-assessment of approved quality systems		
Shall approved quality systems be re-assessed?		
Solution: Yes, at a recommended frequency of every third year, with surveillance audits approval.	s being carried out once per 12 m	onth period following issue of

* PPE * * * *	CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425			PPE-R/00.017 Version 3
	RECOMMEND	ATION FO	R USE	
Number of pages: 5			Approval stage :	Approved on :
Origin : Horizontal Comm	ittee, C2D Ad hoc group		 □ Vertical Group ☑ Horizontal Committee ☑ EU PPE Working Group 	n/a 30/05/2018 22/04/2019
Question related to	☑ PPE Regulation	EN/prE	N:	Other:
Article:	Annex: VIII	Clause:		ISO 9001:2008
	Module D			
Key words:				
Module D minimum requi	rements			
Question:				
	quirements that systems based upon ISC)9001 2008 cd	omplying with module D have to	o cover?
Solution:				
I ne minimum requiremen	its are as attached pages, 2 to 5.			

The system shall be documented in the form of a manual, procedures and supporting work instructions or similar. The scope of the system shall be specified.

Generally the manual submitted in support of an application will be a policy document, which will outline the systems operated, and reference the on-site detailed procedures.

The system requirements are limited to category III PPE, CE marked under the PPE Regulation (EU) 2016/425

Heading, with reference to ISO9001:2008	Comments
4 Quality management system	Shall include or reference quality objectives.
4.1 General requirements	Clear identification and control mechanisms for
Comply with Clause 4.1 of ISO 9001:2008 The quality system ensures compliance of the product with the product described in	any outsourced processes to be documented,
the EC / EU type-examination certificate(s). System shall be documented in the form of manuals, procedures and work instructions.	especially applicable where the company does not manufacture the PPE. Cross reference clause 7.4.1
4.2 Documentation requirements Comply with Clause 4.2 of ISO 9001:20084.2.1 General	To include technical file documents, certificates
4.2.1 General Complies with Clause 4.2.1 of ISO 9001:2008 4.2.2 Quality manual	and external standards, e.g. ENs. To include any
Complies with Clause 4.2.2 of ISO 9001 :2008 4.2.3 Control of documents	external documents that are relevant to the PPE in question, e.g. standards.
Complies with Clause 4.2.3 of ISO 9001:2008 4.2.4 Control of quality records Complies with Clause 4.2.4 of ISO 9001:2008 At least the following documents are retained for at least 10 years after supply of the last item: Those arising from regulatory requirements, to include module D.6 (a) the documentation referred to in point 3.1 of module D (b) the information related to the change referred to in point 3.5 of module D (c) the decisions and reports of the notified body referred to in points 3.5, 4.3 and 4.4 of module D Declaration of conformity, 5.2 module D Training records Inspection and test data Calibration data 5 Management responsibility	Retention period to clearly specify period after supply of the last production item.
5 Management responsibility	
5.1 Management commitment Complies with Clause 5.1 of ISO 9001 :2008	
5.3 Quality policy Complies with Clause 5.3 of ISO 9001:2008	
5.4 Planning	
5.4.1 Quality objectives Complies with Clause 5.4.1 of ISO 9001:2008	

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5.4.2 Quality planning	
Complies with Clause 5.4.2 of ISO 9001:2008	
The quality system ensures compliance of the product with the EC / EU type- examination certificate(s) All adopted elements, requirements and provisions are documented in a systematic and orderly manner in the form of written policies, procedures and instruction.	
5.5 Responsibility, authority and communication	
5.5.1 Responsibility and authority	
Complies with Clause 5.5.1 of ISO 9001:2008	
The following shall be defined:	Position(s) with
A. Need to liaise with notified body responsible for the EC / EU type-examination in case	responsibility and
of changes to the design defined in the EC / EU type-examination certificate and the technical documentation	authority for product quality and contact /
B. Need to liaise with notified body responsible for the assessment of the quality system with respect to updating of the quality system in case of changes of quality system.	advising notified body of any quality system or product problems to be
C. Need to liaise with notified body responsible for the assessment of the quality system with respect to updating of the quality system in case of changes made to the product or technical file	specified.
5.5.2 Management representative Complies with Clause 5.5.2 of ISO 9001 :2008	
5.5.3 Internal communication Complies with Clause 5.5.3 of ISO 9001:2008	
 5.6 Management review 5.6.1 General Complies with Clause 5.6.1 of ISO 9001:2008 A. Intervals should be at least every 12 months, but with a maximum of 14 months B. Top management chairs the review C. The authorized person(s) participate(s) in the review 	The review and audit systems must include those departments /
5.6.2 Review input Complies with Clause 5.6.2 of ISO 9001:2008	positions responsible for compliance with the PPE Regulation.
5.6.3 Review output Complies with Clause 5.6.3 of ISO 9001 :2008	
6 Resource management	
6.1 Provision of resources Complies with Clause 6.1 of ISO 9001 :2008	
6.2 Human resources 6.2.1 General	To include all personnel
Complies with Clause 6.2.1 of ISO 9001:2008	involved in those system
6.2.2.Competence, awareness and training	elements covered by these
Complies with Clause 6,2.2 of ISO 9001 :2008	requirements.
6.3 Infrastructure	
Complies with Clause 6.3 of ISO 9001 :2008	
6.4 Work environment	
Complies with Clause 6.4 of ISO 9001 :2008	

7 Product realization	
7.1 Planning of product realization Complies with Clause 7.1 of ISO 9001:2008	
7.4 Purchasing.	
7.4.1 Purchasing process	
Complies with Clause 7.4.1 of ISO 9001:2008	
Where the processes of manufacture, tests and final inspection are sub-contracted the following shall apply: (the responsibility to ensure compliance to specific requirements cannot be sub-contracted)	
A. Subcontractor has been selected after an evaluation has demonstrated the capability of ensuring compliance with all specific requirements	The Notified Body is
B. The evaluation has been performed by one of the following methods; - third party quality system certification	responsible for ensuring that the manufacturer's
 documented evaluation which provides objective evidence of the capabilities 	quality system complies with module D
- documented site assessment to ensure all relevant capabilities	requirements, and this may include on-site audits
C. Where the features affecting the type of protection cannot be verified at a later stage, the evaluation shall include initial and periodic site assessments at the suppliers premises to ensure that relevant controls are available, documented, understood and effective	of any sub-contracted activities which potentially impact upon conformity
D. Suppliers not used for a period of one year are re-evaluated prior to placing of the contract	with the EC / EU type- examination and / or
E. Ability of supplier is reviewed at least once a year	module D.
7.4.2 Purchasing information Complies with Clause 7.4.2 of ISO 9001:2008	
7.4.3 Verification of purchased products Complies with Clause 7.4.3 of ISO 9001:2008	
A. Verification arrangements are implemented if purchased product can compromise the type of protection	
B. Routine tests or inspections confirmed with declaration of conformity.	
7.5 Production and service operations	
7.5.1 Control of production and service provision Complies with Clause 7.5.1 of ISO 9001:2008	Tracoability is not
Requirements contained in the EC / EU type-examination Certificates are considered.	Traceability is not required. Identification of product is required to
7.5.2 Validation of processes for production and service provision Complies with Clause 7.5.2 of ISO 9001:2008	cover type, batch or serial number, reference
7.5.3 Identification and traceability Complies with Clause 7.5.3 of ISO 9001:2008	Article 8.5
Procedures for product identification during all stages of production, final equipment inspection, testing and placing on the market are established and maintained	
7.5.4 Customer property Complies with Clause 7.5.4 of ISO 9001:2008	
7.5.5 Preservation of product Complies with Clause 7.5.5 of ISO 9001 :2008	

i'	
7.6 Control of measuring and monitoring devices Complies with Clause 7,6 of ISO 9001 :2008	
If the certificate of the calibrated instrument does not bear the accreditation logo of a national authority then the certificate shall contain the following:	
-an unambiguous identification of the item calibrated -traceability to (inter)national standards -the method of calibration -a statement of compliance with any relevant specification	
-the calibration results -the uncertainty of measurement, where relevant -the environmental conditions, where relevant	
-the date of calibration -the signature of the person under whose authority the certificate was issued -the name and address of issuing organization and the date of the certificate -a unique identification of the calibration certificate	
8 Measurement, analyses and improvement	
8.1 General	
Complies with Clause 8.1 of ISO 9001:2008	
8.2 Measuring and monitoring	
8.2.2 Internal audit Complies with Clause 8.2.2 of ISO 9001:2008 The audit program addresses the effectiveness of the elements of the quality system described in this standard. The audits should be carried out at least every 12 months, but within a maximum of 14 months	
8.2.3 Monitoring and measurement of processes Complies with Clause 8.2.3 of ISO 9001:2008	
8.2.4 Measurement and monitoring of product Complies with Clause 8.2.4 of ISO 9001 :2008	
The system shall include inspection and testing from purchased material to finished	
product, to the extent necessary to demonstrate continued compliance with the	
type-examined and the reference product specification, normally a standard. These	
activities shall be carried out by the manufacturer on a regular basis and shall be	
linked to production volumes, time or both.	
To include correct marking of the product, including the CE marking format and user	
information to include NB details.	
 8.3 Control of nonconformity Complies with Clause 8.3 of ISO 9001 :2008 a) There shall be a system for the customer to be identified b) The manufacturer takes action if nonconforming product has been supplied to a supplied to a supplication. 	
customer c) In case of b) the manufacturer informs the customer and the Notified Body responsible for module D supervision. d) In case of b) and the nature of the nonconformity could affect user protection,	
and if not possible to trace the product, then additional actions shall be taken to inform the users, for example, notices placed in appropriate publications e) Product concessions that could affect user safety are not permitted. All concessions	
to be documented and authorised.	
8.4 Analyses of data Complies with Clause 8.4 of ISO 9001:2008	
	To include customer
8.5 Improvement	complaints, warranty
8.5.2 / 8.5.3 Corrective action / Preventive action Complies with Clause 8.5.2 of ISO 9001:2008	returns and returned products
	1

* PPE * * * * *	CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425				
	RECOMMENDATION FOR USE				
Number of pages: 5			Approval stage :	Approved on :	
Origin : Horizontal Comm	ittee, C2D Ad hoc group		 Vertical Group Horizontal Committee EU PPE Working Group 	n/a 30/05/2018 22/04/2019	
Question related to	☑ PPE Regulation	EN/prEl	N:	Other:	
Article:	Annex: VIII Module D	Clause:		ISO 9001:2015	
Key words: Module D minimum requi	rements				
Question: What are the minimum re	Question: What are the minimum requirements that systems based upon ISO9001 2015 complying with module D have to cover??				
Solution:					
The minimum requirement	nts are as attached pages, 2 to 5.				

ISO 9001:2015 Applicability to Module D PPE Regulation

The system shall be documented in the form of a manual, procedures and supporting work instructions or similar. The scope of the system shall be specified.

Generally the manual submitted in support of an application will be a policy document, which will outline the systems operated, and reference the on-site detailed procedures.

The system requirements are limited to category III PPE, CE marked under the PPE Regulation (EU) 2016/425

ISO 9001 2015 clause reference. General compliance plus any specific requirements	Comments / Notes
4.4 Quality management system and its processes	System shall be documented in the form of manuals, procedures and work instructions.
5 Leadership (Section title)	
5.1 Leadership and commitment	
5.2 Policy	
5.3 Organizational roles, responsibilities and authorities	
The following shall be defined:	
A. Need to liaise with notified body responsible for the EU type- examination in case of changes to the design defined in the EU-type examination certificate and the technical documentation	Position(s) with responsibility and authority for product quality and contact /
B. Need to liaise with notified body responsible for the assessment of the quality system with respect to updating of the quality system in case of changes of quality system.	advising notified body of any quality system or product problems to be specified.
C. Need to liaise with notified body responsible for the assessment of the quality system with respect to updating of the quality system in case of changes made to the product or technical file	
6 Planning (Section title)	
6.2 Quality objectives and planning to achieve them	
6.3 Planning of changes	
7 Support (Section title)	
7.1 Resources	
7.1.1 General	
7.1.2 People	To include all personnel involved in those system elements covered by these requirements.
7.1.3 Infrastructure	
7.1.4 Environment for the operation of processes	

7.1.5 Monitoring and measuring resources If the certificate of the calibrated instrument does not bear the accreditation logo of a national authority then the certificate shall contain the following:	
 -an unambiguous identification of the item calibrated -traceability to (inter)national standards -the method of calibration -a statement of compliance with any relevant specification -the calibration results -the uncertainty of measurement, where relevant -the environmental conditions, where relevant -the date of calibration -the signature of the person under whose authority the certificate was issued -the name and address of issuing organization and the date of the certificate -a unique identification of the calibration certificate 	
7.2 Competence	
7.3 Awareness	
7.4 Communication	
7.5 Documented Information	
7.5.1 General	
7.5.2 Creating and Updating	To include technical documentation,
 7.5.3 Control of Documented Information At least the following documents are retained for at least 10 years after supply of the last item: 1.Those arising from regulatory requirements, to include module D.6 (a) the documentation referred to in point 3.1 of module D (b) the information related to the change referred to in point 3.5 of 	certificates and external standards, e.g. ENs. To include any external documents that are relevant to the PPE in question, e.g. standards. Retention period to clearly specify period after supply of the last production item.
 module D (c) the decisions and reports of the notified body referred to in points 3.5, 4.3 and 4.4 of module D Declaration of conformity, 5.2 module D 2. Training records 3. Inspection and test data 	Items 2, 3, 4 to be held for 10 years relative to each production lot / batch.
4. Calibration data	
8 Operation (Section title)	
8.1 Operational planning and control	

8.4 Control of externally provided processes, products and services		
If manufacture, tests and / or final inspection is sub-contracted the responsibility to ensure compliance to specific requirements cannot be sub-contracted.		
Controls to be applied where manufacture or testing or inspection is subcontracted:		
A. Subcontractor has been selected after an evaluation has demonstrated the capability of ensuring compliance with all specific requirements	The Notified Body is responsible for ensuring that the manufacturer's quality	
B. The evaluation has been performed by one of the following methods;	system complies with Module D requirements, and this may include on-site assessment of any subcontracted activities	
 third party quality system certification documented evaluation which provides objective evidence of the capabilities 	which potentially impact upon conformity with the EU type-examination and / or Module D.	
- documented site assessment to ensure all relevant capabilities		
C. Where the features affecting the type of protection cannot be verified at a later stage, the evaluation shall include initial and periodic site assessments at the suppliers premises to ensure that relevant controls are available, documented, understood and effective		
D. Suppliers not used for a period of one year are re-evaluated prior to placing of the contract		
E. Ability of supplier is reviewed at least once a year		
8.4.1 General		
8.4.2 Type and extent of control	A. Verification arrangements are implemented if purchased product can compromise the type of protection	
8.4.3 Information for external providers	B. Routine tests or inspections confirmed with declaration of conformity.	
8.5 Production and service provision		
8.5.1 Control of production and service provision		
8.5.2 Identification and traceability	Traceability is not required.	
Procedures for product identification during all stages of production, final equipment inspection, testing and placing on the market are established and maintained	Identification of product is required to cover type, batch or serial number, reference Article 8.5	
8.5.3 Property belonging to customers or external providers		
8.5.4 Preservation		
8.5.5 Post-delivery activities		
8.5.6 Control of changes		
8.6 Release of products and services		
8.7 Control of nonconforming outputs		

9 Performance evaluation (Section title)		
9.1 Monitoring, measurement, analysis and evaluation		
The system shall include inspection and testing from purchased material to finished product, to the extent necessary to demonstrate continued compliance with the type-examined and the reference product specification, normally a standard. These activities shall be carried out by the manufacturer on a regular basis and shall be linked to production volumes, time or both. To include correct marking of the product, including the CE marking format and user information to include NB details.		
9.1.1 General		
9.1.3 Analysis and evaluation		
9.2 Internal audit		
The audit program addresses the effectiveness of the elements of the quality system described in this standard. The audits should be carried out at least every 12 months, but within a maximum of 14 months		
9.3 Management review		
A. Intervals should be at least every 12 months, but with a maximum of 14 months	The review and audit systems must include those departments / positions responsible for compliance with the PPE	
B. Top management chairs the review	Regulation.	
C. The authorized person(s) participate(s) in the review		
10 Improvement (Section title)		
10.1 General		
10.2 Nonconformity and Corrective Actiona) There shall be a system for the customer to be identified		
b) The manufacturer takes action if nonconforming product has been supplied to a customer		
c) In case of b) the manufacturer informs the customer and the Notified Body responsible for module D supervision.	To include customer complaints, warranty returns and returned	
d) In case of b) and the nature of the nonconformity could affect user protection, and if not possible to trace the product, then additional actions shall be taken to inform the users, for example, notices placed in appropriate publications	products.	
e) Product concessions that could affect user safety are not permitted. All concessions to be documented and authorised.		

* PPE * * * * *	CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425				PPE-R/00.019 Version 2
~	RECOMMEND				
Number of pages: 1			Appro	val stage :	Approved on :
Horizontal Committee			n/a 30/05/2018 22/04/2019		
Question related to	PPE Regulation	EN/prE	۷:		Other:
Article: 26	Annex:	Clause:			
Key words: sub-contracting, accredita	ation, acceptance of test results, competer	nce of laborato	ories		
Are test reports from auth If this is so, what are the What quality control meth Can the notified body use	ation body to accept test data obtained by norities outside the European Union accep minimum criteria to be used in judging the nods should be applied to sub-contracting test reports on materials, items or compo e reports on tests carried out by the manuf	table for the p ir competency laboratories? onents carried	urpose and h out by	e of CE marking? low should they be monit r other specialised labora	
Solution:					
Under all circumstances,	the notified body takes on the responsibili	ty for test resu	lts/tes	t reports it accepts as the	e basis for certification.
Therefore, it should gene	rally be recommended to use test results f	from accredite	d test	laboratories only.	
	possible, other sources of testing have to 7025, if this is not the case, the notified bo				
	ill have to specify the conditions for the ac eets the requirements set out in Article 26				ut the tests. It shall ensure
Quality control measures proceed with this.	for sub-contracting test laboratories are in	nportant, the r	notified	I body itself is responsibl	e for deciding how to

* PPE * *	CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425			PPE-R/00.020 Version 3
	RECOMMEND	ATION FOR USE		
Number of pages: 1		Approv	al stage :	Approved on :
Origin : Horizontal Comr	nittee	🖂 Hoi	rtical Group rizontal Committee I PPE Working Group	n/a 30/05/2018 22/04/2019
Question related to	PPE Regulation	EN/prEN:		Other:
Article:	Annex: III, k	Clause:		
Key words: user information, conform	nity assessment			
certification procedures Solution: The notified body can ch	of the user information will be assessed in for foreign manufacturers?	issessing. Any translat	tion is the responsibility	of the manufacturer /

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Number of pages: 1	RECOMMEND		USE pproval stage :	Approved on :	
Origin : Horizontal Comm	ittee				
☐ Vertical Group ☑ Horizontal Committee			n/a 30/05/2018 22/04/2019		
	☑ PPE Regulation	EN/prEN:		Other:	
Article:	Annex: III, g; V, 4f	Clause:			
Key words:					
•	cedure, harmonised standards				
	Question: What is the procedure to be applied to the EU type-examination in the absence of approved harmonised standards covering product requirements and / or test methods?				
Solution:					
The manufacturer has to	decide what will be the basis for testing ag set the specification for the product and as ications of the manufacturer will remain st	sk for certificatio	on against this specification.		
The notified body is respo	onsible for assessing whether or not the sp ot the submitted PPE does comply with the	ecification mee		ts of annex II and	
	er to existing standards (e.g. national or IS				
	notified body should identify the objectives ppropriate for the EU type-examination.	s to be reached	in testing for conformity with	the requirements and	
	ay be discussed with the other notified bod bject should be brought into the Europear			nterest in a harmonization of	
Note: The references of to	echnical specifications must be included ir	n the EU type-ex	camination certificate and EU	I declaration of conformity.	

* * * * * * * *				PPE-R/00.024 Version 3
Number of pages: 1	RECOMM		al stage :	Approved on :
Origin : Horizontal Comm	nittee		a stage .	
		🖂 Hor	tical Group izontal Committee PPE Working Group	n/a 30/05/2018 22/04/2019
Question related to	☑ PPE Regulation	EN/prEN:		Other:
Article:	Annex:	Clause:		
Key words: standards, deficiencies				
Question: What action should be ta	ken if deficiencies or mistakes in sta	ndards are detected?		
Solution: Deficiencies and mistakes in standards always have to be discussed in the relevant CEN or CENELEC TC or WG. Therefore, as soon as any such mistake is recognised, the appropriate body has to be informed and asked to take action for a possible revision of the standard as soon as possible. In addition to that, the problem should be discussed within the vertical group so that a general approach to the problem is laid down and the notified bodies can agree how to proceed with the testing before a revision of the standard is published. The relevant TC or WG should be informed of any such interim solution. If the problem is of general interest, the Horizontal Committee should be informed so that the subject can be discussed at Horizontal Committee level and, if necessary, with the relevant CEN or CENELEC authorities. The European Commission will receive the existing Recommendation for Use sheets for approval.				

PPE Regu	PPE-R/00.025 Version 1	
KECOWWENI	Approval stage :	Approved on :
nittee		
	Horizontal Committee	
⊠ PPE Regulation	EN/prEN:	Other:
Annex:	Clause:	
		nstead of carrying out tests on
rer attests (in writing) that it is strictly ider by examination of the reference PPE and ferring to high cost PPE produced in sma ply one example of the PPE submitted to	ntical to that used in the construction to the the samples supplied. This procedure sho all quantities. EU type-examination so that the notified	e PPE and if the notified body ould be limited to a specific case
	PPE Regu RECOMMENT inittee PPE Regulation Annex: out tests on materials, parts or component at are the conditions to be met for type-ex- tests on materials described in the stand rer attests (in writing) that it is strictly iden by examination of the reference PPE and ferring to high cost PPE produced in small ply one example of the PPE submitted to	nittee □ Vertical Group ⊠ Horizontal Committee ⊠ EU PPE Working Group ⊠ PPE Regulation □ EN/prEN:

* * * * * * * * *	CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425 RECOMMENDATION FOR USE			
Number of reveal 4	RECOMMEND			
Number of pages: 1			Approval stage :	Approved on :
Origin : Horizontal Comr	nittee		 □ Vertical Group ☑ Horizontal Committee ☑ EU PPE Working Group 	n/a 30/05/2018 22/04/2019
Question related to	PPE Regulation	EN/prEl	N:	Other:
Article:	Annex:	Clause:		
Key words: identification of test sam	ples			
Question: What measures should I	be adopted to ensure traceability and ident	tification of spe	cific PPE product to the origina	al type?
examination.	uity regarding the identification of the PPE at are the subject of the tested type declar	-		a notified body for EU type-
The following is recomm	ended:			
 the alphanume 	eric reference of the models must be provid	ded by the mar	nufacturer with an indication of	its meaning
	ns needed for correct identification of the F ved with the file by the notified body	PE must acco	mpany the certificate and a co	py of these photographs
 an example of 	the PPE model can be archived by the no	tified body whe	en this is possible.	

* * * * * PPE * * * *	CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425 RECOMMENDATION FOR USE			PPE-R/00.027 Version 1
Number of pages: 1	RECOMMEND			Approved on :
Number of pages: 1 Origin : Horizontal Com	mittee	Ap	proval stage :	Approved on :
ongin . Honzontai oon		\square	Vertical Group Horizontal Committee EU PPE Working Group	12/06/2017 23/01/2018
Question related to	☑ PPE Regulation	EN/prEN:		Other:
Article:	Annex:	Clause:		
Key words: components from differ	ent manufacturers			
Question: Should a notified body agree to issue an EU type-examination for a product submitted by manufacturer "A" which includes interchangeable components produced by a manufacturer "B" where the product requires to be tested as a complete device? for example: a) filters for an air powered device b) chemical protective clothing without a hood and/or boots c) helmet mounted ear muffs				
Provided the client's do necessary tests to be o Note: It is the manufa	onsible for reviewing the technical documentation submitted covers all the applica arried out and if found satisfactory issue an E acturer "A"'s responsibility to monitor that eac on and that the product manufactured by "B"	ble requirements EU type-examina n subsequent pro	the notified body may perf tion certificate. oduct is in conformance with	form or arrange for the
(see also RfUs 00.035	and 00.045, 00.046)			

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Number of pages 1	RECOMME	NDATION FOR U		Approved on t
Number of pages: 1	·····	Aļ	oproval stage :	Approved on :
Origin : Horizontal Comm	IIIee		 Vertical Group Horizontal Committee EU PPE Working Group 	n/a 30/05/2018 22/04/2019
Question related to	☑ PPE Regulation	EN/prEN:		Other:
Article:	Annex:	Clause:		
Key words: Marking, standard referen	nce, testing according to prEN			
Question: If only a prEN is available at the time of EU type-examination, can the product be marked with the standard number "EN"? When a product is marked with a prEN, can EN be marked on the product, once the standard is ratified?				nber "EN"?
Where a manufacturer de As long as no final standa If the ratified EN is idention Where the ratified EN is r Marking with a prEN is no examination then it should	reference is not mandatory by the PPE ecides to mark a standard or prEN on I ard exists or the final standard is not ic cal to the prEN, then "EN" may be m not identical to the prEN, then "EN" (ot recommended. However, where a m d be fully identified by year and/or issu or EN do not give presumption of conf gulation).	his product, the follow dentical with the prEM narked on the product cannot be marked or nanufacturer decides ue.	N, the marking cannot be "El ct, subject to confirmation by n the product. s to mark with the prEN used	the notified body. for the EU type-

* PPE * * * *	CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425			PPE-R/00.029 Version 1
	RECOMMEND			
Number of pages: 1		A	pproval stage :	Approved on :
Origin : Horizontal Comm	ittee	D] Vertical Group ☑ Horizontal Committee ☑ EU PPE Working Group	12/06/2017 23/01/2018
Question related to	☑ PPE Regulation	EN/prEN:		Other:
Article:	Annex: II, 1.4	Clause:		
Key words: use of pictograms				
	oduct with a pictogram described in an El I standard or other technical specification		the verification of essential	requirements has been
The notified body, in revi meaning of the pictogram	ictogram even if the standard used is not ewing the manufacturer's instructions for a is clearly defined in respect of the essen to the pictorial presentation; this does no for testing.	use (information tial health and sa	supplied by the manufacture afety requirements of the PP	r), must ensure that the E Regulation.

* * * * * * * * *	CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425		PPE-R/00.030 Version 1	
Number of pages 1	RECOMMENDA			Approved on t
Number of pages: 1			Approval stage :	Approved on :
Origin : Horizontal Comm	ittee		 Vertical Group Horizontal Committee EU PPE Working Group 	12/06/2017 23/01/2018
Question related to	PPE Regulation	EN/prEN	J:	Other:
Article:	Annex:	Clause:		
Key words: test reports, designation of	of materials			
Question: In test reports, materials are often only referred to by a single, mostly commercial reference name. In many cases, however, this name covers a variety of materials different by structure and weight (for fabrics) or by origin and thickness (for leather). Is it possible to have a uniform and clear "finger print designation" of materials in test reports in order to make an evaluation easier? For this purpose, we propose to use the elements as given below: - aramid twill 2/1 - 270 g/m² - cow split 1.3 - 1.5 mm.				
	er or name identifying the material must be tion should contain a documentation of the			

* * * * * PPE * * *	CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425			PPE-R/00.031 Version 1
×	RECOMMENT	DATION FOR US	E	
Number of pages: 1		Appro	oval stage :	Approved on :
Origin : Horizontal Comm	ittee	🖂 H	/ertical Group lorizontal Committee EU PPE Working Group	12/06/2017 23/01/2018
Question related to	PPE Regulation	EN/prEN:		Other:
Article:	Annex:	Clause:		
Key words: Slip resistance, type exan	nination certificate			
Question: Does slip resistance have	to be considered an essential requireme	ent for safety, protect	tive and occupational foo	twear?
Notified bodies have to ca	al feature of safety, protective and occup arry out slip resistance testing, unless the ear does not meet this requirement.		y claims in his product sp	ecification and in the user

* PPE * * * * *	CO-ORDINATION O PPE Regula	PPE-R/00.032 Version 3	
\uparrow \star \uparrow	RECOMMEND	ATION FOR USE	
Number of pages: 1		Approval stage :	Approved on :
Origin : Horizontal Comn	nittee	 □ Vertical Group ☑ Horizontal Committee ☑ EU PPE Working Group 	n/a 30/05/2018 22/04/2019
Question related to	☑ PPE Regulation	EN/prEN:	Other:
Article:	Annex:	Clause:	
Key words:			
type-examination for cate	egory I PPE		
Could PPE which do not	belong to categories II or III be submitted to	o an EU type-examination on a voluntary l	pasis?
Solution: No, only PPE belonging examination certificate.	to categories II or III can be submitted to ar	n EU type-examination procedure leading	to the issue of an EU type-

* PPE * * * * *		N OF NOTIFIED BODIES Julation 2016/425	PPE-R/00.034 Version 2
	RECOMME	NDATION FOR USE	
Number of pages: 1		Approval stage :	Approved on :
Origin : Horizontal Com	mittee	 □ Vertical Group ⊠ Horizontal Committee ▲ EU PPE Working Group 	
Question related to	PPE Regulation	EN/prEN:	Other:
Article:	Annex: II, 1.4	Clause:	
Key words: information to users			
Question: What is the responsibili	ty of the notified body in checking the inf	formation to users?	
body shall check that the specification used and information as required these requirements and manufacturer has the fin Note : Claims of comp	e claims of the manufacturer on the area with the relevant essential safety require by annex II, clauses 1.4, 2 and 3. The n I that it does not contain misleading state nal responsibility for the accuracy of the liance with standards other than harmon xamination or claims that are not related	complete safety for its intended purpose. In a and limits of protection of the product are in ments. One of the essential safety requirem otified body must check that the information ements and obvious mistakes concerning the content including translations. ised European standards that have the sam to user protection, e.g. value for money etc	n line with the technical ents is to supply all relevant is given in accordance with e protection provided. The e scope as those used as a

* PPE * * * * *	CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425		PPE-R/00.035 Version 3		
$\sim \sim \times \sim$	RECOMMEN	NDATION FOR L	JSE		
Number of pages: 1		Ар	oproval stage :	Approved on :	
Origin : Horizontal Comm	nittee		•	n/a 30/05/2018 22/04/2019	
Question related to	☑ PPE Regulation	EN/prEN:		Other:	
Article: 3 (1) (b)	Annex:	Clause:			
Key words: interchangeable compon	ents, EU type-examination				
Question:					
Should interchangeable of Should interchange	Should interchangeable components be subject to a separate EU type-examination?				
Yes, an EU type-examination certificate must be issued in accordance with Article 3 (1) (b). The notified body shall carry out sufficient evaluation and/or testing to verify their suitability for the stated equipment in its final assembly.				nent in its final assembly.	
(See also RfUs 00.027, 0	00.045, 00.046)				

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Number of pages: 1	·	A	pproval stage :	Approved on :
Origin : Horizontal Comm	ittee		☐ Vertical Group ☑ Horizontal Committee ☑ EU PPE Working Group	12/06/2017 23/01/2018
Question related to	☑ PPE Regulation	EN/prEN:	· · · · · · · · · · · · · · · · · · ·	Other:
Article: 25	Annex:	Clause:		
Key words:				
harmonised standards, e	ssential requirements, EU typ	pe-examination		
Question:				
	l type-examination, what is th relevant essential health and		d body when the applicable h	armonised product standard
Solution:				
must identify those not ac is responsible for confirm	ised product standard does r ddressed in the standard and ing that all the relevant esser eir review, inspection and tes	also state how these are deantial health and safety require	alt with in his technical docur ements have been identified,	nentation. The notified body
	product standard gives a pres e product and addresses.	sumption of conformity with the	nose essential health and saf	ety requirements which it
	embered that the Regulation is nay demonstrate his complian			one means by which a

* * * * * * * * * *				PPE-R/00.037 Version 1
	RECOMMEND			
Number of pages: 1		A	Approval stage :	Approved on :
Origin : Horizontal Com	mittee		 ☐ Vertical Group ☑ Horizontal Committee ☑ EU PPE Working Group 	12/06/2017 23/01/2018
Question related to	PPE Regulation	EN/prEN	:	Other:
Article:	Annex:	Clause:		
Key words: technical documentation	n			
Question: How should the notified	body "verify" that the model is the product $\boldsymbol{\theta}$	described in the	e manufacturer's technical doc	sumentation?
	action in order to verify that a PPE model h			
	Iduct a visual comparison between an exameted at the product is as de that, in general terms, the product is as de			
Note: The description descriptions, etc	of the model may take various forms, e. g. c.	general assemb	oly drawings, component draw	vings, photographs, material

* * * * * PPE * * *	CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425			PPE-R/00.038 Version 1
××	RECOMMENDATION	FOR	USE	
Number of pages: 1		/	Approval stage :	Approved on :
Origin : Horizontal Comm	ittee		 □ Vertical Group ☑ Horizontal Committee ☑ EU PPE Working Group 	12/06/2017 23/01/2018
Question related to	PPE Regulation	l/prEN	l:	Other:
Article:	Annex: II, 1.2.1.1 Claus	ə:		
Key words: innocuousness of PPE				
Question:	es require from the manufacturer to demonstrate			
Solution:				
Compliance may be demo that are known to, or susp	onstrated by a written declaration confirming that to bected to, adversely affect user hygiene or health, as required by harmonised standards will not be	if pres	ent; a list of these substances	

* * * * PPE * * * * *				PPE-R/00.039 Version 1
Number of pages: 1			Approval stage :	Approved on :
Origin : Horizontal Comn	nittee			
			 □ Vertical Group ☑ Horizontal Committee ☑ EU PPE Working Group 	12/06/2017 23/01/2018
Question related to	☑ PPE Regulation	EN/prEN	l:	Other:
Article:	Annex:	Clause:		
Key words: conformity to standard				
Question:				
	product in compliance with a standard when	re one or more	e requirements of the standard	are not satisfied?
Solution: No.				
NOTE: The product may	be certified in compliance with the essent	ial health and s	safety requirements of the Reg	gulation.

* * * * * PPE * * *	CO-ORDINATION OI PPE Regulat	PPE-R/00.040 Version 1			
· · × × ·	RECOMMENDA	TION FOR USE			
Number of pages: 1		Approval stage :	Approved on :		
Origin : Horizontal Comn	nittee	☐ Vertical Group⊠ Horizontal Committee☑ EU PPE Working Group	12/06/2017 23/01/2018		
Question related to	☑ PPE Regulation	EN/prEN:	Other:		
Article:	Annex:	Clause:			
Key words: CE marking, separate ite	ems of PPE, technical documentation				
1. Is it possible to					
1. It is possible to	e submit one technical documentation only fo luct must be CE marked.	or all products.			

* P * P * P	CO-ORDINATION OF NOT PPE Regulation 20	PPE-R/00.041 Version 2		
× ×	RECOMMENDATION	FOR USE		
Number of pages: 1		Approval stage :	Approved on :	
Origin : Horizontal Comm	ittee	 □ Vertical Group ⊠ Horizontal Committee ⊠ EU PPE Working Group 	n/a 30/05/2018 22/04/2019	
Question related to	☑ PPE Regulation	/prEN:	Other:	
Article:	Annex: II, 1.2.1.1 Clause): :		
information supplied by the manufacturer; sensitising or allergenic substances Question: Should the manufacturer of PPE display all substances with sensitising or allergenic potential in the "information supplied by the manufacturer", if the PPE is designed to get (even if only partly) in close skin contact with the user, or if parts of the PPE may be released and taken up via the inhalation route by the user?				
Solution: Yes. Annex II, 1.2.1.1 (suitable constituent materials) requires that "PPE materials and parts, including any of their decomposition products must not adversely affect user hygiene or health". In case that PPE contains substances which are known to be potentially sensitising or allergenic, the manufacturer has to display each individual relevant substance in the information supplied by the manufacturer to give a warning to all potentially concerned users.				

PPE Regulation 2016	CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425				
RECOMMENDATION FO					
Number of pages: 1	Approval stage :	Approved on :			
Origin : Horizontal Committee	 □ Vertical Group ⊠ Horizontal Committee ⊠ EU PPE Working Group 	n/a 30/05/2018 22/04/2019			
Question related to PPE Regulation I EN/pre	EN: 17025	Other:			
Article: Annex: Clause: 5.	10.3.1 c)				
Key words: uncertainty of measurement					
Question: When notified bodies commission testing on test laboratories complying with EN/ISO/IEC 17025, and the reference specification includes pass / fail criteria, does the notified body have to make a specific request for uncertainty of measurement to be included in the test report?					
Solution:					
No.					
EN/ISO/IEC 17025 includes a clear requirement for uncertainties of measureme affect compliance with pass / fail criteria. In such cases, the test laboratory has t		where the uncertainty might			

* * * * * * * * *	CO-ORDINATION OF PPE Regulat RECOMMENDA	PPE-R/00.044 Version 3			
Number of pages: 1		A	pproval stage :	Approved on :	
Origin : Horizontal Comm	ittee] Vertical Group] Horizontal Committee] EU PPE Working Group	n/a 30/05/2018 22/04/2019	
Question related to	PPE Regulation	EN/prEN:		Other:	
Article:	Annex:	Clause:			
Key words:					
dedicated test method sta	andards				
Question:					
	refer to specific standards or other sources regard to the interpretation of test results fo				
What should notified bod	ies do when a test method standard is revis	ed?			
Solution:					
As long as the product st has to be used.	andard has not been revised and there is a	n undated refer	rence in the standard, the lat	est version of the test method	
	NOTE: If a test method standard has been revised, the consequences for the interpretation of test results should be discussed in the PPE notified body coordination group and an amendment to the product standard be proposed as quickly as possible, if necessary.				

RECOMMENDATION FOR USE Number of pages: 1 Approval stage : Approval stage :	red on :				
	ed on :				
Origin : Horizontal Committee □ Vertical Group ⊠ Horizontal Committee 16/05/2023 ⊠ EU PPE Expert Group 31/01/2024					
Question related to PPE Regulation PPE Guidelines EN/prEN:					
Article: : 3 (1) (b) Annex: Clause:					
Key words: Article 3(1)(b); interchangeable components for equipment referred to in point (a) which are essential for its protective function	on				
Question: Who can apply for EU type-examination of interchangeable components in the meaning of Article 3, (1) (b) of the PPE Regula	tion?				
Solution:					
The situation will depend on the PPE presented. Some performance standards have design restrictions that present a common/standardised approach (e.g. EN 148-1 "Respiratory protective devices - Threads for facepieces - Part 1: Standard connection) - such that the PPE continues to perform provided each Manufacturer produces in conformity with the standard. any applicant may apply.					
Where no such standard exists, the Notified Body must assess the risk of a change by one manufacturer, on the potential conformity of the second manufacturer proposing an interchangeable part. In this case the manufacturer of the interchangeable component must ensure it is identical to the manufacturer's original components in safety performance, and there must be a contractual agreement between them, which authorises the manufacturer of the interchangeable component.					
(see also RfU 00.027, 00.035, 00.046)					

* * * * * * * *				PPE-R/00.046 Version 3		
Number of pages: 1	KEGOWW		oroval stage :	Approved on :		
Origin : Horizontal Comm	ittee		Vertical Group Horizontal Committee EU PPE Working Group	n/a 30/05/2018 22/04/2019		
Question related to	PPE Regulation	EN/prEN:		Other:		
Article: 3 (1) (b)	Annex:	Clause:				
Key words: interchangeable compone	ents					
	Question: Do interchangeable components of protective equipment that was placed on the market before the PPE Directive became effective fall under the scope of the PPE Regulation?					
The suitability of the comp body must have access to	nent is not CE marked, such interch ponent for the intended use of the F to the complete documentation conc ming equivalence with the part to be	PPE in the protective equerning the whole equipn	ipment must be assessed nent (test reports and certif	and certified. The notified		

* * * * * PPE * * * * *	CO-ORDINATION O PPE Regula	PPE-R/00.047 Version 1	
Number of pages: 2	RECOMMENDA	ATION FOR USE Approval stage :	Approved on :
Origin : Horizontal Comm	nittee		
		 □ C2 / D ad hoc group ☑ Horizontal Committee ☑ EU PPE Working Group 	12/06/2017 23/01/2018
Question related to	☑ PPE Regulation	EN/prEN:	Other:
Article:	Annex:	Clause:	
Key words:			
own brand certificates			
Question:			
How should applications	for own brand certificates be dealt with?		
Solution:			
See attached			

Own brand manufacturers type-examination certificates, Module B.

Any person / company / organisation placing a product on to the market in their own name is the manufacturer of that product within the terms of the PPE Regulation. It then follows that the manufacturer must make an application in their own name and be issued with certificates that support the CE marking of the PPE.

It is common practice for original manufacturers to offer their product to one or more companies who wish to sell the product as their own. There will be no identifiable link back to the original manufacturer in the market place.

The product offered for sale by the own brand manufacturer will be identical to the original product except for marking and probably user instructions. All other elements of the technical documentation can be applied to the own brand product.

The own brand manufacturer will be required to raise and sign a declaration of conformity before placing CE marked product on the market. This will include a statement covering modules C2 or D for category III PPE.

The own brand certificate will only remain valid while the cross-referenced original certificate remains valid.

As the own brand manufacturer is legally responsible for ensuring that the product(s) meet the requirements of the PPE Regulation, it is necessary for the minimum acceptable level of control and responsibilities to be established with the PPE notified bodies.

Proposed conditions to be fulfilled before the granting of certification for an own brand product: -

- 1. The original manufacturer to hold a valid type-examination certificate and if category III, to provide evidence of current supervision in line with module C2 or D.
- 2. Written agreement to be submitted, signed by both parties (original manufacturer and own brand manufacturer), covering the following:
 - Confirmation that the PPE subject to the current application is physically identical to product xxx, which is covered by typeexamination certificate yyy.
 - Any difference between the original submission and this application to be listed.
 - Confirmation from the original manufacturer that only product fully compliant with type-examination certificate xxx will be supplied to own brand manufacturer yyy as the specified model.
 - Confirmation that the original manufacturer will advise the own brand manufacturer of any changes affecting the validity of either the type-examination certificate and for category III PPE, the supervision in line with module C2 or D.
 - Any proposed changes to the product will be sent to both the notified body and the own brand manufacturer before proceeding with the change.
 - Confirmation that the original technical documentation will be made available to the own brand manufacturer's notified body to support their application for certification and, for category III PPE, module C2 or D supervision.
 - Confirmation that both the original manufacturer and own brand manufacturer will inform each other of any incidents involving the products covered by the agreement
- 3. A copy of the EU type-examination certificate from the original manufacturer plus any documents that differ from the original technical documentation, e.g. marking and user information and access to the original technical documentation.

The notified body should review the user information and labelling of the own brand manufacturer in order to confirm it meets the requirements of the PPE Regulation.

A copy of the technical documentation amendments is then to be maintained by the own brand manufacturer and will be subject to review by the notified body during any ongoing surveillance activities.

- 4. For category III PPE, the notified body carrying out module C2 or D supervision will decide during the review of the own brand manufacturer's submission, activities etc., whether or not the premises of the own brand manufacturer need to be visited in the module C2 or D supervision.
- 5. The type-examination certificate issued to the own brand manufacturer will identify them as the manufacturer and will only reference the model identity as used by the own brand manufacturer. It is proposed that the original certificate does not need to be referenced on the certificate, but the NB holds this information, should it be required.

* * * * * * * * *	CO-ORDINA ⁻ PPE I RECOM	PPE-R/00.048 Version 1				
Number of pages: 1	RECOM		oroval stage :	Approved on :		
Origin : Horizontal Comm	nittee		-			
		\square	Vertical Group Horizontal Committee EU PPE Working Group	12/06/2017 23/01/2018		
Question related to	☑ PPE Regulation	EN/prEN:		Other:		
Article:	Annex:	Clause:				
Key words: sizing						
	Question: A manufacturer declares sizes or size ranges for a PPE he submits for EU type-examination. What action does the notified body have to take?					
Solution:						
If a manufacturer submit	s a PPE for certification, declaring ct. The test report shall state the te s or size ranges.					
	size ranges covered by the EU type	e-examination must not be	e CE marked.			

* PPE *	 CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425 				PPE-R/00.049 Version 1
× × ×	RECOMMENDA	ATION FO	R U	SE	
Number of pages: 1			Арр	proval stage :	Approved on :
Origin : Horizontal Committee			\boxtimes	Vertical Group Horizontal Committee EU PPE Working Group	12/06/2017 23/01/2018
Question related to PPE Regu	ulation	EN/prE	N:		Other:
Article: Anne VIII, 3	x: III, V 3. (c), VII, 3. (a), 3. (a)	Clause:			
Key words:					
traceability of technical documentation	n				
Question:					
What are the minimum criteria to guar EU type examination certificate?	rantee the traceability / identific	cation of doc	ume	nts within the technical doc	umentation approved for an
Solution: In order to assure the notified body th the information supplied by the manut correspond to the documents assesses back to the certificate holder at least a documents must be dated and identifi	facturer, which are part of the t ed during the EU type examina a copy of the marking of the PF	echnical doc tion, the noti	ume ified	ntation that must be presen body that carries out the E	nted by the manufacturer, U type examination will send

* PPE * * * *	CO-ORDINATION OF NOTIL PPE Regulation 201	PPE-R/00.050 Version 05		
^ * ^	RECOMMENDATION F	OR USE		
Number of pages: 1		Approval stage :	Approved on :	
Origin : Horizontal Comm	ittee	 □ Vertical Group ⊠ Horizontal Committee ⊠ EU PPE Expert Group 	22/11/2023 26/05/2024	
Question related to 🛛	PPE Regulation PPE Guidelines EN/	orEN:	Other:	
Article:	Annex: V Clause:			
Key words: Module C2 or D assessment, EU type-examination certificate Question: Should the notified body that carries out initial EU-type-examination for a category III product check that module C2 or D assessment is present or in process?				
Solution:				
No. In the regulation it is the manufacturer's responsibility to comply with all requirements before placing product on the market, and therefore they can decide upon the timing of applying for modules B and C2/D.				
Where a C2 / D agreeme	nt is not in place, manufacturers should have a place	marker for their labels and declar	ration, e.g. NB XXXX.	
The module B notified body approving the general presentation and content of conformity marking does not convey that a manufacturer has the right to use any notified body number shown.				
It shall be understood that the place marker 'NB XXXX' cannot be used to place product on the market, and that the manufacturer shall take action to contract with a Module C2 or D body, and following agreement use a valid notified body number in the place of 'XXXX' when production starts and before placement on the market, under a legally enforceable contract with that notified body.				

* * * * * PPE * * * *	CO-ORDINATION O PPE Regula	PPE-R/00.052 Version 1		
	RECOMMENDA	ATION FO		Annered
Number of pages: 1			Approval stage :	Approved on :
Origin : Horizontal Comr	nittee		 Vertical Group Horizontal Committee EU PPE Working Group 	12/06/2017 23/01/2018
Question related to	PPE Regulation	EN/prE	N:	Other:
Article:	Annex:	Clause:		
Key words:				
product marking; referer	ce to standards			
Question:				
Is it allowed to use a def	ined term of a standard (e.g. FFP3) for mar	king a produ	ct without any reference to the s	standard?
Solution:				
No.				

* * * *	CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425			PPE-R/00.053 Version 2	
× *	RECOMMEN	DATION FOR	RUSE		
Number of pages: 1			Approval stage :	Approved on :	
Origin : Horizontal Commi	itee		 Vertical Group Horizontal Committee EU PPE Working Group 	n/a 30/05/2018 22/04/2019	
Question related to] PPE Regulation	EN/prEN	N:	Other:	
Article:	Annex:	Clause:			
Key words: instructions for use					
Question: What can notified bodies do to ensure that the information supplied by the manufacturer is legible?					
Solution:					
	nation supplied by the manufacturer, no ay that it is legible for the user.	tified bodies sho	uld point out to the manufactur	er that the printed version	
They should make the ma	nufacturer aware of relevant document	s such as			
 IEC 82079-1 "Preparation of instructions for use – structuring, content and presentation – Part 1: General principles and detailed requirements", that specifies requirements for the presentation of instructions of use, e.g. font sizes; 					
ISO IEC Guide 3	37:2012 "Instructions for use of product	s by consumers'	n. ,		
• "Guideline on the readability of the labelling and package leaflet of medical products for human use" (version of 12/01/2009).					

* * * * * PPE * * *	CO-ORDINATION O PPE Regula	PPE-R/00.054 Version 1			
~ * * ~	RECOMMENDA	ATION FOI	R USE		
Number of pages: 1			Approval stage :	Approved on :	
Origin : Horizontal Comn	nittee		 Vertical Groups Horizontal Committee EU PPE Working Group 	12/06/2017 23/01/2018	
Question related to	☑ PPE Regulation	EN/prE	N:	Other:	
Article:	Annex: V, VII, VIII	Clause:			
Key words: Modules C2 and D, non- Question:	conform product, unsafe design				
What procedure should be followed during module C2 / D examinations in the event of a non-conforming product where the non-conformity is related to the design of that product?					
	oforming product where the non-conformity is modules C2 or D has to inform the notified y.				

* * * * * PPE * * *	CO-ORDINATION O PPE Regula	PPE-R/00.055 Version 1				
· · · ★ · · ·	RECOMMEND	ATION FOR	USE			
Number of pages: 1			Approval stage :	Approved on :		
Origin : Horizontal Comm	ittee		 ☐ Vertical Group ☐ Horizontal Committee ⊠ EU PPE Working Group 	12/06/2017 23/01/2018		
Question related to	PPE Regulation	EN/prEN	l:	Other:		
Article:	Annex: V	Clause:				
Key words: control systems						
Question: Which information shall be assessed during the EU type-examination if a control system is necessary to ensure a required protection of PPE?						
following:	nnical documentation shall include detailed					
	on of the safety function(s). These are function of the safety function (sound attenuation, light attenu		trol system which ensure a rec	quired protection (e.g.		
 the assessment and definition of the safety relevant parameters of the safety function (e.g. value of minimum airflow, value of min./max. O₂ concentration); 						
 the definition of the required performance level of the safety function. The definition shall cover the determination, verification and validation of the performance level (e.g. using the methods of EN ISO 13849-1 / IEC 62061, EN ISO 12100); 						
 the considerat 	ion of a possible loss or deviation of the ne	ecessary energ	gy supply;			
The manufacturer shall declare that he controls the adequacy and effectiveness of the control system. The notified body shall assess the technical documentation to verify the content accurately reflects the above requirements.						
like e.g. electro-optical f	echatronic components, so called control s illers for welding, electronic hearing protec If-contained re-breathing diving apparatus	tion devices (I				

RECOMMENDATION FOR USE Number of pages: 1 Approval stage : Approved on : Origin : Horizontal Committee Image: Committee Image: Committee Image: Committee Image: Committee Image: Committee Image: Committee Image: Committee Image: Committee <t< th=""><th></th><th colspan="3">CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425</th></t<>		CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425		
Origin : Horizontal Committee □ Vertical Group n/a ☑ Horizontal Committee 30/05/2018				
☐ Vertical Group n/a ☑ Horizontal Committee 30/05/2018	Number of pages: 1	Approval stage :	Approved on :	
	Origin : Horizontal Committee	 □ Vertical Group ☑ Horizontal Committee ☑ EU PPE Working Group 		
Question related to \square PPE Regulation \square EN/prEN: \square Other:	Question related to PPE Regulation	l/prEN:	Other:	
Article: Annex: II, 2.4 Clause:	Article: Annex: II, 2.4 Clause):		
Key words: date of manufacture / obsolescence on the product marking	-			
Question: Is it necessary to include in the label / marking of each product of PPE the date of manufacture or obsolescence?				
Solution: Not in all cases. It has to be marked if annex II, 2.4 paragraph I of the PPE Regulation applies or it is required by the relevant standard or specification.				

* PPE * * * *	CO-ORDINATI PPE R	PPE-R/00.058 Version 3				
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Number of pages: 1		Approval stage :	Approved on :			
Origin : French coordinat	ion	 □ Vertical Group ☑ Horizontal Committee ☑ EU PPE Working Group 	30/05/2018 p 22/04/2019			
Question related to	PPE Regulation	EN/prEN:	Other:			
Article:	Annex:	Clause:				
	ificate / re-certification / transitional	period				
Question: Can the simplified procedure (Annex V, 7.6) be a basis for the conversion of EC type-examination certificates (which comply with the PPE Directive) into EU type-examination certificates (in compliance with the PPE Regulation)?						
Solution: Yes, unless: - the manufacturer is not able to declare that no modification on the type of the product or on the technical file has occurred since the EC type-examination certificate has been issued or; - the generally acknowledged state of the art (standard, RfUs, etc.) has changed since the EC type-examination occurred, in such a way that the product may no longer comply with the applicable essential health and safety requirements of the PPE Regulation or; - data arising from Annex III, point 2 of the PPE Directive is not submitted by the manufacturer (notified bodies can ask for representative test data covering the life of the certificate). In this case they have to follow the full conformity assessment procedure (Article 19 PPE Regulation).						

* * * * PPE * * *	CO-ORDINATION C PPE Regula	PPE-R/00.061 Version 2				
× *						
Number of pages: 1		Approval stage :	Approved on :			
Origin : Horizontal Committee		☐ Vertical Group☑ Horizontal Committee☑ EU PPE Working Grout	30/05/2018 p 22/04/2019			
Question related to	☑ PPE Regulation	EN/prEN:	Other:			
Article:	Annex: Annex III, b), Annex V, 2	Clause:				
Key words: Risk assessment						
Question: What is required by the manufacturer and the notified body as far as risk assessment is concerned?						
Solution: The adequate analysis and assessment of the risk(s) is the responsibility of the manufacturer. The manufacturer describes the identified risks and shows the related sections of standards / specification plus the associated assessment method, e.g. analysis, inspection, test. The notified body reviews the documentation to ensure that - the risks are correctly identified with respect to the application made and the PPE presented - the submitted user information reflects the identified risks and includes associated limitations of use as far as the requirements of the PPE Regulation are concerned.						

* * * * * * * *	CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425			PPE-R/00.069 Version 04	
~	RECOMMENDA	ATION FO	RUSE		
Number of pages: 2			Approval stage :	Approved on :	
Origin: Horizontal Committee Advisory Panel			 □ Vertical Group ☑ Horizontal Committee ☑ EU PPE Expert Group 	22/11/2023 26/05/2024	
Question related to \square F	PPE Regulation PPE Guidelines	EN/prE	N:	Other:	
Article: Article 34(2)	Annex: V, VII, VIII	Clause:			
Key words:					
Issues relating to negative	e conformity assessment results / refused /	/ withdrawn /	suspended / restricted		
Question:					
How does a notified body negative conformity asses	fulfil the obligation to provide information t ssment results?	o other notifi	ed bodies conducting similar ad	ctivities on issues relating to	
Solution:					
	a forum for all notified bodies to participate membership available to provide information 4(2)).				
Therefore, information shall be sent by the deciding notified body declaring the negative conformity assessment results to all the relevant notified bodies carrying out similar conformity assessment activities via the 'Notified Body representative' CIRCABC sub-members of the applicable VG groups.					
The common approach adopted should be (the reason provided should be factual): "Dear Notified Bodies,					
Notified Body XXXX has refused / withdrawn / suspended / otherwise restricted certification / approval decisions for:					
Manufacturer: Address:					
Certificate(s) / Approval Decisions Affected / Types:					
Product Standard(s):					
Reason:					
- non-compliance in type testing (Annex V, Module B).					
- non-compliance in product checks (Annex VII, Module C2).					
- non-fulfilment of obligations arising out of the approved quality system (Annex VIII, Module D)					
- European Union safeguard procedure - failure of the PPE to meet requirements (Article 38(5)(a)) discovered in Market Surveillance					
- European Union safeguard procedure - shortcomings in harmonised standard (Article 38(5)(b)) discovered in Market Surveillance					
- other reasons to be specified.					
Effective from: DD Month	YYYY"				

ANNEX TO PPE-R/00.069 INSTRUCTIONS ON HOW TO USE CIRCABC

- 1. Go to the applicable Vertical Group main page;
- 2. Click on "Forum";
- 3. Click on "Add";
- 4. Click on "Topic";
- 5. Write the Topic title and click on "Create";
- 6. Click on "Add a new comment".

Note: When you make a new topic, do not fill in the box called 'Description'. This will not be sent to subscribers. Make the Topic (title only) and then close the dialog. Then click on the Topic title, and put your question into a new comment. This message will be sent by email to subscribers.

* * * * * PPE * * * *	CO-ORDINATION OF NOTIFIED BODIES PPE Regulation 2016/425				PPE-R/00.075 Version 01
100 🛪 100	RECO	MMENDATION	FOR	USE	
Number of pages: 1			Ар	proval stage:	Approved on:
Origin: NB 2849				Vertical Group	
_			X	Horizontal Committee EU PPE Expert Group	22/11/2023 26/05/2024
Question related to	PPE Regulation	L EN/p	orEN	:	U Other:
Article:	Annex: V: 6.1, 6.2(i), 7.	5, 7.6 Clause:			
Key words:					
Renewal date, renew	val, review, simplified review				
Question:					
	Body represent issue and ren				liance with Annex
	re the renewal process has c	omplied with Ai	nnex	(V 7.5 and 7.6?	
Solution:	· · · · · · · · · · · · · · · · · · ·	· · · · · · · · · · · · · · · · · · ·			
	nanufacturers who are early i ne continued use of the origi priod.			-	
Where a change has occurred, the notified body must consider if the originating certificate may retain validity, or must be suspended, reduced or withdrawn.					
A post-dated type-examination certificate may be issued following positive renewal, inclusive of a new period of validity of up to 5 years.					
The post-dated issue	d renewal type-examination	certification sho	buld	contain the following info	rmation:
- the date of first issue,					
- the date of expiration of the previous period of validity, starting the new period of validity,					
- the date on which the period of validity of renewal expires.					
Additionally, there could be an explanation of the type of renewal, for example:					
- "Reissued following simplified review procedure."					
- "Reissued following review procedure, inclusive of Amendments considering Change in the State of the					
Art."					
- "Reissued following review procedure, inclusive of Amendments to technical documentation."					
- "Reissued following review procedure, inclusive of Amendments to product."					
Note 1: While ISO 17065:2012, Cl 7.7.1 does require that the date certification is granted does not precede the date on which the decision is completed, it does not exclude specifying an effective granting date which succeeds the date on which the decision was completed.					
Note 2: ISO 17020:2012 does not exclude specifying an effective granting date which succeeds the date on which					

the decision was completed.