Refractive development in children with Down’s syndrome: a population based, longitudinal study

Olav H Haugen, Gunnar Høvding, Isa Lundström

Abstract

Aims—To study the refractive development in children with Down’s syndrome longitudinally.

Methods—An unselected population of 60 children with Down’s syndrome was followed with repeated retinoscopies in cycloplegia for 2 years or more (follow up 55 (SD 23) months). Accommodation was assessed with dynamic retinoscopy.

Results—From longitudinal spherical equivalent values of the right eye, three main categories of refraction were defined: stable hypermetropia (≤1.5 D difference between the first and last visit) (n=34), increasing hypermetropia (“hypermetropic shift”; ≥1.5 D difference) (n=11), and decreasing hypermetropia/development of myopia (“myopic shift”; ≥1.5 D difference) (n=9). Patients with anisometropia (n=6) were evaluated separately. In the stable hypermetropia group three sublevels were chosen: low (≤+2.0 D at the last visit), moderate (+2.25 to + 4.0 D), and high (>+4.0 D). An accommodation weakness was found in 55% of the children. Accommodation weakness was significantly less frequent in the stable, low grade hypermetropia group (22%) than in all the other groups (p=0.008). The frequency of astigmatism ≥1.0 D at the last visit was 57%, the direction of axis being predominantly “with the rule.” All the eyes with oblique astigmatism had a side specific direction of axis; the right eyes belonging to the 135° axis group and the left eyes to the 45° axis group.

Conclusion—A stable, low grade hypermetropia was significantly correlated with a normal accommodation. Accommodation weakness may be of actiological importance to the high frequency of refractive errors encountered in patients with Down’s syndrome. A striking right-left specificity in the oblique astigmatic eyes suggests that mechanical factors on the cornea from the upward slanting palpebral fissures may be a major actiological factor in the astigmatism.

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The increased frequency of refractive errors in individuals with Down’s syndrome has been documented by many authors. However, most studies have been based on selected populations. In addition, all reports so far have been cross sectional studies.

Hoping to elucidate new aspects of this issue, we have studied the refractive development in an unselected Down’s syndrome childhood population by repeated examinations during the past 10 years. To our knowledge, this is the first longitudinal study on refractive errors in children with Down’s syndrome.

Subjects and methods

STUDY POPULATION

Seventy seven children with Down’s syndrome born during the years 1988–99 in Hordaland County, Norway (population 416 000, annual births 6000), were referred from Vestlund Habilitation Resource Center to our department for an ophthalmological examination. This centre coordinates the habilitation of all Down’s syndrome children in the county. To ensure a population based study design, the files of the regional laboratory for cytogenetics were examined. In this way we found 16 dropouts. Thus, the total number of patients with Down’s syndrome born in our county during these years was 93, giving a mean annual incidence of 1.25 per 1000 live births (range 0.63–2.12).

Among the 16 dropouts, four had died and three had moved to other parts of the country. The other nine children were invited for an eye examination, which six of them attended. Valid data on previous refraction by other ophthalmologists could be obtained, and they were included in the study. Another six children moved to our region during the study period and were included in the study with successive examinations. Thus, 89 children with Down’s syndrome born in the years 1988–99 were examined. Among these, 40 had their first examination during the first year of life (mean age at examination 7.1 (SD 3.0) months, range 3–12). Cross sectional data from these 40 infants will be presented separately.

Patients with follow up time <2 years (n=29) were excluded from the longitudinal study. The group of children with Down’s syndrome thus followed for ≥2 years with repeated eye examinations consisted of 60 children (30 girls and 30 boys). Mean follow up time was 55 (SD 23) months (range 24–115). With very few exceptions, all the examinations were done by one of the authors (OHH). In the whole longitudinal study group the mean age at the first examination was 21 (SD 14) months (range 3–61).

Informed consent was obtained from the parents, and the study was approved by the regional committee for medical research ethics.
REFRACTIVE MEASUREMENTS

Retinoscopy in cycloplegia was performed using cyclopentolate 1% eye drops twice 30 minutes before the examination. Astigmatism was recorded as minus cylinders. The axis of astigmatism was classified as follows: 180° (SD 15°) (“180° meridian” or “with the rule”), 90° (SD 15°) (“90° meridian” or “against the rule”), 16–74° (“45° meridian”), and 106–164° (“135° meridian”).

To evaluate the axis of oblique astigmatism in a mentally normal population, we used the preoperative refractive data from the excimer laser clinic in our department. Only patients referred for primary, uncomplicated refractive errors were included. Eyes with cylindrical power of <1.0 D were excluded. Thus, our control group included 365 eyes (172 right eyes, 193 left eyes) with astigmatism >1.0 D.

ACCOMMODATION

During the past 2–3 years of the study, each examination included an evaluation of the preoperative refractive data from the excimer laser clinic in our department. Only patients referred for primary, uncomplicated refractive errors were included. Eyes with cylindrical power of <1.0 D were excluded. Thus, our control group included 365 eyes (172 right eyes, 193 left eyes) with astigmatism ≥1.0 D.

Table 1 Longitudinal refractive development in 60 children with Down’s syndrome

<table>
<thead>
<tr>
<th>Group</th>
<th>Definition</th>
<th>Number of patients</th>
</tr>
</thead>
<tbody>
<tr>
<td>Isometropia</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Group 1</td>
<td>Stable hypermetropia (spherical equivalent right eye &lt;1.5 D difference between the first and last measurements)</td>
<td>34</td>
</tr>
<tr>
<td></td>
<td>(la) low grade hypermetropia (spherical equivalent right eye &lt;+2.0 at the last examination)</td>
<td>18</td>
</tr>
<tr>
<td></td>
<td>(lb) moderate hypermetropia (spherical equivalent right eye +2.25–+4.0 D at the last examination)</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>(lc) high grade hypermetropia (spherical equivalent right eye &gt;+4.0 D at the last examination)</td>
<td>4</td>
</tr>
<tr>
<td>Group 2</td>
<td>Increasing hypermetropia (spherical equivalent right eye ≥1.5 D difference between the first and last measurements)</td>
<td>11</td>
</tr>
<tr>
<td>Group 3</td>
<td>Decreasing hypermetropia (spherical equivalent right eye ≥1.5 D difference between the first and last measurements)</td>
<td>9</td>
</tr>
<tr>
<td></td>
<td>or development of myopia (spherical equivalent right eye &lt;0 at the last examination)</td>
<td></td>
</tr>
<tr>
<td>Anisometropia</td>
<td>Difference between right and left eye at the last examination &gt;1.0 D spherical and/or &gt;1.5 D cylinder power</td>
<td>6</td>
</tr>
<tr>
<td>Total</td>
<td></td>
<td>60</td>
</tr>
</tbody>
</table>

Figure 1 Individual curves of spherical equivalent values from children with Down’s syndrome and stable, low grade hypermetropia (A), stable, moderate hypermetropia (B), stable, high grade hypermetropia (C), increasing hypermetropia (D), and decreasing hypermetropia or development of myopia (E).
accommodative function with dynamic retinoscopy. Several authors have found dynamic retinoscopy valuable in assessing the accommodative function, especially in children and in the mentally retarded.10–13 In most of these studies, the fixation target is mounted on the retinoscope, and the observer looks for the “breakdown” of the neutral retinoscopic reflex while constantly moving closer to the child. Because many of our Down’s children became scared as the examiner moved close to them, the technique was modified as follows. In case of myopia, or hypermetropia > +2.0 D, the child should wear his/her distant glass correction. Sitting about 50 cm in front of the child, the examiner observes the retinoscopic streak light movement while the child is looking straight ahead with both eyes open. A small picture that attracts interest (Lang’s cube) is then introduced 20–30 cm in front of the child. The child is constantly encouraged to fixate the near target. If normal accommodation is present, the examiner observes a very distinct shift from “with” movements to “against” movements. Such a response was classified as “normal.” If, when presenting the accommodative target, this clear shift did not take place in spite of a cooperative child, the accommodation response was classified as “accommodation weakness.” The test was repeated three or more times.

VISUAL ACUITY

In infants and small children the visual acuity was tested with the Teller acuity card test. In older children we used an optotype test, mostly the Østerberg chart or the LH chart. In the most cooperative children, ordinary Snellen optotypes were used. Amblyopia was defined as a difference in visual acuity between the two eyes of more than one line on the acuity chart.

STATISTICS

The data were analysed statistically in the SPSS 9.0 program, using the χ² test and the Fisher’s exact test. p = 0.05 was chosen as the level of significance.

Results

CROSS SECTIONAL DATA IN INFANTS

Twenty one (53%) of the infants were emmetropic or hypermetropic <=+2.0 D, while 16 (40%) were hypermetropic between +2.0 and +5.25 D. There were three myopic children, all within –1.5 D. Clinically significant astigmatism (>1.0 D) was present in 21 of 40 infants (53%). In all but one case the astigmatism was bilateral. In one patient, the axis was “against the rule” bilaterally, in all the other cases (95%) there was a “with the rule” astigmatism.

LONGITUDINAL STUDY

Six patients with clinically significant anisometropia at the last examination were regarded as one group. According to the development of the spherical equivalent of the right eye, the other 54 patients were classified into three main groups, presented in Table 1: group 1 stable hypermetropia (at different levels (n=34); group 2 increasing hypermetropia (n=11), and group 3 decreasing hypermetropia or development of myopia (n=9) (see Table 1 for the detailed definitions). The individual
Table 3 Accommodative ability in different refractive groups among 60 children with Down’s syndrome

<table>
<thead>
<tr>
<th>Refractive group</th>
<th>Accommodation</th>
<th>Normal</th>
<th>Weak</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Isometropia</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Group 1a: stable, low grade hypermetropia</td>
<td></td>
<td>14 (78%)</td>
<td>4 (22%)</td>
<td>18</td>
</tr>
<tr>
<td>Group 1b: stable, moderate hypermetropia</td>
<td></td>
<td>6 (50%)</td>
<td>6 (50%)</td>
<td>12</td>
</tr>
<tr>
<td>Group 1c: stable, high grade hypermetropia</td>
<td></td>
<td>0 (0%)</td>
<td>4 (50%)</td>
<td>11</td>
</tr>
<tr>
<td>Group 2: increasing hypermetropia</td>
<td></td>
<td>4 (36%)</td>
<td>7 (64%)</td>
<td>11</td>
</tr>
<tr>
<td>Group 3: decreasing hypermetropia/development of myopia</td>
<td></td>
<td>2 (22%)</td>
<td>7 (78%)</td>
<td>9</td>
</tr>
<tr>
<td>Anisometropia</td>
<td></td>
<td>1 (17%)</td>
<td>5 (83%)</td>
<td>6</td>
</tr>
<tr>
<td>Total</td>
<td></td>
<td>27 (45%)</td>
<td>33 (55%)</td>
<td>60 (100%)</td>
</tr>
</tbody>
</table>

The study period (upper part of Table 2). Three patients with “against the rule” astigmatism also had stable axes (middle part of Table 2). The lower part of Table 2 shows data from the 11 patients in group C with an oblique astigmatism at some point of the study period. All eyes that changed axis during the follow up period (n=8) belonged to this subgroup (in bold). At the last examination, nine patients had an oblique astigmatism in one or both eyes. All the right eyes with oblique astigmatism had their axes in the 135° meridian, while all the left eyes had their axes in the 45° meridian. Figure 2 shows the axes of all the 12 patients with oblique astigmatism at any point of the study.

In the control group of patients referred to our excimer laser clinic, the right and left eyes with oblique astigmatism were equally distributed to the 45° axis and the 135° axis.

ACCOMMODATION
Accommodation weakness was found in 33 patients (55%) (Table 3). However, the frequency was lower in the stable, low grade hypermetropic group than in the other groups ($\chi^2 = 14.7; \text{df}=5; p = 0.008$). Among those tested at two or more visits (n=40), six changed from “normal” to “weak,” while one child changed from “weak” to “normal.”

VISUAL ACUITY
Visual acuity could be evaluated in 53 children (88%) in the longitudinal study group (Table 4). In 27 children visual acuity could be recorded in each eye separately, while in 26 cases only binocular testing was possible. Only one of the children tested monocularly had a difference between the two eyes of more than one line on the acuity chart.

Discussion
DEVELOPMENT OF SPHERICAL EQUIVALENT
This study shows that about one third of the children with Down’s syndrome (group 1a) had stable refractive values around emmetropia or low hypermetropia throughout preschool and early school age. Stable, but higher, values of hypermetropia were demonstrated in about one fourth of the children. Of special interest are the “hypermetropic shift” and “myopic shift” groups, leading to a persistent wide distribution of refractive values. In a normal refractive development, a low to medium grade hypermetropia with a wide
OBL = oblique astigmatism.
ATR = "against the rule."
WTR = "with the rule."

*Astigmatism \( \geq 1.0 \text{ D} \) was one of the inclusion criteria in this study.
WTR = "with the rule."
ATR = "against the rule."
OBL = oblique astigmatism.

The prevalence of astigmatism in our Down’s syndrome infant group (53%) was comparable with that of normal infants (Table 5). However, it was considerably higher than that reported in Down’s syndrome infants by Woodhouse et al. The present study strongly indicates that "with the rule" astigmatism is the predominant type of astigmatism in infants with Down’s syndrome. This contrasts with the findings in normal infants, where "against the rule" astigmatism seems to be the most common type (Table 5).

At the last examination, 57% of the whole longitudinal study group had a clinically significant astigmatism. This frequency (55%) of accommodation weakness among children with Down’s syndrome reported by others is also supported by clinical experience and reported by others. Interestingly, our data demonstrate that the stable, low grade hypermetropia group (corresponding to a normal refractive development) had a significantly lower frequency of accommodation weakness than the other refractive groups. This association between an accommodation weakness and a failing emmetropisation does not prove a causative relation. However, several animal studies have shown that optical defocus on the retina induces compensating eye growth. This supports the view that reduced accommodation in early age, causing a blurred retinal image for objects at near, may be of aetiological importance for the abnormal refractive development in children with Down’s syndrome. Obviously, there must be additional factors, as a reduced accommodation would always shift the optical focus behind the retina and thus induce a myopic shift. This mechanism, therefore, does not explain the cases with increasing hypermetropia.

In addition to a high proportion of accommodation weakness in our material, six of the 40 patients who were repeatedly examined for accommodation showed a change from normal to defect accommodative function. This could indicate that an accommodative ability initially present might later be weakened or lost.

The underlying mechanisms accounting for the reduced accommodation remain unclear. Such factors may be central or peripheral in origin.

**Table 5** Astigmatism in infants: comparison between the present and previous studies

<table>
<thead>
<tr>
<th>Author</th>
<th>Year</th>
<th>Method of refraction</th>
<th>No of patients (age)</th>
<th>Astigmatism ( \geq 1.0 \text{ D} )</th>
<th>Axis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal infants</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mohindra et al*</td>
<td>1978</td>
<td>Non-cycloplegic</td>
<td>276 (0–50 weeks)</td>
<td>45%</td>
<td>WTR</td>
</tr>
<tr>
<td>Ingram et al*</td>
<td>1979</td>
<td>Cycloplegic</td>
<td>148 (1 year)</td>
<td>30%</td>
<td>ATR</td>
</tr>
<tr>
<td>Fulton et al*</td>
<td>1980</td>
<td>Cycloplegic</td>
<td>75 (0–60 weeks)</td>
<td>19%</td>
<td>OBL</td>
</tr>
<tr>
<td>Gwiazda et al*</td>
<td>1984</td>
<td>Cycloplegic</td>
<td>521 (0–11 months)</td>
<td>52%</td>
<td></td>
</tr>
<tr>
<td>Dobson et al*</td>
<td>1984</td>
<td>Cycloplegic</td>
<td>43 (0–18 months)</td>
<td>20%</td>
<td></td>
</tr>
<tr>
<td>Abrahamsson et al*</td>
<td>1988</td>
<td>Cycloplegic</td>
<td>299 (1 year)</td>
<td>100%*</td>
<td></td>
</tr>
<tr>
<td>Ehrlich et al*</td>
<td>1997</td>
<td>Cycloplegic</td>
<td>254 (9 months)</td>
<td>35%</td>
<td></td>
</tr>
<tr>
<td>Down syndrome infants</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Woodhouse et al*</td>
<td>1997</td>
<td>Non-cycloplegic</td>
<td>23 (3–12 months)</td>
<td>26%</td>
<td></td>
</tr>
<tr>
<td>Present study</td>
<td>2001</td>
<td>Cycloplegic</td>
<td>40 (3–12 months)</td>
<td>53%</td>
<td></td>
</tr>
</tbody>
</table>

*Inclusion criteria: \( \geq 1.0 \text{ D} \).

The left eyes were in the 45° meridian, while all the oblique astigmatism in the left eyes was in the 45° meridian. To our knowledge, this right-left specificity of oblique astigmatism in Down’s syndrome has not previously been pointed out or commented on. The astigmatic axes are usually only reported as "with the rule," "against the rule," or oblique. We have only found one publication reporting the axes separately for the right and the left eyes in patients with Down’s syndrome. In this report, 14 out of 15 eyes with oblique astigmatism showed the same right-left specific pattern as in our study.
We have had the opportunity to review the refractive data on 50 teenagers (100 eyes) with Down’s syndrome reported by Doyle et al. Fifty-two eyes had cylindrical power ≥ 1.0 D. Among these, 21 had an oblique axis according to our definition. The direction of the axes showed the same right-left specificity as described above in 18 of these 21 eyes (86%). These data on the right-left specific axes were not reported in the published article.

In normal adults, McKendrick and Brennan found that the right and the left eyes with oblique astigmatism were equally distributed to the 135° and the 45° meridian. This finding corresponds to the results of our preoperative measurements on the excimer laser patients.

We suggest that this right-left specific direction of oblique astigmatism may be caused by the upward slanting of the palpebral fissure, first described in the original publication by Down. Not specifically for Down’s syndrome, but as a general hypothesis, pressure from the eyelids has already been pointed out as a major aetiological factor of corneal astigmatism. Eyelid pressure may cause an inward rotation of the globe and an anterior displacement of the lens axis, resulting in an oblique astigmatism.

Shapiro and France found that the angle of the palpebral fissure in Down’s syndrome patients showed the same right-left specificity as described above. In addition, Lowe reported a widening of the angle between the two anterior-posterior orbital axes from the normal 45° to 75° in four skulls from Down’s syndrome patients. Both these factors might contribute to the right-left specificity in oblique astigmatism in Down’s syndrome.

In conclusion, our longitudinal study confirms an abnormal refractive development in children with Down’s syndrome. A possible causative association to a poor accommodation needs to be further explored. A high frequency of accommodation weakness in children with Down’s syndrome suggests a more liberal use of bifocal or progressive glasses than practised today. At present, however, it must be emphasised that although improving visual performance at near, it is uncertain whether the use of progressive or bifocal glasses will prevent an unfavourable refractive development in children with Down’s syndrome.

We are grateful to Dr Stephen J Doyle and co-workers at Manchester Royal Eye Hospital, who put their data on teenagers with Down’s syndrome at our disposal. The Medical Birth Registry of Norway (MBR) and the Laboratory for Cytogenetics, Center for Medical Genetics and Molecular Medicine, Haukeland University Hospital, Bergen, are gratefully acknowledged for their efforts to provide necessary information. We thank Birgitte Eisehaug for helpful work with the statistical analyses.

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