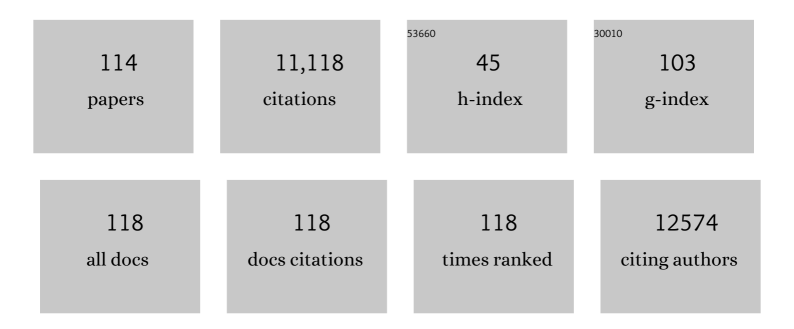
David P Kelsell

List of Publications by Year in descending order

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| # | Article | IF | CITATIONS |
|----|---|-----|-----------|
| 1 | iRHOM2: A Regulator of Palmoplantar Biology, Inflammation, and Viral Susceptibility. Journal of Investigative Dermatology, 2021, 141, 722-726. | 0.3 | 7 |
| 2 | Modelling of temporal exposure to the ambient environment and eczema severity. JID Innovations, 2021, 2, 100062. | 1.2 | 1 |
| 3 | Celebrating the 50th Anniversary of ESDR. Journal of Investigative Dermatology, 2020, 140, S145-S146. | 0.3 | Ο |
| 4 | The Future of ESDR. Journal of Investigative Dermatology, 2020, 140, S192-S193. | 0.3 | 0 |
| 5 | Clinical variability of the <i>GJB4:</i> c.35GÂ>ÂA gene variant <i>:</i> a study of a large Brazilian erythrokeratodermia pedigree. International Journal of Dermatology, 2020, 59, 722-725. | 0.5 | 2 |
| 6 | 3D model of harlequin ichthyosis reveals inflammatory therapeutic targets. Journal of Clinical Investigation, 2020, 130, 4798-4810. | 3.9 | 31 |
| 7 | Definition and treatment of arrhythmogenic cardiomyopathy: an updated expert panel report. European Journal of Heart Failure, 2019, 21, 955-964. | 2.9 | 84 |
| 8 | Lossâ€ofâ€function desmoplakin I and II mutations underlie dominant arrhythmogenic cardiomyopathy with a hair and skin phenotype. British Journal of Dermatology, 2019, 180, 1114-1122. | 1.4 | 41 |
| 9 | A Novel Mechanism for Activation of GLI1 by Nuclear SMO That Escapes Anti-SMO Inhibitors. Cancer Research, 2018, 78, 2577-2588. | 0.4 | 12 |
| 10 | Cellular biomechanics impairment in keratinocytes is associated with a C-terminal truncated desmoplakin: An atomic force microscopy investigation. Micron, 2018, 106, 27-33. | 1.1 | 8 |
| 11 | Recessive Mutation in FAM83G Associated with Palmoplantar Keratoderma and Exuberant Scalp Hair. Journal of Investigative Dermatology, 2018, 138, 984-987. | 0.3 | 10 |
| 12 | p63 is a key regulator of iRHOM2 signalling in the keratinocyte stress response. Nature Communications, 2018, 9, 1021. | 5.8 | 23 |
| 13 | A novel de novo activating mutation in STAT3 identified in a patient with common variable immunodeficiency (CVID). Clinical Immunology, 2018, 187, 132-136. | 1.4 | 19 |
| 14 | Exome Sequencing and Rare Variant Analysis RevealsÂMultiple Filaggrin Mutations in BangladeshiÂFamilies with Atopic Eczema andÂAdditional Risk Genes. Journal of Investigative Dermatology, 2018, 138, 2674-2677. | 0.3 | 37 |
| 15 | Rhomboid family member 2 regulates cytoskeletal stress-associated Keratin 16. Nature Communications, 2017, 8, 14174. | 5.8 | 36 |
| 16 | A profile of lipid dysregulation in harlequin ichthyosis. British Journal of Dermatology, 2017, 177, e217-e219. | 1.4 | 2 |
| 17 | Biallelic Mutations in KDSR Disrupt Ceramide Synthesis and Result in aÂSpectrum of Keratinization Disorders Associated with Thrombocytopenia. Journal of Investigative Dermatology, 2017, 137, 2344-2353. | 0.3 | 53 |
| 18 | Cardiomyopathy diagnosed in the eldest child harbouring p.S24X mutation in <i>JUP</i> . British Journal of Dermatology, 2016, 175, 644-646. | 1.4 | 10 |

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|----|--|-----|-----------|
| 19 | ADAM17/EGFR axis promotes transglutaminase-dependent skin barrier formation through phospholipase C γ1 and protein kinase C pathways. Scientific Reports, 2016, 6, 39780. | 1.6 | 18 |
| 20 | Loss-of-Function Mutations in SERPINB8 Linked to Exfoliative Ichthyosis with Impaired Mechanical Stability of Intercellular Adhesions. American Journal of Human Genetics, 2016, 99, 430-436. | 2.6 | 27 |
| 21 | Cover image: Unpeeling the layers of harlequin ichthyosis. British Journal of Dermatology, 2016, 174, 1160-1161. | 1.4 | 1 |
| 22 | New ANTXR1 Gene Mutation for GAPO Syndrome: A Case Report. Molecular Syndromology, 2016, 7, 160-163. | 0.3 | 9 |
| 23 | DNAJC21 Mutations Link a Cancer-Prone Bone Marrow Failure Syndrome to Corruption in 60S Ribosome Subunit Maturation. American Journal of Human Genetics, 2016, 99, 115-124. | 2.6 | 85 |
| 24 | Genomic Landscape of Esophageal Squamous Cell Carcinoma inÂa Japanese Population. Gastroenterology, 2016, 150, 1171-1182. | 0.6 | 265 |
| 25 | Tylosis with oesophageal cancer: Diagnosis, management and molecular mechanisms. Orphanet Journal of Rare Diseases, 2015, 10, 126. | 1.2 | 55 |
| 26 | Mutations inEDAandEDARGenes in a Large Mexican Hispanic Cohort with Hypohidrotic Ectodermal Dysplasia. Annals of Dermatology, 2015, 27, 474. | 0.3 | 10 |
| 27 | Cell Cycle- and Cancer-Associated Gene Networks Activated by Dsg2: Evidence of Cystatin A Deregulation and a Potential Role in Cell-Cell Adhesion. PLoS ONE, 2015, 10, e0120091. | 1.1 | 22 |
| 28 | Loss-of-Function Mutations in CAST Cause Peeling Skin, Leukonychia, Acral Punctate Keratoses, Cheilitis, and Knuckle Pads. American Journal of Human Genetics, 2015, 96, 440-447. | 2.6 | 36 |
| 29 | iASPP, a previously unidentified regulator of desmosomes, prevents arrhythmogenic right ventricular cardiomyopathy (ARVC)-induced sudden death. Proceedings of the National Academy of Sciences of the United States of America, 2015, 112, E973-E981. | 3.3 | 37 |
| 30 | A severe collodion phenotype in the newborn period associated with a homozygous missense mutation in <i>ALOX12B</i> . British Journal of Dermatology, 2015, 173, 285-287. | 1.4 | 2 |
| 31 | Evolution of Electrocardiographic and Structural Features Over 3 Decades in Arrhythmogenic Cardiomyopathy. Circulation, 2015, 131, 2233-2235. | 1.6 | 2 |
| 32 | A novel frameshift MSX1 mutation in a Saudi family with autosomal dominant premolar and third molar agenesis. Archives of Oral Biology, 2015, 60, 982-988. | 0.8 | 12 |
| 33 | Deletions in the cytoplasmic domain of iRhom1 and iRhom2 promote shedding of the TNF receptor by the protease ADAM17. Science Signaling, 2015, 8, ra109. | 1.6 | 60 |
| 34 | Novel ABCA12 mutations in harlequin ichthyosis: A journey from photo diagnosis to prenatal diagnosis. Gene, 2015, 556, 254-256. | 1.0 | 3 |
| 35 | Discovery in Genetic Skin Disease: The Impact of High Throughput Genetic Technologies. Genes, 2014, 5, 615-634. | 1.0 | 10 |
| 36 | Cryptogenic multifocal ulcerating stenosing enteritis associated with homozygous deletion mutations in cytosolic phospholipase A2-1±. Gut, 2014, 63, 96-104. | 6.1 | 62 |

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|----|--|------|-----------|
| 37 | Exoming into Rare Skin Disease: EGFR Deficiency. Journal of Investigative Dermatology, 2014, 134, 2486-2488. | 0.3 | 6 |
| 38 | iRHOM2-dependent regulation of ADAM17 in cutaneous disease and epidermal barrier function. Human Molecular Genetics, 2014, 23, 4064-4076. | 1.4 | 67 |
| 39 | Defective channels lead to an impaired skin barrier. Journal of Cell Science, 2014, 127, 4343-50. | 1.2 | 25 |
| 40 | Insights into Desmosome Biology from Inherited Human Skin Disease and Cardiocutaneous Syndromes. Cell Communication and Adhesion, 2014, 21, 129-140. | 1.0 | 27 |
| 41 | Rhomboid proteins: a role in keratinocyte proliferation and cancer. Cell and Tissue Research, 2013, 351, 301-307. | 1.5 | 23 |
| 42 | Connexin 26 facilitates gastrointestinal bacterial infection in vitro. Cell and Tissue Research, 2013, 351, 107-116. | 1.5 | 21 |
| 43 | Current insights into protease dynamics in human epithelial disease and barrier function. Cell and Tissue Research, 2013, 351, 213-215. | 1.5 | 0 |
| 44 | Mutations in AQP5, Encoding a Water-Channel Protein, Cause Autosomal-Dominant Diffuse Nonepidermolytic Palmoplantar Keratoderma. American Journal of Human Genetics, 2013, 93, 330-335. | 2.6 | 82 |
| 45 | Recessive oligodontia linked to a homozygous loss-of-function mutation in the SMOC2 gene. Archives of Oral Biology, 2013, 58, 462-466. | 0.8 | 36 |
| 46 | Targeted Sequence Capture and High-Throughput Sequencing in the Molecular Diagnosis of Ichthyosis and Other Skin Diseases. Journal of Investigative Dermatology, 2013, 133, 573-576. | 0.3 | 25 |
| 47 | A Missense Mutation in the MBTPS2 Gene Underlies the X-Linked Form of Olmsted Syndrome. Journal of Investigative Dermatology, 2013, 133, 571-573. | 0.3 | 47 |
| 48 | The DSPII splice variant is critical for desmosome-mediated HaCaT keratinocyte adhesion. Journal of Cell Science, 2012, 125, 2853-61. | 1.2 | 24 |
| 49 | Connexins in epidermal homeostasis and skin disease. Biochimica Et Biophysica Acta - Biomembranes, 2012, 1818, 1952-1961. | 1.4 | 61 |
| 50 | Cell–cell connectivity: desmosomes and disease. Journal of Pathology, 2012, 226, 158-171. | 2.1 | 153 |
| 51 | Metastatic cutaneous squamous cell carcinoma shows frequent deletion in the protein tyrosine phosphatase receptor Type D gene. International Journal of Cancer, 2012, 131, E216-26. | 2.3 | 17 |
| 52 | RHBDF2 Mutations Are Associated with Tylosis, a Familial Esophageal Cancer Syndrome. American Journal of Human Genetics, 2012, 90, 340-346. | 2.6 | 162 |
| 53 | Inflammatory Skin and Bowel Disease Linked to <i>ADAM17</i> Deletion. New England Journal of Medicine, 2011, 365, 1502-1508. | 13.9 | 285 |
| 54 | p63 Mediates an Apoptotic Response to Pharmacological and Disease-Related ER Stress in the Developing Epidermis. Developmental Cell, 2011, 21, 492-505. | 3.1 | 45 |

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|----|--|-----|-----------|
| 55 | Key functions for gap junctions in skin and hearing. Biochemical Journal, 2011, 438, 245-254. | 1.7 | 49 |
| 56 | Mutations in CSTA, Encoding Cystatin A, Underlie Exfoliative Ichthyosis and Reveal a Role for This Protease Inhibitor in Cell-Cell Adhesion. American Journal of Human Genetics, 2011, 89, 564-571. | 2.6 | 89 |
| 57 | Harlequin Ichthyosis. Archives of Dermatology, 2011, 147, 681. | 1.7 | 145 |
| 58 | SNPing at the Epidermal Barrier. Journal of Investigative Dermatology, 2011, 131, 1593-1595. | 0.3 | 11 |
| 59 | Identification and characterization of DSPIa, a novel isoform of human desmoplakin. Cell and Tissue Research, 2010, 341, 121-129. | 1.5 | 17 |
| 60 | Homozygous Mutations in the 5′ Region of the JUP Gene Result in Cutaneous Disease but Normal Heart Development in Children. Journal of Investigative Dermatology, 2010, 130, 1543-1550. | 0.3 | 49 |
| 61 | EKV mutant connexin 31 associated cell death is mediated by ER stress. Human Molecular Genetics, 2009, 18, 4734-4745. | 1.4 | 53 |
| 62 | Filaggrin mutations are associated with ichthyosis vulgaris in the Bangladeshi population. British Journal of Dermatology, 2009, 160, 1113-1115. | 1.4 | 14 |
| 63 | Premature Terminal Differentiation and a Reduction in Specific Proteases Associated with Loss of ABCA12 in Harlequin Ichthyosis. American Journal of Pathology, 2009, 174, 970-978. | 1.9 | 51 |
| 64 | Connexins in Skin Biology. , 2009, , 307-321. | | 2 |
| 65 | Novel and recurring ABCA12 mutations associated with harlequin ichthyosis: implications for prenatal diagnosis. British Journal of Dermatology, 2008, 158, 611-613. | 1.4 | 28 |
| 66 | Mutations in R-Spondin 4 (RSPO4) Underlie Inherited Anonychia. Journal of Investigative Dermatology, 2008, 128, 867-870. | 0.3 | 43 |
| 67 | A novel M163L mutation in connexin 26 causing cell death and associated with autosomal dominant hearing loss. Hearing Research, 2008, 240, 87-92. | 0.9 | 28 |
| 68 | R-Spondins in Cutaneous Biology: Nails and Cancer. Cell Cycle, 2007, 6, 895-897. | 1.3 | 11 |
| 69 | Role for WNT16B in human epidermal keratinocyte proliferation and differentiation. Journal of Cell Science, 2007, 120, 917-917. | 1.2 | 4 |
| 70 | Tissue-specific effects of wild-type and mutant connexin 31: a role in neurite outgrowth. Human Molecular Genetics, 2007, 16, 165-172. | 1.4 | 14 |
| 71 | Allelic imbalances and microdeletions affecting thePTPRDgene in cutaneous squamous cell carcinomas detected using single nucleotide polymorphism microarray analysis. Genes Chromosomes and Cancer, 2007, 46, 661-669. | 1.5 | 82 |
| 72 | Keratitis?ichthyosis?deafness syndrome: disease expression and spectrum of connexin 26 (GJB2) mutations in 14 patients. British Journal of Dermatology, 2007, 156, 1015-1019. | 1.4 | 119 |

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|----|---|-----|-----------|
| 73 | A Deafness-Associated Mutant Human Connexin 26 Improves the Epithelial Barrier In Vitro. Journal of Membrane Biology, 2007, 218, 29-37. | 1.0 | 45 |
| 74 | A novel ABCA12 mutation underlying a case of Harlequin ichthyosis. British Journal of Dermatology, 2006, 155, 204-206. | 1.4 | 18 |
| 75 | The gene encoding R-spondin 4 (RSPO4), a secreted protein implicated in Wnt signaling, is mutated in inherited anonychia. Nature Genetics, 2006, 38, 1245-1247. | 9.4 | 173 |
| 76 | Early Death from Cardiomyopathy in a Family with Autosomal Dominant Striate Palmoplantar Keratoderma and Woolly Hair Associated with a Novel Insertion Mutation in Desmoplakin. Journal of Investigative Dermatology, 2006, 126, 1651-1654. | 0.3 | 69 |
| 77 | ABCA12 Is the Major Harlequin Ichthyosis Gene. Journal of Investigative Dermatology, 2006, 126, 2408-2413. | 0.3 | 88 |
| 78 | Properties of human connexin 31, which is implicated in hereditary dermatological disease and deafness. Proceedings of the National Academy of Sciences of the United States of America, 2006, 103, 5213-5218. | 3.3 | 31 |
| 79 | Clinical and Genetic Heterogeneity of Erythrokeratoderma Variabilis. Journal of Investigative Dermatology, 2005, 125, 920-927. | 0.3 | 56 |
| 80 | Clinical Features of Multiple Cutaneous and Uterine Leiomyomatosis. Archives of Dermatology, 2005, 141, 199-206. | 1.7 | 176 |
| 81 | Connexin interaction patterns in keratinocytes revealed morphologically and by FRET analysis. Journal of Cell Science, 2005, 118, 1505-1514. | 1.2 | 45 |
| 82 | Genomewide Single Nucleotide Polymorphism Microarray Mapping in Basal Cell Carcinomas Unveils Uniparental Disomy as a Key Somatic Event. Cancer Research, 2005, 65, 8597-8603. | 0.4 | 145 |
| 83 | Mutations in ABCA12 Underlie the Severe Congenital Skin Disease Harlequin Ichthyosis. American Journal of Human Genetics, 2005, 76, 794-803. | 2.6 | 302 |
| 84 | Missense Mutations in Fumarate Hydratase in Multiple Cutaneous and Uterine Leiomyomatosis and Renal Cell Cancer. Journal of Molecular Diagnostics, 2005, 7, 437-443. | 1.2 | 56 |
| 85 | Hereditary 'white nails': a genetic and structural study. British Journal of Dermatology, 2004, 151, 65-72. | 1.4 | 26 |
| 86 | Connexin mutations in human disease. Experimental Dermatology, 2004, 13, 661-662. | 1.4 | 15 |
| 87 | p16INK4a and p14ARF Tumor Suppressor Genes Are Commonly Inactivated in Cutaneous Squamous Cell Carcinoma. Journal of Investigative Dermatology, 2004, 122, 1284-1292. | 0.3 | 145 |
| 88 | An autosomal recessive exfoliative ichthyosis with linkage to chromosome 12q13. British Journal of Dermatology, 2003, 149, 174-180. | 1.4 | 16 |
| 89 | Cellular Mechanisms of Mutant Connexins in Skin Disease and Hearing Loss. Cell Communication and Adhesion, 2003, 10, 347-351. | 1.0 | 17 |
| 90 | Defective trafficking and cell death is characteristic of skin disease-associated connexin 31 mutations. Human Molecular Genetics, 2002, 11, 2005-2014. | 1.4 | 68 |

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|-----|---|------|-----------|
| 91 | Intermediate filament–membrane attachments function synergistically with actin-dependent contacts to regulate intercellular adhesive strength. Journal of Cell Biology, 2002, 159, 1005-1017. | 2.3 | 134 |
| 92 | Functional studies of human skin disease- and deafness-associated connexin 30 mutations. Biochemical and Biophysical Research Communications, 2002, 298, 651-656. | 1.0 | 53 |
| 93 | Double jeopardy: Ras and CDK4 co-expression in skin cancer. Trends in Molecular Medicine, 2002, 8, 548. | 3.5 | 1 |
| 94 | A mutation in GJB3 is associated with recessive erythrokeratodermia variabilis (EKV) and leads to defective trafficking of the connexin 31 protein. Human Molecular Genetics, 2002, 11, 1311-1316. | 1.4 | 73 |
| 95 | Diagnosis and confirmation of epidermolytic palmoplantar keratoderma by the identification of mutations in keratin 9 using denaturing high-performance liquid chromatography. British Journal of Dermatology, 2002, 146, 952-957. | 1.4 | 18 |
| 96 | Germline mutations in FH predispose to dominantly inherited uterine fibroids, skin leiomyomata and papillary renal cell cancer. Nature Genetics, 2002, 30, 406-410. | 9.4 | 1,426 |
| 97 | Connexin Mutations in Skin Disease and Hearing Loss. American Journal of Human Genetics, 2001, 68, 559-568. | 2.6 | 156 |
| 98 | Gene expression analysis of EpiDermâ"¢ following exposure to SLS using cDNA microarrays. Toxicology in Vitro, 2001, 15, 393-398. | 1.1 | 28 |
| 99 | Multiple Epidermal Connexins are Expressed in Different Keratinocyte Subpopulations Including Connexin 31. Journal of Investigative Dermatology, 2001, 117, 958-964. | 0.3 | 138 |
| 100 | Whats new in genodermatoses?. Keio Journal of Medicine, 2001, 50, 35-38. | 0.5 | 1 |
| 101 | Association between loss of heterozygosity ofBRCA1 andBRCA2 and morphological attributes of sporadic breast cancer. International Journal of Cancer, 2000, 88, 204-208. | 2.3 | 25 |
| 102 | Mutations in GJB6 cause hidrotic ectodermal dysplasia. Nature Genetics, 2000, 26, 142-144. | 9.4 | 270 |
| 103 | Connexin mutations associated with palmoplantar keratoderma and profound deafness in a single family. European Journal of Human Genetics, 2000, 8, 141-144. | 1.4 | 73 |
| 104 | Clouston hidrotic ectodermal dysplasia (HED): genetic homogeneity, presence of a founder effect in the French Canadian population and fine genetic mapping. European Journal of Human Genetics, 2000, 8, 372-380. | 1.4 | 43 |
| 105 | Identification of a Novel Mutation R42P in the Gap Junction Protein Î ² -3 Associated with Autosomal Dominant Erythrokeratoderma Variabilis. Journal of Investigative Dermatology, 1999, 113, 1119-1122. | 0.3 | 71 |
| 106 | The palmoplantar keratodermas: much more than palms and soles. Trends in Molecular Medicine, 1999, 5, 107-113. | 2.6 | 53 |
| 107 | N-terminal deletion in a desmosomal cadherin causes the autosomal dominant skin disease striate palmoplantar keratoderma. Human Molecular Genetics, 1999, 8, 971-976. | 1.4 | 205 |
| 108 | Connexin mutations in deafness. Nature, 1998, 394, 630-631. | 13.7 | 119 |

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|-----|---|------|-----------|
| 109 | Mutations and alternative splicing of theBRCA1 gene in UK breast/ovarian cancer families. , 1997, 18, 102-110. | | 43 |
| 110 | Combined loss of BRCA1/BRCA2 in grade 3 breast carcinomas. Lancet, The, 1996, 347, 1554-1555. | 6.3 | 28 |
| 111 | Plectin deficiency results in muscular dystrophy with epidermolysis bullosa. Nature Genetics, 1996, 13, 450-457. | 9.4 | 394 |
| 112 | Identification of the breast cancer susceptibility gene BRCA2. Nature, 1995, 378, 789-792. | 13.7 | 3,230 |
| 113 | Genetic linkage studies in non-epidermolytic palmoplantar keratoderma: evidence for heterogeneity. Human Molecular Genetics, 1995, 4, 1021-1025. | 1.4 | 36 |
| 114 | Identifying Mutations in Single Gene Disorders. , 0, , 145-164. | | 0 |