


Schedule of Accreditation

issued by

United Kingdom Accreditation Service

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

 6906 Accredited to ISO/IEC 17021-1: 2015 to provide management systems certification	LRQA Apave Limited	
	Issue No: 001 Issue date: 28 November 2023	
	1 Trinity Park Bickenhall Lane Solihull West Midlands B37 7ES	Contact: Simon Emeny Tel: +44(0)121 817 4418 E-Mail: simon.emeny@lrqa.com Website: www.lrqa.com

SUMMARY OF ACCREDITED SCOPE

Accredited to provide certification of the following Management Systems Standards and related Sector Schemes as detailed in this schedule:

- Accreditation for the purpose of UK Approved Body Activity in accordance with UKCA Requirements and UKAS Publication GEN 5
- Accreditation for the purpose of Notified Body Activity relating to the Northern Ireland market (CE + UKNI) taking into account EA-2/17



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LRQA Apave Limited

Issue No: 001 **Issue date:** 28 November 2023

KEY LOCATION ADDRESS	QMS
1 Trinity Park Bickenhall Lane Solihull West Midlands B37 7ES	✓

This Certification Body has demonstrated to UKAS that it has the systems and processes in place to provide the competence and capability, including understanding of local requirements, to manage and issue accredited management systems certification in the country in which it is established and in any other country, for the standards and scopes detailed on this schedule, unless specifically detailed in the individual Management System scope table (*denoted by **).



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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
Pressure Equipment (Safety) Regulations 2016, SI 2016 No 1105 as amended	Conformity assessment procedures in accordance with Regulation 42 of the SI Schedule 1A Part 11 – Module H Conformity based on full quality assurance	Supply chain to new build nuclear pressure equipment and assemblies. Category III Equipment	Schedule 2, assessment of technical documentation and quality system

Note: The inspection of items marked thus * are subject to legislative requirements including the appointment of 'persons or bodies' to carry out conformity assessment. Reference should be made to the relevant Government Department BEIS for PER for information on this and listings of bodies recognised under UK legislation.

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	Methods / Procedures 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.11 Module H Conformity based on full quality assurance	Supply chain to new build nuclear pressure equipment and assemblies. Category III Equipment	Annex 1, assessment of technical documentation and quality systems

Note: The inspection of items marked thus * are subject to legislative requirements including the appointment of 'persons or bodies' to carry out conformity assessment. Reference should be made to the relevant Government Department BEIS for PER for information on this and listings of bodies recognised under UK legislation.

END