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Published journal articles indexed by SCI, SSCI, and AHCI

- I. **Enhancing Genetic Insight: Chromosomal Microarray Enhances Understanding of Genetics in Rubinstein-Taybi Syndrome**
ERKAN D. D., SOĞUKPINAR M., ÜREL DEMİR G., ÜTİNE G. E., ŞİMŞEK KİPER P. Ö.
MOLECULAR SYNDROMOLOGY, 2024 (SCI-Expanded)
- II. **Coexistence of Two Rare Conditions Complicating the Other's Management: Propionic Acidemia and Apert Syndrome**
ENSERT CİHAN C. K., AKAR H. T., YILDIZ Y., SOĞUKPINAR M., ÜTİNE G. E., ÇELİK H. T.
Molecular Syndromology, vol.15, no.1, pp.83-88, 2024 (SCI-Expanded)
- III. **Review of patients with achondroplasia: a single-center's experience with follow-up and associated morbidities**
SOĞUKPINAR M., ÜREL DEMİR G., ÜTİNE G. E., GÖNÇ E. N., ÖZÖN Z. A., ŞİMŞEK KİPER P. Ö.
EUROPEAN JOURNAL OF PEDIATRICS, no.9, pp.3819-3829, 2024 (SCI-Expanded)
- IV. **Further defining the molecular spectrum and long-term follow-up of 17 patients with Dyggve-Melchior-Clausen and Smith-McCort dysplasia type 2**
AKALIN A., Ayaz E., SOĞUKPINAR M., Avci-Durmusalioglu E., ÜREL DEMİR G., YILDIZ A. E., ATİK T., ELÇİOĞLU H. N., Eda Utine G., ŞİMŞEK KİPER P. Ö.
AMERICAN JOURNAL OF MEDICAL GENETICS PART A, 2024 (SCI-Expanded)
- V. **A spectrum of TP63-related disorders with eight affected individuals in five unrelated families**
SOĞUKPINAR M., ÜTİNE G. E., Boduroğlu K., ŞİMŞEK KİPER P. Ö.
European Journal of Medical Genetics, vol.68, 2024 (SCI-Expanded)
- VI. **Delineation of <i>ADPRHL2</i> Variants: Report of Two New Patients with Review of the Literature**
Yildiz S. O., Yalnizoglu D., ŞİMŞEK KİPER P. Ö., GÖÇMEN R., Sogukpinar M., Utine G. E., Haliloglu G.
NEUROPEDIATRICS, no.02, pp.117-123, 2024 (SCI-Expanded)

Articles Published in Other Journals

- I. **Clinical Evaluation of the Five Patients with Mosaic Trisomy 8 Syndrome: Case Series**
DAŞAR T. N., Soğukpinar M., Simsek-Kiper P. O., ÜTİNE G. E., BODUROĞLU O. K.
Türkiye Klinikleri Pediatri Dergisi, vol.32, no.2, pp.91-95, 2023 (Scopus)

Refereed Congress / Symposium Publications in Proceedings

- I. **A Novel ZBTB20 Variant In A Patient With Primrose syndrome: A rare clinical entity**
SOĞUKPINAR M., KARAOSMANOĞLU B., ÜTİNE G. E., Boduroglu K., Simsek-Kiper P.
56th Annual Conference of the European-Society-of-Human-Genetics (ESHG), Glasgow, England, 10 - 13 June 2023,

pp.185

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Publication: 8