RESEARCH ARTICLE

The head shaft angle is associated with hip displacement in children at GMFCS levels III-V - a population based study

L Finlayson¹, T Czuba², M S Gaston³, G Hägglund⁴* and J E Robb⁵

Abstract

Background: An increased Head Shaft Angle (HSA) has been reported as a risk factor for hip displacement in children with cerebral palsy (CP) but opinions differ in the literature. The purpose of this study was to re-evaluate the relationship between HSA and hip displacement in a different population of children with CP.

Methods: The Cerebral Palsy Integrated Pathway Scotland surveillance programme includes 95% of all children with CP in Scotland. The pelvic radiographs from 640 children in GMFCS levels III-V were chosen. The most displaced hip was analysed and the radiographs used were those taken at the child’s first registration in the database to avoid the potential effects of surveillance on subsequent hip centration. A logistic regression model was used with hip displacement (migration percentage [MP] ≥40%) as outcome and HSA, GMFCS, age and sex as covariates.

Results: The MP was ≥40% in 118 hips with a mean HSA of 164° (range 121–180°) and < 40% in 522 hips with a mean HSA of 160° (range 111–180°). The logistic regression analysis showed no significant influence of age and sex on MP in this population but a high GMFCS level was strongly associated with hip displacement. An increased HSA was also associated with hip displacement, a 10° difference in HSA for children adjusted for age, sex, and GMFCS gave an odds ratio of 1.26 for hip displacement equal or above 40% (p = 0.009) and hips with HSA above 164.5 degrees had an odds ratio of 1.96 compared with hips with HSA below 164.5 degrees (p = 0.002).

Conclusion: These findings confirm that HSA is associated with hip displacement in children in GMFCS levels III-V.

Keywords: Children, Cerebral palsy, Hip displacement, Head shaft angle

Background

For hip surveillance in cerebral palsy (CP) it is important to identify risk factors for hip displacement both for the treatment of the individual child and to optimise the monitoring programme. Young age and severe limitation of gross motor function as measured with the Gross Motor Function Classification System (GMFCS) are known risk factors.

The measurement of the head-shaft angle (HSA, Fig. 1) was popularised by Southwick [1] to measure the degree of slip in slipped upper femoral epiphysis. Foroohar et al. [2] evaluated the relationship between the femoral shaft and head in children with cerebral palsy (CP) and compared the HSA in children with CP with the HSA in typically developing children. They also evaluated the HSA in children with CP who had significant hip displacement. They concluded that the HSA was greater in children with CP and more pronounced in those at risk for displacement and that the evaluation of HSA may be prudent in children with CP.

Lee et al. [3] analysed HSA, Migration Percentage (MP) and the neck-shaft angle (NSA) in 384 patients with CP. They found a higher correlation between NSA and MP than between HSA and MP and concluded that the NSA appeared to be more clinically relevant than the HSA in evaluating proximal femoral deformity in patients with CP. However, the material included children up to 17 years of age some of whom did not have a well demarcated growth plate, making HSA measurement less reliable. Subsequently, van der List et al. [4] reported from a retrospective study that the HSA decreased over...
time in normal hips and children in GMFCS levels II-III but not in GMFCS levels IV-V, the two groups most at risk of developing hip subluxation. In the same year, van der List et al. [5] concluded from a retrospective cohort that, at age two years, the GMFCS and HSA were valuable predictors for hip displacement, but at the age of four years, only the MP should be used in the prediction of hip displacement. These studies seemed to confirm that the HSA was higher in GMFCS IV-V children at least up to the age of two years, but that the effect of age on the HSA as a predictor of hip displacement was less clear.

Hermanson et al. [6] reported on a total population of children with CP in a defined geographical area followed up for five years or until the development of a MP ≥ 40% of either hip within five years. The use of MP 40% as cut-off was chosen as hips with MP ≥ 40 have a high risk for further displacement, indicating the need for surgical intervention [7]. They concluded that a high HSA was a risk factor for hip displacement in children with CP. However, in 2016 Chougule et al. [8] reported no statistically significant correlation between HSA and hip migration in children with CP aged 3–18 years in GMFCS levels III-V.

These seemingly contradictory reports on the relationship between the HSA, age and GMFCS as a risk factor for hip displacement prompted the present study from another defined population of children with CP. The Cerebral Palsy Integrated Pathway Scotland (CPIPS) surveillance programme includes 95% of all children with CP in Scotland. We have used radiographs from this total population to analyse the effect of HSA on hip displacement defined as MP ≥ 40%.

Methods

Radiological data from all children in GMFCS levels III-V was used in the analysis. Children receive a pelvic radiograph on first registration in the programme. This is taken in a standardised position which is used across Scotland [9]. Subsequent radiographs are taken according to a protocol based on age and GMFCS. In this study all radiographs taken at first registration into the programme of children in GMFCS levels III-V were used for analysis to avoid potential effects of surveillance on subsequent hip centration. Radiographs showing femoral osteotomy before inclusion in the programme were excluded. Hips were defined as displaced if the MP was ≥40%. The most displaced hip from each child was analysed. The HSA for all radiographs were measured by one observer (LF). The MP for the index radiograph had been previously been recorded in the database by clinicians responsible for the individual care of the children.

Two logistic regression models were used with hip displacement (MP ≥40%) as the outcome. In the first model the following covariates were used: HSA and age (both continuous), GMFCS (categorical, level III as reference), sex (categorical, female as reference). For the second model a ROC curve analysis between HSA and MP ≥40% was done. The ideal cut-off point was selected as the point that maximizes the product of specificity and sensitivity. HSA was dichotomised according to the ideal cutoff point and in the second model the HSA was analysed as a categorical covariate related to this cutoff value. The other covariates were similar to the first model.

The statistical analyses were performed using STATA 13 software.

Results

There were 640 children with a mean age of 8.2 years (range 0–19 years, 271 female and 369 male) fulfilling the inclusion criteria (Fig. 2). Their distribution by GMFCS
level was as follows: III - 160, IV - 184 and V - 296. One hundred and eighteen hips had a MP ≥40% in which the mean HSA was 164° (range 121–180°). Five hundred and twenty two hips with a MP < 40% had a mean HSA of 160° (range 111–180°). The first logistic regression analysis using HSA as a continuous variable showed no influence of age and sex had on MP but a high GMFCS level was strongly associated with hip displacement (Table 1). HSA was also statistically significantly associated with hip displacement. A 10° difference in HSA for patients adjusted for age, sex, and GMFCS gave an odds ratio of 1.26 for hip displacement to a MP ≥40% (p = 0.009) (Table 1). The ROC curve analysis between HSA and MP ≥40% showed that the ideal cutoff point was HSA =164.5 degrees. The second logistic regression analysis showed that HSA above 164.5 degrees had an odds ratio of 1.96 for MP ≥40% (Table 2).

Discussion
This study has confirmed Hermanson et al’s [6] findings that a high HSA is a risk factor for hip displacement in children with CP. Both studies used defined populations of children with CP and thus probably avoided selection biases. The present study does not give prospective information on HSA changes over time, unlike Hermanson et al’s study, because radiological data was taken from radiographs at the time of first registration of children into the surveillance programme. It does, however, provide information on the HSA without the potentially confounding influence of surveillance on hip centration.

The findings of the present study differ from those of Chougule et al. [8] who reported no statistically significant correlation between HSA and hip migration in children with CP aged 3–18 years in GMFCS levels III-V. There are some differences between the two studies which may explain the respective conclusions. Chougule et al. [8] used a linear regression model in their statistical analysis and randomised hips by laterality in the study design. There is a floor and ceiling effect for MP at 0% and 100% that may limit the usefulness of linear regression analysis which is not seen with logistic regression, where a binary outcome of hip ‘displaced’ or ‘not displaced’ was used and set at MP ≥40%. Randomisation

| Table 1 | Logistic regression estimates on the effect of HSA, GMFCS-level and age on MP using HAS as continuous variable |
|---------|--------------------------------------------------------------------------------------------------|
| HSA°   | Odds Ratio | 95% CI | P       |
| 1.26   | 1.06  | 1.50 | 0.009   |
| GMFCS IV | 2.86 | 1.39  | 5.90 | 0.004 |
| GMFCS V | 4.17 | 2.13  | 8.16 | 0.000 |
| Age°  | 1.03 | 0.99  | 1.08 | 0.174 |
| Sex°  | 0.94 | 0.62  | 1.42 | 0.763 |

| Table 2 | Logistic regression estimates on the effect of HSA, GMFCS-level and age on MP with HSA dichotomized |
|---------|--------------------------------------------------------------------------------------------------|
| HSA° < 164.5 degrees | Odds Ratio | 95% CI | P       |
| 1.96   | 1.29  | 3.00 | 0.002   |
| GMFCS IV | 2.78 | 1.35  | 5.74 | 0.006 |
| GMFCS V | 3.98 | 2.03  | 7.80 | 0.000 |
| Age°  | 1.03 | 0.99  | 1.08 | 0.134 |
| Sex°  | 0.94 | 0.62  | 1.42 | 0.755 |

°degrees, °years, °female as reference
by laterality may also dilute a study population of hips at risk for displacement. Children in higher GMFCS levels may have pelvic obliquity and the abducted hip would most probably have a MP within normal limits and the opposite, adducted, hip an abnormal MP. For this reason we analysed the most displaced hip in each child. Van der List et al. [5] used both hips in their analyses and also did a logistic regression analysis for displacement with MP ≥40%.

The present study does have limitations. A single observer measured the HSA in all 640 hips. Training in the measurement was given and, reassuringly, Hermanson et al. [10] have reported excellent observer reliability for HSA.

The present study is based on the radiographs of children at the time of registration into the CPIPS programme and effects of time or intervention on hip centration or HSA were not considered. The results are only valid for the age range and range of HSA-angle in the material.

Conclusion
This study, which was based on a total CP population, has confirmed that HSA is associated with hip displacement in children in GMFCS levels III-V.

Abbreviations
CI: Confidence Interval; CP: Cerebral palsy; GMFCS: Gross Motor Function Classification Scale; HSA: Head Shaft Angle; MP: Migration Percentage; ROC: Receiver operation characteristic.

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Availability of data and materials
The datasets used and/or analysed during the current study are available from the corresponding author on reasonable request.

Authors’ contributions
JER, MG and GH conceived the research question, MG the data collection, LF the radiographic measurements, and TC the statistical analyses. JER was the major contributor in writing the manuscript to which all authors also contributed. All authors have read and approved the final manuscript.

Ethics approval and consent to participate
The data reported in this study has not involved any research on human subjects. The data collected on the database is purely observational - there is absolutely no change to their treatment or clinical course as part of having this data recorded. The database is a country-wide, national level record for children with cerebral palsy in Scotland. Institutional ethical approval would not apply to the data base as it is a country-wide, national level programme, under the auspices of NHS Scotland, rather than a single institution. We do however have Scottish national level approval to collect the data in the form of an “NHS Scotland Caldicott Guardian Application to use Multiple Boards or Scotland-Wide Personal Health Information for Health Research or Audit” document, with Data Protection Registration No: Z6650266 - this document can be supplied if required. With regard to consent, all parents have given consent for their data to be used in this way and this is recorded in writing in their record.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

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Author details
1University of Edinburgh, Edinburgh, Scotland. 2Epidemiology and Register Center South, Lund, Sweden. 3Department of Orthopaedic Surgery, Royal Hospital for Sick Children, Edinburgh, Scotland. 4Lund University Department of Clinical Sciences, Orthopedics, Lund, Sweden. 5School of Medicine, University of St Andrews, St. Andrews, Scotland.

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