

Asst. Prof. ŞULE ALTINER

Personal Information

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International Researcher IDs

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Yoksis Researcher ID: 191387

Education Information

Expertise In Medicine, Ankara University, Turkey 2012 - 2017

Undergraduate, Ankara University, Tıp Fakültesi, Turkey 2003 - 2009

Dissertations

Expertise In Medicine, Molecular karyotyping in patients with multiple congenital anomalies / mental retardation, Ankara University, 2017

Research Areas

Medical Genetics

Academic Titles / Tasks

Assistant Professor, Ankara University, Tıp Fakültesi, Dahili Tıp Bilimleri Bölümü, 2022 - Continues

Lecturer, Ankara University, Tıp Fakültesi, Dahili Tıp Bilimleri Bölümü, 2020 - 2022

Research Assistant, Katholieke Universiteit Leuven, Genetik, İnsan Genetiği, 2017 - 2017

Research Assistant, Ankara University, Tıp Fakültesi, Dahili Tıp Bilimleri Bölümü, 2012 - 2017

Courses

Genetics of thyroid and parathyroid diseases, MEN syndromes, Undergraduate, 2024 - 2025, 2023 - 2024, 2022 - 2023, 2021 - 2022

Genetic mechanisms of development and disorders of excretory system, Undergraduate, 2024 - 2025, 2023 - 2024, 2022 - 2023, 2021 - 2022

Bağ dokusu gelişimi ve bozukluklarında genetik mekanizmalar, Undergraduate, 2024 - 2025, 2023 - 2024, 2022 - 2023, 2021 - 2022

Mendel kuralları ve kalıtım kavramı, Undergraduate, 2024 - 2025, 2023 - 2024, 2022 - 2023, 2021 - 2022

Genetics of diabetes mellitus and obesity, Undergraduate, 2024 - 2025, 2023 - 2024, 2022 - 2023, 2021 - 2022

Genetic mechanisms of connective tissue development and disorders , Undergraduate, 2024 - 2025, 2023 - 2024, 2022 - 2023, 2021 - 2022

Hereditary breast cancer, Undergraduate, 2024 - 2025, 2023 - 2024, 2022 - 2023

Rare Diseases: General information and diagnostic approaches from clinic to research, Undergraduate, 2024 - 2025

Evolution of the nervous system, Undergraduate, 2024 - 2025

Kromozomların yapısı, Undergraduate, 2024 - 2025, 2023 - 2024, 2022 - 2023, 2021 - 2022

I choose my career; being a member of medical genetics team, Undergraduate, 2024 - 2025, 2023 - 2024, 2022 - 2023

Güncel genom düzenleme yöntemleri , Undergraduate, 2024 - 2025, 2023 - 2024

Undergraduate, 2024 - 2025

Undergraduate, 2024 - 2025

Noncoding RNAs and regulation of gene expression, Undergraduate, 2024 - 2025, 2023 - 2024, 2022 - 2023, 2021 - 2022

Current genome editing methods , Undergraduate, 2024 - 2025, 2023 - 2024

Gen ekspresyon ve işlevinin incelenmesi, Undergraduate, 2024 - 2025, 2023 - 2024, 2022 - 2023, 2021 - 2022

Undergraduate, 2024 - 2025

Undergraduate, 2024 - 2025

Published journal articles indexed by SCI, SSCI, and AHCI

- I. **Perception and management of cancer predisposition in pediatric cancer centers: A European-wide questionnaire-based survey**
Lazic J., Haas O. A., Özbek U., Ripperger T., Byrjalsen A., te Kronnie G., Sayitoğlu M., Ng O. H., Agaoglu N. B., Erbilgin Y., et al.
Pediatric Blood and Cancer, vol.70, no.5, 2023 (SCI-Expanded)
- II. **Contribution of genotypes in Prothrombin and Factor V Leiden to COVID-19 and disease severity in patients at high risk for hereditary thrombophilia.**
Kiraz A., Sezer O., ALEMDAR A., Canbek S., Duman N., BİŞGİN A., Cora T., Ruhi H., Ergoren M. C., GEÇKİNLİ B. B., et al.
Journal of medical virology, vol.95, no.2, 2023 (SCI-Expanded)
- III. **Germline landscape of BRCA by 7-site collaborations as a BRCA consortium in Turkey**
BİŞGİN A., ÖZEMRİ SAĞ Ş., DOĞAN M. E., Yildirim M. S., Gumus A. A., Akkus N., Balasar O., Durmaz C. D., Ersoz R., Altiner S., et al.
BREAST, vol.65, pp.15-22, 2022 (SCI-Expanded)
- IV. **Phenotypic and molecular characterization of five patients with *PIK3CA*-related overgrowth spectrum (PROS)**
Ili E., Tasdelen E., Durmaz C. D., ALTINER Ş., TUNCALI T., Martinez-Glez V., KARABULUT H. G., Vural S., Ceylaner S., ACAR M. O., et al.
AMERICAN JOURNAL OF MEDICAL GENETICS PART A, vol.188, no.6, pp.1792-1800, 2022 (SCI-Expanded)
- V. **A Paternal "Balanced" Chromosome 2 and 4 Translocation with Chromosome 21q Insertion Leading to Duplication of 2q22.1q24.1 in Two Siblings**
Gökoğlu M., Kutlay N., Altiner Ş.
no.162, pp.297-305, 2022 (SCI-Expanded)
- VI. **Extending Phenotypic Spectrum of 17q22 Microdeletion: Growth Hormone Deficiency**
Durmaz C. D., ALTINER Ş., Tasdelen E., KARABULUT H. G., ILGIN RUHİ H.
FETAL AND PEDIATRIC PATHOLOGY, vol.40, no.5, pp.486-492, 2021 (SCI-Expanded)
- VII. ***MASP1*-related 3MC syndrome in a patient from Turkey**
Durmaz C. D., Altiner Ş.
AMERICAN JOURNAL OF MEDICAL GENETICS PART A, vol.185, no.7, pp.2267-2270, 2021 (SCI-Expanded)
- VIII. **Severe skin fragility with postnatal lethal outcome due to a biallelic *KRT5* mutation**
Altiner Ş., Hekimoglu B., Livaoglu A., Has C.
JOURNAL DER DEUTSCHEN DERMATOLOGISCHEN GESELLSCHAFT, vol.19, no.3, pp.440-442, 2021 (SCI-Expanded)
- IX. **Application of Chromosome Microarray Analysis in the Investigation of Developmental Disabilities and Congenital Anomalies: Single Center Experience and Review of *NRXN3* and**

<i>NEDD4L</i> Deletions

ÇEBİ A. H., Altiner Ş.

MOLECULAR SYNDROMOLOGY, no.4, pp.197-206, 2020 (SCI-Expanded)

- X. **Mosaic Small Supernumerary Marker Chromosome Derived from Five Discontinuous Regions of Chromosome 8 in a Patient with Neutropenia and Oral Aphthous Ulcer**
ALTINER Ş., KUTLAY N., İLGIN RUHİ H.
CYTOGENETIC AND GENOME RESEARCH, vol.160, no.1, pp.11-17, 2020 (SCI-Expanded)
- XI. **Extending the Phenotypic Spectrum of Huntington Disease: Hypothermia**
Altiner Ş., Ardic S., ÇEBİ A. H.
MOLECULAR SYNDROMOLOGY, vol.11, no.1, pp.56-58, 2020 (SCI-Expanded)
- XII. **A Novel Atrx Mutation Presenting with Intellectual Disability and Severe Kyphoscoliosis**
Altiner Ş., Raymond L.
FETAL AND PEDIATRIC PATHOLOGY, vol.39, no.6, pp.539-543, 2020 (SCI-Expanded)
- XIII. **Cytogenetic, Molecular, and Phenotypic Characterization of a Patient with de novo Derivative Chromosome 18 and Review of the Literature**
Gokpınar E., Altiner Ş., KARABULUT H. G.
CYTOGENETIC AND GENOME RESEARCH, no.2, pp.74-80, 2019 (SCI-Expanded)
- XIV. **Importance of patient selection criteria in determining diagnostic copy number variations in patients with multiple congenital anomaly/mental retardation**
Altiner Ş., KUTLAY N.
MOLECULAR CYTOGENETICS, 2019 (SCI-Expanded)
- XV. **A novel TWIST1 gene mutation in a patient with Saethre-Chotzen syndrome**
ALTINER Ş., KARABULUT H. G., Yararbas K., Tukun A., Collet C., Kocaay P., BERBEROĞLU M., İLGIN RUHİ H.
CLINICAL DYSMORPHOLOGY, vol.26, no.3, pp.175-178, 2017 (SCI-Expanded)
- XVI. **Prognostic impact of RUNX1 and ETV6 gene copy number on pediatric B-cell precursor acute lymphoblastic leukemia with or without hyperdiploidy**
KUTLAY N., Pekpak E., ALTINER Ş., İLERİ D. T., Vicdan A. N., DİNÇASLAN H., İNCE E., Tukun F. A.
INTERNATIONAL JOURNAL OF HEMATOLOGY, vol.104, no.3, pp.368-377, 2016 (SCI-Expanded)
- XVII. **Constitutional Trisomy 8 Mosaicism with Persistent Macrocytosis**
ALTINER Ş., KUTLAY N., İLHAN O.
CYTOGENETIC AND GENOME RESEARCH, vol.150, no.1, pp.35-39, 2016 (SCI-Expanded)

Articles Published in Other Journals

- I. **Molecular and Clinical Overview of Type 1 Neurofibromatosis: Single Center Study and Mini Review on NF1-Associated Vasculopathy and Juvenile Myelomonocytic Leukemia**
Altiner Ş., Çebi A. H.
GAZI MEDICAL JOURNAL, vol.35, no.4, pp.422-432, 2024 (ESCI)
- II. **Genotype/Phenotype Correlation of Cases with PTPN11 Gene Mutation: Eastern Black Sea Experience**
ALTINER Ş., ÇEBİ A. H., ÇELİK S., GÖKCÜ M.
Ankara Üniversitesi Tıp Fakültesi Mecmuası, vol.75, no.3, pp.368-372, 2022 (Peer-Reviewed Journal)
- III. **Genetic diagnosis of maturity-onset diabetes of the young (MODY) in northeast Turkey**
ALTINER Ş., ÇELİK S., YARAR M. H., ÇEBİ A. H.
İnönü Üniversitesi Tıp Fakültesi Dergisi, vol.29, no.4, pp.325-328, 2022 (Peer-Reviewed Journal)
- IV. **Genotype Phenotype Correlation of A Case Having Chromosome 3 Imbalance**
Taşdelen E., Gökpınar İli E., Altiner Ş., Ceylan A. C., Tuncalı T.
Ankara Üniversitesi Tıp Fakültesi Mecmuası, vol.74, no.3, pp.365-369, 2021 (Peer-Reviewed Journal)
- V. **Molecular Testing for Thalassemia: Mutation Detection According to Referral Reasons and Demographic Data**

ALTINER Ş., KARABULUT H. G., EKİNCİ S., Vicdan A., KUTLAY N., TUNCALI T., Tukun A., ILGIN RUHİ H.
ERCIYES MEDICAL JOURNAL, vol.43, no.5, pp.449-451, 2021 (ESCI)

- VI. **Genetic analysis of BCR-ABL negative chronic myeloproliferative diseases at initial diagnosis and their clinical effects**
UYSAL A., ALTINER Ş., ÇELİK S., UYSAL S., ÇEBİ A. H.
CUKUROVA MEDICAL JOURNAL, vol.45, no.3, pp.933-939, 2020 (ESCI)
- VII. **Type I plasminogen deficiency with unexpected clinical aspects: Could be more than coexistence?**
Altner Ş., Klammt J., Bernhard M., Schuster V., KARABULUT H. G.
COGENT MEDICINE, vol.4, 2017 (Peer-Reviewed Journal)

Books & Book Chapters

- I. **Down Syndrome Phenotype, Underlying Mechanisms and Treatment Strategies**
Altner Ş.
in: Down Syndrome, Hatice İlgin Ruhi, Editor, Türkiye Klinikleri Yayınevi, Ankara, pp.92-97, 2024
- II. **Gen Tedavisi ve Genom Düzenlenmesi: Malign Hastalıklarda Gen Tedavisi Uygulamaları**
Altner Ş.
in: Laboratuvaradan Kliniğe Hematoloji Alanında Yenilikçi Tedaviler, Kurt Yüksel M, Yürür Kutlay N, Editor, Türkiye Klinikleri Yayınevi, Ankara, pp.118-124, 2023
- III. **COVID-19 ve Genetik Yatkınlık**
EKİNCİ S., ALTINER Ş., ILGIN RUHİ H.
in: COVID-19, Osman Memikoğlu, Volkan Genç, Editor, Ankara Üniversitesi Basımevi, pp.29-36, 2021
- IV. **LH Reseptör Bozukluğuna Bağlı 46XY Cinsiyet Gelişim Bozukluğu**
ALTINER Ş.
in: Çocuk ve Ergenlerde Gonad Hastalıkları, Yusuf Kenan Haspolat, Fesih Aktar, Serhat Ege, Salim Bilici, Editor, Orient Yayınları, Ankara, pp.311-317, 2019
- V. **17 Alfa Hidroksilaz / 17-20 Liyaz Enzim Eksikliği .**
ALTINER Ş.
in: Çocuk ve Ergenlerde Gonad Hastalıkları, Yusuf Kenan Haspolat, Fesih Aktar, Serhat Ege, Salim Bilici, Editor, Orient Yayınları, Ankara, pp.301-309, 2019

Refereed Congress / Symposium Publications in Proceedings

- I. **A rare initial presentation of Li-Fraumeni syndrome: mesothelioma**
Leblebici C. B., Altner Ş., Karabulut H. G., Serbes E., Savaş B., Unal E. C.
56th Annual Conference of the European-Society-of-Human-Genetics (ESHG), Glasgow, England, 10 - 13 June 2023, pp.592-593
- II. **Long overdue diagnosis for Turner syndrome: evaluation of two cases**
İlgin Ruhi H., Durmaz C. D., Leblebici C. B., Altner Ş.
56th Annual Conference of the European-Society-of-Human-Genetics (ESHG), Glasgow, England, 10 - 13 June 2023, pp.245-246
- III. **A GHOSAL HEMATODIAPHYSEAL DYSPLASIA CASE; EXCELLENT RESPONSE TO NON-STEROIDAL ANTI-INFLAMMATORY DRUG TREATMENT**
ÇAKMAKLI H. F., MUTLU H., ALTINER Ş., AYDIN F., İLERİ D. T., İNCE E., ERTEM M.
14th Eurasian Hematology Oncology Congress, İstanbul, Turkey, 13 - 14 October 2023, vol.45, no.3, pp.23-24
- IV. **Yetişkin Akut Miyeloid Lösemi/Miyelodisplastik Sendrom Hastalarında Konvansiyonel Sitogenetik İncelemenin Kompleks Karyotip Saptamadaki Gücü**
Altner Ş., Vicdan N. A., Kutlay N., Tuncali T., Karabulut H. G., İlgin Ruhi H.
15.Ulusal Tıbbi Genetik Kongresi, Muğla, Turkey, 9 - 13 November 2022, pp.102-103

- V. **Beaulieu-Boycott-Innes Sendromu: İki Olgu Sunumu**
ÖZBOLAT S., YURTDAŞ A. K., SEVİM B., ALTINER Ş., MUTLU H., HAVAN M., KENDİRLİ T., YÜRÜR KUTLAY N., KARABULUT H. G.
15. ULUSAL TIBBİ GENETİK KONGRESİ, Muğla, Turkey, 9 - 13 November 2022, pp.151
- VI. **A novel mutation in a patient with KIDAR syndrome: tenth patient in the literature**
ALTINER Ş., YURTDAŞ A. K., DOĞULU N., KÖSE E., EMİNOĞLU F. T., KARABULUT H. G.
European Human Genetics Conference, Viyana, Austria, 11 - 14 June 2022
- VII. **Cytogenetic evaluation in myelodysplastic syndrome**
LEBLEBİCİ C. B., ALTINER Ş., VİCDAN N. A., KUTLAY N., TUNCALI T., KARABULUT H. G., EKİNCİ S., ILGIN RUHİ H.
European Human Genetics Conference, Viyana, Austria, 11 - 14 June 2022
- VIII. **A case of PTEN hamartoma tumor syndrome; a family study**
İlgin Ruhi H., Gökpınar İli E., Güleray Lafcı N., Taşdelen E., Altner Ş.
54th European Society of Human Genetics (ESHG) Conference, 28 - 31 August 2021, pp.424-425
- IX. **Evaluation of Plasma Cell Molecular Cytogenetic Findings of Myeloma Patients: One-Year Single-Center Experience**
KAPLAN İ., CESUR BALTACI H. N., ALTINER Ş., EKİNCİ S., VİCDAN N. A., KARABULUT H. G., TUNCALI T., ILGIN RUHİ H., KUTLAY N.
ESHG 2021, 28 August 2022, pp.413
- X. **Famlyal Adenomatöz Polipozis’de Genetik Test Ve Danışmanın Önemi: Bir Aile Örneği**
ALTINER Ş., TUNCALI T.
1. Ulusal Hemotoonkogenetik kongresi, Antalya, Turkey, 25 - 28 November 2021
- XI. **Multiple Miyelom Hastalarının Plazma Hücrelerindeki Moleküler Sitogenetik İnceleme Sonuçları: Ankara Üniversitesi Deneyimi**
KAPLAN İ., CESUR BALTACI H. N., ALTINER Ş., EKİNCİ S., VİCDAN N. A., KARABULUT H. G., TUNCALI T., ILGIN RUHİ H., KUTLAY N.
1. Ulusal HematoOnkoGenetik Kongresi, Antalya, Turkey, 25 November 2021, pp.29
- XII. **Son İki Yılda Kliniğimize Başvuran Yeni Tanı AML Hastalarında Genetik Parametrelerin Değerlendirilmesi**
SEVİM B., KURTÇU O., ALTINER Ş., EKİNCİ S., VİCDAN N. A., TUNCALI T., KARABULUT H. G., ILGIN RUHİ H., KUTLAY N.
1. Ulusal HematoOnkoGenetik Kongresi, Antalya, Turkey, 25 - 28 November 2021
- XIII. **Fanconi anemisi: FANCA Geninde Compound Heterozigot Mutasyonlu İki Olgu**
ACAR M. O., KAPLAN İ., TUNÇEZ E., ALTINER Ş., KARABULUT H. G.
1. Ulusal HematoOnkoGenetik Kongresi, Antalya, Turkey, 25 - 28 November 2021
- XIV. **Kütanöz T Hücreli Lenfomada Görülen Konvansiyonel Sitogenetik ve Akım Sitometri Bulguları**
LEBLEBİCİ C. B., ALTINER Ş., VİCDAN N. A., KUTLAY N., TUNCALI T., KARABULUT H. G., ILGIN RUHİ H.
1. Ulusal HematoOnkoGenetik Kongresi, Antalya, Turkey, 25 - 28 November 2021
- XV. **Clinical findings in 22q11.2 microdeletion syndrome: case series**
ALTINER Ş., TUNCALI T., KUTLAY N., KARABULUT H. G., ILGIN RUHİ H.
European Human Genetics Virtual Conference, 28 - 31 August 2021
- XVI. **Çift Anöploidi Saptanan İki Olgu**
Altner Ş., Sevim B., Karabulut H. G., İlgin Ruhi H., Şahinarslan A.
14.Ulusal Tıbbi Genetik Kongresi “Uluslararası Katılımlı”, 20 - 22 November 2020, vol.31, pp.1-93
- XVII. **Maternal rcp(15;21) nedeniyle oluşan Down sendromu olgusu**
Sevim B., Altner Ş., Kaplan İ., Okulu E., İlgin Ruhi H.
14. Ulusal Tıbbi Genetik Kongresi, 20 - 22 November 2020
- XVIII. **14q32.31q32.33 Delesyon Olgu Sunumu**
ACAR M. O., ALTINER Ş., KENDİRLİ T., KUTLAY N., TUNCALI T.
14.Ulusal Tıbbi Genetik Kongresi “Uluslararası Katılımlı”, Ankara, Turkey, 14 October 2020, vol.31
- XIX. **A case with complex small supernumerary marker chromosome consisting 19p and 22q.**
ALTINER Ş., KUTLAY N.

European Human Genetics Conference, Berlin, Germany, 6 - 09 June 2020

- XX. **Basal Cell Nevus Syndrome: A Case With 9Q22.3 Microdeletion**
ALTINER Ş., ÇEBİ A. H.
1st Bursa International Genetics Days: Dermatogenetics Symposium, 09 January 2020
- XXI. **Interstitial microdeletion of 17q22 in a patient with de novo apparently balanced t(117)**
TAŞDELEN E., DURMAZ C. D., ALTINER Ş., ILGIN RUHİ H.
13. Balkan Congress of Human Genetics, 17 - 20 April 2019
- XXII. **ATN1 gene mutation in patients with Huntington disease-like phenotype**
KARABULUT H. G., GÖKOĞLU M., ALTINER Ş., DURMAZ C. D., GÖKPINAR İLİ E., VİCDAN N. A., KUTLAY N., ILGIN RUHİ H., TUNCALI T., TÜKÜN F. A.
13th Balkan Congress of Human Genetics, Edirne, Turkey, 17 - 20 April 2019
- XXIII. **Clinical and Genetic Analysis of the Bcr-Abl Negative Chronic Myeloproliferative Diseases in Initial Diagnosis: Single Central Experience**
UYSAL A., ALTINER Ş., ÇELİK S., UYSAL S., ÇEBİ A. H.
5th Aegean Hematology Oncology Symposium, PORTO CHELI, Greece, 20 - 23 September 2018, pp.31
- XXIV. **SDR5A2 Gen Mutasyonu Saptanan İki Olgu**
Gökpinar İli E., Altiner Ş., Gökoğlu M., Şıklar Z., Berberoğlu M., İlgin Ruhi H.
XII. Ulusal Tıbbi Genetik Kongresi, İzmir, Turkey, 5 - 09 October 2016, pp.157
- XXV. **Multiple Konjenital anomali/mental retardasyonlu üç olguda subtelomerik FISH bulguları**
ALTINER Ş., GÖKOĞLU M., TUNCALI T., KUTLAY N.
12. ULUSAL TIBBİ GENETİK KONGRESİ, İzmir, Turkey, 5 - 09 October 2016
- XXVI. **Huntington s disease mimicking bipolar disorder a case report**
HosgorenAlici Y., Ozel Kızıl E. T., Kırıcı S., BİLGİN B., BİÇER Ş., KUTLAY N.
27th Congress of the European-College-of-Neuropsychopharmacology, 18 - 21 October 2014, vol.24, pp.639
- XXVII. **Talasemi ön tanısı ile incelenen 206 hastada saptanan mutasyonların başvuru nedenlerine göre dağılımı**
BİÇER Ş., KARABULUT H. G., Ekinci S., Mutlu M. B., VİCDAN N. A., KUTLAY N., TUNCALI T., TÜKÜN F. A., ILGIN RUHİ H.
I. Hematolojik Genetik Sempozyumu, Turkey, 2 - 04 December 2013
- XXVIII. **Multiple Myelomda konvansiyonel sitogenetik ve FISH yöntemlerinin birlikte kullanımının klinik yararı: Tek merkezden 1150 olgunun sonuçları**
Mutlu M. B., VİCDAN N. A., BİÇER Ş., Ekinci S., KUTLAY N., KARABULUT H. G., TUNCALI T., TÜKÜN F. A., ILGIN RUHİ H.
I. Hematolojik Genetik Sempozyumu, İzmir, Turkey, 2 - 04 December 2013, pp.53
- XXIX. **Akut lenfoblastik lösemide konvansiyonel sitogenetik, FISH ve real-time PCR yöntemlerinin birlikte kullanımının klinik yararı: tek merkezden 1050 olgunun sonuçları**
Ekinci S., VİCDAN N. A., BİÇER Ş., Mutlu M. B., KUTLAY N., KARABULUT H. G., TUNCALI T., ILGIN RUHİ H., TÜKÜN F. A.
I. Hematolojik Genetik Sempozyumu, İzmir, Turkey, 2 - 04 December 2013, pp.52
- XXX. **Inherited retinal diseases and recent advances in therapy**
ALTINER Ş.
V. Uluslararası Katılımlı Erciyes Tıp Genetik Günleri Kongresi, 20 - 22 February 2020, vol.31, pp.1-94

Supported Projects

Altiner Ş., Tuncali T., Kurtçu O., Yurtdaş A. K., Project Supported by Other Official Institutions, Two new cases diagnosed with KIDAR (keratitis, ichthyosis, deafness, autosomal recessive) syndrome and literature review, 2024 - 2025
Altiner Ş., Çağdaş Ayvaz D. N., Ulum B., TÜBİTAK Project, KROMOZOM KIRIKLARI İLE GİDEN KOMBİNE İMMÜN YETMEZLİKLERDE DEB VEH2AX TESTLERİNİN KARSILASTIRILMALI OLARAK DEĞERLENDİRİLEREK HASTALIĞIÇIN TANISAL ALGORITMA AMACIYLA KULLANIMI, 2023 - 2024

Kutlay N., Altner Ş., Project Supported by Higher Education Institutions, Multipl konjenitalanomali/Mental retardasyon Hastalarında Moleküler Karyotipleme, 2015 - 2017

Kutlay N., Altner Ş., TUBITAK Project, En Sık Hastalık Nedeni Olan Dokuz Mutasyonun Saptanmadığı Konjenital Adrenal Hiperplazi Hastalarında Cyp21 Genindeki Mutasyonların Araştırılması, 2015 - 2016

Memberships / Tasks in Scientific Organizations

Tıbbi Genetik Derneği, Board Member, 2021 - Continues, Turkey

Tasks In Event Organizations

Altner Ş., Fall School of Medical Genetics - Applied Hematologic Malignancies Course, Science / Art Camp or Summer School Organization, Ankara, Turkey, Ekim 2024

Altner Ş., Spring School of Medical Genetics , Science / Art Camp or Summer School Organization, Ankara, Turkey, Mayıs 2024

Altner Ş., İleri Tedaviler Sempozyumu, Scientific Congress, Gaziantep, Turkey, Aralık 2023

Altner Ş., Tıbbi Genetik Derneği - Asistan Okulu, Science / Art Camp or Summer School Organization, Ankara, Turkey, Ekim 2023

Altner Ş., 2. Ulusal Hematoonkogenetik Kongresi, Scientific Congress, İskele, Cyprus (Kktc), Mayıs 2023

Altner Ş., 15. Ulusal Tıbbi Genetik Kongresi, Scientific Congress, Muğla, Turkey, Kasım 2022

Altner Ş., 1. Ulusal Hematoonkogenetik Kongresi, Scientific Congress, Antalya, Turkey, Kasım 2021

Metrics

Publication: 61

Citation (WoS): 14

Citation (Scopus): 40

H-Index (WoS): 2

H-Index (Scopus): 4