## Sophie Saunier

## List of Publications by Year in Descending Order

Source: https://exaly.com/author-pdf/2906946/sophie-saunier-publications-by-year.pdf

Version: 2024-04-28

This document has been generated based on the publications and citations recorded by exaly.com. For the latest version of this publication list, visit the link given above.

The third column is the impact factor (IF) of the journal, and the fourth column is the number of citations of the article.

71	5,768	42	75
papers	citations	h-index	g-index
77	6,662 ext. citations	11.8	4.46
ext. papers		avg, IF	L-index

#	Paper	IF	Citations
71	Agonists of prostaglandin E receptors as potential first in class treatment for nephronophthisis and related ciliopathies <i>Proceedings of the National Academy of Sciences of the United States of America</i> , <b>2022</b> , 119, e2115960119	11.5	1
70	Renal Ciliopathies: Sorting Out Therapeutic Approaches for Nephronophthisis. <i>Frontiers in Cell and Developmental Biology</i> , <b>2021</b> , 9, 653138	5.7	6
69	Bi-allelic pathogenic variations in DNAJB11 cause Ivemark II syndrome, a renal-hepatic-pancreatic dysplasia. <i>Kidney International</i> , <b>2021</b> , 99, 405-409	9.9	7
68	Cystic kidney diseases associated with mutations in phosphomannomutase 2 promotor: a large spectrum of phenotypes. <i>Pediatric Nephrology</i> , <b>2021</b> , 36, 2361-2369	3.2	2
67	Mouse genetics reveals Barttin as a genetic modifier of Joubert syndrome. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , <b>2020</b> , 117, 1113-1118	11.5	13
66	Novel nephronophthisis-associated variants reveal functional importance of MAPKBP1 dimerization for centriolar recruitment. <i>Kidney International</i> , <b>2020</b> , 98, 958-969	9.9	4
65	Human IFT52 mutations uncover a novel role for the protein in microtubule dynamics and centrosome cohesion. <i>Human Molecular Genetics</i> , <b>2019</b> , 28, 2720-2737	5.6	7
64	Loss-of-function mutations in KIF14 cause severe microcephaly and kidney development defects in humans and zebrafish. <i>Human Molecular Genetics</i> , <b>2019</b> , 28, 778-795	5.6	22
63	Cell type-specific regulation of ciliary transition zone assembly in vertebrates. <i>EMBO Journal</i> , <b>2018</b> , 37,	13	31
62	Functional characterization of tektin-1 in motile cilia and evidence for TEKT1 as a new candidate gene for motile ciliopathies. <i>Human Molecular Genetics</i> , <b>2018</b> , 27, 266-282	5.6	14
61	Whole-genome sequencing in patients with ciliopathies uncovers a novel recurrent tandem duplication in IFT140. <i>Human Mutation</i> , <b>2018</b> , 39, 983-992	4.7	10
60	Targeted exon skipping of a mutation rescues Joubert syndrome phenotypes in vitro and in a murine model. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , <b>2018</b> , 115, 12489-12494	11.5	27
59	Casein kinase 1년and 1년as novel players in polycystic kidney disease and mechanistic targets for (R)-roscovitine and (S)-CR8. <i>American Journal of Physiology - Renal Physiology</i> , <b>2018</b> , 315, F57-F73	4.3	3
58	KIF13B establishes a CAV1-enriched microdomain at the ciliary transition zone to promote Sonic hedgehog signalling. <i>Nature Communications</i> , <b>2017</b> , 8, 14177	17.4	37
57	Mutations in MAPKBP1 Cause Juvenile or Late-Onset Cilia-Independent Nephronophthisis. <i>American Journal of Human Genetics</i> , <b>2017</b> , 100, 323-333	11	19
56	Targeted Exome Sequencing Identifies as Involved in Monogenic Congenital Anomalies of the Kidney and Urinary Tract. <i>Journal of the American Society of Nephrology: JASN</i> , <b>2017</b> , 28, 2901-2914	12.7	59
55	Fifteen years of research on oral-facial-digital syndromes: from 1 to 16 causal genes. <i>Journal of Medical Genetics</i> , <b>2017</b> , 54, 371-380	5.8	58

## (2014-2017)

54	QMPSF is sensitive and specific in the detection of NPHP1 heterozygous deletions. <i>Clinical Chemistry and Laboratory Medicine</i> , <b>2017</b> , 55, 809-816	5.9	5
53	Mutations in GREB1L Cause Bilateral Kidney Agenesis in Humans and Mice. <i>American Journal of Human Genetics</i> , <b>2017</b> , 101, 803-814	11	41
52	A human patient-derived cellular model of Joubert syndrome reveals ciliary defects which can be rescued with targeted therapies. <i>Human Molecular Genetics</i> , <b>2017</b> , 26, 4657-4667	5.6	40
51	TMEM107 recruits ciliopathy proteins to subdomains of the ciliary transition zone and causes Joubert syndrome. <i>Nature Cell Biology</i> , <b>2016</b> , 18, 122-31	23.4	81
50	Novel NEK8 Mutations Cause Severe Syndromic Renal Cystic Dysplasia through YAP Dysregulation. <i>PLoS Genetics</i> , <b>2016</b> , 12, e1005894	6	53
49	Mutation of Growth Arrest Specific 8 Reveals a Role in Motile Cilia Function and Human Disease. <i>PLoS Genetics</i> , <b>2016</b> , 12, e1006220	6	24
48	DCDC2 Mutations Cause Neonatal Sclerosing Cholangitis. <i>Human Mutation</i> , <b>2016</b> , 37, 1025-9	4.7	32
47	DCDC2 mutations cause a renal-hepatic ciliopathy by disrupting Wnt signaling. <i>American Journal of Human Genetics</i> , <b>2015</b> , 96, 81-92	11	66
46	Mutations in KIAA0586 Cause Lethal Ciliopathies Ranging from a Hydrolethalus Phenotype to Short-Rib Polydactyly Syndrome. <i>American Journal of Human Genetics</i> , <b>2015</b> , 97, 311-8	11	57
45	TMEM231, mutated in orofaciodigital and Meckel syndromes, organizes the ciliary transition zone. <i>Journal of Cell Biology</i> , <b>2015</b> , 209, 129-42	7.3	71
44	IFT81, encoding an IFT-B core protein, as a very rare cause of a ciliopathy phenotype. <i>Journal of Medical Genetics</i> , <b>2015</b> , 52, 657-65	5.8	27
43	Mutations in TRAF3IP1/IFT54 reveal a new role for IFT proteins in microtubule stabilization. <i>Nature Communications</i> , <b>2015</b> , 6, 8666	17.4	65
42	The oral-facial-digital syndrome gene C2CD3 encodes a positive regulator of centriole elongation. <i>Nature Genetics</i> , <b>2014</b> , 46, 905-11	36.3	88
41	Integrin Alpha 8 Recessive Mutations Are Responsible for Bilateral Renal Agenesis in Humans. <i>American Journal of Human Genetics</i> , <b>2014</b> , 94, 799	11	78
40	Integrin alpha 8 recessive mutations are responsible for bilateral renal agenesis in humans. <i>American Journal of Human Genetics</i> , <b>2014</b> , 94, 288-94	11	71
39	Mutations of CEP83 cause infantile nephronophthisis and intellectual disability. <i>American Journal of Human Genetics</i> , <b>2014</b> , 94, 905-14	11	59
38	A homozygous PDE6D mutation in Joubert syndrome impairs targeting of farnesylated INPP5E protein to the primary cilium. <i>Human Mutation</i> , <b>2014</b> , 35, 137-46	4.7	88
37	A homozygous missense mutation in the ciliary gene TTC21B causes familial FSGS. <i>Journal of the American Society of Nephrology: JASN</i> , <b>2014</b> , 25, 2435-43	12.7	68

36	ANKS6 is a central component of a nephronophthisis module linking NEK8 to INVS and NPHP3. <i>Nature Genetics</i> , <b>2013</b> , 45, 951-6	36.3	144
35	Genetic bases and pathogenic mechanisms of nephronophthisis. <i>Drug Discovery Today Disease Mechanisms</i> , <b>2013</b> , 10, e143-e151		2
34	Combined NGS approaches identify mutations in the intraflagellar transport gene IFT140 in skeletal ciliopathies with early progressive kidney Disease. <i>Human Mutation</i> , <b>2013</b> , 34, 714-24	4.7	89
33	Defects in the IFT-B component IFT172 cause Jeune and Mainzer-Saldino syndromes in humans. <i>American Journal of Human Genetics</i> , <b>2013</b> , 93, 915-25	11	155
32	Septins 2, 7 and 9 and MAP4 colocalize along the axoneme in the primary cilium and control ciliary length. <i>Journal of Cell Science</i> , <b>2013</b> , 126, 2583-94	5.3	93
31	Inversin/Nephrocystin-2 is required for fibroblast polarity and directional cell migration. <i>PLoS ONE</i> , <b>2013</b> , 8, e60193	3.7	37
30	Exome capture reveals ZNF423 and CEP164 mutations, linking renal ciliopathies to DNA damage response signaling. <i>Cell</i> , <b>2012</b> , 150, 533-48	56.2	266
29	TCTN3 mutations cause Mohr-Majewski syndrome. American Journal of Human Genetics, 2012, 91, 372-8	11	96
28	Mainzer-Saldino syndrome is a ciliopathy caused by IFT140 mutations. <i>American Journal of Human Genetics</i> , <b>2012</b> , 90, 864-70	11	134
27	Dishevelled stabilization by the ciliopathy protein Rpgrip1l is essential for planar cell polarity. Journal of Cell Biology, <b>2012</b> , 198, 927-40	7-3	48
26	INF2 mutations in Charcot-Marie-Tooth disease with glomerulopathy. <i>New England Journal of Medicine</i> , <b>2011</b> , 365, 2377-88	59.2	200
25	Mutation analysis of 18 nephronophthisis associated ciliopathy disease genes using a DNA pooling and next generation sequencing strategy. <i>Journal of Medical Genetics</i> , <b>2011</b> , 48, 105-16	5.8	112
24	TTC21B contributes both causal and modifying alleles across the ciliopathy spectrum. <i>Nature Genetics</i> , <b>2011</b> , 43, 189-96	36.3	271
23	Ciliopathies with skeletal anomalies and renal insufficiency due to mutations in the IFT-A gene WDR19. <i>American Journal of Human Genetics</i> , <b>2011</b> , 89, 634-43	11	180
22	Control of the Wnt pathways by nephrocystin-4 is required for morphogenesis of the zebrafish pronephros. <i>Human Molecular Genetics</i> , <b>2011</b> , 20, 2611-27	5.6	45
21	Mutations in TMEM216 perturb ciliogenesis and cause Joubert, Meckel and related syndromes. <i>Nature Genetics</i> , <b>2010</b> , 42, 619-25	36.3	210
20	Candidate exome capture identifies mutation of SDCCAG8 as the cause of a retinal-renal ciliopathy. <i>Nature Genetics</i> , <b>2010</b> , 42, 840-50	36.3	257
19	The ciliary pocket: an endocytic membrane domain at the base of primary and motile cilia. <i>Journal of Cell Science</i> , <b>2010</b> , 123, 1785-95	5.3	197

## (1995-2010)

18	Nephrocystin-1 forms a complex with polycystin-1 via a polyproline motif/SH3 domain interaction and regulates the apoptotic response in mammals. <i>PLoS ONE</i> , <b>2010</b> , 5, e12719	3.7	22
17	Mutations of NPHP2 and NPHP3 in infantile nephronophthisis. <i>Kidney International</i> , <b>2009</b> , 75, 839-47	9.9	86
16	CC2D2A mutations in Meckel and Joubert syndromes indicate a genotype-phenotype correlation. <i>Human Mutation</i> , <b>2009</b> , 30, 1574-82	4.7	66
15	Nephronophthisis. <i>Pediatric Nephrology</i> , <b>2009</b> , 24, 2333-44	3.2	122
14	Nephrocystin-1 and nephrocystin-4 are required for epithelial morphogenesis and associate with PALS1/PATJ and Par6. <i>Human Molecular Genetics</i> , <b>2009</b> , 18, 4711-23	5.6	76
13	Matrix metalloproteinase 13 (MMP13) and tissue inhibitor of matrix metalloproteinase 1 (TIMP1), regulated by the MAPK pathway, are both necessary for Madin-Darby canine kidney tubulogenesis. <i>Journal of Biological Chemistry</i> , <b>2008</b> , 283, 4272-82	5.4	40
12	Targeting of beta-arrestin2 to the centrosome and primary cilium: role in cell proliferation control. <i>PLoS ONE</i> , <b>2008</b> , 3, e3728	3.7	35
11	The ciliary gene RPGRIP1L is mutated in cerebello-oculo-renal syndrome (Joubert syndrome type B) and Meckel syndrome. <i>Nature Genetics</i> , <b>2007</b> , 39, 875-81	36.3	379
10	High NPHP1 and NPHP6 mutation rate in patients with Joubert syndrome and nephronophthisis: potential epistatic effect of NPHP6 and AHI1 mutations in patients with NPHP1 mutations. <i>Journal of the American Society of Nephrology: JASN</i> , <b>2007</b> , 18, 1566-75	12.7	128
9	The Meckel-Gruber syndrome gene, MKS3, is mutated in Joubert syndrome. <i>American Journal of Human Genetics</i> , <b>2007</b> , 80, 186-94	11	193
8	Pleiotropic effects of CEP290 (NPHP6) mutations extend to Meckel syndrome. <i>American Journal of Human Genetics</i> , <b>2007</b> , 81, 170-9	11	221
7	Nephronophthisis. Current Opinion in Genetics and Development, 2005, 15, 324-31	4.9	69
6	Solution NMR structure of the SH3 domain of human nephrocystin and analysis of a mutation-causing juvenile nephronophthisis. <i>Proteins: Structure, Function and Bioinformatics</i> , <b>2005</b> , 59, 347-55	4.2	6
5	Characterization of the nephrocystin/nephrocystin-4 complex and subcellular localization of nephrocystin-4 to primary cilia and centrosomes. <i>Human Molecular Genetics</i> , <b>2005</b> , 14, 645-56	5.6	130
4	The gene mutated in juvenile nephronophthisis type 4 encodes a novel protein that interacts with nephrocystin. <i>Nature Genetics</i> , <b>2002</b> , 32, 300-5	36.3	186
3	Characterization of the NPHP1 locus: mutational mechanism involved in deletions in familial juvenile nephronophthisis. <i>American Journal of Human Genetics</i> , <b>2000</b> , 66, 778-89	11	103
2	Familial juvenile nephronophthisis. <i>Journal of Molecular Medicine</i> , <b>1998</b> , 76, 310-6	5.5	14
1	A 11 Mb YAC-based contig spanning the familial juvenile nephronophthisis region (NPH1) located on chromosome 2q. <i>Genomics</i> , <b>1995</b> , 30, 514-20	4.3	14