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International Researcher IDs

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Education Information

Expertise In Medicine, İstanbul University, İstanbul Medical Faculty, İç Hastalıkları, Turkey 1987 - 1993

Postgraduate, İstanbul University, İstanbul Medical Faculty, Turkey 1980 - 1987

Foreign Languages

English, B2 Upper Intermediate

Certificates, Courses and Trainings

Health&Medicine, Cancer Genetics, Europan School of Medical Genetics, 1999

Dissertations

Doctorate, Parafin içinde saklanan malign melanom biyopsi örneklerinde p53 geninin DGGE ve dizi analizi; p16, retinoblastoma ve CDK4 genlerinin FISH yöntemi ile incelenmesi, İstanbul University, Health Sciences Institute, İç Hastalıkları Anabilim Dalı, 2002

Expertise In Medicine, PROPAFENON'UN VENTRİKÜLER ARİTMİLER VE SİNYAL ORTALAMALI EKG PARAMETRELERİ ÜZERİNDEKİ ETKİSİNİN ARAŞTIRILMASI, İstanbul University, İstanbul Medical Faculty, İç Hastalıkları, 1993

Research Areas

Medicine, Health Sciences, Internal Medicine Sciences, Medical Genetics

Academic Titles / Tasks

Professor, İstanbul University, İstanbul Medical Faculty, Division of Medical Sciences , 1996 - Continues

Advising Theses

- ÇEFLE K., Kronik Lenfositik Lösemide Kromozomal Aberasyonlar ve p53 Yolağındaki Gen Ekspresyonları Arasındaki İlişki, Doctorate, G.ÖZTAN(Student), Continues
- ÇEFLE K., AML' de APAF-1 Promotör Metilasyonu, Kardeş Kromatid Değişimi, Kromozomal Anomaliler ile Klinik ve Laboratuvar Parametreleri Arasındaki İlişki, Postgraduate, Ö.Özgen(Student), 2012
- ÇEFLE K., Kronik Lenfositik Lösemili Hastalarda XRCC1 (X-Ray Cross Complementing Group 1) Geninde Arg399Gln ve Arg194Trp Polimorfizmlerinin ve Kardeş Kromatid Değişimi Sıklığı ile Korelasyonlarının Araştırılması, Doctorate, N.Duman(Student), 2008

Published journal articles indexed by SCI, SSCI, and AHCI

- I. **Long-term efficacy of canakinumab in hyperimmunoglobulin D syndrome**
Ozdemir Isik O., Karadag D. T., Tekeoglu S., YAZICI A., Cefle K., ÇEFLE A.
International Journal of Rheumatic Diseases, vol.27, no.1, 2024 (SCI-Expanded)
- II. **miR-145-5p suppresses cell proliferation by targeting <i>IGF1R </i>and <i>NRAS </i>genes in multiple myeloma cells**
Kaya M., Suer İ., Ozgur E., Capik O., Karatas O. F., Ozturk Ş., Gezer U., Palanduz Ş., Cefle K.
TURKISH JOURNAL OF BIOCHEMISTRY-TURK BIYOKIMYA DERGİSİ, vol.48, pp.563-569, 2023 (SCI-Expanded)
- III. **Heme oxygenase-1 deficiency as an extremely rare cause of AA-type renal amyloidosis: Expanding the clinical features and review of the literature.**
Dirim A. B., Kalayci T., Safak S., Garayeva N., Gultekin B., Hurdogan O., Solakoglu S., Yazici H., Cefle K., Ozturk Ş., et al.
Clinical rheumatology, vol.42, no.2, pp.597-606, 2023 (SCI-Expanded)
- IV. **The effect of Anzer honey on X-ray induced genotoxicity in human lymphocytes: An in vitro study**
Bagatir G., Kaya M., Suer İ., Çefle K., Palanduz A., Palanduz Ş., Becerir H. B., Koçyiğit Avcı M., Öztürk Ş.
MICROSCOPY RESEARCH AND TECHNIQUE, vol.85, no.6, pp.2241-2250, 2022 (SCI-Expanded)
- V. **OCT-1 Expression in Patients with Chronic Myeloid Leukemia: A Comparative Analysis with Respect to Response to Imatinib Treatment**
Bozkurt Bulakçı B., Aday A., Gürtekin B., Yavuz A. S., Öztürk Ş., Çefle K., Palanduz A., Palanduz Ş.
INDIAN JOURNAL OF HEMATOLOGY AND BLOOD TRANSFUSION, vol.1, no.1, pp.1-7, 2022 (SCI-Expanded)
- VI. **Clinical Characteristics and Mutation Spectrum of Neurofibromatosis Type 1 in 27 Turkish Families**
Sharifi S., Kalayci T., Palanduz S., Ozturk S., Cefle K.
BALKAN MEDICAL JOURNAL, vol.38, no.6, pp.365-373, 2021 (SCI-Expanded)
- VII. **Skeletal and molecular findings in 51 Cleidocranial dysplasia patients from Turkey**
Berkay E. G., Elkanova L., Kalayci T., ULUDAĞ ALKAYA D., Altunoglu U., Cefle K., Mihci E., NUR B., Tasdelen E., Bayramoglu Z., et al.
AMERICAN JOURNAL OF MEDICAL GENETICS PART A, vol.185, no.8, pp.2488-2495, 2021 (SCI-Expanded)
- VIII. **Dysregulation of MS4A3 and PRDX5 Gene Expression in Multiple Myeloma Patients**
Suer İ., Aday A., Sariman M., Ayer M., Hindilerden I. Y., Ekmekci S. S., Abaci N., Palanduz Ş., Çefle K., Öztürk Ş.
UHOD-ULUSLARARASI HEMATOLOJİ-ONKOLOJİ DERGİSİ, vol.31, no.4, pp.205-213, 2021 (SCI-Expanded)
- IX. **RELATIONSHIP BETWEEN CHROMOSOMAL ABERRATIONS AND GENE EXPRESSIONS IN THE p53 PATHWAY IN CHRONIC LYMPHOCYTIC LEUKEMIA**
ÖZTAN G., Aktan M., Palanduz Ş., İŞSEVER H., ÖZTÜRK Ş., Nikerel E., Ucur A., Bagatir G., BAYRAK A. G., ÇEFLE K.
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- X. **DNA damage effects of inhalation anesthetics in human bronchoalveolar cells**
ÇUKUROVA Z., Cetingok H., Ozturk S., Gedikbasi A., HERGÜNSEL O., Ozturk D., Don B., Cefle K., Palanduz S., Ertem D. H.
MEDICINE, vol.98, no.32, 2019 (SCI-Expanded)
- XI. **Investigation of Gene Expressions of Myeloma Cells in the Bone Marrow of Multiple Myeloma Patients by Transcriptome Analysis**
Sariman M., Abaci N., Ekmekci S., Cakiris A., Pacal F., Ustek D., Ayer M., Yenerel M. N., Besisik S., Cefle K., et al.

- Balkan medical journal, vol.36, no.1, pp.23-31, 2019 (SCI-Expanded)
- XII. **Clinical features and molecular genetic analysis in a Turkish family with oral white sponge nevus**
Kurklu E., Ozturk S., Cassidy A. J., Ak G., Koray M., Cefle K., Palanduz S., Gulluoglu M., Tanyeri H., McLean W.
MEDICINA ORAL PATOLOGIA ORAL Y CIRUGIA BUCAL, no.2, 2018 (SCI-Expanded)
- XIII. **REST Final-Exon-Truncating Mutations Cause Hereditary Gingival Fibromatosis**
BAYRAM Y., WHITE J. J., Elcioglu N., CHO M. T., ZADEH N., Gedikbasi A., Palanduz S., Ozturk S., Cefle K., Kasapcopur O., et al.
AMERICAN JOURNAL OF HUMAN GENETICS, vol.101, no.1, pp.149-156, 2017 (SCI-Expanded)
- XIV. **WRN Mutation Update: Mutation Spectrum, Patient Registries, and Translational Prospects**
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HUMAN MUTATION, vol.38, no.1, pp.7-15, 2017 (SCI-Expanded)
- XV. **The frequency of C609T polymorphism in the NQO1 gene and its relation to cytogenetic abnormalities in patients with myelodysplastic syndrome.**
Bagatir G., Sirma S. Ö., Palanduz S., Ozturk S., Cefle K., Ozbek U., Yenerel M. N., Nalcaci M.
Cellular and molecular biology (Noisy-le-Grand, France), vol.62, no.7, pp.61-5, 2016 (SCI-Expanded)
- XVI. **Mutations in RAD21 Disrupt Regulation of APOB in Patients With Chronic Intestinal Pseudo-Obstruction**
BONORA E., BIANCO F., Cordeddu L., Bamshad M., Francescato L., Dowless D., STANGHELLINI V., COGLIANDRO R. F., Lindberg G., Mungan Z., et al.
GASTROENTEROLOGY, vol.148, no.4, pp.771-793, 2015 (SCI-Expanded)
- XVII. **Evaluation of micronuclear frequencies in both circulating lymphocytes and buccal epithelial cells of patients with oral lichen planus and oral lichenoid contact reactions**
Saruhanoglu A., Ergun S., Kaya M. O., Warnakulasuriya S., Erbagci M., Öztürk S., Deniz E., Ozel S., Cefle K., Palanduz S., et al.
ORAL DISEASES, vol.20, no.5, pp.521-527, 2014 (SCI-Expanded)
- XVIII. **Genotoxicity of fixation devices analyzed by the frequencies of sister chromatid exchange**
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- XIX. **A Turkish trichothiodystrophy patient with homozygous XPD mutation and genotype-phenotype relationship**
Pehlivan D., Cefle K., Raams A., Ozturk S., Baykal C., Kleijer W. J., Palanduz S., Jaspers N. G. J.
JOURNAL OF DERMATOLOGY, vol.39, no.12, pp.1016-1021, 2012 (SCI-Expanded)
- XX. **Prostaglandin transporter mutations cause pachydermoperiostosis with myelofibrosis**
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- XXI. **A novel two bases deletion in the albumin gene causes analbuminaemia in a young Turkish man**
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Clinica Chimica Acta, vol.413, pp.950-951, 2012 (SCI-Expanded)
- XXII. **A novel two bases deletion in the albumin gene causes analbuminaemia in a young Turkish man.**
CEFLE K.
CLINICA CHIMICA ACTA, vol.413, pp.950-1, 2012 (SCI-Expanded)
- XXIII. **Investigation of Arg399Gln and Arg194Trp Polymorphisms of the XRCC1 (X-Ray Cross-Complementing Group 1) Gene and Its Correlation to Sister Chromatid Exchange Frequency in Patients with Chronic Lymphocytic Leukemia**
Duman N., Aktan M., Ozturk S., Palanduz S., Cakiris A., Ustek D., Ozbek U., Nalcaci M., Cefle K.
GENETIC TESTING AND MOLECULAR BIOMARKERS, vol.16, no.4, pp.287-291, 2012 (SCI-Expanded)
- XXIV. **A novel frameshift deletion in the albumin gene causes analbuminemia in a young Turkish woman**
Dagnino M., Caridi G., Aydin Z., Ozturk S., Karaali Z., Kazancioglu R., Cefle K., Gursu M., Campagnoli M., Galliano M., et al.

- al.
Clinica Chimica Acta, vol.411, pp.1711-1715, 2010 (SCI-Expanded)
- XXV. **WRN mutations in Werner syndrome patients: genomic rearrangements, unusual intronic mutations and ethnic-specific alterations**
Friedrich K., Lee L., Leistritz D. F., Nuernberg G., Saha B., Hisama F. M., Eyman D. K., Lessel D., Nuernberg P., Li C., et al.
HUMAN GENETICS, vol.128, no.1, pp.103-111, 2010 (SCI-Expanded)
- XXVI. **NILOTINIB EFFICACY IN 21 IMATINIB-RESISTANT OR-INTOLERANT T (9;22) POSITIVE CHRONIC MYELOID LEUKEMIA PATIENTS WITH AND WITHOUT ADDITIONAL CHROMOSOMAL CHANGES**
Yavuz A. S., Elcioglu O. C., Akpinar T. S., Cosan F., Ucur A., Bayrak A., Cefle K., Oeztuerk S., Palanduz S., Yenerel M. N., et al.
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- XXVII. **A POSSIBLE DELETERIOUS EFFECT OF INCREASED SERUM COPPER ON MYOCARDIAL FUNCTION IN PATIENTS WITH DILATED CARDIOMYOPATHY AWAITING TRANSPLANTATION**
Cefle K., Ercag E., Gezertas S., Uzer A., Oeztuerk S., Cefle A., Palanduz S., Gueler K.
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- XXVIII. **TRANSPLANTASYON BEKLEYEN DİLATE KARDİYOMİOPATİLİ HASTALARDA YÜKSEK SERUM BAKIR DÜZEYİNİN MİYOKARD İŞLEVİ ÜZERİNDEKİ MUHTEMEL KÖTÜ ETKİSİ**
PALANDUZ S., ÇEFLE K.
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- XXIX. **İMATİNİBE DİRENÇLİ VEYA ENTOLERANS GÖSTEREN, KROMOZOMAL DEĞİŞİKLİKLERİ OLAN VE OLMAYAN T(9;22) POZİTİF KRONİK MYELOİD LÖSEMİLİ 21 HASTADA NİLOTİNİB'İN ETKİNLİĞİ**
PALANDUZ S., ÇEFLE K.
NOBEL MEDICUS, no.6, pp.57-62, 2010 (SCI-Expanded)
- XXX. **Cytogenetic Analysis and Examination of SOS1 Gene Mutation in a Turkish Family with Hereditary Gingival Fibromatosis**
Pehlivan D., Abe S., Ozturk S., Kayhan K., Gunduz E., Cefle K., Bayrak A. G., Ark N., Gunduz M., Palanduz S.
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- XXXI. **Micronuclear and sister chromatid exchange analyses in peripheral lymphocytes of patients with oral lichen planus - a pilot study**
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ORAL DISEASES, vol.15, no.7, pp.499-504, 2009 (SCI-Expanded)
- XXXII. **Comparison of the Cytogenetic and Molecular Analyses in the Assessment of Imatinib Response in Chronic Myelocytic Leukemia**
Palanduz S., Bayrak A., Sirma S., Vural B., Cefle K., Ucur A., Ozturk S., Yenerel M. N., Besisik S., Yavuz S., et al.
GENETIC TESTING AND MOLECULAR BIOMARKERS, vol.13, no.5, pp.599-602, 2009 (SCI-Expanded)
- XXXIII. **Left Ventricular Thickness Is Increased in Nonhypertensive Turner's Syndrome**
Sozen A. B., Cefle K., Kudat H., Ozturk S., Oflaz H., Akkaya V., Palanduz S., Demirel S., Özcan M., Goren T., et al.
ECHOCARDIOGRAPHY-A JOURNAL OF CARDIOVASCULAR ULTRASOUND AND ALLIED TECHNIQUES, vol.26, no.8, pp.943-949, 2009 (SCI-Expanded)
- XXXIV. **ALTERATIONS IN LYMPHOCYTE MEMBRANE PROTEIN CONTENT AND INCREASED LYMPHOCYTE RIGIDITY IN CATS WITH DIABETES MELLITUS**
CEFLE K.
JOURNAL OF PHYSIOLOGICAL SCIENCES, no.59, pp.505, 2009 (SCI-Expanded)
- XXXV. **The effect of parental consanguinity on the clinical and laboratory findings of rheumatoid arthritis**
Cefle K., ÇEFLE A., YAZICI A., SELEK A.
INTERNATIONAL JOURNAL OF CLINICAL PRACTICE, vol.63, no.7, pp.1056-1060, 2009 (SCI-Expanded)
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Tamer S. A., Cefle K., Kaymaz A. A., Albeniz I.
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- XXXVII. **The effects of etodolac, nimesulid and naproxen sodium on the frequency of sister chromatid exchange after enclosed third molars surgery.**
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- XXXVIII. **Atrial and ventricular arrythmogenic potential in Turner syndrome**
Sozen A. B., Cefle K., Kudat H., Ozturk S., Oflaz H., Pamukcu B., Akkaya V., Isguvan P., Palanduz S., Özcan M., et al.
PACE-PACING AND CLINICAL ELECTROPHYSIOLOGY, vol.31, no.9, pp.1140-1145, 2008 (SCI-Expanded)
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- XL. **A novel locus for syndromic chronic idiopathic intestinal pseudo-obstruction maps to chromosome 8q23-q24**
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- XLI. **The genotoxic effects in lymphocyte cultures of children treated with radiosynovectomy by using yttrium-90 citrate colloid**
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- XLII. **Definition of C282Y mutation in a hereditary hemochromatosis family from Turkey**
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- XLIV. **Initial maternal meiotic I error leading to the formation of a maternal i(2q) and a paternal i(2p) in a healthy male**
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- XLV. **Comparison of rheological parameters in patients with post hepatic and alcoholic cirrhosis**
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- XLVI. **Lens opacities in Bloom syndrome: Case report and review of the literature**
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- XLVII. **Increased sister chromatid exchange frequency in young women with breast cancer and in their first-degree relatives.**
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- XLVIII. **A case of progressive pseudorheumatoid arthropathy of 'childhood' with the diagnosis delayed to the fifth decade**
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- L. **A different approach to telomere analysis with ddPRINS in chronic lymphocytic leukemia**
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- LI. **Two siblings with distal pachydermodactyly**
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- LII. **A solitary calvarial lytic lesion with typical histopathological findings of juvenile hyaline fibromatosis.**
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- LIII. **Clinical and molecular characterization of two adults with autosomal recessive Robinow syndrome**
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- LIV. **Genotoxicity and sister chromatid exchange in patients with myelodysplastic disorders.**
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- LV. **Alterations in rheological properties and erythrocyte membrane proteins in cats with diabetes mellitus.**
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- LVII. **Peutz-Jeghers syndrome: report of 6 cases in a family and management of polyps with intraoperative endoscopy.**
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- LVIII. **A case of mandibuloacral dysplasia presenting with features of scleroderma**
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- LX. **Sister chromatid exchange and mitotic index in patients with cirrhosis related to hepatitis B and C viruses and in chronic carriers**
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- LXII. **Relationship between insulin-like growth factor-I and bone mineral density in men aged over 65 years**
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- LXIII. **Molecular diagnosis of analbuminemia: a novel mutation identified in two Amerindian and two Turkish families.**
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Metrics

Publication: 129
Citation (WoS): 690
Citation (Scopus): 795
H-Index (WoS): 14
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Congress and Symposium Activities

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11.Uluslararası Tıbbi Genetik Kongresi / Kleidokranial Displazi: Olgu Sunumu, Attendee, İstanbul, Turkey, 2014
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