Munis Dundar

List of Publications by Year in descending order

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202 papers

2,490 citations

393982 19 h-index 243296 44 g-index

235 all docs

235
docs citations

times ranked

235

3415 citing authors

#	Article	IF	CITATIONS
1	Neonatal Diabetes, Congenital Hypothyroidism, and Congenital Glaucoma Coexistence: A Case of GLIS3 Mutation. JCRPE Journal of Clinical Research in Pediatric Endocrinology, 2023, 15, 426-430.	0.4	3
2	Clinical and molecular evaluation of MEFV gene variants in the Turkish population: a study by the National Genetics Consortium. Functional and Integrative Genomics, 2022, 22, 291-315.	1.4	7
3	A teenager boy with a novel variant of Sitosterolemia presented with pancytopenia. Clinica Chimica Acta, 2022, 529, 61-66.	0.5	3
4	A novel missense mutation outside the <scp>DNAJ</scp> domain of <scp><i>DNAJC21</i></scp> is associated with <scp>Shwachman–Diamond</scp> syndrome. British Journal of Haematology, 2022, 197, .	1.2	4
5	A very rare cause of arthrogryposis multiplex congenita: a novel mutation in <i>TOR1A</i> . Journal of Pediatric Endocrinology and Metabolism, 2022, 35, 845-850.	0.4	1
6	Diagnosing Alström syndrome in a patient followed up with syndromic obesity for years. Intractable and Rare Diseases Research, 2022, 11, 84-86.	0.3	1
7	Germline landscape of BRCAs by 7-site collaborations as a BRCA consortium in Turkey. Breast, 2022, 65, 15-22.	0.9	3
8	Possible Role of the <i>RORC</i> Gene in Primary and Secondary Lymphedema: Review of the Literature and Genetic Study of Two Rare Causative Variants. Lymphatic Research and Biology, 2021, 19, 129-133.	0.5	5
9	<i>NOTCH1</i> : Review of its role in lymphatic development and study of seven families with rare pathogenic variants. Molecular Genetics & Enomic Medicine, 2021, 9, e1529.	0.6	4
10	Are new genome variants detected in SARS-CoV-2 expected considering population dynamics in viruses?. The EuroBiotech Journal, 2021, 5, 1-3.	0.5	1
11	A brief overview of global biotechnology. Biotechnology and Biotechnological Equipment, 2021, 35, S5-S14.	0.5	14
12	The Story of a Ship Journey, Malaria, and the HBB Gene IVS-II-745 Mutation: Circassian Immigration to Cyprus. Global Medical Genetics, 2021, 08, 069-071.	0.4	0
13	Current and Future Therapeutic Strategies for Limb Girdle Muscular Dystrophy Type R1: Clinical and Experimental Approaches. Pathophysiology, 2021, 28, 238-249.	1.0	4
14	Detection of mutations in CML patients resistant to tyrosine kinase inhibitor: imatinib mesylate therapy. Medical Oncology, 2021, 38, 120.	1.2	0
15	COVID-19 vaccine candidates and vaccine development platforms available worldwide. Journal of Pharmaceutical Analysis, 2021, 11, 675-682.	2.4	8
16	COVID-19 vaccines: Where do we stand?. The EuroBiotech Journal, 2021, 5, 4-7.	0.5	1
17	BRCA Variations Risk Assessment in Breast Cancers Using Different Artificial Intelligence Models. Genes, 2021, 12, 1774.	1.0	3
18	Propranolol significantly reduced DNA polymerase \hat{l}^2 expression in patients with essential tremor. Universa Medicina, 2021, 40, 207-215.	0.1	0

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19	<i>CDH5</i> , a Possible New Candidate Gene for Genetic Testing of Lymphedema. Lymphatic Research and Biology, 2021, , .	0.5	3
20	Segregation Analysis of Rare NRP1 and NRP2 Variants in Families with Lymphedema. Genes, 2020, 11, 1361.	1.0	4
21	Two rare <i>PROX1</i> variants in patients with lymphedema. Molecular Genetics & amp; Genomic Medicine, 2020, 8, e1424.	0.6	4
22	TIE1 as a Candidate Gene for Lymphatic Malformations with or without Lymphedema. International Journal of Molecular Sciences, 2020, 21, 6780.	1.8	11
23	Mutations in the ARAP3 Gene in Three Families with Primary Lymphedema Negative for Mutations in Known Lymphedema-Associated Genes. International Journal of Genomics, 2020, 2020, 1-9.	0.8	1
24	Comparing expression levels of PERIOD genes PER1, PER2 and PER3 in chronic insomnia patients and medical staff working in the night shift. Sleep Medicine, 2020, 73, 101-105.	0.8	7
25	The role of androgen receptor CAG repeat polymorphism in androgen excess disorder and idiopathic hirsutism. Journal of Endocrinological Investigation, 2020, 43, 1271-1281.	1.8	10
26	The Age Structure, Stringency Policy, Income, and Spread of Coronavirus Disease 2019: Evidence From 209 Countries. Frontiers in Psychology, 2020, 11, 632192.	1.1	21
27	Natural compounds as inhibitors of SARS-CoV-2 endocytosis: A promising approach against COVID-19. Acta Biomedica, 2020, 91, e2020008.	0.2	14
28	Pilot study for the evaluation of safety profile of a potential inhibitor of SARS-CoV-2 endocytosis. Acta Biomedica, 2020, 91, e2020009.	0.2	8
29	Comparison between American and European legislation in the therapeutical and alimentary bacteriophage usage. Acta Biomedica, 2020, 91, e2020023.	0.2	6
30	A pilot study on the preventative potential of alpha-cyclodextrin and hydroxytyrosol against SARS-CoV-2 transmission. Acta Biomedica, 2020, 91, e2020022.	0.2	14
31	Bacteriophages presence in nature and their role in the natural selection of bacterial populations. Acta Biomedica, 2020, 91, e2020024.	0.2	16
32	Reflections on Emerging Technologies in Nanomedicine. Erciyes Medical Journal, 2020, , .	0.0	0
33	Enhancer of zeste homolog 2 (EZH2) gene inhibition via 3-Deazaneplanocin A (DZNep) in human liver cells and it is relation with fibrosis. Turkish Journal of Biochemistry, 2020, 45, 737-745.	0.3	0
34	The effects of O ⁶ -methyl guanine DNA-methyl transferase promotor methylation and CpG1, CpG2, CpG3 and CpG4 methylation on treatment response and their prognostic significance in patients with glioblastoma. Balkan Journal of Medical Genetics, 2020, 23, 33-41.	0.5	3
35	Propranolol decreases DRD3 and SLC1A2 gene expression in patients with essential tremor. Universa Medicina, 2020, 39, 105-112.	0.1	1
36	Bacteriophages in food supplements obtained from natural sources. Acta Biomedica, 2020, 91, e2020025.	0.2	0

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37	Genetic testing for autonomic dysfunction or dysautonomias. Acta Biomedica, 2020, 91, e2020002.	0.2	4
38	Ethics committees for clinical experimentation at international level with a focus on Italy. Acta Biomedica, 2020, 91, e2020016.	0.2	4
39	The association of endothelin-1 levels with renal survival in polycystic kidney disease patients. Journal of Nephrology, 2019, 32, 83-91.	0.9	11
40	The molecular basis and genotype–phenotype correlations of congenital adrenal hyperplasia (CAH) in Anatolian population. Molecular Biology Reports, 2019, 46, 3677-3690.	1.0	9
41	Comprehensive genotyping of Turkish women with hirsutism. Journal of Endocrinological Investigation, 2019, 42, 1077-1087.	1.8	10
42	Current state and prospects of biotechnology in Central and Eastern European countries. Part I: Visegrad countries (CZ, H, PL, SK). Critical Reviews in Biotechnology, 2019, 39, 114-136.	5.1	10
43	Current state and prospects of biotechnology in Central and Eastern European countries. Part II: new and preaccession EU countries(CRO, RO, B&H, SRB). Critical Reviews in Biotechnology, 2019, 39, 137-155.	5.1	5
44	Future Biotechnology. The EuroBiotech Journal, 2019, 3, 53-56.	0.5	4
45	A Potential Method to Help Predict Genetic Diseases and Arrange Healthcare: Copy Number Variations Analysis. Erciyes Medical Journal, 2019, , .	0.0	0
46	Genetic background, nutrition and obesity: a review. European Review for Medical and Pharmacological Sciences, 2019, 23, 1751-1761.	0.5	17
47	Genetic testing for aortic valve stenosis. The EuroBiotech Journal, 2018, 2, 61-63.	0.5	1
48	Increased vitamin D receptor gene expression and rs11568820 and rs4516035 promoter polymorphisms in autistic disorder. Molecular Biology Reports, 2018, 45, 541-546.	1.0	16
49	Editorial. Journal of Biotechnology, 2018, 280, S1-S2.	1.9	0
50	Genetic testing for tetralogy of Fallot. The EuroBiotech Journal, 2018, 2, 71-73.	0.5	1
51	Quality assurance of genetic laboratories and the EBTNA practice certification, a simple standardization assurance system for a laboratory network. The EuroBiotech Journal, 2018, 2, 215-222.	0.5	1
52	The frequencies of Y chromosome microdeletions in infertile males. Turkish Journal of Urology, 2018, 44, 389-392.	1.3	20
53	Prof. Mariapia Viola-Magni – An Appreciation. The EuroBiotech Journal, 2018, 2, 1-1.	0.5	0
54	Genetic testing for ventricular septal defect. The EuroBiotech Journal, 2018, 2, 51-54.	0.5	0

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55	Genetic testing for Marfan syndrome. The EuroBiotech Journal, 2018, 2, 35-37.	0.5	O
56	Genetic testing for vascular Ehlers-Danlos syndrome and other variants with fragility of the middle arteries. The EuroBiotech Journal, 2018, 2, 42-44.	0.5	0
57	Genetic testing for Marfan-like disorders. The EuroBiotech Journal, 2018, 2, 38-41.	0.5	0
58	Genetic testing for atrial septal defect. The EuroBiotech Journal, 2018, 2, 45-47.	0.5	0
59	THE ASSOCIATION OF BRAIN-DERIVED NEUROTROPHIC FACTOR GENE POLYMORPHISM WITH OBSTRUCTIVE SLEEP APNEA SYNDROME and OBESITY. , $2018, , .$		0
60	Genetic testing for Ebstein anomaly. The EuroBiotech Journal, 2018, 2, 55-57.	0.5	2
61	Genetic testing for cerebral cavernous malformations. The EuroBiotech Journal, 2018, 2, 83-85.	0.5	0
62	Genetic testing for atrioventricular septal defect. The EuroBiotech Journal, 2018, 2, 48-50.	0.5	0
63	Genetic testing for bicuspid aortic valve. The EuroBiotech Journal, 2018, 2, 67-70.	0.5	0
64	Genetic testing for coarctation of aorta. The EuroBiotech Journal, 2018, 2, 64-66.	0.5	0
65	Genetic testing for pulmonary stenosis. The EuroBiotech Journal, 2018, 2, 58-60.	0.5	0
66	Genetic testing for hereditary hemorrhagic telangiectasia. The EuroBiotech Journal, 2018, 2, 32-34.	0.5	0
67	The effect of parental 5,10-methylenetetrahydrofolate reductase 677C/T and 1298A/C gene polymorphisms on response to single-dose methotrexate in tubal ectopic pregnancy. Journal of Maternal-Fetal and Neonatal Medicine, 2017, 30, 1232-1237.	0.7	0
68	Diagnosis of intracranial calcification and hemorrhage in pediatric patients: Comparison of quantitative susceptibility mapping and phase images of susceptibility-weighted imaging. Diagnostic and Interventional Imaging, 2017, 98, 707-714.	1.8	27
69	Prenatal diagnosis of a foetus with partial monosomy 4p and partial trisomy 13q. Journal of Biotechnology, 2017, 256, S76.	1.9	0
70	The effect of CYP2C19 * 2 polymorphism on clopidogrel resistance in COPD patients. Journal of Biotechnology, 2017, 256, S80.	1.9	0
71	Frequency of chromosome variants in families with recurrent pregnancy loss and statistical analysis of infertility. Journal of Biotechnology, 2017, 256, S76.	1.9	0
72	Developments in biotechnology. Journal of Biotechnology, 2017, 256, S7.	1.9	1

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73	Neurological Manifestations in Familial Mediterranean Fever: Results of 22 Children from a Reference Center in Kayseri, an Urban Area in Central Anatolia, Turkey. Neuropediatrics, 2017, 48, 079-085.	0.3	10
74	Prenatal diagnosis of upper extremity malformations with ultrasonography: Diagnostic features and perinatal outcome. Journal of Clinical Ultrasound, 2017, 45, 267-276.	0.4	3
75	Genetic testing for congenital stationary night blindness. The EuroBiotech Journal, 2017, 1, 38-40.	0.5	О
76	Research of genetic bases of hereditary non-syndromic hearing loss. Turk Pediatri Arsivi, 2017, 52, 122-132.	0.9	7
77	Genetic testing in translational ophthalmology. The EuroBiotech Journal, 2017, 1, 1-5.	0.5	0
78	Genetic testing for Norrie disease. The EuroBiotech Journal, 2017, 1, 77-79.	0.5	0
79	Genetic testing for Usher syndrome. The EuroBiotech Journal, 2017, 1, 108-110.	0.5	1
80	Pharmacologically active fractions of Sideritis spp. and their use in inherited eye diseases. The EuroBiotech Journal, 2017, 1, 6-10.	0.5	5
81	Genetic tests for low- and middle-income countries: a literature review. Genetics and Molecular Research, 2017, 16, .	0.3	12
82	Genetic testing for Bietti crystalline dystrophy. The EuroBiotech Journal, 2017, 1, 20-22.	0.5	1
83	Genetic testing for familial exudative vitreoretinopathy. The EuroBiotech Journal, 2017, 1, 51-53.	0.5	2
84	Genetic testing for pattern dystrophies. The EuroBiotech Journal, 2017, 1, 86-88.	0.5	2
85	Genetic testing for Stargardt macular dystrophy. The EuroBiotech Journal, 2017, 1, 105-107.	0.5	1
86	Advances in biotechnology: Genomics and genome editing. The EuroBiotech Journal, 2017, 1, 2-9.	0.5	1
87	Genetic testing for gyrate atrophy of the choroid and retina. The EuroBiotech Journal, 2017, 1, 54-56.	0.5	2
88	Genetic testing for infantile nystagmus. The EuroBiotech Journal, 2017, 1, 57-59.	0.5	0
89	Genetic testing for retinitis punctata albescens/fundus albipunctatus. The EuroBiotech Journal, 2017, 1, 96-98.	0.5	0
90	Genetic testing for color vision deficiency. The EuroBiotech Journal, 2017, 1, 32-34.	0.5	1

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91	Genetic testing for X-linked juvenile retinoschisis. The EuroBiotech Journal, 2017, 1, 111-113.	0.5	O
92	Genetic testing for central areolar choroidal dystrophy. The EuroBiotech Journal, 2017, 1, 23-25.	0.5	3
93	Genetic testing for Sorsby's fundus dystrophy. The EuroBiotech Journal, 2017, 1, 102-104.	0.5	0
94	Genetic testing for non syndromic retinitis pigmentosa. The EuroBiotech Journal, 2017, 1, 92-95.	0.5	1
95	Genetic testing for Bardet-Biedl syndrome. The EuroBiotech Journal, 2017, 1, 14-16.	0.5	0
96	Genetic testing for inherited eye misalignment. The EuroBiotech Journal, 2017, 1, 60-62.	0.5	0
97	Genetic testing for corneal dystrophies and other corneal Mendelian diseases. The EuroBiotech Journal, 2017, 1, 41-44.	0.5	0
98	Genetic testing for optic atrophy. The EuroBiotech Journal, 2017, 1, 83-85.	0.5	0
99	Genetic testing for Mendelian myopia. The EuroBiotech Journal, 2017, 1, 74-76.	0.5	0
100	Genetic testing for Refsum disease. The EuroBiotech Journal, 2017, 1, 89-91.	0.5	1
101	Genetic testing for achromatopsia. The EuroBiotech Journal, 2017, 1, 11-13.	0.5	0
102	Genetic testing for Mendelian cataract. The EuroBiotech Journal, 2017, 1, 66-69.	0.5	0
103	Genetic testing for Leber congenital amaurosis. The EuroBiotech Journal, 2017, 1, 63-65.	0.5	0
104	Genetic testing for Mendelian glaucoma. The EuroBiotech Journal, 2017, 1, 70-73.	0.5	0
105	Genetic testing for Best vitelliform macular dystrophy. The EuroBiotech Journal, 2017, 1, 17-19.	0.5	0
106	Genetic testing for enhanced S-cone syndrome. The EuroBiotech Journal, 2017, 1, 48-50.	0.5	0
107	Genetic testing for Senior-Loken syndrome. The EuroBiotech Journal, 2017, 1, 99-101.	0.5	0
108	Genetic testing for cone rod dystrophies. The EuroBiotech Journal, 2017, 1, 35-37.	0.5	0

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109	Genetic testing for ocular coloboma. The EuroBiotech Journal, 2017, 1, 29-31.	0.5	4
110	Genetic testing for choroideremia. The EuroBiotech Journal, 2017, 1, 26-28.	0.5	0
111	Genetic testing for ocular albinism and oculocutaneous albinism. The EuroBiotech Journal, 2017, 1, 80-82.	0.5	0
112	Genetic testing for Doyne honeycomb retinal dystrophy. The EuroBiotech Journal, 2017, 1, 45-47.	0.5	0
113	Evaluation of aortic intima-media thickness in newborns with Down syndrome. Advances in Clinical and Experimental Medicine, 2017, 26, 1253-1256.	0.6	0
114	Expression of Ghrelin and GHSR-1a in Long Term Diabetic Rat's Kidney. Brazilian Archives of Biology and Technology, 2016, 59, .	0.5	1
115	A novel nonsense mutation in GALNS gene in family with MPS4A diagnosed child. Journal of Biotechnology, 2016, 231, S108.	1.9	0
116	A case of XYY male patient with micropenis. Journal of Biotechnology, 2016, 231, S109.	1.9	0
117	Genetic expressions of thrombophilic factors in patients with Sheehan's syndrome. Gynecological Endocrinology, 2016, 32, 908-911.	0.7	3
118	Perspectives of biotechnology. Journal of Biotechnology, 2016, 231, S4.	1.9	0
119	Editorial. Journal of Biotechnology, 2016, 231, S1-S3.	1.9	0
120	The Association of Brain-Derived Neurotrophic Factor Gene Polymorphism with Obstructive Sleep Apnea Syndrome and Obesity. Lung, 2016, 194, 839-846.	1.4	5
121	Ameliorative effects of pentoxifylline on NOS induced by diabetes in rat kidney. Renal Failure, 2016, 38, 605-613.	0.8	12
122	Nitric oxide synthase in diabetic rat testicular tissue and the effects of pentoxifylline therapy. Systems Biology in Reproductive Medicine, 2016, 62, 22-30.	1.0	19
123	Genetic Disorders of Pituitary Development in Patients with Sheehan'S Syndrome. Acta Endocrinologica, 2016, 12, 413-417.	0.1	1
124	Clinical Characteristics of Cases with Spinal Muscular Atrophy. Guncel Pediatri, 2016, 14, 18-22.	0.1	2
125	Genetic background of supernumerary teeth. European Journal of Dentistry, 2015, 09, 153-158.	0.8	54
126	A Glutamine Repeat Variant of the RUNX2 Gene Causes Cleidocranial Dysplasia. Molecular Syndromology, 2015, 6, 50-53.	0.3	15

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127	Innovations in biotechnology. Journal of Biotechnology, 2015, 208, S5.	1.9	O
128	A case of SRY positive 46, XX male with speaking disorder. Journal of Biotechnology, 2015, 208, S85.	1.9	1
129	Is idiopathic hirsutism (IH) really idiopathic? mRNA expressions of skin steroidogenic enzymes in women with IH. European Journal of Endocrinology, 2015, 173, 447-454.	1.9	9
130	Genotoxic Effects of some Antituberculosis Drugs and Mixtures in Rats. Drug Research, 2015, 65, 219-222.	0.7	7
131	The Effects of Long-Term Diabetes on Ghrelin Expression in Rat Stomachs. Advances in Clinical and Experimental Medicine, 2015, 24, 401-407.	0.6	4
132	The effects of streptozotocin-induced diabetes on ghrelin expression in rat testis: biochemical and immunohistochemical study. Folia Histochemica Et Cytobiologica, 2015, 53, 26-34.	0.6	12
133	Autozygosity in a Turkish family with scoliosis, blindness, and arachnodactyly syndrome. Annals of Saudi Medicine, 2015, 35, 462-467.	0.5	0
134	A c.1244G A (p.Arg415Gln) mutation in SH3BP2 gene causes cherubism in a Turkish family: Report of a family with review of the literature. Medicina Oral, Patologia Oral Y Cirugia Bucal, 2014, 19, e340-e344.	0.7	5
135	Etiopathogenesis of Sheehan's Syndrome: Roles of Coagulation Factors and TNF-Alpha. International Journal of Endocrinology, 2014, 2014, 1-6.	0.6	7
136	Circulating microRNAs in patients with non-alcoholic fatty liver disease. World Journal of Hepatology, 2014, 6, 613.	0.8	67
137	Is there relation between COL4A1/A2 mutations and antenatally detected fetal intraventricular hemorrhage?. Child's Nervous System, 2014, 30, 419-424.	0.6	12
138	Loss of dermatan sulfate epimerase (DSE) function results in musculocontractural Ehlers–Danlos syndrome. Human Molecular Genetics, 2013, 22, 3761-3772.	1.4	78
139	The role of TNF-α and PAI-1 gene polymorphisms in familial Mediterranean fever. Modern Rheumatology, 2013, 23, 140-145.	0.9	11
140	Analysing the role of MDM2 SNP309 in patients with glioblastoma multiforme. Current Opinion in Biotechnology, 2013, 24, S98.	3.3	0
141	Atypical presentation and a novel mutation in ALMS1: implications for clinical and molecular diagnostic strategies for Alström syndrome. Clinical Genetics, 2013, 83, 96-98.	1.0	9
142	A Novel <i>COL4A3</i> Mutation Causes Autosomal-Recessive Alport Syndrome in a Large Turkish Family. Genetic Testing and Molecular Biomarkers, 2013, 17, 260-264.	0.3	10
143	Progress towards the â€~Golden Age' of biotechnology. Current Opinion in Biotechnology, 2013, 24, S6-S13.	3.3	32
144	The role of TNF- \hat{l}_{\pm} and PAI-1 gene polymorphisms in familial Mediterranean fever. Modern Rheumatology, 2013, 23, 140-145.	0.9	9

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145	Evaluation of the Results of Cases Prenatally Diagnosed as VSD. Erciyes Tip Dergisi, 2012, 34, 111-115.	0.1	1
146	General Report & Description of the European Association for Predictive, Preventive and Personalised Medicine 2012: White Paper of the European Association for Predictive, Preventive and Personalised Medicine. EPMA Journal, 2012, 3, 14.	3.3	218
147	A new syndrome of microtia with unilateral renal agenesis and short stature. American Journal of Medical Genetics, Part A, 2012, 158A, 1837-1840.	0.7	1
148	A molecular analysis of familial Mediterranean fever disease in a cohort of Turkish patients. Annals of Saudi Medicine, 2012, 32, 343-348.	0.5	9
149	Overview of the Healthcare System in Turkey. Advances in Predictive, Preventive and Personalised Medicine, 2012, , 167-187.	0.6	0
150	Idiopathic hirsutism: local and peripheral expression of aromatase (CYP19A) and 5α-reductase genes (SRD5A1 and SRD5A2). Fertility and Sterility, 2011, 96, 479-482.	0.5	11
151	Common Familial Mediterranean Fever gene mutations in a Turkish cohort. Molecular Biology Reports, 2011, 38, 5065-5069.	1.0	56
152	Expression of Biologically Active Human Interferon Gamma in the Milk of Transgenic Mice Under the Control of the Murine Whey Acidic Protein Gene Promoter. Biochemical Genetics, 2011, 49, 251-257.	0.8	11
153	The increasing importance of Medical Genetics in Turkey. Current Opinion in Biotechnology, 2011, 22, S42-S43.	3.3	O
154	Prediction, prevention and personalisation of medication for the prenatal period: genetic prenatal tests for both rare and common diseases. EPMA Journal, 2011, 2, 181-195.	3.3	4
155	Loss of dermatanâ€4â€sulfotransferase 1 (D4ST1/ <i>CHST14</i>) function represents the first dermatan sulfate biosynthesis defect, "dermatan sulfateâ€deficient adducted thumb–clubfoot syndromeâ€. Human Mutation, 2011, 32, 484-485.	1.1	15
156	Prenatally detected de novo 46, XX, $t(21;21)(p12;p12)$ at chorionic villus sampling. Current Opinion in Biotechnology, 2011, 22, S107.	3.3	0
157	Biotechnology worldwide and the  European Biotechnology Thematic Network' Association (EBTNA). Current Opinion in Biotechnology, 2011, 22, S7-S14.	3.3	15
158	Current State of Biotechnology in Turkey. Current Opinion in Biotechnology, 2011, 22, S3-S6.	3.3	6
159	Healthcare in overview of Turkey. EPMA Journal, 2010, 1, 587-594.	3.3	8
160	Unbalanced 3;22 translocation with 22q11 and 3p deletion syndrome. American Journal of Medical Genetics, Part A, 2010, 152A, 2791-2795.	0.7	10
161	Maternal uniparental isodisomy is responsible for serious molybdenum cofactor deficiency. Developmental Medicine and Child Neurology, 2010, 52, 868-872.	1.1	15
162	The prevalence of non-classic adrenal hyperplasia among Turkish women with hyperandrogenism. Gynecological Endocrinology, 2010, 26, 139-143.	0.7	24

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163	Cytogenetic results of patients with infertility in middle anatolia, Turkey: do heterochromatin polymorphisms affect fertility?. Journal of Reproduction and Infertility, 2010, 11, 179-81.	1.0	10
164	The Frequency of CYP 21 Gene Mutations in Turkish Women with Hyperandrogenism. Experimental and Clinical Endocrinology and Diabetes, 2009, 117, 205-208.	0.6	10
165	Detection of p16 promotor hypermethylation in "Maras powder―and tobacco users. Cancer Epidemiology, 2009, 33, 47-50.	0.8	11
166	Loss of Dermatan-4-Sulfotransferase 1 Function Results in Adducted Thumb-Clubfoot Syndrome. American Journal of Human Genetics, 2009, 85, 873-882.	2.6	134
167	Frank-ter Haar syndrome with unusual clinical features. European Journal of Medical Genetics, 2009, 52, 247-249.	0.7	9
168	Inherited diseases and syndromes leading to aortic aneurysms and dissections. European Journal of Cardio-thoracic Surgery, 2009, 35, 931-940.	0.6	42
169	Lack of association between the Glu298Asp polymorphism of endothelial nitric oxide synthase and slow coronary flow in the Turkish population. Canadian Journal of Cardiology, 2009, 25, e69-e72.	0.8	10
170	The Deletion Polymorphism of the Angiotensin-Converting Enzyme Gene Is Associated with Acute Aortic Dissection. Tohoku Journal of Experimental Medicine, 2009, 219, 33-37.	0.5	13
171	Genotype–phenotype correlation in children with familial Mediterranean fever in a Turkish population. Pediatrics International, 2008, 50, 208-212.	0.2	64
172	An autosomal recessive adducted thumb-club foot syndrome observed in Turkish cousins. Clinical Genetics, 2008, 51, 61-64.	1.0	42
173	The Effect of Maras Powder on DNA Methylation and Micronucleus Formation in Human Buccal Tissue. Journal of Toxicology and Environmental Health - Part A: Current Issues, 2008, 71, 396-404.	1.1	10
174	Apolipoprotein E3/E3 Genotype Decreases the Risk of Pituitary Dysfunction after Traumatic Brain Injury due to Various Causes: Preliminary Data. Journal of Neurotrauma, 2008, 25, 1071-1077.	1.7	71
175	ICR1 Epimutations in 11p15 are Restricted to Patients with Silver-Russell Syndrome Features. Journal of Pediatric Endocrinology and Metabolism, 2008, 21, 59-62.	0.4	8
176	Megarbane syndrome. Indian Journal of Human Genetics, 2008, 14, 27.	0.7	0
177	Holt-Oram syndrome in two generations with translocation $t(9;15)(p12;q11.2)$. Annals of Saudi Medicine, 2008, 28, 209.	0.5	1
178	Holt-Oram syndrome in two generations with translocation $t(9;15)(p12;q11.2)$. Annals of Saudi Medicine, 2008, 28, 209-212.	0.5	1
179	Can the classical euchromatic variants of $9q12/qh+$ cause recurrent abortions?. Genetic Counseling, 2008, 19, 281-6.	0.1	3
180	Prenatal Diagnosis of a Fetus with Partial Trisomy 7p. Fetal Diagnosis and Therapy, 2007, 22, 229-232.	0.6	10

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181	Sexing Greater Flamingo Chicks from Feather Bulb DNA. Waterbirds, 2007, 30, 450-453.	0.2	15
182	Sacrococcygeal teratoma in a fetus with prenatally diagnosed partial trisomy 10q (10q24.3â†'qter) and partial monosomy 17p (p13.3â†'pter). Prenatal Diagnosis, 2007, 27, 365-368.	1.1	19
183	How the I1307K adenomatous polyposis coli gene variant contributes in the assessment of risk of colorectal cancer, but not stomach cancer, in a Turkish population. Cancer Genetics and Cytogenetics, 2007, 177, 95-97.	1.0	9
184	Adrenal axis functions in patients with familial Mediterranean fever. Clinical Rheumatology, 2006, 25, 458-461.	1.0	9
185	Familial Mediterranean Fever (FMF) in Turkey. Medicine (United States), 2005, 84, 1-11.	0.4	651
186	5,10-Methylenetetrahydrofolate reductase C677T gene polymorphism in Behcet's patients with or without ocular involvement. British Journal of Ophthalmology, 2005, 89, 1634-1637.	2.1	26
187	Isolated congenital anonychia cases with coincident chromosomal fragility. Annales De Génétique, 2004, 47, 381-386.	0.4	3
188	A Turner patient with a 45,X,t(1;2) (q41;p11.2) karyotype. Annales De Génétique, 2002, 45, 181-183.	0.4	3
189	A family with two different chromosomal translocations. Annales De Génétique, 2002, 45, 185-187.	0.4	4
190	A case with Waardenburg syndrome presenting with two separate translocations ??? one reciprocal and one complex. Clinical Dysmorphology, 2001, 10, 65-66.	0.1	0
191	A case with adducted thumb and club foot syndrome. Clinical Dysmorphology, 2001, 10, 291-293.	0.1	27
192	A case of ambiguous genitalia presenting with a 45,X/46,Xr(Y)(p11.2;q11.23)/47,X,idic(Y)(p11.2),idic(Y)(p11.2) karyotype. Annales De GÃ@nÃ@tique, 2001, 44, 5-8.	0.4	7
193	A novel acropectoral syndrome maps to chromosome 7q36. Journal of Medical Genetics, 2001, 38, 304-309.	1.5	21
194	Patient with Weismann-Netter and Stuhl (Toxopachyosteosis) Syndrome with Communicant Hydrocephalus and Arachnoid Cyst. Journal of Pediatric Endocrinology and Metabolism, 2000, 13, 211-5.	0.4	4
195	Female-to-male transsexual with 47,XXX karyotype. Biological Psychiatry, 2000, 48, 1116-1117.	0.7	18
196	Polycystic kidney disease, biliary dysgenesis in a patient with Larsen's syndrome. Clinical Genetics, 1997, 51, 408-411.	1.0	4
197	Congenital alacrima in a patient with G (Opitz Frias) syndrome. Human Genetics, 1996, 97, 540-542.	1.8	2
198	Congenital alacrima in a patient with G (Opitz Frias) syndrome. Human Genetics, 1996, 97, 540-542.	1.8	0

#	Article	IF	CITATIONS
199	Scottish frequency of the common G985 mutation in the medium-chain acyl-CoA dehydrogenase (MCAD) gene and the role of MCAD deficiency in sudden infant death syndrome (SIDS). Journal of Inherited Metabolic Disease, 1993, 16, 991-993.	1.7	22
200	Anadolu'daki Tıbbın DoÄŸuÅŸu, DÃ⅓nyadaki İlk Tıp Okulu Olarak: Gevher Nesibe Tıp Medresesi v Bilimname: Düşünce Platformu, 0, , 79-103.	e DarüÅ	\ΫÅΫ́jfası.
201	Evaluation of Utilizing the Distinct Genes as Predictive Biomarkers in Late-Onset Alzheimer's Disease. Global Medical Genetics, 0, , .	0.4	O
202	INVESTIGATION OF CDKL5 GENE MUTATIONS IN AUTISTIC PATIENTS ACCOMPANIED WITH INTRACTABLE SEIZURES, AUTISTIC DISORDER AND SEIZURE IN INFANCY AND EARLY CHILDHOOD. Cumhuriyet Medical Journal, $0, , .$	0.1	0