

Kişisel Bilgiler

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SCI, SSCI ve AHCI İndekslerine Giren Dergilerde Yayınlanan Makaleler

- I. **Long-term clinical evaluation of patients with alpha-mannosidosis – A multicenter study**
KÖSE E., KASAPKARA Ç. S., İNCİ A., YILDIZ Y., Sürücü Kara İ., Kahraman A. B., TÜMER L., DURSUN A., EMİNOĞLU F. T.
European Journal of Medical Genetics, cilt.68, 2024 (SCI-Expanded)
- II. **Predictors of eventual requirement of phenylalanine-restricted diet in young infants with phenylalanine hydroxylase deficiency initially managed with sapropterin monotherapy**
ÇIKI K., YILDIZ Y., KAHRAMAN A. B., ÖZGÜL R. K., COŞKUN T., DURSUN A., TOKATLI A., SİVRİ S.
Molecular Genetics and Metabolism, cilt.140, sa.3, 2023 (SCI-Expanded)
- III. **COVID-19 in inherited metabolic disorders: Clinical features and risk factors for disease severity**
KAHRAMAN A. B., YILDIZ Y., ÇIKI K., Erdal I., AKAR H. T., DURSUN A., TOKATLI A., SİVRİ S.
Molecular Genetics and Metabolism, cilt.139, sa.2, 2023 (SCI-Expanded)
- IV. **Successful management of rhabdomyolysis with triheptanoin in a child with severe long-chain 3-hydroxyacyl-coenzyme A dehydrogenase (LCHAD) deficiency**
KAHRAMAN A. B., YILDIZ Y., GÖKMEN ÖZEL H., KADAYIFÇILAR S., SİVRİ S.
Neuromuscular Disorders, cilt.33, sa.4, ss.315-318, 2023 (SCI-Expanded)
- V. **Two tales of LPIN1 deficiency: from fatal rhabdomyolysis to favorable outcome of acute compartment syndrome**
KAHRAMAN A. B., Karakaya B., YILDIZ Y., KAMACI S., KESİCİ S., ŞİMŞEK KİPER P. Ö., KURT ŞÜKÜR E. D., BAYRAKCI B., Haliloglu G.
Neuromuscular Disorders, cilt.32, sa.11-12, ss.931-934, 2022 (SCI-Expanded)
- VI. **Diagnostic distribution and postnatal evaluation of prenatally detected short femur: A single center experience**
KAHRAMAN A. B., ŞİMŞEK KİPER P. Ö., ÜTİNE G. E., BODUROĞLU O. K.
AMERICAN JOURNAL OF MEDICAL GENETICS PART A, cilt.188, sa.8, ss.2367-2375, 2022 (SCI-Expanded)
- VII. **Novel Cranial Imaging Findings and a Splice-Site Variant in a Patient with Tyrosinemia Type III, and a Summary of Published Cases**
KAHRAMAN A. B., AKAR H. T., Lafci N. G., YILDIZ Y., Tokatli A.
MOLECULAR SYNDROMOLOGY, cilt.13, sa.3, ss.193-199, 2022 (SCI-Expanded)

Metrikler

Yayın: 7

Atıf (Scopus): 3

H-İndeks (Scopus): 1