Pathomorphology and treatment of congenital anterolateral bowing of the tibia associated with duplication of the hallux

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Pathological congenital bowing of the tibia in the newborn is rare. Posteromedial and anteromedial bowing may be treated by ‘benign neglect’, but anterolateral bowing should alert the surgeon to the possibility of the development of pseudarthrosis of the tibia and impending fracture.1,2 In contrast to other types of tibial bowing, anterolateral bowing in association with congenital tibial pseudarthrosis is rarely diagnosed at birth and usually develops during the first decade of life.3

In 1994, Tuncay, Johnston and Birch3 reported five cases which seemed to represent a ‘benign’ form of congenital pseudarthrosis of the tibia. Anterolateral bowing of the tibia was observed, with the fibula remaining structurally normal and straight and the deformity of the tibia resolved spontaneously without any surgical treatment. Congenital anterolateral bowing of the tibia in association with a bifid ipsilateral great toe has been reported as a new entity.4-8 None of the reported cases was associated with neurofibromatosis or pseudarthrosis of the tibia. Spontaneous resolution of the axial deformity has been described. In all cases the anterolateral bowing was exclusively in the tibia, the fibula remaining straight and uninvolved.

In previous reports, surgery was only performed for duplication of the ipsilateral great toe and metatarsal involvement. Leg-length discrepancy was noted by all previous authors but little attention was paid to persistent severe subluxation of the proximal tibiofibular joint due to the axial deformity and discrepancy of tibiofibular length.

We now report three further cases of severe anterolateral bowing of the tibia occurring in association with ipsilateral bifidity of the great toe.

Patients and Methods

We describe three patients (two girls and one boy) with congenital anterolateral bowing of the tibia in association with duplication of the hallux who presented between 1990 and 1999. All three were between 14 and 29 days of age when first seen and all the deformities were unilateral. Pregnancy and birth histories were normal and family histories were negative for deformities of the lower limb and neurofibromatosis. No other deformities were present at birth or developed later.

All three patients had the same deformities. At birth the tibia was markedly bowed in the anterolateral direction. The apex of the deformity was in the middle of the diaphysis of the tibia. Plain radiography showed some irregularity which suggested localised duplication of the tibial medullary canal. Duplication of the great toe was present in all three patients and treated by early resection and reconstruction during the first year of life (Fig. 1).

Congenital unilateral anterolateral tibial bowing in combination with a bifid ipsilateral great toe is a very rare deformity which resembles the anterolateral tibial bowing that occurs in association with congenital pseudarthrosis of the tibia. However, spontaneous resolution of the deformity without operative treatment and with a continuously straight fibula has been described in all previously reported cases. We report three additional cases and discuss the options for treatment. We suggest that this is a specific entity within the field of anterolateral bowing of the tibia and conclude that it has a much better prognosis than congenital pseudarthrosis of the tibia, although conservative treatment alone may not be sufficient.
During the first four years of life the tibial deformity spontaneously corrected to some extent. Despite treatment by orthoses after the age of four years, further spontaneous resolution of the tibial bow was not seen. A leg-length discrepancy up to 5.5 cm also developed. The area of major tibial bowing was found in the middle third of the diaphysis in all three patients. Plain radiography and MRI showed signs of focal tibial duplication in the area of the bowing (Fig. 2). No signs of pseudarthrosis or other clinical or radiological features suggestive of neurofibromatosis were seen. The fibula appeared to be structurally normal and straight but clinically the fibula head was prominent proximally causing recurrent localised pain due to overgrowth of the fibula and subluxation of the proximal tibiofibular joint (Fig. 3) (Table I).

Reconstructive surgery using the Ilizarov method was performed at the age of five to six years. The aim was to correct the leg-length discrepancy with simultaneous axial correction and treatment of the subluxation of the proximal tibiofibular joint. All patients were treated by the same step-by-step treatment. The Ilizarov ring fixator was applied, and tibial osteotomy performed at the site of maximal tibial bowing.

Lengthening of the tibia alone, without fibular osteotomy, allowed gradual distalisation of the fibular head. After completion of step one, gradual correction of the bowing and simultaneous further lengthening were performed. Slight overlengthening was carried out to anticipate further leg-length discrepancy with growth (Table II).

Illustrative case report. A five-year-old girl was born at 42 weeks of gestation to non-consanguineous parents. The family history was negative for neurofibromatosis and bowing of the tibia. At birth severe anterolateral bowing of the midshaft and shortening of the right tibia compared with the ipsilateral fibula and contralateral tibia were present. Ipsilateral hexadactyly with duplication of the great toe was evident clinically and radiologically and led to severe foot problems. Surgery was first performed at the age of one year when the duplicated medial anlage of the great toe was resected. Metatarsal hypoplasia and duplication of the medial cuneiform and navicular led to marked shortening of the first ray. The tibial bowing was treated by an orthosis until the age of five years. Partial but not complete spontaneous resolution of the tibial malalignment was seen.

At the age of five years the indications for operative reconstructive surgery were severe proximal fibular overgrowth and gait problems because of marked varus of the ankle and leg-length discrepancy. Plain radiography showed malalignment of 20° in the frontal and 5° in the
sagittal plane. Another striking feature was the marked posteromedial thickening of the tibial cortical bone, with no sign of cystic or degenerative change suggestive of congenital tibial pseudarthrosis. MRI showed signs of tibial duplication. The tibial medullary canal was continuously open but narrow in the area of the bowing. The distal tibial epiphyseal height was 0.6 cm compared with 0.9 cm on the contralateral side and the distal tibial epiphyseal width was 3.1 cm compared with 3.6 cm on the contralateral side suggesting slight tibial hypoplasia. Orthoradiography showed a deficiency of tibial length of 17% compared with the contralateral side, and a leg-length discrepancy of 3.1 cm. The fibula was straight and showed no pathological bony findings suggestive of pseudarthrosis. Proximal overgrowth in comparison with the ipsilateral tibia was 2.8 cm, leading to subluxation of the proximal tibiofibular joint. The fibula was hyperplastic distally.

Treatment using the Ilizarov external fixator was undertaken in two steps. Initially, tibial osteotomy in the diaphyseal area with minimal partial resection and partial acute correction was performed. Intra-operatively, a well-defined duplication of the medullary canal of the tibia was seen (Fig. 4). At the same operation, proximal tibial percutaneous osteotomy was performed to allow tibial lengthening and simultaneous lowering of the fibular head with distal fixation only of the fibula to the frame. After lengthening of the proximal tibia of 2.6 cm, the fibular head had migrated to its correct anatomical position. The second step of the procedure was then undertaken fixing the fibula to the proximal ring with a fine wire together with midshaft fibular osteotomy to allow further tibial lengthening and axial correction. Distraction was completed with slight overlengthening (0.7 cm) and complete correction of the axial deformity. After full consolidation of the lengthened bone, the apparatus was removed at 140 days with a healing index of 37 days per cm. At the latest follow-up six years after surgery, no recurrent tibial deformity was present but the duplication of the navicular, cuneiform, and first metatarsal bones were causing pain and difficulties with gait (Fig. 5).

Results
The results of reconstructive surgery in the three patients are summarised in Table II.

The fibula remained straight during axial correction and lengthening of the tibia, with no sign of thinning or cystic

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**Table I.** Radiological measurements showing spontaneous relief of the anterolateral bow whereas leg-length discrepancy and proximal overlength of the fibula are severe.

<table>
<thead>
<tr>
<th>Case</th>
<th>Age at surgery (yrs)</th>
<th>Lateral bow at birth/at surgery (˚)</th>
<th>Anterior bow at birth/at surgery (˚)</th>
<th>LLD* at surgery (cm)</th>
<th>Proximal overlength of fibula (cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>6</td>
<td>45/22</td>
<td>40/15</td>
<td>5.5</td>
<td>2.3</td>
</tr>
<tr>
<td>2</td>
<td>5</td>
<td>45/15</td>
<td>38/14</td>
<td>4.4</td>
<td>1.8</td>
</tr>
<tr>
<td>3</td>
<td>5</td>
<td>60/23</td>
<td>60/5</td>
<td>3.1</td>
<td>2.8</td>
</tr>
</tbody>
</table>

*leg-length discrepancy

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**Table II.** The end results of operative lengthening of the tibia show that slight overlengthening was achieved. The healing indices were within the normal range.

<table>
<thead>
<tr>
<th>Case</th>
<th>LLD* at surgery (cm)</th>
<th>Amount of lengthening (cm)</th>
<th>Resulting overlength (cm)</th>
<th>Days in frame</th>
<th>Healing index (days/cm)</th>
<th>Recurrence of deformity at follow-up</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>5.5</td>
<td>6.5</td>
<td>1.0</td>
<td>273</td>
<td>42</td>
<td>No</td>
</tr>
<tr>
<td>2</td>
<td>4.4</td>
<td>5.0</td>
<td>0.6</td>
<td>205</td>
<td>41</td>
<td>No</td>
</tr>
<tr>
<td>3</td>
<td>3.1</td>
<td>3.8</td>
<td>0.7</td>
<td>140</td>
<td>37</td>
<td>No</td>
</tr>
</tbody>
</table>

*leg-length discrepancy
Discussion

Congenital anterolateral bowing of the tibia is a rare deformity reported to occur in one in 140,000 live births. In 1949, Heyman and Herndon classified tibial bowing according to the direction of the deformity. Although congenital anteromedial and posteromedial tibial bowing tends to resolve spontaneously during growth, anterolateral bowing is a more serious deformity. Congenital pseudarthrosis of the tibia is rarely observed at birth. It usually develops during the first decade of life and may present as a fracture through the highly sclerotic or atrophic bone in the tibia. Involvement of the fibular bone in the deformity is reported to occur in 62% of cases and can be important in the diagnosis of congenital pseudarthrosis of the tibia when the radiological findings in the tibia are inconclusive.

Congenital anterolateral tibial bowing with spontaneous resolution of the deformity with or without involvement of the fibula has been reported. Medial endosteal thickening extending across the tibial bow has commonly been reported. None of our patients showed cystic, extensive sclerotic, or other degenerative signs. No associated deformities of the foot have been reported.

The association of ipsilateral duplication of the hallux with anterolateral tibial bowing was first reported by Kardon et al in 1986. Subsequently, Adamsbaum et al, Weaver et al, Bressers and Castelein, and Kitoh et al added similar cases with greater detail of the radiological changes in the tibia. In cases of anterolateral tibial bowing without associated deformities of the foot, reactive postero-medial endosteal thickening was commonly reported. The
The deformity was noted to be most severe at birth and spontaneous resolution occurred with growth. Nevertheless, the apex of the tibial deformity showed some irregularity. Watanabe, Fujita and Oka\textsuperscript{17} and Adamsbaum et al\textsuperscript{7} reported a short incomplete tibial duplication based on their CT findings. Weaver et al\textsuperscript{3} and Bressers and Castelein\textsuperscript{8} thought that the radiological signs of duplication were the result of remedullarisation of the tibial canal. Other authors observed a partial cleft in the tibia in the area of the bow on CT\textsuperscript{6,7,15} and suspected incomplete duplication. In the long term, this area of uncertainty at the apex of the tibial bow disappears during resolution of the bow. In our patients MRI revealed a partial cleft, and plain radiography showed possible tibial duplication, but could not confirm true duplication. However, during surgery, true tibial duplication was seen with two distinct cortices surrounding both tibial medullary canals (Fig. 4). This pattern of duplication regressed during growth, and a single well-defined medullary canal developed.

In our cases as in all previous reports of congenital anterolateral tibial bowing in association with duplication of the hallux, the fibula was not involved.\textsuperscript{4,6,8} It was straight and appeared structurally uninvolved in the pathomorphological changes affecting the tibia and foot.

Proximal overgrowth of the fibula compared with the ipsilateral tibia led to subluxation of the proximal tibiofibular joint but overall the fibula was slightly shorter than the contralateral fibula. This may be related to ligamentous tethering due to the relative shortening of the tibia.

In our three patients, the tibia was hypoplastic distally with a diminished width of the epiphysis. The association of preaxial polydactyly and tibial deficiencies is well known.\textsuperscript{18} Lewin and Opitz\textsuperscript{19} described a tibial and fibular field during the development of the lower limb. The tibial field involved the distal femur, tibia, and preaxial toes.

With regard to previous reports of associated foot deformities, our cases showed similar duplication of the hallux. We agree with other authors that surgical treatment of this duplication is difficult.\textsuperscript{3} Foot problems persisted in our patients because of pes planus and severe shortening of the first ray. Our patients also had persistent problems with shoe wear due to widening of the midfoot associated with partial duplication of the navicular, cuneiform, and metatarsal bones. This deformity has not been previously described but may be typical of developmental disorders affecting the tibial field.

Previous authors have recommended conservative treatment since spontaneous complete resolution of the anterolateral bowing during growth can occur.

However, this does not take account of the leg-length discrepancy and subluxation of the proximal tibiofibular joint. In our patients, spontaneous resolution of the tibial bow was observed to occur during the first four years of life, but then persisted to a certain degree leading to foot problems. All our patients complained of intermittent pain in the foot and at the proximal tibiofibular joint before surgery. Reconstructive surgery was indicated for a leg-length discrepancy of up to 5.5 cm, persistent anterolateral bowing and proximal subluxation of the fibula. Even when corrective osteotomy was performed in the area of the bow, healing indices were comparable with those in the literature for lengthening procedures in structurally normal bone. The deformity had not recurred at the latest follow-up, but intermittent pain in the foot persisted. We are therefore in agreement with Weaver et al\textsuperscript{8} that the deformity of the tibia is relatively benign whereas the deformity of the foot is not and may be the most important factor affecting the long-term outcome in this condition.

No benefits in any form have been received or will be received from a commercial party related directly or indirectly to the subject of this article.

References