

Genomics Research Centre Diagnostic Testing

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NATA/RCPA accreditation: 14979

Address:

Genomics Research Centre
 Institute of Health and Biomedical Innovation
 Queensland University of Technology
 60 Musk Ave
 Kelvin Grove, Brisbane, QLD
 AUSTRALIA, 4059

Costs quoted include analysis and interpretation for specified disorders along with preparation and sending of report to referring clinician or pathology laboratory.

The GRC is NATA accredited to provide sequencing diagnostic services for the indications as outlined below. Sequencing services for other conditions employ identical methods but are not NATA accredited. Please contact the GRC directly to enquire about sequencing for other conditions.

Next Generation Sequencing – Whole Exome Sequencing

The GRC is NATA accredited to undertake Whole Exome Sequencing for identification of genetic mutation for the following disorders:

- Familial Hemiplegic Migraine type 1 (FHM1)
- Familial Hemiplegic Migraine type 2 (FHM2)
- Familial Hemiplegic Migraine type 3 (FHM3)
- CADASIL and small vessel disease
- Episodic Ataxia
- Spinocerebellar Ataxia
- Epilepsy
- Neuromuscular Disorders

Cost: \$1100 per sample (including GST)

Timeframe: 6-8 weeks from sample receipt

Next Generation Sequencing – Targeted Gene Panel

The Targeted Gene Panel uses NGS technology to simultaneously interrogate five genes that are implicated in neurogenetic disorders. This approach is cost-effective in cases where causative mutation may exist in a number of known genes which can be analysed concurrently reducing the need for consecutive (and costly) testing.

Cost: \$580 per sample (including GST)

Timeframe: 6-8 weeks from sample receipt

Gene	NATA accredited Indications*	Gene Coverage	
		Full Coverage	Partial Coverage
<i>CACNA1A</i>	FHM1 EA2 SCA6	Exons: 2-28, 30-41 UTR: 5'UTR	Exons: 1, 29, Introns: 11, 20, 25 UTR: 3'UTR
<i>ATP1A2</i>	FHM2	Exons: 1-7, 9-23 UTR: 5'UTR	Exons: 8 UTR: 3'UTR
<i>SCN1A</i>	FHM3 Epilepsy	Exons: 1, 3-5, 7-14, 17, 19-27 UTR: 3'UTR	Exons: 2, 6, 15, 16, 18 Introns: 21 UTR: 5'UTR
<i>NOTCH3</i>	CADASIL	Exons: 1-9, 11-29, 32, 33 UTR: 5'UTR,	Exons: 10, 30, 31, UTR: 3'UTR
<i>TRESK</i>	FHM	Exons: 1-3 UTR: 5'UTR, 3'UTR	

GRC	Forms Folder	Diagnostic Request Information Version 3.1
Amended: 08/04/19 Review By: 08/04/22	Printed: 13/5/19	Authorised: R. Smith

Targeted Mutation Testing

NATA accredited Sanger sequencing method is used for targeted mutation testing of single exons in the genes listed, or a known mutation point for any of our accredited disorders. This approach is cost-effective for a suspected specific genetic mutation.

Cost: \$220 per sample per exon (including GST)

MTHFR C677T Polymorphism - \$79.20 per sample (including GST)

Gene	NATA accredited Indications
<i>CACNA1A</i>	FHM1; EA2; SCA6
<i>ATP1A2</i>	FHM2
<i>SCN1A</i>	FHM3; Epilepsy
<i>NOTCH3</i>	CADASIL
<i>MTHFR</i>	MTHFR C677T Polymorphism
<i>Other Gene</i>	FHM, EA, SCA, Epilepsy, small vessel disease, other accredited disorder

Timeframe: 6-8 weeks from sample receipt

How to Order

1. Discuss testing with the patient
2. Complete a patient consent form for Specialised/DNA Testing and provide a copy to us with your request
3. Offer the patient genetic counselling – Genetic Counselling services can be located on the HGSA website <https://www.hgsa.org.au/asgc/asgc>
4. Complete the Request form – available on the GRC website: <https://research.qut.edu.au/grc/diagnostic-testing/diagnostic-testing-pricing/>
5. Collect and send sample
 - a. 5mls whole EDTA blood or 5mls of blood in lithium Heparin is required sent on ice for testing;
OR
DNA extracted from 10mls blood collected in EDTA or lithium heparin.
Please indicate:
 1. Storage conditions of the extracted DNA - TE or Water
 2. Method of DNA extraction
 3. 260/280 ratios (acceptable range 1.6-2.1)
 4. 260/230 ration (acceptable range 1.7-2.3)
 - b. Specimen labelling must include:
 - patient first name and surname
 - patient date of birth
 - date and time of collection
 - patient signature; and
 - collector's initials
 - c. Send samples packed on ice by courier the same day or overnight to:
Genomics Research Centre (GRC) Clinic
Institute of Health and Biomedical Innovation
Queensland University of Technology
5 School Street (loading bay)
Kelvin Grove, Brisbane, QLD
AUSTRALIA, 4059

Please note: Any specimens arriving at the GRC after seven (7) days from collection date will be rejected

Availability of Clinical Advice

The GRC may give advice on the nature of any mutations identified and their likely effect on protein production. Where information is available in the scientific literature, clinical advice regarding the nature of genetic variations can also be provided.

Protection of Personal Information

The GRC is committed to protecting the personal identification of the patients using the service. Thus, personal information, including the discussion of results will only take place with the patient themselves, clinicians authorised on the initial request, or those subsequently authorised in writing by the patient or authorised clinicians.