


Schedule of Accreditation

issued by

United Kingdom Accreditation Service

2 Pine Trees, Chertsey Lane, Staines-upon-Thames, TW18 3HR, UK

 <p>Accredited to ISO/IEC 17065:2012 to provide product conformity certification</p>	DNV UK Limited	
	Issue No: 005 Issue date: 05 January 2026	
	30 Stamford Street Vivo Building London SE1 9LQ	Contact: Mark Hayward Tel: +44 (0) 203 816 5682 E-Mail: Mark.hayward@dnv.com Website: www.dnv.com

Locations where certification activities covered by the above Accreditation Standard are undertaken	
Location A 30 Stamford Street Vivo Building London SE1 9LQ Head Office	Location B Building 3 International Business Park, Dyce Dr Dyce Aberdeen AB21 0BR Functional Office



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DNV UK Limited

Issue No: 005 Issue date: 05 January 2026

DETAIL OF ACCREDITATION

Accreditation for the purpose of UK Approved Body Activity in accordance with UKCA Requirements and UKAS Publication GEN 5			
Directive / Regulation	Conformity Assessment procedure/ Module/article	Category of products or individual products	Essential requirements: Product specification / Properties/Standards
Pressure Equipment (Safety) Regulations 2016, SI 2016 No 1105 as amended	<p>Conformity assessment procedures in accordance with Regulation 42 of the SI</p> <p>Schedule 1A Part 6 – Module D1 Quality assurance of the production process</p> <p>Schedule 1A Part 8 – Module E1 Quality assurance of final pressure equipment inspection and testing</p>	Category II Equipment	Schedule 2, Essential safety requirements and assessment of technical documentation, quality system and/or products in accordance with the relevant Module.
	<p>Schedule 1A Part 3 – Module B Design Type Examination</p> <p>Schedule 1A Part 4 – Module C2 Conformity to type based on internal production control plus supervised pressure equipment checks at random intervals</p> <p>Schedule 1A Part 5 – Module D Conformity to type based on quality assurance in the production process</p>	Category III Equipment	Schedule 2, Essential safety requirements and assessment of technical documentation, quality system and/or products in accordance with the relevant Module.



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Directive / Regulation	Conformity Assessment procedure/ Module/article	Category of products or individual products	Essential requirements: Product specification / Properties/Standards
Pressure Equipment (Safety) Regulations 2016, SI 2016 No 1105 as amended (cont'd)	Schedule 1A Part 7 – Module E Conformity to type based on pressure equipment quality assurance	Category III Equipment (cont'd)	Schedule 2, Essential safety requirements and assessment of technical documentation, quality system and/or products in accordance with the relevant Module.
	Schedule 1A Part 9 – Module F Conformity to type based on pressure equipment verification		
	Part 3 – Module B Production Type Examination	Category IV Equipment	Schedule 2, Essential safety requirements and assessment of technical documentation, quality system and/or products in accordance with the relevant Module.
	Part 5 – Module D Conformity to type based on quality assurance in the production process		
	Part 9 – Module F Conformity to type based on pressure equipment verification		
	Part 10 – Module G Conformity based on unit verification		
	Part 12 – Module H1 Conformity based on full quality assurance plus design examination		



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**Accreditation for the purpose of Notified Body Activity relating to the Northern Ireland market (CE + UKNI)
taking into account EA-2/17**

Directive / Regulation	Conformity Assessment procedure/ Module/article	Category of products or individual products	Essential requirements: Product specification / Properties/Standards
Pressure Equipment (Directive 2014/68/EU) as implemented in Northern Ireland by the Pressure Equipment (Safety) Regulation 2016, SI 2016 No 1105 as amended	Conformity assessment procedures in accordance with Article 14 of the Directive Annex III.6 Module D1 Quality assurance of the production process Annex III.1 Module E1 Quality assurance of final pressure equipment inspection and testing	Category II Equipment	Annex 1, Essential safety requirements and assessment of technical documentation, quality system and/or products in accordance with the relevant Module.
	Annex III.3 Module B Design Type Examination S Annex III.4 Module C2 Conformity to type based on internal production control plus supervised pressure equipment checks at random intervals Annex III.5 Module D Conformity to type based on quality assurance in the production process Annex III.7 Part 7 – Module E Conformity to type based on pressure equipment quality assurance	Category III Equipment	Annex 1, Essential safety requirements and assessment of technical documentation, quality system and/or products in accordance with the relevant Module.



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**Accreditation for the purpose of Notified Body Activity relating to the Northern Ireland market (CE + UKNI)
taking into account EA-2/17**

Directive / Regulation	Conformity Assessment procedure/ Module/article	Category of products or individual products	Essential requirements: Product specification / Properties/Standards
Pressure Equipment (Directive 2014/68/EU) as implemented in Northern Ireland by the Pressure Equipment (Safety) Regulation 2016, SI 2016 No 1105 as amended (cont'd)	Annex III.9 Module F Conformity to type based on pressure equipment verification	Category III Equipment (cont'd)	Annex 1, Essential safety requirements and assessment of technical documentation, quality system and/or products in accordance with the relevant Module.
	Annex III.3 Module B Production Type Examination Annex III.5 Module D Conformity to type based on quality assurance in the production process Annex III.9 Module F Conformity to type based on pressure equipment verification Annex III.10 Module G Conformity based on unit verification Annex III.12 Module H1 Conformity based on full quality assurance plus design examination	Category IV Equipment	Annex 1, Essential safety requirements and assessment of technical documentation, quality system and/or products in accordance with the relevant Module.



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DETAIL OF ACCREDITATION

Accreditation for the purpose of UK Approved Body Activity in accordance with UKCA Requirements and UKAS Publication GEN 5			
Directive/Regulation	Conformity Assessment procedure/Module/Article	Category of products or individual products	Methods / Procedures Essential requirements: Product specification/ Properties / Standards
Marine Equipment Regulations SI 2025 Location B	Assessment Procedure as defined in Schedule 2 of the Regulations: Part 1: Type examination (Module B) Part 2: Conformity to type based on quality assurance of the production process (Module D) Part 3: Conformity to type based on product quality assurance (Module E) Part 4: Conformity to type based on product verification (Module F) Part 5: Conformity to type based on unit v	For the following equipment categories as defined in MSN 1874, as amended: UK 1.1 to 1.43 UK 2.1 to 2.10 UK 3.1 to 3.71 UK 4.1 to 4.65 UK 5.1 to 5.22 UK 6.1 UK 7.1 UK 8.1	Assessment and Issue of Type Approval in accordance with paragraphs 11 and 12 of MSN 1874, as amended, and assessment of technical documentation, quality system and/or products in accordance with the relevant Module.
Note: The inspection of items marked thus * are subject to legislative requirements including the appointment of bodies' to carry out conformity assessment. Reference should be made to the relevant Government Departments 'BEIS' for MSMER for information on this and listings of bodies recognised under UK legislation			
END			