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## Published journal articles indexed by SCI, SSCI, and AHCI

- I. **A case report of a patient with neurodevelopmental disorder with impaired speech and hyperkinetic movements: A biallelic variant in the ZNF142 gene**  
Kaya D., CEYLAN KÖSE C., AKCAN M. B., SILAN F.  
American Journal of Medical Genetics, Part A, vol.194, no.9, 2024 (SCI-Expanded)
- II. **Anemia and thrombocytopenia due to a novel BRPF1 variant in a family from Çanakkale with intellectual disability and dysmorphic facies: Case report and review of the literature**  
Kose C. C., Kaya D., Akcan M. B., Silan F.  
AMERICAN JOURNAL OF MEDICAL GENETICS, PART A, vol.191, no.6, pp.2209-2214, 2023 (SCI-Expanded)

## Refereed Congress / Symposium Publications in Proceedings

- I. **Importance of genetic diagnosis in demyelinating diseases; adult-onset Alexander Disease case with novel GFAP mutation**  
CEYLAN KÖSE C., OCAK Ö., GÜNDÜZ O. R., SILAN F.  
15. Ulusal Tıbbi Genetik Kongresi, Muğla, Turkey, 9 - 13 November 2022
- II. **A novel BRPF1 variant in a family with intellectual disability and dysmorphic face, from Çanakkale**  
CEYLAN KÖSE C., KAYA D., AKCAN M. B., SILAN F.  
EUROPEAN BIOTECHNOLOGY CONGRESS 2022, PRAG, Czech Republic, 5 - 07 October 2022
- III. **A rare Malpuech, Mignarelli, Michels, Carnevale Syndrome (3MC Syndrome); Novel variant in the MASP1 gene**  
GÜNDÜZ O. R., CEYLAN KÖSE C., SILAN F.  
EUROPEAN BIOTECHNOLOGY CONGRESS 2022, PRAG, Czech Republic, 5 - 07 October 2022
- IV. **A novel homozygous mutation in CC2D1A gene: two case from a Turkish family in Canakkale**  
KAYA D., CEYLAN KÖSE C., SILAN F.  
EUROPEAN BIOTECHNOLOGY CONGRESS 2022, PRAG, Czech Republic, 5 - 07 October 2022
- V. **A rare syndrome from Canakkale; Kindler Syndrome with loss of heterozygosity of FERMT1 gene region**  
GÜNDÜZ O. R., IŞIK MERMUTLU S., CEYLAN KÖSE C., SILAN F.  
7. Uluslararası Erciyes Tıp Tıbbi Genetik Kongresi, Kayseri, Turkey, 26 - 28 May 2022
- VI. **P-28 Meme Ve Over Kanserinde Kombine Mutasyonlar Tedavi Etkinliğini Değiştirebilir Mi? Ailesinde Meme Ve Over Kanseri Öyküsü Olan Vakada Brca1 Ve Sdhb Genleri Patolojik/muhtemel Patolojik Varyant Birlikteliği**  
CEYLAN KÖSE C., SILAN F., TURAN N. E., SÖNMEZ V., SILAN C.  
1. Ulusal HematoOnkoGenetik Kongresi, Antalya, Turkey, 25 - 28 November 2021

## Metrics

Publication: 14

Citation (WoS): 2

Citation (Scopus): 2

H-Index (WoS): 1

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