Long-term developmental follow-up in children with nonsyndromic craniosynostosis

Maggie Bellew, PhD,1 and Paul Chumas, FRCS(SN)2

Departments of 1Plastic, Reconstructive, and Hand Surgery and 2Neurosurgery, Leeds General Infirmary, Leeds, United Kingdom

OBJECT The aim of this study was to determine the distribution of Full Scale IQ (FSIQ) by type of craniosynostosis and to verify the finding that at long-term follow-up, verbal IQ (VIQ) is significantly higher than performance IQ (PIQ) in patients with single-suture sagittal synostosis (SS) despite falling within the “average” range for intelligence. Whether this also occurs in other types of craniosynostosis and whether surgery and sex are relevant were also determined. The relationship between age at time of surgery and later IQ was ascertained.

METHODS The data for 91 children with craniosynostosis (47 sagittal, 15 unicoronal, 13 metopic, 9 multisuture, and 7 bicoronal) were collected at their routine, 10 years of age IQ assessment (mean age 123.8 months). The patients included 61 males and 30 females; 62 patients had undergone surgery and 29 had not.

RESULTS The mean FSIQ for all types of craniosynostosis combined (96.2) fell within the average range for the general population. Some variation was evident across the different types of craniosynostosis: the SS group showed the highest FSIQs and a “normal” distribution of bandings; the other types had a higher proportion of FSIQs in the lower bandings. The data confirmed the finding that VIQ is greater than PIQ despite falling within the average range for intelligence, with a difference of 5.0 for all types of craniosynostosis combined (p = 0.001), 7.6 for the SS group (p = 0.001), and 6.9 for the unicoronal group (p = 0.029). This VIQ > PIQ effect was not found with multisuture craniosynostosis. The VIQ > PIQ discrepancy occurred regardless of whether the patient had undergone surgery and occurred more often in males than females. In the SS group and the bicoronal group, FSIQ (p = 0.036 and p = 0.017, respectively) and PIQ (p = 0.012 and p = 0.017, respectively), though not VIQ, were higher when surgery had been performed early.

CONCLUSIONS The study confirms that at long-term follow-up, although children with nonsyndromic craniosynostosis fall within the normal range for intelligence, there is a VIQ > PIQ discrepancy above what would be expected in the normal population, which may be indicative of more subtle difficulties in achievement. This discrepancy is affected by type of craniosynostosis, sex, and age at time of surgery.

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KEY WORDS craniosynostosis; long-term follow up; developmental attainment; IQ; craniofacial surgery; children

In 2002, Magge et al.16 found that with a small-sample study of 16 patients of varied ages (6.4–15.9 years, mean 10.3 years) with nonsyndromic single-suture sagittal synostosis (SS), verbal IQ (VIQ) was statistically significantly higher (p = 0.021) than performance IQ (PIQ), as measured by the Wechsler Intelligence Scale for Children 3rd edition (WISC-III).25 This was despite falling within the “average” range for intelligence. Whether this also occurs in other types of craniosynostosis and whether surgery and sex are relevant were also determined. The roles of surgery and sex were also examined. There is a paucity of information on long-term follow-up in craniosynostosis, which this study goes some way to address.

Methods

The Leeds (East) Research Ethics Committee confirmed that “ethics committee approval is not needed for the collation and analysis of patient data collected over a period of time on a routine basis during normal clinical care, on the understanding that the data is pooled and therefore nonidentifiable.”

All children attending the Leeds Regional craniofacial center are invited for regular developmental review.
according to a standardized protocol: preoperatively; 6 months postoperatively if receiving surgery (or at a similar interval if not receiving surgery); and at 4, 10, and 15 years of age. Additional assessments may be conducted according to clinical need.

The data from 91 consecutive patients with craniosynostosis who had attended their routine 10 years of age developmental assessment at the Leeds Craniofacial Unit were collated, from the first patient to reach 10 years of age (February 2003) until January 2014. One patient underwent early assessment, at 8.02 years, due to clinical requirements, and 4 patients attended just prior to their tenth birthdays. Eight patients attended at 11 years of age due to difficulties in attending appointments. A further 81 patients did not attend their 10-year review or had been lost to follow-up (SS 46.9%, metopic 24.6%, unicoronal 11.1%, bicoronal 3.7%, multi/other 13.6%).

All the assessments were carried out by clinical psychologists who were blinded to the VIQ > PIQ hypothesis and had not read the Magge et al.16 paper. Table 1 presents the number of patients with each type of craniosynostosis along with their sexes and their mean age at the time of assessment. The table also shows whether patients had undergone surgery or not and their ages at the time of surgery. The mothers of 2 children (1 with metopic craniosynostosis, 1 with SS) had taken sodium valproate during pregnancy.

Plain radiographs were obtained to confirm the clinical diagnosis of craniosynostosis. CT scanning or MRI were also performed in cases of metopic synostosis (to look for associated brain malformations) or in more complex multisuture cases. Children who did not undergo surgery usually had cosmetically mild craniosynostosis. Those patients with more marked disease whose parents did not wish them to undergo surgery were closely observed clinically (ophthalmic and developmental review and serial head circumference measurements), with a low threshold for intracranial pressure (ICP) monitoring if there was any concern. Patients under 6 months of age with SS underwent a modified Renier’s “H” procedure.8 Older patients (> 6 months and usually around 1 year of age) with significant occipital bullets or frontal bossing had these aspects corrected in addition to undergoing the basic Renier technique. Patients with metopic, unicoronal, or bicoronal types were treated by frontal advancement (using either a Marchac frontal template or reversal of the frontal bone).

All the patients were referred for fundus examination by an expert pediatric ophthalmologist.

All patients had been assessed using the WISC 4th edition, UK edition (WISC-IV UK) or the shorter version of this scale, The Wechsler Abbreviated Scale of Intelligence (WASI),24 which correlates well with the WISC-IV UK. Both are individually administered, standardized measures of intelligence, and they yield Full Scale IQ (FSIQ), verbal IQ (VIQ) and performance IQ (PIQ). The FSIQ is the overall estimate of an individual’s general level of intellectual functioning. The VIQ is a measure of acquired knowledge, verbal reasoning, and attention to verbal information. The PIQ provides an indication of nonverbal skills and is a measure of fluid reasoning, spatial processing, attentiveness to detail, and visual-motor integration.

Statistical analyses were conducted using SPSS version 19.0 for Windows (IBM). Independent sample t-tests were undertaken to compare FSIQs between different types of craniosynostosis, and the distribution of FSIQs within types was also examined. Paired t-tests were performed to compare VIQ with PIQ. This was undertaken for all types of craniosynostosis combined, and then for each type separately. The data were also analyzed separately for surgical and nonsurgical patients and for male and female patients. One-way ANOVAs were used to examine whether age at time of surgery was relevant.

### Results

**FSIQ Data**

Table 2 shows that for all types of craniosynostosis combined, the mean FSIQ (96.2) fell within the average range for the child population. There was some variation across the different types of craniosynostosis, with the SS group showing the highest mean FSIQ for simple craniosynostosis (100.5). The other types all had significantly lower mean FSIQs than the SS group, falling within the “low average” range of ability: unicoronal (p = 0.036), metopic (p = 0.043), and bicoronal (p = 0.039).

Table 3 shows that the distribution of FSIQs varied according to type of craniosynostosis, with the SS group having the most “normal” distribution and most similar to the published WASI norms. The other types had a higher proportion of FSIQs in the “borderline learning difficulties” or “learning difficulties” range of ability compared

<table>
<thead>
<tr>
<th>Type of Craniosynostosis</th>
<th>No. of Pts</th>
<th>M, F</th>
<th>Surgery</th>
<th>No Surgery</th>
<th>Age at Time of Surgery*</th>
<th>Age at Time of Assessment*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>&lt;7 7–18 &gt;19 Mean (SD) Range</td>
<td>Mean (SD) Range</td>
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<tr>
<td>Sagittal</td>
<td>47</td>
<td>39, 8</td>
<td>33</td>
<td>14</td>
<td>14 14 5</td>
<td>123.1 (6.57) 96.2–138.6</td>
</tr>
<tr>
<td>Unicoronal</td>
<td>15</td>
<td>5, 10</td>
<td>11</td>
<td>4</td>
<td>1 9 1</td>
<td>124.2 (3.81) 120.6–132.0</td>
</tr>
<tr>
<td>Metopic</td>
<td>13</td>
<td>9, 4</td>
<td>5</td>
<td>8</td>
<td>1 4 0</td>
<td>124.3 (5.79) 120.1–142.0</td>
</tr>
<tr>
<td>Bicoronal</td>
<td>7</td>
<td>4, 3</td>
<td>6</td>
<td>1</td>
<td>0 5 1</td>
<td>127.3 (7.36) 122.2–142.5</td>
</tr>
<tr>
<td>Multisuture</td>
<td>9</td>
<td>4, 5</td>
<td>7</td>
<td>2</td>
<td>1 4 2</td>
<td>122.8 (2.63) 120.2–128.4</td>
</tr>
<tr>
<td>Total</td>
<td>91</td>
<td>61, 30</td>
<td>62</td>
<td>29</td>
<td>17 36 9</td>
<td>123.8 (5.87) 96.2–142.5</td>
</tr>
</tbody>
</table>

* Pts = patients.

* Values are months.

Table 1. Type of craniosynostosis, sex, age at time of surgery, and age at time of developmental assessment

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with the SS group, with very few children scoring in the higher bandings.

There was no significant difference when male patients were compared with female patients in terms of FSIQ data, either for all types of craniosynostosis combined or for each type separately.

**VIQ and PIQ Data**

**Impact of Diagnosis**

The data also confirm Magge et al.'s VIQ > PIQ finding for SS craniosynostosis (Table 2). In our study, there was a difference of 5.0 for all types of craniosynostosis combined (p = 0.001), 7.6 for the SS group (p = 0.001), and 6.9 for the unicoronal group (p = 0.029).

VIQ ≥ PIQ only appeared in the single-suture groups. There was a trend toward it being the other way round (i.e., PIQ > VIQ) in the bicoronal and combined multisuture synostosis groups.

**Impact of Surgery**

Further analysis suggested that a statistically significant effect only occurred in surgical patients (p = 0.001 for all types of craniosynostosis combined, p = 0.002 for the SS group, p = 0.026 for the unicoronal group). This may, of course, reflect condition severity rather than whether surgery had taken place, surgery being less likely to be undertaken in milder cases. However, VIQ > PIQ was found in 22 (66.7%) of the 33 cases in the SS surgical group and in 10 (71.4%) of the 14 cases in the SS nonsurgical group. The lack of statistical significance may, therefore, just reflect small sample size.

**Impact of Sex**

Table 4 shows that VIQ > PIQ only seemed to occur within groups of male patients, with a difference of 6.5 for all types of craniosynostosis combined (p = 0.0001) and a difference of 7.9 for the SS group male patients (p = 0.001). While 72% of the male SS patients demonstrated VIQ > PIQ, only 50% of female SS patients showed this IQ discrepancy (68.1% for male and female combined). There were no significant effects for the female groups (though there were no significant differences).

When male patients were compared directly with female patients in terms of their actual VIQ or PIQ (rather than the difference between VIQ and PIQ), either for all types of craniosynostosis combined or for each type separately, there were no significant differences.

**Age at Time of Surgery**

The data were also examined to establish whether age at time of surgery was relevant. This was first determined for all types of craniosynostosis combined. In terms of FSIQ, PIQ, and VIQ, 1-way ANOVAs comparing surgery...
performed at ≤ 6 months of age with that performed at 7–18 months of age and ≥ 19 months showed no significant results.

There were, however, significant results when patient FSIQs were examined by type of craniosynostosis. One-way ANOVAs showed a significant difference in FSIQ for SS patients (p = 0.036), with the highest mean FSIQ being obtained when surgery was performed at ≤ 6 months of age (FSIQ 108.7), followed by surgery at 7–18 months (FSIQ 95.2), and lowest when surgery was performed at ≥ 19 months (FSIQ 88.6). There were similar results for patients with bicoronal synostosis (p = 0.046), with those patients undergoing surgery at 7–18 months obtaining a higher mean FSIQ (93.2) compared with those undergoing surgery at ≥ 19 months (FSIQ 57.0). The number of patients with bicoronal synostosis was, however, very small, and no one in this group had undergone surgery at ≤ 6 months of age.

One-way ANOVAs for PIQ across the surgery age groups of ≤ 6, 7–18, and ≥ 19 months were very similar, with a significant result for the SS group (p = 0.012) (PIQs of 104.2, 90.1, and 84.8, respectively) and for the bicoronal group (p = 0.017) (PIQs of 96.2 for surgery at 7–18 months and 56.0 for surgery at ≥ 19 months). Again, there were no significant results for VPIQ.

Discussion

Long-Term Developmental Outcome

Some previous studies\(^{13,14,22,23}\) have suggested that there may be long-term developmental and educational difficulties in children with craniosynostosis, even in SS, generally considered to be the most benign of the craniosynostotic conditions. Kapp-Simon et al.\(^{14}\) concluded that the literature suggests that children with single-suture craniosynostosis have “mild but persistent neuropsychological deficits, which become more significant as cognitive demands increase at school age.” Speltz et al.\(^{22}\) found that isolated craniosynostosis was associated with a 3- to 5-fold increase in risk for cognitive deficits or learning/language disabilities.

In a long-term follow-up of 65 adolescents (approximately 14 years of age) who had undergone surgery for SS or unicoronal craniosynostosis when they were under 12 months of age, Chieffo et al.\(^{17}\) found that 7% of their sample with SS demonstrated visuospatial and constructual ability defects with associated visual memory recall deficits; 17% also exhibited selective and sustained attention deficits. Approximately one-third (30%) of their sample with anterior plagiocephaly had deficits in processing and planning speech. Their data, therefore, supported the hypothesis that children with SS or unicoronal craniosynostosis may still manifest lower than average results at long-term selective neuropsychological evaluations, even if they have undergone early surgical treatment.

Maggi et al.\(^{16}\) argued that a VIQ > PIQ discrepancy can represent a risk of developing a learning disability and is suggestive of visual-motor deficiencies, and the authors go on to identify a higher incidence (50%) of learning disabilities (reading, spelling, and visual-motor problems) in their patient group compared with the general population (5%). This is despite the fact that their IQs fall within the normal range, with the mean PIQ being in the average range and the mean VIQ and FSIQ being in the high average range. They argued that despite their normal IQ, this is not borne out in measures of achievement or performance, which are detecting more subtle difficulties (underachievement in literacy).

The current study replicates Magge et al.’s\(^{16}\) finding of a VIQ > PIQ discrepancy in children with nonsyndromic craniosynostosis who fall within the normal range for intelligence (with a much larger sample size and more uniform age). However, although both studies find this discrepancy to be statistically significant when the difference in scores is analyzed for a group of patients combined, at an individual level, a difference of 8.3 in the Magge et al.\(^{16}\) study would only be significant at a p = 0.15 level, and the degree of difference in the current study would also not be considered statistically significant. There is also the issue that a statistically significant difference is not the same as a clinically meaningful difference. It is possible for the difference between an individual’s VIQ and PIQ to be significant in the statistical sense, but for it not to be rare among the general population. For instance, 25% of the population would show a VIQ > PIQ discrepancy of 10 points within the ability range of Magge et al.’s\(^{16}\) patient population. Unfortunately, those authors did not report how many of the patient group showed the VIQ > PIQ discrepancy. However, in the current study, 68.1% of the SS group showed the VIQ > PIQ discrepancy, and this is clearly above what would be expected in the general population.
Furthermore, in the later study by Chieffo et al., a difference of more than 12 points was found between VIQ (mean 111.4) and PIQ (mean 99) in SS patients. However, there was no difference in right coronal patients (mean VIQ 99.5, mean PIQ 95.6), and the difference was reversed in left coronal patients (mean VIQ 92.6, mean PIQ 105).

The current data are consistent with the literature and also with our earlier findings of specific areas of locomotor difficulty at younger ages (Bellew et al.4,5). These earlier data indicated that children with SS have significantly poorer gross locomotor function than the normal controls. However, following surgical intervention, the deficit was shown to have resolved and, consistent with this, a lesser improvement in eye-hand coordination and performance skills was shown. Overall developmental attainment also improved postoperatively. The children with SS who did not undergo surgery did not show any improvement in development. These findings were maintained and improved upon by 5 years of age. The current data suggest that despite this finding, verbal skills remain superior to performance skills and may indicate nonverbal learning difficulties.

This is consistent with (noncraniofacial) work by Neumann and Walker,17 who argued that neuromotor soft signs (e.g., motor coordination problems) at a young age may later persist as subtle perceptual-motor deficits in older children and adults and may be predictive of cognitive, behavioral, and psychiatric problems in later childhood and adolescence. It is not, however, consistent with previous studies that indicate that those with SS are at increased risk of speech and language impairment, particularly expressive language impairment, despite no global cognitive impairment.20

IQ in Different Types of Craniosynostosis

The current data also revealed some variation across the different types of craniosynostosis, with children in the SS group showing the highest FSIQ and the most normal distribution. This is consistent with the view that SS is usually associated with normal development and represents more of a cosmetic problem.1 Children with other types of craniosynostosis had significantly lower FSIQs, which did not show a normal distribution. (Chieffo et al. had a similar finding with their SS and unicoronal craniosynostosis patients.) The bicoronal synostosis group had the lowest attainment, and this is consistent with the view that developmental delay is more likely the more sutures involved.12 This was not, however, borne out by the FSIQ of the multisuture group, although it should be noted that this group was disadvantaged in the study by having a small sample size, a range of individual diagnoses, and a large range of FSIQs (69–134). The data for the metopic patients is also consistent with the literature, which suggests that metopic craniosynostosis is associated with a higher rate of developmental delay than other types of nonsyndromic craniosynostosis. Cognitive and behavioral abnormalities have been reported in about a third of metopic patients.6,21

Surgery and Developmental Outcome

Magge et al.16 also pointed out that previous studies12...
have asserted that children operated on at a younger age (under 1 year) have better outcomes. They did not find any evidence of this in their study, but felt the issue had not been adequately explored, as none of the children in their study had undergone cranial reconstruction after 1 year of age. In our current study, the age range at time of surgery was much wider (3.1–60.3 months), with 21 children in the group overall (10 children in the SS group alone) above the age of 1 year. Our data showed a significant effect for age at time of surgery on later IQ, with SS and bicoronal craniosynostosis patients undergoing surgery at ≤ 6 months having higher FSIQs and PIQs than those undergoing surgery later, and patients receiving surgery at ≥ 19 months performing the least well.

It did, however, seem that the VIQ > PIQ discrepancy occurred regardless of whether patients had undergone surgery. Sixty-eight percent of the patients in this study had been operated on. Limited published data are available with which to compare this figure as the majority of papers on craniosynostosis focus on groups of patients who have undergone, or are due to undergo, surgical correction. In one study of patients with SS, 8 (44%) of 18 cases had undergone surgery, and in another study, 63 (75%) of 84 patients with single-suture craniosynostosis had been operated on. In our patient group, the apparently low number of patients electing to undergo surgery is, at least in part, because the developmental data were collected from every patient attending the service, and this will have included those with mild presentations. In our previous study (Bellew et al.), we identified that 11 of 23 nonsurgical SS cases had had mild presentations. It is interesting, therefore, that the VIQ > PIQ discrepancy seems to occur regardless of whether surgery has taken place, as it suggests that the impact on development occurs even in mild cases.

Sex

The data on sex suggested that while males did not differ from females in terms of FSIQ, VIQ, or PIQ, they did differ from females in terms of showing the VIQ > PIQ pattern of development. This only reached significance when all types of craniosynostosis were combined and in the SS group, but the lack of significance with the other types of craniosynostosis may have reflected small sample size. However, it was striking that the female patients showed the opposite pattern (PIQ > VIQ) in about half of all cases, whereas the majority of male patients (67%) showed VIQ > PIQ. Unfortunately, neither the Chieffo et al. or Magge et al. studies looked at the effects of sex, and so no comparison with the previous data can be made.

Causal Basis

Although there is now a growing body of evidence showing an increase in risk for cognitive deficits in patients with single-suture craniosynostosis, the causal basis for this remains unclear. There is a suggested inverse relationship between raised ICP and IQ, particularly in untreated cases, but the incidence has been shown to increase with the number of sutures involved, and it is generally accepted that raised ICP is unusual in single-suture craniosynostosis. It has been reported to occur in only 7%–13% of patients with SS, and in our previous study of developmental attainment in 5-year-old children with SS, only 1 child in a sample of 32 had confirmed raised ICP. In the current study, only 2 patients had confirmed raised ICP (> 20 mm Hg). One had a diagnosis of pan craniosynostosis, and the other had SS and unilateral lambdoid craniosynostosis. It therefore seems unlikely that the current findings are related to raised ICP although this was not formally assessed in every case. All the patients, however, received regular developmental and ophthalmology reviews, to indirectly assess for raised ICP, with a low threshold to offer ICP monitoring to those parents not wanting to have surgery.

It is, of course, possible that there is a sample bias, as it could be hypothesized that children with problems are more likely to be brought to their follow-up appointments, and certainly by 10 years of age there is a large attrition rate. However, as most of the children had an IQ within the normal range, this seems unlikely to entirely account for the findings. Furthermore, it was striking that the numbers of patients who presented at 10 years were different proportionally to those who were operated on, or seen for developmental assessment, at less than 5 years of age. Typically, patient distribution is SS > metopic > unicominal > bicoronal > multiple, but in the current sample there were fewer surgical metopic cases than would be expected. Traditionally these would be the patients expected to have more cognitive and behavioral problems and therefore more likely to turn up for assessment, but clearly this was not the case in the current study, and so bias cannot account for the research findings.

Kapp-Simon et al. note that anatomical studies of children without single-suture craniosynostosis have suggested that disruption to specific brain structures may be associated with particular cognitive deficits. It could, therefore, be hypothesized that an abnormally shaped skull, as present in craniosynostotic conditions, would produce a secondary deformation of the brain that could result in the disruption of normal neuropsychological development. Lin et al. conducted a comparative analysis of their “shape descriptors” in an attempt to understand the impact of skull deformations on neurobehavior. They found that scaphocephaly severity indices and symbolic shape signatures were predictive of mental ability and psychomotor functions, respectively, which suggests the possibility that secondary deformation could influence neurodevelopmental status. More recently, Beckett et al. have used 3-T MRI and diffusion tensor imaging to show altered neocortical structural and functional connectivity in SS, which they feel, in part, may explain the neuropsychological deficits commonly reported in this patient population.

Conclusions

The results of this study confirm the finding that at long-term follow-up, although children with nonsyndromic craniosynostosis fall within the normal range for intelligence, there is a VIQ > PIQ discrepancy above what would be expected in the normal population, and which may therefore be indicative of more subtle difficulties in achievement.
This is also affected by type of craniosynostosis, sex, and age at time of surgery.

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References


Author Contributions

Conception and design: Bellew. Acquisition of data: Bellew. Analysis and interpretation of data: both authors. Drafting the article: Bellew. Critically revising the article: both authors. Reviewed submitted version of manuscript: both authors. Approved the final version of the manuscript on behalf of both authors: Bellew. Statistical analysis: Bellew.

Correspondence

Maggie Bellew, Department of Plastic, Reconstructive, and Hand Surgery, Leeds General Infirmary, Belmont Grove, Leeds LS2 9NS, United Kingdom. email: maggie.bellew@nhs.net.