Jörg Schmidtke

List of Publications by Year in descending order

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Version: 2024-02-01

623734 580821 32 658 14 25 citations g-index h-index papers 36 36 36 1220 docs citations times ranked citing authors all docs

#	Article	IF	CITATIONS
1	Natal dispersal in rhesus macaques is related to serotonin transporter gene promoter variation. Behavior Genetics, 2000, 30, 295-301.	2.1	91
2	Genetic studies on the Cayo Santiago rhesus macaques: A review of 40 years of research. American Journal of Primatology, 2016, 78, 44-62.	1.7	80
3	The current role of next-generation DNA sequencing in routine care of patients with hereditary cardiovascular conditions: a viewpoint paper of the European Society of Cardiology working group on myocardial and pericardial diseases and members of the European Society of Human Genetics. European Heart Journal. 2015. 36. 1367-1370.	2.2	75
4	Observational Cohort Study of Ventricular Arrhythmia in Adults with Marfan Syndrome Caused by FBN1 Mutations. PLoS ONE, 2013, 8, e81281.	2.5	45
5	Detection of mutations in the COL4A5 gene by SSCP in X-linked Alport syndrome. Human Mutation, 2001, 18, 141-148.	2.5	38
6	New EuroGentest/ESHG guidelines and a new clinical utility gene card format for NGS-based testing. European Journal of Human Genetics, 2016, 24, 1-1.	2.8	37
7	The main pulmonary artery in adults: a controlled multicenter study with assessment of echocardiographic reference values, and the frequency of dilatation and aneurysm in Marfan syndrome. Orphanet Journal of Rare Diseases, 2014, 9, 203.	2.7	34
8	Clinical utility gene card for: Hereditary thoracic aortic aneurysm and dissection including next-generation sequencing-based approaches. European Journal of Human Genetics, 2016, 24, 146-150.	2.8	28
9	Points to consider for prioritizing clinical genetic testing services: a European consensus process oriented at accountability for reasonableness. European Journal of Human Genetics, 2015, 23, 729-735.	2.8	26
10	A murine TSPY. Chromosome Research, 1998, 6, 35-40.	2.2	24
11	Criteria for fairly allocating scarce health-care resources to genetic tests: which matter most?. European Journal of Human Genetics, 2014, 22, 25-31.	2.8	23
12	DNA-based Genetic Testing Is Rising Steeply in a National Health Care System with Open Access to Services: A Survey of Genetic Test Use in Germany, 1996–2002. Genetic Testing and Molecular Biomarkers, 2005, 9, 80-84.	1.7	20
13	Value judgments for priority setting criteria in genetic testing: A discrete choice experiment. Health Policy, 2015, 119, 164-173.	3.0	17
14	Adopting Quality Criteria for Websites Providing Medical Information About Rare Diseases. Interactive Journal of Medical Research, 2016, 5, e24.	1.4	17
15	The EuroGentest Clinical Utility Gene Cards. European Journal of Human Genetics, 2010, 18, 1071-1071.	2.8	13
16	NCAM2 deletion in a boy with macrocephaly and autism: Cause, association or predisposition?. European Journal of Medical Genetics, 2016, 59, 493-498.	1.3	13
17	DNA sequence polymorphisms in genes involved in the regulation of dopamine and serotonin metabolism in rhesus macaques. Electrophoresis, 1999, 20, 1771-1777.	2.4	11
18	Education in Medical Genetics for Physicians: Germany. Public Health Genomics, 2006, 9, 235-239.	1.0	10

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19	The future of Clinical Utility Gene Cards in the context of next-generation sequencing diagnostic panels. European Journal of Human Genetics, 2014, 22, 1247-1247.	2.8	9
20	Validation of a scale for assessing attitudes towards outcomes of genetic cancer testing among primary care providers and breast specialists. PLoS ONE, 2017, 12, e0178447.	2.5	9
21	Contentious ethical issues in community genetics: let's talk about them. Journal of Community Genetics, 2020, 11, 5-6.	1.2	8
22	The Decision Theory of Paternity Disputes: Optimization Considerations Applied to Multilocus DNA Fingerprinting. Journal of Forensic Sciences, 1992, 37, 1525-1533.	1.6	8
23	General Practitioners and Breast Surgeons in France, Germany, Netherlands and the UK show variable breast cancer risk communication profiles. BMC Cancer, 2015, 15, 243.	2.6	6
24	Transgenic Mouse Studies to Understand the Regulation, Expression and Function of the Testis-Specific Protein Y-Encoded (TSPY) Gene. Genes, 2010, 1, 244-262.	2.4	5
25	Variants in TSPYL1 are not associated with sudden infant death syndrome in a cohort of deceased infants from Switzerland. Molecular and Cellular Probes, 2015, 29, 31-34.	2.1	5
26	A new decade of community genetics: old and new challenges. Journal of Community Genetics, 2020, 11, 1-3.	1.2	2
27	Rare diseases and sports: A pilot project to improve physical activity in patients with mucopolysaccharidosis. Translational Sports Medicine, 2018, 1, 184-190.	1.1	1
28	Conceptualization and Implementation of the Central Information Portal on Rare Diseases: Protocol for a Qualitative Study. JMIR Research Protocols, 2018, 7, e112.	1.0	1
29	Who owns the human genome? Ethical and legal aspects. Journal of Pharmacy and Pharmacology, 1992, 44, 205-10.	2.4	1
30	PanelDesign: Integrating Epidemiological Information into the Design of Diagnostic NGS Gene Panels. Genes, 2022, 13, 684.	2.4	1
31	Upcoming special issues in the Journal of Community Genetics. Journal of Community Genetics, 2012, 3, 55-55.	1.2	0
32	NGS-Based genetic testing for heritable cardiovascular diseases. Specific requirements for obtaining informed consent. Molecular and Cellular Probes, 2019, 45, 70-78.	2.1	0