Marcus A Mall

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	Fibroblast Activation Protein–Specific PET/CT Imaging in Fibrotic Interstitial Lung Diseases and Lung Cancer: A Translational Exploratory Study. Journal of Nuclear Medicine, 2022, 63, 127-133.	5.0	72
2	Efficacy and safety of inhaled ENaC inhibitor BI 1265162 in patients with cystic fibrosis: BALANCE-CF 1, a randomised, phase II study. European Respiratory Journal, 2022, 59, 2100746.	6.7	5
3	Final results of the southwest German pilot study on cystic fibrosis newborn screening – Evaluation of an IRT/PAP protocol with IRT-dependent safety net. Journal of Cystic Fibrosis, 2022, 21, 422-433.	0.7	8
4	Impact of lockdown during the COVID-19 pandemic on health status in patients with cystic fibrosis: a mono-centre observational study. ERJ Open Research, 2022, 8, 00588-2021.	2.6	6
5	Complement activation induces excessive T cell cytotoxicity in severe COVID-19. Cell, 2022, 185, 493-512.e25.	28.9	122
6	Age-Related Differences in Structure and Function of Nasal Epithelial Cultures From Healthy Children and Elderly People. Frontiers in Immunology, 2022, 13, 822437.	4.8	5
7	Effects of Elexacaftor/Tezacaftor/Ivacaftor Therapy on CFTR Function in Patients with Cystic Fibrosis and One or Two <i>F508del</i> Alleles. American Journal of Respiratory and Critical Care Medicine, 2022, 205, 540-549.	5.6	78
8	A PI3KÎ ³ mimetic peptide triggers CFTR gating, bronchodilation, and reduced inflammation in obstructive airway diseases. Science Translational Medicine, 2022, 14, eabl6328.	12.4	6
9	Efficacy and safety of elexacaftor plus tezacaftor plus ivacaftor versus tezacaftor plus ivacaftor in people with cystic fibrosis homozygous for F508del-CFTR: a 24-week, multicentre, randomised, double-blind, active-controlled, phase 3b trial. Lancet Respiratory Medicine,the, 2022, 10, 267-277.	10.7	66
10	Comparison of the Lung Clearance Index in Preschool Children With Primary Ciliary Dyskinesia and Cystic Fibrosis. Chest, 2022, 162, 534-542.	0.8	11
11	Magnetic resonance imaging detects improvements of pulmonary and paranasal sinus abnormalities in response to elexacaftor/tezacaftor/ivacaftor therapy in adults with cystic fibrosis. Journal of Cystic Fibrosis, 2022, 21, 1053-1060.	0.7	39
12	Changes in Microbiome Dominance Are Associated With Declining Lung Function and Fluctuating Inflammation in People With Cystic Fibrosis. Frontiers in Microbiology, 2022, 13, .	3.5	6
13	Drug allergy to CFTR modulator therapy associated with lumacaftor-specific CD4+ T lymphocytes. Journal of Allergy and Clinical Immunology, 2021, 147, 753-756.	2.9	7
14	Safety and Efficacy of Elexacaftor/Tezacaftor/Ivacaftor for 24 Weeks or Longer in People with Cystic Fibrosis and One or More <i>F508del</i> Alleles: Interim Results of an Open-Label Phase 3 Clinical Trial. American Journal of Respiratory and Critical Care Medicine, 2021, 203, 381-385.	5.6	116
15	CFTR Modulator Therapy with Lumacaftor/Ivacaftor Alters Plasma Concentrations of Lipid-Soluble Vitamins A and E in Patients with Cystic Fibrosis. Antioxidants, 2021, 10, 483.	5.1	19
16	Therapeutic Inhibition of Cathepsin S Reduces Inflammation and Mucus Plugging in Adult βENaC-Tg Mice. Mediators of Inflammation, 2021, 2021, 1-10.	3.0	3
17	Proteases, Mucus, and Mucosal Immunity in Chronic Lung Disease. International Journal of Molecular Sciences, 2021, 22, 5018.	4.1	15
18	Potential of Intestinal Current Measurement for Personalized Treatment of Patients with Cystic Fibrosis. Journal of Personalized Medicine, 2021, 11, 384.	2.5	9

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19	Quantification of Phenotypic Variability of Lung Disease in Children with Cystic Fibrosis. Genes, 2021, 12, 803.	2.4	6
20	Chronic rhinosinusitis with nasal polyps is associated with impaired TMEM16A-mediated epithelial chloride secretion. Journal of Allergy and Clinical Immunology, 2021, 147, 2191-2201.e2.	2.9	9
21	Congenital Deletion of Nedd4-2 in Lung Epithelial Cells Causes Progressive Alveolitis and Pulmonary Fibrosis in Neonatal Mice. International Journal of Molecular Sciences, 2021, 22, 6146.	4.1	12
22	Effects of Lumacaftor–Ivacaftor on Lung Clearance Index, Magnetic Resonance Imaging, and Airway Microbiome in Phe508del Homozygous Patients with Cystic Fibrosis. Annals of the American Thoracic Society, 2021, 18, 971-980.	3.2	65
23	A Phase 3 Open-Label Study of Elexacaftor/Tezacaftor/Ivacaftor in Children 6 through 11 Years of Age with Cystic Fibrosis and at Least One <i>F508del</i> Allele. American Journal of Respiratory and Critical Care Medicine, 2021, 203, 1522-1532.	5.6	146
24	Magnetic Resonance Imaging Detects Progression of Lung Disease and Impact of Newborn Screening in Preschool Children with Cystic Fibrosis. American Journal of Respiratory and Critical Care Medicine, 2021, 204, 943-953.	5.6	41
25	Linking Fibrotic Remodeling and Ultrastructural Alterations of Alveolar Epithelial Cells after Deletion of Nedd4-2. International Journal of Molecular Sciences, 2021, 22, 7607.	4.1	5
26	Triple Therapy for Cystic Fibrosis <i>Phe508del</i> –Gating and –Residual Function Genotypes. New England Journal of Medicine, 2021, 385, 815-825.	27.0	140
27	Cross-reactive CD4 ⁺ T cells enhance SARS-CoV-2 immune responses upon infection and vaccination. Science, 2021, 374, eabh1823.	12.6	221
28	SARS-CoV-2 infection and transmission in school settings during the second COVID-19 wave: a cross-sectional study, Berlin, Germany, November 2020. Eurosurveillance, 2021, 26, .	7.0	32
29	Increased Inflammatory Markers Detected in Nasal Lavage Correlate with Paranasal Sinus Abnormalities at MRI in Adolescent Patients with Cystic Fibrosis. Antioxidants, 2021, 10, 1412.	5.1	8
30	Prevalence of SARS-CoV-2 Infections Among Students, Teachers, and Household Members During Lockdown and Split Classes in Berlin, Germany. JAMA Network Open, 2021, 4, e2127168.	5.9	9
31	Relationship between airway dysbiosis, inflammation and lung function in adults with cystic fibrosis. Journal of Cystic Fibrosis, 2021, 20, 754-760.	0.7	25
32	Reply to: Contrast Enhanced Magnetic Resonance Imaging Does Not Detect a Progression in Lung Morphological Score in Preschool Children with Cystic Fibrosis. American Journal of Respiratory and Critical Care Medicine, 2021, , .	5.6	0
33	Visualization of Ectopic Serine Protease Activity by Förster Resonance Energy Transfer-Based Reporters. ACS Chemical Biology, 2021, 16, 2174-2184.	3.4	1
34	Epigenetic reprogramming of airway macrophages promotes polarization and inflammation in muco-obstructive lung disease. Nature Communications, 2021, 12, 6520.	12.8	38
35	A Volatile and Dynamic Longitudinal Microbiome Is Associated With Less Reduction in Lung Function in Adolescents With Cystic Fibrosis. Frontiers in Cellular and Infection Microbiology, 2021, 11, 763121.	3.9	5
36	The future of cystic fibrosis care: a global perspective. Lancet Respiratory Medicine,the, 2020, 8, 65-124.	10.7	573

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37	Targeting Proteases in Cystic Fibrosis Lung Disease. Paradigms, Progress, and Potential. American Journal of Respiratory and Critical Care Medicine, 2020, 201, 141-147.	5.6	43
38	Lack of IL-1 Receptor Signaling Reduces Spontaneous Airway Eosinophilia in Juvenile Mice with Muco-Obstructive Lung Disease. American Journal of Respiratory Cell and Molecular Biology, 2020, 62, 300-309.	2.9	7
39	Cystic Fibrosis: Emergence of Highly Effective Targeted Therapeutics and Potential Clinical Implications. American Journal of Respiratory and Critical Care Medicine, 2020, 201, 1193-1208.	5.6	137
40	Comparison of Organoid Swelling and <i>In Vivo</i> Biomarkers of CFTR Function to Determine Effects of Lumacaftor–Ivacaftor in Patients with Cystic Fibrosis Homozygous for the F508del Mutation. American Journal of Respiratory and Critical Care Medicine, 2020, 202, 1589-1592.	5.6	23
41	Protease FRET Reporters Targeting Neutrophil Extracellular Traps. Journal of the American Chemical Society, 2020, 142, 20299-20305.	13.7	28
42	ENaC inhibition in cystic fibrosis: potential role in the new era of CFTR modulator therapies. European Respiratory Journal, 2020, 56, 2000946.	6.7	33
43	Multicentre feasibility of multiple-breath washout in preschool children with cystic fibrosis and other lung diseases. ERJ Open Research, 2020, 6, 00408-2020.	2.6	18
44	Compromised <scp>DNA</scp> repair is responsible for diabetesâ€associated fibrosis. EMBO Journal, 2020, 39, e103477.	7.8	49
45	Normative data for multiple breath washout outcomes in school-aged Caucasian children. European Respiratory Journal, 2020, 55, 1901302.	6.7	79
46	CRISPRi-mediated functional analysis of lung disease-associated loci at non-coding regions. NAR Genomics and Bioinformatics, 2020, 2, Iqaa036.	3.2	7
47	Studying the pathophysiology of coronavirus disease 2019: a protocol for the Berlin prospective COVID-19 patient cohort (Pa-COVID-19). Infection, 2020, 48, 619-626.	4.7	79
48	New method for rapid and dynamic quantification of elastase activity on sputum neutrophils from patients with cystic fibrosis using flow cytometry. European Respiratory Journal, 2020, 55, 1902355.	6.7	4
49	Magnetic Resonance Imaging Detects Chronic Rhinosinusitis in Infants and Preschool Children with Cystic Fibrosis. Annals of the American Thoracic Society, 2020, 17, 714-723.	3.2	23
50	The value of chest magnetic resonance imaging compared to chest radiographs with and without additional lung ultrasound in children with complicated pneumonia. PLoS ONE, 2020, 15, e0230252.	2.5	18
51	Rhinovirus Infection Is Associated With Airway Epithelial Cell Necrosis and Inflammation via Interleukin-1 in Young Children With Cystic Fibrosis. Frontiers in Immunology, 2020, 11, 596.	4.8	16
52	Pseudomonas aeruginosa Modulates the Antiviral Response of Bronchial Epithelial Cells. Frontiers in Immunology, 2020, 11, 96.	4.8	16
53	Echo Timeâ€Dependence of Observed Lung <scp>T₁</scp> in Patients With Cystic Fibrosis and Correlation With Clinical Metrics. Journal of Magnetic Resonance Imaging, 2020, 52, 1645-1654.	3.4	17
54	Neutrophil Adaptations upon Recruitment to the Lung: New Concepts and Implications for Homeostasis and Disease. International Journal of Molecular Sciences, 2020, 21, 851.	4.1	67

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55	At the forefront of cystic fibrosis Basic Science research: 16th ECFS Basic Science Conference. Journal of Cystic Fibrosis, 2020, 19, 169-170.	0.7	1
56	CRISPR-Based Adenine Editors Correct Nonsense Mutations in a Cystic Fibrosis Organoid Biobank. Cell Stem Cell, 2020, 26, 503-510.e7.	11.1	136
57	Intravital microscopic optical coherence tomography imaging to assess mucus-mobilizing interventions for muco-obstructive lung disease in mice. American Journal of Physiology - Lung Cellular and Molecular Physiology, 2020, 318, L518-L524.	2.9	11
58	Conditional deletion of Nedd4-2 in lung epithelial cells causes progressive pulmonary fibrosis in adult mice. Nature Communications, 2020, 11, 2012.	12.8	52
59	SARS-CoV-2-reactive T cells in healthy donors and patients with COVID-19. Nature, 2020, 587, 270-274.	27.8	1,115
60	Lack of Kcnn4 improves mucociliary clearance in muco-obstructive lung disease. JCI Insight, 2020, 5, .	5.0	11
61	Elexacaftor–Tezacaftor–Ivacaftor for Cystic Fibrosis with a Single Phe508del Allele. New England Journal of Medicine, 2019, 381, 1809-1819.	27.0	1,231
62	Efficacy and safety of the elexacaftor plus tezacaftor plus ivacaftor combination regimen in people with cystic fibrosis homozygous for the F508del mutation: a double-blind, randomised, phase 3 trial. Lancet, The, 2019, 394, 1940-1948.	13.7	804
63	Targeting of cathepsin S reduces cystic fibrosis-like lung disease. European Respiratory Journal, 2019, 53, 1801523.	6.7	31
64	Clinical development of triple-combination CFTR modulators for cystic fibrosis patients with one or two <i>F508del</i> alleles. ERJ Open Research, 2019, 5, 00082-2019.	2.6	72
65	Ten years of chest MRI for patients with cystic fibrosis. Der Radiologe, 2019, 59, 10-20.	1.7	14
66	Midterm Reproducibility of Chest Magnetic Resonance Imaging in Adults with Clinically Stable Cystic Fibrosis and Chronic Obstructive Pulmonary Disease. American Journal of Respiratory and Critical Care Medicine, 2019, 200, 103-107.	5.6	25
67	Cathepsin G Activity as a New Marker for Detecting Airway Inflammation by Microscopy and Flow Cytometry. ACS Central Science, 2019, 5, 539-548.	11.3	21
68	Authors' response: Letter to the Editor †Comparison of lung clearance index determined by washout of N2 and SF6 in infants and preschool children with cystic fibrosis'. Journal of Cystic Fibrosis, 2019, 18, e28-e29.	0.7	0
69	Mucus obstruction and inflammation in early cystic fibrosis lung disease: Emerging role of the ILâ€1 signaling pathway. Pediatric Pulmonology, 2019, 54, S5-S12.	2.0	48
70	Preventive Inhalation of Hypertonic Saline in Infants with Cystic Fibrosis (PRESIS). A Randomized, Double-Blind, Controlled Study. American Journal of Respiratory and Critical Care Medicine, 2019, 199, 1238-1248.	5.6	96
71	Comparison of lung clearance index determined by washout of N2 and SF6 in infants and preschool children with cystic fibrosis. Journal of Cystic Fibrosis, 2019, 18, 399-406.	0.7	21
72	Antisense oligonucleotide eluforsen improves CFTR function in F508del cystic fibrosis. Journal of Cystic Fibrosis, 2019, 18, 536-542.	0.7	41

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73	Elastase Exocytosis by Airway Neutrophils Is Associated with Early Lung Damage in Children with Cystic Fibrosis. American Journal of Respiratory and Critical Care Medicine, 2019, 199, 873-881.	5.6	68
74	Lung disease phenotypes caused by over-expression of combinations of alpha, beta, and gamma subunits of the epithelial sodium channel in mouse airways. American Journal of Physiology - Lung Cellular and Molecular Physiology, 2018, 314, ajplung.00382.2.	2.9	10
75	Non-contrast enhanced magnetic resonance imaging detects mosaic signal intensity in early cystic fibrosis lung disease. European Journal of Radiology, 2018, 101, 178-183.	2.6	26
76	One time quantitative PCR detection of Pseudomonas aeruginosa to discriminate intermittent from chronic infection in cystic fibrosis. Journal of Cystic Fibrosis, 2018, 17, 348-355.	0.7	29
77	Effects of Lumacaftor–Ivacaftor Therapy on Cystic Fibrosis Transmembrane Conductance Regulator Function in Phe508del Homozygous Patients with Cystic Fibrosis. American Journal of Respiratory and Critical Care Medicine, 2018, 197, 1433-1442.	5.6	95
78	Mucopurulent Triggering of the Airway Epithelium. Implications in Health and Cystic Fibrosis. American Journal of Respiratory and Critical Care Medicine, 2018, 197, 418-420.	5.6	3
79	Elastase activity on sputum neutrophils correlates with severity of lung disease in cystic fibrosis. European Respiratory Journal, 2018, 51, 1701910.	6.7	67
80	Three-center feasibility of lung clearance index in infants and preschool children with cystic fibrosis and other lung diseases. Journal of Cystic Fibrosis, 2018, 17, 249-255.	0.7	33
81	Cystic fibrosis research topics featured at the 14th ECFS Basic Science Conference: Chairman's summary. Journal of Cystic Fibrosis, 2018, 17, S1-S4.	0.7	5
82	Reply to Verbanck and Vanderhelst: The Respective Roles of Lung Clearance Index and Magnetic Resonance Imaging in the Clinical Management of Patients with Cystic Fibrosis. American Journal of Respiratory and Critical Care Medicine, 2018, 197, 410-411.	5.6	1
83	Emerging Concepts and Therapies for Mucoobstructive Lung Disease. Annals of the American Thoracic Society, 2018, 15, S216-S226.	3.2	37
84	Role of the SLC26A9 Chloride Channel as Disease Modifier and Potential Therapeutic Target in Cystic Fibrosis. Frontiers in Pharmacology, 2018, 9, 1112.	3.5	32
85	VX-659–Tezacaftor–Ivacaftor in Patients with Cystic Fibrosis and One or Two Phe508del Alleles. New England Journal of Medicine, 2018, 379, 1599-1611.	27.0	280
86	VX-445–Tezacaftor–Ivacaftor in Patients with Cystic Fibrosis and One or Two Phe508del Alleles. New England Journal of Medicine, 2018, 379, 1612-1620.	27.0	509
87	Multicentre standardisation of chest MRI as radiation-free outcome measure of lung disease in young children with cystic fibrosis. Journal of Cystic Fibrosis, 2018, 17, 518-527.	0.7	68
88	Ductal Mucus Obstruction and Reduced Fluid Secretion Are Early Defects in Chronic Pancreatitis. Frontiers in Physiology, 2018, 9, 632.	2.8	13
89	Expression and function of Anoctamin 1/TMEM16A calcium-activated chloride channels in airways of in vivo mouse models for cystic fibrosis research. Pflugers Archiv European Journal of Physiology, 2018, 470, 1335-1348.	2.8	12
90	Interleukin-1 is associated with inflammation and structural lung disease in young children with cystic fibrosis. Journal of Cystic Fibrosis, 2018, 17, 715-722.	0.7	47

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91	Ion Channel Modulators in Cystic Fibrosis. Chest, 2018, 154, 383-393.	0.8	128
92	Validation of automated lobe segmentation on paired inspiratory-expiratory chest CT in 8-14 year-old children with cystic fibrosis. PLoS ONE, 2018, 13, e0194557.	2.5	25
93	Comparison of Lung Clearance Index and Magnetic Resonance Imaging for Assessment of Lung Disease in Children with Cystic Fibrosis. American Journal of Respiratory and Critical Care Medicine, 2017, 195, 349-359.	5.6	169
94	MRI accelerating progress in functional assessment of cystic fibrosis lung disease. Journal of Cystic Fibrosis, 2017, 16, 165-167.	0.7	12
95	Hypoxia and sterile inflammation in cystic fibrosis airways: mechanisms and potential therapies. European Respiratory Journal, 2017, 49, 1600903.	6.7	90
96	Airway mucus, inflammation and remodeling: emerging links in the pathogenesis of chronic lung diseases. Cell and Tissue Research, 2017, 367, 537-550.	2.9	128
97	Gene signature driving invasive mucinous adenocarcinoma of the lung. EMBO Molecular Medicine, 2017, 9, 462-481.	6.9	79
98	Protean proteases: at the cutting edgeÂofÂlung diseases. European Respiratory Journal, 2017, 49, 1501200.	6.7	49
99	Disruption of the Hepcidin/Ferroportin Regulatory System Causes Pulmonary Iron Overload and Restrictive Lung Disease. EBioMedicine, 2017, 20, 230-239.	6.1	45
100	Bypassing CFTR dysfunction in cystic fibrosis with alternative pathways for anion transport. Current Opinion in Pharmacology, 2017, 34, 91-97.	3.5	58
101	Homeostatic nuclear RAGE–ATM interaction is essential for efficient DNA repair. Nucleic Acids Research, 2017, 45, 10595-10613.	14.5	66
102	Cellular distribution and function of ion channels involved in transport processes in rat tracheal epithelium. Physiological Reports, 2017, 5, e13290.	1.7	13
103	Impaired mucus clearance exacerbates allergen-induced type 2 airway inflammation in juvenile mice. Journal of Allergy and Clinical Immunology, 2017, 140, 190-203.e5.	2.9	17
104	An informative intragenic microsatellite marker suggests the IL-1 receptor as a genetic modifier in cystic fibrosis. European Respiratory Journal, 2017, 50, 1700426.	6.7	8
105	Chronic but not intermittent infection with <i>Pseudomonas aeruginosa</i> is associated with global changes of the lung microbiome in cystic fibrosis. European Respiratory Journal, 2017, 50, 1701086.	6.7	33
106	Early detection and sensitive monitoring of CF lung disease: Prospects of improved and safer imaging. Pediatric Pulmonology, 2016, 51, S49-S60.	2.0	44
107	Cigarette smoke causes acute airway disease and exacerbates chronic obstructive lung disease in neonatal mice. American Journal of Physiology - Lung Cellular and Molecular Physiology, 2016, 311, L602-L610.	2.9	22
108	Quantification of heterogeneity in lung disease with image-based pulmonary function testing. Scientific Reports, 2016, 6, 29438.	3.3	50

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109	A product of immunoreactive trypsinogen and pancreatitis-associated protein as second-tier strategy in cystic fibrosis newborn screening. Journal of Cystic Fibrosis, 2016, 15, 752-758.	0.7	14
110	Neutrophil elastase and matrix metalloproteinase 12 in cystic fibrosis lung disease. Molecular and Cellular Pediatrics, 2016, 3, 25.	1.8	37
111	A Protease Inhibitor Tackles Epithelial Sodium Channels in Cystic Fibrosis. American Journal of Respiratory and Critical Care Medicine, 2016, 194, 650-652.	5.6	1
112	Generation and functional characterization of epithelial cells with stable expression of SLC26A9 Cl ^{â^'} channels. American Journal of Physiology - Lung Cellular and Molecular Physiology, 2016, 310, L593-L602.	2.9	36
113	Optical coherence tomography detects structural abnormalitiesof the nasal mucosa in patients with cystic fibrosis. Journal of Cystic Fibrosis, 2016, 15, 216-222.	0.7	19
114	Unplugging Mucus in Cystic Fibrosis and Chronic Obstructive Pulmonary Disease. Annals of the American Thoracic Society, 2016, 13 Suppl 2, S177-85.	3.2	41
115	Finding new drugs to enhance anion secretion in cystic fibrosis: Toward suitable systems for better drug screening. Report on the pre-conference meeting to the 12th ECFS Basic Science Conference, Albufeira, 25–28 March 2015. Journal of Cystic Fibrosis, 2015, 14, 700-705.	0.7	2
116	Imaging modalities in cystic fibrosis. Current Opinion in Pulmonary Medicine, 2015, 21, 609-616.	2.6	34
117	Five years of experience with biochemical cystic fibrosis newborn screening based on IRT/PAP in Germany. Pediatric Pulmonology, 2015, 50, 655-664.	2.0	62
118	Free DNA in Cystic Fibrosis Airway Fluids Correlates with Airflow Obstruction. Mediators of Inflammation, 2015, 2015, 1-11.	3.0	100
119	Airway Surface Dehydration Aggravates Cigarette Smoke-Induced Hallmarks of COPD in Mice. PLoS ONE, 2015, 10, e0129897.	2.5	21
120	Comparison of Microbiomes from Different Niches of Upper and Lower Airways in Children and Adolescents with Cystic Fibrosis. PLoS ONE, 2015, 10, e0116029.	2.5	133
121	Supplementation with Red Palm Oil Increases <i>î²</i> -Carotene and Vitamin A Blood Levels in Patients with Cystic Fibrosis. Mediators of Inflammation, 2015, 2015, 1-7.	3.0	10
122	miRNA-221 is elevated in cystic fibrosis airway epithelial cells and regulates expression of ATF6. Molecular and Cellular Pediatrics, 2015, 2, 1.	1.8	27
123	miR-17 overexpression in cystic fibrosis airway epithelial cells decreases interleukin-8 production. European Respiratory Journal, 2015, 46, 1350-1360.	6.7	64
124	Targeting ion channels in cystic fibrosis. Journal of Cystic Fibrosis, 2015, 14, 561-570.	0.7	126
125	The serine protease inhibitor SerpinA3N attenuates neuropathic pain by inhibiting T cell–derived leukocyte elastase. Nature Medicine, 2015, 21, 518-523.	30.7	182
126	The role of chitin, chitinases, and chitinase-like proteins in pediatric lung diseases. Molecular and Cellular Pediatrics, 2015, 2, 3.	1.8	52

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127	Hypoxic Epithelial Necrosis Triggers Neutrophilic Inflammation via IL-1 Receptor Signaling in Cystic Fibrosis Lung Disease. American Journal of Respiratory and Critical Care Medicine, 2015, 191, 902-913.	5.6	78
128	Intestinal Current Measurements Detect Activation of Mutant CFTR in Patients with Cystic Fibrosis with the G551D Mutation Treated with Ivacaftor. American Journal of Respiratory and Critical Care Medicine, 2015, 192, 1252-1255.	5.6	59
129	Bronchoalveolar Sublineage Specification of Pluripotent Stem Cells: Effect of Dexamethasone Plus cAMP-Elevating Agents and Keratinocyte Growth Factor. Tissue Engineering - Part A, 2015, 21, 669-682.	3.1	7
130	Lung arginase expression and activity is increased in cystic fibrosis mouse models. Journal of Applied Physiology, 2014, 117, 284-288.	2.5	11
131	Airway Mucus Obstruction Triggers Macrophage Activation and Matrix Metalloproteinase 12–Dependent Emphysema. American Journal of Respiratory Cell and Molecular Biology, 2014, 51, 709-720.	2.9	76
132	A New Player in the Game: Epithelial Cathepsin S in Early Cystic Fibrosis Lung Disease. American Journal of Respiratory and Critical Care Medicine, 2014, 190, 126-127.	5.6	10
133	Multiple Breath Washout Is Feasible in the Clinical Setting and Detects Abnormal Lung Function in Infants and Young Children with Cystic Fibrosis. Respiration, 2014, 87, 357-363.	2.6	48
134	Lack of Neutrophil Elastase Reduces Inflammation, Mucus Hypersecretion, and Emphysema, but Not Mucus Obstruction, in Mice with Cystic Fibrosis–like Lung Disease. American Journal of Respiratory and Critical Care Medicine, 2014, 189, 1082-1092.	5.6	116
135	Early cystic fibrosis lung disease: Role of airway surface dehydration and lessons from preventive rehydration therapies in mice. International Journal of Biochemistry and Cell Biology, 2014, 52, 174-179.	2.8	28
136	Comparison of different IRT-PAP protocols to screen newborns for cystic fibrosis in three central European populations. Journal of Cystic Fibrosis, 2014, 13, 15-23.	0.7	39
137	Chronic ivacaftor treatment: Getting F508del-CFTR into more trouble?. Journal of Cystic Fibrosis, 2014, 13, 605-607.	0.7	8
138	Nanoparticle uptake by airway phagocytes after fungal spore challenge in murine allergic asthma and chronic bronchitis. BMC Pulmonary Medicine, 2014, 14, 116.	2.0	14
139	CFTR: cystic fibrosis and beyond. European Respiratory Journal, 2014, 44, 1042-1054.	6.7	207
140	Magnetic Resonance Imaging Detects Changes in Structure and Perfusion, and Response to Therapy in Early Cystic Fibrosis Lung Disease. American Journal of Respiratory and Critical Care Medicine, 2014, 189, 956-965.	5.6	228
141	FRETâ€based and other fluorescent proteinase probes. Biotechnology Journal, 2014, 9, 266-281.	3.5	46
142	CFTR functional measurements in human models for diagnosis, prognosis and personalized therapy. Journal of Cystic Fibrosis, 2014, 13, 363-372.	0.7	34
143	Pathophysiology of cystic fibrosis lung disease. , 2014, , 1-13.		5
144	Cellular uptake and localization of inhaled gold nanoparticles in lungs of mice with chronic obstructive pulmonary disease. Particle and Fibre Toxicology, 2013, 10, 19.	6.2	74

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145	Computerized image analysis of iron-stained macrophages. Annals of Hematology, 2013, 92, 1195-1199.	1.8	3
146	Keratinocyte Growth Factor and Dexamethasone Plus Elevated cAMP Levels Synergistically Support Pluripotent Stem Cell Differentiation into Alveolar Epithelial Type II Cells. Tissue Engineering - Part A, 2013, 19, 938-951.	3.1	23
147	Validation of Fourier decomposition MRI with dynamic contrast-enhanced MRI using visual and automated scoring of pulmonary perfusion in young cystic fibrosis patients. European Journal of Radiology, 2013, 82, 2371-2377.	2.6	99
148	Hypertonic Saline Is Effective in the Prevention and Treatment of Mucus Obstruction, but Not Airway Inflammation, in Mice with Chronic Obstructive Lung Disease. American Journal of Respiratory Cell and Molecular Biology, 2013, 49, 410-417.	2.9	45
149	Automatic Airway Analysis on Multidetector Computed Tomography in Cystic Fibrosis. Journal of Thoracic Imaging, 2013, 28, 104-113.	1.5	66
150	Pulmonary Emphysema in Cystic Fibrosis Detected by Densitometry on Chest Multidetector Computed Tomography. PLoS ONE, 2013, 8, e73142.	2.5	40
151	mCLCA3 Does Not Contribute to Calcium-Activated Chloride Conductance in Murine Airways. American Journal of Respiratory Cell and Molecular Biology, 2012, 47, 87-93.	2.9	22
152	Morphologic and functional scoring of cystic fibrosis lung disease using MRI. European Journal of Radiology, 2012, 81, 1321-1329.	2.6	163
153	BPIFB1 (LPLUNC1) is upregulated in cystic fibrosis lung disease. Histochemistry and Cell Biology, 2012, 138, 749-758.	1.7	31
154	Hfe Deficiency Impairs Pulmonary Neutrophil Recruitment in Response to Inflammation. PLoS ONE, 2012, 7, e39363.	2.5	14
155	CFTR Regulates Early Pathogenesis of Chronic Obstructive Lung Disease in βENaC-Overexpressing Mice. PLoS ONE, 2012, 7, e44059.	2.5	44
156	Spatially Resolved Monitoring of Neutrophil Elastase Activity with Ratiometric Fluorescent Reporters. Angewandte Chemie - International Edition, 2012, 51, 6258-6261.	13.8	75
157	Homozygous CFTR mutation M348K in a boy with respiratory symptoms and failure to thrive. Disease-causing mutation or benign alteration?. European Journal of Pediatrics, 2012, 171, 1039-1046.	2.7	4
158	SLC26A9-mediated chloride secretion prevents mucus obstruction in airway inflammation. Journal of Clinical Investigation, 2012, 122, 3629-3634.	8.2	83
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