Magnetic resonance imaging confirmed clinical diagnosis of amynoplasia in two infants with arthrogryposis multiplex congenita

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ABSTRACT

Introduction: We present two cases of arthrogryposis multiplex congenita (AMC) with involvement of the lower extremities. In both cases amynoplasia was confirmed by a magnetic resonance imaging (MRI). The degree of amynoplasia correlated with the severity of arthrogryposis and determined the child’s prognosis. Case Series: Case 1 was a 16-month-old male child with prenatally diagnosed Klinefelter syndrome who was born at 36 weeks gestation. Brain MRI was normal. Joint rigidity was detected in the upper and lower extremities. Amynoplasia was suspected at nine months of age since the lower limb muscles were hardly palpable. Case 2 was a 5 1/2-month-old female child and the first child of non-consanguinuous parents was noticed to have rigid right calcaneovalgus and left equinovarus feet deformities as well as knee rigidity with limitation of knee extension. Bilateral hip displacement was also diagnosed. Absence of muscles on thigh palpation prompted MRI study. Conclusion: Although amynoplasia is the most common type of arthrogryposis multiplex congenita, muscle underdevelopment in these patients remains puzzling for pediatric practitioners. Amynoplasia congenita is usually symmetrical and involves either all extremities or selectively only the lower or upper extremities. Absence of muscle groups on MRI confirms diagnosis of amynoplasia. Early recognition of amynoplasia in children with arthrogryposis multiplex congenita can help in tailoring their treatment and prognosis.

Keywords: Amynoplasia, Arthrogryposis multiplex congenita (AMC)

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INTRODUCTION

Arthrogryposis multiplex congenita (AMC) affects about 1/3000 birth in North America [1]. Amynoplasia (A=no, myo=muscle and plasia=growth) is the most common type of arthrogryposis seen clinically [2]. We present two cases of arthrogryposis multiplex congenita with prominent involvement of the lower extremities. In both cases amynoplasia was confirmed by magnetic resonance imaging (MRI) of the thighs. The degree of amynoplasia correlated with the severity of arthrogryposis and determined the child’s prognosis. MRI may be helpful in the differentiation between amynoplasia and other congenital myopathies and muscular dystrophies [3].
CASE SERIES

Case 1: The patient was a 16-month-old male baby, prenatally diagnosed with Klinefelter syndrome born at 36 weeks gestation from a non-consanguineous marriage. The delivery was by emergency cesarean section due to fetal heart rate deceleration. Apgar scores were 1, 2 and 5 at 1, 5 and 10 minutes. During the first two days of life the baby developed seizures. Brain MRI was reported as normal. Joint contractures were recognized on the fourth day of the baby’s life. Amyoplasia was suspected at nine months of age since the lower limb muscles were barely palpable. MRI of the thigh revealed paucity of muscles (Figures 1 and 2). On examination, the baby presented with microcephaly (head circumference 44 cm, <2%), bilateral epicanthus and clinodactyly of the fifth fingers. The child was able to hold sitting position, however, he was not able to sit up by himself. There was stiffness in the wrists with limitation of wrist extension. Hand grasp was weak. The child was unable to hold a spoon. Hip abduction was limited to 30°. Knee flexion was limited to 90°. There was also equinovarus feet deformity. The child moved the upper extremities well, but only slightly raised the extended lower extremities and minimally flexed them at the knees. He supported his weight on his legs when in standing position with support. In the lower extremities only hip adductors were slightly palpable. Gluteus muscle contraction was evident. Biceps, brachioradial and knee reflexes were normal. Plantar response was down-going. There was no deficiencies in sensation to the touch and pinprick.

Case 2: A 5'/2-month-old female was the first child of non-consanguineous parents was born after full term uneventful pregnancy by normal vaginal delivery. Apgar score was 9 and 9 at 1 and 5 minutes. Birth weight was 3.8 kg. At birth the baby was noticed to have rigid right calcaneovalgus and left equinovarus feet deformities, as well as knee rigidity with limitation of knee extension. Bilateral hip displacement was also diagnosed. Current neurological examination revealed peripheral right facial palsy and torticollis due to shortened right sternocleidomastoid muscle. The upper extremities were without any neurological deficiencies. The lower extremities were in fixed frog-leg position with rigid feet deformity. Active movements manifested with very slight hip adduction and minimal toe movements. The muscles were not detectable on leg palpation. Knee and ankle jerks were absent. Plantar response was mute. Sensation to pinprick and touch was preserved throughout. The anus was closed and anal blink reflex was brisk. The baby demonstrates good head control, raising the head and chest while being in prone position; grasping and transferring an object from one hand to another with good visual attention to the grasped object. The baby had positive stranger anxiety. MRI of the hips showed absence of musculature with preserved fascial planes, vessels, and adipose tissue (Figure 3).

As part of management, both patients underwent surgical correction of their tulip equinovarus. On follow-up at three years of age, their neurological evaluation showed that they both were able to sit up, but not stand up. Only the first patient was able to stand with support.

DISCUSSION

Despite the fact that amyoplasia is the most common type of arthrogryposis multiplex congenital [2, 3], recognition of considerable muscle underdevelopment in these patients remains surprising and puzzling for pediatric practitioners. Amyoplasia congenita is usually

Figure 1: Magnetic resonance imaging of the thigh revealed paucity of muscles in patient from Case 1.

Figure 2: Another magnetic resonance imaging of the thigh revealed paucity of muscles in patient from Case 1.

Figure 3: Magnetic resonance imaging of the hips showed absence of musculature in patient from Case 2.
symmetrical and involves either all extremities or selectively only the lower or upper extremities [4]. The muscle mass of the limbs with arthrogryposis is diminished. In a study by Hall et al., histologic examination of muscles showed the replacement of muscle with fibrofatty scar tissue [5]. Studies of spinal cord and muscles in patients with amynplasia suggests two possible affected areas: either anterior horn cells or muscles [6, 7]. Intrathecal vascular insult of spinal motor neurons or limb muscles was hypothesized [8, 9].

It is also known that arthrogryposis can be a genetically heterogeneous disorder as for instance, in distal arthrogryposis. Our first case can be considered as an arthrogryposis with all limb involvement even though the wrist stiffness is the only finding in the upper extremities. The presence of a brisk knee jerk does not support the spinal origin of amynplasia. Decrease of femoral muscle mass on thigh MRI is significant, but the leg extensors remained strong enough to bear body weight. Klínefelder syndrome diagnosed in this child most likely does not have any causal relation with the amynplasia. Our second case presented with arthrogryposis only in the lower extremities. MRI showed absence of femoral muscles bilaterally. Congenital facial palsy in this child is considered as a partial Moebius anomaly, which is a known comorbidity of amynplasia congenita.

Muscle weakness and multiple joint contractures in both patients could have suggested congenital muscle dystrophy (CMD) and congenital myopathy (CM). However, MRI revealed absence of whole muscle groups, with fibrofatty tissue instead and preserved fibrous planes. In patients with CMD and CM, MRI shows peculiar selectivity of muscle involvement, with decreased muscle volume and abnormal MRI signals from the muscles [3].

CONCLUSION

Early recognition of amynplasia with an assessment of muscle underdevelopment via limb magnetic resonance imaging in children with arthrogryposis multiplex congenita elucidates their clinical presentation and can help in tailoring their treatment and in the prognostication of the degree of their future disability.

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Author Contributions

Ariam Diaz – Conception and design, Acquisition of data, Analysis and interpretation of data, Drafting the article, Critical revision of the article, Final approval of the version to be published

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Valerie May G Sia – Conception and design, Acquisition of data, Analysis and interpretation of data, Drafting the article, Critical revision of the article, Final approval of the version to be published

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Guarantor

The corresponding author is the guarantor of submission.

Conflict of Interest

Authors declare no conflict of interest.

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