Genetic and metabolic investigations for neurodevelopmental disorders: position statement of the Canadian College of Medical Geneticists (CCMG)

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ABSTRACT

Purpose and scope The aim of this position statement is to provide recommendations for clinicians regarding the use of genetic and metabolic investigations for patients with neurodevelopmental disorders (NDDs), specifically, patients with global developmental delay (GDD), intellectual disability (ID) and/or autism spectrum disorder (ASD). This document also provides guidance for primary care and non-genetics specialists caring for these patients while awaiting consultation with a clinical geneticist or metabolic specialist.

Methods of statement development A multidisciplinary group reviewed existing literature and guidelines on the use of genetic and metabolic investigations for the diagnosis of NDDs and synthesised the evidence to make recommendations relevant to the Canadian context. The statement was circulated for comment to the Canadian College of Medical Geneticists (CCMG) membership-at-large and to the Canadian Pediatric Society (Mental Health and Developmental Disabilities Committee); following incorporation of feedback, it was approved by the CCMG Board of Directors on 1 September 2022.

Results and conclusions Chromosomal microarray is recommended as a first-tier test for patients with GDD, ID or ASD. Fragile X testing should also be done as a first-tier test when there are suggestive clinical features or family history. Metabolic investigations should be done if there are clinical features suggestive of an inherited metabolic disease, while the patient awaits consultation with a metabolic physician. Exome sequencing or a comprehensive gene panel is recommended as a second-tier test for patients with GDD or ID. Genetic testing is not recommended for patients with NDDs in the absence of GDD, ID or ASD, unless accompanied by clinical features suggestive of a syndromic aetiology or inherited metabolic disease.

INTRODUCTION

Neurodevelopmental disorders (NDDs) are a group of conditions first manifesting early in childhood, characterised by impairments of personal, social, academic or occupational functioning. Examples of NDDs include autism spectrum disorder (ASD), global developmental delay (GDD), intellectual disability (ID), learning disability (LD) and attention-deficit hyperactivity disorder (ADHD). NDDs have a combined prevalence of ~17% in children aged 3–17 years, making them the most common chronic medical conditions encountered in paediatric primary care. Given their clinical and aetiological heterogeneity, the cause of NDDs can be challenging to diagnose. Comprehensive clinical care includes investigations aimed at elucidating the underlying cause, such as brain imaging and laboratory tests, including genetic testing.

Identifying the genetic aetiology of an NDD is of clinical and personal utility. Increasingly, accurate genetic diagnosis can lead to tailored therapy (and may provide information about prognosis that in turn can guide medical care and therapy). Definitive diagnosis may avoid unnecessary testing or treatments, enable surveillance for known comorbidities and help families access additional disease-specific support. Accurate diagnosis of the cause of NDDs facilitates genetic counseling and informs recurrence risks and that guides reproductive decision making and enables prenatal diagnosis. When an inherited aetiology is identified, this can provide diagnoses to other family members with NDD. Finally, by ending the diagnostic odyssey for families, identifying an aetiology can also provide psychological benefits (eg, alleviation of guilt, acceptance, closure, empowerment) and improve parental quality of life.

The diagnostic process begins with a thorough clinical assessment of the patient, which includes a medical, developmental and family history with physical examination. The presence or absence of certain clinical features may provide clues as to the underlying cause and therefore direct the appropriate investigations. If a known genetic condition is recognised, targeted testing should be done first. For example, if a 2-year-old girl presents with classical features of Rett syndrome, it is prudent to do MECP2 sequencing prior to other investigations. However, it is more often the case that the clinical presentation is non-specific, and thus there are a large number of potential genetic aetiologies. It is therefore important to broaden the testing approach, which could result in earlier diagnosis.

When a known syndrome is not readily apparent, there are a variety of testing options, including chromosomal microarray (CMA), fragile X syndrome...
IMDs: genetic disorders that result in metabolic defects due to deficiency of enzymes, membrane transporters or other functional proteins.  

GDD: an NDD characterised by significant delay (at least two SD below the mean using standardised testing) in achieving at least two developmental domains, including gross or fine motor, speech and language, cognition, social and personal functioning, and activities of daily living. A diagnosis of GDD applies to individuals under the age of 5 years who are too young to participate in standardised testing to assess intellectual functioning.  

ID: an NDD characterised by deficits in intellectual and adaptive functioning. Intellectual functioning includes reasoning, problem solving, planning, abstract thinking, judgement, academic learning and experiential learning. Adaptive functioning are the skills needed to live independently and responsibly and include communication, social skills, personal independence at home or in the community and school or work functioning.  

LD: also known as specific learning disorder; an NDD characterised by deficits in the ability to learn or use foundational academic skills (reading, writing or arithmetic), which is not better accounted for by ID.  

NDD: a group of conditions manifesting early in development, characterised by impairments of personal, social, academic or occupational functioning.  

Syndromic: for an individual with an NDD, refers to the presence of additional clinical features such as dysmorphisms, congenital malformations, medical comorbidities etc, which may suggest an underlying genetic cause (refer to table 1).

DEFINITIONS
Clinical definitions
► ADHD: a persistent pattern of inattention and/or hyperactivity impulsivity that interferes with functioning or development.
► ASD: an NDD characterised by persistent deficits in social communication and social interaction and the presence of restricted, repetitive patterns of behaviour, interests or activities. May be diagnosed with or without accompanying intellectual or language impairment.
► Dysmorphic features: visible morphological findings that differ from those commonly seen in the general population or same genetic ancestry.
► GDD: an NDD characterised by significant delay (at least two SD below the mean using standardised testing) in achieving at least two developmental domains, including gross or fine motor, speech and language, cognition, social and personal functioning, and activities of daily living. A diagnosis of GDD applies to individuals under the age of 5 years who are too young to participate in standardised testing to assess intellectual functioning.
► IMDs: genetic disorders that result in metabolic defects due to deficiency of enzymes, membrane transporters or other functional proteins.
► ID: an NDD characterised by deficits in intellectual and adaptive functioning. Intellectual functioning includes reasoning, problem solving, planning, abstract thinking, judgement, academic learning and experiential learning. Adaptive functioning are the skills needed to live independently and responsibly and include communication, social skills, personal independence at home or in the community and school or work functioning.
► LD: also known as specific learning disorder; an NDD characterised by deficits in the ability to learn or use foundational academic skills (reading, writing or arithmetic), which is not better accounted for by ID.  

Technical definitions
► Next-generation sequencing (NGS): high-throughput sequencing technology used to determine nucleotide sequences and genome dosage at numerous loci simultaneously.
► Genome-wide sequencing (GWS): sequence analysis of all or a large part of the genome performed by NGS; includes exome and ES.
  - ES: an NGS approach that determines the DNA sequence of most of the protein-coding exons found in the genome of an individual. As of 2022, this test is available in most provinces for patients meeting specific clinical criteria.
  - Genome sequencing (GS): an NGS approach that determines the sequence of most of the DNA content encompassing the entire genome of an individual. As of 2022, this test is only available on a research basis or through clinical pilots within Canada.
► Multigene panel: simultaneous sequencing of multiple genes associated with a specific clinical presentation.
► First-tier test: a laboratory test or other diagnostic investigation that should be pursued prior to all others.
► Second-tier test: a laboratory test or other diagnostic investigation that should be pursued if first-tier tests are negative or inconclusive.
► Metabolic testing: blood and/or urine tests that measure analytes (such as amino and organic acids) in order to identify patterns indicating an underlying metabolic disease.

METHODS
To develop recommendations, the Canadian College of Medical Geneticists (CCMG) formed an NDDs working group of physicians and clinical scientists from medical genetics and genomics,
CONSIDERATIONS

Target audience and scope

These recommendations apply to patients with NDDs for whom no specific aetiological diagnosis is suspected after a thorough history and physical examination. If a specific genetic disorder is suspected with a high degree of clinical certainty, targeted testing should be done first, and if negative, then one should proceed with the testing recommended in this position statement. If no diagnosis is reached after the recommended investigations, periodic re-evaluation of the patient by a geneticist is recommended, as testing technology and genomic knowledge is rapidly evolving. The time interval for re-evaluation is specific to the patient’s unique situation (eg, age and testing done) and should be determined by the geneticist.

We acknowledge that the demand for genetic consultation is high, and waitlists are often long for non-urgent referrals. Thus, we provide recommendations for first-tier testing that primary care and/or non-genetic specialist clinics can order while the patient awaits a clinical genetics or metabolic assessment to ensure faster access to genetic testing. The non-geneticist should have a good understanding of the benefits and limitations of first-tier tests and should obtain informed consent for testing from the patient or their guardian(s). Should first-tier testing be abnormal, a referral to the appropriate specialist is indicated, and many genetic/metabolic clinics will expedite consultation for patients referred with an abnormal result. We also provide guidance as to which patients should be referred to genetics if first-tier testing is not diagnostic. The role of the geneticist is to determine whether second-tier testing is indicated to ensure that the patient and family have an opportunity to discuss the test results and to make recommendations for follow-up care if a diagnosis is made.

The evidence reviewed for the development of this position statement applies specifically to patients with GDD, ID and/or ASD. The existing evidence was highly heterogeneous with respect to the clinical phenotypes of the patients studied, so it was not possible to make separate recommendations for patients with and without syndromic features, for patients with different severities of impairment or for patients in different age groups (paediatric or adult). Evidence was also reviewed for patients with other NDDs (eg, ADHD and LD), but there are no high-quality studies that enable distinction between patients with and without comorbid GDD or ID. Therefore, until such evidence emerges, we do not recommend genetic investigations for patients without GDD, ID or ASD unless there are other clinical features suggestive of a genetic (‘syndromic’; see table 1) or metabolic aetiology.

The recommendations herein apply to probands of all ages. The urgency of testing is typically higher for younger children, particularly if the parents are planning a pregnancy. This does not preclude testing and/or genetic/metabolic specialist referral for adolescents and adults, particularly if the patient has not been tested or evaluated in 5 years or more.

The recommendations herein also apply to probands with ID regardless of severity, as there was insufficient evidence to exclude probands from such testing based on level of disability. Further studies addressing this are required.

There may be specific clinical scenarios that demand a different approach to that suggested herein, so clinicians should always use their judgement as to whether the recommended investigations are appropriate. Patient and family preferences should always be considered. This position statement is not intended to be a comprehensive guide to the medical investigation of individuals with NDDs, and as such, we do not make recommendations regarding neuroimaging or health surveillance (eg, thyroid screening, vision assessment, etc).

The Canadian context

Canada’s population of nearly 39 million is concentrated in the southern parts of Ontario, Quebec and Alberta, with relatively sparse population density in the rest of the country. Twenty-one per cent of the population lives in rural or remote areas with limited access to specialist medical care (Canadian Medical Association Policy Statement, 2014). The Canada Health Act stipulates that all Canadians should have universal insurance coverage for medically necessary healthcare service. Healthcare is a provincial and territorial responsibility, and nationwide there is variability regarding the types of genetic and metabolic tests that are publicly funded. There is also some variability in the conditions screened for by newborn screening. The most expensive and resource-intensive genetic tests, such as ES, are not always accessible due to provincial healthcare insurance exclusions or limited access to specialists qualified to order and interpret this testing. The working group has taken these inequities into consideration, and where possible, we have offered alternatives, such as recommending multigene panel testing when ES is not available.

Limitations

The working group formulated the recommendations using the best available evidence at the time of writing (2022). The quality of the evidence, however, generally falls at a level II–IV (moderate) based on the GRADE system,10 so these recommendations should be considered conditional on completion of higher quality studies. Expert opinion was used to fill in the gaps where evidence was lacking.

It was beyond the scope of the working group to conduct an independent cost-effectiveness analysis as part of the development of this position statement. Recognising that resources are finite, these recommendations attempt to strike a balance between avoiding over-testing and the risk of missing a diagnosis.
that may have treatment or recurrence risk implications for the patient and their family.

Finally, input from patients and their families was not solicited in the development of these recommendations, but for future iterations of this statement, this will be an important addition.

The sections further will briefly summarise the evidence for each test modality and list recommendations for use of the test as first or second tier for patients with NDDs. It should be understood that first tier tests are, for the most part, able to be ordered by any qualified clinician and therefore should generally be done prior to (or at the same time as) referral for genetics consultation. Figure 1 summarises the recommendations and provides guidance as to when a patient should be referred to genetics or metabolics for further evaluation.

**Fragile X testing (FMR1 CGG repeat analysis)**

FXS is an X-linked condition caused by the unstable expansion of a CGG repeat in the 5′UTR of the FMR1 gene and subsequent hypermethylation, preventing gene expression. It is a common monogenic cause of NDDs, with a prevalence of 1.4 per 10 000 males and 0.9 per 10 000 females. FXS testing is widely available and can be ordered by a variety of clinicians. FXS cannot, at this time, be diagnosed by CMA or ES.

The ‘classic’ FXS phenotype in males includes moderate-to-severe ID, macro-orchidism and distinctive facial features (long face, large ears and prominent jaw). Common comorbid conditions include ASD, anxiety and hyperactivity. Deviation from the classical phenotype is influenced by age and sex (with females and prepubescent children manifesting fewer physical characteristics) and the presence of DNA methylation or repeat size mosaicism. The phenotype in females is highly variable, with IQ ranging from normal to moderate ID, likely due to the effects of random X-inactivation. FXS has a high recurrence risk in families, as children with a full mutation have mothers with a premutation that confers a risk for expansion in the oocyte, or with a full mutation. If the diagnosis of FXS is delayed, parents may not have the opportunity to have prenatal diagnosis in subsequent pregnancies, which may result in multiple affected children.

For these reasons, a variety of professional organisations recommend FXS testing as a first-line diagnostic test (concurrent with CMA) for GDD, ID and ASD. These recommendations were largely based on expert opinion and/or published prior to publication of larger studies examining the diagnostic yield of NGS-based testing for NDDs.

The diagnostic yield of fragile X testing varies between studies, based on the study design and characteristics of the study population. Taken together, the sex-specific prevalence of FXS in individuals with NDD is 2.2%–2.5% for males and 1.3%–1.6% in females. Retrospective reviews of laboratory databases demonstrate a diagnostic yield of 0%–2.5% for FXS testing. The diagnostic yield of FXS testing increases (9.5%–17%) when inclusion criteria are restricted to males with NDD and characteristic physical and behavioural features or family history suggestive of FXS. Use of a diagnostic ‘checklist’ to determine likelihood of FXS in males generally has high sensitivity, but only if a low threshold is used. There is insufficient data to determine whether this approach yields a higher diagnostic rate in females with NDD. No studies have evaluated the cost-effectiveness of selective FXS testing based on clinical criteria.

**Recommendations for fragile X testing**

1. FXS testing is recommended as a first-tier diagnostic test for individuals presenting with:
   - GDD, ID or ASD and a clinical presentation or family history suggestive of FXS (see table 2).
   - Any NDD and a family history of FXS or other FMR1-related disorder.
2. FXS testing is not recommended for individuals with GDD or ID who do not meet the above criteria and have a complex clinical presentation that is not consistent with FXS (eg, multiple congenital anomalies, profound neurological impairment).

**Chromosomal microarray**

CMA uses comparative genomic hybridisation or SNP array to detect gains and losses of chromosomal material, known as CNVs. SNP-based arrays are also able to detect areas of long contiguous stretches of homozygosity, which may suggest parental consanguinity or uniparental disomy. While CMA can detect CNVs as small as approximately 20–50 kb, resolution varies depending on the technology used and density of DNA probes. CMA is widely available and can be ordered by a variety of clinicians.

CNVs can be classified as pathogenic, likely pathogenic, uncertain significance, likely benign or benign. CNVs of uncertain
significance require interpretation from a clinical geneticist and may warrant additional investigations. CMA may detect ‘susceptibility CNVs’: recurrent CNVs with variable expressivity and/or incomplete penetrance.26–68 The phenotype resulting from these CNVs is variable and likely influenced by other genetic and environmental factors. Detection of these variants in an individual with a neurodevelopmental disorder may not entirely explain the observed phenotypes, so additional investigations may be warranted; however, this is not well studied.

A variety of professional organisations recommend CMA as a first-tier test for individuals with GDD, ID or ASD,27 32 33 49 69 based on its high diagnostic yield compared with standard karyotype.20 70 In general, the diagnostic yield of CMA is lower in individuals with ADHD,76–78 they lack sufficient clinical information on the probands to determine whether ADHD was isolated or associated with GDD/ID or syndromic features. Furthermore, in these studies, many of the detected rare CNVs were inherited and the pathogenicity and clinical impact of these CNVs were unclear. Another study found that children with isolated ADHD had similar detection of rare CNVs compared with controls.79

There are no studies that investigate the diagnostic yield of CMA for isolated speech or motor delay or LD. Although three studies report a detection rate of 8%–9% for rare CNVs in individuals with ADHD,76–78 they lack sufficient clinical information about the probands to determine whether ADHD was isolated or associated with GDD/ID or syndromic features. Furthermore, in these studies, many of the detected rare CNVs were inherited and the pathogenicity and clinical impact of these CNVs were unclear. Another study found that children with isolated ADHD had similar detection of rare CNVs compared with controls.79

There is strong evidence for the utility of CMA as a first-tier test for individuals with GDD, ID or ASD. There is insufficient evidence for the utility of CMA as a first-tier test for other NDDs such as ADHD without GDD, ID or ASD unless there are other clinical features suggestive of a syndromic aetiology (see table 1).

Recommendaions for CMA testing
1. CMA is recommended as a first-tier diagnostic test for individuals presenting with:
   - GDD, ID or ASD.
   - Other NDDS when syndromic features are present (see table 1).

Metabolic testing
Inherited metabolic disorders (IMDs) comprise a group of genetic conditions that can present with NDDs as part of the clinical spectrum. Metabolic testing can detect the presence or absence of biochemical markers that lead to a diagnosis of IMD. While individually rare, there can be clinical value in diagnosing IMDs for some disorders, early identification and treatment can improve outcomes, and most of the conditions are autosomal recessive and therefore have a high recurrence risk. According to a systematic literature review, over 100 IMDs with ID as a major feature have potential treatments.80 81 Examples of IMDs causing NDD that have disease-modifying treatments are urea cycle disorders,82–83 maple syrup urine disease,84 homocystinurias, including cobalamin-related conditions,85–86 and some creatine disorders including guanidinoacetate methyltransferase deficiency.82–84 86–87

Some authors suggest that all individuals presenting with GDD/ID/ASD should be screened for IMDs,82 83 86 87 while others recommend screening only for individuals with additional clinical features suggestive of an IMD.82 84 86 There is also a lack of consensus among experts as to which metabolic investigations comprise a comprehensive screen for IMDs. Indeed, there is wide variability in Canadian clinical practice.36

Both the Canadian Pediatric Society,27 and the American Academy of Pediatrics28 reference a systematic literature review,80 as the rationale to screen for treatable IMDs in all children presenting with GDD/ID; however, the diagnostic yield of this approach was unknown at the time. The protocol suggested by the Treatable Intellectual Disability Endeavor (TIDE) (www.tidebc.org) is extensive; yet, not all the tests in this protocol are widely available, and clinical utility and cost-effectiveness have not been evaluated.

The reported diagnostic yield of metabolic testing for patients with NDDs can vary depending on the population studied. However, using the best evidence available (more recent publications in populations with expanded newborn screening, larger case series with clear case definitions), the overall diagnostic yield of metabolic testing for patients with NDDs is 0.25%–0.42% for GDD/ID in non-consanguineous populations.84 87 Retrospective reviews of the medical records of patients with diagnosed IMDs show that most of these patients have clinical features suggestive of an IMD in addition to GDD/ID.84 87 90 In a prospective study by Campistol et al83 in 2016, an IMD diagnosis was rarely identified in children with non-syndromic ASD. A review of the pilot implementation of the TIDE protocol for patients with GDD/ID, with and without suggestive neurological features,84 demonstrated no significant increase in diagnoses of IMD during time periods before and after implementation despite a greater than fourfold increase in test volumes.84 Metabolic testing early in the diagnostic process should theoretically improve ‘time to diagnosis’ and clinical outcomes for IMDs not detected by newborn screening; however, this has not been rigorously studied.

Reviews, commentaries and guidelines that quote diagnostic yields of metabolic investigations for NDDs as high as 5% should be interpreted with caution. Literature examining the diagnostic yield of metabolic testing for patients with NDDs is generally of low quality due to lack of peer review, small sample sizes, variability in tests performed and lack of robust case definitions. Furthermore, some publications pre-date the expansion of newborn screening to detect more treatable IMDs.87 Although most children with treatable IMDs are now diagnosed by routine provincial newborn screening programmes in Canada, newborn screening cannot detect all treatable IMDs, neonatal screening programmes vary among provinces/territories and false-negative screens are possible.87 84

For children with NDDs, most IMD diagnoses are made in those with clinical features such as regression or plateauing of development, acute encephalopathy or altered level of
consciousness, movement disorders, intractable seizures, multi-
systemic involvement, specific ophthalmological findings, organomegaly and/or features suggestive of a storage disease. Patients with such ‘red flags’ should be referred for specialist consultation without delay. Table 3 provides guidance on addi-
tional investigations that should be ordered, where possible, for patients with these features while they await consultation.

Given its very low yield, routine metabolic testing is not recommended for patients with GDD/ID/ASD without suggest-
tive clinical features listed in table 3. Rarely, a treatable IMD may be missed by not performing metabolic testing for such patients; for example, patients with homocystinurias, including cobalamin-related conditions, can occasionally present with isolated NDDs. However, these conditions are very rare, and those with time-sensitive treatments are detected by newborn screening. Creatine transporter disorder, which is X-linked, may rarely present with isolated NDD. Treatment does not alter clinical outcomes for this condition; however, diagnosis might be helpful for future family planning.

Clinicians should be aware of the conditions tested for via newborn screening in their jurisdiction. Children who have not had newborn screening may be at increased risk for having an undiagnosed IMD, so clinicians should determine what newborn screening was done and, if limited, consider referring to a metabolic specialist.

**Table 3** Clinical features suggestive of inherited metabolic disorders (IMDs) and suggested metabolic testing that could be done while awaiting metabolics (or other specialist) consultation

<table>
<thead>
<tr>
<th>Clinical feature</th>
<th>Major groups of IMDs to consider</th>
<th>Targeted metabolic testing if URGENT, clearly indicate on the requisitions when ordering tests</th>
</tr>
</thead>
<tbody>
<tr>
<td>Developmental plateau or regression in the context of an abnormal neurological exam</td>
<td>LSD, peroxisomal disorders (X-ALD), UCD, HCYS</td>
<td>Ammonia, blood gas, lactate, PAA, UOA, TPH, urine MPS, VLCFA†</td>
</tr>
<tr>
<td>Altered level of consciousness, especially if episodic; stroke-like episodes</td>
<td>UCD, MSUD, HCYS, organic acidurias, mitochondrial disorders</td>
<td>Ammonia, blood gas, glucose, lactate, electrolytes, anion gap, PAA, TPH, ACP, UOA</td>
</tr>
<tr>
<td>Movement disorder (ataxia, dystonia, choreatetohisis, myoclonus, tremor)</td>
<td>Organic acidurias, HCYS, creatine disorders, LSD, Wilson disease</td>
<td>Blood gas, lactate, glucose, ACP, PAA, UOA, TPH, urine MPS, creatine panel†, copper, ceruloplasmin</td>
</tr>
<tr>
<td>MRI/MRS brain abnormality (eg, white matter change, abnormal cerebellum / basal ganglia)</td>
<td>Peroxisomal disorders, organic acidurias, LSD</td>
<td>Metabolic testing tailored to MRI findings by metabolic/genetic specialist</td>
</tr>
<tr>
<td>Hepatomegaly, splenomegaly*</td>
<td>LSD, peroxisomal disorders</td>
<td>Urine MPS (if other systemic abnormalities noted*), VLCFA†</td>
</tr>
<tr>
<td>Specific food aversions: avoiding high protein foods</td>
<td>UCD, MSUD and organic acidurias</td>
<td>PAA, ammonia, UOA, ACP</td>
</tr>
<tr>
<td>Ophthalmological findings (most common: cataracts, dislocated lens, corneal deposits, retinopathy, cherry red spot)</td>
<td>Galactosaemia, Lowe disease, sulfite oxidase deficiency, HCYS, LSD, peroxisomal disorders</td>
<td>Metabolic testing tailored to ophthalmological findings by metabolic/genetic specialist</td>
</tr>
<tr>
<td>Seizures: drug resistant, myoclonic or neonatal</td>
<td>UCD, organic acidurias, HCYS, creatine disorders, vitamin-dependent epilepsies, peroxisomal disorders, Menkes disorder</td>
<td>Ammonia, blood gas, glucose, lactate, electrolytes, anion gap, PAA, UOA, TPH, creatine panel†, VLCFA†</td>
</tr>
<tr>
<td>Abnormal tone (hypotonia or spasticity)</td>
<td>Organic acidurias, HCYS, creatine disorders, UCD (eg, arginase deficiency and HHH), biotinidase deficiency, peroxisomal disorders, LSD</td>
<td>Ammonia, blood gas, glucose, lactate, electrolytes, anion gap, PAA, UOA, TPH, VLCFA†, creatine panel†</td>
</tr>
<tr>
<td>Coarse facial features and/or skeletal abnormalities on X-ray*</td>
<td>MPS and other storage disorders</td>
<td>Urine MPS</td>
</tr>
<tr>
<td>Multisystemic involvement</td>
<td>LSD, peroxisomal disorders, peroxisomal disorders, CDG, SLOS</td>
<td>Metabolic testing tailored to specific findings by metabolic/genetic specialist</td>
</tr>
</tbody>
</table>

X-linked ALD (adrenoleukodystrophy). Highly specialised tests that are not recommended for initial workup are not included.

*For mucopolysaccharidoses (MPS), look for noisy breathing, short stature, macrocephaly and sensorineural hearing impairment. For MPS III (Sanfilippo syndrome), aggressive behaviour, sleep disturbance and hirsutism may be early features.

†Test ordering may be restricted to certain specialists.

ACP, acylcarnitine profile; CDG, congenital disorders of glycosylation; HCYS, homocystinuria and remethylation/cobalamin disorders; HHH, hyperornithinaemia hyperammonaemia homocitrullinuria syndrome; LSD, lysosomal storage disorder; MRS, Magnetic resonance spectroscopy; MSUD, maple syrup urine disease; PAA, plasma amino acids; SLOS, Smith-Lemli-Opitz syndrome; TPH, total plasma homocysteine; UCD, urea cycle disorder; UOA, urine organic acids; VLCFA, very long chain fatty acids.

**ES and multigene panels**

Massively parallel ‘next-generation’ sequencing technology has allowed for the simultaneous analysis of hundreds to thousands of genes from a single DNA sample. Currently, ES is widely available as a clinical diagnostic test. An alternative to ES is the use of comprehensive multigene panels, which are targeted to a specific phenotype. For NDDs, commercially available panels vary widely in the number of genes analysed; the largest ones include over 2500 genes. In future, GS is poised to replace both ES and CMA, but as it is not yet publicly funded in Canada, these recommendations consider the use of panels and ES only.

The main advantage of ES and large panels is the ability to simultaneously interrogate large numbers of genes, which means that the clinician need not have a particular genetic condition in mind to direct the diagnosis. However, they have important limitations that means that a negative test does not rule out a genetic cause. Depending on the methodology used, all coding exons may not be completely sequenced and, thus, some variants may be missed. ES is less able to detect


designation (see table 3) and a referral made to a metabolic specialist promptly.

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1. Metabolic testing is recommended as a first-tier diagnostic test for individuals presenting with:

- GDD, ID, or ASD and clinical features suggestive of an IMD. Metabolic testing should be tailored to the presentation (see table 3) and a referral made to a metabolic specialist promptly.

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mosaism or exon-level deletions compared with GS or panels that do not reliably assess repetitive DNA sequences (such as trinucleotide repeats), intronic or non-coding variants, methylation, epigenetic or mitochondrial DNA variants or balanced chromosomal rearrangements.94

These tests can identify multiple rare variants in a patient’s sample, the clinical relevance of which can be challenging to decipher. They can identify misattributed parentage (when done as a ‘trio’ with both declared parents),95 96 as well as incidental and secondary findings (pathogenic variants in medically actionable genes unrelated to the indication for testing).97 Further, they can identify misattributed parentage (when done, as they often are, as a ‘trio’ with both declared parents) and secondary findings (pathogenic variants in medically actionable genes unrelated to the indication for testing).95 97 Testing of the proband only (vs proband-parent trio) results in a lower diagnostic yield98 99 and more labour-intensive downstream analysis and/or additional testing to resolve variants, so a ‘trio’ approach is the most useful, time-efficient and informative approach.98 100

Given these considerations, these tests require robust pretest and post-test genetic counselling and input from a clinical geneticist.97 101

ES and comprehensive multigene panels were not widely available as routine clinical tests when the practice guidelines for genetic testing for patients with NDDs were published by the American Association of Neurology in 201126 and the American Association of Pediatrics in 2014.28 However, a 2015 CCMG position statement recommended clinical GWS (either exome or genome sequencing) for patients with moderate-to-severe or syndromic ID, when appropriate first-tier genetic testing (such as CMA) is non-diagnostic.102 The 2021 ACGM practice guidelines103 strongly recommend that GWS be considered as a first-tier or second-tier test for paediatric patients with developmental delays and/or ID.

Diagnostic yield of ES for patients with NDDs

The reported diagnostic yield of ES for patients with GDD/ID demonstrates wide variation (8%–68%).9 16 74 98 103–118 likely owing to differences in cohort size and composition, whether ES was performed clinically or under a research protocol and whether a proband-parent trio or proband-only approach was used. When considering the largest studies from clinical laboratories,105 106 the diagnostic yield of ES is 26%–31%, consistent with results from a meta-analysis that showed pooled estimates of 29%.104 It is difficult to tease out the diagnostic yield in specific patient subgroups (such as ASD without GDD/ID, or GDD/ID without syndromic features). A few studies dedicated to patients with ASD, most of whom had ID (with or without other features), reported that ES had a diagnostic yield of 8%–25.8%,74 106 109–111 119 120 The few studies that have compared the diagnostic yield of ES for GDD/ID patients with and without syndromic features have not found statistically significant differences.107 120 121

Diagnostic yield of NGS panels for patients with NDDs

A few studies have examined comprehensive panels for patients with NDDs, reporting a diagnostic yield of 11%–39%,108 122 124 Two studies directly compared a targeted NDD gene panel to ES, and both demonstrated slightly lower diagnostic yields for the panel.117 120

Clinical utility of ES for individuals with NDDs

The clinical utility of ES for the diagnosis of patients with suspected monogenic disorders has been well established. A 2020 Ontario Health Technology Assessment examined the clinical utility of ES in unexplained developmental disabilities and multiple congenital anomalies, concluding that ES allows for changes in clinical management, provides insight into the natural history of the patient’s condition, informs reproductive planning and optimises the patient’s ability to access disease-specific supports.99 Similar findings were reported in a 2020 ACMG systematic evidence review that examined the health, clinical, reproductive and psychosocial outcomes resulting from ES for the investigation of congenital anomalies, developmental delay or ID.125 It also considered possible negative impacts such as insurance discrimination, financial burden and psychological impact on patients and found that return of ES test results incur no clinically significant psychological harms with low levels of test-related distress and positive psychological effects. Genetic counselling and an appropriate consenting process can help mitigate any associated risks.103

Cost-effectiveness of ES

The significant cost of ES has historically been a barrier to its implementation within the Canadian healthcare system as standard-of-care testing for NDDs. Recent evidence shows that performing ES early in the diagnostic testing pathway is cost-effective.99 113 126–131 although other studies have been inconclusive.132–134 In Ontario, a 2020 cost analysis of ES testing before or concurrently with CMA for patients with developmental disabilities supported ES as a second-tier test (when CMA is non-diagnostic) as the most cost-effective approach99; this study is based on ES analysis that does not include duplication/deletion analysis.

In summary, the evidence strongly supports using ES as a diagnostic test for patients with ID/GDD due to its high diagnostic yield, cost-effectiveness and demonstrable clinical utility. The diagnostic yield for patients with GDD/ID is significant even in the absence of syndromic features. For patients with ASD without GDD/ID, there is insufficient data to make evidence-based recommendations, although evidence suggests a higher diagnostic yield for patients with syndromic features (table 1). A trio-based testing approach is recommended. While ES provides a higher diagnostic yield, a comprehensive multigene panel for NDDs is a suitable substitute when ES is not available, especially when done as a trio-based test. When GS becomes a widely available clinical test, it is anticipated to replace fragile X, CMA and ES as a single test of choice for NDDs.

In most cases ES is recommended as a second-tier test after CMA, as the latter has a high diagnostic yield, is more widely available, less costly and can be ordered by non-geneticists. However, it may be appropriate to do GWS before or concurrently with CMA; for example, when the proband is highly likely to have an autosomal recessive condition (based on parental consanguinity and/or affected siblings), or when a diagnosis must be rapidly obtained, such as to inform the management of an ongoing pregnancy (for a sibling of the proband), or to guide medical care in a patient who is acutely ill or whose clinical status is deteriorating (eg, neurological regression).

Recommendations for ES or multigene panels

1. Recommended as a second-tier diagnostic test for individuals presenting with:
   - GDD or ID (with or without ASD).
CONCLUSIONS

Patients with NDDs in Canada require a thorough evaluation for the underlying cause of their condition. The CCMG Neurodevelopmental Disorders working group recommends an approach to the diagnostic evaluation that is tailored to the patient and based on current evidence. These recommendations are based on best available evidence as of 2022 and may evolve as new evidence emerges. Therefore, they should be reviewed every 2–3 years.

An overview of these recommendations are found in figure 1. We recommend a tiered testing strategy, such that first-tier testing is ordered by non-geneticists (family doctors, paediatricians, neurologists, etc), to allow patients faster access to testing and potentially a rapid diagnosis. The provision of genetic testing by non-specialists should be supported with enhanced genetics education at all levels of medical training. We provide guidelines for when a patient should be referred to a specialist in genetics or metabolics: specifically, anyone with GDD/ID (regardless of other features), as well as patients with ASD or other NDDs if they have syndromic features (table 1) or features suggestive of an inherited metabolic disease (table 3).

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Position statement


