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Öğrenim Bilgisi

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The University of Michigan, Medical School, Pediatric Nephrology, Amerika
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2000 - 2002

Hacettepe Üniversitesi, Tıp Fakültesi, Çocuk Nefrolojisi, Türkiye

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1994 - 2000

Hacettepe Üniversitesi, Tıp Fakültesi, Çocuk Sağlığı Ve Hastalıkları, Türkiye

Lisans
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Ege Üniversitesi, Tıp Fakültesi, Türkiye

Yabancı Diller

İngilizce, B2 Orta Üstü

Akademik Unvanlar / Görevler

Prof. Dr.
2018 - Devam Ediyor

Hacettepe Üniversitesi, Tıp Fakültesi (İngilizce), Dahili Tıp Bilimleri Bölümü

Doç. Dr.
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2005 - 2006

Hacettepe Üniversitesi, Tıp Fakültesi (İngilizce), Dahili Tıp Bilimleri Bölümü

Desteklenen Projeler

1. ÖZALTIN F., TOPALOĞLU R., TÜRKOĞLU Ö., ATAYAR E., GÜLHAN B., Yükseköğretim Kurumları Destekli Proje,

- Çocukluk Çağı Otozomal Dominant ve Otozomal Resesif Polikistik Böbrek Hastalıklarının Genetik ve Klinik Özelliklerinin Araştırılması, 2019 - 2021
2. ÖZALTIN F., TOPALOĞLU R., EROĞLU İ., NEMUTLU E., ÖZBAY H. S., YABANOĞLU ÇİFTÇİ S., Yükseköğretim Kurumları Destekli Proje, CoQ10 Nefropatisinde Koenzim Q10 İçerikli Nanoterapötiklerin Tedavi Etkiliğinin Araştırılması ve İn Vitro Modelde Mitokondriyal Fonksiyonlarının Karşılaştırılması, 2018 - 2021
 3. ÖZALTIN F., BEŞBAŞ N., Yükseköğretim Kurumları Destekli Proje, Atipik Hemolitik Üremik Sendromlu Hastalarda Kompleman Düzenleyici Genlerdeki Mutasyonların Saptanması, 2011 - 2016

SCI, SSCI ve AHCI İndekslerine Giren Dergilerde Yayınlanan Makaleler

1. **Calcineurin inhibitor-related hyperkalemia is caused by hyporeninemic hypoaldosteronism and fludrocortisone is an effective treatment: Report of a case series and review of the literature**
ÜNSAL Y., BALTU D., GÜLHAN B., OKUR F. V., ÖZALTIN F., DÜZOVA A., TOPALOĞLU R., ÖZÖN Z. A., GÖNC E. N. PEDIATRIC TRANSPLANTATION, sa.4, 2024 (SCI-Expanded)
2. **Implication of transcription factor FOXD2 dysfunction in syndromic congenital anomalies of the kidney and urinary tract (CAKUT)**
Riedhammer K. M., Nguyen T. T., KOŞUKCU C., Calzada-Wack J., Li Y., Assia Batzir N., SAYGILI S. K., Wimmers V., Kim G., Chrysanthou M., et al.
Kidney International, cilt.105, sa.4, ss.844-864, 2024 (SCI-Expanded)
3. **Steroid-Resistant Nephrotic Syndrome due to NPHS2 Variants Is Not Associated With Posttransplant Recurrence**
Kachmar J., Boyer O., Lipska-Ziętkiewicz B., Morinière V., Gribouval O., Heidet L., Balasz-Chmielewska I., Benetti E., Cloarec S., Csaicsich D., et al.
Kidney International Reports, cilt.9, sa.4, ss.973-981, 2024 (SCI-Expanded)
4. **Acute kidney injury in children with moderate-severe COVID-19 and multisystem inflammatory syndrome in children: a referral center experience**
TAŞTEMEL ÖZTÜRK T., DÜZOVA A., OYGAR P. D., BALTU D., ÖZÇİLİNİR P., LAÇİNEL GÜRLEVİK S., KURT ŞÜKÜR E. D., AYKAN H. H., ÖZEN S., ERTUĞRUL İ., et al.
Pediatric Nephrology, cilt.39, sa.3, ss.867-877, 2024 (SCI-Expanded)
5. **Omic Studies on In Vitro Cystinosis Model: siRNA-Mediated CTNS Gene Silencing in HK-2 Cells**
Baysal İ., YABANOĞLU ÇİFTÇİ S., NEMUTLU E., EYLEM C. C., Gök-Topak E. D., ULUBAYRAM K., KIR S., GÜLHAN B., UÇAR G., ÖZALTIN F., et al.
Laboratory investigation; a journal of technical methods and pathology, cilt.104, sa.1, ss.100287, 2024 (SCI-Expanded)
6. **Management of pediatric hemolytic uremic syndrome**
GÜLHAN B., ÖZALTIN F., Fidan K., ÖZÇAKAR Z. B., Söylemezoğlu O.
Turkish Journal of Pediatrics, cilt.66, sa.1, ss.1-16, 2024 (SCI-Expanded)
7. **Genotype/phenotype relationship in mild congenital nephrotic syndrome**
Mulic B., Peco-Antic A., ÖZALTIN F.
SRPSKI ARHIV ZA CELOKUPNO LEKARSTVO, sa.1-2, ss.81-84, 2024 (SCI-Expanded)
8. **A novel homozygous missense variant in TBC1D31 in a consanguineous family with congenital anomalies of the kidney and urinary tract (CAKUT)**
SAYGILI S. K., KOŞUKCU C., BAŞTUĞ T., DOĞAN EKİCİ A. İ., YILMAZ E. M., Kalyoncu A. U., Ağbaş A., CANPOLAT N., Çalışkan S., ÖZALTIN F.
Clinical Genetics, cilt.104, sa.6, ss.679-685, 2023 (SCI-Expanded)
9. **Long-term kidney follow-up after pediatric acute kidney support therapy for children less than 15 kg**
GÜLÇEK Ö. N., GÜLHAN B., KESİCİ S., KURT ŞÜKÜR E. D., HAYRAN K. M., ÖZALTIN F., DÜZOVA A., BAYRAKCİ B., TOPALOĞLU R.
Pediatric Nephrology, cilt.38, sa.11, ss.3811-3821, 2023 (SCI-Expanded)

10. **COVID-19 associated thrombotic microangiopathy**
YILMAZ E. M., Cebi M. N., Karahan I., SAYGILI S. K., Gulmez R., Demirgan E. B., Durak C., AYGÜN F., ÖZALTIN F., Caliskan S., et al.
Nephrology, cilt.28, sa.10, ss.557-560, 2023 (SCI-Expanded)
11. **Outcomes of steroid-resistant nephrotic syndrome in children not treated with intensified immunosuppression**
Trautmann A., Seide S., Lipska-Zietkiewicz B. S., ÖZALTIN F., Szczepanska M., Azocar M., Jankauskiene A., Zurowska A., ÇALIŞKAN S., Saeed B., et al.
PEDIATRIC NEPHROLOGY, cilt.38, sa.5, ss.1499-1511, 2023 (SCI-Expanded)
12. **A rare cause of nephrotic syndrome-sphingosine-1-phosphate lyase (SGPL1) deficiency: 6 cases and a review of the literature**
TAŞTEMEL ÖZTÜRK T., CANPOLAT N., SAYGILI S. K., BAYRAKCİ U. S., Soylemezoglu O., ÖZALTIN F., TOPALOĞLU R.
PEDIATRIC NEPHROLOGY, cilt.38, sa.3, ss.711-719, 2023 (SCI-Expanded)
13. **HOXA11 is another monogenic cause of congenital anomalies of the kidney and urinary tract**
SAYGILI S. K., ÇALIŞKAN S., ÖZALTIN F.
PEDIATRIC NEPHROLOGY, cilt.38, sa.3, ss.933-934, 2023 (SCI-Expanded)
14. **Metabolomic Analyses to Identify Candidate Biomarkers of Cystinosis**
NEMUTLU E., ÖZALTIN F., YABANOĞLU ÇİFTÇİ S., GÜLHAN B., Eylem C. C., BAYSAL İ., Gök-Topak E. D., ULUBAYRAM K., SEZERMAN O. U., UÇAR G., et al.
International Journal of Molecular Sciences, cilt.24, sa.3, 2023 (SCI-Expanded)
15. **Predictors of kidney complications and analysis of hypertension in children with allogeneic hematopoietic stem cell transplantation**
Gurbanov A., GÜLHAN B., KUŞKONMAZ B. B., OKUR F. V., ÖZALTIN F., DÜZOVA A., ÇETİNKAYA F. D., TOPALOĞLU R.
PEDIATRIC NEPHROLOGY, cilt.38, sa.2, ss.461-469, 2023 (SCI-Expanded)
16. **Hemoglobin cast nephropathy: a rare but serious complication of hemolysis in a pediatric patient**
BALTU D., Oral N. A., KESİCİ S., TOPALOĞLU R., ÖZCEBE O. İ., AKSU T., ORHAN D., ÖZALTIN F.
Turkish Journal of Pediatrics, cilt.65, sa.5, ss.874-880, 2023 (SCI-Expanded)
17. **The Clinical and Mutational Spectrum of 69 Turkish Children with Autosomal Recessive or Autosomal Dominant Polycystic Kidney Disease: A Multicenter Retrospective Cohort Study**
Tutal O., GÜLHAN B., Atayar E., Yuksel S., ÖZÇAKAR Z. B., Soylemezoglu O., SAYGILI S. K., ÇALIŞKAN S., Inozu M., Baskin E., et al.
Nephron, 2023 (SCI-Expanded)
18. **Response to Dr. Spizzirri et al**
TAŞTEMEL ÖZTÜRK T., ÖZALTIN F., TOPALOĞLU R.
Pediatric Nephrology, cilt.38, sa.1, ss.309-310, 2023 (SCI-Expanded)
19. **Hearing Loss Related to Gene Mutations in Distal Renal Tubular Acidosis**
Ay E., GÜRSES E., ASLAN F., GÜLHAN B., Alniacik A., DÜZOVA A., BAJİN M. D., SENNAROĞLU L., GENÇ G. A., ÖZALTIN F., et al.
Audiology & neuro-otology, cilt.28, sa.5, ss.350-359, 2023 (SCI-Expanded)
20. **Oral Coenzyme Q10 supplementation leads to better preservation of kidney function in steroid-resistant nephrotic syndrome due to primary Coenzyme Q10 deficiency**
Drovandi S., Lipska-Zietkiewicz B. S., ÖZALTIN F., Emma F., GÜLHAN B., Boyer O., Trautmann A., Xu H., Shen Q., Rao J., et al.
KIDNEY INTERNATIONAL, cilt.102, sa.3, ss.604-612, 2022 (SCI-Expanded)
21. **Variation of the clinical spectrum and genotype-phenotype associations in Coenzyme Q10 deficiency associated glomerulopathy**
Drovandi S., Lipska-Zietkiewicz B. S., ÖZALTIN F., Emma F., GÜLHAN B., Boyer O., Trautmann A., Zietkiewicz S., Xu H., Shen Q., et al.
KIDNEY INTERNATIONAL, cilt.102, sa.3, ss.592-603, 2022 (SCI-Expanded)
22. **A broad clinical spectrum of PLC epsilon 1-related kidney disease and intrafamilial variability**
Yilmaz E. K., Saygili S., GÜLHAN B., Canpolat N., KARABAY BAYAZIT A., KILIÇ B. D., Akinci N., Benzer M., GÖKNAR N.,

- KAVAZ TUFAN A., et al.
PEDIATRIC NEPHROLOGY, cilt.37, sa.8, ss.1855-1866, 2022 (SCI-Expanded)
23. Long-term renal survival of paediatric patients with lupus nephritis
Demir S., GÜLHAN B., ÖZEN S., Çelēgen K., BATU AKAL E. D., Taş N., ORHAN D., BİLGİNER Y., DÜZOVA A., ÖZALTIN F., et al.
Nephrology Dialysis Transplantation, cilt.37, sa.6, ss.1069-1077, 2022 (SCI-Expanded)
24. Eculizumab treatment and discontinuation in pediatric patients with atypical hemolytic uremic syndrome: a multicentric retrospective study
Baskin E., Fidan K., GÜLHAN B., Gulleroglu K., CANPOLAT N., Yilmaz A., Parmakiz G., ÖZÇAKAR Z. B., ÖZALTIN F., Soylemezoglu O.
JOURNAL OF NEPHROLOGY, cilt.35, sa.4, ss.1213-1222, 2022 (SCI-Expanded)
25. Clinically Different Presentations of Family Members With the Same Homozygote Diacylglycerol Kinase Epsilon Mutation: Case Report
Çelikkaya E., Güngör T., Karakaya D., Çakıcı E. K., Yazılıtaş F., ÖZALTIN F., Bülbül M.
Experimental and Clinical Transplantation, cilt.20, sa.5, ss.45-48, 2022 (SCI-Expanded)
26. Mitochondria-targeted CoQ(10) loaded PLGA-b-PEG-TPP nanoparticles: Their effects on mitochondrial functions of COQ8B(-/-) HK-2 cells
Sena Ozbay H., Yabanoglu-Ciftci S., Baysal I., Gultekinoglu M., Can Eylem C. C., Ulubayram K., Nemutlu E., Topaloglu R., Ozaltin F.
EUROPEAN JOURNAL OF PHARMACEUTICS AND BIOPHARMACEUTICS, cilt.173, ss.22-33, 2022 (SCI-Expanded)
27. A splice site mutation in the TSEN2 causes a new syndrome with craniofacial and central nervous system malformations, and atypical hemolytic uremic syndrome
CANPOLAT N., Liu D., Atayar E., SAYGILI S. K., Kara N. S., Westfall T. A., Ding Q., Brown B. J., Braun T. A., Slusarski D., et al.
CLINICAL GENETICS, cilt.101, ss.346-358, 2022 (SCI-Expanded)
28. Mitigation of portal fibrosis and cholestatic liver disease in ANKS6-deficient livers by macrophage depletion
Airik M., McCourt B., TAŞTEMEL ÖZTÜRK T., Huynh A. B., Zhang X., Tometich J. T., TOPALOĞLU R., ÖZEN H., ORHAN D., Nejak-Bowen K., et al.
FASEB JOURNAL, cilt.36, sa.2, 2022 (SCI-Expanded)
29. Glomerulonephritis with crescents in childhood; etiologies and significance of M2 macrophages
KAYKI G., ORHAN D., GÜLHAN B., TOPALOĞLU R., AKÇÖREN Z., DÜZOVA A., ÖZALTIN F., ÖZEN S., BİLGİNER Y., Gucer S.
TURKISH JOURNAL OF PEDIATRICS, cilt.64, sa.1, ss.59-68, 2022 (SCI-Expanded)
30. Could plasma based therapies still be considered in selected cases with atypical hemolytic uremic syndrome?
ÖZLÜ S. G., GÜLHAN B., Aydog O., Atayar E., Delibas A., Parmaksiz G., Ozdogan E. B., ÇOMAK E., Tasdemir M., Acar B., et al.
TURKISH JOURNAL OF PEDIATRICS, cilt.63, sa.6, ss.986-993, 2021 (SCI-Expanded)
31. An international cohort study spanning five decades assessed outcomes of nephropathic cystinosis
Emma F., van't Hoff W., Hohenfellner K., TOPALOĞLU R., Greco M., Ariceta G., Bettini C., Bockenhauer D., Veys K., Pape L., et al.
KIDNEY INTERNATIONAL, cilt.100, sa.5, ss.1112-1123, 2021 (SCI-Expanded)
32. AGTR1-RELATED RENAL TUBULAR DYSGENESSES MAY NOT BE FATAL
Demirgan E. B., Saygili S., CANPOLAT N., Sever L., Kilicaslan I., Taylan D., ÇALIŞKAN S., ÖZALTIN F.
PEDIATRIC NEPHROLOGY, cilt.36, sa.10, ss.3455, 2021 (SCI-Expanded)
33. TAKAYASU ARTERITIS WITH RENAL ARTERY INVOLVEMENT IN CHILDREN: 12 YEARS EXPERIENCE OF A TERTIARY CENTER
TAŞTEMEL ÖZTÜRK T., BALTU D., KURT ŞÜKÜR E. D., GÜLHAN B., ÖZEN S., DÜZOVA A., ÖZALTIN F., TOPALOĞLU R.
PEDIATRIC NEPHROLOGY, cilt.36, sa.10, ss.3441, 2021 (SCI-Expanded)
34. PRIMARY COQ10 DEFICIENCY: CLINICAL SPECTRUM AND GENOTYPE-PHENOTYPE CORRELATIONS

- Drovandi S., Lipska-zietkiewicz B. S., ÖZALTIN F., Emma F., GÜLHAN B., Boyer O., Trautmann A., Shen Q., Rao J., Riedhammer K. M., et al.
PEDIATRIC NEPHROLOGY, cilt.36, sa.10, ss.3445-3446, 2021 (SCI-Expanded)
35. **CHARACTERISTICS AND OUTCOME OF BK VIRUS INFECTION IN PEDIATRIC RENAL TRANSPLANT RECIPIENT**
KURT ŞÜKÜR E. D., Ozdemir G., TAŞTEMEL ÖZTÜRK T., BALTU D., GÜLHAN B., ÖZALTIN F., DÜZOVA A., TOPALOĞLU R.
PEDIATRIC NEPHROLOGY, cilt.36, sa.10, ss.3460, 2021 (SCI-Expanded)
36. **EFFECTS OF RAAS INHIBITION AND IMMUNOSUPPRESSIVE THERAPY IN PEDIATRIC PATIENTS WITH X-LINKED ALPORT SYNDROME**
Ozdemir G., GÜLHAN B., KURT ŞÜKÜR E. D., Atayar E., DURSUN İ., ÖZÇAKAR Z. B., Saygili S., SOYLU A., Soylemezoglu O., Yilmaz A., et al.
PEDIATRIC NEPHROLOGY, cilt.36, sa.10, ss.3366, 2021 (SCI-Expanded)
37. **INFANTILE NEPHROTIC SYNDROME WITH PLCE1 MUTATION; TREATMENT OF CYCLOSPORINE-3-YEAR FOLLOW-UP**
Tekcan D., Nalcacioglu H., Onal H. G., ÖZALTIN F., AYDOĞ Ö.
PEDIATRIC NEPHROLOGY, cilt.36, sa.10, ss.3475-3476, 2021 (SCI-Expanded)
38. **LONG TERM FOLLOW UP IN RENAL SCARRING OF URINARY TRACT INFECTION: ALBUMINURIA, DIASTOLIC BLOOD PRESSURE**
BALTU D., VOLCAN SALANCI B., GÜLHAN B., TAŞTEMEL ÖZTÜRK T., KURT ŞÜKÜR E. D., ÖZALTIN F., DÜZOVA A., TOPALOĞLU R.
PEDIATRIC NEPHROLOGY, cilt.36, sa.10, ss.3434, 2021 (SCI-Expanded)
39. **NEPHROTIC SYNDROME WITH MUTATION IN SPHINGOSINE-1-PHOSPHATE LYASE: 6 CASES**
TAŞTEMEL ÖZTÜRK T., CANPOLAT N., Saygili S., BAYRAKCİ U. S., Soylemezoglu O., ÖZALTIN F., TOPALOĞLU R.
PEDIATRIC NEPHROLOGY, cilt.36, sa.10, ss.3476, 2021 (SCI-Expanded)
40. **CLINICAL AND MUTATIONAL SPECTRUM OF CHILDREN WITH AUTOSOMAL RECESSIVE AND AUTOSOMAL DOMINANT POLYCYSTIC KIDNEY DISEASE**
Tutal O., GÜLHAN B., Atayar E., YÜKSEL S., Ozcakar B., Soylemezoglu O., Saygili S., Inozu M., Baskin E., DÜZOVA A., et al.
PEDIATRIC NEPHROLOGY, cilt.36, sa.10, ss.3421-3422, 2021 (SCI-Expanded)
41. **AUDIOLOGICAL FINDINGS IN DISTAL RENAL TUBULAR ACIDOSIS**
Ay E., GÜRSES E., Arslan F., GÜLHAN B., Alniacik A., DÜZOVA A., BAJİN M. D., SENNAROĞLU L., GENÇ G. A., ÖZALTIN F., et al.
PEDIATRIC NEPHROLOGY, cilt.36, sa.10, ss.3428, 2021 (SCI-Expanded)
42. **A HOMOZYGOUS HOXA11 VARIATION AS A POTENTIAL NOVEL CAUSE OF AUTOSOMAL RECESSIVE CONGENITAL ANOMALIES OF THE KIDNEY AND URINARY TRACT**
Saygili S., Atayar E., CANPOLAT N., ELİÇEVİK M., KURUĞOĞLU S., Sever L., ÇALIŞKAN S., ÖZALTIN F.
PEDIATRIC NEPHROLOGY, cilt.36, sa.10, ss.3404-3405, 2021 (SCI-Expanded)
43. **DGKE MUTATION IN A CHILD TREATED WITH ECULIZUMAB**
Tekcan D., Nalcacioglu H., Karadag S. I. K., Onal H. G., ÖZALTIN F., AYDOĞ Ö.
PEDIATRIC NEPHROLOGY, cilt.36, sa.10, ss.3403, 2021 (SCI-Expanded)
44. **Outcome of diacylglycerol kinase epsilon-mediated hemolytic uremic syndrome in an infant**
KOYUN M., KAYA AKSOY G., ÇOMAK E., ÖZALTIN F., AKMAN S.
KIDNEY INTERNATIONAL, cilt.99, sa.6, ss.1500-1501, 2021 (SCI-Expanded)
45. **Transplantation in pediatric aHUS within the era of eculizumab therapy**
ÖZÇAKAR Z. B., ÖZALTIN F., GÜLHAN B., ÇOMAK E., Parmaksız G., Baskin E., TOPALOĞLU R., Kasap Demir B., Canpolat N., Yuruk Yildirim Z., et al.
PEDIATRIC TRANSPLANTATION, cilt.25, sa.3, 2021 (SCI-Expanded)
46. **EARLY URINARY SYSTEM COMPLICATIONS IN CHILDREN WITH HEMATOPOETIC STEM CELL TRANSPLANTATION**
Gurbanov A., GÜLHAN B., KUŞKONMAZ B. B., OKUR F. V., ÇETİNKAYA F. D., Ozdemir G., TAŞ N., Celegen K., ÖZALTIN

- F., DÜZOVA A., et al.
NEPHROLOGY DIALYSIS TRANSPLANTATION, cilt.36, 2021 (SCI-Expanded)
47. **CLINICAL AND MUTATIONAL SPECTRUM OF CHILDREN WITH AUTOSOMAL RECESSIVE AND AUTOSOMAL DOMINANT POLYCYSTIC KIDNEY DISEASE**
Tutal O., GÜLHAN B., Atayar E., YÜKSEL S., ÖZÇAKAR Z. B., Soylemezoglu O., Saygili S., Inozu M., Baskin E., DÜZOVA A., et al.
NEPHROLOGY DIALYSIS TRANSPLANTATION, cilt.36, 2021 (SCI-Expanded)
48. **Management of congenital nephrotic syndrome: consensus recommendations of the ERKNet-ESPN Working Group**
Boyer O., Schaefer F., Haffner D., Bockenhauer D., Holtta T., Berody S., Webb H., Heselden M., Lipska-Zietkiewicz B. S., ÖZALTIN F., et al.
NATURE REVIEWS NEPHROLOGY, cilt.17, sa.4, ss.277-289, 2021 (SCI-Expanded)
49. **AGTR1-related Renal Tubular Dysgeneses May Not Be Fatal**
Demirgan E. B., SAYGILI S. K., CANPOLAT N., Sever L., Kilicaslan I., Taylan D., ÇALIŞKAN S., ÖZALTIN F.
KIDNEY INTERNATIONAL REPORTS, cilt.6, sa.3, ss.846-852, 2021 (SCI-Expanded)
50. **Predictors for the use of herbal and dietary supplements in children and adolescents with kidney and urinary tract diseases**
Ozturk T. T., KANBUR N., ÖZMERT E. N., GÜLHAN B., ÖZALTIN F., TOPALOĞLU R., DÜZOVA A.
EUROPEAN JOURNAL OF PEDIATRICS, cilt.180, sa.1, ss.253-262, 2021 (SCI-Expanded)
51. **Acute kidney injury in a patient with COVID-19: Answers**
TAŞTEMEL ÖZTÜRK T., BALTU D., KURT ŞÜKÜR E. D., ÖZSÜREKCİ Y., Gucer S., BAŞARAN H. Ö., GÜLHAN B., ÖZALTIN F., DÜZOVA A., TOPALOĞLU R.
PEDIATRIC NEPHROLOGY, cilt.36, sa.12, ss.4111-4113, 2020 (SCI-Expanded)
52. **Acute kidney injury in a patient with COVID-19: Questions**
Tastemel Ozturk T., Baltu D., Kurt Sukur E. D., Ozsurekci Y., Gucer S., Basaran O., Gulhan B., Ozaltin F., Duzova A., Topaloglu R.
PEDIATRIC NEPHROLOGY, cilt.36, sa.12, ss.4109-4110, 2020 (SCI-Expanded)
53. **Determinants of outcomes in chronic pediatric peritoneal dialysis: a single center experience**
Asi T., DÜZOVA A., DOĞAN H. S., Karakurt G., BAHADIR Ö. F., Bozaci A. C., GÜLHAN B., ÖZALTIN F., AKI F. T., TEKGÜL S., et al.
TURKISH JOURNAL OF PEDIATRICS, cilt.62, sa.6, ss.940-948, 2020 (SCI-Expanded)
54. **Genetic aspects of congenital nephrotic syndrome: a consensus statement from the ERKNet-ESPN inherited glomerulopathy working group**
Lipska-Zietkiewicz B. S., ÖZALTIN F., Holtta T., Bockenhauer D., Berody S., Levchenko E., Vivarelli M., Webb H., Haffner D., Schaefer F., et al.
EUROPEAN JOURNAL OF HUMAN GENETICS, cilt.28, sa.10, ss.1368-1378, 2020 (SCI-Expanded)
55. **Renal Biopsy Prognostic Findings in Children With Atypical Hemolytic Uremic Syndrome**
YÜKSEL S., GÖNÜL İ. İ., Canpolat N., GÖKCE İ., Ozlu S. G., ÖZÇAKAR Z. B., ÖZALTIN F., SÖYLEMEZOĞLU O.
PEDIATRIC AND DEVELOPMENTAL PATHOLOGY, cilt.23, sa.5, ss.362-371, 2020 (SCI-Expanded)
56. **COL4A3 mutation is an independent risk factor for poor prognosis in children with Alport syndrome**
Ozdemir G., GÜLHAN B., Atayar E., Saygili S., Soylemezoglu O., Ozcakar Z. B., Eroglu F. K., Candan C., Demir B. K., Soylu A., et al.
PEDIATRIC NEPHROLOGY, cilt.35, sa.10, ss.1941-1952, 2020 (SCI-Expanded)
57. **A homozygous HOXA11 variation as a potential novel cause of autosomal recessive congenital anomalies of the kidney and urinary tract**
Saygili S., Atayar E., Canpolat N., ELİÇEVİK M., KURUĞOĞLU S., Sever L., Caliskan S., ÖZALTIN F.
CLINICAL GENETICS, cilt.98, sa.4, ss.390-395, 2020 (SCI-Expanded)
58. **Clinical characteristics of children with congenital anomalies of the kidney and urinary tract and predictive factors of chronic kidney disease**
Cetinkaya P. G., GÜLHAN B., DÜZOVA A., Besbas N., HAYRAN K. M., TOPALOĞLU R., ÖZALTIN F.
TURKISH JOURNAL OF PEDIATRICS, cilt.62, sa.5, ss.746-755, 2020 (SCI-Expanded)

59. **Cystinosis beyond kidneys: gastrointestinal system and muscle involvement**
TOPALOĞLU R., GÜLTEKİNGİL KESER A., GÜLHAN B., ÖZALTIN F., DEMİR H., Ciftci T., DEMİR N., TEMUÇİN Ç. M., YÜCE A., Akhan O.
BMC GASTROENTEROLOGY, cilt.20, sa.1, 2020 (SCI-Expanded)
60. **CD80 expression and infiltrating regulatory T cells in idiopathic nephrotic syndrome of childhood**
Eroglu F. K., ORHAN D., Inozu M., DÜZOVA A., GÜLHAN B., ÖZALTIN F., TOPALOĞLU R.
PEDIATRICS INTERNATIONAL, cilt.61, sa.12, ss.1250-1256, 2019 (SCI-Expanded)
61. **Surgical management of renovascular hypertension in children and young adults: a 13-year experience**
Peker O., AKI F. T., Kumbasar U., GÜVENER M., YILMAZ M., DOĞAN R., ÖZALTIN F., DÜZOVA A., TOPALOĞLU R., PEYNİRCİOĞLU B., et al.
INTERACTIVE CARDIOVASCULAR AND THORACIC SURGERY, cilt.29, sa.5, ss.746-752, 2019 (SCI-Expanded)
62. **Long-term follow-up results of patients with ADCK4 mutations who have been diagnosed in the asymptomatic period: effects of early initiation of CoQ10 supplementation**
ATMACA M., GÜLHAN B., Atayar E., KARABAY BAYAZIT A., CANDAN C., ARICI M., TOPALOĞLU R., ÖZALTIN F.
TURKISH JOURNAL OF PEDIATRICS, cilt.61, sa.5, ss.657-663, 2019 (SCI-Expanded)
63. **Rituximab for Children With Difficult-to-Treat Nephrotic Syndrome: Its Effects on Disease Progression and Growth**
TOPALOĞLU R., GÜLHAN B., Celegen K., Inozu M., Hayran M., Duzoya A., ÖZALTIN F.
FRONTIERS IN PEDIATRICS, cilt.7, 2019 (SCI-Expanded)
64. **An immunohistochemical approach to detect oncogenic CTNNB1 mutations in primary neoplastic tissues**
AKYOL A., GUNER G., OZSEKER H. S., İŞIK A., Atci O., UZUN S., ATAYAR E., ÖZALTIN F., GEDIKOGLU G., SÖKMENSÜER C., et al.
LABORATORY INVESTIGATION, cilt.99, sa.1, ss.128-137, 2019 (SCI-Expanded)
65. **Persistent hypoglycemic attacks during hemodialysis sessions in an infant with congenital nephrotic syndrome: Questions and Answers**
Saygili S., CANPOLAT N., Sever L., ÇALIŞKAN S., Atayar E., ÖZALTIN F.
PEDIATRIC NEPHROLOGY, cilt.34, sa.1, ss.75-79, 2019 (SCI-Expanded)
66. **BK virus associated nephropathy and severe pneumonia in a kidney transplanted adolescent with Schimke immune-osseous-dysplasia**
DÜZOVA A., GÜLHAN B., TOPALOĞLU R., ÖZALTIN F., CENGİZ A. B., Yetimakman A. F., Dogru D., Gucer S., Besbas N.
TURKISH JOURNAL OF PEDIATRICS, cilt.61, sa.1, ss.111-116, 2019 (SCI-Expanded)
67. **Response to Early Coenzyme Q10 Supplementation Is not Sustained in CoQ10 Deficiency Caused by CoQ2 Mutation**
Eroglu F. K., ÖZALTIN F., Gonç N., Nalcacioglu H., Ozcakar Z. B., YALNIZOĞLU D., Gucer S., ORHAN D., Eminoglu F. T., GÖÇMEN R., et al.
PEDIATRIC NEUROLOGY, cilt.88, ss.71-74, 2018 (SCI-Expanded)
68. **EARLY USE OF ECULIZUMAB IN A CASE OF STEC-HUS BEFORE THE DEVELOPMENT OF CLINICAL SYMPTOMS**
BAYRAM M. T., DEMİR B. K., SOYLU A., KAVUKÇU S., ÖZALTIN F.
PEDIATRIC NEPHROLOGY, cilt.33, sa.10, ss.1845, 2018 (SCI-Expanded)
69. **LONG-TERM FOLLOW-UP RESULTS OF PATIENTS WITH ADCK4 MUTATIONS WHO HAVE BEEN DIAGNOSED IN ASYMPTOMATIC PERIOD: EFFECTS OF EARLY INITIATION OF COQ10 SUPPLEMENTATION**
ATMACA M., GÜLHAN B., KARABAY BAYAZIT A., CANDAN C., ARICI M., TOPALOĞLU R., ÖZALTIN F.
PEDIATRIC NEPHROLOGY, cilt.33, sa.10, ss.1821-1822, 2018 (SCI-Expanded)
70. **GASTROINTESTINAL SYSTEM INVOLVEMENT IN ATYPICAL HEMOLYTIC UREMIC SYNDROME**
Fidan K., Yıldırım Z. Y., Goknar N., GÜLHAN B., Gulleroglu K., Ozcakar Z. B., Baskin E., Hayran M., ÖZALTIN F., Soylemezoglu O.
PEDIATRIC NEPHROLOGY, cilt.33, sa.10, ss.1843, 2018 (SCI-Expanded)

71. **ACUTE HEMODIALYSIS EXPERIENCE IN CHILDREN WEIGHING LESS THAN 15 KG**
TAŞ N., GÜLHAN B., Celegen K., Ozdemir G., DÜZOVA A., ÖZALTIN F., TOPALOĞLU R.
PEDIATRIC NEPHROLOGY, cilt.33, sa.10, ss.1893, 2018 (SCI-Expanded)
72. **GASTRIC DUPLICATION CYST IN AN INFANT WITH FINNISH-TYPE CONGENITAL NEPHROTIC SYNDROME: CONCURRENCE OR COINCIDENCE?**
Eroglu T. G. F. K., Can G. G., Cakici E. K., Yazilitas F., Celikkaya E., ÖZALTIN F., Bulbul M.
PEDIATRIC NEPHROLOGY, cilt.33, sa.10, ss.1931-1932, 2018 (SCI-Expanded)
73. **LUPUS NEPHRITIS: LONG TERM FOLLOW-UP AND EFFECT OF TREATMENT ON GROWTH**
TOPALOĞLU R., TAŞ N., Celegen K., Ozdemir G., GÜLHAN B., DÜZOVA A., ÖZALTIN F., ORHAN D.
PEDIATRIC NEPHROLOGY, cilt.33, sa.10, ss.1947, 2018 (SCI-Expanded)
74. **CLINICAL AND PATHOLOGIC CHARACTERISTICS OF GENETICALLY CONFIRMED ALPORT SYNDROME PATIENTS**
TOPALOĞLU R., Ozdemir G., GÜLHAN B., Atayar E., DÜZOVA A., SOYLU A., Ozcakar B., SÖYLEMEZOĞLU O., ÖZALTIN F.
PEDIATRIC NEPHROLOGY, cilt.33, sa.10, ss.1826-1827, 2018 (SCI-Expanded)
75. **CLINICAL AND MUTATIONAL SPECTRUM OF CHILDREN WITH AUTOSOMAL RECESSIVE AND AUTOSOMAL DOMINANT POLYCYSTIC KIDNEY DISEASE**
TÜRKOĞLU Ö., GÜLHAN B., YÜKSEL S., Caliskan S., DÜZOVA A., ÇAKAR N., SÖYLEMEZOĞLU O., TOPALOĞLU R., ÖZALTIN F.
PEDIATRIC NEPHROLOGY, cilt.33, sa.10, ss.1814, 2018 (SCI-Expanded)
76. **MYCETOMA OF THE URINARY TRACT IN AN INFANT WITH HORSESHOE KIDNEY: USEFULNESS OF MECHANICAL REMOVAL AND AMPHOTERICIN-B IRRIGATION**
OĞUZ B., Celegen K., ÇİFTÇİ A. Ö., ÜNAL E., ŞENOCAK M. E., ÖZKALE YAVUZ Ö., TAŞ N., GÜLHAN B., ÖZALTIN F., TOPALOĞLU R., et al.
PEDIATRIC NEPHROLOGY, cilt.33, sa.10, ss.1984, 2018 (SCI-Expanded)
77. **TURKISH ATYPICAL HEMOLYTIC UREMIC SYNDROME REGISTRY: ECULIZUMAB TREATMENT IN AHUS PATIENTS**
Baskin E., Canpolat N., Gulleroglu K., Yilmaz A., MELEK E., Yuksel S., GÜLHAN B., KALYONCU M., Parmaksiz G., Ozcakar B., et al.
PEDIATRIC NEPHROLOGY, cilt.33, sa.10, ss.1832, 2018 (SCI-Expanded)
78. **Familial Mediterranean fever patients homozygous for E148Q variant may have milder disease**
TOPALOĞLU R., BATU AKAL E. D., Yildiz C., Korkmaz E., ÖZEN S., BEŞBAŞ N., ÖZALTIN F.
INTERNATIONAL JOURNAL OF RHEUMATIC DISEASES, cilt.21, sa.10, ss.1857-1862, 2018 (SCI-Expanded)
79. **ADRENAL INSUFFICIENCY IN AN INFANT WITH CONGENITAL NEPHROTIC SYNDROME: NEPHROTIC SYNDROME-14**
Saygili S., Canpolat N., Caliskan S., Atayar E., Sever L., ÖZALTIN F.
PEDIATRIC NEPHROLOGY, cilt.33, sa.10, ss.1929, 2018 (SCI-Expanded)
80. **Extra-Renal manifestations of atypical hemolytic uremic syndrome in children**
Fidan K., Goknar N., GÜLHAN B., MELEK E., Yildirim Z. Y., Baskin E., HAYRAN K. M., Gulleroglu K., Ozcakar Z. B., ÖZALTIN F., et al.
PEDIATRIC NEPHROLOGY, cilt.33, sa.8, ss.1395-1403, 2018 (SCI-Expanded)
81. **Effect of IGF-1 and HGF induced bone marrow mesenchymal stem cells on focal segmental glomerulosclerosis in Sprague-Dawley rats: a pilot study**
ŞAHAN Ö., KORKMAZ E., ONBAŞILAR İ., Gucer S., KAYMAZ F. F., ÖZALTIN F., ÖZCAN A.
FEBS OPEN BIO, cilt.8, ss.152, 2018 (SCI-Expanded)
82. **Atypical Hemolytic Uremic Syndrome in Children Aged < 2 Years**
ÇAKAR N., ÖZÇAKAR Z. B., ÖZALTIN F., KOYUN M., ACAR B. Ç., BAHAT ÖZDOĞAN E., GÜLHAN B., Korkmaz E., Yurt A., Yilmaz S., et al.
NEPHRON, cilt.139, sa.3, ss.211-218, 2018 (SCI-Expanded)
83. **Nephropathic Cystinosis Mimicking Bartter Syndrome A Novel Mutation**
Bastug F., Nalcacioglu H., ÖZALTIN F., Korkmaz E., Yel S.

- IRANIAN JOURNAL OF KIDNEY DISEASES, cilt.12, sa.1, ss.61-63, 2018 (SCI-Expanded)
84. **Whole Exome Sequencing of Patients with Steroid-Resistant Nephrotic Syndrome**
Warejko J. K., Tan W., Daga A., Schapiro D., Lawson J. A., Shril S., Lovric S., Ashraf S., Rao J., Hermle T., et al.
CLINICAL JOURNAL OF THE AMERICAN SOCIETY OF NEPHROLOGY, cilt.13, sa.1, ss.53-62, 2018 (SCI-Expanded)
85. **Hemolytic uremic syndrome and IgA nephropathy in a child: Coincidence or not?**
Surmeli-Doven S., Delibas A., Gurses I., Kayacan U. R., Coskun-Yilmaz B., Esen K., Korkmaz E., ÖZALTIN F.
TURKISH JOURNAL OF PEDIATRICS, cilt.60, sa.1, ss.81-85, 2018 (SCI-Expanded)
86. **Mutations in KEOPS-complex genes cause nephrotic syndrome with primary microcephaly**
Braun D. A., Rao J., Mollet G., Schapiro D., Daugeron M., Tan W., Gribouval O., Boyer O., Revy P., Jobst-Schwan T., et al.
NATURE GENETICS, cilt.49, sa.10, ss.1529-1541, 2017 (SCI-Expanded)
87. **Long-Term Outcome of Steroid-Resistant Nephrotic Syndrome in Children**
Trautmann A., Schnaidt S., Lipska-Zietkiewicz B. S., Bodria M., ÖZALTIN F., Emma F., Anarat A., Melk A., Azocar M., Oh J., et al.
JOURNAL OF THE AMERICAN SOCIETY OF NEPHROLOGY, cilt.28, sa.10, ss.3055-3065, 2017 (SCI-Expanded)
88. **IS RAAS BLOCKADE EFFECTIVE IN NPHS2 GLOMERULOPATHY?**
Trautmann A., ÖZALTIN F., Zurowksa A., Helena H., Saeed B., Azocar M., Yilmaz A., Anarat A., Balat A., Caliskan S., et al.
PEDIATRIC NEPHROLOGY, cilt.32, sa.9, ss.1661, 2017 (SCI-Expanded)
89. **CLINICAL CHARACTERISTICS OF PATIENTS WITH MTORI CONVERSION AND EFFECTS ON RENAL OUTCOME**
TAŞ N., GÜLHAN B., Inozu M., Celegen K., DÜZOVA A., ÖZALTIN F., TOPALOĞLU R.
PEDIATRIC NEPHROLOGY, cilt.32, sa.9, ss.1803, 2017 (SCI-Expanded)
90. **DOES C.3979 G > A/P.VAL1327MET VARIANT OF COL4A4 HAS ANY PATHOGENIC EFFECT IN TURKISH PATIENTS WITH ALPORT SYNDROME?**
YILDIZ N., ATA P., ALPAY H., GÖKCE İ., ÖZALTIN F.
PEDIATRIC NEPHROLOGY, cilt.32, sa.9, ss.1773, 2017 (SCI-Expanded)
91. **LONG TERM RESULTS OF RITUXIMAB IN A TERTIARY REFERRAL CENTER FOR DIFFICULT-TO-TREAT NEPHROTIC SYNDROME**
Celegen K., GÜLHAN B., Inozu M., TAŞ N., ÖZALTIN F., DÜZOVA A., TOPALOĞLU R.
PEDIATRIC NEPHROLOGY, cilt.32, sa.9, ss.1769-1770, 2017 (SCI-Expanded)
92. **RISK FACTORS FOR CARDIOVASCULAR COMORBIDITIES IN CHILDREN WITH STEROID-SENSITIVE AND STEROID-RESISTANT NEPHROTIC SYNDROME**
Eroglu F. K., TOPALOĞLU R., KARAGÖZ T., OĞUZ B., ÖZALTIN F., GÜLHAN B., DÜZOVA A.
PEDIATRIC NEPHROLOGY, cilt.32, sa.9, ss.1661, 2017 (SCI-Expanded)
93. **USE OF HERBAL AND DIETARY SUPPLEMENTS IN CHILDREN WITH KIDNEY AND URINARY TRACT DISEASE**
TAŞTEMEL ÖZTÜRK T., Ozmert E., KANBUR N., GÜLHAN B., ÖZALTIN F., TOPALOĞLU R., DÜZOVA A.
PEDIATRIC NEPHROLOGY, cilt.32, sa.9, ss.1675, 2017 (SCI-Expanded)
94. **GENETIC AND CLINICAL CHARACTERISTICS OF PATIENTS WITH C3 GLOMERULOPATHY**
TOPALOĞLU R., GÜLHAN B., Korkmaz E., DÜZOVA A., ÖZALTIN F.
PEDIATRIC NEPHROLOGY, cilt.32, sa.9, ss.1669, 2017 (SCI-Expanded)
95. **Low renal but high extrarenal phenotype variability in Schimke immuno-osseous dysplasia**
Lipska-Zietkiewicz B. S., Gellermann J., Boyer O., Gribouval O., Zietkiewicz S., Kari J. A., Shalaby M. A., ÖZALTIN F., Dusek J., Melk A., et al.
PLOS ONE, cilt.12, sa.8, 2017 (SCI-Expanded)
96. **Follow-up results of patients with ADCK4 mutations and the efficacy of CoQ10 treatment**
Atmaca M., GÜLHAN B., KORKMAZ E., INOZU M., Soylemezoglu O., CANDAN C., Bayazit A. K., Elmaci A. M., Parmaksiz G., DÜZOVA A., et al.
PEDIATRIC NEPHROLOGY, cilt.32, sa.8, ss.1369-1375, 2017 (SCI-Expanded)
97. **Tocilizumab treatment in childhood Takayasu arteritis: Case series of four patients and systematic**

review of the literature

- Batu E. D., Sonmez H. E., HAZIROLAN T., ÖZALTIN F., BİLGİNER Y., ÖZEN S.
SEMINARS IN ARTHRITIS AND RHEUMATISM, cilt.46, sa.4, ss.529-535, 2017 (SCI-Expanded)
98. **Turkish pediatric atypical hemolytic uremic syndrome registry: initial analysis of 146 patients**
BESBAS N., GULHAN B., SOYLEMEZOGLU O., OZCAKAR Z. B., KORKMAZ E., HAYRAN M., Ozaltin F.
BMC NEPHROLOGY, cilt.18, 2017 (SCI-Expanded)
99. **Timing of renal replacement therapy does not influence survival and growth in children with congenital nephrotic syndrome caused by mutations in NPHS1: data from the ESPN/ERA-EDTA Registry**
Holtta T., Bonthuis M., Van Stralen K. J., Bjerre A., TOPALOĞLU R., ÖZALTIN F., Holmberg C., Harambat J., Jager K. J., Schaefer F., et al.
PEDIATRIC NEPHROLOGY, cilt.31, sa.12, ss.2317-2325, 2016 (SCI-Expanded)
100. **First-Line, Early and Long-Term Eculizumab Therapy in Atypical Hemolytic Uremic Syndrome: A Case Series in Pediatric Patients**
YÜKSEL S., Evrengul H., ÖZÇAKAR Z. B., Becerir T., Yalcin N., Korkmaz E., ÖZALTIN F.
PEDIATRIC DRUGS, cilt.18, sa.6, ss.413-420, 2016 (SCI-Expanded)
101. **Genetic and clinical features of cryopyrin-associated periodic syndromes in Turkish children**
EROGLU F. K., Kasapcopur O., BESBAS N., ÖZALTIN F., BİLGİNER Y., Barut K., MENSA-VILARO A., Nakagawa K., Heike T., Nishikomori R., et al.
CLINICAL AND EXPERIMENTAL RHEUMATOLOGY, cilt.34, sa.6, 2016 (SCI-Expanded)
102. **Turkish Atypical Hemolytic Uremic Syndrome Registry: Evaluation of 146 Patients**
Besbas N., Soylemezoglu O., GÜLHAN B., Ozcakar Z. B., KORKMAZ E., HAYRAN K. M., ÖZALTIN F.
PEDIATRIC NEPHROLOGY, cilt.31, sa.10, ss.1884, 2016 (SCI-Expanded)
103. **Loss of diacylglycerol kinase epsilon in mice causes endothelial distress and impairs glomerular Cox-2 and PGE(2) production**
Zhu J., Chaki M., Lu D., Ren C., Wang S., Rauhauser A., Li B., Zimmerman S., Jun B., Du Y., et al.
AMERICAN JOURNAL OF PHYSIOLOGY-RENAL PHYSIOLOGY, cilt.310, sa.9, 2016 (SCI-Expanded)
104. **Mutations in nuclear pore genes NUP93, NUP205 and XPO5 cause steroid-resistant nephrotic syndrome**
Braun D. A., Sadowski C. E., Kohl S., Lovric S., Astrinidis S. A., Pabst W. L., Gee H. Y., Ashraf S., Lawson J. A., Shril S., et al.
NATURE GENETICS, cilt.48, sa.4, ss.457-467, 2016 (SCI-Expanded)
105. **Whole exome sequencing identifies causative mutations in the majority of consanguineous or familial cases with childhood-onset increased renal echogenicity**
Braun D. A., Schueler M., Halbritter J., Gee H. Y., Porath J. D., Lawson J. A., Airik R., Shril S., Allen S. J., Stein D., et al.
KIDNEY INTERNATIONAL, cilt.89, sa.2, ss.468-475, 2016 (SCI-Expanded)
106. **Normal 25-Hydroxyvitamin D Levels Are Associated with Less Proteinuria and Attenuate Renal Failure Progression in Children with CKD**
Shroff R., Aitkenhead H., Costa N., Trivelli A., Litwin M., Picca S., Anarat A., Sallay P., ÖZALTIN F., Zurowska A., et al.
JOURNAL OF THE AMERICAN SOCIETY OF NEPHROLOGY, cilt.27, sa.1, ss.314-322, 2016 (SCI-Expanded)
107. **ADCK4-Associated Glomerulopathy Causes Adolescence-Onset FSGS**
Korkmaz E., Lipska-Zietkiewicz B. S., Boyer O., Gribouval O., Fourrage C., Tabatabaei M., Schnaidt S., Gucer S., Kaymaz F., ARICI M., et al.
JOURNAL OF THE AMERICAN SOCIETY OF NEPHROLOGY, cilt.27, sa.1, ss.63-68, 2016 (SCI-Expanded)
108. **Lupus in a patient with cystinosis: is it drug induced?**
Eroglu F. K., BESBAS N., ÖZALTIN F., TOPALOĞLU R., ÖZEN S.
LUPUS, cilt.24, sa.13, ss.1452-1454, 2015 (SCI-Expanded)
109. **MCP1 2518 A/G polymorphism affects progression of childhood focal segmental glomerulosclerosis**
Besbas N., KALYONCU M., Cil O., ÖZGÜL R. K., Bakkaloglu A., ÖZALTIN F.
RENAL FAILURE, cilt.37, sa.9, ss.1435-1439, 2015 (SCI-Expanded)
110. **NEPHROPATHIC CYSTINOSIS MIMICKING BARTTER SYNDROME: NOVEL MUTATION**

- BaStuG F., Yel S., ÖZALTIN F., Korkmaz E., Uytun S., DÜŞÜNSEL R.
PEDIATRIC NEPHROLOGY, cilt.30, sa.9, ss.1681, 2015 (SCI-Expanded)
111. **RESPONSE TO INTENSIFIED IMMUNOSUPPRESSIVE THERAPY AND IDENTIFICATION OF GENETIC DISEASE ARE HIGHLY PREDICTIVE OF LONG-TERM RENAL OUTCOME IN CHILDREN WITH STEROID RESISTANT NEPHROTIC SYNDROME (SRNS)**
Trautmann A., Schnaitt S., Ghiggeri G. M., ÖZALTIN F., Saeed B., Drozdz D., Caliskan S., Anarat A., Oh J., Bogdanovic R., et al.
PEDIATRIC NEPHROLOGY, cilt.30, sa.9, ss.1556-1557, 2015 (SCI-Expanded)
112. **ATYPICAL HEMOLYTIC UREMIC SYNDROME UNRESPONSIVE TO ECULIZUMAB THERAPY**
Inozu M., BODUR İ., DÜZOVA A., BİLGİNER Y., ÖZALTIN F., BEŞBAŞ N.
PEDIATRIC NEPHROLOGY, cilt.30, sa.9, ss.1648, 2015 (SCI-Expanded)
113. **RESULTS OF TURKISH MULTICENTRIC NATIONAL CYSTINOSIS REGISTRY**
TOPALOĞLU R., GÜLHAN B., ÖZALTIN F., BODUR İ., BEŞBAŞ N., Dursun H., Yilmaz A., GÜRGÖZE M. K., GÖKCE İ., Akinci N., et al.
PEDIATRIC NEPHROLOGY, cilt.30, sa.9, ss.1668-1669, 2015 (SCI-Expanded)
114. **RISK FACTORS FOR POST-TRANSPLANT RECURRENCE OF STEROID RESISTANT NEPHROTIC SYNDROME (SRNS): RESULTS FROM THE PODONET REGISTRY**
Trautmann A., Ghiggeri G. M., Azocar M., Remuzzi G., ÖZALTIN F., Melk A., Jankauskiene A., Peco-antic A., Gellermann J., Emma F., et al.
PEDIATRIC NEPHROLOGY, cilt.30, sa.9, ss.1557-1558, 2015 (SCI-Expanded)
115. **CHARACTERISTICS OF RENAL TRANSPLANT CHILDREN WITH CHRONIC ALLOGRAFT NEPHROPATHY: EXPERIENCE OF A TERTIARY REFERRAL CENTER**
GÜLHAN B., INOZU M., BİLGİNER Y., ÖZALTIN F., DÜZOVA A., ÖZEN S., TOPALOĞLU R., AKI F. T., BeSbaS N.
PEDIATRIC NEPHROLOGY, cilt.30, sa.9, ss.1721, 2015 (SCI-Expanded)
116. **SEVERE PNEUMONIA ASSOCIATED WITH BK VIRUS NEPHROPATHY IN A KIDNEY TRANSPLANTED ADOLESCENT WITH SCHIMKE IMMUNO-OSSEOUS-DYSPLASIA**
DÜZOVA A., GÜLHAN B., ÖZALTIN F., TOPALOĞLU R., BİLGİNER Y., BEŞBAŞ N.
PEDIATRIC NEPHROLOGY, cilt.30, sa.9, ss.1619-1620, 2015 (SCI-Expanded)
117. **RITUXIMAB EXPERIENCE OF A TERTIARY REFERRAL CENTER FOR DIFFICULT-TO-TREAT NEPHROTIC SYNDROME**
GÜLHAN B., TOPALOĞLU R., ÖZALTIN F., DÜZOVA A., EROGLU F. K., Bodur I., BİLGİNER Y., ÖZEN S., BesbaS N.
PEDIATRIC NEPHROLOGY, cilt.30, sa.9, ss.1692, 2015 (SCI-Expanded)
118. **Genetic abnormalities and prognosis in patients with congenital and infantile nephrotic syndrome**
Cil O., Besbas N., DÜZOVA A., TOPALOĞLU R., PEKO-ANTIC A., KORKMAZ E., ÖZALTIN F.
PEDIATRIC NEPHROLOGY, cilt.30, sa.8, ss.1279-1287, 2015 (SCI-Expanded)
119. **Thrombophilic Risk Factors And The Efficiency Of Prophylactic Anticoagulation Therapy In Children Who Underwent Renal Transplantation**
GÜLHAN B., Tavil B., DÜZOVA A., ÖZALTIN F., ÖZEN S., TOPALOĞLU R., BİLGİNER Y., GÜMRÜK F., BEŞBAŞ N.
PEDIATRIC NEPHROLOGY, cilt.29, sa.9, ss.1694, 2014 (SCI-Expanded)
120. **Mutations In Anks6 Cause A Nephronophthisis-like Phenotype With End Stage Renal Disease**
Taskiran E., Korkmaz E., Gucer S., KOŞUKCU C., Kaymaz F., Koyunlar C., Bryda E. C., Chaki M., Lu D. D., Vadnagara K., et al.
PEDIATRIC NEPHROLOGY, cilt.29, sa.9, ss.1820, 2014 (SCI-Expanded)
121. **A novel CFHR5 mutation associated with C3 glomerulonephritis in a Turkish girl**
BEŞBAŞ N., GÜLHAN B., GÜÇER K. Ş., Korkmaz E., ÖZALTIN F.
JOURNAL OF NEPHROLOGY, cilt.27, sa.4, ss.457-460, 2014 (SCI-Expanded)
122. **Diagnostic validity of colchicine in patients with Familial Mediterranean fever**
ÖZALTIN F., BİLGİNER Y., GÜLHAN B., Bajin I., Erdogan O., HAYRAN K. M., Yilmaz E., ÖZEN S.
CLINICAL RHEUMATOLOGY, cilt.33, sa.7, ss.969-974, 2014 (SCI-Expanded)
123. **Post-transplant hypertension in pediatric kidney transplant recipients**
GÜLHAN B., TOPALOĞLU R., KARABULUT E., ÖZALTIN F., AKI F. T., BİLGİNER Y., Besbas N.

- PEDIATRIC NEPHROLOGY, cilt.29, sa.6, ss.1075-1080, 2014 (SCI-Expanded)
124. Primary coenzyme Q(10) (CoQ(10)) deficiencies and related nephropathies
ÖZALTIN F.
PEDIATRIC NEPHROLOGY, cilt.29, sa.6, ss.961-969, 2014 (SCI-Expanded)
125. Mutations in EMP2 Cause Childhood-Onset Nephrotic Syndrome
Gee H. Y., Ashraf S., Wan X., Vega-Warner V., Esteve-Rudd J., Lovric S., Fang H., Hurd T. W., Sadowski C. E., Allen S. J., et al.
AMERICAN JOURNAL OF HUMAN GENETICS, cilt.94, sa.6, ss.884-890, 2014 (SCI-Expanded)
126. Genotype-phenotype associations in WT1 glomerulopathy
Lipska-Zietkiewicz B. S., Ranchin B., Iatropoulos P., Gellermann J., Melk A., ÖZALTIN F., Caridi G., Seeman T., Tory K., Jankauskiene A., et al.
KIDNEY INTERNATIONAL, cilt.85, sa.5, ss.1169-1178, 2014 (SCI-Expanded)
127. NPHS2 Mutations in Steroid-Resistant Nephrotic Syndrome: A Mutation Update and the Associated Phenotypic Spectrum
Bouchireb K., Boyer O., Gribouval O., Nevo F., Huynh-Cong E., Moriniere V., Campait R., Ars E., Brackman D., Dantal J., et al.
HUMAN MUTATION, cilt.35, sa.2, ss.178-186, 2014 (SCI-Expanded)
128. Proteinuria in Frasier Syndrome
Peco-Antic A., ÖZALTIN F., Parezanovic V., Milosevski-Lomic G., Zdravkovic V.
SRPSKI ARHIV ZA CELOKUPNO LEKARSTVO, cilt.141, ss.685-688, 2013 (SCI-Expanded)
129. CONTRIBUTORY RISK FACTORS FOR DEVELOPMENT OF THROMBOSIS IN CHILDREN WITH NEPHROTIC SYNDROME
KaraEroglu F., Tavil B., ÖZALTIN F., BEŞBAŞ N., ÖZEN S., ÇETİN M., GÜMRÜK F., TOPALOĞLU R.
PEDIATRIC NEPHROLOGY, cilt.28, sa.8, ss.1584, 2013 (SCI-Expanded)
130. The Effect Of Fibroblast Growth Factor 23 On Left Ventricular Function In Peritoneal Dialysis Children
Hacihamdioglu D. O., BEŞBAŞ N., ALEHAN D., OĞUZ B., ÖZALTIN F., DÜZOVA A., ÖZEN S., TOPALOĞLU R., Gok F.
PEDIATRIC NEPHROLOGY, cilt.28, sa.8, ss.1643, 2013 (SCI-Expanded)
131. Genetic screening in adolescents with steroid-resistant nephrotic syndrome
Lipska-Zietkiewicz B. S., Iatropoulos P., Maranta R., Caridi G., ÖZALTIN F., Anarat A., Balat A., Gellermann J., Trautmann A., Erdogan O., et al.
KIDNEY INTERNATIONAL, cilt.84, sa.1, ss.206-213, 2013 (SCI-Expanded)
132. DGKE Variants Cause a Glomerular Microangiopathy That Mimics Membranoproliferative GN
ÖZALTIN F., Li B., Rauhauser A., An S., Soylemezoglu O., Gonul I. I., Taskiran E. Z., Ibsirlioglu T., Korkmaz E., BİLGİNER Y., et al.
JOURNAL OF THE AMERICAN SOCIETY OF NEPHROLOGY, cilt.24, sa.3, ss.377-384, 2013 (SCI-Expanded)
133. Neonatal onset atypical hemolytic uremic syndrome successfully treated with eculizumab
Besbas N., Gulhan B., Karpman D., Topaloglu R., Duzova A., Korkmaz E., ÖZALTIN F.
PEDIATRIC NEPHROLOGY, cilt.28, sa.1, ss.155-158, 2013 (SCI-Expanded)
134. Circulating suPAR in Two Cohorts of Primary FSGS
Wei C., Trachtman H., Li J., Dong C., Friedman A. L., Gassman J. J., McMahan J. L., Radeva M., Heil K. M., Trautmann A., et al.
JOURNAL OF THE AMERICAN SOCIETY OF NEPHROLOGY, cilt.23, sa.12, ss.2051-2059, 2012 (SCI-Expanded)
135. ATYPICAL HUS AND ECOLIZUMAB TREATMENT: EXPERIENCE OF A TERTIARY CENTER
BEŞBAŞ N., GÜLHAN B., ÖZEN S., TOPALOĞLU R., DÜZOVA A., Yıldız C., ÖZALTIN F.
PEDIATRIC NEPHROLOGY, cilt.27, sa.9, ss.1630, 2012 (SCI-Expanded)
136. Response to Intensified Immunosuppressive Therapy Predicts Long-Term Prognosis in Steroid Resistant Nephrotic Syndrome (SRNS)
Trautmann A., ÖZALTIN F., Bodria M., Anarat A., Saeed B., Azocar M., Gellermann J., Drozdz D., Bogdanovic R., Krmar R., et al.
PEDIATRIC NEPHROLOGY, cilt.27, sa.9, ss.1639-1640, 2012 (SCI-Expanded)

137. **WT1 screening in nephrotic syndrome - lessons from PodoNet**
Lipska-Zietkiewicz B. S., Iatropoulos P., Ranchin B., Dusek J., ÖZALTIN F., Melk A., Gellermann J., Jankauskiene A., Peco-Antic A., Zurowska A., et al.
PEDIATRIC NEPHROLOGY, cilt.27, sa.9, ss.1617-1618, 2012 (SCI-Expanded)
138. **Takayasu arteritis in a 4-year-old girl: case report and brief overview of the pediatric literature**
Aypar E., Celebi-Tayfur A., Keser M., Odabas D., ÖZALTIN F., PAKSOY Y., ÖZEN S.
TURKISH JOURNAL OF PEDIATRICS, cilt.54, sa.5, ss.536-539, 2012 (SCI-Expanded)
139. **THE BENEFITS OF CYCLOSPORINE TREATMENT OF THE PATIENT WITH NPHS2 MUTATION**
Hacihamdioglu D. O., ÖZALTIN F., Zeybek C., Kalman S., Demirkaya E., Gok F.
PEDIATRIC NEPHROLOGY, cilt.27, sa.9, ss.1703-1704, 2012 (SCI-Expanded)
140. **Rituximab in steroid-dependent and resistant nephrotic syndrome patients**
TOPALOĞLU R., GÜLHAN B., ÖZALTIN F., DÜZOVA A., BEŞBAŞ N.
PEDIATRIC NEPHROLOGY, cilt.27, sa.9, ss.1710-1711, 2012 (SCI-Expanded)
141. **PERITONEAL DIALYSIS IN CHILDREN UNDER TWO YEARS OF AGE**
GÜLHAN B., DÜZOVA A., ÖZALTIN F., TOPALOĞLU R., ÖZEN S., BİLGİNER Y., Tayfur A. C., Yıldız C., BEŞBAŞ N.
PEDIATRIC NEPHROLOGY, cilt.27, sa.9, ss.1799-1800, 2012 (SCI-Expanded)
142. **C1q deficiency: identification of a novel missense mutation and treatment with fresh frozen plasma**
TOPALOĞLU R., TASKIRAN E. Z., TAN Ç., Erman B., ÖZALTIN F., SANAL O.
CLINICAL RHEUMATOLOGY, cilt.31, sa.7, ss.1123-1126, 2012 (SCI-Expanded)
143. **NPHS2 gene mutation in an Iranian family with familial steroid-resistant nephrotic syndrome**
Ameli S., Mazaheri M., Zare-Shahabadi A., ÖZALTIN F., Asgarian F., Monajemzadeh M., Bazargani B., Ataei N., Hajezadeh N., Madani A., et al.
NEFROLOGIA, cilt.32, sa.5, ss.674-676, 2012 (SCI-Expanded)
144. **Genetic basis of cystinosis in Turkish patients: a single-center experience**
TOPALOĞLU R., Vilboux T., COŞKUN T., ÖZALTIN F., Tinloy B., Gunay-Aygun M., BAKKALOĞLU A., BEŞBAŞ N., van den Heuvel L., Kleta R., et al.
PEDIATRIC NEPHROLOGY, cilt.27, sa.1, ss.115-121, 2012 (SCI-Expanded)
145. **OUTCOME OF NEPHROPATHIC CYSTINOSIS IN PEDIATRIC RENAL TRANSPLANTATION: A SINGLE CENTRE EXPERIENCE**
TOPALOĞLU R., BİLGİNER Y., ÖZALTIN F., DÜZOVA A., Aki T., ÖZEN S., BAKKALOĞLU A., BEŞBAŞ N.
PEDIATRIC TRANSPLANTATION, cilt.15, ss.82, 2011 (SCI-Expanded)
146. **Disruption of PTPRO Causes Childhood-Onset Nephrotic Syndrome**
ÖZALTIN F., Ibsirlioglu T., Taskiran E. Z., Baydar D. E., Kaymaz F., Buyukcelik M., Kilic B. D., Balat A., Iatropoulos P., Asan E., et al.
AMERICAN JOURNAL OF HUMAN GENETICS, cilt.89, sa.1, ss.139-147, 2011 (SCI-Expanded)
147. **Respiratory-chain deficiency presenting as diffuse mesangial sclerosis with NPHS3 mutation**
Baskin E., Bayrakci U. S., Alehan F., Ozdemir H., Oner A., Horvath R., Vega-Warner V., Hildebrandt F., ÖZALTIN F.
PEDIATRIC NEPHROLOGY, cilt.26, sa.7, ss.1157-1161, 2011 (SCI-Expanded)
148. **MYO1E Mutations and Childhood Familial Focal Segmental Glomerulosclerosis**
Mele C., Iatropoulos P., Donadelli R., Calabria A., Maranta R., Cassis P., Buelli S., Tomasoni S., Piras R., Krendel M., et al.
NEW ENGLAND JOURNAL OF MEDICINE, cilt.365, sa.4, ss.295-306, 2011 (SCI-Expanded)
149. **COQ6 mutations in human patients produce nephrotic syndrome with sensorineural deafness**
Heeringa S. F., Chernin G., Chaki M., Zhou W., Sloan A. J., Ji Z., Xie L. X., Salviati L., Hurd T. W., Vega-Warner V., et al.
JOURNAL OF CLINICAL INVESTIGATION, cilt.121, sa.5, ss.2013-2024, 2011 (SCI-Expanded)
150. **Follow-Up of Patients With Juvenile Nephronophthisis After Renal Transplantation: A Single Center Experience**
Tayfur A. C., Besbas N., BİLGİNER Y., ÖZALTIN F., DÜZOVA A., Bakkaloglu M., AKI F. T., ÖZEN S., TOPALOĞLU R., Bakkaloglu A.
TRANSPLANTATION PROCEEDINGS, cilt.43, sa.3, ss.847-849, 2011 (SCI-Expanded)
151. **Three cases of a rare disease, congenital chloride diarrhea, summons up the variation in the clinical**

course and significance of early diagnosis and adequate treatment in the prevention of intellectual disability

Gurakan F., Baysoy G., Wedenoja S., Uslu N., ÖZEN H., ÖZALTIN F., Hoglund P.

TURKISH JOURNAL OF PEDIATRICS, cilt.53, sa.2, ss.194-198, 2011 (SCI-Expanded)

152. **Spondyloenchondrodyplasia with Systemic Lupus Erythematosus: a report of three cases**

BİLGİNER Y., ALANAY Y., DÜZOVA A., TOPALOĞLU R., Superti-Furga A., ÖZEN S., ÖZALTIN F., BEŞBAŞ N.

CLINICAL AND EXPERIMENTAL RHEUMATOLOGY, cilt.29, sa.2, ss.430, 2011 (SCI-Expanded)

153. **Macrophage Activating Syndrome complicating Wegener Granulomatosis: treatment with plasma exchange**

Eroglu F. K., GÜLHAN B., ÜNAL S., ÖZALTIN F., ORHAN D., ÖZEN S.

CLINICAL AND EXPERIMENTAL RHEUMATOLOGY, cilt.29, sa.2, ss.463, 2011 (SCI-Expanded)

154. **The bone and mineral disorder of children undergoing chronic peritoneal dialysis**

Borzych D., Rees L., Ha I. S., Chua A., Valles P. G., Lipka M., Zambrano P., Ahlenstiel T., BAKKALOĞLU EZGÜ S. A., Spizzirri A. P., et al.

KIDNEY INTERNATIONAL, cilt.78, sa.12, ss.1295-1304, 2010 (SCI-Expanded)

155. **Neuroendocrine immune system in familial Mediterranean fever**

TOPALOĞLU R., BİLGİNER Y., ALİKAŞIFOĞLU A., ÖZALTIN F., BEŞBAŞ N., ÖZEN S., BAKKALOĞLU A.

TURKISH JOURNAL OF PEDIATRICS, cilt.52, sa.6, ss.588-593, 2010 (SCI-Expanded)

156. **Genotype/Phenotype Correlation in Nephrotic Syndrome Caused by WT1 Mutations**

Chernin G., Vega-Warner V., Schoeb D. S., Heeringa S. F., Ovunc B., Saisawat P., Cleper R., ÖZALTIN F., Hildebrandt F. CLINICAL JOURNAL OF THE AMERICAN SOCIETY OF NEPHROLOGY, cilt.5, sa.9, ss.1655-1662, 2010 (SCI-Expanded)

157. **New syndrome - Situs inversus totalis with cystic dysplasia of kidneys, pancreas and bowing**

Balci S., ÖZALTIN F., Bostanoglu S.

CLINICAL DYSMORPHOLOGY, cilt.19, sa.3, ss.173-174, 2010 (SCI-Expanded)

158. **Clinical course of primary focal segmental glomerulosclerosis (FSGS) in Turkish children: a report from the Turkish Pediatric Nephrology FSGS Study Group**

BEŞBAŞ N., ÖZALTIN F., Emre S., Anarat A., ALPAY H., BAKKALOĞLU A., Baskin E., BUYAN N., DÖNMEZ O., DÜŞÜNSEL R., et al.

TURKISH JOURNAL OF PEDIATRICS, cilt.52, sa.3, ss.255-261, 2010 (SCI-Expanded)

159. **Risk factors in community-acquired urinary tract infections caused by ESBL-producing bacteria in children**

TOPALOĞLU R., Er I., Dogan B. G., BİLGİNER Y., ÖZALTIN F., Besbas N., ÖZEN S., Bakkaloglu A., GÜR D.

PEDIATRIC NEPHROLOGY, cilt.25, sa.5, ss.919-925, 2010 (SCI-Expanded)

160. **A Case Report of Thrombocytopenia-associated Multiple Organ Failure Secondary to Salmonella enterica Serotype Typhi Infection in a Pediatric Patient: Successful Treatment With Plasma Exchange**

Yıldırım I., CEYHAN M., BAYRAKİ B., Uysal M., Kuskonmaz B. B., ÖZALTIN F.

THERAPEUTIC APHERESIS AND DIALYSIS, cilt.14, sa.2, ss.226-229, 2010 (SCI-Expanded)

161. **Toll-like receptors 2 and 4 cell surface expression reflects endotoxin tolerance in Henoch-Schönlein purpura.**

CANPINAR H., ÖZALTIN F., BILGINER Y., BAKKALOĞLU A., Ozen S.

The Turkish journal of pediatrics, cilt.52, ss.22-7, 2010 (SCI-Expanded)

162. **A novel CLCN7 mutation resulting in a most severe form of autosomal recessive osteopetrosis**

BEŞBAŞ N., Draaken M., Ludwig M., DEREN Ö., ORHAN D., BİLGİNER Y., ÖZALTIN F.

EUROPEAN JOURNAL OF PEDIATRICS, cilt.168, sa.12, ss.1449-1454, 2009 (SCI-Expanded)

163. **Treatment of severe Henoch-Schönlein nephritis: justifying more immunosuppression**

Altugan F. S., ÖZEN S., Aktay-Ayaz N., Guçer S., TOPALOĞLU R., DÜZOVA A., ÖZALTIN F., Besbas N.

TURKISH JOURNAL OF PEDIATRICS, cilt.51, sa.6, ss.551-555, 2009 (SCI-Expanded)

164. **Thirteen novel NPHS1 mutations in a large cohort of children with congenital nephrotic syndrome**

Heeringa S. F., Vlangos C. N., Chernin G., Hinkes B., Gbadegesin R., Liu J., Hoskins B. E., ÖZALTIN F., Hildebrandt F.

NEPHROLOGY DIALYSIS TRANSPLANTATION, cilt.23, sa.11, ss.3527-3533, 2008 (SCI-Expanded)

165. **Evaluation of intima media thickness of the common and internal carotid arteries with inflammatory**

- markers in familial Mediterranean fever as possible predictors for atherosclerosis**
BİLGİNER Y., ÖZALTIN F., Basaran C., DÜZOVA A., Besbas N., TOPALOĞLU R., ÖZEN S., Bakkaloglu A.
RHEUMATOLOGY INTERNATIONAL, cilt.28, sa.12, ss.1211-1216, 2008 (SCI-Expanded)
166. **Mutations in PLCE1 are a major cause of isolated diffuse mesangial sclerosis (IDMS)**
Gbadegenin R., Hinkes B. G., Hoskins B. E., Vlangos C. N., Heeringa S. F., Liu J., Loirat C., ÖZALTIN F., Hashmi S., Ulmer F., et al.
NEPHROLOGY DIALYSIS TRANSPLANTATION, cilt.23, sa.4, ss.1291-1297, 2008 (SCI-Expanded)
167. **Eye involvement in children with primary focal segmental glomerulosclerosis**
ÖZALTIN F., Heeringa S., Poyraz C. E., BİLGİNER Y., KADAYIFÇILAR S., BEŞBAŞ N., TOPALOĞLU R., ÖZEN S., Hildebrandt F., BAKKALOĞLU A.
PEDIATRIC NEPHROLOGY, cilt.23, sa.3, ss.421-427, 2008 (SCI-Expanded)
168. **Right atrial thrombosis complicating renal transplantation in a child**
BİLGİNER Y., ÖZALTIN F., DÜZOVA A., Erdogan I., AKI F. T., DEMİRCİN M., Bakkaloglu M., Bakkaloglu A.
PEDIATRIC TRANSPLANTATION, cilt.12, sa.2, ss.251-255, 2008 (SCI-Expanded)
169. **Specific podocin mutations correlate with age of onset in steroid-resistant nephrotic syndrome**
Hinkes B., Vlangos C., Heeringa S., Mucha B., Gbadegenin R., Liu J., Hasselbacher K., ÖZALTIN F., Hildebrandt F.
JOURNAL OF THE AMERICAN SOCIETY OF NEPHROLOGY, cilt.19, sa.2, ss.365-371, 2008 (SCI-Expanded)
170. **Triple immunosuppression with tacrolimus in pediatric renal transplantation: Single-center experience**
DÜZOVA A., AKI F. T., Bakkaloglu A., Besbas N., TOPALOĞLU R., ÖZEN S., ÖZALTIN F., BİLGİNER Y., Demirkaya E., Bakkaloglu M.
TRANSPLANTATION PROCEEDINGS, cilt.40, sa.1, ss.132-134, 2008 (SCI-Expanded)
171. **Cerebral sinovenous thrombosis in a child with steroid sensitive nephrotic syndrome**
Isik Balci Y., Tavil B., Fidan G., Ozaltin F.
EUROPEAN JOURNAL OF PEDIATRICS, cilt.166, sa.7, ss.757-758, 2007 (SCI-Expanded)
172. **Nephrotic syndrome in the first year of life: Two thirds of cases are caused by mutations in 4 genes (NPHS1, NPHS2, WT1, and LAMB2)**
Hinkes B. G., Mucha B., Vlangos C. N., Gbadegenin R., Liu J., Hasselbacher K., Hangan D., Ozaltin F., Zenker M., Hildebrandt F.
PEDIATRICS, cilt.119, sa.4, 2007 (SCI-Expanded)
173. **Childhood vasculitides in Turkey: a nationwide survey**
Ozen S., Bakkaloglu A., Dusunsel R., Soylemezoglu O., Ozaltin F., Poyrazoglu H., KASAPÇOPUR Ö., Ozkaya O., Yalcinkaya F., Balat A., et al.
CLINICAL RHEUMATOLOGY, cilt.26, sa.2, ss.196-200, 2007 (SCI-Expanded)
174. **Positional cloning uncovers mutations in PLCE1 responsible for a nephrotic syndrome variant that may be reversible**
Hinkes B., Wiggins R. C., Gbadegenin R., Vlangos C. N., Seelow D., Nuernberg G., Garg P., Verma R., Chaib H., Hoskins B. E., et al.
NATURE GENETICS, cilt.38, sa.12, ss.1397-1405, 2006 (SCI-Expanded)
175. **Novel OCRL1 mutations in patients with the phenotype of dent disease**
Utsch B., Boekenkamp A., Benz M. R., Besbas N., Doetsch J., Franke I., Fruend S., Gok F., Hoppe B., Karle S., et al.
AMERICAN JOURNAL OF KIDNEY DISEASES, cilt.48, sa.6, ss.942-954, 2006 (SCI-Expanded)
176. **Serum IgD concentrations in patients with ataxia telangiectasia and with selective IgA deficiency**
SANAL O., Ozaltin F., TEZCAN İ., ERSOY F.
INTERNATIONAL ARCHIVES OF ALLERGY AND IMMUNOLOGY, cilt.116, sa.3, ss.246, 1998 (SCI-Expanded)

Düger Dergilerde Yayınlanan Makaleler

1. **BK Virus Infections in Pediatric Kidney Transplant Recipients: A Single-Center Experience**
KURT ŞÜKÜR E. D., GÜLHAN B., Özdemir G., Öztürk T. T., BALTU D., ÖZALTIN F., DÜZOVA A., TOPALOĞLU R.

- Turkish Journal of Nephrology, cilt.32, sa.4, ss.353-360, 2023 (ESCI)
- 2. **Nefrotik Düzey Proteinürü ile Başvurunun Çocukluk Çağı Iga Vaskülit Nefriti Seyrine Etkileri**
Kurt Şükür E. D., METE O., Yılmaz T., Gülhan B., Orhan D., Ozaltin F., DÜZOVA A., TOPALOĞLU R.
Osmangazi Tıp Dergisi, cilt.45, sa.4, ss.581-589, 2023 (Hakemli Dergi)
 - 3. **Albuminuria is associated with 24-hour and night-time diastolic blood pressure in urinary tract infection with renal scarring**
BALTU D., Salancı B. V., GÜLHAN B., ÖZALTIN F., DÜZOVA A., TOPALOĞLU R.
The Turkish journal of pediatrics, cilt.65, sa.4, ss.620-629, 2023 (Scopus)
 - 4. **Hemolytic Uremic Syndrome in Children**
GÜLHAN B., ÖZALTIN F.
TURKISH ARCHIVES OF PEDIATRICS, cilt.56, sa.5, ss.415-422, 2021 (ESCI)
 - 5. **Clinicopathological and immunohistological features in childhood IgA nephropathy: a single-centre experience**
TOPALOĞLU R., Orhan D., BİLGİNER Y., KARABULUT E., ÖZALTIN F., DÜZOVA A., KALE G., BEŞBAŞ N.
CLINICAL KIDNEY JOURNAL, cilt.6, sa.2, ss.169-175, 2013 (ESCI)

Kitap & Kitap Bölümleri

- 1. **Primer Podositopatiler**
KURT ŞÜKÜR E. D., ÖZALTIN F.
Genetik böbrek hastalıkları, , Editör, Güneş Tıp Kitabevleri, İstanbul, ss.59-71, 2022
- 2. **Çocukluk Çağında Mitokondriyal Hastalıklar**
KURT ŞÜKÜR E. D., ÖZALTIN F.
Çocukluk Çağında Mitokondriyal Hastalıklar, Ünal Uzun Özlem, Editör, Türkiye Klinikleri, Ankara, ss.46-52, 2022
- 3. **Nefrotik Sendrom ve Genetik**
GÜLHAN B., ÖZALTIN F.
Çocuklarda Nefrotik Sendrom, Prof. Dr. Lale Sever, Editör, Türkiye Klinikleri (Ortadoğu Reklam Tanıtım Yayıncılık Turizm Eğitim İnşaat Sanayi ve Ticaret A.Ş.), ss.28-38, 2021
- 4. **The Kidney in Mitochondrial Diseases**
ÖZALTIN F., Salviati L., Rahman S.
Pediatric Nephrology, Emma Francesco, Goldstein Stuart, Bagga Arvind, Bates Carlton M., Shroff Rukshana, Editör, Springer, Heidelberg, ss.1-13, 2021
- 5. **Nefrotik Sendrom ve Genetik**
GÜLHAN B., ÖZALTIN F.
Çocuklarda Nefrotik Sendrom - 2021, Lale Sever, Editör, Türkiye Klinikleri (Ortadoğu Reklam Tanıtım Yayıncılık Turizm Eğitim İnşaat Sanayi ve Ticaret A.Ş.), Ankara, ss.28-38, 2021
- 6. **İmmün Kompleks Aracılı Glomerülonefritler: Membranoproliferatif Glomerülonefrit, Yeni Sınıflandırma, C3 Glomerülopati**
TAŞTEMEL ÖZTÜRK T., ÖZALTIN F.
Türkiye Klinikleri, Rezan Topaloğlu, Editör, Türkiye Klinikleri (Ortadoğu Reklam Tanıtım Yayıncılık Turizm Eğitim İnşaat Sanayi ve Ticaret A.Ş.), ss.29-37, 2021

Hakemli Kongre / Sempozyum Bildiri Kitaplarında Yer Alan Yayınlar

- 1. **CLINICAL CHARACTERISTICS AND OUTCOME OF BK VIRUS INFECTION IN PEDIATRIC RENAL TRANSPLANT RECIPIENTS**
KURT ŞÜKÜR E. D., ÖZDEMİR E. G., TAŞTEMEL ÖZTÜRK T., BALTU D., GÜLHAN B., ÖZALTIN F., DÜZOVA A., TOPALOĞLU R.
53rd European society of pediatric nephrology congress 2021, Amsterdam, Hollanda, 16 Eylül 2021 - 16 Eylül

2022

2. ANALYSIS OF LATE RENAL COMPLICATIONS AND RISK FACTORS IN CHILDREN WITH HEMATOPOIETIC STEM CELL TRANSPLANTATION
Gurbanov A., GÜLHAN B., KUŞKONMAZ B. B., OKUR F. V., Cetinkaya D. U., Ozdemir G., Tas N., Celegen K., ÖZALTIN F., DÜZOVA A., et al.
57th ERA-EDTA Congress, ELECTR NETWORK, 6 - 09 Haziran 2020, cilt.35, ss.2140
3. EARLY URINARY SYSTEM COMPLICATIONS IN CHILDREN WITH HEMATOPOIETIC STEM CELL TRANSPLANTATION
Gurbanov A., GÜLHAN B., KUŞKONMAZ B. B., OKUR F. V., Cetinkaya D. U., Ozdemir G., Tas N., Celegen K., ÖZALTIN F., DÜZOVA A., et al.
57th ERA-EDTA Congress, ELECTR NETWORK, 6 - 09 Haziran 2020, cilt.35, ss.2123
4. CLINICAL CHARACTERISTICS OF PATIENTS WITH GENETICALLY CONFIRMED ALPORT SYNDROME
Ozdemir G., GÜLHAN B., Atayar E., Canpolat N., SÖYLEMEZOĞLU O., ÖZÇAKAR Z. B., Eroglu F. K., Candan C., Demir B. K., SOYLU A., et al.
57th ERA-EDTA Congress, ELECTR NETWORK, 6 - 09 Haziran 2020, cilt.35, ss.359
5. Hematopoietik Kök Hücre Nakli Yapılan Çocuk Hastalarda Geç Dönem Renal Komplikasyonların Ve RiskFaktörlerinin Analizi
Gurbanov A., GÜLHAN B., KUŞKONMAZ B. B., OKUR F. V., Uçkan Çetinkaya D., Özdemir G., TAŞ N., Celegen K., ÖZALTIN F., DÜZOVA A., et al.
10. Çocuk Nefrolojisi Kongresi, Türkiye, 1 - 04 Mayıs 2019
6. Hematopoietik Kök Hücre Nakli Yapılan Çocuk Hastalarda Erken Dönem Üriner Sistem Komplikasyonları
Gurbanov A., GÜLHAN B., KUŞKONMAZ B. B., OKUR F. V., Uçkan Çetinkaya D., Özdemir G., Taş N., Celegen K., ÖZALTIN F., DÜZOVA A., et al.
10. Çocuk Nefrolojisi Kongresi, Türkiye, 1 - 04 Mayıs 2019
7. Çocukluk Çağı Preemptif Ve Preemptif Olmayan Böbrek Nakli Hastalarında Nakil Sonrası İlk Bir Yıldaki Greft Fonksiyonlarının Değerlendirilmesi
TAŞ N., AKI F. T., GÜLHAN B., ÖZALTIN F., DÜZOVA A., TEKGÜL S., TOPALOĞLU R.
10. Çocuk Nefrolojisi Kongresi, Türkiye, 1 - 04 Mayıs 2019
8. A REFRACTORY CLINICAL CASE OF LUPUS NEPHRITIS WHO UNDERWENT BONE MARROW TRANSPLANTATION,
TAŞ N., DEMİR S., ÇAĞDAŞ AYVAZ D. N., ÖZALTIN F., KUŞKONMAZ B. B., TEZCAN F. İ., ÖZEN S., ORHAN D., TOPALOĞLU R.
European Society for Paediatric Nephrology, 51th Annual Meeting, 3 - 06 Ekim 2018
9. CLINICAL AND PATHOLOGIC CHARACTERISTICS OF GENETICALLY CONFIRMED ALPORT SYNDROME PATIENTS
TOPALOĞLU R., ÖZDEMİR E. G., GÜLHAN B., ATAYAR E., DÜZOVA A., SOYLU A., ÖZÇAKAR Z. B., SÖYLEMEZOĞLU H. O., ÖZALTIN F.
51st Annual Scientific Meeting of the European Society for the Paediatric Nephrology, 3 - 06 Ekim 2018
10. TURKISH ATYPICAL HEMOLYTIC UREMIC SYNDROME REGISTRY: ECULIZUMAB TREATMENT IN AHUS PATIENTS
BASKIN S. E., CANPOLAT N., GÜLLEROĞLU K. S., YILMAZ A., MELEK E., YÜKSEL S., GÜLHAN B., KALYONCU M., parmaksız g., ÖZÇAKAR Z. B., et al.
51st Annual Scientific Meeting of the European Society for the Paediatric Nephrology, 3 - 06 Ekim 2018
11. MYCETOMA OF THE URINARY TRACT IN AN INFANTWITH HORSESHOE KIDNEY: USEFULNESS OF MECHANICAL REMOVAL AND AMPHOTERICIN-BIRRIGATION
OĞUZ B., CELEGEN K., ÇİFTÇİ A. Ö., ÜNAL E., ŞENOCAK M. E., ÖZKALE YAVUZ Ö., TAŞ N., GÜLHAN B., ÖZALTIN F., TOPALOĞLU R., et al.
51st Annual Scientific Meeting of the European Society for the Paediatric Nephrology, 3 - 06 Ekim 2018
12. Lupus nephritis: Long term follow-up and effect of treatment on growth
TOPALOĞLU R., TAŞ N., celegen k., ÖZDEMİR E. G., GÜLHAN B., DÜZOVA A., ÖZALTIN F., ORHAN D.

- 51st Annual Scientific Meeting of the European Society for the Paediatric Nephrology, 3 - 06 Ekim 2018
13. **Long-term follow-up results of patients with ADCK4 mutations who have been diagnosed in asymptomatic period: effects of early initiation of CoQ10 supplementation**
ATMACA M., GÜLHAN B., KARABAY BAYAZIT A., CANDAN C., ARICI M., TOPALOĞLU R., ÖZALTIN F.
51st Annual Scientific Meeting of the European Society for the Paediatric Nephrology, 3 - 06 Ekim 2018
14. **Clinical and Mutational Spectrum of Children with Autosomal Recessive and Autosomal Dominant Polycystic Kidney Disease**
TÜRKÖĞLU Ö., GÜLHAN B., YÜKSEL S., ÇALIŞKAN S., DÜZOVA A., ÇAKAR N., SÖYLEMEZOĞLU H. O., TOPALOĞLU R., ÖZALTIN F.
51st Annual Scientific Meeting of the European Society for the Paediatric Nephrology, 3 - 06 Ekim 2018
15. **ACUTE HEMODIALYSIS EXPERIENCE IN CHILDRENWEIGHING LESS THAN 15 KG**
TAŞ N., GÜLHAN B., çelegen k., ÖZDEMİR E. G., DÜZOVA A., ÖZALTIN F., TOPALOĞLU R.
51st Annual Scientific Meeting of the European Society for the Paediatric Nephrology, 3 - 06 Ekim 2018
16. **Effect of IGF-1 and HGF Induced Bone Marrow Mesenchymal Stem Cells on Focal Segmental Glomerulosclerosis in Sprague-Dawley Rats: A pilot study**
Sahan Ö. B., Korkmaz E., Onbaşilar İ., Güçer K. Ş., Kaymaz F. F., Özaltın F., Günel-Özcan A.
43rd Febs Congress, Praha, Çek Cumhuriyeti, 7 - 12 Temmuz 2018
17. **PEPTIDE BIOMARKER SIGNATURES IN STEROID-RESISTANT NEPHROTIC SYNDROME**
Zurbig P., ÖZALTIN F., Anarat A., Paripovic D., Yilmaz A., Caliskan S., Jankauskiene A., Trautmann A., Mischak H., Schaefer F.
55th Congress of the European-Renal-Association (ERA) and European-Dialysis-and-Transplantation-Association (EDTA), Copenhagen, Danimarka, 24 - 27 Mayıs 2018, cilt.33
18. **Naif ve İnduklenmis Mezenkimal Kök Hücrelerin Fokal Segmental Glomeruloskleroz Üzerine Etkisinin Sprague-Dawley Modelinde İncelenmesi**
Sahan Ö. B., Korkmaz E., ONBAŞILAR İ., GÜÇER K. Ş., KAYMAZ F. F., ÖZALTIN F., ÖZCAN A.
4. Klinik Immunoloji Kongresi, Antalya, Türkiye, 11 - 14 Nisan 2018
19. **Çocuk ve genç erişkinlerde renovasküler hipertansiyonda cerrahi tedaviler.**
PEKER R. O., AKI F. T., KUMBASAR U., GÜVENER M., YILMAZ M., ÖZALTIN F., DÜZOVA A., TOPALOĞLU R., PEYNİRCİOĞLU B., DEMİRCİN M.
18. Ulusal Vasküler ve Endovasküler Cerrahi Kongresi 9. Ulusal Fleboloji Kongresi, Girne, Kıbrıs (Kktc), 9 - 12 Aralık 2017
20. **Çocuk ve genç erişkinlerde renovasküler hipertansiyonda cerrahi tedaviler**
PEKER R. O., aki F. T., KUMBASAR U., GÜVENER M., YILMAZ M., DOĞAN R., ÖZALTIN F., DÜZOVA A., TOPALOĞLU R., PEYNİRCİOĞLU B., et al.
18. Ulusal Vasküler ve Endovasküler Cerrahi Kongresi 9. Ulusal Fleboloji Kongresi, Türkiye, 9 - 12 Kasım 2017
21. **Çocuk ve Genç Erişkinlerde Renovasküler Hipertansiyonda Cerrahi Tedavi**
PEKER R. O., AKI F. T., KUMBASAR U., GÜVENER M., YILMAZ M., DOĞAN R., ÖZALTIN F., DÜZOVA A., DEMİRCİN M., TOPALOĞLU R., et al.
18. Ulusal Vasküler ve Endovasküler Cerrahi Kongresi, Türkiye, 9 - 12 Kasım 2017
22. **USE OF HERBAL AND DIETARY SUPPLEMENTS IN CHILDREN WITH KIDNEY AND URINARY TRACT DISEASE**
Öztürk T. T., ÖZMERT E. N., KANBUR N., GÜLHAN B., ÖZALTIN F., TOPALOĞLU R., DÜZOVA A.
PEDIATRIC NEPHROLOGY, 6 - 09 Eylül 2017, cilt.32, ss.1675
23. **Use of herbal and dietary supplements in children with kidney and urinary tract disease**
Taştemel Öztürk T., KANBUR N., ÖZMERT E. N., GÜLHAN B., ÖZALTIN F., TOPALOĞLU R., DÜZOVA A.
50th Anniversary Meeting of the ESPN, 6 - 09 Eylül 2017, cilt.32, ss.1675
24. **Risk factors for cardiovascular comorbidities in children with steroid-sensitiveand steroid-resistant nephrotic syndrome.**
Kara Eroğlu F., TOPALOĞLU R., KARAGÖZ T., OĞUZ B., ÖZALTIN F., GÜLHAN B., DÜZOVA A.
50th Anniversary Meeting of the ESPN, 6 - 09 Eylül 2017, cilt.32, ss.1661
25. **Does C.3979 G A/P.VAL1327MET variant of COL4A4 has any pathogenic effect in Turkish patients**

- with Alport Syndrome ?**
 YILDIZ N., ATA P., ALPAY H., GÖKCE İ., ÖZALTIN F.
 48th Anniversary Meeting of the European Society for Paediatric Nephrology, 6 - 09 Eylül 2017
- 26. Genetic and clinical characteristics of patients with C3 glomerulopathy.**
 TOPALOĞLU R., GÜLHAN B., KORKMAZ E., DÜZOVA A., ÖZALTIN F.
 50th Anniversary Meeting of the European Society for Paediatric Nephrology, 6 - 09 Eylül 2017
- 27. GENETIC AND CLINICAL CHARACTERISTICS OF PATIENTS WITH C3 GLOMERULOPATHY**
 TOPALOĞLU R., GÜLHAN B., korkmaz e., DÜZOVA A., ÖZALTIN F., SOYLU A., Study Group C. G.
 The 50th Anniversary Meeting of the European Society for Pediatric Nephrology, 6 - 09 Eylül 2017
- 28. OC-7 Crescentic glomerulonephritis (CGN) in childhood classification of aetiology and clinicopathological importance of cd163 positive (M2) macrophages**
 KAYKI G., ORHAN D., ÖZALTIN F., TALİM B., DÜZOVA A., AKÇAÖREN z., TOPALOĞLU R., GÜÇER K. S.
 Oral Communications, 7 - 10 Haziran 2017
- 29. Tocilizumab Treatment in childhood takayasu arteritis case series of four patients and systematic review of the literature**
 BATU E. D., SÖNMEZ H. E., HAZIROLAN T., ÖZALTIN F., BİLGİNER Y., ÖZEN S.
 PRES 2016 Genova, 28 Eylül - 01 Ekim 2016
- 30. NEXT GENERATION GENE PANEL SCREENING IN STEROID-RESISTANT NEPHROTIC SYNDROME**
 Lipska-Zietkiewicz B. S., Boyer O., Gribouval O., Tabatabaei M., Fourrage C., Nischke P., Bole-Feysot C., Rotthier A., ÖZALTIN F., Noris M., et al.
 53rd ERA-EDTA Congress, Vienna, Avusturya, 21 - 24 Mayıs 2016, cilt.31, ss.353
- 31. SMARCAL1 SCREENING IN NEPHROTIC SYNDROME - LESSONS FROM PODONET**
 Lipska-Zietkiewicz B. S., Gellermann J., Boyer O., Shalaby M. A., ÖZALTIN F., Dusek J., Melk A., KARABAY BAYAZIT A., Massella L., Schaefer F.
 53rd ERA-EDTA Congress, Vienna, Avusturya, 21 - 24 Mayıs 2016, cilt.31, ss.353-354
- 32. NORMAL 25-HYDROXYVITAMIN D LEVELS ARE ASSOCIATED WITH LESS PROTEINURIA AND ATTENUATE RENAL FAILURE PROGRESSION IN CHILDREN WITH CHRONIC KIDNEY DISEASE**
 Shroff R., Trivelli A., Litwin M., Picca S., Anarat A., Sallay P., ÖZALTIN F., Zurowska A., Jankauskiene A., Montini G., et al.
 52nd Congress of the European-Renal-Association-European-Dialysis-and-Transplant-Assocation, London, Kanada, 28 - 31 Mayıs 2015, cilt.30
- 33. YENİDOĞAN DÖNEMİNDE EPİDERMOLİZİS BÜLLOSA, PİLOR ATREZİSİ VE APLASİA KUTİS: BİR VAKA TAKDİMİ**
 KAYKI G., BOZKAYA D., ÖZALTIN F., ORHAN D., AYGÜN D., KAYMAZ F., KORKMAZ E., YİĞİT Ş.
 23. ULUSAL NEONATOLOJİ KONGRESİ, Antalya, Türkiye, 19 - 22 Nisan 2015
- 34. CONVERSION TO SIROLIMUS IN PEDIATRIC RENAL TRANSPLANT RECIPIENTS**
 TOPALOĞLU R., GÜLHAN B., BİLGİNER Y., Tayfur A. C., Yıldız C., ÖZALTIN F., DÜZOVA A., ÖZEN S., Aki T., BEŞBAŞ N.
 50th European-Renal-Association - European-Dialysis-and-Transplant-Association Congress, İstanbul, Türkiye, 18 - 21 Mayıs 2013, cilt.28, ss.294
- 35. Eye involvement in children with primary FSGS**
 BİLGİNER Y., Poyraz C. E., ÖZALTIN F., KADAYIFÇILAR S., BEŞBAŞ N., Topaloglu R., ÖZEN S., Hildebrandt F., BAKKALOĞLU A.
 44th ERA-EDTA Congress, Barcelona, İspanya, 22 - 24 Haziran 2007, cilt.22, ss.292
- 36. Influence of monocyte chemoattractant protein-1-2518A > G polymorphism on childhood focal segmental glomerulosclerosis**
 Kalyoncu M., Besbas N., Ozgul K., Ozaltin F., Duzova A., Topaloglu R., Ozen S., Ozguc M., Bakkaloglu A.
 43rd ERA-EDTA Congress, Glasgow, Birleşik Krallık, 15 - 18 Temmuz 2006, cilt.21, ss.328

SODYUM VE SU DENGESİ VE BOZUKLUKLARI (DÖNEM III, TÜRKÇE TIP), Ön Lisans, 2016 - 2017
AKUT BÖBREK YETMEZLİĞİ (DÖNEM III, TÜRKÇE TIP), Ön Lisans, 2016 - 2017
SIVI-ELEKTROLİT (DÖNEM IV, PDÖ), Ön Lisans, 2016 - 2017
DEHİDRATASYON VE TEDAVİSİ (DÖNEM IV, TÜRKÇE TIP), Ön Lisans, 2016 - 2017
SODYUM VE SU DENGESİ BOZUKLUKLARI (DÖNEM III, İNGİLİZCE TIP), Ön Lisans, 2016 - 2017
AKUT VE KRONİK BÖBREK YETMEZLİĞİNE YAKLAŞIM (DÖNEM IV), Ön Lisans, 2016 - 2017
POTASYUM DENGESİ VE BOZUKLUKLARI (DÖNEM III, TÜRKÇE TIP), Ön Lisans, 2016 - 2017
Akut Böbrek Yetmezliği, Lisans, 2016 - 2017
EKSTREMİTE MUAYENESİ VE KAN BASINCI ÖLÇÜMÜ (DÖNEM IV), Ön Lisans, 2016 - 2017
DEHİDRATASYON VE TEDAVİSİ (DÖNEM IV, İNGİLİZCE TIP), Ön Lisans, 2016 - 2017
POTASYUM DENGESİ VE BOZUKLUKLARI (DÖNEM III, İNGİLİZCE TIP), Ön Lisans, 2016 - 2017
İDRAR KAÇIRAN ÇOCUĞA YAKLAŞIM, Ön Lisans, 2016 - 2017

Yönetilen Tezler

Özaltın F., Çocukluk çağı otozomal dominant ve otozomal resesif polikistik böbrek hastalıklarının klinik ve genetik özelliklerinin araştırılması, Tıpta Uzmanlık, Ö.TUTAL(Öğrenci), 2021
Özaltın F., ADCK4 mutasyonu saptanan hastaların uzun dönem izlem sonuçları, Tıpta Uzmanlık, M.ATMACA(Öğrenci), 2017

Metrikler

Yayın: 234
Atıf (WoS): 3571
Atıf (Scopus): 4056
H-İndeks (WoS): 31
H-İndeks (Scopus): 31

Araştırma Alanları

Tıp, Sağlık Bilimleri, Dahili Tıp Bilimleri, Çocuk Sağlığı ve Hastalıkları, Pediatrik Nefroloji

Akademi Dışı Deneyim

Hacettepe Üniversitesi Tıp Fakültesi