J E Mellerio

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#	Paper	IF	Citations
140	The classification of inherited epidermolysis bullosa (EB): Report of the Third International Consensus Meeting on Diagnosis and Classification of EB. <i>Journal of the American Academy of Dermatology</i> , 2008 , 58, 931-50	4.5	690
139	Inherited epidermolysis bullosa: updated recommendations on diagnosis and classification. <i>Journal of the American Academy of Dermatology</i> , 2014 , 70, 1103-26	4.5	596
138	Lipoid proteinosis maps to 1q21 and is caused by mutations in the extracellular matrix protein 1 gene (ECM1). <i>Human Molecular Genetics</i> , 2002 , 11, 833-40	5.6	192
137	Consensus reclassification of inherited epidermolysis bullosa and other disorders with skin fragility. <i>British Journal of Dermatology</i> , 2020 , 183, 614-627	4	179
136	Potential of fibroblast cell therapy for recessive dystrophic epidermolysis bullosa. <i>Journal of Investigative Dermatology</i> , 2008 , 128, 2179-89	4.3	176
135	Extracutaneous manifestations and complications of inherited epidermolysis bullosa: part I. Epithelial associated tissues. <i>Journal of the American Academy of Dermatology</i> , 2009 , 61, 367-84; quiz 385-6	4.5	165
134	Extracutaneous manifestations and complications of inherited epidermolysis bullosa: part II. Other organs. <i>Journal of the American Academy of Dermatology</i> , 2009 , 61, 387-402; quiz 403-4	4.5	155
133	Harlequin ichthyosis: a review of clinical and molecular findings in 45 cases. <i>Archives of Dermatology</i> , 2011 , 147, 681-6		111
132	A consensus approach to wound care in epidermolysis bullosa. <i>Journal of the American Academy of Dermatology</i> , 2012 , 67, 904-17	4.5	103
131	A homozygous nonsense mutation within the dystonin gene coding for the coiled-coil domain of the epithelial isoform of BPAG1 underlies a new subtype of autosomal recessive epidermolysis bullosa simplex. <i>Journal of Investigative Dermatology</i> , 2010 , 130, 1551-7	4.3	101
130	An unusual N-terminal deletion of the laminin alpha3a isoform leads to the chronic granulation tissue disorder laryngo-onycho-cutaneous syndrome. <i>Human Molecular Genetics</i> , 2003 , 12, 2395-409	5.6	100
129	Fibroblast cell therapy enhances initial healing in recessive dystrophic epidermolysis bullosa wounds: results of a randomized, vehicle-controlled trial. <i>British Journal of Dermatology</i> , 2013 , 169, 102	25 ⁴ 33	92
128	Skin fragility and hypohidrotic ectodermal dysplasia resulting from ablation of plakophilin 1. <i>British Journal of Dermatology</i> , 1999 , 140, 297-307	4	92
127	Potential of Systemic Allogeneic Mesenchymal Stromal Cell Therapy for Children with Recessive Dystrophic Epidermolysis Bullosa. <i>Journal of Investigative Dermatology</i> , 2015 , 135, 2319-2321	4.3	89
126	Gastrointestinal complications of epidermolysis bullosa in children. <i>British Journal of Dermatology</i> , 2008 , 158, 1308-14	4	80
125	Management of cutaneous squamous cell carcinoma in patients with epidermolysis bullosa: best clinical practice guidelines. <i>British Journal of Dermatology</i> , 2016 , 174, 56-67	4	70
124	Prenatal diagnosis for severe inherited skin disorders: 25 yearsRexperience. <i>British Journal of Dermatology</i> , 2006 , 154, 106-13	4	64

123	The international dystrophic epidermolysis bullosa patient registry: an online database of dystrophic epidermolysis bullosa patients and their COL7A1 mutations. <i>Human Mutation</i> , 2011 , 32, 110	10 ⁴ 7 ⁷	63
122	Comparative mutation detection screening of the type VII collagen gene (COL7A1) using the protein truncation test, fluorescent chemical cleavage of mismatch, and conformation sensitive gel electrophoresis. <i>Journal of Investigative Dermatology</i> , 1999 , 113, 673-86	4.3	63
121	Desmosomal genodermatoses. British Journal of Dermatology, 2012, 166, 36-45	4	59
120	Allelic heterogeneity of dominant and recessive COL7A1 mutations underlying epidermolysis bullosa pruriginosa. <i>Journal of Investigative Dermatology</i> , 1999 , 112, 984-7	4.3	59
119	Ectodermal dysplasia-skin fragility syndrome. <i>Dermatologic Clinics</i> , 2010 , 28, 125-9	4.2	55
118	Whole-exome sequencing improves mutation detection in a diagnostic epidermolysis bullosa laboratory. <i>British Journal of Dermatology</i> , 2015 , 172, 94-100	4	54
117	HB-EGF induces COL7A1 expression in keratinocytes and fibroblasts: possible mechanism underlying allogeneic fibroblast therapy in recessive dystrophic epidermolysis Bullosa. <i>Journal of Investigative Dermatology</i> , 2011 , 131, 1771-4	4.3	54
116	Plectin defects in epidermolysis bullosa simplex with muscular dystrophy. <i>Muscle and Nerve</i> , 2007 , 35, 24-35	3.4	54
115	Moderation of phenotypic severity in dystrophic and junctional forms of epidermolysis bullosa through in-frame skipping of exons containing non-sense or frameshift mutations. <i>Journal of Investigative Dermatology</i> , 1999 , 113, 314-21	4.3	53
114	Alpha 6 beta 4 integrin abnormalities in junctional epidermolysis bullosa with pyloric atresia. <i>British Journal of Dermatology</i> , 2001 , 144, 408-14	4	52
113	Epithelial inflammation resulting from an inherited loss-of-function mutation in EGFR. <i>Journal of Investigative Dermatology</i> , 2014 , 134, 2570-2578	4.3	51
112	Germline Mutation in EXPH5 Implicates the Rab27B Effector Protein Slac2-b in Inherited Skin Fragility. <i>American Journal of Human Genetics</i> , 2012 , 91, 1115-21	11	48
111	Revertant mosaicism in recessive dystrophic epidermolysis bullosa. <i>Journal of Investigative Dermatology</i> , 2010 , 130, 1937-40	4.3	47
110	Recurrent mutations in the type VII collagen gene (COL7A1) in patients with recessive dystrophic epidermolysis bullosa. <i>Journal of Investigative Dermatology</i> , 1997 , 109, 246-9	4.3	47
109	Bone mineralization in children with epidermolysis bullosa. <i>British Journal of Dermatology</i> , 2006 , 154, 959-62	4	46
108	A spectrum of mutations in keratins K6a, K16 and K17 causing pachyonychia congenita. <i>Journal of Dermatological Science</i> , 2007 , 48, 199-205	4.3	45
107	Thalidomide in the management of epidermolysis bullosa pruriginosa. <i>British Journal of Dermatology</i> , 2005 , 152, 1332-4	4	44
106	Pyloric atresia-junctional epidermolysis bullosa syndrome: mutations in the integrin beta4 gene (ITGB4) in two unrelated patients with mild disease. <i>British Journal of Dermatology</i> , 1998 , 139, 862-71	4	42

105	Homozygous mutations in the 5R region of the JUP gene result in cutaneous disease but normal heart development in children. <i>Journal of Investigative Dermatology</i> , 2010 , 130, 1543-50	4.3	40
104	Infection and colonization in epidermolysis bullosa. <i>Dermatologic Clinics</i> , 2010 , 28, 267-9, ix	4.2	40
103	Autosomal recessive epidermolysis bullosa simplex due to loss of BPAG1-e expression. <i>Journal of Investigative Dermatology</i> , 2012 , 132, 742-4	4.3	39
102	A heterozygous frameshift mutation in the V1 domain of keratin 5 in a family with Dowling-Degos disease. <i>Journal of Investigative Dermatology</i> , 2007 , 127, 298-300	4.3	39
101	Recessive epidermolysis bullosa simplex associated with plectin mutations: infantile respiratory complications in two unrelated cases. <i>British Journal of Dermatology</i> , 1997 , 137, 898-906	4	39
100	Homozygous nonsense mutation in helix 2 of K14 causes severe recessive epidermolysis bullosa simplex. <i>Human Mutation</i> , 1998 , 11, 279-85	4.7	36
99	Severe palmo-plantar hyperkeratosis in Dowling-Meara epidermolysis bullosa simplex caused by a mutation in the keratin 14 gene (KRT14). <i>Journal of Investigative Dermatology</i> , 1998 , 111, 893-5	4.3	36
98	Congenital muscular dystrophy, myasthenic symptoms and epidermolysis bullosa simplex (EBS) associated with mutations in the PLEC1 gene encoding plectin. <i>Neuromuscular Disorders</i> , 2010 , 20, 709-	1 1 .9	35
97	Heterozygous germline missense mutation in the p63 gene underlying EEC syndrome. <i>Clinical and Experimental Dermatology</i> , 2000 , 25, 441-3	1.8	33
96	PORCN gene mutations and the protean nature of focal dermal hypoplasia. <i>British Journal of Dermatology</i> , 2009 , 160, 1103-9	4	32
95	Molecular basis of EEC (ectrodactyly, ectodermal dysplasia, clefting) syndrome: five new mutations in the DNA-binding domain of the TP63 gene and genotype-phenotype correlation. <i>British Journal of Dermatology</i> , 2010 , 162, 201-7	4	31
94	Phase I study protocol for ex vivo lentiviral gene therapy for the inherited skin disease, Netherton syndrome. <i>Human Gene Therapy Clinical Development</i> , 2013 , 24, 182-90	3.2	30
93	Target proteins in inherited and acquired blistering skin disorders. <i>Clinical and Experimental Dermatology</i> , 2006 , 31, 252-9	1.8	30
92	Rapp-Hodgkin and Hay-Wells ectodermal dysplasia syndromes represent a variable spectrum of the same genetic disorder. <i>British Journal of Dermatology</i> , 2010 , 163, 624-9	4	29
91	New glycine substitution mutations in type VII collagen underlying epidermolysis bullosa pruriginosa but the phenotype is not explained by a common polymorphism in the matrix metalloproteinase-1 gene promoter. <i>Acta Dermato-Venereologica</i> , 2009 , 89, 6-11	2.2	29
90	Autosomal dominant junctional epidermolysis bullosa. British Journal of Dermatology, 2009, 160, 1094-7	4	28
89	Identical glycine substitution mutations in type VII collagen may underlie both dominant and recessive forms of dystrophic epidermolysis bullosa. <i>Acta Dermato-Venereologica</i> , 2011 , 91, 262-6	2.2	28
88	A recurrent glycine substitution mutation, G2043R, in the type VII collagen gene (COL7A1) in dominant dystrophic epidermolysis bullosa. <i>British Journal of Dermatology</i> , 1998 , 139, 730-7	4	28

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87	Complete maternal isodisomy of chromosome 3 in a child with recessive dystrophic epidermolysis bullosa but no other phenotypic abnormalities. <i>Journal of Investigative Dermatology</i> , 2006 , 126, 2039-4	34.3	28
86	Dilated cardiomyopathy in epidermolysis bullosa: a retrospective, multicenter study. <i>Pediatric Dermatology</i> , 2010 , 27, 238-43	1.9	27
85	Fluoroscopically guided dilation of esophageal strictures in patients with dystrophic epidermolysis bullosa: long-term results. <i>American Journal of Roentgenology</i> , 2012 , 199, 208-12	5.4	26
84	ADULT ectodermal dysplasia syndrome resulting from the missense mutation R298Q in the p63 gene. <i>Clinical and Experimental Dermatology</i> , 2004 , 29, 669-72	1.8	26
83	Complete paternal uniparental isodisomy of chromosome 1 resulting in Herlitz junctional epidermolysis bullosa. <i>Clinical and Experimental Dermatology</i> , 2005 , 30, 71-4	1.8	26
82	Dilemmas in distinguishing between dominant and recessive forms of dystrophic epidermolysis bullosa. <i>British Journal of Dermatology</i> , 2003 , 149, 810-8	4	25
81	Mutations in AEC syndrome skin reveal a role for p63 in basement membrane adhesion, skin barrier integrity and hair follicle biology. <i>British Journal of Dermatology</i> , 2012 , 167, 134-44	4	22
80	Serum levels of high mobility group box 1 correlate with disease severity in recessive dystrophic epidermolysis bullosa. <i>Experimental Dermatology</i> , 2013 , 22, 433-5	4	22
79	Under-recognition of acral peeling skin syndrome: 59 new cases with 15 novel mutations. <i>British Journal of Dermatology</i> , 2014 , 171, 1206-10	4	21
78	The inversa type of recessive dystrophic epidermolysis bullosa is caused by specific arginine and glycine substitutions in type VII collagen. <i>Journal of Medical Genetics</i> , 2011 , 48, 160-7	5.8	21
77	Novel and recurring ABCA12 mutations associated with harlequin ichthyosis: implications for prenatal diagnosis. <i>British Journal of Dermatology</i> , 2008 , 158, 611-3	4	21
76	The management of general and disease specific ENT problems in children with Epidermolysis Bullosaa retrospective case note review. <i>International Journal of Pediatric Otorhinolaryngology</i> , 2007 , 71, 385-91	1.7	21
75	Molecular pathology of the cutaneous basement membrane zone. <i>Clinical and Experimental Dermatology</i> , 1999 , 24, 25-32	1.8	21
74	Clinical features and WNT10A mutations in seven unrelated cases of Schpf-Schulz-Passarge syndrome. <i>British Journal of Dermatology</i> , 2014 , 171, 1211-4	4	20
73	E210K mutation in the gene encoding the beta3 chain of laminin-5 (LAMB3) is predictive of a phenotype of generalized atrophic benign epidermolysis bullosa. <i>British Journal of Dermatology</i> , 1998 , 139, 325-31	4	20
72	Dominant dystrophic epidermolysis bullosa presenting as familial nail dystrophy. <i>Clinical and Experimental Dermatology</i> , 2001 , 26, 93-6	1.8	20
71	Neonatal diagnosis of Kindler syndrome. <i>Journal of Dermatological Science</i> , 2005 , 39, 183-5	4.3	19
70	Genotype-phenotype correlation in a large English cohort of patients with autosomal recessive ichthyosis. <i>British Journal of Dermatology</i> , 2020 , 182, 729-737	4	19

69	The challenges of meeting nutritional requirements in children and adults with epidermolysis bullosa: proceedings of a multidisciplinary team study day. <i>Clinical and Experimental Dermatology</i> , 2011 , 36, 579-83; quiz 583-4	1.8	18
68	Anaesthetic management of two different modes of delivery in patients with dystrophic epidermolysis bullosa. <i>International Journal of Obstetric Anesthesia</i> , 2008 , 17, 153-8	2.1	18
67	Immunofluorescence antigen mapping for hereditary epidermolysis bullosa. <i>Indian Journal of Dermatology, Venereology and Leprology</i> , 2012 , 78, 692-7	0.8	17
66	The missense mutation p.R1303Q in type XVII collagen underlies junctional epidermolysis bullosa resembling Kindler syndrome. <i>Journal of Investigative Dermatology</i> , 2014 , 134, 845-849	4.3	16
65	Gastrostomy tube feeding in children with epidermolysis bullosa: consideration of key issues. <i>Pediatric Dermatology</i> , 2012 , 29, 277-84	1.9	16
64	Osteopenia and osteoporosis in epidermolysis bullosa. <i>Dermatologic Clinics</i> , 2010 , 28, 353-5, xi	4.2	16
63	Epidermolysis bullosa. British Journal of Hospital Medicine (London, England: 2005), 2006, 67, 188-91	0.8	16
62	EBGene trial: patient preselection outcomes for the European GENEGRAFT ex vivo phase I/II gene therapy trial for recessive dystrophic epidermolysis bullosa. <i>British Journal of Dermatology</i> , 2020 , 182, 794-797	4	15
61	Novel and recurrent COL7A1 mutations in Chilean patients with dystrophic epidermolysis bullosa. <i>Journal of Dermatological Science</i> , 2012 , 65, 149-52	4.3	14
60	An unusual patient with Rothmund-Thomson syndrome, porokeratosis and bilateral iris dysgenesis. <i>Clinical and Experimental Dermatology</i> , 2006 , 31, 401-3	1.8	14
59	Three cases of de novo dominant dystrophic epidermolysis bullosa associated with the mutation G2043R in COL7A1. <i>Clinical and Experimental Dermatology</i> , 2001 , 26, 97-9	1.8	14
58	MBTPS2 mutation in a British pedigree with keratosis follicularis spinulosa decalvans. <i>Clinical and Experimental Dermatology</i> , 2012 , 37, 631-4	1.8	12
57	Genetics and Genodermatoses 2010 , 1-97		12
56	Frameshift mutations in the type VII collagen gene (COL7A1) in five Mexican cousins with recessive dystrophic epidermolysis bullosa. <i>British Journal of Dermatology</i> , 1998 , 138, 852-8	4	12
55	Prevalence, pathophysiology and management of itch in epidermolysis bullosa. <i>British Journal of Dermatology</i> , 2021 , 184, 816-825	4	12
54	Homozygous variegate porphyria presenting with developmental and language delay in childhood. <i>Clinical and Experimental Dermatology</i> , 2013 , 38, 737-40	1.8	11
53	Epidermolysis bullosa pruriginosa in association with lichen planopilaris. <i>Clinical and Experimental Dermatology</i> , 2009 , 34, e825-8	1.8	10
52	Prognostic implications of determining 180 kDa bullous pemphigoid antigen (BPAG2) gene/protein pathology in neonatal junctional epidermolysis bullosa. <i>British Journal of Dermatology</i> , 1998 , 138, 661-6	4	10

51	Congenital anetoderma in a preterm infant. <i>Pediatric Dermatology</i> , 2008 , 25, 626-9	1.9	10
50	Multiple dermatofibromas associated with lupus profundus. <i>Clinical and Experimental Dermatology</i> , 2005 , 30, 128-30	1.8	10
49	A recurrent COL7A1 mutation, R2814X, in British patients with recessive dystrophic epidermolysis bullosa. <i>Clinical and Experimental Dermatology</i> , 1999 , 24, 37-9	1.8	10
48	Mutations in EXPH5 result in autosomal recessive inherited skin fragility. <i>British Journal of Dermatology</i> , 2014 , 170, 196-9	4	9
47	Genitourinary tract involvement in epidermolysis bullosa. <i>Dermatologic Clinics</i> , 2010 , 28, 343-6, xi	4.2	9
46	Foot care in epidermolysis bullosa: evidence-based guideline. <i>British Journal of Dermatology</i> , 2020 , 182, 593-604	4	9
45	Recessive epidermolysis bullosa simplex associated with plectin mutations: infantile respiratory complications in two unrelated cases. <i>British Journal of Dermatology</i> , 1997 , 137, 898-906	4	9
44	Recurrent molecular abnormalities in type VII collagen in Southern Italian patients with recessive dystrophic epidermolysis bullosa. <i>Clinical and Experimental Dermatology</i> , 1999 , 24, 232-5	1.8	8
43	Meeting Report: The First Global Congress on Epidermolysis Bullosa, EB2020 London: Toward Treatment and Cure. <i>Journal of Investigative Dermatology</i> , 2020 , 140, 1681-1687	4.3	8
42	Considerations in surgical management of a Buschke-Lowenstein tumor in Netherton syndrome: A case report. <i>Pediatric Dermatology</i> , 2017 , 34, e328-e330	1.9	7
41	Recessive epidermolysis bullosa simplex associated with plectin mutations: infantile respiratory complications in two unrelated cases. <i>British Journal of Dermatology</i> , 1997 , 137, 898-906	4	7
40	A recurrent frameshift mutation in exon 19 of the type VII collagen gene (COL7A1) in Mexican patients with recessive dystrophic epidermolysis bullosa. <i>Experimental Dermatology</i> , 1999 , 8, 22-9	4	7
39	Autosomal dominant diffuse nonepidermolytic palmoplantar keratoderma due to a recurrent mutation in aquaporin-5. <i>British Journal of Dermatology</i> , 2016 , 174, 430-2	4	7
38	Mutations in EXPH5 underlie a rare subtype of autosomal recessive epidermolysis bullosa simplex. <i>British Journal of Dermatology</i> , 2016 , 174, 452-3	4	7
37	Predictive phenotyping of inherited ichthyosis by next-generation DNA sequencing. <i>British Journal of Dermatology</i> , 2017 , 176, 249-251	4	6
36	Ichthyosis Prematurity Syndrome: From Fetus to Adulthood. <i>JAMA Dermatology</i> , 2016 , 152, 1055-8	5.1	6
35	Pain, purpura and curly hairs. Clinical and Experimental Dermatology, 2013, 38, 940-2	1.8	6
34	Wound healing in epidermolysis bullosa. <i>British Journal of Dermatology</i> , 2017 , 177, e193-e195	4	6

33	Epidermolysis bullosa care in the United Kingdom. <i>Dermatologic Clinics</i> , 2010 , 28, 395-6, xiv	4.2	6
32	Histopathological features of gastrointestinal mucosal biopsy specimens in children with epidermolysis bullosa. <i>Journal of Clinical Pathology</i> , 2007 , 60, 843-4	3.9	6
31	Early-onset dermatosis papulosa nigra. British Journal of Dermatology, 2016, 174, 1148-50	4	6
30	Prenatal diagnosis of Herlitz junctional epidermolysis bullosa in nonidentical twins. <i>Clinical and Experimental Dermatology</i> , 2005 , 30, 180-2	1.8	5
29	Frequency of the CCR5 gene 32-basepair deletion in Hispanic Mexicans. <i>Clinical and Experimental Dermatology</i> , 1999 , 24, 127-9	1.8	5
28	Skin disease in Gulf war veterans. <i>QJM - Monthly Journal of the Association of Physicians</i> , 2002 , 95, 671-6	5 2.7	4
27	Homozygous Nonsense Mutation in DSC3 Resulting in Skin Fragility and Hypotrichosis. <i>Journal of Investigative Dermatology</i> , 2020 , 140, 1285-1288	4.3	4
26	Focal dermal hypoplasia: inheritance from father to daughter. <i>Clinical and Experimental Dermatology</i> , 2017 , 42, 457-459	1.8	3
25	Pityriasis rubra pilaris with histologic features of lichen nitidus. <i>Journal of the American Academy of Dermatology</i> , 2015 , 73, 336-7	4.5	3
24	Maxillary alveolar process fracture complicating intubation in a patient with epidermolysis bullosa. <i>Paediatric Anaesthesia</i> , 2009 , 19, 706-7	1.8	3
23	Dominant dystrophic epidermolysis bullosa presenting as familial nail dystrophy. <i>Clinical and Experimental Dermatology</i> , 2001 , 26, 93-96	1.8	3
22	Potential therapeutic targeting of inflammation in epidermolysis bullosa simplex. <i>British Journal of Dermatology</i> , 2019 , 180, 258-260	4	3
21	Heterogeneous addiction to transforming growth factor-beta signalling in recessive dystrophic epidermolysis bullosa-associated cutaneous squamous cell carcinoma. <i>British Journal of Dermatology</i> , 2021 , 184, 697-708	4	3
20	Homozygous nonsense mutation in helix 2 of K14 causes severe recessive epidermolysis bullosa simplex 1998 , 11, 279		3
19	A pyrexial unwell child with a papular eruption. Clinical and Experimental Dermatology, 2012, 37, 811-3	1.8	2
18	Growth Impaired Children with Epidermolysis Bullosa have Increased Serum Markers of Inflammation and Reduced Circulating IGF-1/IGFBP-3. <i>Pediatric Research</i> , 2011 , 70, 294-294	3.2	2
17	An unusual case of epidermolysis bullosa complicated by persistent oligoarticular juvenile idiopathic arthritis; lessons to be learned. <i>Pediatric Rheumatology</i> , 2011 , 9, 13	3.5	2
16	Bullous pemphigoid in a patient with suspected non-Herlitz junctional epidermolysis bullosa. <i>Clinical and Experimental Dermatology</i> , 2010 , 35, 881-4	1.8	2

LIST OF PUBLICATIONS

15	The psychological functioning of children with epidermolysis bullosa and its relationship with specific aspects of disease. <i>British Journal of Dermatology</i> , 2020 , 182, 789-790	4	2
14	Practical management of epidermolysis bullosa: consensus clinical position statement from the European Reference Network for Rare Skin Diseases. <i>Journal of the European Academy of Dermatology and Venereology</i> , 2021 , 35, 2349-2360	4.6	2
13	G45 Haematocolpos due to a transverse vaginal septum in 15 year old girl with junctional epidermolysis bullosa. <i>Archives of Disease in Childhood</i> , 2014 , 99, A19-A19	2.2	1
12	Recalcitrant generalized eruption and low alkaline phosphatase: think zinc. <i>Clinical and Experimental Dermatology</i> , 2011 , 36, 225-6	1.8	1
11	A recurrent laminin 5 mutation in British patients with lethal (Herlitz) junctional epidermolysis bullosa: evidence for a mutational hotspot rather than propagation of an ancestral allele. <i>British Journal of Dermatology</i> , 1997 , 136, 674-677	4	1
10	Generalized pustular eruption in a 5-year-old boy. Clinical and Experimental Dermatology, 2008, 33, 79-8	8 0 1.8	1
9	Dominant dystrophic epidermolysis bullosa presenting as familial nail dystrophy. <i>Clinical and Experimental Dermatology</i> , 2001 , 26, 93-96	1.8	1
8	Characteristics of children with Netherton syndrome: a review of 21 patients. <i>Journal of the European Academy of Dermatology and Venereology</i> , 2021 , 35, e466-e469	4.6	1
7	The epidemiology of epidermolysis bullosa in England and Wales: data from the national epidermolysis bullosa database <i>British Journal of Dermatology</i> , 2021 ,	4	1
6	Beta blockers for infantile haemangiomas: where should we go from here?. <i>British Journal of Dermatology</i> , 2019 , 180, 450-451	4	O
5	Neonatal aggressive systemic mastocytosis. <i>British Journal of Dermatology</i> , 2017 , 177, 1167-1168	4	
4	Heterogeneous Mutations of the Type VII Collagen Gene (COL7A1) in Dystrophic Epidermolysis Bullosa. <i>Clinical Science</i> , 2000 , 98, 18P-18P		
3	Pseudoporphyria induced by ultraviolet radiation. Australasian Journal of Dermatology, 2020, 61, 177-1	79 .3	
2	PLACK syndrome: the penny dropped. Clinical and Experimental Dermatology, 2020, 45, 1091-1092	1.8	
1	Itch and scratch: could pain be the catch?. British Journal of Dermatology, 2018, 179, 1029	4	