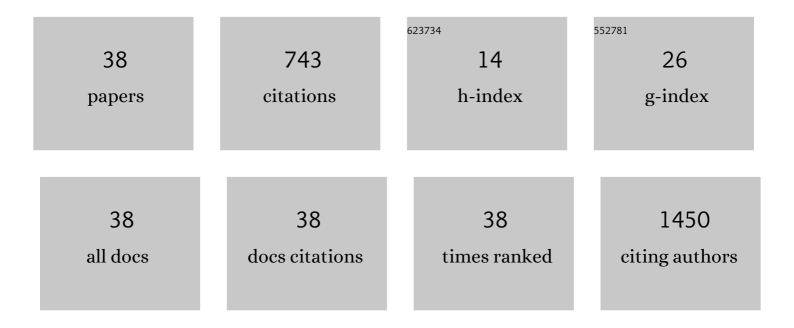
## Anne-Marie Laberge

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/7761938/publications.pdf Version: 2024-02-01



#	Article	IF	CITATIONS
1	The clinical application of genome-wide sequencing for monogenic diseases in Canada: Position Statement of the Canadian College of Medical Geneticists. Journal of Medical Genetics, 2015, 52, 431-437.	3.2	187
2	BCL11B mutations in patients affected by a neurodevelopmental disorder with reduced type 2 innate lymphoid cells. Brain, 2018, 141, 2299-2311.	7.6	81
3	Experiences of caregivers of children with inherited metabolic diseases: a qualitative study. Orphanet Journal of Rare Diseases, 2016, 11, 168.	2.7	38
4	The health system impact of false positive newborn screening results for medium-chain acyl-CoA dehydrogenase deficiency: a cohort study. Orphanet Journal of Rare Diseases, 2016, 11, 12.	2.7	38
5	A Novel Recurrent <i>COL5A1</i> Genetic Variant Is Associated With a Dysplasia-Associated Arterial Disease Exhibiting Dissections and Fibromuscular Dysplasia. Arteriosclerosis, Thrombosis, and Vascular Biology, 2020, 40, 2686-2699.	2.4	30
6	Child and family experiences with inborn errors of metabolism: a qualitative interview study with representatives of patient groups. Journal of Inherited Metabolic Disease, 2016, 39, 139-147.	3.6	26
7	Canadian Pregnant Women's Preferences Regarding NIPT for Down Syndrome: The Information They Want, How They Want to Get It, and With Whom They Want to Discuss It. Journal of Obstetrics and Gynaecology Canada, 2019, 41, 782-791.	0.7	25
8	The value of non-invasive prenatal testing: preferences of Canadian pregnant women, their partners, and health professionals regarding NIPT use and access. BMC Pregnancy and Childbirth, 2019, 19, 22.	2.4	23
9	Scoping review of patient- and family-oriented outcomes and measures for chronic pediatric disease. BMC Pediatrics, 2015, 15, 7.	1.7	20
10	Cross-cultural perspectives on decision making regarding noninvasive prenatal testing: A comparative study of Lebanon and Quebec. AJOB Empirical Bioethics, 2018, 9, 99-111.	1.6	19
11	Homozygous/compound heterozygote <i>RYR1</i> gene variants: Expanding the clinical spectrum. American Journal of Medical Genetics, Part A, 2019, 179, 386-396.	1.2	19
12	SecondaryÂfindings from next-generation sequencing: what does actionable in childhood really mean?. Genetics in Medicine, 2019, 21, 124-132.	2.4	18
13	Long-term outcomes of the "Genetics in Primary Care" faculty development initiative. Family Medicine, 2009, 41, 266-70.	0.5	18
14	When to test fetuses for RASopathies? Proposition from a systematic analysis of 352 multicenter cases and a postnatal cohort. Genetics in Medicine, 2021, 23, 1116-1124.	2.4	17
15	Use of Factor V Leiden genetic testing in practice and impact on management. Genetics in Medicine, 2009, 11, 750-756.	2.4	16
16	Obstetric and cardiac outcomes in women with Marfan syndrome and an aortic root diameter ≤45mm. European Journal of Obstetrics, Gynecology and Reproductive Biology, 2018, 230, 68-72.	1.1	14
17	Improving recommendations for genomic medicine: building an evolutionary process from clinical practice advisory documents to guidelines. Genetics in Medicine, 2019, 21, 2431-2438.	2.4	13
18	Genetic Testing in Thoracic Aortic Disease—When, Why, and How?. Canadian Journal of Cardiology, 2016, 32, 131-134.	1.7	12

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19	Genetic burden linked to founder effects in Saguenay–Lac-Saint-Jean illustrates the importance of genetic screening test availability. Journal of Medical Genetics, 2021, 58, 653-665.	3.2	12
20	Experience of carrier couples identified through a populationâ€based carrier screening pilot program for four founder autosomal recessive diseases in Saguenay–Lacâ€5aintâ€Jean. Prenatal Diagnosis, 2018, 38, 67-74.	2.3	12
21	Diagnostic and Therapeutic Misconception: Parental Expectations and Perspectives Regarding Genetic Testing for Developmental Disorders. Journal of Autism and Developmental Disorders, 2019, 49, 363-375.	2.7	11
22	Retrospective analysis of fetal vertebral defects: Associated anomalies, etiologies, and outcome. American Journal of Medical Genetics, Part A, 2020, 182, 664-672.	1.2	11
23	Implementation challenges for an ethical introduction of noninvasive prenatal testing: a qualitative study of healthcare professionals' views from Lebanon and Quebec. BMC Medical Ethics, 2020, 21, 15.	2.4	11
24	Noninvasive Prenatal Testing: Views of Canadian Pregnant Women and Their Partners Regarding Pressure and Societal Concerns. AJOB Empirical Bioethics, 2021, 12, 53-62.	1.6	10
25	Health services use among children diagnosed with medium-chain acyl-CoA dehydrogenase deficiency through newborn screening: a cohort study in Ontario, Canada. Orphanet Journal of Rare Diseases, 2019, 14, 70.	2.7	9
26	Metabolically healthy obesity in children enrolled in the <scp>CANadian</scp> Pediatric Weight management Registry ( <scp>CANPWR</scp> ): An exploratory secondary analysis of baseline data. Clinical Obesity, 2022, 12, e12490.	2.0	9
27	Pre-implantation Genetic Diagnosis: The Road Forward in Canada. Journal of Obstetrics and Gynaecology Canada, 2019, 41, 68-71.	0.7	7
28	The Serious Factor in Expanded Prenatal Genetic Testing. American Journal of Bioethics, 2022, 22, 23-25.	0.9	7
29	Paediatricians underuse recommended genetic tests in children with global developmental delay. Paediatrics and Child Health, 2018, 23, e156-e162.	0.6	5
30	A qualitative study of women and partners from Lebanon and Quebec regarding an expanded scope of noninvasive prenatal testing. BMC Pregnancy and Childbirth, 2021, 21, 54.	2.4	5
31	Screening Children for Familial Aortopathies: Tread With Caution. Canadian Journal of Cardiology, 2016, 32, 60-65.	1.7	4
32	Toward Broader Genetic Contextualism: Genetic Testing Enters the Age of Evidence-Based Medicine. American Journal of Bioethics, 2019, 19, 77-79.	0.9	4
33	Recommending inclusion of HFE C282Y homozygotes in the ACMG actionable gene list: cop-out or stealth move toward population screening?. Genetics in Medicine, 2018, 20, 400-402.	2.4	3
34	Variability in How Canadian Pediatric Weight Management Clinics Deliver Care: Evidence from the CANadian Pediatric Weight Management Registry. Childhood Obesity, 2021, 17, 420-426.	1.5	3
35	Expanded Prenatal Testing: Maintaining a Non-Directive Approach to Promote Reproductive Autonomy. American Journal of Bioethics, 2022, 22, 39-42.	0.9	3
36	Individual and family characteristics associated with health indicators at entry into multidisciplinary pediatric weight management: findings from the CANadian Pediatric Weight management Registry (CANPWR). International Journal of Obesity, 2021, , .	3.4	2

#	Article	IF	CITATIONS
37	Family Experiences with Care for Children with Inherited Metabolic Diseases in Canada: A Cross-Sectional Survey. Patient, 2022, 15, 171-185.	2.7	1
38	MG-114â€First 2 years of experience of an integrated multidisciplinary clinic for adults with aortopathies in a canadian context. Journal of Medical Genetics, 2015, 52, A3.1-A3.	3.2	0