

MENU



Calendar

EQA Catalogue Groups

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|  Genomic and Inherited Disorders |  Preimplantation Genetic Testing (PGT) |
|  Haematological Neoplasms |  Reproductive Genomics |
|  Individual Competency Assessment |  Sample Handling |
|  Molecular Newborn Screening |  Technical (Next Generation Sequencing) |
|  Molecular Pathology |  Variant Classification |
|  Multidisciplinary team (MDT) working |  Viral Sequencing |

Other Events

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|  GenQA EQA Enrolment Event |  GenQA workshop |
|  GenQA EQA Distribution Event |  GenQA Participant meeting |
|  GenQA webinar |  GenQA office closed |
|  Conference |  Other |

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■ 2026 Familial colorectal cancer and polyposis

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■ 2026 Familial endocrine tumour predisposition disorders

- CLOSES

■ 2026 Familial Hypercholesterolaemia (FH)

- CLOSES

■ 2026 Gastroenterology and hepatology disorders

- CLOSES

■ 2026 Hereditary Breast and Ovarian Cancer (HBOC) disorders

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					CNVs (pilot) - CLOSES  2026 Next Generation Sequencing (NGS) for germline SNVs and indels - CLOSES  2026 Next Generation Sequencing (NGS) for somatic SNVs and indels - tumour testing only - CLOSES  2026 Next Generation Sequencing (NGS) for somatic SNVs and indels - tumour with germline subtraction analysis - CLOSES	
Sun 15	Mon 16  2026 Distribution 23  2026 DNA extraction from saliva - OPENS [more]	Tue 17	Wed 18	Thu 19	Fri 20  2026 Bone Marrow Failure syndromes (pilot) - CLOSES  2026 Inherited cytopenias (pilot) - CLOSES  2026 DNA extraction from formalin-fixed paraffin-embedded (FFPE) tissue - CLOSES  2026 Reproductive carrier screening (pilot) - CLOSES	Sat 21
Sun 22	Mon 23	Tue 24	Wed 25	Thu 26	Fri 27  2026 cfDNA extraction from plasma - prenatal (pilot) - CLOSES  2026 cfDNA extraction from plasma - somatic (pilot)	Sat 28

					round 2 - CLOSES	
Sun 29	Mon 30	Tue 1	Wed 2	Thu 3	Fri 4 2026 Homologous recombination repair (HRD) testing - CLOSES 2026 Solid tumour FISH - CLOSES 2026 DNA extraction from saliva - CLOSES	Sat 5

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Sun	Mon	Tue	Wed	Thu	Fri	Sat
Sun 28	Mon 29	Tue 30	Wed 31  GENie ISCN and HGVS nomenclature modules CLOSE	Thu 1	Fri 2	Sat 3
Sun 4	Mon 5	Tue 6	Wed 7	Thu 8	Fri 9  2025 SARS-CoV-2 sequencing - Distribution 5938 - round 3 - CLOSES	Sat 10
Sun 11	Mon 12  Distribution 23 OPEN  2025 Optical Genome Mapping (OGM) - haematological neoplasms (pilot) - OPENS [more]  2025 Cystic Fibrosis (CF) Molecular Newborn screening (bloodspots) round 4 - OPENS [more]  2025 MCADD Molecular Newborn Screening (bloodspots) round 4 - OPENS [more]  2025 SCID Molecular Newborn Screening (bloodspots) round 4 - OPENS [more]  2025 Spinal Muscular Atrophy (SMA) Molecular Newborn screening (bloodspots) round 4 - OPENS [more]	Tue 13	Wed 14	Thu 15	Fri 16	Sat 17
Sun 18	Mon 19  Distribution 24 OPEN 	Tue 20	Wed 21	Thu 22	Fri 23  2025 Genomic molecular tumour board	Sat 24

	2025 Central Nervous System (CNS) tumours - OPENS [more] 2025 Cholangiocarcinoma - OPENS [more] 2025 Endometrial tumours - OPENS [more] 2025 Molecular Tissue Identification - OPENS [more] 2025 NTRK fusions - OPENS [more] 2025 Renal tumours - OPENS [more]				(MTB) for lung cancer (pilot) - CLOSES 2025 Genomic multidisciplinary team (MDT) working for rare disease (pilot) - CLOSES	
Sun 25	Mon 26	Tue 27	Wed 28	Thu 29	Fri 30	Sat 31