




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Case Report**Craniosynostosis, Ptosis, Hypodontia, Prominent and Everted Lower Lip, Mental Retardation; Is it a Second Case of Mehta-Lewis-Patton Syndrome?****Naeimeh Tayebi¹, MD, Genetic Counselor; Seyed Mahmood Akhavi Mirabbashi¹, MD, Fellowship of Pediatric Neurology; Hossein Khodae¹, MSc of Molecular Genetics**

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Abstract:

We present an unknown case of an 11-year-old boy with mental retardation, microcephaly, prominent ears, unilateral ptosis, long philtrum, prominent and everted lower lip, abnormally shaped teeth and developmental delay. This is an unknown case with special facial features and mental retardation which can probably be the second case of Mehta-Lewis-Patton syndrome.

Keywords:[Craniosynostosis](#) . [Mental retardation](#) . [Microcephaly](#) . [Ptosis](#) . [Everted lip](#)

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