The current role of skeletal survey in children with suspected skeletal dysplasia

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Introduction

Skeletal dysplasia (SD) is a large, heterogeneous group of inherited disorders of cartilage/bones that affect their growth, morphology and integrity. They can be classified as osteochondrodysplasias and dysostoses (1). Osteochondrodysplasias are conditions associated with abnormalities of the growth (dysplasia) or texture (osteodystrophy) of the bone/cartilage due to genetic mutations. The changes in the affected bones may progress throughout life, changing their phenotype. Dysostoses are conditions secondary to abnormal blastogenesis in utero in the first 6 weeks of foetal life. They remain static throughout life. Currently more than 400 different entities have been described based on radiological, molecular and biochemical criteria. They may manifest in ways ranging from a barely noticeable abnormality to a severe and lethal condition. An overall prevalence of SD 2.3-7.6 per 10,000 births has been reported in different epidemiologic studies (2, 3).
The “International Nomenclature of Constitutional Diseases of Bones” group has classified these disorders intermittently since its first publication in 1970. In the 1970s, the categories were purely clinical and descriptive. The study of the genetics of SDs led to a new nomenclature based on new findings, primarily molecular in nature. Therefore, the combination of clinical, radiological and molecular knowledge has become the basis for new classifications (4, 5). Forty-two groups and 436 disorders were classified in the latest 2015 revision of the Nosology and Classification of Genetic Skeletal Disorders (6).

Families of disorders that share a common molecular basis or pathway were created. In addition, recent findings have demonstrated that identical phenotypes may result from mutations in different genes that act through a similar pathway, and that the same gene can cause different phenotypes.

It is important to remember that an accurate diagnosis of a SD is still based on detailed evaluation of clinical (family history, physical examination) and radiographic findings, followed by molecular and the biochemical tests. In general, the SDs’ common clinical and radiological findings help us to group them in several ways regardless of the specific diagnosis. Individuals with SD have to be identified, because they present with significant morbidities due to the destruction of bone and cartilage caused by defects in growth, bone modelling and regeneration, and other comorbidities (affecting the liver, kidney, lung and brain). The most widely used method for differentiating skeletal disorders showing different skeletal morphological and structural abnormalities, is “old-fashioned” radiography.

In this mini review we will emphasize the importance of skeletal survey, its position in the diagnostic algorithm of SD, and the importance of its systematic analysis.

A diagnostic approach to suspected skeletal dysplasia

Some forms of skeletal dysplasia may be recognized in utero by identifying a skeletal abnormality on antenatal ultrasound. After birth, SD is suspected in children with unexplained short stature, particularly if it is disproportionate, and in children with particularly facial dysmorphism. Dysplasia should be differentiated from other causes of short stature, such as familial short stature, endocrine disorders and metabolic disorders. Clinical evaluation includes various anthropometric measurements, description of the limb involvement, hair quality, cleft palate, eye abnormalities, immunological/hematological data, and even internal organ abnormalities, such as cystic kidney, or hepatosplenomegaly seen during abdominal ultrasonography.

A radiological evaluation is an integral part of the post-natal diagnostic workup of SD. A skeletal survey, composed of a series of radiographs, is performed in a child with a suspected bone disorder. The skeletal survey includes radiography of the skull in 2 views, the thoracolumbar spine in 2 views, the chest, pelvis, one upper limb, one lower limb, and the left hand (including assessment of bone age) (7). In new-borns, due to the small size of the baby, images can usually be joined into an AP and a lateral babygram, with all the upper and lower extremities (Fig.1 and Fig. 2) (8).

Sometimes dedicated views of specific sites may be required for adequate bone visualisation and assessment. It is important to keep in mind that a complete skeletal survey is not necessary in patients with proportionate short stature or the presence of specific local skeletal abnormalities, according to the published guidelines (7). This protects children from unnecessary radiation exposure. In general, there is nothing to be gained from repeating the images; imaging should not be repeated more than once or twice, over a period of less than one year. It is also not
recommended to image previously normal sites unless clinically indicated. In fact, some radiographic abnormalities may only become visible at an older age, making serial radiographic evaluation necessary (3). Recently, Watson et al. proposed the introduction of a standardised protocol that may be adapted in certain specific situations (9). They found that due to the variability of imaging protocols used at different centres, radiographs may be inadequate or incomplete, resulting in diagnostic delays. The proposed standardised protocol could optimize images, particularly if referred for an expert opinion, reduce the radiation dose from unnecessary exposure, and enable faster diagnosis. The radiographs should be of the high quality necessary for precise analysis. They also recommended common indications for timing of skeletal dysplasia imaging.

Fig. 1. “Babygram” of a new-born girl with asphyxiating thoracic dystrophy (Jeune syndrome): long and barrel-shaped thorax, handlebar clavicles, short horizontal ribs with bulbous anterior ends, and normal spine. Morphologically altered pelvis with small, flared iliac wings, narrowed sacrosciatic notches, and dysplastic trident acetabular roof (from our own archives).

Fig. 2. Radiogram of the upper and lower extremities in a new-born girl with Jeune syndrome: general limb shortening and rhizomelia, but without the typical precocious proximal femoral epiphyseal ossification in this case (from our own archives).
The interpretation of the skeletal survey is a challenge for paediatric radiologists with an untrained eye for skeletal disorders. The radiologist needs to analyse the skeletal survey radiographs in a systematic way, and without an organized approach the task may seem overwhelming. Two main approaches, Offiah’s and Alany’s, are suggested in order not to miss important details regarding the morphological or structural features of the bones (10-12). Offiah and Hall suggested the ABC evaluation of suspected SD, which is easy to remember and easy to use (Table 1) (10).

“A” describes the anatomical localisation and alignment. The axial skeleton is often involved and manifests as a short spine, either due to a generalised reduction in the vertebral body (platspondyly) or due to an angular deformity of the spine (scoliosis, kyphosis). When the limbs (appendicular skeleton) are involved, the type of bone shortening should be determined, as well as the predominant location of the morphological changes. Predominant involvement of the epiphyses might lead to premature osteoarthritis, of the diaphysis to deformities of the affected bones, and of the metaphysis to short bones. Descriptions of the bone alignment include dislocation, subluxation, scoliosis and kyphosis.

“B” describes different features of the individual bone with five “S”: structure (focused on bone density, but also include a description of wormian bones, exostoses, enchondromata), shape (using descriptive radiological terms such as angel-shaped, cone-shaped, stippled, flared, scalloped, trident, etc.), size (larger, smaller), sum (too few or too many bones, fused, absent or with delayed ossification), and soft tissues (wasted or excessive, contractures, calcification).

“C” describes the complications which could be part of a condition (mostly pathological fractures, compression of adjacent structures, malalignment, malignant alteration) or secondary to treatment.

“D” describes dead or alive. If the SD is lethal, this might help in diagnosis (affect the subtype) and even change the mode of inheritance.

On the other hand, Alanay et al. suggested three steps of assessment, which are similar, but less precise than the ABC evaluation (11, 12). As the first step, assessment of the disproportion is performed (a quick look at the spine – short trunk and at the extremities may help to define rhizomelia, mesomelia, and acromelia). The second step is assessment of epiphyseal, metaphyseal, and diaphyseal ossification. The third step, is the differentiation of normal variants from pathological abnormalities by an experienced paediatric radiologist.

We should be aware that sometimes no specific, firm radiological diagnosis can be reached. No diagnosis is better than a wrong diagnosis, particularly if it leads to inappropriate and expensive genetic testing, and inap-
propriate genetic counselling (1). Veeramani et al. (13) found that a clear diagnosis of SD is not possible in a third of cases (skeletal abnormalities were presented but a clear diagnosis could not be reached). In addition, their research suggested that there are differences in diagnosis rates for patients who have had a full skeletal survey compared to a limited survey (79% v. 44%). There is a need for widely accepted standardization of skeletal surveys and for access to multidisciplinary, highly specialized teams (13). Expert opinion may be sought either face-to-face (multidisciplinary meetings) or via web-based consultation (the European Skeletal Dysplasia Network or the International Skeletal Dysplasia Society).

Conclusion

SDs are not as uncommon as once thought. Without doubt, the diagnosis and management of SDs needs teamwork and multidisciplinary management, including paediatric (endocrine) radiologists, geneticists, and orthopaedists. A full skeletal survey should be the first line of investigation when a SD is suspected, followed by targeted molecular, biochemical and genetic tests, based on the radiological and clinical findings.

Conflict of interest: The author declares that she has no conflict of interest.

References


