

Prof. ERCAN DEMİR

Personal Information

Office Phone: [+90 312 202 6006](tel:+903122026006)

Email: ercandemir@gazi.edu.tr

Web: <https://avesis.gazi.edu.tr/ercandemir>

International Researcher IDs

ScholarID: I0uGRvQAAAAJ

ORCID: 0000-0003-3431-7316

Publons / Web Of Science ResearcherID: CAA-1553-2022

ScopusID: 7004473185

Yoksis Researcher ID: 54961

Education Information

Post Doctorate of Medicine, Hacettepe University, Tıp Fakültesi, Dahili Tıp Bilimleri Bölümü, Turkey 1997 - 2000

Expertise In Medicine, Hacettepe University, Tıp Fakültesi, Dahili Tıp Bilimleri Bölümü, Turkey 1991 - 1996

Undergraduate, Hacettepe University, Tıp Fakültesi, Tip Pr., Turkey 1984 - 1991

Research Areas

Health Sciences

Academic Titles / Tasks

Associate Professor, Gazi University, Tıp Fakültesi, Dahili Tıp Bilimleri, 2006 - 2010

Associate Professor, Karadeniz Technical University, Tıp Fakültesi, Dahili Tıp Bilimleri Bölümü, 2005 - 2006

Assistant Professor, Karadeniz Technical University, Tıp Fakültesi, Dahili Tıp Bilimleri Bölümü, 2003 - 2005

Courses

Undergraduate

PARALİTİK HASTALIKLAR (Dönem4), Undergraduate, 2017 - 2018

Hereditary and acquired motor neuron diseases (Phase3), Undergraduate, 2017 - 2018

PARALYTIC DISORDERS (Phase4), Undergraduate, 2017 - 2018

APPROACH TO HEADACHE (Phase4), Undergraduate, 2017 - 2018

BAŞAĞRISINA YAKLAŞIM (Dönem4), Undergraduate, 2017 - 2018

Primary muscle disorders (Phase3), Undergraduate, 2017 - 2018

Heredititer ve akkiz motor nöron hastalıkları (Dönem3), Undergraduate, 2017 - 2018

Primer kas hastalıkları (Dönem3), Undergraduate, 2017 - 2018

Supervised Theses

DEMİR E., Investigation of primary headache, comorbid diseases and their effects on the quality of life in childhood, Expertise In Medicine, C.TOSUN(Student), 2014

DEMİR E., Investigation of the relation between oxidative stress and carnitine levels in epileptic children treated with valproic acid or carbamazepine, Expertise In Medicine, A.NEŞE(Student), 2012

DEMİR E., Investigation of the etiological aspects of pediatric patients with acute impairment of consciousness, Expertise In Medicine, S.ELMAOĞULLARI(Student), 2012

Published journal articles indexed by SCI, SSCI, and AHCI

- I. **Thymic carcinoma presenting with overlap polyarthritis and myositis: A rare paraneoplastic syndrome in childhood**
Yıldız Ç., Türkcan B. T., Vural Ö., Gezgin Yıldırım D., İnan M. A., Poyraz A., Pınarlı F. G., Taştepe İ., Demir E., Sunar Yayla E. N., et al.
INTERNATIONAL JOURNAL OF RHEUMATIC DISEASES, vol.27, no.5, 2024 (SCI-Expanded)
- II. **An ultra-rare cause of severe hypotonia mimicking Pompe disease in an infant: RRM2B related mitochondrial DNA depletion syndrome with a novel mutation**
İNCİ A., OKUR İ., DEMİR E., BİBEROĞLU G., TÜMER L., SERDAROĞLU A., EZGÜ F. S.
NEUROLOGY ASIA, vol.27, no.1, pp.199-202, 2022 (SCI-Expanded)
- III. **Hypophosphatasia: is it an underdiagnosed disease even by expert physicians?**
İnci A., Ergin F. B., Yüce B. T., Çiftçi B., Demir E., Buyan N., Okur İ., Biberoglu G., Öktem R. M., Tümer L., et al.
JOURNAL OF BONE AND MINERAL METABOLISM, vol.39, no.4, pp.598-605, 2021 (SCI-Expanded)
- IV. **Milk of calcium: A rare manifestation of juvenile dermatomyositis**
Yıldırım D., AKDULUM İ., TALİM B., DEMİR E., BUYAN N., BAKKALOĞLU EZGÜ S. A.
ARCHIVES OF RHEUMATOLOGY, vol.36, no.2, pp.302-304, 2021 (SCI-Expanded)
- V. **Evaluation of state and trait anxiety levels of parents and children before electroencephalography procedures: A prospective study from a tertiary epilepsy center**
Tekin L. O., Cebeci D., Unver E., Acar A. Ş., Demir E., Güçüyener K., Uğraş Dikmen A., Serdaroglu A., Arhan E.
EPILEPSY & BEHAVIOR, vol.112, 2020 (SCI-Expanded)
- VI. **Temporary Seizure in an Infant Who Had Been Exposed to G-Amino Butyric Acid Receptor Antagonist Thiocolchicoside**
Havah C., GÜCÜYENER K., Gurkas E., DEMİR E.
PEDIATRIC EMERGENCY CARE, vol.35, no.5, 2019 (SCI-Expanded)
- VII. **A child with atypically subtle clinical presentation of acute arterial ischaemic stroke in the middle cerebral artery**
Orgun L. T., Derinkuyu B. E., Havalı C., Boyunaga Ö. L., Yenicesu I., DEMİR E., GÜCÜYENER K.
INTERVENTIONAL NEURORADIOLOGY, vol.24, no.6, pp.684-687, 2018 (SCI-Expanded)
- VIII. **Internal carotid artery dissection without intracranial infarct following a minor shoulder trauma: The second pediatric case and review of the literature**
Cebeci D., ARHAN E., DEMİR E., UÇAR M., Ucar H. K., Serdaroglu A., Ozturk Z.
JOURNAL OF CLINICAL NEUROSCIENCE, vol.56, pp.172-175, 2018 (SCI-Expanded)
- IX. **Homozygous Mutations in TBC1D23 Lead to a Non-degenerative Form of Pontocerebellar Hypoplasia**
Marin-Valencia I., Gerondopoulos A., Zaki M. S., Ben-Omran T., Almureikhi M., Demir E., Guemez-Gamboa A., Gregor A., Issa M. Y., Appelhof B., et al.
AMERICAN JOURNAL OF HUMAN GENETICS, vol.101, no.3, pp.441-450, 2017 (SCI-Expanded)
- X. **Regional brain volume reduction and cognitive outcomes in preterm children at low risk at 9 years of age**
ARHAN E., GÜCÜYENER K., Soysal S., Salvarli S., Gurses M. A., SERDAROĞLU A., DEMİR E., Ergenekon E., TÜRKYILMAZ C., Onal E. E., et al.
CHILDS NERVOUS SYSTEM, vol.33, no.8, pp.1317-1326, 2017 (SCI-Expanded)

- XI. **Sacral aneurysmal bone cyst in a child presenting with radiculopathy.**
Derinkuyu B. E., BOYUNAGA Ö. L., Tekin-Orgun L., Ozdemir-Gokce A., Firat H., DEMİR E.
The spine journal : official journal of the North American Spine Society, vol.16, no.6, 2016 (SCI-Expanded)
- XII. **Opsoclonus-myoclonus syndrome following rotavirus gastroenteritis**
Gurkas E., GÜCÜYENER K., Yilmaz U., Havalı C., DEMİR E.
PEDIATRICS INTERNATIONAL, vol.56, no.6, 2014 (SCI-Expanded)
- XIII. **The Effect of Topiramate on Body Weight and Ghrelin, Leptin, and Neuropeptide-Y Levels of Prepubertal Children With Epilepsy**
Ozcelik A. A., SERDAROĞLU A., Bideci A., ARHAN E., Soysal S., DEMİR E., GÜCÜYENER K.
PEDIATRIC NEUROLOGY, vol.51, no.2, pp.220-224, 2014 (SCI-Expanded)
- XIV. **Matrix Metalloproteinase-7, Matrix Metalloproteinase-9, and Disease Activity in Pediatric Multiple Sclerosis Response**
Yilmaz U., GÜCÜYENER K., Gurkas E., DEMİR E., SERDAROĞLU A., Atak A., Aral A., Oner Y. A.
PEDIATRIC NEUROLOGY, vol.48, no.3, pp.255-256, 2013 (SCI-Expanded)
- XV. **Reduced Retinal Nerve Fiber Layer Thickness and Macular Volume in Pediatric Multiple Sclerosis**
Yilmaz U., GÜCÜYENER K., Erin D. M., Yazar Z., Gurkas E., SERDAROĞLU A., Tepe N., DEMİR E.
JOURNAL OF CHILD NEUROLOGY, vol.27, no.12, pp.1517-1523, 2012 (SCI-Expanded)
- XVI. **Retinal Thickness in Pediatric Multiple Sclerosis**
Yilmaz U., GÜCÜYENER K., Erin D. M., Yazar Z., Gurkas E., SERDAROĞLU A., Tepe N., DEMİR E.
JOURNAL OF CHILD NEUROLOGY, vol.27, no.12, pp.1621-1622, 2012 (SCI-Expanded)
- XVII. **Matrix Metalloproteinase-7 and Matrix Metalloproteinase-9 in Pediatric Multiple Sclerosis**
Unsal Y., Kivilcim G., Aysegul A., Arzu A., Esra G., Ercan D., Ayse S.
PEDIATRIC NEUROLOGY, vol.47, no.3, pp.171-176, 2012 (SCI-Expanded)
- XVIII. **Molecular Genetic Analysis of the PLP1 Gene in 38 Families with PLP1-related disorders: Identification and Functional Characterization of 11 Novel PLP1 Mutations**
Grossi S., Regis S., Biancheri R., Mort M., Lualdi S., Bertini E., Uziel G., Boespflug-Tanguy O., Simonati A., Corsolini F., et al.
ORPHANET JOURNAL OF RARE DISEASES, vol.6, 2011 (SCI-Expanded)
- XIX. **Clinical, neuroradiological and genetic findings in pontocerebellar hypoplasia**
Yasmin Namavar Y., Barth P. G., Kasher P. R., Van Ruissen F., Brockmann K., Bernert G., Witzel K., Ventura K., Cheng E. Y., Demir E.
BRAIN : A JOURNAL OF NEUROLOGY, vol.134, no.1, pp.143-156, 2011 (SCI-Expanded)
- XX. **Are serum nitric oxide and vascular endothelial growth factor levels affected by packed red blood cell transfusions?**
Ergenekon E., Bozkaya D., Goktas T., Erbas D., ATAŞ YÜCEL A., Turan O., Hirfanoglu I., Onal E. E., TÜRKYILMAZ C., KOÇ E., et al.
HEMATOLOGY, vol.15, no.3, pp.170-173, 2010 (SCI-Expanded)
- XXI. **An unusual presentation of Muscle-Eye-Brain disease: Severe eye abnormalities with mild muscle and brain involvement**
DEMİR E., GÜCÜYENER K., Akturk A., TALİM B., Konus Ö. L., Del Bo R., Ghezzi S., Comi G. P.
NEUROMUSCULAR DISORDERS, vol.19, no.10, pp.692-695, 2009 (SCI-Expanded)
- XXII. **Assessment of mothers' knowledge and perceptions of electroencephalography and determination of the short-term effect of an informational leaflet**
ARHAN E., SERDAROĞLU A., Soysal S., Ozcelik A., GÜCÜYENER K., DEMİR E.
EPILEPSY & BEHAVIOR, vol.15, no.4, pp.491-495, 2009 (SCI-Expanded)
- XXIII. **Atypical Presentations of Subacute Sclerosing Panencephalitis in Two Neurologically Handicapped Cases**
Demir E., Ozcelik A., Arhan E., Serdaroglu A., Gucuyener K.
NEUROPEDIATRICS, vol.40, no.4, pp.195-198, 2009 (SCI-Expanded)
- XXIV. **3-Methylcrotonyl-CoA Carboxylase Deficiency: Phenotypic Variability in a Family**
Eminoglu F. T., Ozcelik A. A., OKUR İ., TÜMER L., BİBEROĞLU G., DEMİR E., Hasanoglu A., Baumgartner M. R.

- JOURNAL OF CHILD NEUROLOGY, vol.24, no.4, pp.478-481, 2009 (SCI-Expanded)
- XXV. Atypical presentations of SSPE: a clinical study in four cases**
Demir E., Aksoy A., Anlar B., Soenmez F. M.
TURKISH JOURNAL OF PEDIATRICS, vol.49, no.3, pp.295-300, 2007 (SCI-Expanded)
- XXVI. Novel truncating and missense mutations of the KCC3 gene associated with Andermann syndrome**
Uyanik G., Elcioglu N., Penzien J., Gross C., Yilmaz Y., Olmez A., Demir E., Wahl D., Scheglmann K., Winner B., et al.
NEUROLOGY, vol.66, no.7, pp.1044-1048, 2006 (SCI-Expanded)
- XXVII. Effect of antiepileptic drugs on plasma lipids, lipoprotein (a), and liver enzymes**
Sonmez F., Demir E., Örem A., Yıldırım S., Orhan F., Aslan A., Topbaş M.
JOURNAL OF CHILD NEUROLOGY, vol.21, no.1, pp.70-74, 2006 (SCI-Expanded)
- XXVIII. Ultrastructural defects of collagen VI filaments in an Ullrich syndrome patient with loss of the alpha 3(VI) N10-N7 domains**
Swarzoni S., Sabatelli P., Bergamin N., Guicheney P., Demir E., Merlini L., Lattanzi G., Ognibene A., Capanni C., Mattioli E., et al.
JOURNAL OF CELLULAR PHYSIOLOGY, vol.206, no.1, pp.160-166, 2006 (SCI-Expanded)
- XXIX. Benign familial infantile convulsions: phenotypic variability in a family.**
Demir E., Turanii G., Yalntzoglu D., Topcu M.
Journal of child neurology, vol.20, no.6, pp.535-8, 2005 (SCI-Expanded)
- XXX. Giant axonal neuropathy: clinical and genetic study in six cases**
Demir E., Bomont P., Erdem S., Cavalier L., Demirci M., Kose G., Muftuoglu S., Cakar A., Tan E., Aysun S., et al.
JOURNAL OF NEUROLOGY NEUROSURGERY AND PSYCHIATRY, vol.76, no.6, pp.825-832, 2005 (SCI-Expanded)
- XXXI. Infantile convulsions and paroxysmal choreoathetosis in a consanguineous family**
Demir E., Prud'homme J., Topcu M.
PEDIATRIC NEUROLOGY, vol.30, no.5, pp.349-353, 2004 (SCI-Expanded)
- XXXII. Collagen VI status and clinical severity in Ullrich congenital muscular dystrophy: Phenotype analysis of 11 families linked to the COL6 loci**
Demir E., Ferreiro A., Sabatelli P., Allamand V., Makri S., Echenne B., Maraldi M., Merlini L., Topaloglu H., Guicheney P.
NEURODIETRICS, vol.35, no.2, pp.103-112, 2004 (SCI-Expanded)
- XXXIII. Andermann syndrome in a Turkish patient**
Demir E., Irobi J., Erdem S., Demirci M., Tan E., Timmerman V., De Jonghe P., Topaloglu H.
JOURNAL OF CHILD NEUROLOGY, vol.18, no.1, pp.76-79, 2003 (SCI-Expanded)
- XXXIV. Exon deletions in the GCH1 gene in two of four Turkish families with dopa-responsive dystonia**
Klein C., Hedrich K., Kabakci K., Mohrmann K., Wiegers K., Landt O., Hagenah J., Schwinger E., Pramstaller P., Ozelius L., et al.
NEUROLOGY, vol.59, no.11, pp.1783-1786, 2002 (SCI-Expanded)
- XXXV. Mutations in COL6A3 cause severe and mild phenotypes of Ullrich congenital muscular dystrophy**
Demir E., Sabatelli P., Allamand V., Ferreiro A., Moghadashadeh B., Makrelouf M., Topaloglu H., Echenne B., Merlini L., Guicheney P.
AMERICAN JOURNAL OF HUMAN GENETICS, vol.70, no.6, pp.1446-1458, 2002 (SCI-Expanded)
- XXXVI. A case of Sandifer's syndrome with hand tremor.**
Demir E., Saka E., Aysun S.
The Turkish journal of pediatrics, vol.43, no.4, pp.348-50, 2001 (SCI-Expanded)
- XXXVII. The gene encoding gigaxonin, a new member of the cytoskeletal BTB/kelch repeat family, is mutated in giant axonal neuropathy**
Bomont P., Cavalier L., Blondeau F., Hamida C., Belal S., Tazir M., Demir E., Topaloglu H., Korinthenberg R., Tuysuz B., et al.
NATURE GENETICS, vol.26, no.3, pp.370-374, 2000 (SCI-Expanded)
- XXXVIII. Giant axonal neuropathy locus refinement to a < 590 kb critical interval**
Cavalier L., BenHamida C., Amouri R., Belal S., Bomont P., Lagarde N., Gressin L., Callen D., Demir E., Topaloglu H., et al.

- EUROPEAN JOURNAL OF HUMAN GENETICS, vol.8, no.7, pp.527-534, 2000 (SCI-Expanded)
- XXXIX. **A cross section of autosomal recessive limb-girdle muscular dystrophies in 38 families**
DİNÇER P. R., Akcoren Z., Demir E., Richard I., Sancak O., Kale G., Ozme S., KARADUMAN A. A., Tan E., Urtzbere J., et al.
JOURNAL OF MEDICAL GENETICS, vol.37, no.5, pp.361-367, 2000 (SCI-Expanded)
- XL. **Weight gain associated with valproate in childhood**
Demir E., Aysun S.
PEDIATRIC NEUROLOGY, vol.22, no.5, pp.361-364, 2000 (SCI-Expanded)
- XLI. **Effect of alendronate treatment on the clinical picture and bone turnover markers in chronic idiopathic hyperphosphatasia**
DEMİR E., Bereket A., Özkan B., Topçu M.
Journal of Pediatric Endocrinology and Metabolism, vol.13, no.2, pp.217-221, 2000 (SCI-Expanded)
- XLII. **Clinical and genetic correlate in childhood onset Friedreich ataxia**
Alikasifoglu M., Topaloglu H., Tuncbilek E., Ceviz N., Anar B., Demir E., Ozme S.
NEUROPEDIATRICS, vol.30, no.2, pp.72-76, 1999 (SCI-Expanded)
- XLIII. **A new case of oculocerebral hypopigmentation syndrome (Cross syndrome) with additional findings**
Tezcan I., Demir E., Asan E., Kale G., Muftuoglu S., Kotiloglu E.
CLINICAL GENETICS, vol.51, no.2, pp.118-121, 1997 (SCI-Expanded)
- XLIV. **Infantile spasms as the initial symptom of biotinidase deficiency**
Kalayci Ö., Coskun T., Tokatli A., Demir E., Erdem G., Güngör C., Yükselen A., Özalp I.
Journal of Pediatrics, vol.124, no.1, pp.103-104, 1994 (SCI-Expanded)

Articles Published in Other Journals

- I. **The Issue of Minorities from Sèvres to Lausanne in the Context of Its Reflections to the Present and Its Practices in the Republican Era Günümüze Yansımaları Bağlamında Sevr'den Lozan'a Azınlıklar Meselesi ve Cumhuriyet Devri'ndeki Uygulamaları**
DEMİR E.
History Studies, vol.17, no.1, pp.187-213, 2025 (Scopus)
- II. **The Relation of Epilepsy Type with Depression and Anxiety in Children**
Ozturk Z., Soysal S., GÜCÜYENER K., DEMİR E., ARHAN E., SERDAROĞLU A.
GAZI MEDICAL JOURNAL, vol.32, pp.273-275, 2021 (ESCI)
- III. **SMN1 gen delesyonu dışlanılmış Spinal Musküler Atrofi ön tanılı çocukların etiyolojinin tüm ekzom dizi analizi verilerine dayanarak retrospektif olarak araştırılması**
SEZER A., DEMİR E., KAYHAN G., ERGÜN M. A., TUĞ E.
Gazi Medical Journal, 2020 (ESCI)
- IV. **Dikkat Eksikliği Hiperaktivite Bozukluğu olan Hastaların Nöropsikolojik Test Performansının Değerlendirilmesi**
GÜCÜYENER K., SOYSAL ACAR A. Ş., öztürk z., DEMİR E., ARHAN E., SERDAROĞLU A.
Gazi Medical Journal, vol.30, no.2, 2019 (Scopus)
- V. **Evaluation of Neuropsychological Test Performance of Patients with Attention Deficit Hyperactivity Disorder**
SOYSAL ACAR A. Ş., Ozturk Z., GÜCÜYENER K., DEMİR E., ARHAN E., SERDAROĞLU A.
GAZI MEDICAL JOURNAL, vol.30, no.2, pp.114-118, 2019 (ESCI)
- VI. **Sepsis-Associated Encephalopathy in a Child with the Torsion of Meckel's Diverticulum**
Gurkas E., Yilmaz U., DEMİR E., GÜCÜYENER K.
JOURNAL OF CLINICAL AND ANALYTICAL MEDICINE, vol.7, pp.158-160, 2016 (ESCI)
- VII. **Case: Valproat-Induced Hyperammonemic Encephalopathy**
Aksu A. U., ARHAN E., DEMİR E., Ozcelik A. A., Serdaroglu A., GÜCÜYENER K.
GAZI MEDICAL JOURNAL, vol.27, no.2, pp.85-86, 2016 (ESCI)

- VIII. **Spastic Paraplegia with SPG11 Gene delE39 in a Turkish Patient**
 Ozturk Z., Havalı C., DEMİR E., GÜCÜYENER K.
 GAZI MEDICAL JOURNAL, vol.27, no.4, pp.205-206, 2016 (ESCI)
- IX. **Autism Spectrum Disorder Management Practices and Level of Knowledge Among General Pediatricians**
 Ozcelik A. A., Soysal S., ARHAN E., DEMİR E., GÜCÜYENER K., SERDAROĞLU A.
 GAZI MEDICAL JOURNAL, vol.26, no.4, pp.158-162, 2015 (ESCI)
- X. **Reactive haemophagocytic syndrome in a child with acalculous cholecystitis**
 ERDURAN E., DEMİR E., SARI A., girişken i.
 Haema, vol.8, no.1, pp.113-117, 2005 (Peer-Reviewed Journal)

Books

- I. **ARTROGRİPOZİS TANI VE TEDAVİ ALGORİTMASI**
 ÖZBUDAK P., DEMİR E.
 in: Pediatrik Nöroloji: Algoritmalar ve İlaç Rehberi, Kumandaş Sefer, Cankurtaran Mehmet, Editor, AKADEMİSYEN KİTABEVİ, Ankara, pp.501-502, 2022
- II. **SMA TANI VE TEDAVİ ALGORİTMASI**
 DEMİR E., ÖZBUDAK P.
 in: Pediatrik Nöroloji: Algoritmalar ve İlaç Rehberi, Kumandaş Sefer, Cankurtaran Mehmet, Editor, AKADEMİSYEN KİTABEVİ, Ankara, pp.509-511, 2022
- III. **Artrogripozis**
 ÖZBUDAK P., DEMİR E.
 in: Temel Pediatrik Nöroloji: Tanı ve Tedavi, Kumandaş Sefer, Cankurtaran Mehmet, Editor, AKADEMİSYEN KİTABEVİ, Ankara, pp.2973-2983, 2022
- IV. **ÖN BOYNUZ MOTOR NÖRON HASTALIKLARI**
 DEMİR E., ÖZBUDAK P.
 in: Temel Pediatrik Nöroloji: Tanı ve Tedavi, Kumandaş Sefer, Cankurtaran Mehmet, Editor, AKADEMİSYEN KİTABEVİ, Ankara, pp.3003-3029, 2022
- V. **Dikkat Eksikliği ve Hiperaktivite Bozukluğu ile Nörolojik Bozukluklar**
 ÖZTÜRK Z., DEMİR E.
 in: Dikkat Eksikliği ve Hiperaktivite Bozukluğu, Azime Şebnem Soysal Acar, Editor, NOBEL AKADEMİK YAYINCILIK EĞİTİM DANIŞMANLIK TİC. LTD. ŞTİ., Ankara, pp.458-467, 2019
- VI. **40. Bölüm. DEHB ve Nörolojik Bozukluklar**
 öztürk z., DEMİR E.
 in: DİKKAT EKSİKLİĞİ HİPERAKTİVİTE BOZUKLUĞU EL KİTABI, A. Şebnem Soysal Acar, Editor, Nobel Akademik Yayıncılık Eğitim Danışmanlık Tic. Ltd. Şti., Ankara, pp.455-464, 2019

Papers Published in Refereed Scientific Meetings

- I. **1-3 Yaş Arası Epilepsili Çocukların Büyüme Durumunun Anneleri Tarafından Değerlendirilmesi**
 Çelik B. İ., KARAKAŞ N. M., DEMİR E., SERDAROĞLU E.
 68. TÜRKİYE MILLİ PEDIATRI KONGRESİ VE 1. ULUSLARARASI TÜRKİYE MILLİ PEDIATRI DERNEĞİ KONGRESİ, Antalya, Turkey, 20 - 24 November 2024, pp.10-11
- II. **Rituksimab'a Bağlı Hipogamaglobulinemi Gelişen Nöromiyelitis Optika Spektrum Bozukluğu Olgu Sunumu**
 Amrahova P., Çetin H., Baskın A. K., Serdaroğlu E., Arhan E., Hırfanoğlu T., Demir E., Öztürk Z.
 25.Uluslararası Çocuk Nöroloji Kongresi, Ankara, Turkey, 22 - 24 May 2024
- III. **The effect of new third-generation antiseizure drugs on seizure recurrence after drug**

- discontinuation (OC-11)**
Cebeci D., Arhan E., Demir E., Hırfanoğlu T., Güçüyener K., Serdaroğlu E., Aydın K., Serdaroğlu A.
25. Ulusal Çocuk Nörolojisi Kongresi, Ankara, Turkey, 22 - 26 May 2024, pp.27
- IV. A Case Report of Neuromyelitis Optica Spectrum Disorder with Rituximab-Induced Hypogammaglobulinemia (PP-091)**
Amrahova P., Çetin H., Baskın A. K., Serdaroğlu E., Arhan E., Hırfanoğlu T., Demir E., Öztürk Z.
25. Ulusal Çocuk Nörolojisi Kongresi, Ankara, Turkey, 22 - 26 May 2024, pp.102
- V. Cerebral Palsy from a Child Neurologist's Perspective (PP-159)**
Taş E. N., Amrahova P., Öztürk Z., Serdaroğlu E., Hırfanoğlu T., Demir E., Arhan E.
25. Ulusal Çocuk Nörolojisi Kongresi, Ankara, Turkey, 22 - 26 May 2024, pp.140
- VI. Lökodistrofi benzeri patern ile başvuran MOGAD olgusu**
ÇETİN H., GÜRSOY M., YAZOL M., ÖZTÜRK Z., SERDAROĞLU E., HIRFANOĞLU T., AYDIN K., DEMİR E., ARHAN E.
25. Ulusal Çocuk Nörolojisi Kongresi, Ankara, Turkey, 22 May 2024
- VII. Optic Neuritis: Familial Mediterranean Fever vs Demyelinating Disease? (PP-090)**
Amrahova P., Gezgin Yıldırım D., Serdaroğlu E., Arhan E., Hırfanoğlu T., Bakkaloğlu Ezgü S. A., Demir E., Öztürk Z.
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- VIII. Applicability of the Diagnostic Criteria of the ILAE 2022 Classification of Childhood Epileptic Syndromes: Retrospective study of 1000 children with epilepsy (OC-10)**
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Congress and Symposium Activities

25. ULUSAL ÇOCUK NÖROLOJİ KONGRESİ, Session Moderator, Ankara, Turkey, 2024
- VI. NÖROMUSKÜLER HASTALIKLAR KONGRESİ, Session Moderator, Antalya, Turkey, 2023
- GENÇ MERAKLILAR İÇİN PEDIATRİK NÖROMUSKÜLER HASTALIKLAR KURSU, Invited Speaker, Antalya, Turkey, 2023
1. Uluslararası Katılımlı Erciyes Türk Dünyası Çocuk Nörolojisi Kongresi, Invited Speaker, Kayseri, Turkey, 2023
- GENÇ MERAKLILAR İÇİN PEDIATRİK NÖROMUSKÜLER HASTALIKLAR KURSU, Invited Speaker, İstanbul, Turkey, 2023
23. ULUSAL ÇOCUK NÖROLOJİ KONGRESİ, Session Moderator, İzmir, Turkey, 2021
- V. NÖROMUSKULER HASTALIKLAR KONGRESİ, Invited Speaker, Samsun, Turkey, 2021

Non Academic Experience

U 523, INSTITUT DE MYOLOGIE, GROUPE HOSPITALIER PITIÉ-SALPÉTRIÈRE, PARIS, FRANSA, Moleküler genetik alanında doktora sonrası fellow
TEPEBAŞI POLİKLİNİĞİ, UZMAN ÇOCUK DOKTORU