Joint hypermobility in school-aged children and adolescents with idiopathic scoliosis – A chance for more accurate screening?

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Abstract

Introduction and Objective. Joint hypermobility (JH) can be diagnosed in the case of increased range of joint mobility and the absence of systemic diseases. Lack of stability of the spinal joints is mentioned as one of the possible causes of scoliosis. Simplicity of hypermobility examination could help identify children at higher risk of developing scoliosis. The aim of this study was to assess JH prevalence in children with idiopathic scoliosis and to analyse coincidence of joint hypermobility, scoliosis features, age, gender and bone maturity.

Materials and method. The case-control study enrolled 125 children aged 7–18 years (mean 13.2 ± 2; Cobb angle range 10°–53°; mean 24.3 ± 11.7) diagnosed with idiopathic scoliosis. The control group included 83 volunteers. The Beighton scale was used to determine joint hypermobility. The relationship between joint hypermobility and scoliosis was tested. Secondly, hypermobility prevalence according to age, gender, curve severity, number of curvatures and Risser sign in scoliosis group was summarised. Data were compared by Student’s t-test, U Mann-Whitney, chi-square test or Fisher’s exact test where appropriate. P < 0.05 was considered statistically significant.

Results. JH was diagnosed in 64 (51.2%) scoliotic patients and in 34 (41%) control group children. The difference found was not statistically significant (p = 0.148). No significant difference was found comparing scoliosis subgroups with curve size cut-off point from 20 degrees, single/double curve scoliosis, male/female gender, age cut-off point 13.2 years of age, and Risser test score 0–2/3–5.

Conclusions. This study shows that children with JH features do not have a statistically significant increased risk of scoliosis co-diagnosis.

Key words

joint instability, joint laxity, Beighton score, posture, spinal curvatures

INTRODUCTION AND OBJECTIVE

Joint hypermobility (JH) should not be considered as a medical problem. It is a condition in which joint range of motion exceeds normal for age, gender, and ethnic background in the absence of any systemic disease [1]. To emphasize the difference, the condition in which the musculoskeletal and non-musculoskeletal complaints without systemic disease occur is known as joint hypermobility syndrome (JHS) [2]. The reported prevalence of JH varies between 2%-57% depending on age, gender, ethnicity, and race. It is more common in young women and of Asian descent. In Europe, the prevalence of JH in children fluctuates around 15–20% [3].

JH is commonly diagnosed with the Beighton score that evaluates the ability to perform a selection of joint manoeuvres [4], with a maximum total of 9 points. A value of 4 or more is typically considered positive for JH. Researchers studying JH confirm, that the Beighton score is a valid instrument to be used in children [5]. JH occurrence correlates with articular pain (mainly knee, foot and spine), postural disorders and planovalgus feet. Previous studies found that JH was more frequent in children and adolescents with idiopathic scoliosis [6, 7].

Scoliosis is defined as 3-dimensional deviation of the spine (lateral with Cobb angle of 10 degrees or higher) and axial rotation [8, 9]. Scoliosis can be defined as secondary (when associated with known medical problem) or idiopathic (without defined cause in otherwise healthy children) [10]. Prevalence of idiopathic scoliosis (IS) have a wide range of 0.93–12%, with value of 2–3% as most often repeated [10]. It is more common in girls (except for infantile idiopathic scoliosis) with the ratio increasing from 1.4:1 with Cobb angle between 10°–20° to 7.2:1 for above 30° [11, 12]. In retrospective studies, Fadzan et al. summarized that the prevalence of scoliosis is higher in adolescents participating in sports activities, such as dancing, ballet, swimming, tennis, table tennis, hurling, javelin, volleyball, and gymnastics [13].

Understanding of IS etiopathogenesis is still limited. There is a pattern of familial occurrence of IS suggesting genetic etiology. Researchers noted that IS may be connected to disorder of estrogen receptor structure and function, mucopolysaccharide and lipoprotein synthesis, melatonin synthesis and co-existent disorders of blood platelet and collagen function. Latest studies point for calmodulin as a protein connecting IS, melatonin level and platelet’s function.
Other researchers suggest that MMP-3 and IL-6 promoter polymorphisms or increased BNC2 expression could possibly be genetic markers for IS [10].

There are many risk factors that may predispose to spinal instability. Joint hypermobility, delayed puberty, asymmetric spinal load, placing the thoracic spine in a lordotic position, growth-related factors (greater body height and growth velocity of more than 2 cm per year), low body mass index and abnormal levels of leptin are also suspected to play a role in scoliosis development [13]. Finding the association of JH with the development of IS is an important part of medical care for children and adolescents especially from rural regions where diagnostic imaging and therapeutic intervention may be delayed due to reduced accessibility [14,15]. The diagnosis of joint hypermobility during a simple physical examination could identify children at higher risk of developing scoliosis.

MATERIALS AND METHOD

Patients were recruited in the Orthopaedic and Rehabilitation Centre for Children and Youth in Konstancin-Jeziorna, Mazowieckie Province, Poland, from 01.2018–07.2019. Admitted patients were pre-screened to generate lists of potential study subjects. The patients and their legal guardians were then approached and both provided informed written consent. Inclusion criteria: age 7–18; IS diagnosis based on anteroposterior radiogram with Cobb angle ≥10° and vertebral rotation, exclusion criteria: secondary scoliosis, connective tissue disease in medical interview. Control group included children and adolescence volunteers examined from April–June 2019. To minimize confounding of gender and age, which are known to affect the Beighton score, the group was matched by average age and gender ratio. Inclusion criteria: age range 7–18, exclusion criteria: previously diagnosed incorrect posture (irregularities in the shape of the spine with suggestion of a dedicated therapeutic intervention) and connective tissue disorder in medical interview. The following information was collected from the medical records: medical history, age and gender. Maximum lateral spinal curve size was measured by BDK (intra-rater reliability 0.941 with 2° error range) using the Cobb method [16]. Number of curves and the US Risser sign [17] were also determined on anteroposterior radiogram of whole spine. Classification of IS depending on angle of lateral deviation used in this study was excerpted from SOSORT guidelines and adopted (Tab. 1) [10].

Joint mobility was examined by BDK using the Beighton score (between 0–9) via the Beighton Scale (Tab. 2) [4]. Transparent plastic goniometer (Saehan Plastic Goniometer 08–030111, Masan, Korea) was used to assess knees and elbow hyperextension >10°. The side of the body on which the bilateral measurements were made first was selected at random. Patient with result of 4 or more was identified as hypermobile.

Statistical analysis. Continuous data were presented as a mean value and standard deviation (SD). Categorical data were presented as a percentage. Normal distribution was verified by Kolmogorov-Smirnov test. Continuous data were compared by Student’s t-test or U Mann-Whitney test where appropriate. Categorical data were compared by chi-square test and Fisher’s exact test. P value less than 0.05 was considered statistically significant. Data were analyzed using IBM SPSS Statistics for Windows, v21 (IBM Corp., Armonk, NY, USA)

RESULTS

Basic statistics of the studied groups. The study group consisted of 125 patients with idiopathic scoliosis and the control group of 83 children and adolescents [16]. Girls statistically more often showed hypermobility in all examined cases (p=0.01) and the control group alone (p=0.01), but not in the study group (p=0.24). Basic parameters comparing both groups are shown in Table 3.

Cobb angle range was 10°–61°, mean 21.85°± 11.99. The quantitative division due to the angular size of the IS curvature is typical and presented in Figure 1.

Comparative analysis of hypermobility in the study group and control group. The differences found between patients and controls for JH were not statistically significant considering all used cut-off points.

Individual comparison of almost all tested manoeuvres in the groups did not show statistical significance. The only exception to this observation was the measurement of left knee hyperextension (Tab. 4).

Scoliosis intra-group analysis. The division of the AIS group into subgroups considering the features of scoliosis,
bone maturity or demographic characteristics, allowed for quantitative and statistical analysis.

Figure 2 shows a scatter plot of the relationship between the scoliosis angle and the Beighton score. The trend line shows a weak relationship consistent with the hypothesis made. With the division considering the angular size of scoliosis, patients with a curvature of more than 20 degrees more often presented hypermobility, but the difference was not statistically significant. Also, while comparing the subgroups with single and double curve scoliosis, no statistically significant difference was observed in the frequency of JH. The study did not show a statistically significant difference in the occurrence of hypermobility in the study group divided into subgroups by gender and by age (cut-off point at the median – 13.2 years), and no correlation with level of bone maturity represented by the Risser Sign. Table 5 summarizes the results and level of statistical significance.

**DISCUSSION**

It is well known that joint hypermobility and scoliosis co-occur in diseases involving connective tissue, especially the Ehlers-Danlos and Marfan syndromes [19, 20]. This study attempted to investigate whether the coexistence of these two conditions is coincidental, or maybe they can also be seen in a population without connective tissue disorder, and whether JH might be a factor in the idiopathic scoliosis origin. Perhaps the very existence of articular hypermobility, regardless of its cause, promotes the occurrence of scoliosis. To the best of the authors' knowledge, this is the second-largest group of IS patients examined in terms of JH, and the only study comparing patients with scoliosis in terms of bone age denoted by the Risser Sign. This topic has already been researched by scientists. The earliest identified article addressing this issue dates back to 1983. Mattson at al. examined 51 girls with untreated scoliosis and 65 controls. The study showed that patients with scoliosis had similar flexibility or were even less flexible than healthy controls [21]. In the same year, Veliskakis described a high incidence of JH in the families
of children with AIS (adolescent idiopathic scoliosis) [22]. In 1988, Binns observed that girls with scoliosis had greater JH defined by a shorter thumb-to-forearm distance in a wrist bending test [23]. The study by Veldhuizen et al. confirmed the observations of Mattson on 10 patients and 10 controls concerning bending stiffness of the trunk – analysis did not show a clear difference between patients and controls [24]. Compared to the presented study, none of those mentioned above used the Beighton score to determine mobility, but were based on different sets of flexibility measurements.

Studies with a similar methodology to the presented study were published relatively recently. Czaprowski et al. renewed the topic by publishing two papers in 2011 and 2014. The first, including 70 patients and 58 controls, showed a statistically significant (p = 0.00015) difference in the incidence of JH in children with scoliosis (51%) compared to healthy control group (19%). Even considering the lack of groups matching in terms of gender additional calculations on the same-gender subgroups, still showed a statistically significant higher incidence of JH in the IS group [6]. The second study repeated the results in a narrow population of Caucasian girls with scoliosis, where JH (defined as Beighton score ≥5), was found in 23.2% of 155 patients and 13.4% of 201 controls – a statistically significant result (p = 0.02) [7]. In the latest study by Bozkurt at al. on a apopulation of 822 Turkish children, 43 of whom were diagnosed with AIS – no significant correlation was found between scoliosis and hypermobility occurrence [3].

In the current study, the incidence of JH was greater in the group of IS patients, but the difference was not statistically significant. In previous studies on healthy populations of similar race, culture, and geographic location, the typical JH (Beighton score ≥5) prevalence was 15–20%, the same as in 2011 article by Czaprowski [3, 6, 7]. In the current study, the incidence of JH in the control group was unusually high. This may have been due to the inclusion of children with previously undiagnosed posture defects in the control group. Unfortunately, there are no studies of JH prevalence in Polish children, adolescents and the general population.

In the presented study, the only measurements that were statistically significantly different between the research and control groups, were left knee hyperextension where the results of both knee hyperextension measurements were at the border of statistical significance. This may be a possible direction for further research.

In terms of JH prevalence within the IS subgroups for scoliosis features, the findings of the current study are consistent with those of Czaprowski and Bozkut who found no association between hypermobility and scoliosis curve size or curve pattern. Demographic features compared (age, gender) showed typical trends, with JH more common in younger females, but without statistical significance [3, 6, 7]. Bone age seems to be even less related to the occurrence of JH than the record age. Perhaps if the correlation between the occurrence of scoliosis and hypermobility is poorly marked, a subsequent study would require more numerous groups. At the same time, various research methods and cut-off points make it difficult to conduct a proper meta-analysis of existing studies.

**Limitations.** The study was carried out on both first-time and previously treated patients, which could have a positive (stretching exercises) and negative (corset treatment) impact on joint mobility. However, some of the earliest articles examined the scoliosis treatment influence on joint mobility, showing that only elbow hyperextension decreased [21].

**CONCLUSIONS**

As there is no statistically significant difference in the incidence of JH in the scoliosis group compared to the control group, special vigilance for IS detection in JH children seems unnecessary.

Population studies on the incidence of articular hypermobility in Polish children and adolescents are needed. Despite the lack of statistical significance of the results of this study, the authors believe that the assessment of joint mobility should be a permanent element of the child’s examination as it affects the stability of the joints and scoliosis treatment options. Hypermobility impact on the scoliosis occurrence remains ambiguous.

**Availability of data and materials.** The datasets [16] used and/or analyzed during the current study are available from the corresponding author on reasonable request.

**REFERENCES**


Barbara Katarzyna Dobies-Krześniak, Beata Tarnacka, Agnieszka Werblińska. Joint hypermobility in school-aged children and adolescents with idiopathic scoliosis...