The FASD Eye Code: a complementary diagnostic tool in fetal alcohol spectrum disorders

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ABSTRACT

Objective To create an easy-to-use complementary ophthalmological tool to support a fetal alcohol spectrum disorder (FASD) diagnosis.

Methods and Analysis The FASD Eye Code was derived from 37 children with FASD evaluated along with 65 healthy age-matched and sex-matched controls. Four ophthalmological categories, which are abnormalities commonly found in children with FASD, were ranked independently on a 4-point scale, with 1 reflecting normal finding and 4 a strong presence of an abnormality: visual acuity, refraction, strabismus/binocular function and ocular structural abnormalities. The tool was validated on 33 children with attention deficit/hyperactivity disorder (ADHD), 57 children born moderate- to low-preterm and none had FASD. Children with ADHD were all born prematurely or small for gestational age (SGA) or diagnosed with FASD. Among children born MLP none was SGA, had a diagnosis of ADHD or FASD, or a history of retinopathy of prematurity. Children with SRS were all born SGA, half were born preterm and none had FASD. Children with SRS were re-examined as young adults.

Results An FASD Eye Code cut-off total score of ≥10 showed an area under the curve (AUC) of 0.79 (95% CI 0.69 to 0.87), with 94% specificity and 43% sensitivity, in discriminating between FASD and controls, MLP and ADHD, corresponding to a positive likelihood ratio (LR+) of 7.5. Between FASD and controls, an AUC of 0.87 (CI 0.80 to 0.95), with 100% specificity and 43% sensitivity, was found; between FASD and SRS, an AUC of 0.60 (CI 0.45 to 0.75) was found, with 88% specificity and 43% sensitivity. A cut-off score of ≥9 showed a specificity of 98% and a sensitivity of 57% for FASD versus controls, corresponding to an LR+ of 36.9. Scores in individuals with FASD were compared with FASD diagnostic criteria.

Conclusion The FASD Eye Code has the potential to serve as a complementary tool and help to strengthen an FASD diagnosis.

INTRODUCTION

Since the recognition of fetal alcohol syndrome (FAS), alcohol has been shown to produce a wide spectrum of physical, neurological, cognitive and ophthalmological abnormalities, now known as fetal alcohol spectrum disorders (FASD). The prototypical features of FAS are small palpebral fissures, a thin upper lip, a smooth philtrum, growth restrictions and central nervous system abnormalities (figure 1). Although previous studies have shown that small palpebral fissures are not the only ophthalmological features occurring frequently in children with FASD, the role of ophthalmological assessment in the work-up of FASD may be underestimated.

Alcohol use during pregnancy is a public health problem worldwide. A meta-analysis estimated that over 100 000 children around the world are born with FAS every year. The incorporation of the diagnosis ‘neurodevelopmental disorder – prenatally exposed’ into the Diagnostic and Statistical Manual of Mental Disorders, Fifth Edition is a recognition of the association between alcohol exposure...
Within this wider spectrum of prenatal alcohol exposure (PAE), there is less agreement on what constitutes an alcohol-related aetiological diagnosis, and several sets of diagnostic criteria are used around the world to diagnose FASD. However, the key features of FAS in each set of criteria are the same: growth deficiency, facial dysmorphology and neurobehavioural impairment (figure 1).

Ophthalmological findings, such as subnormal visual acuity, refractive errors, motility disorders, strabismus, and abnormalities of the retinal vessels and optic head, are frequently found in individuals with FASD, but these ophthalmological findings are also found in children with, for example, attention deficit/hyperactivity disorder (ADHD), as well as prematurely born children. On the other hand, both ADHD and prematurity are frequently found in children diagnosed with FASD. To what extent this co-occurrence is caused by confounding alcohol exposure is unknown, and clinically differentiating between alcohol exposure and other factors remains a challenge. Strabismus and refractive errors may cause amblyopia and visual impairment, which are treatable if found in early childhood, but if not treated may present additional problems for already affected individuals. Moreover, other groups with neurodevelopmental syndromes and genetic disorders may also present with similar ophthalmological abnormalities, although other clinical symptoms are usually present as well.

The medical diagnostic process is probabilistic in nature. Arriving at a seemingly dichotomous ‘yes or no’ answer to a diagnosis is guided by a probabilistic weighing of history, findings and tests, where the overarching question is: ‘What is the probability that the patient has this diagnosis, given the information and results?’ In this process, information and tests unspecific to the diagnosis in question help to adjust the probabilities of the diagnosis in a useful way. Analogous to supportive laboratory investigations, a complementary eye diagnostic tool could provide independent verification for clinicians, thus strengthening and helping FASD diagnostics.

To be of use, findings must differentiate between FASD and children with other conditions presenting with similar symptoms. It should also be considered that anthropometric criteria are less evident in adulthood.

**Clinical criteria of Fetal Alcohol Spectrum Disorders (FASD):**
- Central nervous system abnormalities.
- Face abnormalities: small palpebral fissures, smooth, and thin upper lip and philtrum.
- Growth deficits (pre- and/or postnatally).
while cognitive impairment and psychiatric morbidity are more evident in adults with FASD.4 5 26 27

The aims of this study were threefold: first, to develop and propose a complementary and easy-to-use tool based on ophthalmological findings to support an FASD diagnosis; second, to validate the FASD Eye Code’s capacity to discriminate FASD eye findings from prematurity, growth restriction and ADHD without diagnosed FASD; and third, to test the tool in long-term follow-up of individuals with FASD from childhood to young adulthood. In addition, the tool should be useful both in eye clinics and elsewhere, allowing use in outreach and epidemiological surveys of FASD in different communities, while simultaneously identifying treatable deficits requiring management no matter what the cause.

MATERIALS AND METHODS

Developing the FASD Eye Code

Based on the accrued evidence of ophthalmological studies on FASD and our own clinical experience,4 5 9–14 19 four of the most commonly affected ophthalmological features in children with FASD were chosen to create the four categories constituting the FASD Eye Code. The four categories are (A) best corrected visual acuity (BCVA), (B) refraction, (C) strabismus and binocular function, and (D) ocular structural abnormalities. By design, the selected tests are easy to use both in the clinic and on the field (figure 2).

Among the chosen ophthalmological features, two are structural measures (refraction and ocular structural abnormalities) and two are functional tests (BCVA and strabismus/binocular function). The functional measurements represent both afferent and efferent functions. Normal scores in each category are based on known levels in a normal paediatric population,28 29 and the maximal scores are based on affected ophthalmological features well characterised in children with FASD.9–14 30 31 We have excluded palpebral fissure length in the FASD Eye Code since this variable is included in all FASD diagnostic criteria previously mentioned.2 5 16–18

Definitions and scores of the FASD Eye Code

- **BCVA**: visual acuity was tested at 3m with the best possible correction and with a linear Konstantin Moutakis (KM)-Boks chart.31
- **Refraction**: refraction was tested under cycloplegia and was performed with an autorefractor (Topcon A6300/KR8800; Topcon Corporation, Tokyo, Japan) after a single instillation of a mixture of cyclopentolate (0.85%) and phenylephrine (1.5%).
- **Strabismus and binocular function**: strabismus was diagnosed using a cover test and was defined as a
deviation that is manifested always (heterotropia) or intermittently, or as a latent deviation (heterophoria). Binocular function was performed primarily with the Nederlandse Organisatie voor Toegepast Natuurwetenschappelijk Onderzoek (TNO) random dot stereo test. Lang I was used if the child could not participate in the TNO test.

- Ocular structural abnormalities: ptosis, epicanthal folds and ocular fundus were examined clinically, then later by indirect ophthalmoscopy. Fundus photos were taken.

On a 4-point scale, each feature is ranked independently, with ‘1’ reflecting normal ophthalmological findings and ‘4’ reflecting the presence of an abnormality commonly found in individuals with FASD. Thus, ‘4444’ represents the most severe expression of reduced BCVA, significant refractive errors, manifest strabismus or defect binocular functions, and structural abnormalities of the eye. At the opposite end of the scale, code ‘1111’ represents normal ophthalmological findings. The four categories included in the FASD Eye Code and the definitions of the scores (1–4) are summarised in the protocol used for evaluating the FASD Eye Code (online supplemental file 1).

**Study design and participants**

Altogether, the FASD Eye Code was evaluated on 208 different individuals with FASD (n=37), ADHD (n=33), born moderate-to-late preterm (MLP) (n=57) and Silver-Russell syndrome (SRS) (n=16) and in controls (n=65). Children with FASD were drawn from a population-based study of 71 children adopted from Eastern Europe and examined at a mean age of 7.5 years (online supplemental file 2).

- Group 1 (n=37): 15 female, 22 male, mean age 9.8 years (range 4.9–10.5 years), diagnosed with FASD according to the Institute of Medicine criteria,\(^1\) including the following subgroups: FAS (n=21), partial FAS (PFAS) (n=10) and alcohol-related neurodevelopmental disorder (ARND) (n=6).\(^4\)

- Group 2 (n=65): 27 female, 38 male, mean age 9.9 years (range 4.1–12.3), healthy children who served as a control group and were matched by age and sex. Controls for the FASD group were selected from a group of 143 healthy Caucasian school children living in the same area. No child was born small for gestational age (SGA) or prematurely or had a diagnosis of ADHD or FASD.\(^28\)

The FASD Eye Code was then validated on the following additional groups (online supplemental file 2).

- Group 3 (n=33): 12 female, 21 male, mean age 12.1 years (range 6.3–17.5), diagnosed with ADHD. No child was born SGA or prematurely or diagnosed with FASD.\(^24\)

- Group 4 (n=57): 23 female, 34 male, mean age 5.7 years (range 5.4–5.9), born MLP between 32 and 36 weeks’ gestation. No child was born SGA, had a diagnosis of ADHD or FASD, or had a history of retinopathy of prematurity (ROP). These children were selected from a previous study of 78 children, as described in detail elsewhere.\(^24\)

- Group 5 (n=16): 8 female, 8 male, mean age 11.2 years (range 3.4–18.1), with SRS. Half of the children were born preterm and all were born SGA. None was diagnosed with FASD.\(^33\)

Patients or the public were not involved in the design, or conduct, or reporting or dissemination plans of our research.

**The FASD Eye Code: long-term follow-up**

In total, 30 out of the 37 young adults (81%) with FASD (FAS=19, PFAS=6, ARND=5) participated in a follow-up investigation at a mean age of 23 years (range 19–28 years). Six young adults declined to participate and one discontinued the study (figure 3).\(^12\)

![Figure 3](http://bmjophth.bmj.com/) Individual FASD Eye Code total score in 30 subjects diagnosed with FASD, in both childhood and early adulthood. ARND, alcohol-related neurodevelopmental disorder; FAS, fetal alcohol syndrome; FASD, fetal alcohol spectrum disorders; PFAS, partial FAS.
Table 1 The FASD Eye Code median score in the four categories and the total score among the five groups

<table>
<thead>
<tr>
<th>Groups</th>
<th>Median (IQR)</th>
<th>Median (IQR)</th>
<th>Median (IQR)</th>
<th>Median (IQR)</th>
<th>Median (IQR)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group 1: FASD (n=37)</td>
<td>1.0 (2.0)</td>
<td>2.0 (3.0)</td>
<td>3.0 (3.0)</td>
<td>3.0 (2.0)</td>
<td>9 (16–4), 4.5</td>
</tr>
<tr>
<td>FAS (n=21)</td>
<td>2.0 (1.5)</td>
<td>1.0 (3.0)</td>
<td>3.0 (1.5)</td>
<td>3.0 (1.5)</td>
<td>10 (16–4), 5.5</td>
</tr>
<tr>
<td>PFAS (n=10)</td>
<td>1.5 (1.0)</td>
<td>1.5 (1.5)</td>
<td>1.0 (2.25)</td>
<td>2.5 (2.0)</td>
<td>7.0 (11–4), 3.25</td>
</tr>
<tr>
<td>ARND (n=6)</td>
<td>2.0 (1.25)</td>
<td>1.0 (0.25)</td>
<td>2.5 (3.0)</td>
<td>2.0 (2.0)</td>
<td>7.5 (12–4), 6.5</td>
</tr>
<tr>
<td>Group 2: controls (n=65)</td>
<td>1.0 (0.0)</td>
<td>1.0 (0.0)</td>
<td>1.0 (0.0)</td>
<td>1.0 (0.0)</td>
<td>4.0 (9–4), 0.0</td>
</tr>
<tr>
<td>Group 3: ADHD (n=33)</td>
<td>1.0 (0.0)</td>
<td>1.0 (2.0)</td>
<td>1.0 (2.0)</td>
<td>1.0 (2.0)</td>
<td>7.0 (11–4), 4.5</td>
</tr>
<tr>
<td>Group 4: MLP (n=57)</td>
<td>1.0 (1.0)</td>
<td>1.0 (1.0)</td>
<td>1.0 (1.0)</td>
<td>1.0 (0.0)</td>
<td>5.0 (13–4), 2.5</td>
</tr>
<tr>
<td>Group 5: SRS (n=16)</td>
<td>1.0 (1.0)</td>
<td>1.5 (3.0)</td>
<td>1.0 (1.75)</td>
<td>3.0 (2.0)</td>
<td>7.0 (13–4), 3.0</td>
</tr>
</tbody>
</table>

FASD, fetal alcohol spectrum disorders; ADHD, attention deficit/hyperactivity disorder; ARND, alcohol-related neurodevelopmental disorder; BCVA, best corrected visual acuity; FAS, fetal alcohol syndrome; FASD, fetal alcohol spectrum disorders; MLP, moderate-to-late preterm; PFAS, partial FAS; SRS, Silver-Russell syndrome.

A cut-off total score of ≥10 was chosen to represent an FASD diagnosis based on the total sample (this score was obtained by 16 of 37 participants with FASD, 6 of 33 with ADHD, 3 of 57 with MLP, 3 of 16 with SRS and 0 of 65 controls). Thus, an FASD Eye Code cut-off score of ≥10 has 100% specificity and 43% sensitivity, with an area under the curve (AUC) of 0.87 (95% CI 0.80 to 0.95), in discriminating between FASD and healthy controls. When comparing FASD versus controls, ADHD and MLP, the specificity was 94% and the sensitivity was 43% (AUC=0.78; 95% CI 0.69 to 0.87). This result corresponds to a positive likelihood ratio of 7.5. If comparing FASD versus all groups, the AUC was 0.76 (95% CI 0.67 to 0.86). When comparing FASD versus ADHD, the AUC was 0.66 (95% CI 0.53 to 0.78); for FASD versus MLP, the AUC was 0.75 (95% CI 0.64 to 0.86); and for FASD versus SRS, the AUC was 0.60 (95% CI 0.45 to 0.75). The subgroup FAS versus controls showed the highest AUC value of 0.92 (95% CI 0.85 to 1.0), and for a cut-off score of ≥10 the specificity was 100% and the sensitivity was 62%. A cut-off score of ≥9 showed a specificity of 98% and a sensitivity of 57% for FASD versus healthy controls, corresponding to a positive likelihood ratio of 36.9. Tables of diagnostic indices for ROC curves (cut-off, sensitivity, specificity, accuracy, likelihood ratio) are provided in (figure 4A–G and online supplemental file 3).

DISCUSSION

We created and evaluated a new complementary diagnostic tool, the FASD Eye Code, by comparing children with FASD, age-matched and sex-matched healthy controls, and groups of individuals with ADHD, MLP and SRS. Long-term follow-up (re-examination) of individuals with FASD in young adulthood indicated the
persistence of childhood findings. Our results show that the FASD Eye Code can distinguish among different FASD subgroups and discriminate between different patient groups with similar ophthalmological problems. The FASD Eye Code shows better specificity than sensitivity for FASD, with a total score of ≥10 having an increasing likelihood of ruling in an FASD diagnosis. In addition, our results are of clinical importance and require management no matter what the cause is, and the findings may offer useful clues to potential aetiology. To the best of our knowledge, this is the first ophthalmological tool developed to support an FASD diagnosis in both childhood and young adulthood.

The wide range of scores among children with FASD is not surprising as it reflects the range of eye findings associated with PAE during the different stages of development of the fetus, and in some cases eye findings may be due to PAE without rising to a FASD diagnosis. Thus, children who scored ≥10 were most commonly those with fully developed FAS, and no one among the controls scored 10 or above. Among children with SRS, 2 of 16 children scored 11 and 1 scored 13. Unlike the other evaluated groups (ADHD, MLP and controls), all children with SRS showed growth deficits both prenatally and at the time of assessment, which are among the hallmarks of an FASD diagnosis. However, children with SRS have other symptoms and signs, including ophthalmological findings, which indicate an SRS rather than an FASD diagnosis. The same is mostly true when differentiating FASD from other syndromes. However, genetic testing could also be valuable in these cases as a co-occurrence with PAE is possible.

Six children in the ADHD group (children without diagnosed FASD nor born preterm/SGA) were noted to have an FASD Eye Code total score ≥10; three of these children with ADHD had a score of 11 or above. This may be misclassification due to confounding from undiagnosed PAE and undiagnosed FASD, or due to neurodevelopmental afflictions of other aetiology. The criteria for ADHD, a symptom-based diagnosis, were met by the majority of children/young adults with an aetiological diagnosis of FASD.

Visual acuity and refraction are age-dependent, which must be considered when diagnosing different individuals and age groups. No gender differences between the ophthalmological variables used in the code have been
shown in controls. However, refraction and strabismus may have ethnicity-based variation. Thus, our values are suitable for Caucasian children between 4 and 15 years of age. Since the typical facial feature of FAS seems to diminish with age, we tested whether the FASD Eye Code could still be applicable in young adults with FASD. Out of 37 individuals, 30 were assessed in both childhood and young adulthood, showing no significant differences in the FASD Eye Code total score. Since refraction differs in different age groups, myopia was also evaluated in the young adult group, with a cut-off of ≥2.0 dioptre (D) spherical equivalent (SE). However, no difference was noted in the FASD Eye Code total median score when comparing the two different cut-offs (myopia ≥1 D SE vs ≥2 D SE), as individuals with FASD more often have astigmatism or anisometropia, which gives a higher score in the refraction category.

Strengths and limitations

The strength of the study is that the same ophthalmological methods were used in all groups. All the ophthalmological tests necessary for using the FASD Eye Code are well known, inexpensive and can easily be used in low-resource settings outside the clinic. Furthermore, as technology develops, handheld instruments may provide more accurate investigations in the future. To minimise confounding, no children with ROP were included in the study.

Children with ADHD, children born MLP and children with SRS may present with ophthalmological findings similar to those of children with FASD. Since FASD is an aetiologic diagnosis, it warrants consideration in children presenting with prematurity, SGA and ADHD. Even so, the FASD Eye Code gave these groups lower scores—a result that provides support to the use of the code. The same experienced multidisciplinary team investigated all the children in the different groups, with a meticulous and consistent methodology.

Our study has some limitations: the small size of the groups, the fact that the ophthalmological findings in isolation are unspecific and that the eye code was derived from a group of adopted children from Eastern Europe with FASD. In addition, other confounding factors may have had an impact on the results which must be addressed when planning future studies.

In 35 years working with children exposed to alcohol, we have learnt that PAE is under-recognised and the dosimetry complicated. Thus, our control group may include individuals exposed to alcohol without our knowledge. Irrespective of the FASD diagnosis, this tool may identify treatable eye problems. Refractive errors are treatable with glasses, and if not treated in childhood may affect visual acuity for life; an untreated strabismus can also result in amblyopia and eye strain. The code needs to be independently validated by other assessors examining other FASD cohorts and comparison groups. Further studies are needed to validate the FASD Eye Code longitudinally—that is, within the same individuals with FASD and in other age groups with different ethnicities, as well as in syndromes of other aetiology and in healthy controls with a wider age range. We recommend using the code as a complement when there is suspicion for an FASD diagnosis and when genetic syndrome phenocopies have been ruled out.

In conclusion, in this derivation cohort, an FASD Eye Code total score ≥10 significantly corroborates an FASD diagnosis, although low scores cannot rule out FASD. The FASD Eye Code may support the criteria for diagnosing individuals with suspected FASD and help identify children and young adults with treatable ophthalmological problems. In addition, all the ophthalmological tests involved in the FASD Eye Code are well known, relatively inexpensive and can easily be used outside the clinic. In the future, as technology develops, handheld instruments such as different refractive and imaging devices may provide more accurate investigations. Ophthalmological assessment should be a routine part of the FASD work-up and the FASD Eye Code can help to highlight a need for intervention.

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Acknowledgements We thank Daniel Skorczynski, a biostatistician of Statistiska Konsultgruppen, Gothenburg, and Salmir Nasic, Skaraborg Hospital, Sweden, for statistical analyses.

Contributors EA, EG, VL, LS, ML and MAG were responsible for the concept and design of the study, data interpretation and manuscript drafting and revision, and have approved the final manuscript as submitted. All authors have full access to all study data. EA had the final responsibility for the decision to submit the publication and is responsible for the overall content as the guarantor. All authors attest that they meet the current ICMJE criteria for authorship.

Funding MAG, EA, ML and LS were supported by grants from Gothenburg Medical Society, the W & M Lundgren Vetenskapsfond I, the Swedish Research Society, Research and Development of Region Västra Götaland and the Swedish State under the agreement between the Swedish government and the country councils, the ALF agreement (grants no ALFGBG-11626, ALFGBG-211671, ALFGBG-445021, ALFGBG 509761, ALFGBG 672501 and ALFGBG 71933), EG is funded by grants from research funds at Skaraborg Hospital, Skövde, Sweden (VGSKAS 598651, VGSKAS 688941 and VGSKAS 793611). MAG has received lecture fees from Bayer, EG, VL, EA and ML have no financial disclosures.

Competing interests None declared.

Patient consent for publication Not required.

Ethics approval The study was approved by the Research Ethics Board at the University of Gothenburg, Sweden (no 704-11, 311-99, 043-01, 335-11, 0321-03, 0 125-00; T029-02), and conducted according to the Declaration of Helsinki. The children and their parents were informed of the nature of the study, and written informed consent was then obtained from all the parents and young adults.

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement Data are available upon reasonable request. All data relevant to the study are included in the article or uploaded as supplementary information.
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