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Education Information

Post Doctorate, Universiteit Antwerpen-Universitair Centrum Antwerpen, Gelibolu Piri Reis Meslek Yüksekokulu, Moleküler Genetik, Belgium 1994 - Continues

Post Doctorate, Cukurova University, Tıp Fakültesi, Sitogenetik Genetik, Turkey 1990 - Continues

Doctorate, Cukurova University, Sağlık Bilimleri Enstitüsü, Tıbbi Biyoloji Ve Genetik, Turkey 1988 - 1991

Postgraduate, Cukurova University, Sağlık Bilimleri Enstitüsü, Tıbbi Biyoloji Ve Genetik, Turkey 1984 - 1987

Undergraduate, Hacettepe University, Fen Fakültesi, Biyoloji, Turkey 1977 - 1982

Dissertations

Doctorate, Sıtma Vektörü Anopheles sacharovi Favre'de Konak Tercihi, Cukurova University, Temel Tıp, Tıbbi Biyoloji, 1991

Postgraduate, Konakçı çeşitinin Anopheles sacharovi Favre'nin yumurta verimi ve ömür uzunluğuna etkisi, Cukurova University, Tıp Fakültesi, Tıbbi Biyoloji Ve Genetik, 1987

Research Areas

Medicine, Health Sciences, Fundamental Medical Sciences, Medical Biology

Academic Titles / Tasks

Associate Professor, Cukurova University, Tıp Fakültesi, Tıbbi Biyoloji Ve Genetik, 1999 - Continues

Assistant Professor, Cukurova University, Tıp Fakültesi, Tıbbi Biyoloji Ve Genetik, 1992 - Continues

Medical Doctor, Cukurova University, Tıp Fakültesi, Tıbbi Biyoloji Ve Genetik, 1991 - Continues

Research Assistant, Cukurova University, Tıp Fakültesi, Tıbbi Biyoloji Ve Genetik, 1987 - Continues

Other, Hacettepe University, Fen Fakültesi, Biyoloji, 1986 - Continues

Professor, Cukurova University, Tıp Fakültesi, Temel Tıp, 1983 - Continues

Academic and Administrative Experience

Cukurova University, Tıp Fakültesi, Temel Tıp, 1983 - Continues

Cukurova University, 2014 - 2019

Cukurova University, Tıbbi Biyoloji Ve Genetik , 2004 - 2019

Cukurova University, Tıbbi Biyoloji Ve Genetik , 2006 - 2009

Cukurova University, Ç.Ü. Gen Ve Embriyo Mühendisliği Araştırma Ve Uygulama Merkezi Müdür , 2000 - 2008

Cukurova University, 2002 - 2005

Cukurova University, 1998 - 2003

Advising Theses

- DEMİRHAN O., Girişimsel Kardiyak Radyolojik İşlemlerin İnsan Kromozomları Üzerine Genotoksik Etkisi, Postgraduate, N.Çetinel(Student), 2017
- DEMİRHAN O., Akut Lenfoblastik Lösemili Çocuklarda VEGF-A, MMP-2, MMP-9, TIMP-1 ve TIMP-2 Genlerinin Ekspresyon ve Metilasyon Düzeylerindeki Değişikliklerin İncelenmesi, Doctorate, N.İnandıkloğlu(Student), 2014
- DEMİRHAN O., 900-1800 Mhz Radyofrekans Elektromanyetik Alanın (Rf-Ema) İnsan Fetal Hücre Kültürlerinde Kromozomlar Üzerine Etkileri, Postgraduate, İ.Nur(Student), 2014
- DEMİRHAN O., İnvaziv Olmayan Düşük Malignite Potansiyelli Mesane Tümörleriyle, İnvaziv Yüksek Malignite Dereceli Tümörlerin Genetik Açından Farklılıkları, Expertise In Medicine, D.Abat(Student), 2011
- DEMİRHAN O., Sağlık Sendromlu Hastaların Sitogenetik Analizleri ve Kalsiyuma Duyarlı Reseptör Geni 2. ve 3. Ekzon Dizilerinin Belirlenmesi, Doctorate, E.Tunç(Student), 2011
- DEMİRHAN O., Frajil Sendromu Ön Tanısı Alan Çocuklarda Sitogenetik Değişiklikler ve FMR1 Geni Trinükleotid Tekrar Sayılarının Araştırılması, Postgraduate, O.Özer(Student), 2010
- DEMİRHAN O., Nöroblastoma Hastalarında Kromozom Düzensizlikleri, MYCN ve AURKA Gen Değişikliklerinin İncelenmesi, Postgraduate, N.İnandıkloğlu(Student), 2010
- DEMİRHAN O., Akciğer kanserli hastalarda gözlenen kromozomal düzensizlikler ile hücre ölüm reseptörü-4 genindeki polimorfizmlerin incelenmesi, Doctorate, D.Taştemir(Student), 2008
- DEMİRHAN O., İlk Trimester Spontan Düşüklerde Fetal, Maternal ve Paternal Sitogenetik İncelemeler., Postgraduate, E.Tunç(Student), 2005
- DEMİRHAN O., Nöropsikiyatrik ve Zeka Özürlü Hastaların Sitogenetik İncelenmesi, Postgraduate, D.TAŞTEMİR(Student), 2002

Published journal articles indexed by SCI, SSCI, and AHCI

- I. **Relationship Between Long Arm Duplication of X Chromosome and Lysosomal Storage Disease**
Demirhan O.
STEM CELL RESEARCH AND THERAPY, vol.1, no.1, pp.1-4, 2023 (SCI-Expanded)
- II. **Cytogenetic status of patients with congenital malformations or suspected chromosomal abnormalities in Turkey: a comprehensive cytogenetic survey of 11,420 patients**
Demirhan O., Tunc E.
CHROMOSOMA, vol.131, no.4, pp.225-237, 2022 (SCI-Expanded)
- III. **Modifications of the locomotor system in habitually quadrupedal humans**
Tardieu C., Demirhan O., Akbal E., Ozgozen L., Biçer Ö. S., Delapre A., Cornette R., Herrel A.
JOURNAL OF ANATOMY, vol.241, no.3, pp.765-775, 2022 (SCI-Expanded)
- IV. **STRIKING NOVEL MULTI-MISMUTATIONS ON GNAS1, FGF23 AND FGFR3 GENES IN CKD WITH SECONDARY HYPERPARATHYROIDISM (SH) AND SAGLIKER SYNDROME (SS): A COMBINATION OF BONE DISPLASIAS, CKD, SH**
Sagliker Y., Demirhan O., Arslan A., Akbal E., Ergun S., Bayraktar R., Sagliker H. S., Gunesacar R., Ozkaynak S. P., Paylar N., et al.
NEPHROLOGY, vol.24, pp.41, 2019 (SCI-Expanded)
- V. **Fp472 striking Novel Multi- Mismutations On Gnas1, Fgf23 And Fgfr3 Genes In Ckd With Secondary**

Hyperparathyroidism (Sh). Sagliker Syndrome (Ss). Ss Is A Combination -Compulsion Of Bone Displasias-Hereditary Osteodystrophies And Sh And Ckd

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Nephrology Dialysis Transplantation, vol.34, 2019 (SCI-Expanded)

- VI. **Alteration of Peripheral Blood T-cell Subsets in Patients with Cardiovascular Disease; Exposure to Ionizing Radiation (X-rays) and Contrast Medium**
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INTERNATIONAL JOURNAL OF AUDIOLOGY, vol.5, no.2, pp.104-108, 2018 (SCI-Expanded)
- VII. **Alteration of Peripheral Blood T-cell Subsets in Patients with Cardiovascular Disease; Exposure to Ionizing Radiation (X-rays) and Contrast Medium**
DEMİRHAN O., Çetinel N., ÇETİNER S., ÇAĞLIYAN Ç. E., CÜREOĞLU A., USLU İ. N., DEVECİ O. S., SERTDEMİR Y., DEMİRTAŞ M.
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- VIII. **Alteration of Peripheral Blood T-cell Subsets in Patients with Cardiovascular Disease; Exposure to Ionizing Radiation (X-rays) and Contrast Medium**
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International Journal Of Cardiology, vol.5, no.2, pp.104-108, 2018 (SCI-Expanded)
- IX. **"Analysis of maternal polymorphism of CBS gene and risk of Down syndrome offspring",**
PAZARBAŞI A., ÖZPAK L., DEMİRHAN O.
European Journal Of Human Genetics, vol.26, pp.827, 2018 (SCI-Expanded)
- X. **"Chromosomal alterations in patients with breast cancer",**
DEMİRHAN O., TANRIVERDİ N., PAZARBAŞI A., süleymanova D.
European Journal Of Human Genetics, vol.26, pp.536, 2018 (SCI-Expanded)
- XI. **Frequencies and distributions of sex chromosome abnormalities in females with the Turner phenotype: a long-term retrospective study in the southern region of Turkey**
TANRIVERDİ N., Demirhan O., SÜLEYMANOVA D., PAZARBAŞI A.
Turkish journal of medical sciences, vol.47, pp.1447-1455, 2017 (SCI-Expanded)
- XII. **MULTIPLE GNAS1, FGF23, FGFR3 GENES' STRIKING MUTATIONS IN CKD PATIENTS WITH SH. NEW BONE DISPLASIA-HEREDITARY OSTEODISTROPHY AND UGLIFYING HUMAN FACE APPEARANCES. SAGLIKER SYNDROME**
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NEPHROLOGY DIALYSIS TRANSPLANTATION, vol.32, 2017 (SCI-Expanded)
- XIII. **Chromosomal findings and sequence analysis of target exons of calcium-sensing receptor (CaSR) gene in patients with Sagliker syndrome**
Tunc E., DEMİRHAN O., Sağliker Y., Yıldız I., Paylar N., Guzel A. I.
TURKISH JOURNAL OF MEDICAL SCIENCES, vol.47, no.1, pp.13-21, 2017 (SCI-Expanded)
- XIV. **Frequencies and distributions of sex chromosome abnormalities in females with Turner phenotype in south region of Turkey.**
TANRIVERDİ N., DEMİRHAN O., süleymanova D., PAZARBAŞI A.
Turkish Journal Of Medical Sciences, vol.47, no.5, pp.1445-1457, 2017 (SCI-Expanded)
- XV. **Effects of GSM-like radiofrequency irradiation during the oogenesis and spermiogenesis of Xenopus laevis**
Boga A., EMRE M., SERTDEMİR Y., UNCU I., Binokay S., DEMİRHAN O.
ECOTOXICOLOGY AND ENVIRONMENTAL SAFETY, vol.129, pp.137-144, 2016 (SCI-Expanded)
- XVI. **Chromosomal analyses of 1510 couples who have experienced recurrent spontaneous abortions**
TUNÇ E., TANRIVERDİ N., DEMİRHAN O., SÜLEYMANOVA D., ÇETİNEL N.
REPRODUCTIVE BIOMEDICINE ONLINE, vol.32, no.4, pp.414-419, 2016 (SCI-Expanded)
- XVII. **Report of a new case with pentasomy X and novel clinical findings.**
DEMİRHAN O., TANRIVERDİ N., YILMAZ M. B., KOCATÜRK SEL S., LÜLEYAP H. Ü.
BALKAN JOURNAL OF MEDICAL GENETICS, vol.18, no.1, pp.85-92, 2015 (SCI-Expanded)

- XVIII. **Report of a new case with pentasomy X and novel clinical findings.**
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BALKAN JOURNAL OF MEDICAL GENETICS, vol.18, no.1, pp.85-92, 2015 (SCI-Expanded)
- XIX. **Microchimeric Cells, Sex Chromosome Aneuploidies and Cancer**
TAŞTEMİR KORKMAZ D., DEMİRHAN O., Abat D., Demirberk B., Tunc E., KULECİ S.
PATHOLOGY & ONCOLOGY RESEARCH, vol.21, no.4, pp.1157-1165, 2015 (SCI-Expanded)
- XX. **The effect of 900 and 1800 MHz GSM-like radiofrequency irradiation and nicotine sulfate administration on the embryonic development of *Xenopus laevis***
Boga A., EMRE M., SERTDEMİR Y., Akilhoglu K., Binokay S., DEMİRHAN O.
ECOTOXICOLOGY AND ENVIRONMENTAL SAFETY, vol.113, pp.378-390, 2015 (SCI-Expanded)
- XXI. **Gene Mutations in Chronic Kidney Disease Patients With Secondary Hyperparathyroidism and Sagliker Syndrome**
DEMİRHAN O., Arslan A., Sagliker Y., Akbal E., Ergun S., Bayraktar R., Sagliker H. S., Dogan E., Gunesacar R., Ozkaynak P. S.
JOURNAL OF RENAL NUTRITION, vol.25, no.2, pp.176-186, 2015 (SCI-Expanded)
- XXII. **REPORT OF A NEW CASE WITH PENTASOMY X AND NOVEL CLINICAL FINDINGS**
DEMİRHAN O., TANRIVERDİ N., YILMAZ M., Kocaturk-Sel S., INANDIKLIOĞLU N., Luleyap U., AKBAL E., cömertpay G., tufan t., DUR Ö.
BALKAN JOURNAL OF MEDICAL GENETICS, vol.18, no.1, pp.85-92, 2015 (SCI-Expanded)
- XXIII. **Frequency and Types of Chromosomal Abnormalities in Turkish Women with Amenorrhea**
DEMİRHAN O., TANRIVERDİ N., TUNC E., INANDIKLIOĞLU N., SULEYMANOVA D.
JOURNAL OF PEDIATRIC AND ADOLESCENT GYNECOLOGY, vol.27, no.5, pp.274-277, 2014 (SCI-Expanded)
- XXIV. **The Role Of Glucorticoid Receptor Gene (nr3c1 Gene) Polymorphism On Relapsing Of Idiopathic Nephrotic Syndrome In Children.**
DUR O., Anarat A., DEMİRHAN O., INANDIKLIOĞLU N., BAYAZIT A. K.
PEDIATRIC NEPHROLOGY, vol.29, no.9, pp.1730-1731, 2014 (SCI-Expanded)
- XXV. **Genetic alterations of chromosomes, p53 and p16 genes in low- and high-grade bladder cancer**
Abat D., DEMİRHAN O., Inandiklioglu N., Tunc E., ERDOĞAN Ş., Tastemir D., USLU İ. N., Tansug Z.
ONCOLOGY LETTERS, vol.8, no.1, pp.25-32, 2014 (SCI-Expanded)
- XXVI. **The Role Of Glucorticoid Receptor Gene (nr3c1 Gene) Polymorphism On Relapsing Of Idiopathic Nephrotic Syndrome In Children.**, vol.29, pp.1730-1731, 2014
Dur O., Anarat A., DEMİRHAN O., İNANDIKLIOĞLU N.
PEDIATRIC NEPHROLOGY,, vol.29, pp.1730-1731, 2014 (SCI-Expanded)
- XXVII. **There is no Significant Association Between Death Receptor 4 (DR4) Gene Polymorphisms and Lung Cancer in Turkish Population**
Tastemir-Korkmaz D., DEMİRHAN O., KULECİ S., HASTURK S.
PATHOLOGY & ONCOLOGY RESEARCH, vol.19, no.4, pp.779-784, 2013 (SCI-Expanded)
- XXVIII. **Polychlorinated biphenyls and organochlorine pesticides in amniotic fluids of pregnant women in south-central Turkey**
DAĞLIOĞLU N., AKÇAN R., Efeoglu P., INANDIKLIOĞLU N., GÜLMEN M. K., DEMİRHAN O.
TOXICOLOGICAL AND ENVIRONMENTAL CHEMISTRY, vol.95, no.6, pp.954-961, 2013 (SCI-Expanded)
- XXIX. **Are there fetal stem cells in the maternal brain?**
DEMİRHAN O., ÇEKİN N., Tastemir D., TUNC E., Guzel A. I., MERAL D., DEMİRBEK B.
NEURAL REGENERATION RESEARCH, vol.8, no.7, pp.593-598, 2013 (SCI-Expanded)
- XXX. **Chromosome Imbalances and Alterations in the p53 Gene in Uterine Myomas from the Same Family Members: Familial Leiomyomatosis in Turkey**
Hakverdi S., DEMİRHAN O., TUNC E., INANDIKLIOĞLU N., USLU İ. N., Gungoren A., Erdem D., Hakverdi A. U.
ASIAN PACIFIC JOURNAL OF CANCER PREVENTION, vol.14, no.2, pp.651-658, 2013 (SCI-Expanded)
- XXXI. **Connexin 26 and 30 mutations in paediatric patients with congenital, non-syndromic hearing loss treated with cochlear implantation in Mediterranean Turkey**
TARKAN Ö., SARI P., DEMİRHAN O., KIROĞLU M. M., TUNCER Ü., SÜRME LİOĞLU Ö., ÖZDEMİR S., YILMAZ M., KARA K.

- JOURNAL OF LARYNGOLOGY AND OTOLOGY, vol.127, no.1, pp.33-37, 2013 (SCI-Expanded)
- XXXII. **INHERITANCE OF A CHROMOSOME 3 AND 21 TRANSLOCATION IN THE FETUSES, WITH ONE ALSO HAVING TRISOMY 21, IN THREE PREGNANCIES IN ONE FAMILY**
PAZARBAŞI A., DEMİRHAN O., ALPTEKİN D., ÖZGÜNEN F. T., ÖZPAK L., YILMAZ M., NAZLICAN E., TANRIVERDİ N., LULEYAP U., GÜMÜRDÜLÜ D.
BALKAN JOURNAL OF MEDICAL GENETICS, vol.16, no.2, pp.91-96, 2013 (SCI-Expanded)
- XXXIII. **. Identification of chromosome abnormalities in screening of a family with manic depression and psoriasis: Predisposition to aneuploidy**
Demirhan O., Demirberk B., Çetiner S., Uslu I., Serin A.
ASIAN JOURNAL OF SOCIAL PSYCHOLOGY, vol.274, pp.1-6, 2012 (SSCI)
- XXXIV. **RAPID DETECTION OF FETAL ANEUPLOIDIES BY QUANTITATIVE FLUORESCENT-POLYMERASE CHAIN REACTION FOR PRENATAL DIAGNOSIS IN THE TURKISH POPULATION**
GÜZEL A. İ., YILMAZ M., DEMİRHAN O., PAZARBAŞI A., KOCATURK-SEL S., ERKOC M. A., INANDIKLIOĞLU N., ÖZGÜNEN F. T., SARITURK C.
BALKAN JOURNAL OF MEDICAL GENETICS, vol.15, no.1, pp.11-17, 2012 (SCI-Expanded)
- XXXV. **Chromosome Imbalances and Alterations of AURKA and MYCN Genes in Children with Neuroblastoma**
INANDIKLIOĞLU N., YILMAZ S., DEMİRHAN O., ERDOĞAN Ş., TANYELİ A.
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- XXXVI. **International Evaluation of Unrecognizably Uglifying Human Faces in Late and Severe Secondary Hyperparathyroidism in Chronic Kidney Disease. Sagliker Syndrome. A Unique Catastrophic Entity, Cytogenetic Studies for Chromosomal Abnormalities, Calcium-Sensing Receptor Gene and GNAS1 Mutations. Striking and Promising Missense Mutations on the GNAS1 Gene Exons 1, 4, 10, 4**
Yildiz I., SAGLIKER Y., Demirhan O., TUNC E., INANDIKLIOĞLU N., TAŞDEMİR D., Acharya V., ZHANG L., GOLEA O., SABRY A., et al.
JOURNAL OF RENAL NUTRITION, vol.22, no.1, pp.157-161, 2012 (SCI-Expanded)
- XXXVII. **Association of the Nramp1 gene polymorphisms and clinical forms in patients with tuberculosis**
HANTA İ., Tastemir-Korkmaz D., DEMİRHAN O., HANTA D., KULECİ S., SEYDAOĞLU G.
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- XXXVIII. **The genotoxic effect of nicotine on chromosomes of human fetal cells: The first report described as an important study**
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- XXXIX. **The Reliability of Maternal Serum Triple Test in Prenatal Diagnosis of Fetal Chromosomal Abnormalities of Pregnant Turkish Women**
DEMİRHAN O., PAZARBAŞI A., GÜZEL A. İ., Tastemir D., Yilmaz B., KASAP M., Ozgunen F. T., Evruke C., Demir C., Tunc E., et al.
GENETIC TESTING AND MOLECULAR BIOMARKERS, vol.15, no.10, pp.701-707, 2011 (SCI-Expanded)
- XL. **MTHFR Gene Polymorphisms in Bladder Cancer in the Turkish Population**
Izmirli M., INANDIKLIOĞLU N., ABAT D., ALPTEKİN D., DEMİRHAN O., TANSUG Z., BAYAZIT Y.
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- XLI. **Polymorphisms in NRAMP1 and MBL2 genes and their relations with tuberculosis in Turkish children.**
SOLĞUN H., TASTEMİR D., AKSARAY N., INAN I., Demirhan O.
Tuberkuloz ve toraks, vol.59, pp.48-53, 2011 (SCI-Expanded)
- XLII. **Association of serum sex steroid levels and bone mineral density with CYP17 and CYP19 gene polymorphisms in postmenopausal women in Turkey**
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- XLIII. **Analysis of peripheral blood T-cell subsets, natural killer cells and serum levels of cytokines in children with Down syndrome**
Çetiner S., Demirhan O., İnal T. C., Tastemir D., Sertdemir Y.
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- XLIV. **Alterations in p16 and p53 genes and chromosomal findings in patients with lung cancer: Fluorescence in situ hybridization and cytogenetic studies**
DEMİRHAN O., Tastemir D., HASTUERK S., KULECİ S., HANTA İ.
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- XLV. **CLINICAL MANIFESTATIONS OF PARTIAL TRISOMY 4p**
DEMİRHAN O., ÖZGÜNEN F. T., Tastemir D.
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- XLVI. **The Expression of Folate Sensitive Fragile Sites in Patients with Bipolar Disorder**
DEMİRHAN O., TASTEMİR D., SERTDEMİR Y.
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- XLVII. **Detection of Parental Origin and Cell Stage Errors of a Double Nondisjunction in a Fetus by QF-PCR**
Guzel A. I., DEMİRHAN O., PAZARBAŞI A., Ozgunen F. T., Kocaturk-Sel S., Tastemir D.
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- XLVIII. **PARENTAL ORIGIN AND CELL STAGE ERRORS IN X-CHROMOSOME POLYSOMY 49,XXXXY**
GÜZEL A. B., DEMİRHAN O., PAZARBAŞI A., YÜKSEL B.
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- XLIX. **THE CLINICAL EFFECTS OF ISOCHROMOSOME Xq IN KLINEFELTER SYNDROME: REPORT OF A CASE AND REVIEW OF LITERATURE**
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GENETIC COUNSELING, vol.20, no.3, pp.235-242, 2009 (SCI-Expanded)
- L. **THE EFFECT OF A DE NOVO PERICENTRIC INVERSION (10)(p11.1;q22.1) ON AGGRESSIVE BEHAVIOR AND HYPERACTIVITY**
Demirhan O., PAZARBAŞI A., TUNC E., KARAHAN D., TANRIVERDİ N., AVCI A., Tahiroglu A. Y.
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- LI. **Cerebellar hypoplasia, with quadrupedal locomotion, caused by mutations in the very low-density lipoprotein receptor gene**
Tuerkmen S., Hoffmann K., DEMİRHAN O., Aruoba D., Humphrey N., Mundlos S.
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- LII. **Different segregation of chromosomes 5 and 7 in two generations and related phenotypic findings**
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- LIII. **Correlation of clinical phenotype with a pericentric inversion of chromosome 9 and genetic counseling**
DEMİRHAN O., PAZARBAŞI A., Suleymanova-Karahan D., Tanriverdi N., KILINÇ Y.
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- LIV. **Cytogenetic effects of ethanol on chronic alcohol users**
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- LV. **PRENATAL DIAGNOSIS OF TRANSLOCATION 13;13 PATAU SYNDROME: CLINICAL FEATURES OF TWO CASES**
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- LVI. **Inheritance of pericentric inversion in chromosome 7 through the three progenies and a newborn with congenital hydronephrosis diagnosed prenatally by fetal urine sampling**
DEMİRHAN O., Ozcan K., Tastemir D., Demir C., Tunc E., Solgun H. A., Guzel A. I.
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CXXXIX. . Inheritance of a Translocation between Chromosomes 12 and 16 in a Family with Recurrent Miscarriages and a Newborn with Down Syndrome Carrying the Same Translocation

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CXLV. Bir fetustaki ikili anöploidinin parental orijini ve ayrılmama evresinin QF-PCR yöntemi ile belirlenmesi.

Güzel A. İ., DEMİRHAN O., ark. v.

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CXLVI. VIII. Ulusal Tıbbi Genetik Kongresi

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CXLVII. İnmanitib tedavisi altında kronik myelositik lösemili hastalarda gebelik.

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CXLVIII. Konjenital hidronefrozisli yeni doğan bir olgunun fetal idrar örneğinden prenatal olarak saptanan

ve üç kuşak boyunca gözlenen perisentrik inversiyon 7 inin kalıtımı

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DEMİRHAN O., ark. v.

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CLI. İlk trimester spontan düşüklerde fetal, maternal ve paternal sitogenetik incelemeler.

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CLIII. Cytogenetic Effects of Ethanol in Chronic Alcohol users

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DEMİRHAN O., ark. v.

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CLV. Kasap M, Demirhan O, Pazarbaşı A, Süleymanova-Karahan D, Tunç E, Taştemir D. Frequencies of chromosomal abnormalities at amniocentesis: over 4 years of cytogenetic analyses.

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CLXI. Chromosome aberrations in a schizophrenia population

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Supported Projects

DEMİRHAN O., Project Supported by Higher Education Institutions, Bilimsel Yayın Teşvik Desteği, 2017 - 2018

AKBABA M., ÇELİK S., DEMİRHAN O., USLU İ. N., TUNÇ E., Çetinel n., Project Supported by Higher Education Institutions, Adana ili Ceyhan ilçesi tarım çalışanlarında kromozom bozuklukları araştırması, 2017 - 2018

DEMİRHAN O., Project Supported by Higher Education Institutions, Akut Lenfoblastik Lösemili Çocuklarda VEGF-C Genlerin Ekspresyon ve Metilasyon Düzeylerindeki Değişikliklerinin İncelenmesi, 2013 - 2018

İSTİFLİ E. S., HÜSUNET M. T., İLA H. B., ÇELİK R., DEMİRHAN O., Çetinel N., Project Supported by Higher Education Institutions, Sertralinin Genotoksik Etkisinin Kültüre Edilmiş İnsan Periferal Lenfositlerine Spesifik Pantelomerik ve Pansentromerik Problar İle Karakterizasyonu, 2016 - 2017

DEMİRHAN O., ÇAĞLIYAN Ç. E., DEVECİ O. S., DEMİRTAŞ M., Çetinel N., Project Supported by Higher Education Institutions, Girişimsel Kardiyak Radyolojik İşlemlerin İnsan Kromozomları Üzerine Genotoksik Etkisi, 2015 - 2017

DEMİRHAN O., Project Supported by Higher Education Institutions, Frequency and Types of Chorosomal Abnormalities in Turkish Women with Amenorrhea, 2015 - 2016

DEMİRHAN O., Project Supported by Higher Education Institutions, Uluslararası Yayınları Özendirme Desteği (10 Adet), 2013 - 2015

DEMİRHAN O., Project Supported by Higher Education Institutions, Düşük Frekanslı Manyetik Alanların (ELF-MF) İnsan Fetal Hücre Kültürlerinde Hücre Çoğalması ve Kromozomlar Üzerine Etkileri, 2012 - 2015

DEMİRHAN O., Project Supported by Higher Education Institutions, Akut Lenfoblastik Lösemili Çocuklarda VEGF,MMP-2,MMP-9,TIMP-1 veTIMP-2 Genlerin Ekspresyon ve Metilasyon Düzeylerindeki Değişikliklerinin İncelenmesi, 2012 - 2015

Activities in Scientific Journals

The Journal of Immunology & Clinical Microbiology, Publication Committee Member, 2016 - Continues

Deutscher Wissenschaftsherold German Science Herald , Publication Committee Member, 2016 - Continues

Metrics

Publication: 379

Citation (WoS): 443

Citation (Scopus): 633

H-Index (WoS): 10

H-Index (Scopus): 13

Congress and Symposium Activities

6. Multidisipliner Kanser Arařtırma Kongresi, Attendee, Konya, Turkey, 2016
1.Uluslararası Akdeniz Bilim ve Mühendislik Kongresi (IMSEC 2016), Attendee, Adana, Turkey, 2016
The European Society of Human Genetics, Attendee, Barcelona, Spain, 2016

Awards

- DEMİRHAN O., İnsanda Dört Ayak Üzerinde Yürümenin Bilinmeyen Genetik Alt Yapısının Aydınlatılması., DOKTORCLUB AWARDS 2019, Yılın Sağlık Profesyoneli Ödülleri - Yılın Yenilikçi Sağlık Profesyoneli Finalisti, December 2019
DEMİRHAN O., 2016 Yılı Temel Kanser Arařtırmacısı Ödülü, Moleküler Kanser Arařtırma Derneđi (MOKAD), October 2016
DEMİRHAN O., Chromosomal analyses of 1510 couples who have experienced recurrent spontaneous abortions has been shortlisted as one of the four papers under consideration for the 2016, Robert G. Edwards Prize Paper Award, April 2016
DEMİRHAN O., The genotoxic effect of nicotine on chromosomes of human fetal cells: the first report described as an important study., TUBİTAK, October 2010
DEMİRHAN O., TUBİTAK Bilimsel Yayınları Teşvik Deđerlendirme Komisyon Ödülü, TUBİTAK, June 2009
DEMİRHAN O., bilimsel yayın teşvik, Çukurova Üniversitesi Tıp Fakültesi, January 2009
DEMİRHAN O., Uluslararası Bilimsel Yayın Başarı Onur Ödülü, Çukurova Üniversitesi Tıp Fakültesi, April 2008
DEMİRHAN O., Yılı En Çok Uluslar Arası Yayın Yapan Öğretim Üyesi Başarı Ödülü, Çukurova Üniversitesi Tıp Fakültesi, March 2006
DEMİRHAN O., TUBİTAK Bilimsel Yayınları Teşvik Deđerlendirme Komisyon Ödülü, TUBİTAK, June 2004
DEMİRHAN O., Yurt Dışı Bilimsel Yayın Özel Ödülü, Çukurova Üniversitesi Güçlendirme Vakfı, May 2004
DEMİRHAN O., Poster Ödülü, VI. Ulusal Genetik ve Prenatal Tanı Kongresi, April 2004
DEMİRHAN O., Yurtdışı En fazla Yayın Ödülü,, Ç.Ü. Rektörlüğü,, September 2003
DEMİRHAN O., TUBİTAK Bilimsel Yayınları Teşvik Deđerlendirme Komisyon Ödülü, TUBİTAK, June 2003
DEMİRHAN O., Bilimsel Yayın Teşvik, TUBİTAK, December 2000
DEMİRHAN O., TUBİTAK Bilimsel Yayınları Teşvik Deđerlendirme Komisyon Ödülü, TUBİTAK, June 2000
DEMİRHAN O., Güçlendirme Vakfı Ödülü, Çukurova Üniversitesi, March 2000
DEMİRHAN O., YAYIN, Ç.Ü. Rektörlüğü,, June 1998
DEMİRHAN O., Güçlendirme Vakfı Ödülü, Çukurova Üniversitesi, February 1998
DEMİRHAN O., Güçlendirme Vakfı Ödülü, Çukurova Üniversitesi, March 1996
DEMİRHAN O., TUBİTAK Bilimsel Yayınları Teşvik Deđerlendirme Komisyon Ödülü, TUBİTAK, June 1995
DEMİRHAN O., TUBİTAK Bilimsel Yayınları Teşvik Deđerlendirme Komisyon Ödülü, TUBİTAK, June 1994

Non Academic Experience

- Belçika, Antwerp Üniversitesi,
Osmangazi Üniversitesi,