

Original Article

Clinical Implementation and Outcomes of Genetic Testing for Epilepsy by the Ontario Epilepsy Genetic Testing Program

Tugce B. Balci^{1,2}, Laila C. Schenkel^{3,4}, Cassandra Rastin³, Kevin Jones⁵, Lysa Boissé Lomax⁶, Jacob Turowec³, Jennifer Kerkhof³, Pratibha Bhai³, Olga Jarinova^{7,8}, Stacy Hewson⁹, A. Narayan Prasad¹¹, O. Carter Snead III¹², David A. Dyment¹⁰ and Bekim Sadikovic^{3,4}

¹Division of Genetics, Department of Paediatrics, Western University, London, ON, Canada, ²Medical Genetics Program of Southwestern Ontario, London Health Sciences Centre, London, ON, Canada, ³Molecular Diagnostics Program and the Verspeeten Clinical Genome Centre, London Health Sciences Centre, London, ON, Canada, ⁴Department of Pathology and Laboratory Medicine, Western University, London, ON, Canada, ⁵Division of Pediatric Neurology Department of Pediatrics McMaster University, Hamilton, ON, Canada, ⁶Division of Neurology Department of Medicine Queen's University, Kingston, ON, Canada, ⁷Department of Pathology and Laboratory Medicine, University of Ottawa, ON, Canada, ⁸Department of Genetics, CHEO, Ottawa, ON, Canada, ⁹Department of Genetic Counselling, The Hospital for Sick Children, Toronto; Department of Molecular Genetics, University of Toronto, Toronto, ON, Canada, ¹⁰Children's Hospital of Eastern Ontario Research Institute, University of Ottawa, ON, Canada, ¹¹Clinical Neurological Sciences, Department of Pediatrics, Western University London, ON, Canada and ¹²Department of Paediatrics, University of Toronto, Toronto, ON, Canada

ABSTRACT: Background: Epilepsy is a relatively common condition that affects approximately 4–5 per 1000 individuals in Ontario, Canada. While genetic testing is now prevalent in diagnostic and therapeutic care plans, optimal test selection and interpretation of results in a patient-specific context can be inconsistent and provider dependent. Methods: The first of its kind, the Ontario Epilepsy Genetic Testing Program (OEGTP) was launched in 2020 to develop clinical testing criteria, curate gene content, standardize the technical testing criteria through a centralized testing laboratory, assess diagnostic yield and clinical utility and increase genetics literacy among providers. Results: Here we present the results of the first two years of the program, demonstrating the overall 20.8% diagnostic yield including pathogenic sequence and copy number variation detected by next-generation sequencing panels. Routine follow-up testing of family members enabled the resolution of ambiguous findings. Post-test outcomes were collected as reported by the ordering clinicians, highlighting the clinical benefits of genetic testing. Conclusion: This programmatic approach to genetic testing in epilepsy by OEGTP, together with engagement of clinical and laboratory stakeholders, provided a unique opportunity to gather insight into province-wide implementation of a genetic testing program.

RÉSUMÉ : L'Ontario Epilepsy Genetic Testing Program : mise en œuvre clinique du programme et résultats des tests génétiques, relatifs à l'épilepsie *Contexte* : L'épilepsie est une maladie relativement fréquente, qui touche environ 4 – 5 personnes pour 1000, en Ontario, au Canada. Bien que les tests génétiques soient aujourd'hui d'usage courant dans les plans de soins de diagnostic et de traitement, la sélection optimale des tests et l'interprétation des résultats peuvent varier selon les patients et dépendre des fournisseurs de soins. *Méthode* : Premier en son genre en Ontario, le programme Ontario Epilepsy Genetic Testing Program (OEGTP) a été lancé en 2020 afin d'établir des critères de tests cliniques, d'organiser le contenu génétique, d'uniformiser les critères de tests techniques par l'intermédiaire d'un laboratoire central d'analyse, d'évaluer le rendement diagnostique et l'utilité clinique des tests et d'améliorer la compétence informationnelle en génétique chez les fournisseurs de soins. *Résultats* : D'après les résultats ici présentés, obtenus au cours des deux premières années du programme, le rendement diagnostique global est de 20,8 %; il comprend notamment les variations des séquences pathogènes et du nombre de copies détectées par les panels de séquençage de nouvelle génération. Par ailleurs, les tests de suivi courants, effectués parmi les membres des familles touchées, ont permis de résoudre des résultats équivoques. Enfin, les résultats post-tests recueillis comme ils avaient été déclarés par les médecins prescripteurs ont fait ressortir les avantages cliniques des tests génétiques. *Conclusion* : L'approche programmatique des tests génétiques relatifs à l'épilepsie, élaborée dans le cadre de l'OEGTP, en collaboration avec des intervenants cliniques et de laboratoire, a été l'occasion unique de rassembler de l'information sur la mise en œuvre d'un programme de tests génétiques à la grandeur de la province.

Keywords: diagnostics; epilepsy; next-generation sequencing

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Corresponding author: Bekim Sadikovic; Email: bekim.sadikovic@lhsc.on.ca

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Highlights

- **Unique program:** The OEGTP standardized testing criteria, panel gene curation and testing protocols.
- Diagnostic success: A 20.8% yield in 983 individuals was achieved with next-generation sequencing over 2 years.
- Clinical impact: Post-test outcomes emphasized clinical benefits of genetic testing in epilepsy, fostering provider engagement and genetics literacy province wide.

Introduction

Epilepsy is a chronic neurological disorder characterized by recurrent seizures, affecting approximately 75 000 adults and 15 000 children in the province of Ontario, Canada. ^{1,2} The cause of epilepsy is heterogeneous though genetics plays a significant role, especially in those with an early age of onset, a positive family history or associated co-morbidities. Accurate genetic diagnosis can direct treatment, better inform prognosis, limit further investigations and interventions and improve surveillance measures. ^{4,5} A genetic diagnosis will also inform and enable accurate genetic counseling, provide prenatal testing options and psychosocial benefits to the individual and their families. ⁶ Given these significant impacts on both the individuals, families and the healthcare system, accessing timely and accurate genetic services remains of importance in improving the diagnostic process in epilepsy.

Next-generation sequencing (NGS) technologies have transformed genetic testing in epilepsy clinics by enabling high-throughput, cost-effective and high-precision analysis of multiple gene panels with approximately 15%–20% diagnostic yield.^{7–10} In addition to diagnostic yield, a broader health system impact of NGS testing needs to be considered. While there have been considerable efforts toward a standardized curation of epilepsy-related genes, ¹¹ no clear information exists regardingthe optimal implementation of multigene panels in specific healthcare settings.

To better understand the regional landscape of multigene panel testing in epilepsy in a publicly funded provincial system, the Ontario Ministry of Health and Long-Term Care (MOHLTC) Laboratories and Genetics Branch established the Ontario Epilepsy Genetic Testing Program (OEGTP) in 2020.^{12,13} The program oversees multigene panel testing for epilepsy in the province, which is performed by a single reference laboratory funded by the MOHLTC (London Health Sciences Centre [LHSC] Molecular Genetics Laboratory) for epilepsy genetic testing in the province and is managed by a steering committee with wide representation from across the province that includes regulatory, clinical and laboratory stakeholders. This work summarizes the results from the initial two years (2020-2022) of the OEGTP in a cohort of over 1000 epilepsy patients tested through the province of Ontario. In addition to the diagnostic yield, we describe the various health system impact measures based on the systematic collection and review of pre- and post-test health and patient management information. Overall, the study underscores the importance of integrating genomics into epilepsy care, while highlighting the unique potential of the first of its kind OEGTP model to gather health system evidence for optimization of genomic diagnostic pathways in the care of patients with epilepsy.

Materials and methods

OEGTP - makeup and mandate of steering committee

The OEGTP steering committee comprises of two geneticists, one genetic counselor, two epileptologists (neurologists with

specialized epilepsy training), three molecular geneticists, two molecular technologists and one MOHLTC representative and is co-chaired by a diagnostic laboratory specialist and a clinical specialist involved in care of patients with epilepsy (i.e., epileptologist, neurologist, geneticist). The clinicians and laboratory scientists are from five separate academic health sciences centers in Ontario (in London, Ottawa, Toronto, Kingston, Hamilton), ensuring representation from almost all mid- to large-sized healthcare facilities with epilepsy treatment programs in the province. The committee met quarterly in the first year of implementation and has been meeting at least annually thereafter to perform its mandate.

The committee coordinates the ongoing curation of the multigene panels and active optimization of testing requirements based on updated evidence from literature, as well as feedback from the end-users. Its mandate also covers management of optimal access to testing by ensuring genetics literacy in ordering providers, promoting awareness of epilepsy genetics among healthcare professionals and delivering evidence-based guidelines for the streamlined use of NGS technologies in epilepsy.

OEGTP - design of epilepsy panels

The initial creation of evidence-based epilepsy gene panels for appropriately selected epilepsy patients was guided by the recommendations of a working group appointed by the Genetic Testing Advisory Committee, which had a mandate to review the clinical utility and validity of genetic tests and provide advice to the MOHLTC on the provision of genetic testing in Ontario. Using the principles outlined by the National Institutes of Health funded ClinGen resource (https://www.clinicalgenome.org), more specifically the ClinGen Epilepsy Gene Curation Expert Panel's guidance, the working group developed a "made in Ontario" tailored set of gene panels for epilepsy testing in a way that maximizes clinical validity (appropriateness) and actionability (benefit), by defining and determining parameters.

The first iteration of NGS panel options encompassed the following categories: (1) focal epilepsy (14 genes), (2) progressive myoclonic (20 genes), (3) early infantile epileptic encephalopathy (51 genes), (4) childhood-onset (45 genes), (5) brain malformation (44 genes), (6) actionable (22 genes) and (7) comprehensive panel (167 genes) (https://www.lhsc.on.ca/pathology-and-laboratory-medicine/ontario-epilepsy-genetic-testing-program). The panel content was reviewed and curated on a periodic basis by the steering committee. All genetic testing services were conducted at the provincial reference laboratory within the of Molecular Diagnostics Program, Verspeeten Clinical Genome Centre, Department of Pathology and Laboratory Medicine at LHSC in London, Ontario.

OEGTP – testing and ordering eligibility criteria in Ontario, Canada

Clinical eligibility criteria for epilepsy genetic testing in Ontario had been previously described by the Genetic Testing Advisory Committee. ¹³ The ordering providers included a board-certified medical geneticist, or epileptologist, or a neurologist with at least 6 months of training in epilepsy. Additional specialties were considered with a requirement for completing the "Project ECHO Ontario, Epilepsy Across the Life Span" educational course. This program was delivered by a team of adult and pediatric epileptologists, geneticists and genetic counselor and

laboratory specialists and included didactic sessions on the clinical criteria for genetic testing, pre- and post-test counseling and epilepsy gene panel selection. The program was based on the original hub-and-spoke model created for management of hepatitis C in New Mexico and incorporates a case-based learning program to improve engagement and access to care, called "Extension for Community Healthcare Outcomes" (ECHO). 14

Patient recruitment and testing process

Samples recruited from October 2020 to November 2022 from patients meeting the eligibility criteria for genetic testing¹³ were included in this study. The cohort included both pediatric and adult patients with epilepsy, from the neurology or genetics clinics of tertiary care/academic health centers, community hospitals and other neurology or primary care practices across Ontario, ordered by practitioners fulfilling the above criteria. The OEGTP NGS panel testing required adherence to specific clinical criteria to ensure that all referred patients exhibited clinical indications of suspected genetic forms of epilepsy. Physicians were asked to complete a questionnaire confirming that the age of onset, seizure type and electroclinical syndrome aligned with the criteria for genetic epilepsy diagnosis (Epilepsy Test Questionnaire in Supplementary Material 1). The requisition form included a selection of seven distinct NGS panels to be chosen based on clinical indication following recommendations from the OEGTP. Additionally, physicians were required to complete a post-test questionnaire (Management Impact form) to assess the clinical implications of the genetic testing results (Management Impact Form in Supplementary Material 1). The laboratory periodically recontacted the providers for pending forms to improve the response rate.

Specimens were received in the form of either extracted DNA or underwent DNA isolation from peripheral blood at the Molecular Genetics Laboratory (LHSC) using standard protocols using the MagNA Pure system (Roche Diagnostics, Laval, QC, Canada). Subsequently, DNA quantification was carried out by measuring absorbance with a DTX 880 Multimode Detector (Beckman Coulter, Brea, CA, USA).

NGS targeted panel testing

Gene sequencing was performed using a custom, in-house designed NGS panel protocol.¹⁵ Sequence capture probes were designed for KAPA HyperCap target enrichment encompassing 167 genes linked to epilepsy (Roche Sequencing Solutions, Inc., Santa Clara, CA, USA). These probes were designed to provide full coverage of all targeted coding regions, along with 20 bp of the 5' and 3' flanking intronic regions and untranslated regions (UTRs). Additionally, 100 bp surrounding the 24 Single Nucleotide Polymorphism (SNP) targets of the Agena iPLEX Pro Exome QC panel were included in the design. The detailed gene content of the epilepsy NGS panels is described in Supplemental Table 1 (in Supplementary Material 2). Library preparation was carried out using KAPA HyperChoice Chemistry (KAPA HyperCap Workflow v3, Roche Sequencing Solutions, Inc.), followed by sequencing on either NextSeq 550 or MiSeq instruments (Illumina, San Diego, CA, USA), in accordance with standard protocols. Sequencing was performed with a mean 200x coverage in order to

enable accurate sequence variant, copy number variant (CNV) and mosaicism detection.

NGS alignment, variant identification and sample fidelity

Sequence alignment and coverage distribution were performed using NextGene software version 2.4.2.3 (SoftGenetics, LLC., State College, PA, USA) with standard alignment settings, as previously described. Variants were filtered at a 10% allelic fraction threshold. Subsequently, BAM and VCF files were imported into Geneticist Assistant version 1.8.1 (SoftGenetics, LLC) for databasing and variant quality assessment, focusing on read depth, allelic fraction and read balance.

CNV calling was performed using an in-house developed methodology involving quantile normalization of the read depth as previously described, ^{16,17} with the following modifications: Read depth reports were generated with DNAcopy and exomeCopy R (v3.5.1) packages before normalization. The normalization results were binned by 10 bp and segmented to merge neighboring bins within an average normalized value of 0.1. Segments with ratio thresholds exceeding 1.35 (for duplications) or falling below 0.65 (for deletions) were flagged for manual review. Regions potentially affected by homologous sequence interference were subjected to more stringent ratio thresholds for review (greater than 1.15 for duplications or less than 0.85 for deletions).

The NGS targeted panels also encompass coverage for the 21 SNPs and 3 sex markers of the Agena iPLEX Pro Exome QC panel to evaluate sample fidelity. The Agena panel is processed on each stock DNA specimen according to the manufacturer's recommendations and in parallel with the KAPA HyperCap workflow. The results of both assessments are compared to ensure expected sex and SNP matching as part of the sample fidelity evaluation.

Targeted testing for familial variants

The familial variant testing was conducted using either Sanger sequencing for sequence variants or multiplex ligation-dependent probe amplification (MLPA) for CNVs in duplicate, following standard laboratory protocols. The proband's DNA was used as a positive control. In instances where a commercial MLPA kit was unavailable or more than three familial variants needed investigation simultaneously, targeted NGS was carried out as described above, with all non-familial variants filtered out.

Variant interpretation and diagnostic yield assessment

Variants were interpreted by a certified molecular geneticist in accordance with American College of Medical Genetics (ACMG) guidelines. Variants classified as pathogenic (P), likely pathogenic (LP) or variants of unknown significance (VUS) were included in the reports.

For the proband results, findings were categorized into four main groups: (i) molecular diagnosis, (ii) possible molecular diagnosis, (iii) uncertain or (iv) negative. A molecular diagnosis was established by the presence of one (for autosomal dominant [AD]) or two (for autosomal recessive [AR]) P/LP variants within a single gene or following the confirmation of a *de novo* VUS variant in a clinically related AD gene. A possible diagnosis was categorized by the presence of two VUS in an AR gene or a

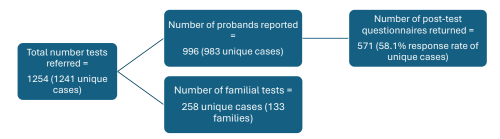


Figure 1. Schematic of samples in Ontario Epilepsy Genetic Testing Program (OEGTP) database from October 2020 to November 2022. In total, there were 1254 patients tested (1241 unique cases). 996 probands were tested by next-generation sequencing, and 258 familial samples were reported. 13 next-generation sequencing (NGS) cases were "add ons" meaning that after a sample was reported with the requested targeted subpanel as negative, an additional requisition was received requesting an expanded panel be reported. 571 post-test questionnaires were received following the reporting of results.

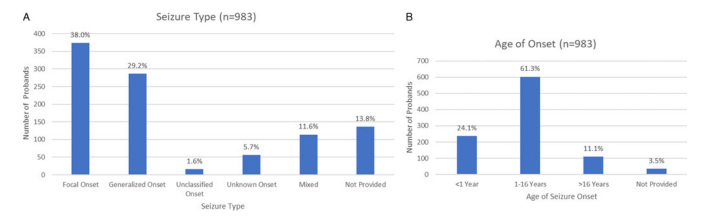


Figure 2. Clinical and demographic information of reported next-generation sequencing probands. (A) Type of seizure and (B) age of seizure onset as reported by the Ontario Epilepsy Genetic Testing Program pre-test questionnaire, available in Supplementary Material 1.

single P/LP variant in a clinically relevant AR gene or following the confirmation of a VUS to be *de novo* variant in an AR gene. Both sequence and CNVs are routinely and simultaneously reported. *Cis* versus *trans* status of multiple variants in the same gene was not available to report based on proband-only testing but was available if follow-up familial testing was performed. Similarly, presumed de novo status could be ascertained through parental testing using Sanger, MLPA or NGS, depending on the specific variant under investigation. Paternity was not confirmed during familial testing during the period of the reported cohort, but this capability was later added to the process based on the suggestions by the steering committee. An uncertain result encompassed any other VUS, while a negative result indicated the absence of reportable variants or a single reported VUS in an AR gene.

OEGTP database

The results from the OEGTP database for NGS testing, follow-up familial test results and pre- and post-test questionnaire responses were stored in an internally curated database at LHSC for further analysis and reference.

Statistical analysis

The standard chi-square test of independence was employed to evaluate the association between the reported diagnostic results in the post-test questionnaires and variables available from the data included in the test requisition. The complete list of features included in the pre-test questionnaires/requisitions can be found in Supplementary Material 1.

Results

During the period from October 2020 to November 2022, a total of 1254 tests were conducted. Among these, 996 probands, representing 983 unique cases, were referred for testing using the NGS panel (Figure 1). Additionally, 258 samples in 133 distinct families were referred for targeted testing of a known variant. Clinicians provided responses to 571 completed post-test questionnaires for NGS probands, a response rate of 58.1% for unique cases (excluding *add-on* testing).

Patient information and NGS panel results

Data from the requisitions revealed that the most referred seizure type was focal, followed by generalized seizures (Figure 2A). The predominant age of onset was between 1 and 16 years of age (n=606,61.3%, Figure 2B). The Comprehensive Epilepsy panel, encompassing 167 genes, was the most frequently requested panel (n=674), accounting for 67.7% of all NGS requests (Figure 3). Additionally, targeted subpanels such as focal epilepsy (n=36,3.6%) of tests), progressive myoclonic (n=13,1.3%), early infantile (n=57,5.7%), childhood-onset (n=76,7.6%), brain malformation (n-42,4.2%) and actionable (n=15,1.5%) were

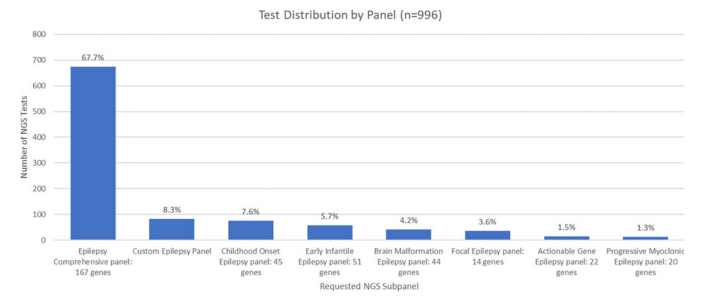


Figure 3. Next-generation sequencing (NGS) test distribution by subpanel of all NGS samples. 996 NGS tests were reported (983 unique cases and 13 samples that received an additional, expanded "add-on" report following a negative result from a smaller subpanel at the request of a clinician). The majority (67.7%) of requests are for the epilepsy comprehensive panel.

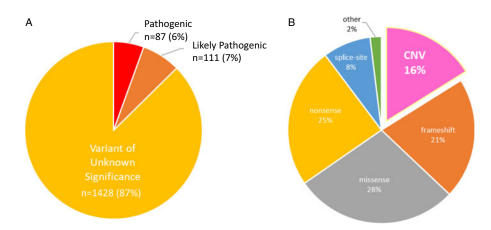


Figure 4. Variants detected by next-generation sequencing (NGS) panel. (A) 1626 total variants were reported by NGS testing (n = 996 reports), with the majority of them being classified as variants of unknown significance (VUS). Note that "add-on" cases where the same variant was included in the original and updated request were only counted once. (B) A breakdown of the types of pathogenic/likely pathogenic variants reported (VUS excluded). A roughly equal distribution of missense, frameshift and nonsense variants was detected, and 16% of the reportable pathogenic/likely pathogenic variants were copy number variants (CNVs). Other variants included inframe deletions and duplications. Benign or likely benign variants are not reported but are available upon request.

utilized, along with custom requests involving any combination of panels or genes (n = 83, 8.3%).

In this cohort, a total of 1626 variants were identified as P/LP/VUS, including 212 P/LP variants reported (13% of all variants), and a comparable frequency of missense, frameshift and nonsense variants was observed (Figure 4B). Notably, 16% of the reportable P/LP variants were CNVs, encompassing subexonic, exon-level or gene-level deletion/duplications (Table 1). A list of CNVs detected is provided in Table 1, and a visualization of some examples is shown Figure 5. VUS were reported in 1414 of 1626 of all variants (87%; Figure 4A).

Diagnostic yield

Overall, 75% (743/983) of probands had at least one reportable variant (VUS/LP/P), and 19.2% (189/983) had at least one P/LP variant reported. The diagnostic rate reported by the laboratory, comprising both molecular and possible molecular diagnoses, for all unique samples (excluding add-on requests) was 20.8% (204/983) (Figure 6A). The early infantile epileptic encephalopathy panel yielded the highest diagnostic rate at 26.7% (15/57), followed by custom requests at 23.7% (19/83) (Figure 6B). However, the comprehensive panel resulted in the highest absolute number of

Table 1. Copy number variants (CNVs) detected using in-house developed algorithm from NGS panels

#	Gene	CNV	Interpretation	Size	Inheritance
1	ALDH7A1	c.(650 + 21_651-21)_(695 + 21_696-21)del	LP	single-exon	AR
2	CACNA1A	c.(293 + 21_294-21)_(539 + 21_540-21)del	VUS	multi-exon	AD
3	CLN3	c.(460 + 21_461-21)_(677 + 21_678-21)del	Р	multi-exon	AR
4	CNTNAP2	c.(97 + 21_98-21)_(402 + 21_403-21)del	Р	multi-exon	AR
5	CTSD	c.(?21)_(*21_?)dup	VUS	whole gene	AR
6	DEPDC5	c.(?21)_(2104 + 21_2105-21)del	LP	multi-exon	AD/AR
7	DEPDC5	c.(2801 + 21_2802-21)_(3021 + 21_3022-21) del	LP	single-exon	AD/AR
8	DEPDC5	c.(?21)_(871 + 21_872-21)del	LP	multi-exon	AD/AR
9	DEPDC5	c.(146 + 21_147-21)_(193 + 21_194-21)del	LP	single-exon	AD/AR
10	GABRA1	c.(?21)_(74 + 21_75-21)del	LP	single-exon	AD
11	GPR56	c.(?21)_(*21_?)del	LP	whole gene	AR
12	GRIN2A	c.(?21)_(414 + 21_415-21)dup	VUS	single-exon	AD
13	KATNB1	c.(?21)_(*21_?)del	LP	whole gene	AR
14	KCNQ2	c.(1148 + 21_1149-21)_(*21_?)del	Р	multi-exon	AD
15	KCNQ2	c.(1763 + 21_1764-21)_(1887 + 21_1888-21) del	LP	single-exon	AD
16	LARGE	${\sf c.(408+21_409-21)_(615+21_616-21)} {\sf del}$	VUS	multi-exon	AR
17	MDH2	c.(?21)_(555 + 21_556-21)dup	VUS	multi-exon	AR
18	MEF2C	c.(?21)_(54 + 21_55-21)del	LP	single-exon	AD
19	NDE1	c.(?21)_(*21_?)del	Р	whole gene	AR – Possible microdeletion syndrome 16p13.11
20	NDE1	c.(?21)_(*21_?)del	Р	whole gene	AR – Possible microdeletion syndrome 16p13.11
21	NDE1	c.(?21)_(*21_?)del	Р	whole gene	AR – Possible microdeletion syndrome 16p13.11
22	NDE1	c.(?21)_(*21_?)del	Р	whole gene	AR – Possible microdeletion syndrome 16p13.11
23	NDE1	c.(?21)_(*21_?)del	Р	whole gene	AR – Possible microdeletion syndrome 16p13.11
24	NDE1	c.(?21)_(*21_?)dup	VUS	whole gene	AR – Possible microdeletion syndrome 16p13.11
25	NPRL3	c.(118 + 21_119-21)_(393 + 21_394-21)del	LP	multi-exon	AD
26	NRXN1	c.(3190 + 21_3191-21)_(3484 + 21_3485-21) del	VUS	multi-exon	AD
27	NRXN1	c.(?21)_(931 + 21_932-21)del	Р	multi-exon	AD
28	PCDH19	c.101_297del, p.(Arg34Hisfs*126)	LP	sub-exon	XL
29	POMK	c.(?21)_(*21_?)dup	VUS	whole gene	AR
30	PRRT2	c.(?21)_(*21_?)del	Р	whole gene	AD – Possible microdeletion syndrome 16p11.2
31	PRRT2	c.(?21)_(*21_?)del	Р	whole gene	AD – Possible microdeletion syndrome 16p11.2
32	PRRT2	c.(?21)_(*21_?)del	Р	whole gene	AD – Possible microdeletion syndrome 16p11.2
33	PRRT2	c.(?21)_(*21_?)del	Р	whole gene	AD – Possible microdeletion syndrome 16p11.2
34	PRRT2	c.(?21)_(*21_?)del	Р	whole gene	AD – Possible microdeletion syndrome 16p11.2

(Continued)

Table 1. Copy number variants (CNVs) detected using in-house developed algorithm from NGS panels (Continued)

#	Gene	CNV	Interpretation	Size	Inheritance
35	PSAT1	c.(740 + 21_741-21)_(*21_?)del	LP	multi-exon	AR
36	RTTN	c.(5541 + 21_5542-21)_(*21_?)del	LP	multi-exon	AR
37	RTTN	c.(5541 + 21_5542-21)_(*21_?)del	LP	multi-exon	AR
38	SGCE	c.(?21)_(*21_?)del	Р	whole gene	AD
39	SNAP29	c.(?21)_(*21_?)dup	VUS	whole gene	AR – Possible microduplication syndrome 22q11.2
40	SNAP29	c.(?21)_(*21_?)del	LP	whole gene	AR – Possible microdeletion syndrome 22q11.2
	TCF4	$c.(195+21_196\text{-}21)_(286+21_287\text{-}21) del$	LP	single-exon	AD
41	TSC1	c.(?21)_(106 + 21_107-21)del	VUS	single-exon	AD
42	TUBB	c.1061_*779dup	VUS	sub-exon	AD
43	TUBB	c.1061_*779dup	VUS	sub-exon	AD
44	TUBB3	c.(166 + 21_167-21)_(*21_?)dup	VUS	multi-exon	AD
45	WWOX	c.(516 + 21_517-21)_(1056 + 21_1057-21)del	LP	multi-exon	AR
46 \$	Chr15	Chr15:(?_25584224)_(27018129_?)dup	LP	multigene	AD – Possible microduplication syndrome 15q11.2-q13

AD = autosomal dominant; AR = autosomal recessive; LP = likely pathogenic; NGS = next-generation sequencing; P = pathogenic; VUS = variant of uncertain significance. Individual #40 had two separate CNVs, involving SNAP29 and TCF4 genes. S: Confirmed by microarray as having 15q11.2q13 microduplication syndrome.

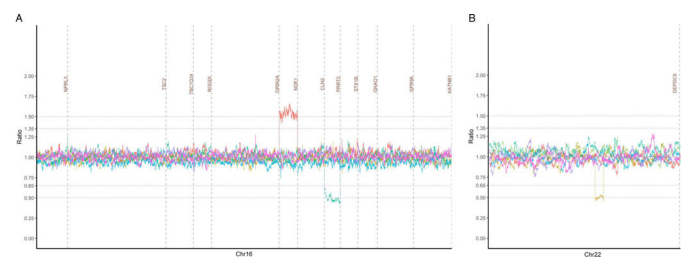


Figure 5. Copy number variants (CNVs) detected by next-generation sequencing (NGS). Each colored line represents a different patient. Samples normalized to a ratio of 1 represent normal diploid copy number, whereas regions normalized to 1.5/0.5 are representative of a heterozygous duplication/deletion, respectively. (A) Shows a section of chromosome 16. The patient represented by the red line has a full gene duplication of NDE1. The patient represented by the green line has a full gene deletion of PRRT2. (B) Shows a section of chromosome 22. The targeted NGS panel can also detect subgene or sub-exon-level CNVs. The patient represented by the yellow line has a deletion of DEPDC5 exon 30.

diagnostic results (148/674; 22.1%) (Figure 6B). Correspondingly, the comprehensive panel also presented the highest (65.6%) proportion of uncertain results.

Subsequent familial testing identified 26 assumed *de novo* variants (but without confirmation of paternity and maternity) (Table 2), with five of these variants identified in *SCN1A*, three in *KCNQ2* and two in *NRXN1*, while the remaining variants were distributed across unique genes. Seven (out of 9) LP variants were confirmed to be P after parental testing. Five (out of 9) VUS were reclassified as LP based on the *de novo* criteria (PM6) and additional clinical information for gene-disease correlation

(PP4).¹⁸ Clinical information was missing in some cases with presumed *de novo* VUS. As a result, some of these variants could still be reclassified as LP if the patient's clinical features match the genetic diagnosis. For a complete summary of familial testing outcomes and variant segregation, refer to Supplemental Table 2 (in Supplementary Material 2).

Post-test diagnostic outcome assessment

The post-test questionnaire included two questions. The first inquired about clinicians' assessment of the reported result,

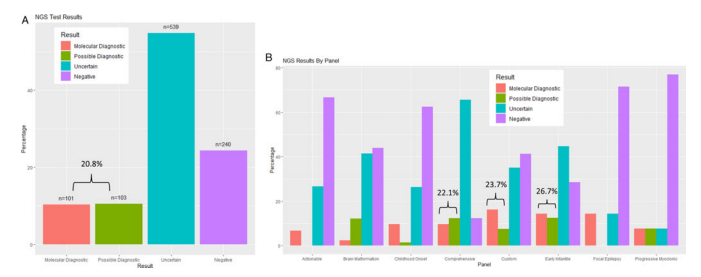


Figure 6. Next-generation sequencing (NGS) epilepsy panel diagnostic rates (n = 983 unique proband reports). A molecular diagnosis was defined as the presence of 1 or 2 pathogenic or likely pathogenic (P/LP) variants in a single gene, depending on the mode of inheritance of the associated disorder, or a confirmed de novo variants of unknown significance (VUS) in an autosomal dominant gene. A possible diagnosis was defined as two VUS in a clinically related autosomal recessive (AR) gene or 1LP/1P variant in an AR gene. An uncertain result is any VUS that does not fit into prior categories. Negative results identified no reportable variants (benign or likely benign variants are not reported but are available upon request) by NGS. (A) NGS test results for the collective database found a combined molecular/possible diagnostic rate of 20.8%. (B) NGS test results using the same classification system broken down by subpanel.

categorizing them as either (i) diagnostic, (ii) non-diagnostic or (iii) uncertain (involving a VUS). The average clinician-reported diagnostic rate in the returned questionnaires was 16% (92/571), the Early Infantile panel exhibited the highest percentage of clinician-reported diagnostic rate at 28% (10/36) (Figure 7).

The second question in the post-test questionnaire asked clinicians to indicate if the testing resulted in any change in patient management and, if so, to select the specific change in management (see questionnaire in Supplementary Material 1). From this, 47% (267/571) of patients were reported to have some form of management benefit post-testing as defined by the ordering provider (Figure 8). Of those cases with reported change in management, the majority of them (70%) received a non-diagnostic or VUS genetic testing result. Excluding cases where the only management change applied to referrals of family members for testing, 24.3% (139/571) of patients underwent a direct change in management: most frequently referrals to other specialists and additional imaging/testing. Medication changes or ketogenic diet institution were reported in 22 (4%) cases, from which the majority 15 (68%) received a diagnostic molecular test result.

A significant association was observed between the age of onset and diagnostic findings (chi 2 (degrees of freedom = 2, N = 557) = 21.43, p = 2.22E-5), indicating a higher diagnostic rate among individuals with an onset of < 1 year of age (Supplemental Table 3 in Supplementary Material 2).

Discussion

The OEGTP model encompasses many unique elements, such as incorporating input from multiple experts from provincial centers across specialties within an oversight committee, who then provides a regular review process to improve all aspects of implementation by periodic curation of both panel gene content

and the testing process, from tailoring the requisition form to implementing variant resolution strategies. The collection of posttest clinician-reported outcomes that highlight clinical benefits, the inclusion of a community-centered training program to educate providers and improve access and the implementation of an NGS panel with CNV detection capabilities were other unique features of the OEGTP that allowed the program to set a provincial standard for integration of genetic testing in epilepsy care. A programmatic, centralized approach can streamline future research initiatives aimed at enhancing diagnostic rates, such as expanded genomic and epigenomic analyses.

Establishment of the OEGTP and implementation of routine NGS panel testing resulted in a diagnostic yield with a direct management impact in a significant proportion of patients with epilepsy. In this comprehensive analysis of the genetic basis of epilepsy in a large cohort of 983 individuals with diverse phenotypes, we demonstrate the diagnostic efficacy along with clinical impact. We identified a molecular diagnostic/possible diagnostic in 20.8% of probands, a rate consistent with previous studies utilizing targeted NGS panels. 19-21

The most ordered clinical testing for epilepsy includes chromosomal microarray (CMA) and NGS testing, with a significant proportion of patients (~20%) having both tests ordered. While there is evidence that NGS (large panels or exome sequencing) has a higher diagnostic yield than CMA in pediatric epilepsy, many medical providers add CMA to the diagnostic process, especially if the patient also has a developmental disability, based on the 2010 American College of Medical Genetics guidelines. A more recent publication by ACMG recommends the use of whole exome sequencing (WES) or whole genome sequencing (WGS) as a first or second tier (after CMA) test for patients with developmental disabilities (including epilepsies). In addition, cost-effectiveness analysis supports a step-wise approach with WES or NGS-panel as first tier and, if negative, CMA for patients with epilepsy of unknown etiology. A NGS-based test

Table 2. Variants reported as assumed *de novo* based on familial testing, resulting in upgrade in classification and diagnostic confirmation in 5/9 VUS. In all cases (except #24), samples from both biological parents were tested; in some cases, extended family members were available

			Interpr	etation		
#	Gene	Variant (all heterozygous)	Before familial testing	After familial testing	Final ACMG code ¹⁸	Mode of inheritance (OMIM)
1	CDKL5	c.545T>C, p.(Leu182Pro)	VUS	LP*	PM2, PM6, PP3, PP4	XLD
2	DCX	c.506C>G, p.(Thr169Arg)	VUS	LP*	PM2, PM6, PP3, PP4	XL
3	DNM1	c.709C>T, p.(Arg237Trp)	Р	Р	PS4, PM2, PP3, PM6, PP5	AD and AR
4	DYNC1H1	c.925C>T, p.(Arg309Cys)	VUS	VUS	PM2, PM6, PP3	AD
5	DYRK1A	c.312C>A, p.(Tyr104*)	Р	Р	PVS1, PM2, PM6	AD
6	FGF12	c.341G>A, p.(Arg114His)	Р	Р	PS4, PS3, PM2, PP3, PM6, PP5	AD
7	FLNA	c.2693delA, p.(Asn898Metfs*48)	LP	P*	PVS1, PM2, PM6	XL
8	GABRB3	c.868A>C, p.(Asn290His)	VUS	VUS	PM2, PM6, PP3	AD
9	HCN1	c.2558_2562dup, p. Gly855Thrfs*23)	VUS	VUS	PVS1_not applied, PM6, PM2	AD
10	KCNQ2	c.830C>T, p.(Thr277Ile)	LP	LP	PM1, PM2, PM5, PM6, PP3	AD
11		c.901G>A, p.(Gly301Ser)	LP	LP	PM2, PS2, PP3	
12		c.941C>T, p.(Ser314Phe)	VUS	LP*	PM2, PM6, PP3, PP4	
13	NRXN1	c.(3190 + 21_3191-21)_(3484 + 21_3485- 21)del	VUS	LP*	PM2, PM6, PVS1_mod	AD
14		c.2824_2827dup. p.(Ile943Asnfs*44)	LP	P*	PVS1, PM2, PM6	
15	SCN1A	c.2849G>T, p.(Gly950Val)	VUS	LP*	PM2, PM6, PP3, PP4	AD
16		c.2921delT, p.(Met974Argfs*4)	LP	P*	PVS1, PM2, PM6	
17		c.580G>A, p.(Asp194Asn)	Р	Р	PS2, PS4, PM2, PM6, PP3	
18		c.664C>T, p.(Arg222*)	Р	Р	PVS1, PS2, PS4, PM2	
19		c.695-2A>G	LP	P*	PVS1, PM2, PM6	
20	SCN2A	c.4446 + 2T>C	LP	P*	PVS1, PM2, PM6	AD
21	SCN8A	c.4850G>A, p.(Arg1617Gln)	Р	Р	PS2, PS3, PS4, PM2, PM6, PP3	AD
22	SLC35A2	c.752G>A, p.(Trp251*)	LP	P*	PVS1, PM2, PM6	XLD
23	SYNGAP1	c.380_383dupGGCC, p.(Ser129Alafs*24)	LP	P*	PVS1, PM2, PM6	AD
24 ^{\$}	TCF4	c.1543G>C, p.(Gly515Arg)	VUS	VUS	PM2, PM6, BP4	AD
25	TSC1	c.733C>T, p.(Arg245*)	Р	Р	PVS1, PS4, PM2, PM6	AD
26	TSC2	c.1832G>A, p.(Arg611Gln)	Р	Р	PS3, PS4, PM2, PM6, PP3	AD

AD = autosomal dominant; AR = autosomal recessive; LP = likely pathogenic; P = pathogenic; VUS = variant of uncertain significance; XLD = X-linked dominant; XL = X-linked recessive. *:cases in "after familial testing" resulted in variant reclassification. \$: Only one parent tested.

capable of detecting gene-level CNVs, such as the OEGTP panel, further enhances the diagnostic yield of NGS by combining the two approaches, making it the optimal first-tier testing option for this population.

Similarly to our previous findings in hereditary cancer, ^{16,26} mitochondrial disorders²⁷ and neuromuscular disorders, ²⁸ we demonstrate that a custom NGS assay that includes CNVs detection significantly increased the diagnostic yield in patients with epilepsy, with 16% of all reported pathogenic variants being large sub-exon, exon and multigene deletions and duplications (Table 1). Some of these CNVs involved the deletion or duplication of a gene within a known microdeletion or microduplication genomic region, such as a *PRRT2* gene deletion within 16p11.2, while the majority were novel, highlighting the

advantage of a comprehensive NGS-based CNV analysis. For cases involving larger CNVs, microarray testing was recommended to confirm the extent of the deletion or duplication, contributing to the complete and accurate identification of the underlying genetic diagnosis.

Limitations of this study include its retrospective design and potential sampling or response bias, given the 58% response rate to the management impact forms. However, the use of standardized forms for both pre- and post-test processes facilitated the uniform collection of data across subjects and centers, providing a reasonable snapshot of the provincial landscape.

Personalized medicine is rapidly evolving in the field of epilepsy. A genetic diagnosis narrows the scope of medications used in certain epilepsies, as exemplified by a diagnostic result with

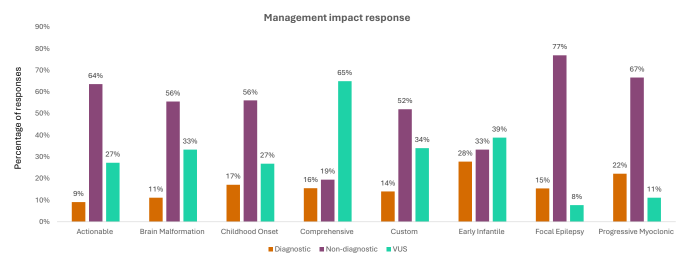


Figure 7. Ontario Epilepsy Genetic Testing Program (OEGTP) management impact form diagnostic responses by panel (n = 571). Following testing, clinicians were asked to respond to the reported test result in a post-test questionnaire and indicate if the result was in their opinion: diagnostic, non-diagnostic or if a variant of unclear clinical significance was identified. The average reported diagnostic rate across all panels was 17.2%. The Early Infantile panel reported the highest percentage of diagnostic findings (27.8%).

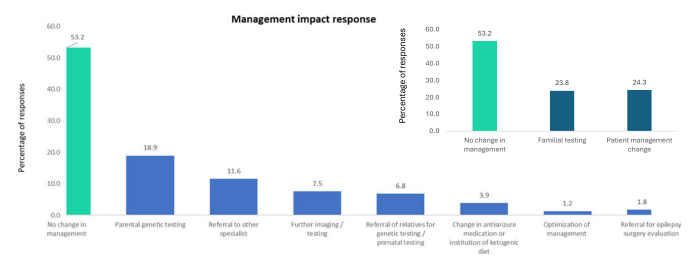


Figure 8. Ontario Epilepsy Genetic Testing Program (OEGTP) management impact form change in management responses by panel (n = 571). Following testing, clinicians indicated on the post-testing questionnaire if the OEGTP test results led to a change in management of their patient. If applicable, it is possible to select multiple responses. Collectively, 24.3% of patients were reported to have some change in management following testing, with $\sim 4\%$ reporting a direct change in patient treatment. Familial testing for parents, cascade diagnostic or prenatal testing was observed in 23.8% of cases.

a LP variant in *SLC2A1*, resulting in a change in treatment strategy. Disease causing variants in *SLC2A1* are associated with glucose transporter 1 (GLUT1) deficiency disorder, which is targeted successfully with institution of ketogenic diet.²⁹ There were further examples of diagnostic results involving SCN1A-related disorders, where sodium channel blocking agents are avoided to prevent seizure exacerbation³⁰ and agents such as stiripentol are considered first-line treatment options following diagnosis,³¹ as well as in KCNQ2-related encephalopathy, where sodium channel blockers are preferred.^{28,29}

As for future directions, the OEGTP will continue to keep up with the advances in epilepsy genetic testing. Recent studies have demonstrated an increased diagnostic rate with WES among epilepsy patients, offering the additional advantage of identifying novel genetic causes.^{8,34} With the expanding list of genes associated with epilepsy

and the rapid pace of novel gene discovery, WES or other genome-wide methods may complement current diagnostic approaches for syndromic epilepsy. Ontario Health's Provincial Genetics Program has recently begun issuing "Genetic Testing Recommendation Documents" for various indications, including neuromuscular diseases and neurodevelopmental disorders, to delineate the appropriate stages for implementing genome-wide sequencing in the diagnostic process. The program also recently endorsed the use of genome-wide sequencing (WES or WGS) as a first-tier diagnostic option in the diagnosis of rare genetic conditions with updated criteria and eligibility (https://www.ontariohealth.ca/providing-health-care/clinical-guidelines-standards/genetics-guidelines). Finally, emerging research demonstrates diagnostic utility of DNA methylation analysis in patients with genetically unsolved pediatric epilepsies, even after extensive genomic profiling including WES and WGS.³⁵ The OEGTP

now has the infrastructure and experience to utilize a programmatic approach to adapt to rapidly changing landscapes and implement the most clinically relevant technology for epilepsy patients in Ontario.

The integrated, dynamic and responsive approach of OEGTP to epilepsy genetic testing, coupled with continued genetic education, could empower healthcare providers to consider genetic testing early in their diagnostic algorithm, confidently order the optimal genetic test with relevant follow-up, ultimately interpret the results and counsel patients accurately.

Conclusion

The OEGTP was established through collaboration among clinical, laboratory and regulatory stakeholders. This partnership created a comprehensive and effective solution that was able to integrate seamlessly into the provincial healthcare system. By enhancing diagnostic capabilities and patient care for syndromic epilepsies, the program ensures a coordinated and impactful approach to delivering high-quality, system-responsive healthcare.

Supplementary material. The supplementary material for this article can be found at https://doi.org/10.1017/cjn.2025.69

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