## Patrick F Augustijns

List of Publications by Year in descending order

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365 papers 19,416 citations

75 h-index

8749

19169 118 g-index

377 all docs

377 docs citations

times ranked

377

16211 citing authors

#	Article	IF	CITATIONS
1	Supersaturating Drug Delivery Systems: The Answer to Solubility-Limited Oral Bioavailability?. Journal of Pharmaceutical Sciences, 2009, 98, 2549-2572.	1.6	778
2	Top-down production of drug nanocrystals: Nanosuspension stabilization, miniaturization and transformation into solid products. International Journal of Pharmaceutics, 2008, 364, 64-75.	2.6	611
3	Physical stabilisation of amorphous ketoconazole in solid dispersions with polyvinylpyrrolidone K25. European Journal of Pharmaceutical Sciences, 2001, 12, 261-269.	1.9	370
4	In vitro models for the prediction of in vivo performance of oral dosage forms. European Journal of Pharmaceutical Sciences, 2014, 57, 342-366.	1.9	297
5	Histamine Receptor H1–Mediated Sensitization of TRPV1 Mediates Visceral Hypersensitivity and Symptoms in Patients With Irritable Bowel Syndrome. Gastroenterology, 2016, 150, 875-887.e9.	0.6	263
6	Favipiravir at high doses has potent antiviral activity in SARS-CoV-2â^infected hamsters, whereas hydroxychloroquine lacks activity. Proceedings of the National Academy of Sciences of the United States of America, 2020, 117, 26955-26965.	3.3	240
7	Ordered Mesoporous Silica Material SBA-15: A Broad-Spectrum Formulation Platform for Poorly Soluble Drugs. Journal of Pharmaceutical Sciences, 2009, 98, 2648-2658.	1.6	237
8	In vivo methods for drug absorption – Comparative physiologies, model selection, correlations with in vitro methods (IVIVC), and applications for formulation/API/excipient characterization including food effects. European Journal of Pharmaceutical Sciences, 2014, 57, 99-151.	1.9	226
9	The mechanisms of pharmacokinetic food-drug interactions – A perspective from the UNGAP group. European Journal of Pharmaceutical Sciences, 2019, 134, 31-59.	1.9	224
10	Successful oral delivery of poorly water-soluble drugs both depends on the intraluminal behavior of drugs and of appropriate advanced drug delivery systems. European Journal of Pharmaceutical Sciences, 2019, 137, 104967.	1.9	222
11	Increasing the oral bioavailability of the poorly water soluble drug itraconazole with ordered mesoporous silica. European Journal of Pharmaceutics and Biopharmaceutics, 2008, 69, 223-230.	2.0	221
12	Comparison of drug transporter gene expression and functionality in Caco-2 cells from 10 different laboratories. European Journal of Pharmaceutical Sciences, 2008, 35, 383-396.	1.9	220
13	Physical State of Poorly Water Soluble Therapeutic Molecules Loaded into SBA-15 Ordered Mesoporous Silica Carriers: A Case Study with Itraconazole and Ibuprofen. Langmuir, 2008, 24, 8651-8659.	1.6	212
14	Evaluation of gastrointestinal drug supersaturation and precipitation: Strategies and issues. International Journal of Pharmaceutics, 2013, 453, 25-35.	2.6	212
15	Enhanced release of itraconazole from ordered mesoporous SBA-15 silica materials. Chemical Communications, 2007, , 1375.	2.2	202
16	A screening study of surface stabilization during the production of drug nanocrystals. Journal of Pharmaceutical Sciences, 2009, 98, 2091-2103.	1.6	191
17	Physicochemical characterization of solid dispersions of the antiviral agent UC-781 with polyethylene glycol 6000 and Gelucire 44/14. European Journal of Pharmaceutical Sciences, 2000, 10, 311-322.	1.9	187
18	Enhanced absorption of the poorly soluble drug fenofibrate by tuning its release rate from ordered mesoporous silica. European Journal of Pharmaceutical Sciences, 2010, 41, 623-630.	1.9	180

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19	Drying of crystalline drug nanosuspensions—The importance of surface hydrophobicity on dissolution behavior upon redispersion. European Journal of Pharmaceutical Sciences, 2008, 35, 127-135.	1.9	179
20	Characterization of Human Duodenal Fluids in Fasted and Fed State Conditions. Journal of Pharmaceutical Sciences, 2016, 105, 673-681.	1.6	178
21	Nasal vaccination with N-trimethyl chitosan and PLGA based nanoparticles: Nanoparticle characteristics determine quality and strength of the antibody response in mice against the encapsulated antigen. Vaccine, 2010, 28, 6282-6291.	1.7	176
22	Physico-chemical characterization of solid dispersions of temazepam with polyethylene glycol 6000 and PVP K30. International Journal of Pharmaceutics, 1998, 164, 67-80.	2.6	169
23	Evidence for a Polarized Efflux System in Caco-2 Cells Capable of Modulating Cyclosporine A Transport. Biochemical and Biophysical Research Communications, 1993, 197, 360-365.	1.0	165
24	A review of drug solubility in human intestinal fluids: Implications for the prediction of oral absorption. European Journal of Pharmaceutical Sciences, 2014, 57, 322-332.	1.9	159
25	Local immune response to food antigens drives meal-induced abdominal pain. Nature, 2021, 590, 151-156.	13.7	153
26	Interaction of HIV protease inhibitors with OATP1B1, 1B3, and 2B1. Xenobiotica, 2010, 40, 163-176.	0.5	148
27	Simulated intestinal fluid as transport medium in the Caco-2 cell culture model. International Journal of Pharmaceutics, 2002, 232, 183-192.	2.6	146
28	Impact of regional differences along the gastrointestinal tract of healthy adults on oral drug absorption: An UNGAP review. European Journal of Pharmaceutical Sciences, 2019, 134, 153-175.	1.9	146
29	Microbiota-Derived Phenylacetylglutamine Associates with Overall Mortality and Cardiovascular Disease in Patients with CKD. Journal of the American Society of Nephrology: JASN, 2016, 27, 3479-3487.	3.0	144
30	Impact of gastrointestinal physiology on drug absorption in special populations––An UNGAP review. European Journal of Pharmaceutical Sciences, 2020, 147, 105280.	1.9	142
31	Effect of simulated intestinal fluid on drug permeability estimation across Caco-2 monolayers. International Journal of Pharmaceutics, 2004, 274, 221-232.	2.6	141
32	Characterization of fasted-state human intestinal fluids collected from duodenum and jejunum. Journal of Pharmacy and Pharmacology, 2010, 58, 1079-1089.	1.2	140
33	Supersaturating Drug Delivery Systems: Fast is Not Necessarily Good Enough. Journal of Pharmaceutical Sciences, 2012, 101, 7-9.	1.6	140
34	Drug permeability profiling using cell-free permeation tools: Overview and applications. European Journal of Pharmaceutical Sciences, 2018, 119, 219-233.	1.9	139
35	Impact of gastrointestinal tract variability on oral drug absorption and pharmacokinetics: An UNGAP review. European Journal of Pharmaceutical Sciences, 2021, 162, 105812.	1.9	137
36	Antiretroviral Efficacy and Pharmacokinetics of Oral Bis(isopropyloxycarbonyloxymethyl)9-(2-Phosphonylmethoxypropyl)adenine in Mice. Antimicrobial Agents and Chemotherapy, 1998, 42, 1568-1573.	1.4	135

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37	In vivo, in vitro and in silico methods for small molecule transfer across the BBB. Journal of Pharmaceutical Sciences, 2009, 98, 4429-4468.	1.6	128
38	Excipient-Mediated Supersaturation Stabilization in Human Intestinal Fluids. Molecular Pharmaceutics, 2011, 8, 564-570.	2.3	119
39	Characterization of physico-chemical properties and pharmaceutical performance of sucrose co-freeze–dried solid nanoparticulate powders of the anti-HIV agent loviride prepared by media milling. International Journal of Pharmaceutics, 2007, 338, 198-206.	2.6	118
40	In Vitro Hepatic Metabolism Explains Higher Clearance of Voriconazole in Children versus Adults: Role of CYP2C19 and Flavin-Containing Monooxygenase 3. Drug Metabolism and Disposition, 2010, 38, 25-31.	1.7	115
41	Effect of pH and Comedication on Gastrointestinal Absorption of Posaconazole. Clinical Pharmacokinetics, 2011, 50, 725-734.	1.6	114
42	Postprandial Evolution in Composition and Characteristics of Human Duodenal Fluids in Different Nutritional States. Journal of Pharmaceutical Sciences, 2009, 98, 1177-1192.	1.6	112
43	Combined use of ordered mesoporous silica and precipitation inhibitors for improved oral absorption of the poorly soluble weak base itraconazole. European Journal of Pharmaceutics and Biopharmaceutics, 2010, 75, 354-365.	2.0	111
44	Self-Assembly of Cyclodextrins and Their Complexes in Aqueous Solutions. Journal of Pharmaceutical Sciences, 2016, 105, 2556-2569.	1.6	111
45	Structure-Based Identification of OATP1B1/3 Inhibitors. Molecular Pharmacology, 2013, 83, 1257-1267.	1.0	110
46	Sandwich-cultured hepatocytes: utility for <i>in vitro </i> exploration of hepatobiliary drug disposition and drug-induced hepatotoxicity. Expert Opinion on Drug Metabolism and Toxicology, 2013, 9, 589-616.	1.5	110
47	Postprandial Changes in Solubilizing Capacity of Human Intestinal Fluids for BCS Class II Drugs. Pharmaceutical Research, 2009, 26, 1456-1466.	1.7	109
48	Drug precipitation–permeation interplay: Supersaturation in an absorptive environment. European Journal of Pharmaceutics and Biopharmaceutics, 2012, 82, 424-428.	2.0	107
49	The Influence of CKD on Colonic Microbial Metabolism. Journal of the American Society of Nephrology: JASN, 2016, 27, 1389-1399.	3.0	106
50	Use of Azo Polymers for Colon-Specific Drug Delivery. Journal of Pharmaceutical Sciences, 1997, 86, 1321-1327.	1.6	103
51	The use of human nasal in vitro cell systems during drug discovery and development. Toxicology in Vitro, 2005, 19, 107-122.	1.1	102
52	Investigation of thermal properties of glassy itraconazole: identification of a monotropic mesophase. Thermochimica Acta, 2001, 376, 175-181.	1.2	100
53	Solubility Increases Associated with Crystalline Drug Nanoparticles: Methodologies and Significance. Molecular Pharmaceutics, 2010, 7, 1858-1870.	2.3	100
54	Thiolated chitosan nanoparticles for the nasal administration of leuprolide: Bioavailability and pharmacokinetic characterization. International Journal of Pharmaceutics, 2012, 428, 164-170.	2.6	100

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55	Intestinal drug solubility estimation based on simulated intestinal fluids: Comparison with solubility in human intestinal fluids. European Journal of Pharmaceutical Sciences, 2011, 43, 260-269.	1.9	97
56	Metabolism of stevioside in pigs and intestinal absorption characteristics of stevioside, rebaudioside A and steviol. Food and Chemical Toxicology, 2003, 41, 1599-1607.	1.8	96
57	Biological, Pharmaceutical, and Analytical Considerations with Respect to the Transport Media Used in the Absorption Screening System, Caco-2. Journal of Pharmaceutical Sciences, 2003, 92, 1545-1558.	1.6	93
58	Evaluation of various PAMPA models to identify the most discriminating method for the prediction of BBB permeability. European Journal of Pharmaceutics and Biopharmaceutics, 2010, 74, 495-502.	2.0	92
59	Supersaturation and Precipitation of Posaconazole Upon Entry in the Upper Small Intestine in Humans. Journal of Pharmaceutical Sciences, 2016, 105, 2677-2684.	1.6	92
60	Oral biopharmaceutics tools $\hat{a} \in \text{``Ime for a new initiative } \hat{a} \in \text{``An introduction to the IMI project OrBiTo.}$ European Journal of Pharmaceutical Sciences, 2014, 57, 292-299.	1.9	91
61	In vitro models for the prediction of in vivo performance of oral dosage forms: Recent progress from partnership through the IMI OrBiTo collaboration. European Journal of Pharmaceutics and Biopharmaceutics, 2019, 136, 70-83.	2.0	91
62	The combined treatment of Molnupiravir and Favipiravir results in a potentiation of antiviral efficacy in a SARS-CoV-2 hamster infection model. EBioMedicine, 2021, 72, 103595.	2.7	91
63	Napsamycins, new Pseudomonas active antibiotics of the mureidomycin family from Streptomyces sp. HIL Y-82, 11372 Journal of Antibiotics, 1994, 47, 595-598.	1.0	88
64	Drug Supersaturation in Simulated Human Intestinal Fluids Representing Different Nutritional States. Journal of Pharmaceutical Sciences, 2010, 99, 4525-4534.	1.6	88
65	Intraluminal drug and formulation behavior and integration in in vitro permeability estimation: A case study with amprenavir. Journal of Pharmaceutical Sciences, 2006, 95, 372-383.	1.6	87
66	Intestinal Absorption Enhancement of the Ester Prodrug Tenofovir Disoproxil Fumarate through Modulation of the Biochemical Barrier by Defined Ester Mixtures. Drug Metabolism and Disposition, 2002, 30, 924-930.	1.7	86
67	The oral protease inhibitor (PF-07321332) protects Syrian hamsters against infection with SARS-CoV-2 variants of concern. Nature Communications, 2022, 13, 719.	5.8	86
68	Aging behavior of pharmaceutical formulations of itraconazole on SBA-15 ordered mesoporous silica carrier material. Microporous and Mesoporous Materials, 2010, 130, 154-161.	2.2	85
69	Food-dependent disintegration of immediate release fosamprenavir tablets: In vitro evaluation using magnetic resonance imaging and a dynamic gastrointestinal system. European Journal of Pharmaceutics and Biopharmaceutics, 2011, 77, 313-319.	2.0	84
70	Current challenges and future perspectives in oral absorption research: An opinion of the UNGAP network. Advanced Drug Delivery Reviews, 2021, 171, 289-331.	6.6	84
71	Role of Flavin-Containing Monooxygenase in Oxidative Metabolism of Voriconazole by Human Liver Microsomes. Drug Metabolism and Disposition, 2008, 36, 1119-1125.	1.7	82
72	Gastrointestinal behavior of nano- and microsized fenofibrate: In vivo evaluation in man and in vitro simulation by assessment of the permeation potential. European Journal of Pharmaceutical Sciences, 2015, 77, 40-47.	1.9	82

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73	Exploring gastrointestinal variables affecting drug and formulation behavior: Methodologies, challenges and opportunities. International Journal of Pharmaceutics, 2017, 519, 79-97.	2.6	81
74	Ordered mesoporous silica induces pH-independent supersaturation of the basic low solubility compound itraconazole resulting in enhanced transepithelial transport. International Journal of Pharmaceutics, 2008, 357, 169-179.	2.6	79
75	Hepatocyte-based in vitro model for assessment of drug-induced cholestasis. Toxicology and Applied Pharmacology, 2014, 274, 124-136.	1.3	79
76	Microcrystalline cellulose, a useful alternative for sucrose as a matrix former during freeze-drying of drug nanosuspensions – A case study with itraconazole. European Journal of Pharmaceutics and Biopharmaceutics, 2008, 70, 590-596.	2.0	78
77	The Influence of Dietary Protein Intake on Mammalian Tryptophan and Phenolic Metabolites. PLoS ONE, 2015, 10, e0140820.	1.1	77
78	Incomplete Desorption of Liquid Excipients Reduces the <i>in Vitro</i> and <i>in Vivo</i> Performance of Self-Emulsifying Drug Delivery Systems Solidified by Adsorption onto an Inorganic Mesoporous Carrier. Molecular Pharmaceutics, 2012, 9, 2750-2760.	2.3	76
79	Sodium fluorescein is a probe substrate for hepatic drug transport mediated by OATP1B1 and OATP1B3. Journal of Pharmaceutical Sciences, 2011, 100, 5018-5030.	1.6	74
80	The Influence of Prebiotic Arabinoxylan Oligosaccharides on Microbiota Derived Uremic Retention Solutes in Patients with Chronic Kidney Disease: A Randomized Controlled Trial. PLoS ONE, 2016, 11, e0153893.	1.1	74
81	The dynamic gastric environment and its impact on drug and formulation behaviour. European Journal of Pharmaceutical Sciences, 2017, 96, 207-231.	1.9	73
82	Multidrug resistance-associated protein 2 (MRP2) affects hepatobiliary elimination but not the intestinal disposition of tenofovir disoproxil fumarate and its metabolites. Xenobiotica, 2005, 35, 1055-1066.	0.5	72
83	Classification of the Crystallization Behavior of Amorphous Active Pharmaceutical Ingredients in Aqueous Environments. Pharmaceutical Research, 2014, 31, 969-982.	1.7	71
84	Inulin hydrogels as carriers for colonic drug targeting: I. Synthesis and characterization of methacrylated inulin and hydrogel formation. Pharmaceutical Research, 1997, 14, 1730-1737.	1.7	69
85	Improvement of the dissolution rate of artemisinin by means of supercritical fluid technology and solid dispersions. International Journal of Pharmaceutics, 2003, 254, 173-181.	2.6	69
86	The angiotensin converting enzyme inhibitory tripeptides lle-Pro-Pro and Val-Pro-Pro show increasing permeabilities with increasing physiological relevance of absorption models. Peptides, 2008, 29, 1312-1320.	1.2	69
87	Validation of Dissolution Testing with Biorelevant Media: An OrBiTo Study. Molecular Pharmaceutics, 2017, 14, 4192-4201.	2.3	69
88	Preoperative administration of the 5-HT4 receptor agonist prucalopride reduces intestinal inflammation and shortens postoperative ileus via cholinergic enteric neurons. Gut, 2019, 68, 1406-1416.	6.1	69
89	Transport of Artemisinin and Sodium Artesunate in Caco-2 Intestinal Epithelial Cells. Journal of Pharmaceutical Sciences, 1996, 85, 577-579.	1.6	68
90	Novel generic UPLC/MS/MS method for high throughput analysis applied to permeability assessment in early Drug Discovery. Journal of Chromatography B: Analytical Technologies in the Biomedical and Life Sciences, 2007, 847, 182-187.	1.2	68

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91	Characterization of glassy itraconazole: a comparative study of its molecular mobility below Tg with that of structural analogues using MTDSC. International Journal of Pharmaceutics, 2001, 213, 163-173.	2.6	67
92	The effect of food components on the absorption of P-gp substrates: a review. Journal of Pharmacy and Pharmacology, 2010, 55, 153-162.	1.2	67
93	Ex vivo permeability experiments in excised rat intestinal tissue and in vitro solubility measurements in aspirated human intestinal fluids support age-dependent oral drug absorption. European Journal of Pharmaceutical Sciences, 2010, 39, 15-22.	1.9	67
94	<i>In Vitro</i> Evaluation of Viability, Integrity, and Inflammation in Genital Epithelia upon Exposure to Pharmaceutical Excipients and Candidate Microbicides. Antimicrobial Agents and Chemotherapy, 2010, 54, 5105-5114.	1.4	65
95	Inulin hydrogels. I. Dynamic and equilibrium swelling properties. International Journal of Pharmaceutics, 1998, 172, 127-135.	2.6	64
96	Alternative matrix formers for nanosuspension solidification: Dissolution performance and X-ray microanalysis as an evaluation tool for powder dispersion. European Journal of Pharmaceutical Sciences, 2008, 35, 344-353.	1.9	63
97	Gastrointestinal transfer: In vivo evaluation and implementation in in vitro and in silico predictive tools. European Journal of Pharmaceutical Sciences, 2014, 63, 233-242.	1.9	63
98	In situ perfusion in rodents to explore intestinal drug absorption: Challenges and opportunities. International Journal of Pharmaceutics, 2015, 478, 665-681.	2.6	63
99	In vitro behavior of a phosphate ester prodrug of amprenavir in human intestinal fluids and in the Caco-2 system: Illustration of intraluminal supersaturation. International Journal of Pharmaceutics, 2007, 336, 302-309.	2.6	62
100	Rapid conversion of the ester prodrug abiraterone acetate results in intestinal supersaturation and enhanced absorption of abiraterone: In vitro, rat in situ and human in vivo studies. European Journal of Pharmaceutics and Biopharmaceutics, 2015, 90, 1-7.	2.0	62
101	Higher clearance of micafungin in neonates compared with adults: role of ageâ€dependent micafungin serum binding. Biopharmaceutics and Drug Disposition, 2011, 32, 222-232.	1.1	61
102	Title is missing!. Magyar Apróvad Közlemények, 2002, 68, 591-601.	1.4	60
103	An in vitro biorelevant gastrointestinal transfer (BioGIT) system for forecasting concentrations in the fasted upper small intestine: Design, implementation, and evaluation. European Journal of Pharmaceutical Sciences, 2016, 82, 106-114.	1.9	60
104	Associations of Soluble CD14 and Endotoxin with Mortality, Cardiovascular Disease, and Progression of Kidney Disease among Patients with CKD. Clinical Journal of the American Society of Nephrology: CJASN, 2015, 10, 1525-1533.	2.2	59
105	<i>In Silico</i> Modeling Approach for the Evaluation of Gastrointestinal Dissolution, Supersaturation, and Precipitation of Posaconazole. Molecular Pharmaceutics, 2017, 14, 4321-4333.	2.3	59
106	Drug absorption studies of prodrug esters using the Caco-2 model: evaluation of ester hydrolysis and transport. International Journal of Pharmaceutics, 1998, 166, 45-53.	2.6	58
107	Use of Caco-2 cells and LC/MS/MS to screen a peptide combinatorial library for permeable structures. International Journal of Pharmaceutics, 1999, 177, 103-115.	2.6	58
108	Formulate-ability of ten compounds with different physicochemical profiles in SMEDDS. European Journal of Pharmaceutical Sciences, 2009, 38, 479-488.	1.9	58

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109	Iron Deficiency After Roux-en-Y Gastric Bypass: Insufficient Iron Absorption from Oral Iron Supplements. Obesity Surgery, 2014, 24, 56-61.	1.1	58
110	Scintigraphic evaluation in rabbits of nasal drug delivery systems based on carbopol 971p $\hat{A}^{\otimes}$ and carboxymethylcellulose. Journal of Controlled Release, 2000, 68, 207-214.	4.8	57
111	Supersaturation in human gastric fluids. European Journal of Pharmaceutics and Biopharmaceutics, 2012, 81, 184-189.	2.0	57
112	Metabolism, Protein Binding, and Renal Clearance of Microbiota–Derived p-Cresol in Patients with CKD. Clinical Journal of the American Society of Nephrology: CJASN, 2016, 11, 1136-1144.	2.2	57
113	Lipid-Based Formulations Solidified Via Adsorption onto the Mesoporous Carrier Neusilin $\hat{A}^{\otimes}$ US2: Effect of Drug Type and Formulation Composition on In Vitro Pharmaceutical Performance. Journal of Pharmaceutical Sciences, 2014, 103, 1734-1746.	1.6	56
114	In Vitro Investigation of the Hepatobiliary Disposition Mechanisms of the Antifungal Agent Micafungin in Humans and Rats. Drug Metabolism and Disposition, 2010, 38, 1848-1856.	1.7	55
115	Proton Pump Inhibitors Reduce Duodenal Eosinophilia, Mast Cells, and Permeability in Patients With Functional Dyspepsia. Gastroenterology, 2021, 160, 1521-1531.e9.	0.6	55
116	Determination of OATP-, NTCP- and OCT-mediated substrate uptake activities in individual and pooled batches of cryopreserved human hepatocytes. European Journal of Pharmaceutical Sciences, 2011, 43, 297-307.	1.9	54
117	Synthesis and characterisation of inulin-azo hydrogels designed for colon targeting. International Journal of Pharmaceutics, 2001, 213, 143-152.	2.6	53
118	Species-dependent and site-specific intestinal metabolism of ester prodrugs. International Journal of Pharmaceutics, 2000, 205, 93-100.	2.6	52
119	Downscaling Drug Nanosuspension Production: Processing Aspects and Physicochemical Characterization. AAPS PharmSciTech, 2009, 10, 44-53.	1.5	52
120	The conflict between in vitro release studies in human biorelevant media and the in vivo exposure in rats of the lipophilic compound fenofibrate. International Journal of Pharmaceutics, 2011, 414, 118-124.	2.6	52
121	Boosting of HIV Protease Inhibitors by Ritonavir in the Intestine: The Relative Role of Cytochrome P450 and P-Glycoprotein Inhibition Based on Caco-2 Monolayers versus In Situ Intestinal Perfusion in Mice. Drug Metabolism and Disposition, 2012, 40, 1473-1477.	1.7	52
122	Itraconazole/TPGS/Aerosil®200 solid dispersions. European Journal of Pharmaceutical Sciences, 2009, 38, 270-278.	1.9	50
123	In-vitro nasal drug delivery studies: comparison of derivatised, fibrillar and polymerised collagen matrix-based human nasal primary culture systems for nasal drug delivery studies. Journal of Pharmacy and Pharmacology, 2010, 53, 1447-1456.	1.2	50
124	Iron deficiency after bariatric surgery: what is the real problem?. Proceedings of the Nutrition Society, 2018, 77, 445-455.	0.4	50
125	In vitrointestinal transport and antihypertensive activity of ACE inhibitory pea and whey digests. International Journal of Food Sciences and Nutrition, 2005, 56, 415-430.	1.3	49
126	An In-Depth View into Human Intestinal Fluid Colloids: Intersubject Variability in Relation to Composition. Molecular Pharmaceutics, 2016, 13, 3484-3493.	2.3	49

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127	Gastrointestinal dissolution, supersaturation and precipitation of the weak base indinavir in healthy volunteers. European Journal of Pharmaceutics and Biopharmaceutics, 2016, 109, 122-129.	2.0	49
128	Adaptations in gastrointestinal physiology after sleeve gastrectomy and Roux-en-Y gastric bypass. The Lancet Gastroenterology and Hepatology, 2021, 6, 225-237.	3.7	49
129	Transport, uptake, and metabolism of the bis(pivaloyloxymethyl)-ester prodrug of 9-(2-phosphonylmethoxyethyl)adenine in an in vitro cell culture system of the intestinal mucosa (Caco-2). Pharmaceutical Research, 1997, 14, 492-496.	1.7	48
130	Cardiovascular disease relates to intestinal uptake of p-cresol in patients with chronic kidney disease. BMC Nephrology, 2014, 15, 87.	0.8	48
131	Parallel Monitoring of Plasma and Intraluminal Drug Concentrations in Man After Oral Administration of Fosamprenavir in the Fasted and Fed State. Pharmaceutical Research, 2007, 24, 1862-1869.	1.7	47
132	Application of PAMPA-models to predict BBB permeability including efflux ratio, plasma protein binding and physicochemical parameters. International Journal of Pharmaceutics, 2010, 395, 182-197.	2.6	46
133	Inulin hydrogels. II. In vitro degradation study. International Journal of Pharmaceutics, 1998, 172, 137-145.	2.6	45
134	HPLC with programmed wavelength fluorescence detection for the simultaneous determination of marker compounds of integrity and P-gp functionality in the Caco-2 intestinal absorption model. Journal of Pharmaceutical and Biomedical Analysis, 2004, 34, 971-978.	1.4	44
135	High-speed digital imaging method for ciliary beat frequency measurement. Journal of Pharmacy and Pharmacology, 2010, 57, 521-526.	1.2	44
136	Stereoselective Pharmacokinetic Properties of Chloroquine and De-Ethyl-Chloroquine in Humans. Clinical Pharmacokinetics, 1993, 24, 259-269.	1.6	43
137	Gastrointestinal and Systemic Monitoring of Posaconazole in Humans After Fasted and Fed State Administration of a Solid Dispersion. Journal of Pharmaceutical Sciences, 2016, 105, 2904-2912.	1.6	43
138	Safety assessment of selected cyclodextrins â€" effect on ciliary activity using a human cell suspension culture model exhibiting in vitro ciliogenesis. International Journal of Pharmaceutics, 2000, 193, 219-226.	2.6	42
139	Evaluation of fasted and fed state simulated and human intestinal fluids as solvent system in the Ussing chambers model to explore food effects on intestinal permeability. International Journal of Pharmaceutics, 2015, 478, 736-744.	2.6	42
140	Micronutrient intake, from diet and supplements, and association with status markers in pre- and post-RYGB patients. Clinical Nutrition, 2017, 36, 1175-1181.	2.3	42
141	Usefulness of a novel Caco-2 cell perfusion system. I. in vitro prediction of the absorption potential of passively diffused compounds. Journal of Pharmaceutical Sciences, 2004, 93, 2507-2521.	1.6	41
142	Tunability of Pore Diameter and Particle Size of Amorphous Microporous Silica for Diffusive Controlled Release of Drug Compounds. Journal of Physical Chemistry C, 2007, 111, 13404-13409.	1.5	41
143	Exploring food effects on indinavir absorption with human intestinal fluids in the mouse intestine. European Journal of Pharmaceutical Sciences, 2013, 49, 27-32.	1.9	41
144	Human and simulated intestinal fluids as solvent systems to explore food effects on intestinal solubility and permeability. European Journal of Pharmaceutical Sciences, 2014, 63, 178-186.	1.9	41

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145	Confocal Imaging with a Fluorescent Bile Acid Analogue Closely Mimicking Hepatic Taurocholate Disposition. Journal of Pharmaceutical Sciences, 2014, 103, 1872-1881.	1.6	41
146	Molecular organization of hydrophobic molecules and co-adsorbed water in SBA-15 ordered mesoporous silica material. Physical Chemistry Chemical Physics, 2011, 13, 2706-2713.	1.3	40
147	A liquid chromatography – tandem mass spectrometry method to measure a selected panel of uremic retention solutes derived from endogenous and colonic microbial metabolism. Analytica Chimica Acta, 2016, 936, 149-156.	2.6	40
148	Gastrointestinal Behavior of Weakly Acidic BCS Class II Drugs in Manâ€"Case Study of Diclofenac Potassium. Journal of Pharmaceutical Sciences, 2016, 105, 687-696.	1.6	40
149	Assessment of Passive Intestinal Permeability Using an Artificial Membrane Insert System. Journal of Pharmaceutical Sciences, 2018, 107, 250-256.	1.6	40
150	Interplay of Supersaturation and Solubilization: Lack of Correlation between Concentration-Based Supersaturation Measurements and Membrane Transport Rates in Simulated and Aspirated Human Fluids. Molecular Pharmaceutics, 2019, 16, 5042-5053.	2.3	40
151	In vivo evaluation of xanthan gum as a potential excipient for oral controlled-release matrix tablet formulation. International Journal of Pharmaceutics, 1998, 169, 105-113.	2.6	39
152	Potential of amorphous microporous silica for ibuprofen controlled release. International Journal of Pharmaceutics, 2010, 397, 84-91.	2.6	39
153	Toxicity and intracellular accumulation of bile acids in sandwich-cultured rat hepatocytes: Role of glycine conjugates. Toxicology in Vitro, 2014, 28, 218-230.	1.1	39
154	Hydration Changes Implicated in the Remarkable Temperature-Dependent Membrane Permeation of Cyclosporin A. Biochemistry, 2000, 39, 7621-7630.	1.2	37
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