Is the Apert foot an overlooked aspect of this rare genetic disease? Clinical findings and treatment options for foot deformities in Apert syndrome

Alexandra Stauffer and Sebastian Farr

Abstract

Background: Apert syndrome is characterised by the presence of craniosynostosis, midface retrusion and syndactyly of hands and feet, thus, synonymously referred to as acrocephalosyndactyly type I. Considering these multidisciplinary issues, frequently requiring surgical interventions at an early age, deformities of the feet have often been neglected and seem to be underestimated in the management of Apert syndrome. Typical Apert foot features range from complete fusion of the toes and a central nail mass to syndactyly of the second to fifth toe with a medially deviated great toe; however, no clear treatment algorithms were presented so far. This article reviews the current existing literature regarding the treatment approach of foot deformities in Apert syndrome.

State-of-the-art topic review: Overall, the main focus in the literature seems to be on the surgical approach to syndactyly separation of the toes and the management of the great toe deformity (hallux varus). Although the functional benefit of syndactyly separation in the foot has yet to be determined, some authors perform syndactyly separation usually in a staged procedure. Realignment of the great toe and first ray can be performed by multiple means including but not limited to second ray deletion, resection of the proximal phalanx delta bone on one side, corrective open wedge osteotomy, osteotomy of the osseous fusion between metatarsals I and II, and metatarsal I lengthening using gradual osteodistraction. Tarsal fusions and other anatomical variants may be present and have to be corrected on an individual basis. Shoe fitting problems are frequently mentioned as indication for surgery while insole support may be helpful to alleviate abnormal plantar pressures.

Conclusion: There is a particular need for multicenter studies to better elaborate surgical indications and treatment plans for this rare entity. Plantar pressure measurements using pedobarography should be enforced in order to document the biomechanical foot development and abnormalities during growth, and to help with indication setting. Treatment options may include conservative means (i.e. insoles, orthopedic shoes) or surgery to improve biomechanics and normalize plantar pressures.

Level of evidence: Level V.

Keywords: Apert syndrome, Foot deformity, Treatment options

* Correspondence: sebastian.farr@oss.at
Department of Pediatric Orthopaedics and Foot and Ankle Surgery, Orthopedic Hospital Speising, Speisingerstrasse 109, 1130 Vienna, Austria

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Background
Apert syndrome [1] is characterised by the presence of multisuture craniosynostosis, midface retrusion and syndactyly of hands and feet, thus, synonymously referred to as acrocephalosyndactyly type I. This disorder occurs in 6 to 15.5 out of 1 million livebirths depending on the study cited, with an almost equally affected sex ratio [2–4]. The prevalence may be higher amongst children born to fathers with advanced paternal age, as the greatest variant of sperm mutations occurs in fathers above 60 years of age. Furthermore, there seems to be a paternal-age effect, as children with Apert syndrome are born to fathers who exhibit an increased frequency of mutant sperm at an earlier age [5]. Apert syndrome results from mutations localised on the FGFR2 caused by specific missense substitutions involving adjacent amino acids, thus resulting in two genotype-phenotypes consisting of Pro253Arg, causing more significant hand and foot involvement, and Ser252Trp [6].

Diagnosis is established in a patient clinically or by genetic testing resulting in the identification of heterozygous pathogenic variant of the FGFR2 and the presences of clinical features consistent with Apert syndrome. Almost all patients present with craniosynostosis commonly involving bilateral coronal sutures, however, a majority also displays a progressive fusion of the sagittal and lambdoid sutures. Unlike other acrocephalosyndactyly syndromes, the midface in Apert syndrome is underdeveloped, contributing to the development of prominent eyes with downslanding palpebral fissures, as well as retruded midface, thus leading to a shorter maxillary bone. A multitude of other abnormalities have been described in Apert syndrome affecting inner organs and other structures, such as spinal fusions, palatal and/or dental abnormalities, ocular abnormalities, hearing loss or inner ear anomalies, as well as multilevel airway obstructions, cardiac abnormalities, gastrointestinal issues, abnormalities of the genitourinary tract and various neurologic findings [7].

Considering these multidisciplinary issues, frequently requiring surgical interventions at an early age, deformities of the feet have been forced into the background and seem to be often overlooked in the management of Apert syndrome. Although these typical clinical appearances in Apert syndrome regarding feet have been described in multiple case reports ranging from complete fusion of the toes and a central nail mass [8] to syndactyly of the second to fifth toe with a medially deviated great toe [9], no treatment algorithms were presented so far. The purpose of this state-of-the-art topic article is to review the current existing literature regarding the approach to the treatment of foot deformities in Apert syndrome.

The Apert foot and its treatment
Overall, the main focus in the literature seems to be on the surgical approach to 1) syndactyly separation of the toes and 2) the management of the great toe deformity. The following part is divided into sections addressing these main topics, as well as joint fusions and anatomical variants.

Syndactyly of the toes
Three types of syndactyly of the toes have been described by Blauth et al. [10]. In a study which included 37 feet, Cohen et al. reported the predominant type being type III which was present in 54% of their patients. In type I, a fusion of the second, third and fourth toe is present, with one and five being separate. Other findings include a broad great toe which may shorten over time and deviate medially. According to Cohen et al. this type occurs in 27% of analysed cases. Type II (19% of cases) consists of a fusion which involves all toes except for the great toe, resulting in a supination of the forefoot (Fig. 1a). The position of the great toe is in varus and dorsiflexion. A fusion of all toes with a possibly hyperextended great toe is present in type III [11].

Fearon et al. proposed a two-stage approach as initial treatment of syndactyly release. Both stages consist of syndactyly release, however, stage one is performed at around 12 months of age, followed by the second stage 3 months later. The digits are separated distally to proximally through Z-plasties, and all distal bony fusions are divided to create normal phalangeal width, the neurovascular bundles are carefully separated, the skin is approximated and full-thickness skin grafts, which were harvested from the groin, transplanted into open spaces along the phalanges if required. The late stage is performed between the ages of 9 to 12 if necessary and includes osteotomies to create a permanent angulation of the digits in order to achieve a better functional position. This pertains to symptomatic foot disorders, which can be corrected by osteotomy of either the first or fifth metatarsal head to narrow the affected area. Elevation of the second metatarsal head through osteotomy and reangulation may be necessary if it becomes overly prominent on the plantar surface [12].

Upton et al. suggested separation of the toes at an early age, as interdigital skin grafts require immobilization to heal. They noticed that the skin transplants did not heal well for active children, regardless of whether or not they were treated with casts or splints. Furthermore, they noticed that the severity of the foot deformity did not correlate with the extent of hand deformity, especially in patients with type II and especially type III hands [13].

As a result of an osseous syndactyly subsequently changes in the nailbed may occur, the most frequent
change being an acneiform eruption. Other occurrences include complete nail fusion with a single wide nail, central indentations, short nails with absent lunulae, and micronychia [14]. Narrowing of the nail may be achieved through fulguration of the lateral germinal matrix with electrocautery [12]. Other cutaneous changes include the formation of painful calluses due to altered weight bearing mechanics, especially over the protuberant second and third metatarsal heads [15]. Osteotomies of the metatarsals for stress relief have been reported in literature [16].

Deformities of the great toe
Deformities of the great toe in patients with Apert syndrome were described by Dell et al. in 1981 [17]. Their study included case reports of three children aged between 2 and 3 years of age who showed significant difficulty in shoe fitting which lead to the decision to
operate rather than to accept the aesthetic appearance of the foot. The first case presented with complex syndac-
tilies of all toes and the presences of five toe-nails.
Weight bearing took place on the lateral four rays (Fig.
1b), as planter pressure of the first ray decreases with
shortening [17]. Radiographs revealed a complex syndac-
tyly between the first and second metatarsal with the
presence of osseous fusion, as well as a delta phalanx proximally and an abnormally large and misshapen distal
phalanx of the great toe. In order to correct this deform-
ity, realignment of the great toe was performed by
means of second ray deletion, resection of the delta
phalanx on one side and corrective open wedge osteot-
omy on the other, furthermore, osteotomy of the osse-
ous fusion and narrowing of the distal phalanx by
longitudinal osteotomies were performed, as well as fix-
ation of realignment of the great toe using K-wire, which
were placed across the metatarsal-tarsal joints, and Z-
plasty.

The second case presented with bilateral hallux varus
deformities with shortening of the great toe, as well as
syndactyly of the second, third and fourth toe, respect-
ively. Radiographic findings included duplication of the
proximal phalanx and a widened distal phalanx in the
great toe. Measures to narrow the feet were undertaken,
which included resection of the great toe and transpos-
tion of the second ray medially.

The last case presented with syndactyly of all toes, bi-
lateral fusion of the first and second metatarsal, duplica-
tion (Fig. 1c) of the distal phalanx of both second toes
and a medial angulation of the great toe. However, no
shoe fitting problems were described by the parents,
thus, no surgical intervention took place [18].

While Dell et al.’s approach to the realignment and
correction of the great toe included a deletion of the
second ray, other authors developed a different tech-
nique which allows for the preservation of the second
ray. In a patient with Apert syndrome who presented
with synostosis between the first and second ray, bi-
lateral hallux varus, as well as metatarsus adductus of
the second, third and fourth ray, Blauth et al. pro-
posed a different surgical approach. They performed a
resection of the osseous fusion between the first and
second metatarsal, carried out an osteotomy of the
second and third metatarsal, completed the correction
of hallux varus by means of resecting the proximal
rudimentary bone fragment and accomplished the re-
alignment of the great toe using K-wire as a treat-
ment option [19].

As the great toe shortens and broadens over time
compared to the rest of the foot and the presence of a
delta phalanx may lead to medial deviation and varus
angulation, thus surgical measures such as correction of
brachymetatarsia and medial angulation may be
necessary. Calis et al. reviewed the long-term outcome
of distraction osteogenesis (Fig. 1d) for this deformity in
14 Apert feet. They performed an osteotomy of the first
metatarsal sparing the epiphyseal plate and inserted two
pins proximal and distal to the osteotomy site, which
were then connected to an external distractor device.
Distraction protocol consisted of initiation 5 days post-
operatively and a distraction rate of 0.5 mm per day. The
mean distraction period observed was 12.6 ± 1.7 days,
after a mean consolidation period of 62.6 ± 5.8 days, dis-
tractors were removed. Even though they observed com-
lications such as pin loosening, pin infection and early
union requiring revision surgery, Calis et al. reported a
correction of brachymetatarsia long-term in all patients
[20]. To this day, some surgical measures as performed
by the authors mentioned above are still considered
when formulating an operative plan for patients with de-
formities of the feet in Apert syndrome (Fig. 1e).

Joint fusions and other anatomical variants
Schauerte et al. reported on bony fusions in 1966, ob-
serving that a fusion between the tarsal bones was the
most constant finding in patients with Apert syndrome.
They reported on cases showing progressive synostosis
of the tarsal bones, in particular a fusion of the calca-
neus and the cuboid, as well as fusions of the 3rd meta-
tarsal with the lateral cuneiform or the navicular with
the medial cuneiform [21].

Upton et al. proposed a classification into two groups,
depending on the extent of fusion between the first two
rays. In type I feet, the separation between the cuneiform
bones and the first metatarsal are present at birth. Type
II feet displayed more severe skeletal deformities, as fu-
sions between the cuneiform bones and the navicular
were present, as well as complete or partial fusions with
the second metatarsal [22].

Collins et al. used computer tomography (CT) scans
to define the spatial dysmorphology of the foot in a
series of case reports of patients with Apert syndrome.
They observed consistent anatomical findings in all three
patients who underwent computer assisted medical im-
aging. All pairs of feet showed abnormal great toes
with phalangeal and metatarsal changes, syndactyly of
the toes 2 to 5, bony fusions between metatarsals, tarsal
coalignments and limitations in shoe fitting. Case 1 was a 2-
year-old boy who showed closely approximated calca-
neocuboidial joints on CT scan which had not yet
reached the stage of definite bony bridging, however, the
approximation might suggest the presence of cartilagin-
ous or fibrous fusion. Case 2, a 4-year-old girl presented
with a bilateral calcaneocuboidal fusion in the hindfoot.
The feet of a 9-year-old boy displayed a synostosis of the
hindfoot with talonavicular, talocalcaneal and calcaneo-
cuboidal fusions on CT scan [23].
Carranza-Bencano et al. described a case of a 9-year-old boy with Apert syndrome who presented with calcaneo-cuboid and cuneonavicular fusion in addition to complex syndactyly of all toes and a shortened great toe with medial deviation [24].

Anderson et al. described other anatomical variants of the Apert foot in various case reports. Radiographs revealed additional metatarsal bones in a patient who appeared to have typical feet associated with Apert syndrome upon clinical examination [25]. Another case presented with asymmetrical findings on clinical examination, as there were only four toes present in the right foot compared to the left; however, complex syndactyly was present in both feet. The radiographs, however, showed all metatarsals present with various phalanges of different toes missing in the right foot. Another case presented with similar findings on clinical examination, radiographs, however, showed a synostosis of the proximal phalanx of the third and fourth toes. These case reports strongly reinforce the need for radiographic evaluation of both hands and feet in patients with Apert syndrome in order to determine the underlying bone composition [26].

Conservative approach
Blauth et al. mentioned orthopaedic arch support insoles of the first and second metatarsal for elevation of pressure points on the third metatarsal head in Apert foot. Furthermore, orthopaedic shoes are an option to help with discomfort caused by lateral weight bearing or great toe deformities [19]. Pedobarography should be used for the identification of locations with elevated pressure in order to make a valid foot care decision regarding support insoles, as Guldemond et al. showed that clinical evaluation overestimated the pressures in the metatarsal region, whereas underestimating the ones in the great toe [27]. No other relevant information concerning further conservative treatment strategies was found in our literature search.

Treatment recommendations
Children with Apert syndrome should receive early pediatric orthopaedic attention to assess major deformities of the hands and feet. In the latter, bone and soft tissue alterations should be noted and addressed in a staged manner (Fig. 2). Given the fact that craniosynostosis are usually given priority in the first year of life, extremity surgery, if necessary, is usually performed thereafter. The main deformities to be considered for treatment are forefoot adduction deformity, syndactylies, first ray/great toe shortening and malalignment, and hindfoot fusions/bony variants. Based on our personal experience, casts, splints and/or orthoses may be helpful in cases with rigid forefoot adduction and/or supination deformity to achieve a plantigrade foot position. This should be corrected ideally before walking age or latest until the end of the second year. We usually separate syndactylies only if the great toe is involved; we do not see any clinical or true cosmetic advantage in syndactyly separation of the other lesser toes, especially since the rate of local infection and keloid formation is rather high. The children are allowed to wear conventional shoes; however, in pre-school age, thorough clinical, radiographic and pedobarographic assessment is pursued to check for abnormal biomechanics and pressures, and anatomical variants. The shortening of the first ray is usually addressed in pre-school to school-aged children if symptoms and/or markedly elevated plantar pressures are present. However, this surgical intervention, whether performed acutely or in a gradual manner by
Pedobarography is an essential tool to help with indication setting for insole prescription, orthopaedic shoe-wear or surgery. Hindfoot deformities and variants such as coalitions or bony fusion may pose a disturbing problem but have to be cautiously addressed on a highly individualized basis. In our experience, many of these findings do not need any specific treatment.

**Conclusions**

Due to the multidisciplinary aspect of Apert syndrome, patients undergo multiple surgical corrections primarily focusing on the correction of craniosynostosis and midface retrusion at an early age [28]. Although various surgical options for the correction of the Apert foot have been described in literature, mainly focusing on the correction of syndactyly of the toes, as well as approaches of the problem caused of anomalous great toes, no management protocol for the Apert foot has been established thus far. In the past, surgical interventions were reserved only for patients complaining of functional impairment or difficulties when fitting shoes, however, as the life expectancy of patients with Apert syndrome increases these problems may become more present in medical practice. Therefore, research regarding treatment options for the feet is essential in providing optimal care and improvement of quality of life for children born with this rare condition. Pedobarography is capable to highlight areas of increased loading during gait and should be used routinely to monitor development during growth. Based on the clinical findings and symptoms during childhood conservative and/or operative means are followed.

**Abbreviations**

FGFR: Fibroblast growth factor receptor; CT: Computer tomography

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Not applicable.

**Consent for publication**

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