## Sebastien Jacquemont

List of Publications by Year in descending order

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151 papers

11,938 citations

52 h-index 29081 104 g-index

175 all docs

175 docs citations

175 times ranked

12786 citing authors

#	Article	IF	CITATIONS
1	Fragile X Premutation Tremor/Ataxia Syndrome: Molecular, Clinical, and Neuroimaging Correlates. American Journal of Human Genetics, 2003, 72, 869-878.	2.6	720
2	Penetrance of the Fragile X–Associated Tremor/Ataxia Syndrome in a Premutation Carrier Population. JAMA - Journal of the American Medical Association, 2004, 291, 460.	3.8	571
3	A new highly penetrant form of obesity due to deletions on chromosome 16p11.2. Nature, 2010, 463, 671-675.	13.7	476
4	Neuronal intranuclear inclusions in a new cerebellar tremor/ataxia syndrome among fragile X carriers. Brain, 2002, 125, 1760-1771.	3.7	474
5	A Higher Mutational Burden in Females Supports a "Female Protective Model―in Neurodevelopmental Disorders. American Journal of Human Genetics, 2014, 94, 415-425.	2.6	457
6	Mirror extreme BMI phenotypes associated with gene dosage at the chromosome 16p11.2 locus. Nature, 2011, 478, 97-102.	13.7	394
7	KCTD13 is a major driver of mirrored neuroanatomical phenotypes of the 16p11.2 copy number variant. Nature, 2012, 485, 363-367.	13.7	363
8	Epigenetic Modification of the <i>FMR1</i> Gene in Fragile X Syndrome Is Associated with Differential Response to the mGluR5 Antagonist AFQ056. Science Translational Medicine, 2011, 3, 64ra1.	5.8	344
9	Fragile-X–Associated Tremor/Ataxia Syndrome (FXTAS) in Females with the FMR1 Premutation. American Journal of Human Genetics, 2004, 74, 1051-1056.	2.6	320
10	Fragile Xâ€essociated tremor/ataxia syndrome: Clinical features, genetics, and testing guidelines. Movement Disorders, 2007, 22, 2018-2030.	2.2	306
11	Fragile X premutation carriers: characteristic MR imaging findings of adult male patients with progressive cerebellar and cognitive dysfunction. American Journal of Neuroradiology, 2002, 23, 1757-66.	1.2	272
12	A 600â€kb deletion syndrome at 16p11.2 leads to energy imbalance and neuropsychiatric disorders. Journal of Medical Genetics, 2012, 49, 660-668.	1.5	251
13	Drug development for neurodevelopmental disorders: lessons learned from fragile X syndrome. Nature Reviews Drug Discovery, 2018, 17, 280-299.	21.5	247
14	Mutations in CTC1, encoding conserved telomere maintenance component 1, cause Coats plus. Nature Genetics, 2012, 44, 338-342.	9.4	234
15	Fragile-X syndrome and fragile X-associated tremor/ataxia syndrome: two faces of FMR1. Lancet Neurology, The, 2007, 6, 45-55.	4.9	212
16	Mavoglurant in fragile X syndrome: Results of two randomized, double-blind, placebo-controlled trials. Science Translational Medicine, 2016, 8, 321ra5.	5.8	210
17	Defining the Effect of the 16p11.2 Duplication on Cognition, Behavior, and Medical Comorbidities. JAMA Psychiatry, 2016, 73, 20.	6.0	195
18	Truncating mutations in the last exon of NOTCH2 cause a rare skeletal disorder with osteoporosis. Nature Genetics, 2011, 43, 306-308.	9.4	181

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19	Detection of genomic imbalances by array based comparative genomic hybridisation in fetuses with multiple malformations. Journal of Medical Genetics, 2005, 42, 121-128.	1.5	174
20	The Fragile X Premutation Presenting as Essential Tremor. Archives of Neurology, 2003, 60, 117.	4.9	162
21	The 16p11.2 locus modulates brain structures common to autism, schizophrenia and obesity. Molecular Psychiatry, 2015, 20, 140-147.	4.1	160
22	Fourteen new cases contribute to the characterization of the 7q11.23 microduplication syndrome. European Journal of Medical Genetics, 2009, 52, 94-100.	0.7	157
23	The Efficacy of Melatonin for Sleep Problems in Children with Autism, Fragile X Syndrome, or Autism and Fragile X Syndrome. Journal of Clinical Sleep Medicine, 2009, 05, 145-150.	1.4	149
24	Copy Number Variations and Cognitive Phenotypes in Unselected Populations. JAMA - Journal of the American Medical Association, 2015, 313, 2044.	3.8	143
25	Progression of tremor and ataxia in male carriers of the FMR1 premutation. Movement Disorders, 2007, 22, 203-206.	2.2	134
26	Age-dependent cognitive changes in carriers of the fragile X syndrome. Cortex, 2008, 44, 628-636.	1.1	133
27	Impairment in the cognitive functioning of men with fragile X-associated tremor/ataxia syndrome (FXTAS). Journal of the Neurological Sciences, 2006, 248, 227-233.	0.3	126
28	Mutations at a single codon in Mad homology 2 domain of SMAD4 cause Myhre syndrome. Nature Genetics, 2012, 44, 85-88.	9.4	125
29	A Comparative Phenotypic Study of Kallmann Syndrome Patients Carrying Monoallelic and Biallelic Mutations in the Prokineticin 2 or Prokineticin Receptor 2 Genes. Journal of Clinical Endocrinology and Metabolism, 2010, 95, 659-669.	1.8	124
30	Clinical and molecular genetic features of ARC syndrome. Human Genetics, 2006, 120, 396-409.	1.8	118
31	Mutations in CEP57 cause mosaic variegated aneuploidy syndrome. Nature Genetics, 2011, 43, 527-529.	9.4	117
32	Neuropathological, clinical and molecular pathology in female fragile X premutation carriers with and without FXTAS. Genes, Brain and Behavior, 2012, 11, 577-585.	1.1	117
33	Insufficient Evidence for "Autism-Specific―Genes. American Journal of Human Genetics, 2020, 106, 587-595.	2.6	110
34	Deletion of the SIM1 gene (6q16.2) in a patient with a Prader-Willi-like phenotype. Journal of Medical Genetics, 2002, 39, 594-596.	1.5	105
35	Size bias of fragile X premutation alleles in late-onset movement disorders. Journal of Medical Genetics, 2006, 43, 804-809.	1.5	99
36	The challenges of clinical trials in fragile X syndrome. Psychopharmacology, 2014, 231, 1237-1250.	1.5	98

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37	Lifespan changes in working memory in fragile X premutation males. Brain and Cognition, 2009, 69, 551-558.	0.8	93
38	Psychiatric disorders in children with $16p11.2$ deletion and duplication. Translational Psychiatry, 2019, 9, 8.	2.4	93
39	A framework for the investigation of rare genetic disorders in neuropsychiatry. Nature Medicine, 2019, 25, 1477-1487.	15.2	90
40	Transmission Disequilibrium of Small CNVs in Simplex Autism. American Journal of Human Genetics, 2013, 93, 595-606.	2.6	87
41	Impairment of executive cognitive functioning in males with fragile X-associated tremor/ataxia syndrome. Movement Disorders, 2007, 22, 645-650.	2.2	84
42	SCRIB and PUF60 Are Primary Drivers of the Multisystemic Phenotypes of the 8q24.3 Copy-Number Variant. American Journal of Human Genetics, 2013, 93, 798-811.	2.6	82
43	The efficacy of melatonin for sleep problems in children with autism, fragile X syndrome, or autism and fragile X syndrome. Journal of Clinical Sleep Medicine, 2009, 5, 145-50.	1.4	82
44	The Number of Genomic Copies at the 16p11.2 Locus Modulates Language, Verbal Memory, and Inhibition. Biological Psychiatry, 2016, 80, 129-139.	0.7	78
45	Measuring and Estimating the Effect Sizes of Copy Number Variants on General Intelligence in Community-Based Samples. JAMA Psychiatry, 2018, 75, 447.	6.0	77
46	The Canadian Partnership for Tomorrow Project: a pan-Canadian platform for research on chronic disease prevention. Cmaj, 2018, 190, E710-E717.	0.9	71
47	Sex differences in brain plasticity: a new hypothesis for sex ratio bias in autism. Molecular Autism, 2015, 6, 33.	2.6	70
48	Chromosomal contacts connect loci associated with autism, BMI and head circumference phenotypes. Molecular Psychiatry, 2017, 22, 836-849.	4.1	68
49	In-vivo probabilistic atlas of human thalamic nuclei based on diffusion- weighted magnetic resonance imaging. Scientific Data, 2018, 5, 180270.	2.4	67
50	CNV-association meta-analysis in 191,161 European adults reveals new loci associated with anthropometric traits. Nature Communications, 2017, 8, 744.	5.8	64
51	Screen for expandedFMR1 alleles in patients with essential tremor. Movement Disorders, 2004, 19, 930-933.	2.2	63
52	The phenotype of recurrent 10q22q23 deletions and duplications. European Journal of Human Genetics, 2011, 19, 400-408.	1.4	63
53	A Genetics-First Approach to Dissecting the Heterogeneity of Autism: Phenotypic Comparison of Autism Risk Copy Number Variants. American Journal of Psychiatry, 2021, 178, 77-86.	4.0	62
54	Penetrance of marked cognitive impairment in older male carriers of the FMR1 gene premutation. Journal of Medical Genetics, 2009, 46, 818-824.	1.5	61

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55	16p11.2 600 kb Duplications confer risk for typical and atypical Rolandic epilepsy. Human Molecular Genetics, 2014, 23, 6069-6080.	1.4	61
56	Dysregulated ADAM10-Mediated Processing of APP during a Critical Time Window Leads to Synaptic Deficits in Fragile X Syndrome. Neuron, 2015, 87, 382-398.	3.8	59
57	Robust thalamic nuclei segmentation method based on local diffusion magnetic resonance properties. Brain Structure and Function, 2017, 222, 2203-2216.	1.2	58
58	Quantifying the Effects of 16p11.2 Copy Number Variants on Brain Structure: A Multisite Genetic-First Study. Biological Psychiatry, 2018, 84, 253-264.	0.7	56
59	Spastic paraparesis, cerebellar ataxia, and intention tremor: a severe variant of FXTAS?. Journal of Medical Genetics, 2005, 42, e14-e14.	1.5	55
60	Association of Copy Number Variation of the 15q11.2 BP1-BP2 Region With Cortical and Subcortical Morphology and Cognition. JAMA Psychiatry, 2020, 77, 420.	6.0	54
61	A Potential Contributory Role for Ciliary Dysfunction in the 16p11.2 600 kb BP4-BP5 Pathology. American Journal of Human Genetics, 2015, 96, 784-796.	2.6	53
62	Brain structure in asymptomatic FMR1 premutation carriers at risk for fragile X-associated tremor/ataxia syndrome. Neurobiology of Aging, 2013, 34, 1700-1707.	1.5	52
63	Protein synthesis levels are increased in a subset of individuals with fragile X syndrome. Human Molecular Genetics, 2018, 27, 2039-2051.	1.4	51
64	Effect Sizes of Deletions and Duplications on Autism Risk Across the Genome. American Journal of Psychiatry, 2021, 178, 87-98.	4.0	50
65	The Growing Importance of CNVs: New Insights for Detection and Clinical Interpretation. Frontiers in Genetics, 2013, 4, 92.	1.1	49
66	Dose response of the $16p11.2$ distal copy number variant on intracranial volume and basal ganglia. Molecular Psychiatry, 2020, 25, 584-602.	4.1	49
67	Translating Molecular Advances in Fragile X Syndrome Into Therapy. Journal of Clinical Psychiatry, 2014, 75, e294-e307.	1.1	46
68	Biallelic mutations in the prokineticin-2 gene in two sporadic cases of Kallmann syndrome. European Journal of Human Genetics, 2008, 16, 865-868.	1.4	44
69	Cognitive Impairment in a 65-year-old Male With the Fragile X-associated Tremor-Ataxia Syndrome (FXTAS). Cognitive and Behavioral Neurology, 2006, 19, 165-171.	0.5	43
70	Estimating the effect size of the 15Q11.2 BP1–BP2 deletion and its contribution to neurodevelopmental symptoms: recommendations for practice. Journal of Medical Genetics, 2019, 56, 701-710.	1.5	43
71	Implication of <i>LRRC4C</i> and <i>DPP6</i> in neurodevelopmental disorders. American Journal of Medical Genetics, Part A, 2017, 173, 395-406.	0.7	40
72	Myhre and LAPS syndromes: clinical and molecular review of 32 patients. European Journal of Human Genetics, 2014, 22, 1272-1277.	1.4	38

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73	Prevalence of <i>FMR1</i> Repeat Expansions in Movement Disorders. Neuroepidemiology, 2006, 26, 151-155.	1.1	37
74	Dissecting autism and schizophrenia through neuroimaging genomics. Brain, 2021, 144, 1943-1957.	3.7	37
75	Mutations associated with neuropsychiatric conditions delineate functional brain connectivity dimensions contributing to autism and schizophrenia. Nature Communications, 2020, 11, 5272.	5.8	35
76	Mavoglurant in Fragile X Syndrome: Results of two open-label, extension trials in adults and adolescents. Scientific Reports, 2018, 8, 16970.	1.6	33
77	Genome-wide analysis of gene dosage in 24,092 individuals estimates that 10,000 genes modulate cognitive ability. Molecular Psychiatry, 2021, 26, 2663-2676.	4.1	33
78	Rare Genomic Structural Variants in Complex Disease: Lessons from the Replication of Associations with Obesity. PLoS ONE, 2013, 8, e58048.	1.1	33
79	Cutaneous lipoma in children: 5 cases with Bannayan-Riley-Ruvalcaba syndrome. Journal of Pediatric Surgery, 2006, 41, 1601-1603.	0.8	32
80	Disease-associated mutations in the actin-binding domain of filamin B cause cytoplasmic focal accumulations correlating with disease severity. Human Mutation, 2012, 33, 665-673.	1.1	31
81	Rationale, design, and methods for Canadian alliance for healthy hearts and minds cohort study (CAHHM) $\hat{a}$ $\in$ a Pan Canadian cohort study. BMC Public Health, 2016, 16, 650.	1.2	31
82	16p11.2 Locus modulates response to satiety before the onset of obesity. International Journal of Obesity, 2016, 40, 870-876.	1.6	31
83	The Immune Signaling Adaptor LAT Contributes to the Neuroanatomical Phenotype of 16p11.2 BP2-BP3 CNVs. American Journal of Human Genetics, 2017, 101, 564-577.	2.6	30
84	Effects of copy number variations on brain structure and risk for psychiatric illness: Largeâ€scale studies from the <scp>ENIGMA &lt; /scp&gt; working groups on <scp>CNVs &lt; /scp&gt;. Human Brain Mapping, 2022, 43, 300-328.</scp></scp>	1.9	30
85	Deleterious de novo variants of Xâ€linked <i>ZC4H2</i> in females cause a variable phenotype with neurogenic arthrogryposis multiplex congenita. Human Mutation, 2019, 40, 2270-2285.	1.1	29
86	Genes To Mental Health (G2MH): A Framework to Map the Combined Effects of Rare and Common Variants on Dimensions of Cognition and Psychopathology. American Journal of Psychiatry, 2022, 179, 189-203.	4.0	29
87	Inherited ring chromosome 8Âwithout loss of subtelomeric sequences. Annales De Génétique, 2004, 47, 289-296.	0.4	25
88	Development of mavoglurant and its potential for the treatment of fragile X syndrome. Expert Opinion on Investigational Drugs, 2014, 23, 125-134.	1.9	24
89	Molecular characterization of 39 de novo sSMC : contribution to prognosis and genetic counselling, a prospective study. Clinical Genetics, 2014, 85, 233-244.	1.0	24
90	Auditory repetition suppression alterations in relation to cognitive functioning in fragile X syndrome: a combined EEG and machine learning approach. Journal of Neurodevelopmental Disorders, 2018, 10, 4.	1.5	24

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91	1q21.1 distal copy number variants are associated with cerebral and cognitive alterations in humans. Translational Psychiatry, 2021, 11, 182.	2.4	24
92	Altered visual repetition suppression in Fragile X Syndrome: New evidence from ERPs and oscillatory activity. International Journal of Developmental Neuroscience, 2017, 59, 52-59.	0.7	23
93	Eosinophilic infiltration related to CAPN3 mutations: a pathophysiological component of primary calpainopathy?. Clinical Genetics, 2011, 80, 398-402.	1.0	22
94	Earlyâ€onset encephalopathy with epilepsy associated with a novel splice site mutation in <i>SMC1A</i> American Journal of Medical Genetics, Part A, 2015, 167, 3076-3081.	0.7	22
95	Lessons Learned From Neuroimaging Studies of Copy Number Variants: A Systematic Review. Biological Psychiatry, 2021, 90, 596-610.	0.7	22
96	Spatial control of nucleoporin condensation by fragile Xâ€related proteins. EMBO Journal, 2020, 39, e104467.	3.5	21
97	Carriers of the fragile X mental retardation 1 (FMR1) premutation allele present with increased levels of cytokine IL-10. Journal of Neuroinflammation, 2012, 9, 238.	3.1	20
98	Investigation of memory, executive functions, and anatomic correlates in asymptomatic FMR1 premutation carriers. Neurobiology of Aging, 2014, 35, 1939-1946.	1.5	20
99	Complete Maxillo-Mandibular Syngnathia in a Newborn with Multiple Congenital Malformations. Pediatrics and Neonatology, 2016, 57, 65-68.	0.3	19
100	Sex chromosome aneuploidies and copy-number variants: a further explanation for neurodevelopmental prognosis variability?. European Journal of Human Genetics, 2017, 25, 930-934.	1.4	19
101	Reciprocal changes in DNA methylation and hydroxymethylation and a broad repressive epigenetic switch characterize FMR1 transcriptional silencing in fragile X syndrome. Clinical Epigenetics, 2016, 8, 15.	1.8	18
102	Effects of eight neuropsychiatric copy number variants on human brain structure. Translational Psychiatry, 2021, 11, 399.	2.4	18
103	Screening for FXTAS. European Journal of Human Genetics, 2005, 13, 2-3.	1.4	15
104	Early neurological impairment and severe anemia in a newborn with Pearson syndrome. European Journal of Pediatrics, 2009, 168, 311-315.	1.3	14
105	Genotype–phenotype correlation at codon 1740 of <scp><i>SETD2</i></scp> . American Journal of Medical Genetics, Part A, 2020, 182, 2037-2048.	0.7	14
106	Structural and functional brain alterations revealed by neuroimaging in CNV carriers. Current Opinion in Genetics and Development, 2021, 68, 88-98.	1.5	14
107	Duplication of 10q24 locus: broadening the clinical and radiological spectrum. European Journal of Human Genetics, 2019, 27, 525-534.	1.4	13
108	Clinical utility gene card for: fragile X mental retardation syndrome, fragile X-associated tremor/ataxia syndrome and fragile X-associated primary ovarian insufficiency. European Journal of Human Genetics, 2011, 19, 1017-1017.	1.4	12

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109	Reporting incidental findings of genomic disorder-associated copy number variants to unselected biobank participants. Personalized Medicine, 2016, 13, 303-314.	0.8	12
110	Clinimetric Properties of the Fragile Xâ€associated Tremor Ataxia Syndrome Rating Scale. Movement Disorders Clinical Practice, 2019, 6, 120-124.	0.8	12
111	Bayonet-shaped language development in autism with regression: a retrospective study. Molecular Autism, 2021, 12, 35.	2.6	12
112	Convergence and Divergence of Rare Genetic Disorders on Brain Phenotypes. JAMA Psychiatry, 2022, 79, 818.	6.0	12
113	Cardiovascular risk scoring and magnetic resonance imaging detected subclinical cerebrovascular disease. European Heart Journal Cardiovascular Imaging, 2020, 21, 692-700.	0.5	11
114	Mitochondrial tRNA <sup>Leu(UUR)</sup> mutation m.3302A > G presenting as childhoodâ€onset se myopathy: threshold determination through segregation study. Journal of Inherited Metabolic Disease, 2010, 33, 219-226.	evere 1.7	10
115	Phenotypic and molecular characterization of a novel case of dyssegmental dysplasia, Silverman-Handmaker type. European Journal of Medical Genetics, 2010, 53, 294-298.	0.7	10
116	Developmental trajectories of neuroanatomical alterations associated with the 16p11.2 Copy Number Variations. NeuroImage, 2019, 203, 116155.	2.1	9
117	Reduced Cognitive Assessment Scores Among Individuals With Magnetic Resonance Imaging–Detected Vascular Brain Injury. Stroke, 2020, 51, 1158-1165.	1.0	9
118	Deletion of Loss-of-Function–Intolerant Genes and Risk of 5 Psychiatric Disorders. JAMA Psychiatry, 2022, 79, 78.	6.0	8
119	Copy Number Variant Risk Scores Associated With Cognition, Psychopathology, and Brain Structure in Youths in the Philadelphia Neurodevelopmental Cohort. JAMA Psychiatry, 2022, 79, 699.	6.0	8
120	The effects of sex on prevalence and mechanisms underlying neurodevelopmental disorders. Handbook of Clinical Neurology / Edited By P J Vinken and G W Bruyn, 2020, 173, 327-339.	1.0	7
121	Conditional canonical correlation estimation based on covariates with random forests. Bioinformatics, 2021, 37, 2714-2721.	1.8	7
122	A GENETIC FIRST APPROACH TO DISSECTING THE HETEROGENEITY OF AUTISM: PHENOTYPIC COMPARISON OF AUTISM RISK COPY NUMBER VARIANTS. European Neuropsychopharmacology, 2019, 29, S783-S784.	0.3	6
123	The psychiatric phenotypes of $1q21$ distal deletion and duplication. Translational Psychiatry, 2021, 11, 105.	2.4	6
124	The Epidemiology of FXTAS. , 2010, , 17-30.		6
125	EEG Signal Complexity Is Reduced During Resting-State in Fragile X Syndrome. Frontiers in Psychiatry, 2021, 12, 716707.	1.3	5
126	Familial aplasia of the trapezius muscle: clinical and MRI findings. Acta Paediatrica, International Journal of Paediatrics, 2011, 100, 464-466.	0.7	4

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127	Preimplantation genetic diagnosis (PGD) for HLA typing: bases for setting up an open international collaboration when PGD is not available. Fertility and Sterility, 2010, 94, 1129-1131.	0.5	3
128	Possible association of $16p11.2$ copy number variation with altered lymphocyte and neutrophil counts. Npj Genomic Medicine, $2022, 7, .$	1.7	3
129	Copy Number Variations and Cognitive Phenotypes in Unselected Populations. Obstetrical and Gynecological Survey, 2015, 70, 559-560.	0.2	2
130	Tinkering with the vasopressin pathway in autism. Science Translational Medicine, 2019, 11, .	5.8	2
131	Functional Connectivity Analyses Suggest Shared Molecular Mechanisms Across 12 Neuropsychiatric Mutations, Autism and Schizophrenia. Biological Psychiatry, 2020, 87, S395.	0.7	2
132	250. Gene Dosage Effects on Neurobehavioral PhenotypesÂand Development: Relevance to Idiopathic Neuropsychiatric Disorders. Biological Psychiatry, 2019, 85, S104.	0.7	1
133	Poster Withdrawn: QUANTIFYING THE EFFECTS OF 16P11.2 CNVs ON BRAIN STRUCTURE, A MULTI-SITE â€~GENETIC-FIRST'MRI STUDY. European Neuropsychopharmacology, 2019, 29, S859-S860.	0.3	1
134	Measuring and Estimating the Effect Size of Rare Non-Recurrent Deletions and Duplications on General Intelligence. Biological Psychiatry, 2020, 87, S196.	0.7	1
135	Estimating the effects of copyâ€number variants on intelligence using hierarchical Bayesian models. Genetic Epidemiology, 2020, 44, 825-840.	0.6	1
136	White Matter Diffusion MRI Findings in Carriers of 16p11.2 Copy Number Variants. Biological Psychiatry, 2021, 89, S40.	0.7	1
137	Reflections on Clinical Trials in Fragile X Syndrome. , 2017, , 419-441.		1
138	Dysregulated ADAM10-Mediated Processing of APP during a Critical Time Window Leads to SynapticDeficits in Fragile X Syndrome. Neuron, 2015, 87, 908.	3.8	0
139	Cover Image, Volume 173A, Number 2, February 2017. American Journal of Medical Genetics, Part A, 2017, 173, i.	0.7	0
140	QUANTIFYING THE EFFECT OF COPY-NUMBER VARIANTS ON GENERAL INTELLIGENCE IN UNSELECTED POPULATIONS. European Neuropsychopharmacology, 2019, 29, S848.	0.3	0
141	5 DIFFERENCES IN THE GENETIC BACKGROUND CONTRIBUTE TO RISK AND RESILIENCE TO AUTISM. European Neuropsychopharmacology, 2019, 29, S61.	0.3	0
142	40 MAPPING THE EFFECT-SIZE OF GENE DOSAGE ON GENERAL INTELLIGENCE ACROSS THE GENOME. European Neuropsychopharmacology, 2019, 29, S81.	0.3	0
143	191. Mirror Effects of 4 Neurodevelopmental CNVs on Functional Connectivity and Implication for Idiopathic Autism. Biological Psychiatry, 2019, 85, S79.	0.7	0
144	251. Measuring and Estimating the Effects of Rare Variants, Genome-Wide, on Cognition. Biological Psychiatry, 2019, 85, S104.	0.7	0

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145	M59 SHARED FUNCTIONAL CONNECTIVITY ALTERATIONS ACROSS NEURODEVELOPMENTAL MUTATIONS, AUTISM AND SCHIZOPHRENIA. European Neuropsychopharmacology, 2019, 29, S197-S198.	0.3	O
146	Estimating the Effect Size of CNVs on Risk for Major Psychopathology, the CAMP Project. Biological Psychiatry, 2020, 87, S392.	0.7	0
147	Omnigenic Impact of Copy Number Variants on Cognition and Psychopathology in the Philadelphia Neurodevelopmental Cohort. Biological Psychiatry, 2021, 89, S320.	0.7	O
148	Analysis of Genomic Copy Number Variation Across Psychiatric Disorders. Biological Psychiatry, 2021, 89, S106-S107.	0.7	0
149	Assessing the Effects of Rare Copy Number Variants on Psychiatric Symptoms and Cognitive Ability in 1M Individuals. Biological Psychiatry, 2021, 89, S46.	0.7	O
150	The General Impact of Haploinsufficiency on Brain Connectivity Underlies the Pleiotropic Effect of Neuropsychiatric CNVS. Biological Psychiatry, 2021, 89, S40.	0.7	0
151	ENIGMA-CNV and Other Initiatives to Understand the Impact of Rare Copy Number Variants on Brain Structure and Other Measures. Biological Psychiatry, 2021, 89, S41.	0.7	O