Developmental dysplasia of hip – An overview

Dr. RK Jain and Dr. Siddharth Patel

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Abstract
Developmental dysplasia of the hip refers to a spectrum of severity ranging from mild acetabular dysplasia with a stable hip, to more severe forms of dysplasia with hip instability, to established hip dysplasia with or without subluxation or dislocation. DDH affects 1-3% of newborns and is responsible for 29% of primary hip replacements in people up to the age of 60 years. Ligamentous laxity is also believed to be associated with hip dysplasia, though this association is less clear. The treatment of DDH is age-related and the goal is to achieve and maintain concentric reduction of the femoral head into the acetabulum. A dislocated hip which can be reduced and a reduced hip that can be subluxated should be treated with the orthosis. All such hips should be treated in orthosis or harness beginning at the time of diagnosis. This results from excessive pressure on the femoral head after reduction in extreme abduction and internal rotation leading to vascular occlusion. Avascular necrosis results from excessive pressure on the femoral head after reduction in extreme abduction and internal rotation leading to vascular occlusion. Developmental dysplasia of the hip is a challenging condition. Formal training in the treatment of various age groups with DDH is mandatory. Successful treatment with Orthoses requires careful counselling of parents. Surgical management requires careful pre-operative planning and adequate follow up to ensure the best clinical outcomes.

Keywords: Avascular necrosis, developmental dysplasia, femoral head, hip

Introduction
Developmental dysplasia of the hip – DDH, (formerly known as congenital dislocation of the hip - CDH) is now the preferred term, reflecting as an ongoing developmental process. The definition of DDH is not universally agreed upon. It refers to a spectrum of severity ranging from mild acetabular dysplasia with a stable hip, to more severe forms of dysplasia with hip instability, to established hip dysplasia with or without subluxation or dislocation. Early detection and treatment of developmental dysplasia of the hip (DDH) is critical for the best chances of obtaining a well-functioning, pain-free hip joint well into adulthood, and research has shown that a child’s age at initial reduction is correlated to radiographic outcomes.

If detected within first six weeks of life is considered as early detection, and one detected beyond three months and into walking age is considered as late presentation DDH. DDH may lead to impaired hip function and premature degenerative joint disease particularly if treatment is delayed Hip instability if detected in the neonatal period can be effectively treated by non-surgical means in a majority of instances, while delay in its diagnosis increases the likelihood of surgical intervention. It has been said that delayed presentation of children with DDH is a common occurrence in India.

Overview
Background
The term congenital dislocation of the hip dates back to the time of Hippocrates. It has been diagnosed and treated for several hundred years. Most notably, Ortolani, an Italian paediatrician in the early 1900s, evaluated, diagnosed, and began treating hip dysplasia. Galeazzi later reviewed more than 12,000 cases of DDH and reported the association between apparent shortening of the flexed femur and hip dislocation. Since then, significant progress has been made in the evaluation and treatment of DDH [1, 2, 3, 4, 5].
Epidemiology

DDH affects 1-3% of newborns and is responsible for 29% of primary hip replacements in people up to the age of 60 years.\(^6\) A systematic review of unscreened populations estimated the prevalence of clinically diagnosed, established hip dysplasia to be 1.3 per 1,000 but in populations screened clinically with Ortolani and Barlow tests, the prevalence is higher at 1.6-28.5 per 1,000 and it is higher still with ultrasound screening.\(^6\)

Predisposing Factors

The exact cause of hip dysplasia is not clear, but this condition does appear to be related to a number of different predisposing factors.\(^7\)

- **Racial background:** In Native Americans and Laplanders, the prevalence of hip dysplasia is much higher (nearly 25-50 cases per 1000 persons) than in other races, and the prevalence is very low in southern Chinese and black populations.\(^6,12\)
- **Genetic predisposition:** Also appears to exist, in that the frequency of hip dysplasia is 10 times higher in children whose parents had DDH than in those whose parents did not.\(^13\)
- **Intrauterine positioning and sex:** Female sex, being the first-born child and breech positioning are all associated with an increased prevalence of DDH. An estimated 80% of persons with DDH are female\(^14\), and the rate of breech positioning in children with DDH is approximately 20% (compared with 2-4% in the general population)\(^15\).
- **Other musculoskeletal disorders of intrauterine mal positioning or crowding, such as metatarsus adductus and torticollis, have been reported to be associated with DDH.**\(^16-19\)
- **Oligohydramnios is also reported to be associated with an increased prevalence of DDH.**\(^20\)
- **The left hip is more commonly associated with DDH than the right hip,** possibly because of the common intrauterine position of the left hip against the mother's sacrum, which forces it into an adducted position.
- **Children in cultures in which the mother swaddles the baby, forcing the infant's hips to be adducted, also have a higher rate of hip dysplasia.**\(^21\)

Hip dysplasia can be associated with underlying neuromuscular disorders, such as cerebral palsy, meningo(myelo)coel, arthrogryposis, and Larsen syndrome, though such cases are not usually considered DDH.

Anatomy

The normal growth of the acetabulum depends on normal epiphyseal growth of the tri-radiate cartilage and on the three ossification centres located within the acetabular portion of the pubis (os acetabulum), ilium (acetabular epiphysis), and Ischiium. Additionally, normal growth of the acetabulum depends on normal interstitial appositional growth within the acetabulum. The presence of the spherical femoral head within the acetabulum is critical for stimulating normal development of the acetabulum.

Abnormal development of the hip includes the osseous structures, such as the acetabulum and the proximal femur, as well as the labrum, capsule, and other soft tissues. This condition may occur at any time, from conception to skeletal maturity.

The anatomy of the dislocated hip, especially after several months, often includes formation of a ridge called the neo-limbus. Closed reduction is often unsuccessful at a later date, secondary to various obstacles to reduction. These include adductor and psoas tendon contraction, ligamentous teres, a transverse acetabular ligament, and pulvinar and capsular constriction. With long-standing dislocations, interposition of the labrum can also interfere with reduction.

Fig 1: Schematic diagram of Normal Neonatal Hip Joint compared to Developmental Dysplasia of Hip Joint in Neonate.

Pathophysiology

DDH involves abnormal growth of the hip. Ligamentous laxity is also believed to be associated with hip dysplasia, though this association is less clear. Children often have ligamentous laxity at birth, yet their hips are not usually unstable; in fact, it takes a great deal of effort to dislocate a child's hip. Therefore, more than just ligamentous laxity may be required to result in DDH.

At birth, white children tend to have a shallow acetabulum\(^6,7\), this may provide a susceptible period in which abnormal positioning or a brief period of ligamentous laxity may result in hip instability. However, this characteristic is not as true for children of African descent, who have a lower rate of DDH.\(^8\)

Diagnostic Tests Recommended

**Ortolani test:** The examiner applies gentle forward pressure to each femoral head in turn, in an attempt to move a posteriorly dislocated femoral head forwards into the acetabulum. Palpable movement suggests that the hip is dislocated or subluxed, but reducible.

**Barlow test:** Gentle backward pressure is applied to the head of each femur in turn, and a subluxable hip is suspected on the basis of palpable partial or complete displacement.

**The Galeazzi sign:** The child is examined lying supine with the hips and knees flexed to 90° and the height of each knee is compared. Unilateral femoral shortening may signify hip dislocation or rarer abnormalities of the femur. False negative results may occur with bilateral hip dysplasia or when the pelvis is not level.

Other physical signs for late dislocation include asymmetry of the gluteal thigh or labral skin folds, discrepancy in leg length, a widened perineum on the affected side, buttock flattening, asymmetrical thigh skin folds and standing or walking with external rotation of the affected leg. Both Barlow and Ortolani tests detect an unstable hip but do
not detect an irreducible dislocated hip, which is best detected by identifying limited abduction of the flexed hip. Nor do they detect a stable hip with abnormal anatomy - eg, acetabular dysplasia. The Barlow and Ortolani tests are useful in neonates but become difficult by 2-3 months of age. Stable hips may be dysplastic. Limited hip abduction (less than 60°) when the hip is flexed to 90° is the most important sign of a dislocated or dysplastic hip [22].

Benign hip clicks, resulting from soft tissues snapping over bony prominences during hip movement, should be distinguished from the clunks produced during the Ortolani manoeuvre as the dislocated femoral head is reduced and from the subluxation felt during the Barlow test.

Children aged 3-6 months: The physical signs are rather different and so are the requirements of examination. Unilateral limitation and asymmetry of hip abduction is the most reliable sign of DDH after eight weeks. If the hip is dislocated it is in a fixed position. Older children there is limited abduction when fully flexed. They may walk on toes on the affected side or present with a painless limp.

**Bilateral dislocation:** It is present in about 20% of cases, it can be quite difficult to diagnose, especially after the neonatal period. There is often a waddling gait with hyperlordosis. The Galeazzi sign for hip shortening is often absent, as are asymmetrical thigh and skin folds, or asymmetrically decreased abduction. Careful examination is needed with a high level of suspicion.

**Crowe classification of Developmental Dysplasia of Hip**

<table>
<thead>
<tr>
<th>Crowe</th>
<th>Description</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Femur and acetabulum show minimal abnormal development.</td>
<td>Less than 50% Dislocation</td>
</tr>
<tr>
<td>II</td>
<td>The Acetabulum shows abnormal development</td>
<td>50% to 75% Dislocation</td>
</tr>
<tr>
<td>III</td>
<td>The Acetabulum is developed without a roof. A false Acetabulum develops opposite dislocated femur head position.</td>
<td>75% to 100% Dislocation</td>
</tr>
<tr>
<td>IV</td>
<td>The Acetabulum insufficiently Developed. Since the femur is positioned high up on the pelvis.</td>
<td>High Hip Dislocation</td>
</tr>
</tbody>
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Investigations

**Radiography:** It become primary imaging modality at 4-6 months after the femoral head begins to ossify. Radiological features in DDH.

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**Ultrasound evaluates for acetabular dysplasia and/or the presence of a hip dislocation.**

- Useful before femoral head ossification (<4-6 months of age).
- May produce spurious results if performed before 4-6 weeks of age.
- Allows view of bony acetabular anatomy, femoral head, labrum, ligamentous teres, hip capsule

**Alpha Angle**

- Angle created by lines along the bony acetabulum and the Ilium
- Normal is greater than 60°

**Beta Angle**

- Angle created by lines along the labrum and the ilium
- Normal is less than 55°
- Femoral head is normally bisected by a line drawn down from the ilium

Normal ultrasound in patients with soft-tissue 'clicks' will have normal acetabular development. It allows for monitoring of reduction during Pavlik harness treatment. It is not cost effective for routine screening

**Arthrogram**

- Used to confirm reduction after closed reduction under anaesthesia
Help identify possible blocks to reduction

Inverted Labrum
- Labrum enhances the depth of the acetabulum by 20% to 50% and contributes to the growth of the acetabular rim
- In the older infant with DDH the labrum may be inverted and may mechanically block concentric reduction of the hip

Inverted Limbus
- Represents a pathologic response of the acetabulum to abnormal pressures caused by superior migration of the head
- Consists of fibrous tissue
- Transverse acetabular ligament
- Hip capsule is constricted by ilio-psoas tendon causing hour-glass deformity of the capsule, pulvinar, ligamentous teres.

CT scan: CT study of choice to evaluate reduction of the hip after closed reduction and spica casting.

MRI: Does not play significant role in primary diagnosis.

Screening Programme for DDH
The objective of any screening programme is to identify “pre-clinical” cases (in the case of DDH, read as “neonatal hip instability”) earlier in their natural history than would occur in the absence of screening.[22]

Two basic assumptions commonly made in evaluating the effectiveness of screening are
1. All or most cases of the disease will pass through a detectable “pre-clinical phase” of illness.
2. In the absence of intervention, all or most cases in the “pre-clinical phase” would progress to a clinical phase.

A Cochrane review found that there was insufficient evidence to give clear recommendations regarding screening for DDH. There was some inconsistent evidence that universal ultrasound results in a significant increase in treatment compared to the use of targeted ultrasound or clinical examination alone. The review concluded that ultrasound strategies have not been demonstrated to improve clinical outcomes including late diagnosis and surgery.[23]

A large long-term randomised controlled study of universal or selective ultrasound versus a well-done physical examination showed higher treatment rates but no reduction in late cases of DDH.[24-29] There was also no difference, at skeletal maturity, in acetabular dysplasia or degenerative change.

Universal ultrasound examination is not recommended by Public Health England but selective ultrasound examination for babies with specific risk factors is recommended. An ultrasound examination of the hips should be performed within 6 weeks of age.[30]

Management of developmental dysplasia of the hip
The treatment of DDH is age-related and the goal is to achieve and maintain concentric reduction of the femoral head into the acetabulum. The best outcome can be expected only if the treatment is started very early.

This has been achieved in the developed world, through improved awareness and training, increased surveillance (use of ultrasound), and quicker access to paediatric orthopaedic surgeons.[31] must be noted that there are very few specialized paediatric orthopaedic surgeons in the developing world, and fellowship programs for paediatric orthopaedic surgery should be initiated to cater for this requirement, in addition to the myriad of other paediatric orthopaedic conditions.[32]

Treatment of a neonate with developmental dysplasia of the hip
Subluxation of hip noted at birth may correct spontaneously. Some physicians may continue with observation before initiating treatment. When observation is chosen, steps should be taken to ensure close follow up because some of these hips will subsequently dislocate if left alone. The child should then be re-evaluated both clinically and by ultrasound at three weeks of age to confirm concentricity.

Hips that are still dislocated need further treatment. Orthoses, such as Erlanger, Thübinger or Pavlik, which promote abduction and flexion of the hip joint can reduce an physiological pressure to the antero lateral acetabulum and are a preferred treatment option in this age group, aiming for development of a normal lateral edge of the cartilaginous acetabulum.

A dislocated hip which can be reduced and a reduced hip that can be subluxated should be treated with the orthosis. All such hips should be treated in orthosis or harness beginning at the time of diagnosis. These harnesses are very dynamic and successful tools in the treatment of DDH, although they are sometimes difficult to handle for the parents.

Once properly applied, they allow motion while the hips are flexed (>90°) as well as abducted and treatment is continued until the hip is stable, as evidenced by negative Barlow and Ortolani tests. The normal degree of vulgus of femoral head and neck requires this degree of flexion to promote spontaneous reduction of dislocation.

Application of an orthosis should be followed by bi-weekly clinical examination, and ultrasonography if required. If the hip is reduced at three weeks following application of orthosis, the patient may continue to wear it for a further three weeks. After six weeks, the orthosis is removed and the hip is examined both clinically and by ultrasound.

If the hip is reduced then the orthosis can be discontinued. The dislocated hip, even after 3–4 weeks of orthosis use, should be evaluated and may be treated with an abduction brace.[33-36] The child needs regular follow up until skeletal maturity to identify the late squeal, such as acetabular dysplasia.[37-40]

The hip should not be placed in a position of hyper flexion and hyper abduction as it may result in high pressure on the femoral head leading to osteo necrosis. In addition, there is evidence of an increased risk of femoral nerve palsy.[41-44] or inferior dislocation of the femoral head due to hyper flexion.[45] On the other hand; inadequate flexion will fail to reduce the hip.

The Pavlik harness is contraindicated ‘when there is major muscle imbalance, as in meningo myelocoele (L2 to L4 functional level); major stiffness, as in arthrogryposis, ligamentous laxity, as in Ehlers-Danlos syndrome’ or where the family situation cannot guarantee careful and consistent use of the harness.

Treatment of a child (one to six months)
Orthesis is the choice of treatment in this age group. The hip should be placed in 90 degree flexion with the proximal femur pointing towards tri-radiate cartilage. The hip which is not reducible at the time of clinical examination may still be treated with orthosis; however, higher dislocations are less likely to reduce than the lower ones.[45, 46] The position of the
hip must be confirmed after application of orthosis with AP X-ray. The child should be examined at regular intervals to ensure the reduction. The orthosis should be continued for at least six weeks after the stability is achieved. If the hip fails to reduce with orthosis then other options should be considered, such as an abduction (Von Rosen) splint. The main aim of treatment is to achieve concentric reduction and to prevent a vascular necrosis. Whatever method is used, it must be ensured that reduction can occur spontaneously. The hips should never be immobilised in a forced position.

Nakamura et al. reported his results of 115 patients with 130 hips. The mean age was 4.8 months and there was a mean follow up of 16 years. Patients were treated with a Pelvic harness with a mean duration of treatment of 6.1 months. Twenty-two hips required supplementary surgery for residual dysplasia, the choice of surgery depending on the state of joint and surgeon’s preference. A satisfactory outcome (Severin classes I and II) was achieved in 119 patients [43].

**Treatment of child (six months to two years)**

The child in this age group may be treated with either closed or open reduction, followed by a spica cast. The aim is to achieve reduction without damaging the femoral head. There are several studies favouring reduction of hips after the appearance of ossific nucleus [48]. Segal et al. reported on hips in children less than 12 months of age [49]. Thirty-eight hips were reduced closed while 17 were reduced by an open method. One patient with bilateral hip dislocation was treated initially by closed means and later treated by open reduction at three months. Avascular necrosis (AVN) developed in only one of 25 patients in which a nucleus was present while 17 of 32 patients developed AVN when reduction was performed before the appearance of an ossific nucleus at a mean follow up of 59 months.

Roposch et al. conducted a meta-analysis including 6 observational studies of a total of 358 patients with a mean age of 9.6 months at the time of reduction. Closed as well as open reduction was performed in 3 studies, open reduction alone was performed in 2 studies and closed reduction alone was performed in one study. They noted that the presence of an ossific nucleus had an insignificant effect on the development of AVN when all grades were considered. Forty-one (19%) patients developed AVN when an ossific nucleus was present as compared to 30 (22%) patients when this was absent. However, the absence of an ossific nucleus was associated with the development of osteonecrosis when grade II or more were considered: 14 (7%) patients with an ossific nucleus and 18 (16%) without an ossific nucleus [50].

On the other hand, there are studies which do not confirm this. Konigsberg et al. presented his results of 40 patients in whom an open reduction through a medial approach had been performed. Average age was 7.7 months, ranging from 2.4 to 18.9 at the time of surgery, with a mean follow up of 10.3 years. Only one of 20 hips reduced before the age of six months developed AVN [51]. Other investigators have reported that hips reduced after the appearance of an ossific nucleus have a higher number of operative procedures [52]. In addition, due to reduced growth potential, hips reduced later will not remodel as well as those reduced earlier.

The use of pre-reduction traction is controversial. It is supported by many studies showing a decreased rate of AVN and relative success of closed reduction [53-55]. On the other hand, there are studies showing a similar rate of open reduction (56) and AVN in which traction was not used [56, 57]. Open reduction may be achieved through a medial or anterior approach. Minimal dissection is required in a medial approach and allows the surgeon to achieve a stable reduction without the risk of AVN [58]. Disadvantages of the medial approach include inadequate exposure, risk to medial circumflex femoral vessels and inability to perform capsulorrhaphy. Post-operatively a cast is applied and changed after six weeks for a total period of three months.

The anterior approach has a better visualization and allows the surgeon to perform a good capsulorrhaphy [59]. The choice of approach depends upon patient age and the surgeon's experience. A medial approach is recommended for children under one year of age with a maximum age limit of 18 months in expert hands.

**Treatment of older child (two years of age and older)**

In older children, the femoral head lies in a more proximal location. Previously, pre-operative traction was used to bring the hip into the normal position, but now femoral shortening has replaced the use of traction. Femoral shortening is usually required after the age of two years to reduce the pressure on the femoral head after reduction. This will decrease the risk of osteonecrosis [60].

Sankar et al. studied the factors predicting the need for femoral shortening in 72 hips (64 patients). All patients underwent open treatment for DDH with a mean age of 35.6 months and a mean follow up of 21.4 months. He concluded that the patients over the age of 36 months and patients with vertical displacement greater than 30% of the width of pelvis were more likely to require femoral shortening [61].

The aim of femoral derotation varus osteotomy is to achieve concentric reduction. Spence et al. compared two groups of patients undergoing open reduction through an anterior approach either with femoral derotation osteotomy (38 patients with 47 hips) or innominate osteotomy (33 patients with 37 hips). Mean age at reduction was 25.3 months in the first group and 21.9 months in the second group. Mean duration of follow up was 6.2 years. Acetabular remodelling and stability was better in the patients with innominate osteotomy compared to femoral derotation osteotomy [62]. Adequate post-reduction coverage of the femoral head is essential. The risk of residual dysplasia at skeletal maturity increases as open reduction is performed in older children. If femoral head coverage is inadequate, then pelvic osteotomy should be considered in this age group. Salter innominate and Pemberton osteotomy are the most commonly used techniques [63].

If closed or open reduction is performed in a child under 18 months of age, the child should be followed till the age of 3.5–4 years. If the femoral head coverage is still inadequate, then pelvic osteotomy should be considered. Bohm et al. studied 73 hips in 61 patients with a mean age of 4.1 years at the time of surgery. He concluded that good clinical results can be expected if normal acetabular anatomy is restored without the development of AVN. However, pelvic osteotomy should preferably be performed at a later stage than at the time of open reduction [64]. Thomas et al. presented the results of 80 hips in 60 patients with a mean follow up of 43.3 years. All patients underwent open reduction, capsulorrhaphy and Salter innominate osteotomy at a mean age of 2.8 years. Failure was defined as the joint requiring replacement surgery. They reported a 54% survival rate with excellent prognosis in two-thirds who were able to function at a high level [65].

El-Sayed et al. reported 71 surgical interventions in 55
patients (ages ranging from two to four years) in whom open reduction, Salter innominate osteotomy, and proximal femoral osteotomy were performed in a single stage. The mean follow up was five years and four months. They reported favourable clinical and radiological results. Mean pre-operative acetabular index was 41.86 while the final mean acetabular index was 16.78 [66].

Late presentation acetabular dysplasia

A number of patients present late in adolescence with complaints of aching pain either in the groin or lateral hip pain which increases after exertion. After detailed physical and radiographic assessment, a treatment strategy is devised for these dislocated hips. The goal of treatment is to attain concentric reduction by realignment osteotomy of the acetabulum to cover the femoral head such as Salter, Pemberton and Dega osteotomy. For hips that cannot be concentrically reduced, the aim is to cover the femoral head with structures that become fibro cartilage. The Chiari osteotomy and the shelf procedure are two such approaches [67].

Complications

Avascular necrosis

This results from excessive pressure on the femoral head after reduction in extreme abduction and internal rotation leading to vascular occlusion. The incidence of AVN varies widely from 0–73% [68] depending on the age, mode of treatment and criteria used to describe AVN. Four percent incidence was reported by Weiner et al. in children under the age of three months. AVN can easily be prevented by performing femoral shortening [69]. Change in femoral head density, failed ossification and widening of the femoral neck suggest a vascular changes.70 The greater trochanter is usually preserved and continues to grow whereas the femoral epiphysis is damaged which disturbs hip biomechanics. There are a number of procedures to correct the mechanical imbalance, such as trochanteric epiphysiodesis [71], trochanteric advancement [72] and a lateral closing wedge along with trochanteric advancement.

Residual dysplasia

Acetabular index measures the severity of the residual dysplasia. In current literature, the definition of acetabular dysplasia is inconsistent [73]. Gwynne et al. defines dysplasia as definite if the acetabular index is more than 30 degrees at 6-month X-rays (corresponding to more than 2 SD above normal at this age) and as mild if greater than 25 degrees [74]. Cashman reported a 3.5% incidence of residual acetabular dysplasia in his study [75]. Acetabulum remodels in response to the pressure exerted by the femoral head after concentric reduction. However, this process may remain inefficient and results in a shallow acetabulum providing inadequate coverage and poor outcome [76]. This can be treated by acetabular reorientation surgery [77].

Prognosis

Overall, the prognosis for children treated for hip dysplasia is very good, especially if the dysplasia is managed with closed treatment. If closed treatment is unsuccessful and open reduction is needed, the outcome may be less favourable although the short-term outcome appears to be satisfactory. If secondary procedures are needed to obtain reduction, then the overall outcome is significantly worse. Some authors believe that patients with bilateral hip dysplasia have a poorer prognosis because of frequent delays in diagnosis and greater treatment requirements [24, 25].

In a study comparing the outcomes of walking-age children with bilateral hip dislocations who underwent open reduction and pelvic osteotomy with or without femoral osteotomy with those of walking-age children with unilateral dislocated hips who underwent the same set of procedures, the radiographic outcomes were similar [26]. In this study, the rate of osteonecrosis was higher in the bilateral group, but this difference was explained by older age at surgery and a greater degree of hip dislocation before surgery. The authors concluded that the clinical outcomes after surgery of the children with bilateral hip dislocations were worse mainly because of asymmetric outcomes.

Conclusion

Developmental dysplasia of the hip is a challenging condition. Formal training in the treatment of various age groups with DDH is mandatory. Paediatricians and family physicians should have a high index of suspicion and referral to a paediatric orthopaedic surgeon should be made early. The outcome of treatment during the first six months of life is much better than in late-diagnosed DDH. Successful treatment with Orthoses requires careful counselling of parents. Surgical management requires careful pre-operative planning and adequate follow up to ensure the best clinical outcomes.

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