TAR syndrome (Case Report)

Abstract
Diagnosis of TAR syndrome usually are made at birth because of the characteristic physical appearance combined with thrombocytopenia. The two essential features of TAR syndrome are hypomegakaryocytic thrombocytopenia and bilateral radial aplasia. The rest of the phenotype varies widely and can manifest with abnormalities involving skeletal, skin, gastrointestinal and cardiac systems. Considering to limited cases of syndrome and variability of phenotypic abnormalities excess two essential features, all cases of this syndrome are reportable. We reported a patient with TAR syndrome with any related malformations.

Key Words: TAR syndrome- Thrombocytopenia- Radius