MURCS association: case report and review

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Abstract
We report on a 25 year old woman with aplasia of the Müllerian duct, unilateral renal agenesis, and anomalies of the cervicothoracic somites (MURCS association). Growth retardation and facial asymmetry were also present. A review of published reports allows MURCS association to be distinguished from related associations, sequences, and syndromes. Moreover, sporadic occurrence, the broad spectrum of associated anomalies, and the involvement of different organ systems closely related in early embryogenesis are arguments for considering MURCS association as the consequence of a developmental field defect.

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Key words: Mayer-Rokitansky-Küster-Hauser (MRKH) syndrome; MURCS; renal agenesis.

In 1979 Duncan et al.[1] reported the non-random association of Müllerian duct aplasia/hypoplasia, renal agenesis/ectopy, cervicothoracic somite dysplasia (MURCS), and various other anomalies in 30 cases. So far, 35 further reports of patients have been published.[2-17] The 25 year old woman is the third child of healthy, non-consanguineous parents. The pregnancy was uneventful and delivery was at term. Birth weight (2750 g) and length (50 cm) were within the normal range. The right kidney was absent and contralateral hydronephrosis required kidney transplantation in childhood.

On examination height was 152 cm (3rd centile), weight was 55 kg (50th centile), and occipitofrontal circumference was 58 cm (97th centile). Endocrinological investigations, karyotyping, psychomotor development, and secondary sexual characteristics were unremarkable. Laparoscopy displayed histologically normal but macroscopic cystic ovaries, a rudimentary uterus duplex, and a cord-like vagina without any lumen.

Facial asymmetry was obvious (fig 1). Both ears were normal in shape, location, and function. The teeth were unremarkable. X rays of the spine showed subtotal fusion of C2 and C3, hemivertebra C7, and bilateral C7 cervical ribs partly fused with the first thoracic ribs (fig 2). Bone age corresponded to the chronological age.

Aplasia of the Müllerian duct is also called Mayer-Rokitansky-Küster-Hauser (MRKH) syndrome. Strübbe et al.[18] found 13 cases with MURCS association out of 56 women with atypical MRKH syndrome, but none out of 44 women with typical MRKH syndrome.

Dysplasia of the cervical somites is also called Klippel-Feil sequence. Associated anomalies of the musculoskeletal, neurological, cardiovascular, and genitourinary system have been described,[19] but only one woman had in addition an anomaly of the upper urinary tract and vaginal agenesis with a bicornuate uterus.[20]

The aetiology of MURCS association is not known. The karyotype was normal in all investigated cases and so far all cases have been sporadic. Therefore, autosomal recessive inheritance is unlikely. X linked inheritance would require a more severe or even lethal phenotype in males. In autosomal dominant inheritance, only fathers would be affected, because reproduction in females is not possible.

In males aplasia/hypoplasia of the Müllerian duct should not have any functional consequence, since only the cranial and the caudal ends persist, the first as the appendix testis, the other as the uterus masculinus in the prostate.[21]

Wellesley and Slaney[22] reported on a man with left renal agenesis, block fusion from C6-T8 with multiple deformities of several neural arches, azoospermia, and normal chromo-

Figure 1  Note facial asymmetry with normal ears at the age of 25 years.
Various other anomalies have been described frequently in women with MURCS association. Abnormalities of the upper extremities or thoracic ribs, scoliosis, and facial asymmetry can be attributed to the dysplasia of the cervical somites. Encephalocoele, hearing loss, corneal anaesthesia, cleft lip/palate, or malformations of the central nervous system are rare and might be by chance. Renal problems as in our case often result in growth retardation as a secondary symptom.

In the original report of Duncan et al. uterine aplasia/hypoplasia, renal agenesis/ectopy, and anomalies of the cervical spine were only present in 16 out of 30 cases. Only 46 out of 65 previously published cases have had all three major manifestations. Therefore, the acronym MURCS should be restricted to women with all typical symptoms. Application to patients with only some of the typical manifestations is confusing and hinders the delineation from atypical cases of Klippel-Feil sequence, MRKH syndrome, or Goldenhar syndrome.

In conclusion, MURCS association should be considered as a developmental field defect and only diagnosed if both aplasia/hypoplasia of the Müllerian duct, renal agenesis/ectopy, and anomalies of the cervicothoracic somites are present.

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