Atypical Carpal Tunnel Syndrome in a Holt Oram Patient: A Case Report and Literature Review

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Abstract: We present a case report of a patient diagnosed with Holt-Oram syndrome (HOS) presenting with clinical and electrophysiologically confirmed carpal tunnel syndrome. Pre-operative Magnetic resonance imaging revealed an abnormal course of the median nerve; as such an atypical incision and approach were carried out to decompress the nerve to excellent post operative clinical effect. To our knowledge this is the first description of abnormal nervous course in a patient with HOS leading to peripheral entrapment. A literature surrounding the important aspects of HOS to the orthopaedic surgeon is presented concomitantly.

Keywords: Carpal tunnel syndrome, congenital hand deformity, holt oram syndrome, median nerve variation, radial deficiency, thumb hypoplasia.

CASE REPORT

A 27 year old gentleman presented to orthopaedic outpatient clinic with the principal complaint of paraesthesia over the right index finger, worse overnight and causing sleep disturbance. In addition he had noted a recent reduction in grip strength. His past medical history was that of congenital bilateral thumb hypoplasia, for which he had undergone left index finger pollicisation and excision of the rudimentary thumb to his right hand. Both his mother and maternal grandfather had been diagnosed with HOS.

Examination revealed decreased sensation over the index finger and hypoplasia of the thenar eminence. Provocation with Tineal’s test over the carpal tunnel and with a modified Phalen’s test were both negative. He was referred for electrophysiologic nerve examination, which revealed marked conduction abnormalities of the median nerve. In light of his abnormal provocative tests and anatomy a magnetic resonance imaging (MRI) scan was requested to delineate the carpal tunnel and median nerve anatomy.

The MRI scan revealed a very radial lying median nerve within the carpal tunnel and scaphoid hypoplasia, it was postulated that the abnormal lie of the median nerve could be in relation to the observed scaphoid hypoplasia (Fig. 1).

The patient underwent carpal tunnel decompression under general anesthesia with a more radial incision. The transverse carpal ligament was released and the nerve explored, where it was found to be located radially within the tunnel and illustrated signs of chronic compression.

At review, 3 months post operatively the patient’s symptoms had dramatically improved with a resultant high satisfaction with the outcome of surgery.

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Fig. (1). Magnetic resonance scan detailing radially located median nerve.

LITERATURE REVIEW

In 1960, Holt and Oram published their description of family members over 4 generations with 9 affected subjects, describing upper limb abnormalities in conjunction with congenital heart abnormalities [1]. Knowledge and understanding of Holt-Oram syndrome (HOS), or Heart-Hand syndrome, has progressed; its diagnosis should be a consideration in the assessment of all patients presenting with radial longitudinal deficiencies [2].

INCIDENCE AND PATHOGENESIS

Population studies reveal congenital hand anomalies have an overall incidence of between 19.7 - 21.5 per 10,000. HOS is still uncommon, estimated to affect 1 in 100 000 live births, shows autosomal dominant inheritance (40% sporadic
Table 1. Common causes of radial deficiencies.

| Syndrome          | Inheritance          | Associated Conditions                        | Diagnosis                                                                 |
|-------------------|----------------------|----------------------------------------------|---------------------------------------------------------------------------|
| Thrombocytopenia absent radius | Complex Autosomal Recessive [13, 14] | Absent radius with normal thumb Thrombocytopenia and anaemia (improves in first year of life) | Absent radii, thumbs present and platelet count < 50x10^10/L |
| Fanconi’s Anaemia | Autosomal Recessive  | Pancytopenia between 5-10y of age            | Early diagnosis cal be made using chromosomal breakage studies (diepoxybutane or mitomycin C) |
| VACTERL           | Sporadic             | V ➔ Verbebral A ➔ Anal atresia C ➔ Cardiac anomalies TE ➔ Tracheosophageal fistula R ➔ Renal agenesis L ➔ Limb anomalies | Clinical Exclusion of overlapping syndromes [15] |
| Holt-Oram         | Autosomal dominant   | Cardiac conduction or structural anomalies Generalised upper limb anomalies | Clinical (TBX5 genotyping (Ch12)*) |

*Up to 74% sensitivity reported [9].

new mutation), variable expression and near complete penetrance [3-8]. Strict clinical criteria are required to reduce the, frequently reported, risk of incorrect or missed diagnosis [4, 9]. Basson et al. described, and validated, diagnostic phenotypes as the presence of pre axial radial ray malformation of at least one upper limb along with a personal or family history of septation defects and/or atriventricular disease, these criteria are widely supported [8-11]. There are several syndromes presenting commonly with preaxial radial malformation patterns, they are summarised in Table 1. It is recommended that all patients should undergo coordination with the patient’s physician/paediatrician a full blood count, renal ultrasound and echocardiogram once a radial longitudinal deficiency is noted [2, 4, 12].

The underlying genetic driver of HOS has been linked to the TBX5 gene of the Chromosome 12q2 region [3, 8-10, 16]. TBX5 is a part of the T-box gene family which serves to encode a number of transcription factors important during embryonic development; in particular the differentiation of cardiomyocytes where it aids the formation of electrical conduction pathways. Alongside this is the regulation of patterning and morphogenesis within the upper limb [3, 9, 10]. Through noting differing phenotypes in monozygotic twins Huang has suggested the relationship of HOS to this gene site may be complex and subject to other, as yet identified, factors ultimately giving rise to individual phenotypic expression [8].

PRESENTATION AND TYPICAL FEATURES OF HOS

The classical presentation is that of a tripahalneal thumb with a secundum atrial septal defect (ASD) [17]; however, the syndrome yields wide phenotypic variability, as previously stated unusually even in monozygotic twins [8]. Upper limb abnormalities can involve the entire extremity, are often bilateral but may not be symmetrical [9, 18]. Abnormal shoulder girdle function, phocomelia, hypoplasia of pectoralis major, clavicular defects, hypoplasia of the humerus, radius, ulnar, metacarpals, syndactaly, carpal anomalies and digital malformations ranging from dysplasia to absence have all been described in the literature [3, 19-23]. The thumb remains the most commonly affected digit, in their 70 year analysis of 77 patients with thumb hypoplasia James et al. found an incidence of HOS in this population of 16% [24]. A series of 55 HOS patients reported by Newbury-Ecob et al. yielded salient clinical observations, they described an 84% rate of thumb abnormality, found ulnar sided defects only in the presence of concomitant radial pathology, stated proximal reduction defects occurred only when associated with distal, with the exception of the shoulder girdle, in which abnormalities were common (77%) [18]. The characteristic shoulder pattern observed is that of narrow sloping shoulders, caused by hypoplasia of the humeral head, clavicle and surrounding musculature [18, 25].

Cardiac abnormalities range from the most common ASD, to include conduction defects and the Teratology of Fallot [8, 18, 26, 27]; often there is no correlation in the severity of cardiac and upper limb abnormalities [26]. Bruneau et al. categorised the cardiac defects in a series of 240 patients; they found 58% of patients had an atrial septal defect, 28% a ventricular septal defect and conduction defects in 18% [16], less common pathologies (including tetralogy of Fallot, truncus arteriosus, mitral valve defects and patent ductus arteriosus) were also present. These often complex cardiac manifestations may require cardiac catheterisation to fully define and subsequent multidisciplinary management prior to considering administration of anaesthesia and subsequent operative intervention [4, 8].

NON SKELETAL AND CARDIAC CONSIDERATIONS

With the majority of literature reporting skeletal upper limb anomalies one must not assume all the soft tissues are normal; muscular deficiencies, nerve anomalies (hypertrophic median nerve) and vascular anomalies have all
been associated with radial longitudinal deficiencies [2]. Frota Filho et al. [28] reported a case of HOS with absent radial and ulnar pulses, but good hand perfusion. In reporting the outcomes of 74 patients undergoing index finger pollicization Lochner et al. reported a higher occurrence of physical arrest in HOS (46%) compared to the entire group (25%); they speculated the vascularity of these digits were at greater risk due to potential aberrant size or location of digital vessels [29]. Shono et al. noted hypoplastic vasculature during femoral venous cannulation and recommended this be performed under ultrasound guidance [21]. These feelings were reinforced by Sing et al. who also cautioned of the reliability of non invasive blood pressure monitoring in the face of abnormal vasculature [30].

There has been only one case report we have found in the literature describing carpal tunnel syndrome in association of HOS, occurring as a result of an aberrant FDS muscle belly found within the carpal tunnel at decompression [31]. Thenar hypoplasia has been noted to occur in HOS in the absence of clinical median nerve dysfunction [18, 25]. Spranger et al. [25] investigated upper limb muscle morphology in 13 patients with HOS using MRI and conducted electrophysiological testing on 4. All patients had normal serum muscle enzyme levels. Whilst clinically observed pattern of muscle weakness varied, paresis of thumb opposition and abduction was universal. MRI scanning revealed isolated hypoplasia, but normal structural pattern in muscles correlating with clinical weakness. In addition normal electrophysiological testing results were reported in the muscles observed to be weak, supporting their clinical use for detecting peripheral compression in the setting of abnormal anatomy and function.

DISCUSSION

Holt Oram syndrome remains rare; the anatomical variations encountered in this population should alert the treating clinicians to be mindful of the often abnormal anatomy. In our case preoperative MRI scanning proved invaluable at demonstrating atypical anatomy of the median nerve and allowed for alteration of usual surgical incisions, approach and prompted the use of general anaesthesia to facilitate prolonged exposure of the median nerve had it become necessary. MRI has been used to define the normal anatomy and variation of the median nerve in 194 patients by Pierre-Jerome et al, they did not report the variation we present [32]. Lindley and Kleinert presented their findings of median nerve anatomy from 526 elective carpal tunnel decompressions, and again did not report a course of the median nerve similar to that observed in our case [33]. Thus we would recommend any patient with Holt Oram syndrome presenting with symptoms of upper limb peripheral nerve entrapment undergo electrophysiological evaluation to clearly define the location of nerve dysfunction, followed by MRI evaluation of the area to define the anatomy prior to surgical intervention. Should general anaesthesia be employed there are extra considerations and demands of the anaesthetic.

CONFLICT OF INTEREST

The authors confirm that this article content has no conflict of interest.

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