Pierre Vabres

List of Publications by Year in descending order

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177 papers 13,326 citations

50276 46 h-index 109 g-index

219 all docs

219 docs citations

219 times ranked

19709 citing authors

#	Article	IF	CITATIONS
1	Clinical and molecular data in cases of prenatal localized overgrowth disorder: major implication of genetic variants in <scp>Pl3Kâ€AKTâ€mTOR</scp> signaling pathway. Ultrasound in Obstetrics and Gynecology, 2022, 59, 532-542.	1.7	6
2	A standard of care for individuals with <scp><i>PIK3CA</i></scp> â€related disorders: An international expert consensus statement. Clinical Genetics, 2022, 101, 32-47.	2.0	21
3	Toward clinical and molecular dissection of frontonasal dysplasia with facial skin polyps: From Pai syndrome to differential diagnosis through a series of 27 patients. American Journal of Medical Genetics, Part A, 2022, 188, 2036-2047.	1.2	1
4	Quality of life of children with capillary malformations of the lower limbs: Evolution and associated factors. Data from the French national paediatric cohort, CONAPE. Annales De Dermatologie Et De Venereologie, 2022, , .	1.0	1
5	Clinical and neuroimaging findings in 33 patients with <scp>MCAP</scp> syndrome: A survey to evaluate relevant endpoints for future clinical trials. Clinical Genetics, 2021, 99, 650-661.	2.0	12
6	Complete agenesis of the corpus callosum in phacomatosis pigmentovascularis cesioflammea: a manifestation of mosaicism?. European Journal of Dermatology, 2021, 31, 248-249.	0.6	0
7	Clinical spectrum of MTOR-related hypomelanosis of Ito with neurodevelopmental abnormalities. Genetics in Medicine, 2021, 23, 1484-1491.	2.4	14
8	Cerebriform sebaceous nevus: a subtype of organoid nevus due to specific postzygotic <i>FGFR2</i> mutations. Journal of the European Academy of Dermatology and Venereology, 2021, 35, 2085-2090.	2.4	6
9	ARP-T1-associated Bazex–Dupré–Christol syndrome is an inherited basal cell cancer with ciliary defects characteristic of ciliopathies. Communications Biology, 2021, 4, 544.	4.4	7
10	Acute acral eruptions in children during the COVID-19 pandemic: Characteristics of 103 children and their family clusters. Annales De Dermatologie Et De Venereologie, 2021, 148, 94-100.	1.0	15
11	Expanding the clinical spectrum of mosaic <i>BRAF</i> skin phenotypes. Journal of the European Academy of Dermatology and Venereology, 2021, 35, e690-e693.	2.4	1
12	Response to Resta et al Genetics in Medicine, 2021, 23, 2225.	2.4	0
13	Health care transition for patients with vascular malformations: a French multicenter cross-sectional study. Orphanet Journal of Rare Diseases, 2021, 16, 352.	2.7	5
14	Safety and efficacy of low-dose PI3K inhibitor taselisib in adult patients with CLOVES and Klippel–Trenaunay syndrome (KTS): the TOTEM trial, a phase 1/2 multicenter, open-label, single-arm study. Genetics in Medicine, 2021, 23, 2433-2442.	2.4	12
15	Skin conditions among 20th century politicians and world leaders. Journal of the European Academy of Dermatology and Venereology, 2021, 35, 2346-2348.	2.4	2
16	Autoantibodies neutralizing type I IFNs are present in \sim 4% of uninfected individuals over 70 years old and account for \sim 20% of COVID-19 deaths. Science Immunology, 2021, 6, .	11.9	357
17	Sirolimus (Rapamycin) for Slow-Flow Malformations in Children. JAMA Dermatology, 2021, 157, 1289.	4.1	51
18	Mosaic NEK9 mutation, fibrous dysplasia and premature puberty in naevus comedonicus syndrome. British Journal of Dermatology, 2021, , .	1.5	1

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19	Fertility in McCune Albright syndrome female: A case study focusing on AMH as a marker of ovarian dysfunction and a literature review. Journal of Gynecology Obstetrics and Human Reproduction, 2021, 50, 102171.	1.3	2
20	Mosaic abnormalities of the skin: review and guidelines from the European Reference Network for rare skin diseases. British Journal of Dermatology, 2020, 182, 552-563.	1.5	45
21	Rituximab is an effective treatment in patients with pemphigus vulgaris and demonstrates a steroidâ€sparing effect. British Journal of Dermatology, 2020, 182, 1111-1119.	1.5	55
22	Reverse Phenotyping in Patients with Skin Capillary Malformations and Mosaic GNAQ or GNA11 Mutations Defines a Clinical Spectrum with Genotype-Phenotype Correlation. Journal of Investigative Dermatology, 2020, 140, 1106-1110.e2.	0.7	30
23	Congenital infiltrating lipomatosis of the face with lingual mucosal neuromas associated with a <i>PIK3CA</i> mutation. Pediatric Dermatology, 2020, 37, 1128-1130.	0.9	9
24	Limb overgrowth associated with a mosaic <scp><i>TSC2</i></scp> secondâ€hit in tuberous sclerosis complex. American Journal of Medical Genetics, Part A, 2020, 182, 2803-2804.	1.2	4
25	Inborn errors of type I IFN immunity in patients with life-threatening COVID-19. Science, 2020, 370, .	12.6	1,749
26	Autoantibodies against type I IFNs in patients with life-threatening COVID-19. Science, 2020, 370, .	12.6	1,983
27	<i>De novo</i> mutations in the X-linked <i>TFE3</i> gene cause intellectual disability with pigmentary mosaicism and storage disorder-like features. Journal of Medical Genetics, 2020, 57, 808-819.	3.2	11
28	Neutralization of HSF1 in cells from PIK3CA-related overgrowth spectrum patients blocks abnormal proliferation. Biochemical and Biophysical Research Communications, 2020, 530, 520-526.	2.1	5
29	Factors Associated With Short-term Relapse in Patients With Pemphigus Who Receive Rituximab as First-line Therapy. JAMA Dermatology, 2020, 156, 545.	4.1	40
30	Kosaki overgrowth syndrome: A novel pathogenic variant in PDGFRB and expansion of the phenotype including cerebrovascular complications. Clinical Genetics, 2020, 98, 19-31.	2.0	17
31	Compassionate use of everolimus for refractory epilepsy in a patient with MTOR mosaic mutation. European Journal of Medical Genetics, 2020, 63, 104036.	1.3	8
32	Epidermolysis bullosa simplex generalized severe induces a T helper 17 response and is improved by apremilast treatment. British Journal of Dermatology, 2019, 180, 357-364.	1.5	34
33	Severe gynaecological involvement in Proteus Syndrome. European Journal of Medical Genetics, 2019, 62, 270-272.	1.3	3
34	Mosaicism due to postzygotic mutations in women with focal dermal hypoplasia. British Journal of Dermatology, 2019, 180, 657-661.	1.5	5
35	Deciphering exome sequencing data: Bringing mitochondrial DNA variants to light. Human Mutation, 2019, 40, 2430-2443.	2.5	11
36	Genetical, clinical, and functional analysis of a large international cohort of patients with autosomal recessive congenital ichthyosis due to mutations in <i>NIPAL4</i> . Human Mutation, 2019, 40, 2318-2333.	2.5	8

3

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37	315 Cerebriform sebaceous nevus is caused by the specific postzygotic FGRF2 p.(Cys382Arg) variation. Journal of Investigative Dermatology, 2019, 139, S268.	0.7	O
38	Follow-Up of Patients With Complete Remission of Locally Advanced Basal Cell Carcinoma After Vismodegib Discontinuation: A Multicenter French Study of 116 Patients. Journal of Clinical Oncology, 2019, 37, 3275-3282.	1.6	41
39	Postzygotic inactivating mutations of RHOA cause a mosaic neuroectodermal syndrome. Nature Genetics, 2019, 51, 1438-1441.	21.4	25
40	Biallelic pathogenic variants in the lanosterol synthase gene LSS involved in the cholesterol biosynthesis cause alopecia with intellectual disability, a rare recessive neuroectodermal syndrome. Genetics in Medicine, 2019, 21, 2025-2035.	2.4	40
41	Secondary actionable findings identified by exome sequencing: expected impact on the organisation of care from the study of 700 consecutive tests. European Journal of Human Genetics, 2019, 27, 1197-1214.	2.8	18
42	Topical sirolimus 0.1% for treating cutaneous microcystic lymphatic malformations in children and adults (TOPICAL): protocol for a multicenter phase 2, within-person, randomized, double-blind, vehicle-controlled clinical trial. Trials, 2019, 20, 739.	1.6	17
43	Lysosomal Signaling Licenses Embryonic Stem Cell Differentiation via Inactivation of Tfe3. Cell Stem Cell, 2019, 24, 257-270.e8.	11.1	97
44	2.5 years' experience of GeneMatcher data-sharing: a powerful tool for identifying new genes responsible for rare diseases. Genetics in Medicine, 2019, 21, 1657-1661.	2.4	14
45	Large International Validation of ABSIS and PDAI Pemphigus Severity Scores. Journal of Investigative Dermatology, 2019, 139, 31-37.	0.7	55
46	Safety and efficacy of low-dose sirolimus in the PIK3CA-related overgrowth spectrum. Genetics in Medicine, 2019, 21, 1189-1198.	2.4	115
47	Clinical and haemodynamic risk factors associated with discrepancies in lower limb length with capillary malformations: data from the national paediatric French cohort CONAPE. British Journal of Dermatology, 2018, 178, 520-526.	1.5	3
48	Clinical whole-exome sequencing for the diagnosis of rare disorders with congenital anomalies and/or intellectual disability: substantial interest of prospective annual reanalysis. Genetics in Medicine, 2018, 20, 645-654.	2.4	146
49	The oculoauriculofrontonasal syndrome: Further clinical characterization and additional evidence suggesting a nontraditional mode of inheritance. American Journal of Medical Genetics, Part A, 2018, 176, 2740-2750.	1.2	6
50	Extending the <i>ALDH18A1</i> clinical spectrum to severe autosomal recessive fetal cutis laxa with corpus callosum agenesis. American Journal of Medical Genetics, Part A, 2018, 176, 2509-2512.	1.2	9
51	Tongue psoriasis: Clinical aspects and analysis of epidemiological associations in 313 children, with a systematic literature review. Annales De Dermatologie Et De Venereologie, 2018, 145, 578-586.	1.0	5
52	LB1539 Genotype-first phenotyping of 32 patients with post-zygotic GNAQ or GNA11 mutations. Journal of Investigative Dermatology, 2018, 138, B12.	0.7	0
53	Search for RASA1 Variants in Capillary Malformations of the Legs in 113 Children: Results from the French National Paediatric Cohort CONAPE. Acta Dermato-Venereologica, 2018, 98, 251-255.	1.3	3
54	Treatment of voluminous and complicated superficial slow-flow vascular malformations with sirolimus (PERFORMUS): protocol for a multicenter phase 2 trial with a randomized observational-phase design. Trials, 2018, 19, 340.	1.6	21

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55	Mosaicâ€activating <i>FGFR2</i> mutation in two fetuses with papillomatous pedunculated sebaceous naevus. British Journal of Dermatology, 2017, 176, 204-208.	1.5	23
56	Molecular diagnosis of PIK3CA-related overgrowth spectrum (PROS) in 162 patients and recommendations for genetic testing. Genetics in Medicine, 2017, 19, 989-997.	2.4	90
57	Sensitivity and specificity of BP180 NC16A enzyme-linked immunosorbent assay for the diagnosis of pemphigoid gestationis. Journal of the American Academy of Dermatology, 2017, 76, 560-562.	1.2	27
58	Lack of interaction between NEMO and SHARPIN impairs linear ubiquitination and NF-κB activation and leads to incontinentia pigmenti. Journal of Allergy and Clinical Immunology, 2017, 140, 1671-1682.e2.	2.9	13
59	Mosaicism for a KITLG Mutation in Linear and Whorled Nevoid Hypermelanosis. Journal of Investigative Dermatology, 2017, 137, 1575-1578.	0.7	18
60	Atypical dermal melanocytosis: a diagnostic clue in constitutional mismatch repair deficiency syndrome. British Journal of Dermatology, 2017, 177, e185-e186.	1.5	7
61	First-line rituximab combined with short-term prednisone versus prednisone alone for the treatment of pemphigus (Ritux 3): a prospective, multicentre, parallel-group, open-label randomised trial. Lancet, The, 2017, 389, 2031-2040.	13.7	438
62	616 Postzygotic mutations of RHOA cause a mosaic neuroectodermal syndrome. Journal of Investigative Dermatology, 2017, 137, S298.	0.7	0
63	Germline Loss-of-Function Mutations in EPHB4 Cause a Second Form of Capillary Malformation-Arteriovenous Malformation (CM-AVM2) Deregulating RAS-MAPK Signaling. Circulation, 2017, 136, 1037-1048.	1.6	204
64	Pain and quality of life evaluation in patients with localized epidermolysis bullosa simplex. Orphanet Journal of Rare Diseases, 2017, 12, 119.	2.7	31
65	The scalp hair collar and tuft signs: A retrospective multicenter study of 78 patients with a systematic review of the literature. Journal of the American Academy of Dermatology, 2017, 76, 478-487.	1.2	19
66	Mutations in ACTRT1 and its enhancer RNA elements lead to aberrant activation of Hedgehog signaling in inherited and sporadic basal cell carcinomas. Nature Medicine, 2017, 23, 1226-1233.	30.7	59
67	Follow-up of patients with complete remission of locally advanced basal cell carcinoma treated with vismodegib after treatment discontinuation: A retrospective multicentric French study Journal of Clinical Oncology, 2017, 35, 9535-9535.	1.6	4
68	Methotrexate in Severe Childhood Alopecia Areata: Long-term Follow-up. Acta Dermato-Venereologica, 2016, 96, 102-103.	1.3	26
69	Active tuberculosis in psoriasis patients treated with <scp>TNF</scp> antagonists: a French nationwide retrospective study. Journal of the European Academy of Dermatology and Venereology, 2016, 30, 1336-1341.	2.4	16
70	Calculation of cutâ€off values based on the Autoimmune Bullous Skin Disorder Intensity Score () Tj ETQq0 0 0 for defining moderate, significant and extensive types of pemphigus. British Journal of Dermatology,	rgBT /Overl	ock 10 Tf 50 1 68
71	2016, 175, 142-149. Mosaic Activating Mutations in GNA11 and GNAQ Are Associated with Phakomatosis Pigmentovascularis and Extensive Dermal Melanocytosis. Journal of Investigative Dermatology, 2016, 136, 770-778.	0.7	144
72	151 Postzygotic KITLG mutation in a congenital non-progressive linear nevoid hyperpigmentation. Journal of Investigative Dermatology, 2016, 136, S186.	0.7	0

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73	186 Mutational spectrum in PIK3CA -Related Overgrowth Spectrum (PROS) and recommendations for molecular testing. Journal of Investigative Dermatology, 2016, 136, S192.	0.7	1
74	A Typical Vascular and Pigmentary Dermoscopic Pattern of Capillary Malformations in Capillary Malformation–Arteriovenous Malformation Syndrome: Report of Four Cases. Pediatric Dermatology, 2016, 33, e337-41.	0.9	17
75	Neonatal Cutis Laxa and Hypertrichosis Lanuginosa in Sotos Syndrome. Pediatric Dermatology, 2016, 33, e351-e352.	0.9	1
76	Mutations in the <i>ERCC2</i> (<i>XPD</i>) gene associated with severe fetal ichthyosis and dysmorphic features. Prenatal Diagnosis, 2016, 36, 1276-1279.	2.3	5
77	Integrating longitudinal serum IL-17 and IL-23 follow-up, along with autoantibodies variation, contributes to predict bullous pemphigoid outcome. Scientific Reports, 2016, 5, 18001.	3.3	66
78	Etiologies and prognostic factors of leukocytoclastic vasculitis with skin involvement. Medicine (United States), 2016, 95, e4238.	1.0	36
79	Oral epigallocatechin-3-gallate for treatment of dystrophic epidermolysis bullosa: a multicentre, randomized, crossover, double-blind, placebo-controlled clinical trial. Orphanet Journal of Rare Diseases, 2016, 11, 31.	2.7	13
80	Postzygotic BRAF p.Lys601Asn Mutation in Phacomatosis Pigmentokeratotica with Woolly Hair Nevus and Focal Cortical Dysplasia. Journal of Investigative Dermatology, 2016, 136, 1060-1062.	0.7	15
81	Genetic Testing for Melanoma—Where Are We With Moderate-Penetrance Genes?. JAMA Dermatology, 2016, 152, 375.	4.1	2
82	Reliability, validity and feasibility of nail ultrasonography in psoriatic arthritis. Joint Bone Spine, 2016, 83, 539-544.	1.6	18
83	Psoriasis and obesity in French children: a case-control, multicentre study. British Journal of Dermatology, 2015, 172, 1593-1600.	1.5	41
84	A prospective study of risk for Sturge-Weber syndrome in children with upper facial port-wine stain. Journal of the American Academy of Dermatology, 2015, 72, 473-480.	1.2	77
85	Oral erythromycin therapy in epidermolysis bullosa simplex generalized severe. British Journal of Dermatology, 2015, 173, 563-564.	1.5	7
86	A Randomized, Controlled Trial of Oral Propranolol in Infantile Hemangioma. New England Journal of Medicine, 2015, 372, 735-746.	27.0	601
87	Burden of Inherited Ichthyosis: A French National Survey. Acta Dermato-Venereologica, 2015, 95, 326-328.	1.3	34
88	Severe Xâ€linked chondrodysplasia punctata in nine new female fetuses. Prenatal Diagnosis, 2015, 35, 675-684.	2.3	15
89	SPRED1, a RAS MAPK pathway inhibitor that causes Legius syndrome, is a tumour suppressor downregulated in paediatric acute myeloblastic leukaemia. Oncogene, 2015, 34, 631-638.	5.9	47
90	Search for <i><scp>ReCQL4</scp></i> mutations in 39 patients genotyped for suspected Rothmund–Thomson/Ballerâ€Gerold syndromes. Clinical Genetics, 2015, 87, 244-251.	2.0	22

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91	Autosomal-recessive SASH1 variants associated with a new genodermatosis with pigmentation defects, palmoplantar keratoderma and skin carcinoma. European Journal of Human Genetics, 2015, 23, 957-962.	2.8	39
92	Successful switch to dabrafenib after vemurafenib-induced toxic epidermal necrolysis. British Journal of Dermatology, 2015, 172, 1454-1455.	1.5	36
93	Factors Associated with Impaired Quality of Life in Adult Patients Suffering from Ichthyosis. Acta Dermato-Venereologica, 2014, 94, 344-346.	1.3	33
94	Prospective Study of the Evolution of Blood Lymphoid Immune Parameters during Dacarbazine Chemotherapy in Metastatic and Locally Advanced Melanoma Patients. PLoS ONE, 2014, 9, e105907.	2.5	14
95	Diagnostic value of nail examination in Hailey-Hailey disease. European Journal of Dermatology, 2014, 24, 628-629.	0.6	5
96	Next-Generation Sequencing of Nevus Spilusâ€"Type Congenital Melanocytic Nevus: Exquisite Genotypeâ€"Phenotype Correlation in Mosaic RASopathies. Journal of Investigative Dermatology, 2014, 134, 2658-2660.	0.7	40
97	Aplasia cutis congenita with dystrophic epidermolysis bullosa: clinical and mutational study. British Journal of Dermatology, 2014, 170, 901-906.	1.5	30
98	Juvenile Xanthogranuloma and Nevus Anemicus in the Diagnosis of Neurofibromatosis Type 1. JAMA Dermatology, 2014, 150, 42.	4.1	63
99	Clinical and Immunologic Factors Associated With Bullous Pemphigoid Relapse During the First Year of Treatment. JAMA Dermatology, 2014, 150, 25.	4.1	71
100	Prevalence of inherited ichthyosis in France: a study using capture-recapture method. Orphanet Journal of Rare Diseases, 2014, 9, 1.	2.7	87
101	Diversity of the clinical presentation of the MMR gene biallelic mutations. Familial Cancer, 2014, 13, 131-135.	1.9	18
102	Treatment of port wine stains with pulsed dye laser and topical timolol: a multicenter randomized controlled trial. British Journal of Dermatology, 2014, 170, 1350-1353.	1.5	29
103	Treatment of granuloma annulare with the 595â€nm pulsed dye laser, a multicentre retrospective study with longâ€term followâ€up. Journal of the European Academy of Dermatology and Venereology, 2013, 27, 785-788.	2.4	9
104	Validation of a clinical evaluation score for irritative dermatitis: SCOREPI. Journal of the European Academy of Dermatology and Venereology, 2013, 27, 1138-1142.	2.4	2
105	Systematic molecular and cytogenetic screening of 100 patients with marfanoid syndromes and intellectual disability. Clinical Genetics, 2013, 84, 507-521.	2.0	23
106	High cumulative risks of cancer in patients with <i>PTEN</i> hamartoma tumour syndrome. Journal of Medical Genetics, 2013, 50, 255-263.	3.2	290
107	Genotypes and Phenotypes of 162 Families with aGlomulinMutation. Molecular Syndromology, 2013, 4, 157-64.	0.8	55
108	Dacarbazine-Mediated Upregulation of NKG2D Ligands on Tumor Cells Activates NK and CD8 T Cells and Restrains Melanoma Growth. Journal of Investigative Dermatology, 2013, 133, 499-508.	0.7	75

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109	Flexural Agminated Eruptive Nevi in Langerhans Cell Histiocytosis. JAMA Dermatology, 2013, 149, 635.	4.1	10
110	Quinoline Yellow dyeâ€induced fixed foodâ€andâ€drug eruption. Contact Dermatitis, 2013, 68, 187-188.	1.4	14
111	Cutaneous B-cell lymphoblastic lymphoma in children: A rare diagnosis. Journal of the American Academy of Dermatology, 2012, 66, 51-57.	1.2	38
112	Systematic search for neutropenia should be part of the first screening in patients with poikiloderma. European Journal of Medical Genetics, 2012, 55, 8-11.	1.3	22
113	Familial melanoma: Clinical factors associated with germline CDKN2A mutations according to the number of patients affected by melanoma in a family. Journal of the American Academy of Dermatology, 2012, 67, 1257-1264.e2.	1.2	26
114	In-Frame Mutations in Exon 1 of SKI Cause Dominant Shprintzen-Goldberg Syndrome. American Journal of Human Genetics, 2012, 91, 950-957.	6.2	95
115	Propranolol for treatment of ulcerated infantile hemangiomas. Journal of the American Academy of Dermatology, 2011, 64, 827-832.	1.2	98
116	A SUMOylation-defective MITF germline mutation predisposes to melanoma and renal carcinoma. Nature, 2011, 480, 94-98.	27.8	466
117	Dramatic response of recalcitrant warts as a side effect of colorectal cancer treatment with oral capecitabine. European Journal of Dermatology, 2011, 21, 789-790.	0.6	0
118	Pathway-Based Analysis of a Melanoma Genome-Wide Association Study: Analysis of Genes Related to Tumour-Immunosuppression. PLoS ONE, 2011, 6, e29451.	2.5	18
119	Efficacy and tolerability of methotrexate in severe childhood alopecia areata. British Journal of Dermatology, 2011, 165, 407-410.	1.5	69
120	Congenital Plaqueâ€Type Glomuvenous Malformations Associated with Fetal Pleural Effusion and Ascites. Pediatric Dermatology, 2011, 28, 528-531.	0.9	15
121	Endemic Treponemal Infections in International Adoptees and Immigrant Children: How Common are they?. Pediatric Dermatology, 2011, 28, 214-215.	0.9	1
122	Medial Fronto-Facial Capillary Malformations. Journal of Pediatrics, 2011, 158, 836-841.	1.8	12
123	Reconstruction of hyperspectral cutaneous data from an artificial neural network-based multispectral imaging system. Computerized Medical Imaging and Graphics, 2011, 35, 85-88.	5.8	48
124	Validation of a 2D multispectral camera: application to dermatology/cosmetology on a population covering five skin phototypes. , 2011 , , .		9
125	Novel FH mutations in families with hereditary leiomyomatosis and renal cell cancer (HLRCC) and patients with isolated type 2 papillary renal cell carcinoma. Journal of Medical Genetics, 2011, 48, 226-234.	3.2	116
126	Multiple anaemic macules and diffuse erythrocyanosis revealing mixed cryoglobulinaemia. European Journal of Dermatology, 2011, 21, 269-270.	0.6	5

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127	â€~Matchstick' eyebrow hairs: a dermoscopic clue to the diagnosis of Netherton syndrome. Journal of the European Academy of Dermatology and Venereology, 2010, 24, 740-741.	2.4	24
128	Source separation on hyperspectral cube applied to dermatology. , 2010, , .		4
129	Anetoderma of Prematurity: An latrogenic Consequence of Neonatal Intensive Care. Archives of Dermatology, 2010, 146, 565-7.	1.4	18
130	Efficacy of Propranolol in Hepatic Infantile Hemangiomas with Diffuse Neonatal Hemangiomatosis. Journal of Pediatrics, 2010, 157, 340-342.	1.8	148
131	Linear atrophoderma of moulin associated with antinuclear antibodies. Journal of the European Academy of Dermatology and Venereology, 2010, 24, 108-109.	2.4	9
132	Dermoscopy of Longitudinal Leukonychia in Hailey-Hailey Disease. Archives of Dermatology, 2010, 146, 1204.	1.4	14
133	Segmental and nonsegmental childhood vitiligo has distinct clinical characteristics: A prospective observational study. Journal of the American Academy of Dermatology, 2010, 62, 945-949.	1.2	69
134	Revised nomenclature and classification of inherited ichthyoses: Results of the First Ichthyosis Consensus Conference in Sorà ze 2009. Journal of the American Academy of Dermatology, 2010, 63, 607-641.	1.2	610
135	Molecular apocrine differentiation is a common feature of breast cancer in patients with germline PTEN mutations. Breast Cancer Research, 2010, 12, R63.	5.0	54
136	Functional, structural, and genetic evaluation of 20 <i>CDKN2A</i> germ line mutations identified in melanoma-prone families or patients. Human Mutation, 2009, 30, 564-574.	2.5	38
137	Fixed eruption due to quinine contained in tonic water: positive patchâ€testing. Contact Dermatitis, 2009, 61, 242-244.	1.4	26
138	Polyethylenimine-mediated in vivo gene transfer of a transmembrane superantigen fusion construct inhibits B16 murine melanoma growth. Cancer Gene Therapy, 2008, 15, 742-749.	4.6	11
139	The contribution of large genomic deletions at the CDKN2A locus to the burden of familial melanoma. British Journal of Cancer, 2008, 99, 364-370.	6.4	25
140	Remembering Marc Larrègue (1935–2007). Pediatric Dermatology, 2008, 25, 581-583.	0.9	1
141	Prospective Multicenter Study of Pegylated Liposomal Doxorubicin Treatment in Patients With Advanced or Refractory Mycosis Fungoides or Sézary Syndrome. Archives of Dermatology, 2008, 144, 727-33.	1.4	88
142	Anétodermie du prématuré. Annales De Dermatologie Et De Venereologie, 2007, 134, 602.	1.0	0
143	Granulomes cutanés idiopathiques au cours d'une ataxie-télangiectasie. Annales De Dermatologie Et De Venereologie, 2007, 134, 617.	1.0	0
144	Vitamine D et soleil : risques et bénéfices chez l'enfant. Annales De Dermatologie Et De Venereologie, 2007, 134, 14-17.	1.0	2

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145	Diagnostic and predictive value of skin testing in platinum salt hypersensitivity. Journal of Allergy and Clinical Immunology, 2007, 119, 726-730.	2.9	111
146	Segmental overgrowth, lipomatosis, arteriovenous malformation and epidermal nevus (SOLAMEN) syndrome is related to mosaic PTEN nullizygosity. European Journal of Human Genetics, 2007, 15, 767-773.	2.8	129
147	The prognostic impact of the extent of lymph node dissection in patients with stage III melanoma. European Journal of Surgical Oncology, 2006, 32, 790-794.	1.0	33
148	Features associated with germline CDKN2A mutations: a GenoMEL study of melanoma-prone families from three continents. Journal of Medical Genetics, 2006, 44, 99-106.	3.2	350
149	Bi-acromial Dimples: A Series of Seven Cases. Pediatric Dermatology, 2005, 22, 412-414.	0.9	18
150	Hereditary mucoepithelial dysplasia: clinical, ultrastructural and genetic study of eight patients and literature review. British Journal of Dermatology, 2005, 153, 310-318.	1.5	37
151	Mutational spectrum of NSDHL in CHILD syndrome. Journal of Medical Genetics, 2005, 42, e17-e17.	3.2	65
152	Childhood Dermatosis due to Microchimerism. Dermatology, 2005, 211, 388-389.	2.1	8
153	Lamin A and ZMPSTE24 (FACE-1) defects cause nuclear disorganization and identify restrictive dermopathy as a lethal neonatal laminopathy. Human Molecular Genetics, 2004, 13, 2493-2503.	2.9	325
154	Primary cutaneous B-cell lymphoma mimicking pyoderma gangrenosum: first-line treatment with rituximab. British Journal of Dermatology, 2004, 151, 250-252.	1.5	9
155	Claudin-1 gene mutations in neonatal sclerosing cholangitis associated with ichthyosis: A tight junction disease. Gastroenterology, 2004, 127, 1386-1390.	1.3	378
156	Retinal angioma in a patient with Cowden disease. American Journal of Ophthalmology, 2003, 135, 400-402.	3.3	20
157	Anetoderma. American Journal of Dermatopathology, 2002, 24, 118-129.	0.6	52
158	Microchimerism from a dizygotic twin in juvenile ulcerative lichen planus. Lancet, The, 2002, 359, 1861-1862.	13.7	22
159	Linear and whorled nevoid hypermelanosis with bilateral giant cerebral aneurysms. American Journal of Medical Genetics Part A, 2002, 112, 95-98.	2.4	9
160	Homozygosity Mapping of a Locus for a Novel Syndromic Ichthyosis to Chromosome 3q27–q28. Journal of Investigative Dermatology, 2002, 119, 70-76.	0.7	58
161	Absence of Lisch Nodules in Sporadic Neurofibromatosis Type 1 May Reflect Somatic Mosaicism. Archives of Dermatology, 2002, 138, 839-840.	1.4	3
162	Pitfalls in Clinical Diagnosis of Female Carriers of X-linked Hypohidrotic Ectodermal Dysplasia. Archives of Dermatology, 2002, 138, 1256-1258.	1.4	6

#	Article	IF	Citations
163	A Novel Missense Mutation, A118E, in the Helix Initiation Motif of the Type II Hair Cortex Keratin hHb6, Causing Monilethrix. Human Heredity, 2000, 50, 322-324.	0.8	18
164	Linkage of Marie-Unna hypotrichosis locus to chromosome 8p21 and exclusion of 10 genes including the hairless gene by mutation analysis. European Journal of Human Genetics, 2000, 8, 273-279.	2.8	30
165	Genomic rearrangement in NEMO impairs NF-κB activation and is a cause of incontinentia pigmenti. Nature, 2000, 405, 466-472.	27.8	709
166	Génétique de la dermatite atopique. Revue Francaise D'allergologie Et D'immunologie Clinique, 2000, 40, 60-64.	0.1	0
167	œdème aigu hémorragique du nourrisson. Journal De Pediatrie Et De Puericulture, 2000, 13, 270-274.	0.0	0
168	Two unusual tumors in a patient with xeroderma pigmentosum: atypical fibroxanthoma and basosquamous carcinoma. Journal of Cutaneous Pathology, 1999, 26, 430-435.	1.3	35
169	Skin elastic fibers in Williams syndrome. American Journal of Medical Genetics Part A, 1999, 87, 134-138.	2.4	28
170	Atypical Lyme borreliosis in an HIV-infected man. British Journal of Dermatology, 1997, 137, 437-439.	1.5	11
171	Atypical Lyme borreliosis in an HIV-infected man. British Journal of Dermatology, 1997, 137, 437-439.	1.5	11
172	Atypical Lyme borreliosis in an HIV-infected man. British Journal of Dermatology, 1997, 137, 437-9.	1.5	3
173	The Gene for Bazex-Dupré-Christol Syndrome Maps to Chromosome Xq. Journal of Investigative Dermatology, 1995, 105, 87-91.	0.7	83
174	Incontinentia Pigmenti: Late Sequelae and Genotypic Diagnosis: A Three-Generation Study of Four Patients. Pediatric Dermatology, 1995, 12, 107-111.	0.9	11
175	Le prurit : aspects pathologiques cliniques et th \tilde{A} ©rapeutiques. Revue Francaise D'allergologie Et D'immunologie Clinique, 1995, 35, 613.	0.1	0
176	The gene for the familial form of incontinentia pigmenti (IP2) maps to the distal part of Xq28. Human Molecular Genetics, 1994, 3, 273-278.	2.9	71
177	Bazex-Dupré-Christol syndrome: A possible diagnosis for basal cell carcinomas, coarse sparse hair, and milia. American Journal of Medical Genetics Part A, 1993, 45, 786-786.	2.4	21