Klippel-Trenaunay Syndrome (KTS) was first described by Klippel and Trenaunay in 1900. It is common in children and young adults and is characterized by soft tissue and bony hypertrophy, varicose venous structures and skin nevus which are unilateral and almost exclusively involve lower extremities. It is rarely bilaterally and involves upper extremities (1-4).

In this paper we present the radiologic findings of KTS in the light of previous findings.

A 30-month-old boy was brought in with the complaint of gradually increased size of his right lower extremity from birth. Soft tissue hypertrophy was detected in his right thigh, right dorsal region and scrotum. On the right leg, the skin had bluish discoloration. The leg was swollen and edematous. On plain radiography, we detected soft tissue hypertrophy, and there was length discrepancy between the right and left legs. The right femur was longer than the left (Figure 1). In color Doppler imaging, deep venous system was absent, multiple varicose veins were demonstrated under subcutaneous fatty tissue at the thigh, near and below the knee. The thickness of subcutaneous fatty tissue was also increased. No arterio-venous fistula was detected. In magnetic resonance imaging, subcutaneous fatty tissue at the dorsal region (Figure 2), and right leg was detected as increased thickness with the equal intensity of fatty tissue on T1 and T2-weighted sequences. Venous varicosities were demonstrated as a signal void on T1-weighted spin-echo sequences (Figure 3) and a bright signal on T2-weighted gradient-echo sequence (Figure 4).

The classic triad of KTS was venous abnormalities, cutaneous hemangiomas and hypertrophy at the involved extremity. In addition to these findings, syndactyly, polydactyly and congenital hip dislocation can be seen as a part of this syndrome (2). The syndrome usually affects the unilateral and lower extremities but occasionally may be bilateral and involve the upper extremities. There is no sex preponderance or familial pattern. There may be hypoplasia, atresia or agenesis of the deep venous system at the involved extremity. So the venous return is provided via the superficial venous system. The superficial venous system becomes dilated and tortuous. The deep venous system is not normal in any case (4,5). In the study of Serville, fibrous cord and aberrant artery compressing the deep venous system was described (6). Parker and Weber described arterio-venous fistula in addition to classic triad and named it Parker-Weber syndrome (7). The hypertrophy of the involved extremity, the important clinic manifestation of the syndrome, is due to increased arterial flow caused by venous stasis and hemangiomatosis. The involved extremity is usually larger and longer than normal. Less frequently, intermittent claudication, varicose ulcers, increased skin temperature, hair loss, and dyskeratosis are seen (4,8).

Plain radiography gives useful information about the limb hypertrophy which is the cardinal manifestation of KTS. It is manifested as extremity length discrepancy. Other osseous anomalies such as syndactyly and polydactyly can be seen. The phleboliths shows the venous malformation and prior intralesional hemorrhage and thrombus.

Color Doppler imaging may be used to identify the deep venous system and tortuous and dilate superficial venous system. Arterio-venous fistula can be detected by its characteristic spectral analysis.
Figure 1. Plain radiography shows marked soft tissue hypertrophy at right leg compared with left. And right femur is also longer than left.

Figure 2. On spin-echo T1-weighted MR section, the thickness of subcutaneous fatty is markedly increased.
Venography and arteriography is performed for treatment planning. Arterio-venous fistula associated with Parker-Weber syndrome can be detected by arteriography.

Magnetic resonance imaging can detect the soft tissue and bony extension of venous malformation. This information is important in surgical planning. Leg discrepancy is also detected by MR imaging. The signal intensity of vascular channels may vary depending on the imaging parameters and flow velocities. Vascular channels are demonstrated as a signal void on spin-echo T1-weighted images and bright signal intensities on gradient echo T2-weighted images.

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Klippel-Trenaunay Syndrome

References