Fetal alcohol spectrum disorder: Canadian guidelines for diagnosis

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Abstract

The diagnosis of fetal alcohol spectrum disorder (FASD) is complex and guidelines are warranted. A subcommittee of the Public Health Agency of Canada’s National Advisory Committee on Fetal Alcohol Spectrum Disorder reviewed, analysed and integrated current approaches to diagnosis to reach agreement on a standard in Canada. The purpose of this paper is to review and clarify the use of current diagnostic systems and make recommendations on their application for diagnosis of FASD-related disabilities in people of all ages. The guidelines are based on widespread consultation of expert practitioners and partners in the field. The guidelines have been organized into 7 categories: screening and referral; the physical examination and differential diagnosis; the neurobehavioural assessment; and treatment and follow-up; maternal alcohol history in pregnancy; diagnostic criteria for fetal alcohol syndrome (FAS), partial FAS and alcohol-related neurodevelopmental disorder; and harmonization of Institute of Medicine and 4-Digit Diagnostic Code approaches. The diagnosis requires a comprehensive history and physical and neurobehavioural assessments; a multidisciplinary approach is necessary. These are the first Canadian guidelines for the diagnosis of FAS and its related disabilities, developed by broad-based consultation among experts in diagnosis.

In this document, we discuss the diagnostic approach to disabilities associated with prenatal alcohol exposure. Fetal alcohol spectrum disorder (FASD), along with its most visible presentation, fetal alcohol syndrome (FAS), is a serious health and social concern to Canadians. FASD is an umbrella term describing the range of effects that can occur in an individual whose mother drank alcohol during pregnancy. These effects may include physical, mental, behavioural and learning disabilities with lifelong implications. The term FASD is not intended for use as a clinical diagnosis. FASD is the result of maternal alcohol consumption during pregnancy and has implications for the affected person, the mother, the family and the community. Since FAS was first described in 1973, it has become apparent that it is complex; affected people exhibit a wide range of expression, from severe growth restriction, intellectual disability, birth defects and characteristic dysmorphic facial features to normal growth, facial features and intellectual abilities, but with lifelong deficits in several domains of brain function. FASD requires a medical diagnosis in the context of a multidisciplinary assessment. FASD itself is not a diagnostic term. The purpose of this paper is to review and clarify the use of the current diagnostic systems and make recommendations on their application for diagnosis of FASD-related disabilities in people of all ages. For a description of the characteristics and the natural course of FASD, consult some of the broader reviews.

Epidemiology of FASD

The prevalence of FAS in the United States has been reported as 1–3 per 1000 live births and the rate of FASD as 9.1 per 1000 live births. However, diagnosis may often be delayed or missed entirely.

There are no national statistics on the rates of FASD in Canada, although studies have estimated its prevalence in small populations. In an isolated Aboriginal community in British Columbia, FASD prevalence was 190 per 1000 live births. In northeastern Manitoba, an incidence of about 7.2 per 1000 live births was found. In another Manitoba study in a First Nations community, the prevalence of FAS and partial FAS was estimated to be 55–101 per 1000. In their survey, Asante and Nelms-Matzke estimated the rate of FAS and related effects at 46 per 1000 native Canadian children in the Yukon and 25 per 1000 in northern British Columbia. Based on referrals to a diagnostic clinic in Saskatchewan, the rate of FAS was estimated at 0.589 per 1000 live births in 1988–1992 and 0.515 per 1000 in 1973–1977. However, none of these data should be generalized to other communities, other populations or the Canadian population in general.

Risk factors

A common misconception is that FASD is associated with ethnocultural background. However, the data suggest that risk factors for prenatal alcohol exposure include higher maternal age and lower education level, prenatal exposure to cocaine and smoking, custody changes, lower socioeconomic status and paternal drinking and...
drug use at the time of pregnancy; and reduced access to prenatal and postnatal care and services, inadequate nutrition and a poor developmental environment (e.g., stress, abuse, neglect). In a 5-year follow-up study of birth mothers of children with full FAS, Astley and colleagues found that these women came from diverse racial, educational and economic backgrounds. They were often challenged by untreated or under-treated mental health concerns, they were socially isolated, they were victims of abuse and they had histories of severe childhood sexual abuse.

Because there are no large-scale studies of risk factors and because risks are interrelated and could be different for different populations, it is difficult to provide accurate figures for relative risk. However, the most important risk factor for FASD is related to high blood-alcohol concentration: the timing of exposure during fetal development, the pattern of consumption, i.e., binge drinking (4 or more drinks per occasion) and the frequency of use. Although there seems to be no definite threshold of exposure, there appears to be a dose-response relation.

**Importance of early diagnosis**

An early diagnosis is essential to allow access to interventions and resources that may mitigate the development of subsequent “secondary disabilities” (e.g., unemployment, mental health problems, trouble with the law, inappropriate sexual behaviour, disrupted school experience) among affected people. Furthermore, an early diagnosis will also allow appropriate intervention, counselling and treatment for the mother and may prevent the birth of affected children in the future. It may also prompt caregivers to seek diagnosis and support for previously undiagnosed siblings. A review of medical and behavioural management of those with FASD can be found in other sources. Astley and Clarren suggest that accurate and timely diagnosis is essential to improve outcome, as misclassification leads to inappropriate patient care, increased risk of secondary disabilities, missed opportunities for prevention and inaccurate estimates of incidence and prevalence. Together, these inaccuracies could hinder efforts to allocate sufficient social and health care services to the vulnerable populations and preclude accurate assessment of primary prevention efforts.

Because of limited capacity and expertise and the need to involve several professionals in a comprehensive multidisciplinary diagnostic evaluation, only a fraction of those affected currently receive a diagnosis. Results from the Canadian national survey regarding knowledge and attitudes of health professionals suggest that standardized guidelines for diagnosis and further professional education and training are needed for practitioners to participate in diagnosis. In response to these concerns, Health Canada’s National Advisory Committee on FASD, along with experts and practitioners in FAS diagnosis and treatment, present the following guidelines for diagnosis.

**Process of guideline development**

These guidelines are the result of more than 10 face-to-face consultations with Canadian and American experts in the diagnosis of FAS and its related disabilities (Appendix 1). Many of the participants are currently providing diagnostic services across Canada. Review and feedback were provided by a diverse group of individuals; professional organizations and societies; and provincial, territorial and federal levels of government. Guidelines are presented in 6 areas related to the diagnostic process: 1. screening and referral; 2. the physical examination and differential diagnosis; 3. neurobehavioural assessment; 4. treatment and follow-up; 5. maternal alcohol history in pregnancy; and 6. diagnostic criteria for FAS, partial FAS and alcohol-related neurodevelopmental disorder. We also include recommendations for harmonization of the 2 main approaches to diagnosis.

There are multiple approaches to diagnosis, and the working group sought to integrate these to achieve consistent diagnoses across Canada. Current knowledge of the complexity of the disabilities associated with prenatal alcohol exposure dictates that a comprehensive, multidisciplinary assessment is necessary to make an accurate diagnosis and provide recommendations for management. We are recommending such a multidisciplinary approach. This approach will also allow for collection of Canadian data for estimating incidence and prevalence of FASD. This information is essential to identify the need for and the development of appropriate prevention and intervention programs and services.

**Background and terminology for the diagnosis of FAS**

The first recognition of a variety of birth defects and developmental disabilities in offspring born to alcoholic parents is attributed to Lemoine and colleagues. A specific pattern of birth defects following maternal alcohol exposure was described in the United States. The specific pattern, referred to as FAS, consists of facial abnormalities (smooth philtrum [the space between the upper lip and the nose], thin vermilion border [the exposed mucosal, or red part, of the upper lip], short palpebral fissures), impaired prenatal or postnatal growth (or both) and central nervous system or neurobehavioural disorders. Alcohol probably acts through multiple mechanisms and a range of disabilities has been observed in the absence of dysmorphic features reflecting varying degrees of damage during fetal development; undoubtedly, timing and degree of exposure are important variables that contribute to the variation. Thus, the term “suspected fetal alcohol effects” (FAE) was created. These “effects” were further delineated by the United States’ Institute of Medicine (IOM), which published recommendations in 1996 for diagnosis of FAS in consultation with a panel of experts.
The diagnostic categories presented were: FAS with and without a confirmed history of alcohol exposure, partial FAS, alcohol-related birth defects (ARBD), and alcohol-related neurodevelopmental disorder (ARND) (Table 1).

In the late 1990s, another diagnostic strategy was developed by Astley and Clarren. They created a 4-Digit Diagnostic Code using data from the Washington State Fetal Alcohol Syndrome Diagnostic and Prevention Network of clinics. The system uses quantitative, objective measurement scales and specific case definitions. The 4 digits in the code reflect the magnitude of expression or severity of the 4 key diagnostic features of FAS in the following order: growth deficiency; the FAS facial phenotype; central nervous system damage or dysfunction; gestational exposure to alcohol. The magnitude of expression of each feature is ranked independently on a 4-point Likert scale with 1 reflecting complete absence of the feature and 4 reflecting its extreme expression. The 4-Digit Diagnostic Code is now being used for diagnosis, screening and surveillance in clinics throughout the United States and Canada. Terminology from Astley’s 2004 revision of the 4-Digit Diagnostic Code are used in this article.

Although the approaches are different, the underlying, fundamental criteria of the IOM and the 4-Digit Diagnostic Code are similar. Some clinics are choosing to integrate the diagnostic tools and precision reflected in the 4-Digit Diagnostic Code with the diagnostic categories and language recommended by the IOM committee. Although both IOM criteria and the 4-Digit Diagnostic Code have been published, many clinicians still use the less desirable language recommended by the IOM committee. Although both IOM criteria and the 4-Digit Diagnostic Code have been published, many clinicians still use the less desirable and potentially misleading gestalt approach (Table 2).

The diagnostic process

The diagnostic process consists of screening and referral, the physical examination and differential diagnosis, the neurobehavioural assessment and treatment and follow-up. Because of the complexity and the range of expression of dysfunction related to prenatal alcohol exposure, a multidisciplinary team is essential for an accurate and comprehensive diagnosis and treatment recommendations. The assessment process begins with recognition of the need for diagnosis and ends with implementation of appropriate recommendations. The multidisciplinary diagnostic team can be geographic, regional or virtual; it can also accept referrals from distant communities and carry out an evaluation using telemedicine.

The core team may vary according to the specific context, but ideally it should consist of the following professionals with appropriate qualifications, training and experience in their particular discipline:

- Coordinator for case management (e.g., nurse, social worker).
- Physician specifically trained in FASD diagnosis.
- Psychologist.
- Occupational therapist.
- Speech-language pathologist.

Additional members may include addiction counsellors, childcare workers, cultural interpreters, mental health workers, parents or caregivers, probation officers, psychiatrists, teachers, vocational counselors, nurses, geneticists or dysmorphologists, neuropsychologists, family therapists.

Comments

Clearly, funding for development, training and maintenance of multidisciplinary diagnostic teams is necessary so that major centres will have the expertise and capacity to serve their communities. To optimize the outcome of the diagnosis, the community and the family must be prepared, ready to participate in, and be in agreement with the diagnostic assessment. The diagnostic process should be sensitive to the family’s and the caregiver’s needs. In each community, referrals must be evaluated and their level of priority established. The family and guardian must be in agreement on the purpose of diagnosis. They must be made aware of the potential psychosocial consequences of a diagnosis of FASD (e.g., increasing a sense of guilt and anger, especially with the birth mother, or potential stigmatization of the child). The family or guardian will likely need help to move confidently through the diagnostic process. This help might include some preparatory education concerning FASD and linking them with community supports and resources.

Information from multiple sources (e.g., school records, hospital records, social services, previous assessments) should be obtained; this might involve meetings with relevant professionals who know the patient (e.g., teachers, physicians, social workers, psychologists). Other relevant documentation would include birth and pregnancy records, medical and hospital records, adoption records, academic records, achievement tests, developmental assessments, psychological and psychometric assessments, legal reports and documentation of the family history.

The comprehensive assessment by the diagnostic team provides important information about the individual’s unique needs and allows interventions to be tailored to his or her strengths and challenges. The post-diagnostic report should state the basis for the diagnosis by including the history of alcohol use, the physical criteria and the psychological data that support it.

Multidisciplinary teams work with community partners and resources to develop and implement management plans to maximize the potential of the affected individual. Following assessment, a report containing recommendations should be made available to caregivers, educators, and biological families, as well as other appropriate indi-

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*Astley SJ. Diagnostic Guide for Fetal Alcohol Spectrum Disorders: The 4-Digit Diagnostic Code (3rd edition). Seattle: University of Washington Publication Services; 2004.
viduals who work with the child (i.e., daycare workers, early intervention workers, social workers, etc). The team findings should be discussed with the guardian. Older children who have the cognitive ability should have the opportunity to learn about their diagnosis from the team. The team might also take on the responsibility for facilitating and providing follow-up with the family and community resources regarding outcomes of the recommendations. Ultimately, the diagnostic process will result in concrete management recommendations to improve the lives of the affected individuals, their families and the communities.

Canada is a large country with vast distances between communities, some of which are remote and isolated. Specialists providing consultation to remote areas require specialized training in FASD assessment and need to link with centres that have multidisciplinary teams to assist in the diagnostic process. A number of tools may be useful for distant diagnosis. More frequent use of telemedicine, for example, will allow assessment of children in distant communities.31 Other examples include the use of digital photographs32,33 and 3-D laser surface scanning34,35 sent electronically to teams in larger centres.

We recognize that there is currently a limited capacity even in some large communities in Canada to provide a multidisciplinary team-based approach to FASD diagnosis. Professionals should make the best use of available resources and expertise to provide an accurate assessment and treatment plan for affected individuals and their families, recognizing the key role of psychology.

1. Screening and referral

Recommendations

1.1 All pregnant and post-partum women should be screened for alcohol use with validated screening tools (i.e., T-ACE, TWEAK) by relevant health care providers. Women at risk for heavy alcohol use should receive early brief intervention (i.e., counselling).

1.2 Abstinence should be recommended to all women during pregnancy, as the mother’s continued drinking during pregnancy will put the fetus at risk for effects related to prenatal alcohol exposure.

1.3 Referral of individuals for a possible FASD-related diagnosis should be made in the following situations:

| Table 1: Institute of Medicine diagnostic criteria for fetal alcohol syndrome and alcohol-related effects |
|----------------------------------------------------------|
| **Fetal alcohol syndrome (FAS)**                        |
| 1. **FAS with confirmed maternal alcohol exposure**      |
| A. Confirmed maternal alcohol exposure*                 |
| B. Evidence of a characteristic pattern of facial anomalies that includes features such as short palpebral fissures and abnormalities in the premaxillary zone (e.g., flat upper lip, flattened philtrum and flat midface) |
| C. Evidence of growth retardation, as in at least one of the following: |
|   • low birth weight for gestational age |
|   • decelerating weight over time not due to nutrition |
|   • disproportionally low weight-to-height ratio |
| D. Evidence of central nervous system neurodevelopmental abnormalities, as in at least one of the following: |
|   • decreased cranial size at birth |
|   • structural brain abnormalities (e.g., microcephaly, partial or complete agenesis of the corpus callosum, cerebellar hypoplasia) |
|   • neurologic hard or soft signs (as age appropriate), such as impaired fine motor skills, neurosensory hearing loss, poor tandem gait, poor eye–hand coordination |
| 2. **FAS without confirmed maternal alcohol exposure**    |
| B, C, and D as above                                      |
| 3. **Partial FAS with confirmed maternal alcohol exposure** |
| A. Confirmed maternal alcohol exposure*                 |
| B. Evidence of some components of the pattern of characteristic facial anomalies |
| Either C or D or E                                       |
| C. Evidence of growth retardation, as in at least one of the following: |
|   • low birth weight for gestational age |
|   • decelerating weight over time not due to nutrition |
|   • disproportionally low weight-to-height ratio |
| D. Evidence of CNS neurodevelopmental abnormalities, e.g., |
|   • decreased cranial size at birth |
|   • structural brain abnormalities (e.g., microcephaly, partial or complete agenesis of the corpus callosum, cerebellar hypoplasia) |
|   • neurologic hard or soft signs (as age appropriate) such as impaired fine motor skills, neurosensory hearing loss, poor tandem gait, poor eye–hand coordination |
| E. Evidence of a complex pattern of behaviour or cognitive abnormalities that are inconsistent with developmental level and cannot be explained by familial background or environment alone: e.g., learning difficulties; deficits in school performance; poor impulse control; problems in social perception; deficits in higher level receptive and expressive language; poor capacity for abstraction or metacognition; specific deficits in mathematical skills; or problems in memory, attention or judgment. |
a. Presence of 3 characteristic facial features (short palpebral fissures, smooth or flattened philtrum, thin vermilion border).
b. Evidence of significant prenatal exposure to alcohol at levels known to be associated with physical or developmental effects, or both.
c. Presence of 1 or more facial features with growth deficits plus known or probable significant prenatal alcohol exposure.
d. Presence of 1 or more facial features with 1 or more central nervous system deficits plus known or probable significant prenatal alcohol exposure.
e. Presence of 1 or more facial features with pre- or postnatal growth deficits, or both (at the 10th percentile or below [1.5 standard deviations below the mean]) and 1 or more central nervous system deficits plus known or probable significant prenatal alcohol exposure.

1.4 Individuals with learning or behavioural difficulties, or both, without physical or dysmorphic features and without known or likely prenatal alcohol exposure should be assessed by appropriate professionals or specialty clinics (i.e., developmental pediatrics, clinical genetics, psychiatry, psychology) to identify and treat their problems.

**Comments**

Screening should not be equated with diagnosis. We know that in some places with no diagnostic services, screening tools have been inappropriately used in lieu of a proper diagnosis. One purpose of screening is to identify and refer pregnant women who may be at risk for an alcohol use disorder and who may place their child at risk for FASD. Several alcohol screening tools have been found to be effective in identifying problem drinking in a primary care setting.

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Table 1: continued

**Alcohol-related effects**

Clinical conditions in which there is a history of maternal alcohol exposure,*† and where clinical or animal research has linked maternal alcohol ingestion to an observed outcome. There are 2 categories, which may co-occur. If both diagnoses are present, then both diagnoses should be rendered.

4. **Alcohol-related birth defects (ARBD)**
   Congenital anomalies, including malformations and dysplasias

   **Cardiac**
   - Atrial septal defects
   - Ventricular septal defects
   - Tetralogy of Fallot

   **Skeletal**
   - Hypoplastic nails
   - Clinodactyly
   - Pectus excavatum and carinatum
   - Klippel-Feil syndrome
   - Hemivertebrae
   - Scoliosis

   **Renal**
   - Aplastic, dysplastic, hypoplastic kidneys
   - Ureteral duplications
   - Horseshoe kidneys
   - Hydronephrosis

   **Ocular**
   - Strabismus
   - Refractive problems secondary to small globes

   **Auditory**
   - Conductive hearing loss
   - Neurosensory hearing loss

   **Other**
   - Virtually every malformation has been described in some patient with FAS. The etiologic specificity of most of these anomalies to alcohol teratogenesis remains uncertain.

5. **Alcohol-related neurodevelopmental disorder (ARND)**

   Presence of A or B or both.

   A. Evidence of CNS neurodevelopmental abnormalities, as in any one of the following:
   - decreased cranial size at birth
   - structural brain abnormalities (e.g., microcephaly, partial or complete agenesis of the corpus callosum, cerebellar hypoplasia)
   - neurologic hard or soft signs (as age appropriate), such as impaired fine motor skills, neurosensory hearing loss, poor tandem gait, poor eye-hand coordination

   B. Evidence of a complex pattern of behaviour or cognitive abnormalities that are inconsistent with developmental level and cannot be explained by familial background or environment alone; e.g., learning difficulties; deficits in school performance; poor impulse control; problems in social perception; deficits in higher level receptive and expressive language; poor capacity for abstraction or metacognition; specific deficits in mathematical skills; or problems in memory, attention or judgment.

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*A pattern of excessive intake characterized by substantial, regular intake or heavy episodic drinking. Evidence of this pattern may include frequent episodes of intoxication, development of tolerance or withdrawal, social problems related to drinking, legal problems related to drinking, engaging in physically hazardous behaviour while drinking or alcohol-related medical problems such as hepatic disease.

†As further research is completed and as, or if, lower quantities or variable patterns of alcohol use are associated with ARBD or ARND, these patterns of alcohol use should be incorporated into the diagnostic criteria.
care setting (e.g., TWEAK, T-ACE, CAGE, AUDIT, S-MAST, B-MAST). There is moderate evidence to support the use of T-ACE and TWEAK to identify women who would benefit from intervention for alcohol use during pregnancy. If the woman cannot abstain, she should receive support and be referred to appropriate counselling and treatment. Stopping drinking at any point during the pregnancy will improve the outcome for the baby. Research is being carried out to develop gender and culturally appropriate instruments for the screening of all women during their child-bearing years.

The purpose of screening individuals at risk for the effects of prenatal alcohol exposure is to determine whether a pattern of learning and behavioural problems may be related to prenatal alcohol exposure. The screening could be conducted through the education system, the mental health system, the judicial system or social services. The purpose of screening should be to facilitate referral to a diagnostic clinic and highlight the need for referral and support for the birth mother.

The FAS Diagnostic and Prevention Network has had encouraging results in applying the FAS facial photographic screening tool in foster children and school-age children populations. However, in the wide array of FASDs, facial dysmorphology is often absent and, in the final analysis, has little importance compared with the impact of prenatal alcohol exposure on brain function. However, it is important to note that the facial phenotype is a midline defect that is the most sensitive and specific marker for alcohol-related brain damage.

All those suspected of having brain dysfunction should be referred to an appropriate professional or clinic for assessment (i.e., developmental pediatrics, clinical genetics, psychiatry, psychology). Because of the specificity of FASD clinics in addressing issues related to prenatal alcohol exposure, those with no prenatal alcohol exposure should be referred to an appropriate professional or clinic for assessment, treatment and follow-up.

2. The physical examination and differential diagnosis

The purpose of dysmorphology assessment is to identify those with features related to prenatal alcohol exposure and also to identify children with dysmorphic features due to other causes. Occasionally, children with prenatal alcohol effects may have another genetic syndrome as a comorbidity. When in doubt and if feasible, a genetic dysmorphology assessment is advisable.

A general physical and neurologic examination, including appropriate measurements of growth and head size, assessment of characteristic findings and documentation of anomalies (e.g., cleft palate, congenital heart defects, epicanthic folds, high arched palate, poorly aligned or abnormal teeth, hypertelorism, micrognathia, abnormal hair pattern, abnormal palmar creases, skin lesions) is required to exclude the presence of other genetic disorders or multifactorial disorders that could lead to features mimicking FAS or partial FAS (Table 3). Some children will have significant neurologic deficits, such as deafness, blindness or seizures, and these should be

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### Table 2: 4-Digit Diagnostic Code criteria for FASD

| Rank | Growth deficiency | FAS facial phenotype | CNS damage or dysfunction | Gestational exposure to alcohol |
|------|-------------------|----------------------|---------------------------|--------------------------------|
| 4    | **Significant**   | **Severe**           | **Definite**              | **High risk**                  |
|      | Height and weight below 3rd percentile | All 3 features: PFL 2 or more SDs below mean | Structural or neurologic evidence | Confirmed exposure to high levels |
|      |                   | Thin lip: rank 4 or 5 Smooth philtrum: rank 4 or 5 |
| 3    | **Moderate**      | **Moderate**         | **Probable**              | **Some risk**                  |
|      | Height and weight below 10th percentile | Generally 2 of the 3 features | Significant dysfunction across 3 or more domains | Confirmed exposure. Level of exposure unknown or less than rank 4 |
| 2    | **Mild**          | **Mild**             | **Possible**              | **Unknown**                    |
|      | Height or weight below 10th percentile | Generally 1 of the 3 features | Evidence of dysfunction, but less than rank 3 | Exposure not confirmed present or absent |
| 1    | **None**          | **Absent**           | **unlikely**              | **No risk**                    |
|      | Height and weight at or above 10th percentile | None of the 3 features | No structural, neurologic or functional evidence of impairment | Confirmed absence of exposure from conception to birth |

Note: PFL = palpebral fissure length; SD = standard deviation.
assessed and documented as essential components of the child’s profile. These features do not discriminate alcohol-exposed from unexposed children. The face of FAS is the result of a specific effect of ethanol teratogenesis altering growth of the midface and brain. Those exposed to other embryotoxic agents may display a similar, but not identical, phenotypic facial development, impaired growth, a higher frequency of anomalies and developmental and behavioural abnormalities (for a review, see Chudley and Longstaffe"). However, because FAS facial criteria have been restricted to short palpebral fissures, smooth philtrum and thin upper lip, there is far less overlap with the facial phenotypes associated with other syndromes. Knowledge of exposure history will decrease the possibility of misdiagnosing FASD.

Children may be found to need other medical assessments to address co-occurring issues. For example, sleep

| Syndrome                                      | Overlapping features                                                                 | Features of this syndrome that differentiate it from FAS                        |
|-----------------------------------------------|--------------------------------------------------------------------------------------|----------------------------------------------------------------------------------|
| Aaruskog syndrome                             | Widely spaced eyes, small nose with anteverted nares, broad philtrum, mid-facial recession | Round face, downslanted palpebral fissures, widow’s peak, prominent “lop” ears, specific contracture of digits on extension. Inherited as an x-linked trait. Molecular defect identified. |
| Brachman-de Lange or Cornelia de Lange syndrome | Long philtrum, thin vermilion border of upper lip, depressed nasal bridge, anteverted nares, microcephaly | Single eyebrow across eyes and forehead (synophrys), long eyelashes, downturned corners of mouth, short upper limbs particularly involving ulnar side, very short stature. Molecular defect identified. |
| Dubowitz syndrome                              | Short palpebral fissures, widely-spaced eyes, epicanthal folds, variable ptosis (droopy eyes) and blepharophimosis, microcephaly | Shallow supraorbital ridges, broad nasal tip, cleft palate                           |
| Fetal anticonvulsant syndrome (includes fetal hydantoin and fetal valproate syndromes) | Widely-spaced eyes, depressed nasal bridge, mid-facial recession, epicanthal folds, long philtrum, thin vermilion border of upper lip | Bowed upper lip, high forehead, small mouth                                           |
| Maternal phenylketonuria (PKU) fetal effects   | Epicanthal folds, short palpebral fissures, long poorly formed philtrum, thin vermilion border of upper lip, microcephaly | Prominent glabella, small up turned nose, round face                                 |
| Noonan syndrome                                | Low nasal bridge, epicanthal folds, wide spaced eyes, long philtrum                   | Down-slanted palpebral fissures, wide mouth with well-formed philtrum, protruding upper lip. Molecular defect identified. |
| Toluene embryopathy                            | Short palpebral fissures, mid face hypoplasia, smooth philtrum, thin vermilion border upper lip, microcephaly | Large anterior fontanelle, hair patterning abnormalities, ear anomalies              |
| Williams syndrome                              | Short palpebral fissures, anteverted nares, broad long philtrum, maxillary hypoplasia, depressed nasal bridge, epicanthic folds, microcephaly | Wide mouth with full lips and pouting lower lip, stellate pattern of iris, periorbital fullness, connective tissue dysplasia, specific cardiac defect of supravalvar aortic stenosis in many. Chromosome deletion on FISH (fluorescent in situ hybridization) probe analysis of 7q. |
| Other chromosome deletion and duplication syndromes | Many have short palpebral fissures, mid-facial hypoplasia, smooth philtrum.          | Chromosomal analysis by standard analysis and some select syndromes by specific FISH probe analysis |
disturbance is common with prenatal alcohol exposure and medical problems related to obstructive sleep apnea may have been overlooked previously. Atypical seizures may also be present and endocrinopathies may exist as a comorbid reason for growth deficiency. These individuals should be assessed by appropriate health professionals.

2a. Growth

**Recommendations**

2.1 Growth should be monitored to detect deficiency. Presence of pre- or post-natal growth deficiency, defined as height or weight at or below the 10th percentile (1.5 standard deviations below the mean) or a disproportionately low weight-to-height ratio (at or below the 10th percentile) using appropriate norms. To determine that a child is growth deficient requires taking into consideration confounding variables such as parental size, genetic potential and associated conditions (e.g., gestational diabetes, nutritional status, illness).

**Comments**

Children affected by prenatal alcohol exposure may have prenatal or postnatal growth deficits. They can be small for gestational age in utero and remain below average throughout their lives with respect to head circumference, weight and height. Many children can have normal growth parameters, but be at risk in later development for clinically significant learning, behavioural and cognitive deficits. If there is no alcohol exposure in the third trimester, the growth parameters can be normal. Gestational diabetes can lead to increased fetal size, which can mask the effects of growth retardation from prenatal alcohol exposure. Furthermore, if the infant is born into a family or a community where “normal” size is above the average for the general population, growth impairment may be masked if the child is compared with standard growth parameters rather than community norms. Growth deficiencies may not persist with age, and infant growth records may not be available for adults coming in for assessment for the first time. There is a need to establish growth norms for the Canadian population and subpopulations that differ from the general population.

2b. Facial features

**Recommendations**

2.2 The 3 characteristic facial features that discriminate individuals with and without FAS are:

- Short palpebral fissures, at or below the 3rd percentile (2 standard deviations below the mean).
- Smooth or flattened philtrum, 4 or 5 on the 5-point Likert scale of the lip-philtrum guide. &nbsp;&n

- Thin vermilion border of the upper lip, 4 or 5 on the 5-point Likert scale of the lip-philtrum guide.

2.3 Associated physical features (abnormalities such as midface hypoplasia, micrognathia, abnormal position or formation of the ears, high arched palate, hypertelorism, epicanthic folds, limb and palmar crease abnormalities and short-upturned nose) should be recorded but do not contribute to establishing the diagnosis.

2.4 Facial features should be measured in all age groups. If a patient’s facial features change with age, the diagnosis of the facial features should be based on the point in time when the features were most severely expressed. When diagnosing adults, it can be helpful to view childhood photographs.

**Comments**

A characteristic craniofacial profile associated with FAS was first described by Jones and Smith in 1975 and later refined by Astley, Clarren and others. Individuals with FAS have short palpebral fissures, a thin upper lip and an...
indistinct philtrum (Fig. 1). Palpebral fissure length, philtrum and upper lip differ with race and age. Growth and facial anthropometric data are needed for the specific population, as sensitivity and specificity of the assessment will be lowered without the use of appropriate norms. Some discriminating characteristic features in FAS (i.e., upper lip or philtrum) may become less recognizable with age, making accurate diagnosis more difficult in older groups, but facial features should always be measured. More longitudinal research is needed to correlate changes in these characteristic physical findings in adolescents and adults diagnosed with FAS or partial FAS.

Palpebral fissure length (Fig. 2) is difficult to measure accurately without training. Thomas and co-workers41 have published norms for palpebral fissure length at 29 weeks gestation to 14 years. There are a number of opinions about which norms are appropriate,41-44 but it is generally agreed that all are flawed in some respect.

Two graphs of palpebral fissure length are presented in Appendix 2. Some discrepancies exist. Both studies used North American white subjects; standards for other populations in Canada are not currently available. Appendix 2-1 may be more reliable when measuring palpebral fissure length using a plastic ruler (in the experience of one of the authors); Appendix 2-2 may be more reliable if slide calipers are used (in the experience of one of the authors). Percentile ranks for both graphs seem to be in agreement until age 7 years, after which Appendix 2-2 shows longer palpebral fissures in older children and adolescents than Appendix 2-1. We believe this may be due to differences in measurement technique. Because calipers are not a common tool in most medical clinics, we recommend the use of a clear flexible plastic ruler.

There is a need to establish updated norms for all ages and subpopulations. Astley and Clarren25,39 have developed norms for the assessment of the lip and philtrum using their pictorial guide. Lip-philtrum guides were developed for use in Caucasian and African-American populations, but no standards are currently available for other populations.

3. Neurobehavioural assessment

Recommendations

3.1 The following domains should be assessed:

a. Hard and soft neurologic signs (including sensory-motor signs).

b. Brain structure (occipitofrontal circumference, magnetic resonance imaging, etc.).

c. Cognition (IQ).

d. Communication: receptive and expressive.

e. Academic achievement.

f. Memory.

g. Executive functioning and abstract reasoning.

h. Attention deficit/hyperactivity.

i. Adaptive behaviour, social skills, social communication.

3.2 The assessment should include and compare basic and complex tasks in each domain, as appropriate.

3.3 The domains should be assessed as though they were independent entities, but where there is overlap experienced clinical judgment is required to decide how many domains are affected.

3.4 A domain is considered “impaired” when on a standardized measure:

a. Scores are 2 standard deviations or more below the mean, or

b. There is a discrepancy of at least 1 standard deviation between subdomains. For example:

i. Verbal v. non-verbal ability on standard IQ tests,

ii. Expressive v. receptive language,

iii. Verbal v. visual memory, or

c. There is a discrepancy of at least 1.5–2 standard deviations among subtests on a measure, taking into account the reliability of the specific measure and normal variability in the population.

3.5 In areas where standardized measurements are not available, a clinical judgment of “significant dysfunction” is made, taking into consideration that important variables, including the child’s age, mental health factors, socioeconomic factors and disrupted family or home environment (e.g., multiple foster placements, history of abuse and neglect), may affect development but do not indicate brain damage.

3.6 Evidence of impairment in 3 domains is necessary for a diagnosis, but a comprehensive assessment requires that each domain be assessed to identify strengths and weaknesses.

3.7 The diagnosis should be deferred for some at-risk children (e.g., preschool-age) who have been exposed to al-
cochlor but may not yet demonstrate measurable deficits in the brain domains or may be too young to be tested in all the domains. However, developmental assessment should identify areas for early intervention.

Examples of tests that are most widely used to assess the domains and their criteria are provided in Appendix 3.

Comments

Research reports have documented a range of cognitive and behavioural outcomes associated with prenatal alcohol exposure. Contemporary studies have reported some of these outcomes in the absence of FAS physical features. Currently, no modal profile of abilities has been found to be unique to alcohol exposure, is observed in all those with prenatal alcohol exposure, or can be distinguished from that observed with some other neurobehavioural disorders. Furthermore, not every deficit that we may identify in a child with prenatal exposure to alcohol may be solely the result of alcohol exposure. An expert analysis of neurodevelopmental deficits caused by a range of teratogens and congenital disorders failed to result in a consensus on core deficits associated only with FASD.4

Research and experience has shown that features of FASD are complex and multifaceted, originating with organic brain damage caused by alcohol, but interacting with genetic and other influences. Over the lifespan of the affected person, these features may be exacerbated or mitigated by environmental experiences.

To make the diagnosis of FAS, features such as microcephaly, structural abnormalities (as may be detected on brain scans) and hard neurologic signs are taken as strong evidence of organic brain damage. We believe that low-average to borderline intelligence and soft neurologic signs alone are insufficient evidence of brain damage because they are frequently found in the general population. Features such as learning difficulties, attention deficit/hyperactivity disorder and deficits in adaptive skills, memory, higher-level language and abstract thinking are frequently seen in children with prenatal alcohol exposure, but also among those with other etiologies. These deficits can be multifactorial in etiology and can also be attributed to genetics or postnatal experiences.

The 4-Digit Diagnostic Code evaluation of the FASD brain is based on levels of certainty, in the judgement of the clinician, that the individual’s cognitive and behavioural problems reflect brain damage. A higher rating may reflect a more severe expression of functional disability, asynchronous patterns across domains or certainty based on deficits in multiple domains. The determination is based on objective evidence of “substantial deficiencies or discrepancies across multiple areas of brain performance.”

The IOM also requires “evidence of a complex pattern of behavior or cognitive abnormalities that are inconsistent with developmental level and cannot be explained by familial background or environment alone, such as learning difficulties; deficits in school performance; poor impulse control; problems in social perception; deficits in higher level receptive and expressive language; poor capacity for abstraction or metacognition; specific deficits in mathematical skills; or problems in memory, attention, or judgment,” but is much less specific than the 4-Digit Diagnostic Code with regard to the criteria for determining the deficit.

We have adapted the method of the 4-Digit Diagnostic Code with regard to identifying domains and severity of impairment or certainty of brain damage. Current research shows overlap between the neurobehavioural outcomes in FAS and ARND groups when neuropsychologic data are compared. In addition, we believe that a single feature such as microcephaly is not a sufficient indicator of brain damage for the purposes of an FAS diagnosis because it may reflect genetic or ethnic differences not reflected in currently available physical norms. Our concern is that there may be an over-diagnosis of FAS if evidence of brain damage is based on a single indicator as allowed by both the 4-Digit Diagnostic Code and the IOM models. An individual showing hard neurologic signs or structural brain abnormalities (i.e., true brain damage) will likely show additional functional deficits in the listed domains. A diagnosis of full FAS will not be denied by combining the criteria for full FAS and ARND in this harmonized system.

Although the domains are considered to be separate and independent entities, there is obviously overlap. For example, a discrepancy between verbal and non-verbal scores on an IQ test (taking into account normal variability in the population) may be reflecting a specific language disability. If language is deficient, can deficits in verbal memory be considered an additional domain? Does a language deficit represent brain damage if the child has experienced a prolonged period of social deprivation? The cut-off of 2 standard deviations below the mean on standardized tests is recommended to increase confidence that abilities in the domain are impaired as a result of brain damage and are scored as “3” (significant dysfunction) on the 4-Digit Diagnostic Code. With 3 such domains, the brain rank is 3: “probable brain dysfunction.”

We realize that in standard neuropsychologic practice, 1.5 standard deviations below the mean may indicate subtle impairments. Using the 4-Digit Diagnostic Code, the domains would be scored as “moderate dysfunction” and may result in a brain rank of 2: “possible brain dysfunction.” These more subtle findings are an important part of the individual’s profile. For the purpose of diagnosis, however, and the certainty that the scores represent injury caused by alcohol, the more extreme cut-off is recommended. The multidisciplinary team, reviewing the data and using experienced clinical judgement, is critical in making an accurate diagnosis as qualitative aspects of performance are also important. The diagnostic profile is dynamic and may change over time; thus individuals affected or suspected to be affected may require several assessments over time. Services should not be based on the diagnosis itself, but rather on the profile of brain function-dysfunction.
4. Treatment and follow-up

Recommendations

4.1 Education of the patient and family members on features of FASD is crucial. The potential psychosocial tensions that might be expected to develop within the family as a result of the diagnosis should also be discussed. This must be done in a culturally sensitive manner using appropriate language.

4.2 A member of the diagnostic team should follow-up outcomes of diagnostic assessments and treatment plans within a reasonable length of time to assure that the recommendations have been addressed.

4.3 Diagnosed individuals and their families should be linked to resources and services that will improve outcome. However, where services are limited in the community, an individual should not be denied an assessment for diagnosis and treatment. Often the diagnosis in the individual is the impetus that leads to the development of resources.

5. Maternal alcohol history in pregnancy

Recommendations

5.1 Prenatal alcohol exposure requires confirmation of alcohol consumption by the mother during the index pregnancy based on reliable clinical observation, self-report, reports by a reliable source or medical records documenting positive blood alcohol, alcohol treatment or other social, legal or medical problems related to drinking during the pregnancy.

5.2 The number and type(s) of alcoholic beverages consumed (dose), the pattern of drinking and the frequency of drinking should all be documented if available.

5.3 Hearsay, lifestyle, other drug use or history of alcohol exposure in previous pregnancies cannot, in isolation, be informative of drinking patterns in the index pregnancy. However, co-occurring disorders, significant psychosocial stressors and prenatal exposure to other substances (e.g., smoking, licit or illicit drugs) in the index and previous pregnancies should still be recorded, based on known interactive effects of these variables on the severity of pregnancy outcomes for both the mother and her offspring.

Comments

Gathering reliable information about maternal drinking is key to establishing an accurate diagnosis. Special attention must be paid to inquiring about maternal alcohol use before the woman recognized that she was pregnant. Some women do not consider that their prior drinking is important and many underreport it. Training is required in how to obtain this information in a non-threatening, non-judgmental way.

Canadian survey data suggest that the number of women who report drinking during pregnancy has decreased. The National Population Health Survey, 1994–1995 and National Longitudinal Survey of Children and Youth, 1994–1995 reported that 17–25% of women drank alcohol at some point during their pregnancy and 7–9% drank alcohol throughout their pregnancy. According to the National Longitudinal Survey of Children and Youth, 1998–1999, 14.4% of women drank at some point during their pregnancy and 4.9% drank throughout their entire pregnancy (3% reported binge drinking during pregnancy). In the Fall 2002 Survey of First Nations People Living on Reserve, 53% of the respondents said that cutting down or stopping alcohol use was important for women to have a healthy baby.

The evaluation of “significant alcohol exposure” is often confusing. The IOM describes significant alcohol exposure as “a pattern of excessive intake characterized by substantial, regular intake or heavy episodic drinking” (the National Institute on Alcohol, Alcoholism, and Alcohol Abuse defines heavy alcohol use as drinking 5 or more drinks per occasion on 5 or more days in the past 30 days). Evidence of this pattern may include frequent episodes of intoxication, development of tolerance or withdrawal, social problems related to drinking, legal problems related to drinking, engaging in physically hazardous behaviour while drinking, or alcohol-related medical problems such as hepatic disease. As further research is completed and as, or if, lower quantities or variable patterns of alcohol use are associated with alcohol-related birth defects (ARBD) or ARND, these patterns of alcohol use should be incorporated into the diagnostic criteria.

6. Diagnostic criteria for FAS, partial FAS and ARND

Recommendations

6.1 The criteria for the diagnosis of fetal alcohol syndrome, after excluding other diagnoses, are:

A. Evidence of prenatal or postnatal growth impairment, as in at least 1 of the following:
   a. Birth weight or birth length at or below the 10th percentile for gestational age.
   b. Height or weight at or below the 10th percentile for age.
   c. Disproportionately low weight-to-height ratio (at 10th percentile).

B. Simultaneous presentation of all 3 of the following facial anomalies at any age:
   a. Short palpebral fissure length (2 or more standard deviations below the mean).
   b. Smooth or flattened philtrum (rank 4 or 5 on the lip-philtrum guide).
c. Thin upper lip (rank 4 or 5 on the lip-philtrum guide).

C. Evidence of impairment in 3 or more of the following central nervous system domains: hard and soft neurologic signs; brain structure; cognition; communication; academic achievement; memory; executive functioning and abstract reasoning; attention deficit/hyperactivity; adaptive behaviour, social skills, social communication.

D. Confirmed (or unconfirmed) maternal alcohol exposure.

6.2 The diagnostic criteria for partial fetal alcohol syndrome, after excluding other diagnoses, are:
A. Simultaneous presentation of 2 of the following facial anomalies at any age:
   a. Short palpebral fissure length (2 or more standard deviations below the mean).
   b. Smooth or flattened philtrum (rank 4 or 5 on the lip-philtrum guide).
   c. Thin upper lip (rank 4 or 5 on the lip-philtrum guide).

B. Evidence of impairment in 3 or more of the following central nervous system domains: hard and soft neurologic signs; brain structure; cognition; communication; academic achievement; memory; executive functioning and abstract reasoning; attention deficit/hyperactivity; adaptive behaviour, social skills, social communication.

C. Confirmed maternal alcohol exposure.

6.3 The diagnostic criteria for alcohol-related neurodevelopmental disorder, after excluding other diagnoses, are:
A. Evidence of impairment in 3 or more of the following central nervous system domains: hard and soft neurologic signs; brain structure; cognition; communication; academic achievement; memory; executive functioning and abstract reasoning; attention deficit/hyperactivity; adaptive behaviour, social skills, social communication.

B. Confirmed maternal alcohol exposure.

6.4 The term alcohol-related birth defects (ARBD) should not be used as an umbrella or diagnostic term, for the spectrum of alcohol effects. ARBD constitutes a list of congenital anomalies, including malformations and dysplasias and should be used with caution (Table 1).

Comments

Our definition of partial FAS differs from the published IOM criteria. Where significant prenatal alcohol exposure is known and there is significant growth retardation and significant indicative facial features but no evidence of brain involvement, a diagnosis of partial FAS could be made using the IOM criteria. It is our view that, using the term partial FAS in the absence of measurable brain deficits could be harmful for the individual because the diagnosis of partial FAS implies brain dysfunction. If some characteristic facial features and growth impairment, without significant developmental or behavioural problems, are found in children under 6 years of age, it would be prudent to say that the child may be at risk of learning and behaviour problems at a later time due to prenatal alcohol exposure. No alcohol-related diagnosis should be made, but the child must be monitored by the family physician or health care worker and deficits should be documented using a neurodevelopmental assessment.

The term “partial” in partial FAS does not imply that these individuals are less severely impaired in day-to-day functioning than those with a diagnosis of FAS, as the deficits in brain function may be similar.

7. Harmonization of the Institute of Medicine (IOM) and 4-Digit Diagnostic Code approaches

Recommendations

7.1 The approach identified in the 4-Digit Diagnostic Code should be used to describe, assess and measure objectively alcohol exposure, growth, facial features and brain damage. The 4-Digit Diagnostic Code should be recorded for each assessment and may be useful for surveillance and research purposes.

7.2 The terminology in the IOM criteria should be used to describe the diagnosis.

Comments

Table 4 and Table 5 illustrate how we recommend harmonizing the IOM and 4-Digit Diagnostic Code criteria. The ARBD category has limited utility in the diagnosis, but we do recognize that alcohol is teratogenic and may be responsible for birth defects if exposure occurs during critical periods of development. However, in the absence of other features of FAS or brain deficits, it is difficult to attribute causation.

Future research related to diagnostic guidelines

The lack or unavailability of evidence and data in key areas limits the effectiveness of the diagnostic process, in general. Such key areas include the development of Canadian growth and anthropometric norms for all ages and ethno-cultural groups. There is also a need for the development and validation of screening tools that are specific and sensitive to prenatal alcohol exposure. These tools should be adaptable for use in various contexts, they should be culturally appropriate and they should lead to accurate referrals for diagnosis and assessment.
Emerging issues

Biomarkers

Often, women will not accurately recall the amount or frequency of alcohol consumption during pregnancy. Some women may also underestimate consumption level or deny that they drank alcohol during pregnancy. Medical records are known to be incomplete with respect to maternal alcohol history. Currently, there are no reliable means to confirm maternal drinking using biochemical markers in pregnancy. High levels of whole blood-associated acetaldehyde, carbohydrate-deficient transferrin, gamma-glutamyl transpeptidase and mean red blood cell volume may be useful markers in pregnant women.50

Studies are underway to determine the utility of fatty acid ethyl esters in meconium as markers for prenatal exposure to alcohol.51-53 This marker will only be useful if it can be established that fatty acid ethyl ester levels in meconium are predictive of developmental outcome. Meconium testing could alert caregivers to infants who might be at risk for alcohol effects and lead to appropriate monitoring, intervention and prevention. Ethical issues regarding informed consent surround the use of biological markers in the baby that may indicate maternal drinking.

Recent innovations have led to the development of laser surface scanning, a non-invasive method for acquiring 3-dimensional images.33,34 This technique is promising in the analysis of facial features associated with prenatal alcohol exposure, but, at present, is a research tool only.

Remote and rural areas

The availability of diagnostic services is limited in rural and remote areas. A community may not have access to a diagnostic team or resources and services. Until regionally based diagnostic teams are established, the use of telemedicine for distant diagnosis, consultation and training may be helpful.31 Recent advances using digital imaging and computer-assisted analysis for the diagnosis of characteristic features of FAS have shown promise for analysis of facial features associated with prenatal alcohol exposure.32,33,44

Table 4: Harmonization of Institute of Medicine (IOM) nomenclature and 4-digit diagnostic code ranks for growth, face, brain and alcohol history

| IOM nomenclature                        | 4-digit diagnostic code ranks |
|-----------------------------------------|------------------------------|
| FAS (with confirmed exposure)           | Growth deficiency 2, 3 or 4  | FAS facial phenotype 3 or 4  | CNS damage or dysfunction 3 or 4  | Gestational exposure to alcohol 3 or 4  |
| FAS (without confirmed exposure)        | 2, 3 or 4                   | 3 or 4                      | 3 or 4                         | 3 or 4                              |
| Partial FAS (with confirmed exposure)*  | 1, 2, 3 or 4                | 2, 3 or 4                   | 3 or 4                         | 3 or 4                              |
| ARND (with confirmed exposure)          | 1, 2, 3 or 4                | 1 or 2                      | 3 or 4                         | 3 or 4 (2 for < 6 years) |

Note: ARND = alcohol-related neurodevelopmental disorder; CNS = central nervous system; FAS = fetal alcohol syndrome.

Source: Developed by Kwadwo Asante and Julianne Conry

*Any 4-digit code that can be made with these combinations of numbers and that is not also an FAS code signifies partial FAS. Combinations of face 2 that include two significant facial features also meet criteria for partial FAS.

Table 5: Comparison of Institute of Medicine (IOM) and 4-Digit Diagnostic Code methods in the diagnosis of FAS

| Feature                                      | IOM                | 4-Digit Diagnostic Code |
|----------------------------------------------|--------------------|-------------------------|
| Facial characteristics                       |                    |                         |
| Number of features required                  | Not specified      | 3 of 3                  |
| Thin (flat) upper lip                        | Yes                | Yes                     |
| Flattened philtrum                          | Yes                | Yes                     |
| Flat midface                                 | Yes                | No                      |
| Short palpebral fissures                     | Yes                | Yes                     |
| Other features                               | ?                  | No                      |
| Growth                                       |                    |                         |
| Number of features required                  | 1                  | 1                       |
| Low birth weight alone                       | Yes, percentile not specified | No                     |
| Decelerating weight over time                | Yes                | No                      |
| Low weight-to-height ratio                   | Yes                | No                      |
| Low height and low weight                    | No                 | Yes, ≤ 10th percentile  |
| Central nervous system dysfunction           |                    |                         |
| Number of features required                  | 1 structural or neurologic feature | 1 structural or neurologic feature OR 3 domains of significant impairment in function |
| Structural features may include:            |                    |                         |
| Microcephaly at birth                        | Yes, percentile not specified | Yes, ≤ 3rd percentile         |
| Structural abnormalities                     | Yes                | Yes                     |
| Hard neurologic signs                        | Yes                | Yes                     |
| Soft neurologic signs                        | Yes                | No                      |
Adult diagnosis

Diagnosis of adults creates special challenges in all aspects of the diagnosis. Physical features may change over time, there may be catch-up growth, and environmental influences may distort the evaluation of brain function. The adult's history may include additional traumatic head injury, alcohol and drug abuse, and mental health problems. Although tests for the various domains are readily available, clinicians working with the adult FASD population find that the tests are often not sensitive to real-life issues. In addition to the data required for the diagnosis, an assessment must include additional components such as functional literacy and numeracy, employability and quality of life, which fall within the domain of adaptive skills. The clinician should not rely solely on the self-report of the individual who is alcohol-affected; the history and abilities of the individual must be verified by a reliable source.

Conclusion

The assessment for prenatal alcohol exposure is a diagnosis for the affected person, the birth mother and possibly affected siblings. Rather than labeling, a diagnosis provides a blueprint for early intervention. Treatment planning and implementation, specifically targeted toward the unique needs of the individual and the family, form a large part of the diagnosis.

These guidelines and recommendations have been developed in parallel and in consultation with a United States committee charged with the same task. The challenges for prevention and diagnosis of FASD and intervention to assist those affected by this disorder are evolving and dynamic. Research is ongoing to determine whether tools, such as novel brain imaging techniques, biomarkers and DNA micro-array techniques, might enhance accurate and reliable alcohol-related diagnoses and treatment.

We hope that these guidelines and recommendations will be used to facilitate training of health professionals, improve access to diagnostic services and facilitate referral for intervention or treatment for all people and families living with this disability.

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# Appendix 1: Participants* in meetings and teleconferences to develop Canadian guidelines for the diagnosis of FAS and its related disabilities

| Participant          | City         | Province | Profession                                           | # Consultations |
|----------------------|--------------|----------|------------------------------------------------------|-----------------|
| Albert Chudley        | Winnipeg     | Man.     | Physician, Clinic for Drug and AlcoholExposed Children | 12              |
| Al Kircher            | Winnipeg     | Man.     | Psychologist                                         | 1               |
| Andrea Moser          | Ottawa       | Ont.     | Correctional Services Canada                         | 1               |
| Anne Fuller           | Vancouver    | BC       | BC Ministry of Children and Family Development        | 1               |
| Annette Lemire        | Edmonton     | Alta.    | Health and Wellness                                  | 2               |
| Arthur Blue           | Brandon      | Man.     | Native Psychologists in Canada                       | 1               |
| Ben Giddard           | Calgary      | Alta.    | Physician, Alberta Children’s Hospital               | 1               |
| Billie Jean Benisty   | Ottawa       | Ont.     | Health Canada                                        | 2               |
| Bob Armstrong         | Vancouver    | BC       | Physician, BC Women’s and Children’s Hospital         | 2               |
| Bonnie Baxter         | Vancouver    | BC       | Speech/Language Pathologist                          | 1               |
| Brad Bell             | Whitehorse   | YT       | Health and Social Services                           | 1               |
| Brian Marder          | Edmonton     | Alta.    | Career Counsellor                                    | 1               |
| Bryce Lark            | Whitehorse   | YT       | Health and Social Services                           | 1               |
| Carol Gregson         | Iqaluit      | Nun.     | Nunavut Dept of Health                                | 1               |
| Carol Woodworth       | Vancouver    | BC       | Speech/Language Pathologist, Asante Centre for FAS   | 1               |
| Cathie Royle          | St. John’s   | Nfld.    | Child Youth and Family Programs, Dept. Health and Community Services | 2               |
| Christine Lilley      | Vancouver    | BC       | Psychologist, BC Women’s and Children’s Hospital     | 1               |
| Christine Loock       | Vancouver    | B.C.     | Physician, BC Women’s and Children’s Hospital        | 1               |
| Claudette Landry      | Fredericton  | N.B.     | Public Health, Dept. Health and Wellness             | 2               |
| Dan Dubovsky          | Washington   | DC       | FAS Specialist, FAS Center of Excellence              | 2               |
| Darlene MacDonnell    | Ottawa       | Ont.     | Health Canada                                        | 2               |
| Darren Joslin         | Edmonton     | Alta.    | Health and Wellness                                  | 2               |
| Dawn Ridd             | Winnipeg     | Man.     | Manitoba Health                                      | 1               |
| Del Nyberg            | BC           |          | BC Health                                            | 2               |
| Diane Fast            | Vancouver    | BC       | Psychiatrist, BC Women’s and Children’s Hospital     | 1               |
| Donna Ludvigsen       | Edmonton     | Alta.    | Health and Wellness                                  | 1               |
| Edward Cross          | Kanhawake    | Que.     | Education Specialist                                 | 2               |
| Elaine Orrbine        | Ottawa       | Ont.     | Canadian Association of Pediatric Health Centres, Canadian Pediatric Chairs | 1               |
|                      |              |          | Psychologist, Toronto Hospital for Sick Children     | 1               |
| Ellen Fantus          | Toronto      | Ont.     | Canadian Nurses Association                          | 1               |
| Faye Brooks           | Ottawa       | Ont.     | Canadian Nurses Association                          | 1               |
| Faye Stark            | Fort Providence | NWT   | Health and Social Services                           | 1               |
| Fjola Hart-Wasekeeksiw | Ottawa     | Ont.     | Aboriginal Nurses Association of Canada              | 2               |
| Fred Boland           | Kingston     | Ont.     | Psychologist, Queen’s University                     | 6               |
### Appendix 1: continued

| Name                  | City          | Province | Position                                           | Institution or Organization                                      | Number |
|-----------------------|---------------|----------|---------------------------------------------------|-------------------------------------------------------------------|--------|
| Gail Andrew           | Edmonton      | Alta     | Physician, Glenrose Rehabilitation Hospital       |                                                                   | 4      |
| Gideon Koren          | Toronto       | Ont.     | Physician, Toronto Hospital for Sick Children     |                                                                   | 6      |
| Graham Robinson       | Ottawa        | Ont.     | RCMP                                              |                                                                   | 1      |
| Guy Burbon            | Ottawa        | Ont.     | Solicitor General                                 |                                                                   | 1      |
| Hasu Rajani           | Cold Lake     | Alta.    | Physician, Lakeland Centre for FAS               |                                                                   | 2      |
| Holly Mackay          | Ottawa        | Ont.     | Health Canada                                     |                                                                   | 1      |
| Irena Nulman          | Toronto       | Ont.     | Physician, Toronto Hospital for Sick Children     |                                                                   | 3      |
| Jacquelyn Bertrand    | Atlanta       | GA       | Psychologist, Centers for Disease Control and Prevention |                                                                   | 1      |
| Jan Lutke             | Vancouver     | BC       | BC FAS Support Network                            |                                                                   | 1      |
| Janice Birney         | Ottawa        | Ont.     | Indian and Northern Affairs Canada                |                                                                   | 1      |
| Jasjeet Sidhu         | Atlanta       | GA       | Medical Epidemiologist, Centers for Disease Control and Prevention |                                                                   | 1      |
| Jo Nanson             | Saskatoon     | Sask.    | Psychologist                                      |                                                                   | 4      |
| Joanne Rovet          | Toronto       | Ont.     | Psychologist, Toronto Hospital for Sick Children  |                                                                   | 2      |
| Joanne Weinberg       | Vancouver     | BC       | Neuroscientist                                     |                                                                   | 1      |
| Jocelynn Cook         | Ottawa        | Ont.     | Health Canada                                     |                                                                   | 11     |
| Jocylene Gauthier     | Whitehorse    | YT       | Health and Social Services                        |                                                                   |        |
| John Amett            | Winnipeg      | Man.     | Psychologist                                      |                                                                   | 1      |
| John Godel            | Campbell River| BC       | Physician                                          |                                                                   | 1      |
| John Service          | Ottawa        | Ont.     | Canadian Psychological Society                    |                                                                   | 1      |
| Julie Conry           | Vancouver     | BC       | Psychologist, Asante Centre for FAS               |                                                                   | 11     |
| Karen Archbell        | Toronto       | Ont.     | Ontario Dept of Health                            |                                                                   | 1      |
| Kathleen Montpetit    | Montreal      | Que.     | Occupational Therapist, Shriners Hospital         |                                                                   | 2      |
| Kathleen Montpetit    | Montreal      | Que.     | Occupational Therapist, Shriners Hospital         |                                                                   | 1      |
| Kathy Home            | Edmonton      | Alta.    | Psychologist, Glenrose Rehabilitation Hospital    |                                                                   | 1      |
| Kathy Jones           | Winnipeg      | Man.     | Psychologist, West Region First Nation Child and Family Centre |                                                                   | 1      |
| Kelly Stone           | Ottawa        | Ont.     | Director, Health Canada                           |                                                                   | 3      |
| Kwardwo Asante        | Vancouver     | BC       | Physician, Asante Centre for FAS                  |                                                                   | 4      |
| Leigh Wincott         | Thompson      | Man.     | Physician, Thompson Diagnostic Clinic for FAS     |                                                                   | 2      |
| Leslie Grob           | Regina        | Sask.    | Saskatchewan Health                               |                                                                   | 2      |
| Margaret Clarke       | Calgary       | Alta     | Physician, Alberta Children's Hospital           |                                                                   | 3      |
| Marie Adele Davis     | Ottawa        | Ont.     | Canadian Pediatric Society                        |                                                                   | 1      |
| Marilou Reeve         | Ottawa        | Ont.     | Youth Justice                                     |                                                                   | 1      |
| Marilyn Van Bibber    | Vancouver     | BC       | BC FAS Resource Network                           |                                                                   | 1      |
| Mary Cox-Millar       | Winnipeg      | Man.     | Coordinator, Clinic for Drug and Alcohol Exposed Children |                                                                   | 1      |
| Mary Ellen Baldwin    | Calgary       | Alta.    | Psychologist, Alberta Children's Hospital         |                                                                   | 1      |
| Mary Johnston         | Ottawa        | Ont.     | Health Canada                                     |                                                                   | 4      |
| Mary Lynch            | Saint John    | NB       | New Brunswick Family Services                      |                                                                   | 1      |
| Mercedes Mompel       | Toronto       | Ont.     | Health and Long-term Care                         |                                                                   | 1      |
## Appendix 1: continued

| Name                | Location       | Province | Organisation/Institution                                      | Count |
|---------------------|----------------|----------|---------------------------------------------------------------|-------|
| Michelle Dubik      | Winnipeg       | Man.     | Healthy Child Manitoba                                        | 2     |
| Nadine Huggins      | Ottawa         | Ont.     | Health Canada                                                 | 3     |
| Nancy Taylor        | Halifax        | NS       |                                                              | 1     |
| Nicole Chatel       | Yellowknife    | NWT      | Stanton Territorial Health Authority                         | 1     |
| Nicole LeBlanc      | Moncton        | NB       | Physician, Georges Dumont Hospital                            | 5     |
| Nikki Bansil        | Ottawa         | Ont.     | Canadian Medical Association                                  | 1     |
| Pamela Massad       | Ottawa         | Ont.     | Health Canada                                                 | 1     |
| Patricia Blakely    | Saskatoon      | Sask.    | Physician, Kinsmen Children's Centre                         | 4     |
| Patricia MacPherson | Montague       | PEI      | Canadian Correctional Services Research Centre                | 1     |
| Pearl Park          | Calgary        | Alta.    | Speech/Language Pathologist, Alberta Children's Hospital      | 1     |
| Peter Waas          | LaCombe        | Alta.    | Psychologist                                                  | 3     |
| Rachelle Deneault   | Whitehorse     | YT       |                                                              | 1     |
| Richard Snyder      | Saskatoon      | Sask.    | Physician, Kinsmen Children's Centre                         | 1     |
| Roxana Vernescu     | St John's      | Nfld.    | Psychologist, Memorial University                             | 1     |
| Samantha            | Ottawa         | Ont.     | First Nations Child and Family Caring Society of Canada       | 1     |
| Nadjiwan            |                |          |                                                              |       |
| Sandy Claren        | Seattle        | WA       | Psychologist, University of Washington                       | 1     |
| Sandy Steinwender   | Iqualuit       | Nun.     | Health and Social Services                                    | 1     |
| Sharon              | Ottawa         | Ont.     | Health Canada                                                 | 2     |
| Bartholomew Soo-Hong Uh | Vancouver   | BC       | Scientist, BC Vital Statistics                                 | 2     |
| Sterling Claren     | Seattle        | WA       | Physician, University of Washington                           | 1     |
| Suzanne Guay        | Ottawa         | Ont.     | National Parole Board                                         | 1     |
| Ted Rosales         | St. John’s     | Nfld.    | Physician, Memorial University                                 | 1     |
| Terry Benoit        | Winnipeg       | Man.     | Physician, Clinic for Drug and Alcohol Exposed Children       | 12    |
| Tim Oberlander      | Vancouver      | BC       | Physician, BC Women’s and Children’s Hospital                 | 1     |
| Val Massey          | Edmonton       | Alta.    | Psychologist, DV Massey and Associates                        | 1     |
| Valerie Flynn       | Ottawa         | Ont.     | Health Canada                                                 | 2     |
| Vyta Senikas        | Ottawa         | Ont.     | Society of Obstetricians and Gynecologists of Canada          | 1     |
| Wendy Sky Delaronde | Kahnawake      | Que.     | Nurse                                                         | 1     |
| Yaya deAndrade      | Vancouver      | BC       | Psychologist, BC Women's and Children's Hospital              | 1     |
| Yeshodara Naidoo    | Ottawa         | Ont.     | Health Canada                                                 | 1     |
Appendix 2: Guides for measurement of palpebral fissure length

Appendix 2-1: Relation between palpebral fissure length and age in both sexes of American white children aged 29 weeks to 14 years.41

Appendix 2-2: Palpebral fissure length for both sexes, birth to 16 years.42
Appendix 3: Examples of tests that are most widely used to assess the domains

* Psychologists, speech-language pathologists and occupational therapists were consulted regarding their widely used tests. Tests for brain function are regularly updated and the most current versions should be used where appropriate.

### Hard and soft neurologic signs (including sensory-motor)

Hard neurologic signs are assessed by the physician according to usual standards. Soft neurologic signs include motor signs that can be elicited on the physical examination, with referral for occupational therapy assessment where appropriate.

**Tests of motor functioning include:**
- Movement Assessment Battery for Children
- Bruininks-Oseretsky Scales of Motor Development
- Alberta Infant Motor Scale
- Peabody Developmental Motor Scales
- Quick Neurological Screening Test-II

**Tests for visual-motor functioning include:**
- Developmental Test of Visual-Motor Integration or Bender Gestalt (simple)
- Rey Complex Figure Test and Recognition Trial (complex)

**Tests of perception include:**
- Gardner Test of Visual Perceptual Skills
- Gardner Test of Auditory Perceptual Skills

**Tests of sensory function include:**
- Dunn Sensory Profile
- University of Washington Sensori-motor Checklist
- Congenital sensory-neural hearing loss as evaluated by audiologist
- Congenital vision anomalies as evaluated by an ophthalmologist

**Tests and observations of articulation, phonology and motor speech if indicated:**
- Goldman-Fristoe –2 Test of Articulation
- Phonological Awareness Test

### Brain structure

Documented measurements of the head circumference (occipitofrontal circumference below the 3rd percentile) adjusted for age and gender (during the physical examination at any age including head circumference at birth) and other evidence of functional or structural CNS dysfunction based on a neurologic examination or findings on imaging techniques (computed tomography scan, magnetic resonance imaging, electroencephalogram). Neurologic problems may include seizures not due to a postnatal insult or other signs such as impaired motor skills, neurosensory hearing loss, memory loss or poor eye–hand coordination.

### Cognition

**Tests of intellectual functioning include:**
- Wechsler Intelligence Scale for Children-III (WISC-IV not yet tested for usefulness with the FASD population)
- Stanford-Binet- Fourth Edition (SB5 not yet tested for usefulness with the FASD population)
- Wechsler Preschool and Primary Scale of Intelligence-III
- Differential Ability Scales
- Bayley Scales of Infant Development

### Communication

Test batteries of language functioning usually combine both receptive and expressive language functions, as well as single-word and complex functions (sentences and paragraphs). Elicited versus recognition ability (multiple-choice) should be distinguished.

- Peabody Picture Vocabulary Test-III
- Expressive Vocabulary Test
- Preschool Language Scale (3 or 4)
- Reynell Developmental Language Scales
- Test of the Auditory Comprehension of Language-3
- Token Test
- Listening Test
- Test of Word Knowledge
- Clinical Evaluation of Language Fundamentals (Preschool, CELF-3, CELF-4)

These measures are complemented by a language sample analysis that includes: length of utterance, use of complex sentences and word retrieval.

### Social Language Observations

- Narrative skill (PLS-E story retell); Renfrew Bus Story, Frog Where are You
- (Note: Language pragmatics are considered in the domain of social/adaptive skills.)
Appendix 3: continued

**Academic achievement**

Tests commonly used include:
- Wechsler Individual Achievement Test-II (most widely used)
- Gray Oral Reading Test
- Woodcock Johnson Achievement Battery
- Wide Range Achievement Test-3 (note: needs to be supplemented by a test that includes reading comprehension)

Note: Avoid relying on group administered achievement test data.

Preschool children present a challenge in this domain; however, concept knowledge as assessed by the Preschool Language Scale, Bracken Test of Basic Concepts and Boehm Basic Concept Scale can be used.

**Memory**

Assessment should include comparisons between visual and auditory memory; short-term memory, delayed recall, and working memory.

Tests commonly used include:
- Children’s Memory Scale-III
- Wechsler Memory Scale-III
- Wide Range Assessment of Memory and Learning
- Rey Complex Figure Test (recall)
- Developmental Neuropsychological Assessment (NEPSY) memory subtests
- Stanford-Binet Fourth Edition memory subtests
- California Verbal Learning Test
- Working memory composites from Wechsler scales

**Executive functioning and abstract reasoning**

- Delis-Kaplan Executive Function System
- Behaviour Rating Inventory of Executive Function (BRIEF): parent and teacher versions
- Verbal Abstract Reasoning and Problem Solving
- Test of Problem Solving (Elementary and Adolescent)
- Semantic Relationships (CELF-3) and Similarities and Differences (LPT-R, TLC-expanded)
- Observation (e.g., answering how and why questions, explanations, inferences)
  (Note: observations made on the IQ test may also apply here)
- Visual Abstract Reasoning and Problem Solving
- Executive function subtests on the NEPSY
- Wisconsin Card Sorting Test

**Attention deficit/hyperactivity**

Tests commonly used include:
- Observation
- Conners’ Rating Scale
- Child Behaviour Checklist
- Continuous Performance Test-2

**Adaptive behaviour, social skills, social communication**

Assessment of social and adaptive skills is considered most important, but the available standardized instruments do not adequately tap the unusual adaptive problems found in FASD.

- Observation and interview, school reports and previous assessments
- Vineland Adaptive Behaviour Scale: often used, but inadequate at higher ages
- Adaptive Behaviour Assessment System: easier to administer and seems to correlate well with other measures and observation
- Informal assessment of language pragmatics (not standardized), social communication