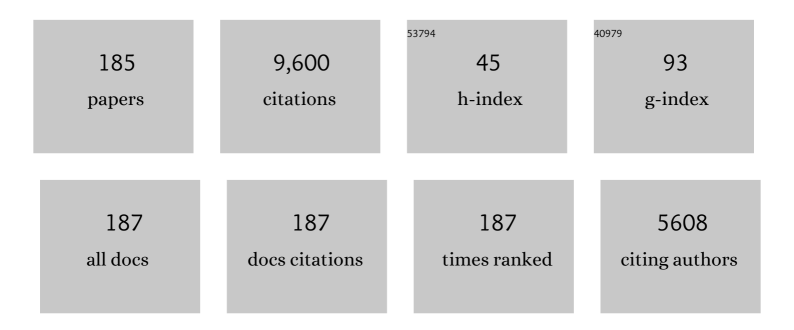
## Samir K Ballas

List of Publications by Year in descending order

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SAMID K RALLAS

#	Article	IF	CITATIONS
1	Maternal and perinatal outcomes in pregnant women with sickle cell disease: an update. Hematology, Transfusion and Cell Therapy, 2022, 44, 369-373.	0.2	9
2	Menopause in Brazilian women with sickle cell anemia with and without hydroxyurea therapy. Hematology, Transfusion and Cell Therapy, 2021, 43, 386-388.	0.2	6
3	Opioids and Sickle Cell Disease: From Opium to the Opioid Epidemic. Journal of Clinical Medicine, 2021, 10, 438.	2.4	9
4	Voxelotor Modulates the Analgesic Effect of Certain Opioids. Journal of Clinical Medicine Research, 2021, 13, 130-132.	1.2	2
5	Opioids are not a major cause of death of patients with sickle cell disease. Annals of Hematology, 2021, 100, 1133-1138.	1.8	5
6	Time to rethink haemoglobin threshold guidelines in sickle cell disease. British Journal of Haematology, 2021, 195, 518-522.	2.5	7
7	Determinants of the Outcome of the Transition of Children with Sickle Cell Disease to Adult Programs. Hemoglobin, 2021, 45, 62-65.	0.8	0
8	Randomized clinical trial of computerized PAINRelievelt® for patients with sickle cell disease: PAINReportIt® and PAINUCope®. Patient Education and Counseling, 2020, 103, 136-144.	2.2	5
9	How I treat acute and persistent sickle cell pain. Mediterranean Journal of Hematology and Infectious Diseases, 2020, 12, e2020064.	1.3	5
10	Risk factors associated with increased emergency department utilization in patients with sickle cell disease: a systematic literature review. Annals of Hematology, 2020, 99, 2483-2495.	1.8	3
11	Case Series and Review of Hematological and Non-Hematological Malignancies in Aging Patients with Sickle Cell Disease in the Hydroxyurea Era. Hemoglobin, 2020, 44, 303-306.	0.8	0
12	Predictors of impending acute chest syndrome in patients with sickle cell anaemia. Scientific Reports, 2020, 10, 2470.	3.3	15
13	THE EVOLVING PHARMACOTHERAPEUTIC LANDSCAPE FOR THE TREATMENT OF SICKLE CELL DISEASE. Mediterranean Journal of Hematology and Infectious Diseases, 2020, 12, e2020010.	1.3	32
14	Review/overview of pain in sickle cell disease. Complementary Therapies in Medicine, 2020, 49, 102327.	2.7	26
15	Transcutaneous electrical nerve stimulation (TENS) for pain management in sickle cell disease. The Cochrane Library, 2020, 2020, CD012762.	2.8	8
16	The vasoâ€occlusive pain crisis in sickle cell disease: Definition, pathophysiology, and management. European Journal of Haematology, 2020, 105, 237-246.	2.2	92
17	SURVIVAL PROBABILITY IN PATIENTS WITH SICKLE CELL ANEMIA USING THE COMPETITIVE RISK STATISTICAL MODEL. Mediterranean Journal of Hematology and Infectious Diseases, 2019, 11, e2019022.	1.3	1
18	Pharmacological interventions for painful sickle cell vaso-occlusive crises in adults. The Cochrane Library, 2019, 2019, .	2.8	13

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19	Treatment of dental complications in sickle cell disease. The Cochrane Library, 2019, 2019, CD011633.	2.8	3
20	AAAPT Diagnostic Criteria for Acute Sickle Cell Disease Pain. Journal of Pain, 2019, 20, 746-759.	1.4	37
21	Indications for RBC Exchange Transfusion in Patients with Sickle Cell Disease: Revisited. Annals of Clinical and Laboratory Science, 2019, 49, 836-837.	0.2	4
22	Blood rheological abnormalities in sickle cell anemia. Clinical Hemorheology and Microcirculation, 2018, 68, 165-172.	1.7	38
23	Sickle cell disease: Classification of clinical complications and approaches to preventive and therapeutic management. Clinical Hemorheology and Microcirculation, 2018, 68, 105-128.	1.7	33
24	Comorbidities in aging patients with sickle cell disease. Clinical Hemorheology and Microcirculation, 2018, 68, 129-145.	1.7	13
25	The effect of iron chelation therapy on overall survival in sickle cell disease and βâ€ŧhalassemia: A systematic review. American Journal of Hematology, 2018, 93, 943-952.	4.1	47
26	Mortality in children, adolescents and adults with sickle cell anemia in Rio de Janeiro, Brazil. Hematology, Transfusion and Cell Therapy, 2018, 40, 37-42.	0.2	26
27	Opioid Utilization by Pregnant Women with Sickle Cell Disease and the Risk of Neonatal Abstinence Syndrome. Journal of the National Medical Association, 2018, 110, 163-168.	0.8	8
28	Rheological properties of sickle erythrocytes in patients with sickleâ€cell anemia: The effect of hydroxyurea, fetal hemoglobin, and αâ€thalassemia. European Journal of Haematology, 2018, 101, 798-803.	2.2	19
29	Opioid utilization patterns in United States individuals with sickle cell disease. American Journal of Hematology, 2018, 93, E345-E347.	4.1	29
30	Microglia-like Cells Derived from Hematopoietic Stem and Progenitor Cells Are a Model System to Investigate Chronic Pain in Sickle Cell Disease. Blood, 2018, 132, 1071-1071.	1.4	0
31	Defining Sickle Cell Disease Acute Painful Episodes: The Pisces Project. Blood, 2018, 132, 3510-3510.	1.4	0
32	Re-Assessing the Red Blood Cell Lifespan in Sickle Cell Anemia: Does Size Matter?. Blood, 2018, 132, 4877-4877.	1.4	0
33	From total blood exchange to erythrocytapheresis and back to treat complications of sickle cell disease. Transfusion, 2017, 57, 2277-2280.	1.6	7
34	Beyond the transition of adolescents and young adults with sickle cell disease to adult care: Role of geography. American Journal of Hematology, 2017, 92, E110-E112.	4.1	2
35	Transcutaneous electrical nerve stimulation (TENS) for pain management in sickle cell disease. The Cochrane Library, 2017, , .	2.8	4
36	Plasmapheresis may improve clinical condition in sickle cell disease through its effects on red blood cell rheology. American Journal of Hematology, 2017, 92, E629-E630.	4.1	16

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37	The Use of Cannabis by Patients with Sickle Cell Disease Increased the Frequency of Hospitalization due to Vaso-Occlusive Crises. Cannabis and Cannabinoid Research, 2017, 2, 197-201.	2.9	19
38	Perinatal Maternal Mortality in Sickle Cell Anemia: Two Case Reports and Review of the Literature. Hemoglobin, 2017, 41, 225-229.	0.8	8
39	Red blood cell exchange: 2015 American Society for Apheresis consensus conference on the management of patients with sickle cell disease. Journal of Clinical Apheresis, 2017, 32, 342-367.	1.3	25
40	Of pools, oceans, and the Dead Sea. Blood, 2017, 130, 2578-2579.	1.4	2
41	The Sixth Vital Sign: Body Mass Index in Patients With Sickle Cell Disease. Journal of Clinical Medicine Research, 2017, 9, 889-890.	1.2	12
42	The paradox of the serrated sickle erythrocyte: The importance of the red bloodÂcell membrane topography. Clinical Hemorheology and Microcirculation, 2016, 63, 149-152.	1.7	7
43	Case series of octogenarians with sickle cell disease. Blood, 2016, 128, 2367-2369.	1.4	16
44	Pfaffia paniculata extract improves red blood cell deformability in sickle cell patients. Clinical Hemorheology and Microcirculation, 2016, 62, 327-333.	1.7	2
45	Original Research: A case-control genome-wide association study identifies genetic modifiers of fetal hemoglobin in sickle cell disease. Experimental Biology and Medicine, 2016, 241, 706-718.	2.4	21
46	Amputations in Sickle Cell Disease: Case Series and Literature Review. Hemoglobin, 2016, 40, 150-155.	0.8	4
47	Comment on: negative health implications of sickle cell trait in high income countries: from the football field to the laboratory. British Journal of Haematology, 2016, 175, 349-350.	2.5	1
48	High mortality among children with sickle cell anemia and overt stroke who discontinue blood transfusion after transition to an adult program. Transfusion, 2016, 56, 1014-1021.	1.6	22
49	The Opioid Drug Epidemic and Sickle Cell Disease: Guilt by Association. Pain Medicine, 2016, 17, 1793-1798.	1.9	69
50	The role of blood rheology in sickle cell disease. Blood Reviews, 2016, 30, 111-118.	5.7	142
51	Safety and efficacy of blood exchange transfusion for priapism complicating sickle cell disease. Journal of Clinical Apheresis, 2016, 31, 5-10.	1.3	31
52	From Individualized Treatment of Sickle Cell Pain to Precision Medicine: A 40-Year Journey. Journal of Clinical Medicine Research, 2016, 8, 357-360.	1.2	4
53	Use of Marijuana in Patients with Sickle Cell Disease Increased the Frequency of Hospitalization for Acute Painful Vaso-Occlusive Crises. Blood, 2016, 128, 2498-2498.	1.4	0
54	Pathophysiology and principles of management of the many faces of the acute vasoâ€occlusive crisis in patients with sickle cell disease. European Journal of Haematology, 2015, 95, 113-123.	2.2	29

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55	Hydroxyurea treatment does not increase blood viscosity and improves red blood cell rheology in sickle cell anemia. Haematologica, 2015, 100, e383-e386.	3.5	46
56	Deferasirox in patients with iron overload secondary to hereditary hemochromatosis: results of a 1â€yr Phase 2 study. European Journal of Haematology, 2015, 95, 545-550.	2.2	34
57	Deep venous thrombosis in children with sickle cell disease. Pediatric Blood and Cancer, 2015, 62, 838-841.	1.5	17
58	Morphine for the Treatment of Pain in Sickle Cell Disease. Scientific World Journal, The, 2015, 2015, 1-10.	2.1	35
59	Deafness and Sickle Cell Disease: Three Case Reports and Review of the Literature. Journal of Clinical Medicine Research, 2015, 7, 189-192.	1.2	8
60	Clinical Utility of Lactate Dehydrogenase in Determining the Severity of Hemolysis in Sickle Cell Anemia. American Journal of Clinical Pathology, 2015, 144, 173-174.	0.7	1
61	Is the Medical Home for Adult Patients with Sickle Cell Disease a Reality or an Illusion?. Hemoglobin, 2015, 39, 130-133.	0.8	3
62	Emerging drugs for sickle cell anemia. Expert Opinion on Emerging Drugs, 2015, 20, 47-61.	2.4	18
63	Risk Factors of Pulmonary Hypertension in Brazilian Patients with Sickle Cell Anemia. PLoS ONE, 2015, 10, e0137539.	2.5	5
64	Management of Sickle Cell Disease. JAMA - Journal of the American Medical Association, 2014, 312, 1033.	7.4	1,189
65	A predecessor of the current blood bank pneumatic tube delivery system. Transfusion, 2014, 54, 3035-3035.	1.6	1
66	Thinking beyond sickling to better understand pain in sickle cell disease. European Journal of Haematology, 2014, 93, 89-95.	2.2	53
67	Alphaâ€ŧhalassaemia and response to hydroxyurea in sickle cell anaemia. European Journal of Haematology, 2014, 92, 341-345.	2.2	14
68	Newborn screening program for hemoglobinopathies in Rio de Janeiro, Brazil. Pediatric Blood and Cancer, 2014, 61, 34-39.	1.5	46
69	Characteristics of Acute Care Utilization of a Delaware Adult Sickle Cell Disease Patient Population. Population Health Management, 2014, 17, 60-65.	1.7	4
70	Primary Stroke in a Woman With Sickle Cell Anemia Responsive to Hydroxyurea Therapy. Hemoglobin, 2014, 38, 373-375.	0.8	0
71	Treatment of the acute sickle cell vasoâ€occlusive crisis in the Emergency Department: a Brazilian method of switching from intravenous to oral morphine. European Journal of Haematology, 2014, 93, 34-40.	2.2	12
72	Leg Amputation for an Extensive, Severe and Intractable Sickle Cell Anemia Ulcer in a Brazilian Patient. Hemoglobin, 2014, 38, 95-98.	0.8	6

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73	Deferiprone versus Deferoxamine in Sickle Cell Disease: Results from a 5-year long-term Italian multi-center randomized clinical trial. Blood Cells, Molecules, and Diseases, 2014, 53, 265-271.	1.4	17
74	Dental Complications of Sickle Cell Disease. JBR Journal of Interdisciplinary Medicine and Dental Science, 2014, 02, .	0.1	5
75	Neuropathy, neuropathic pain, and sickle cell disease. American Journal of Hematology, 2013, 88, 927-929.	4.1	27
76	Lactate dehydrogenase and hemolysis in sickle cell disease. Blood, 2013, 121, 243-244.	1.4	44
77	The first Cardeza donor center: attracting donors who do not wish to see blood. Transfusion, 2013, 53, 13-13.	1.6	0
78	Long-term persistency and costs associated with the use of iron chelation therapies in the treatment of Sickle cell disease within Medicaid programs. Journal of Medical Economics, 2013, 16, 10-18.	2.1	8
79	Is Sickle Cell Disease a Hematologic Disorder?. Journal of the National Medical Association, 2012, 104, 463.	0.8	1
80	Sickle cell pain: a critical reappraisal. Blood, 2012, 120, 3647-3656.	1.4	333
81	Beyond the Definitions of the Phenotypic Complications of Sickle Cell Disease: An Update on Management. Scientific World Journal, The, 2012, 2012, 1-55.	2.1	125
82	More definitions in sickle cell disease: Steady state <i>v</i> base line data. American Journal of Hematology, 2012, 87, 338-338.	4.1	73
83	Defining the Phenotypes of Sickle Cell Disease. Hemoglobin, 2011, 35, 511-519.	0.8	21
84	The Association Between Hydroxyurea Treatment and Pain Intensity, Analgesic Use, and Utilization in Ambulatory Sickle Cell Anemia Patients. Pain Medicine, 2011, 12, 697-705.	1.9	31
85	Healthâ€related quality of life in adults with sickle cell disease (SCD): A report from the comprehensive sickle cell centers clinical trial consortium. American Journal of Hematology, 2011, 86, 203-205.	4.1	111
86	Update on Pain Management in Sickle Cell Disease. Hemoglobin, 2011, 35, 520-529.	0.8	48
87	Sickle Cell Disease and Pregnancy: Does Outcome Depend on Genotype or Phenotype?. International Journal of Clinical Medicine, 2011, 02, 313-317.	0.2	3
88	Sickle Cell Pain. , 2011, , 243-248.		0
89	Definitions of the phenotypic manifestations of sickle cell disease. American Journal of Hematology, 2010, 85, 6-13.	4.1	291
90	Self-management of Sickle Cell Disease: A New Frontier. Journal of the National Medical Association, 2010, 102, 1042-1044.	0.8	9

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91	The Dark Side of the Discovery of Sickle Cell Disease in the United States in 1910. Journal of the National Medical Association, 2010, 102, 1100-1101.	0.8	0
92	The Impact of Hydroxyurea on Career and Employment of Patients With Sickle Cell Anemia. Journal of the National Medical Association, 2010, 102, 993-999.	0.8	27
93	New era dawns on sickle cell pain. Blood, 2010, 116, 311-312.	1.4	10
94	Hydroxyurea and Acute Painful Crises in Sickle Cell Anemia: Effects on Hospital Length of Stay and Opioid Utilization During Hospitalization, Outpatient Acute Care Contacts, and at Home. Journal of Pain and Symptom Management, 2010, 40, 870-882.	1.2	65
95	The risks and benefits of longâ€ŧerm use of hydroxyurea in sickle cell anemia: A 17.5 year followâ€up. American Journal of Hematology, 2010, 85, 403-408.	4.1	385
96	Utilization of analgesics in the multicenter study of hydroxyurea in sickle cell anemia: Effect of sex, age, and geographical location. American Journal of Hematology, 2010, 85, 613-616.	4.1	59
97	Early Detection of Response to Hydroxyurea Therapy in Patients with Sickle Cell Anemia. Hemoglobin, 2010, 34, 424-429.	0.8	10
98	Neurocognitive Complications of Sickle Cell Anemia in Adults. JAMA - Journal of the American Medical Association, 2010, 303, 1862.	7.4	10
99	Effects of a Single Sickling Event on the Mechanical Fragility of Sickle Cell Trait Erythrocytes. Hemoglobin, 2010, 34, 24-36.	0.8	14
100	The impact of hydroxyurea on career and employment of patients with sickle cell anemia. Journal of the National Medical Association, 2010, 102, 993-9.	0.8	8
101	Investigational drugs in sickle cell anemia. Expert Opinion on Investigational Drugs, 2009, 18, 1817-1828.	4.1	13
102	Exposure to Hydroxyurea and Pregnancy Outcomes in Patients With Sickle Cell Anemia. Journal of the National Medical Association, 2009, 101, 1046-1051.	0.8	94
103	Climatic and geographic temporal patterns of pain in the Multicenter Study of Hydroxyurea. Pain, 2009, 146, 91-98.	4.2	43
104	The cost of health care for patients with sickle cell disease. American Journal of Hematology, 2009, 84, 320-322.	4.1	82
105	Corticosteroids and Sickle Cell Disease. Journal of the National Medical Association, 2009, 101, 283.	0.8	3
106	Definition of the Responder to Hydroxyurea Therapy: Revisited Blood, 2009, 114, 1513-1513.	1.4	3
107	Developmental Outcomes of Offspring of Adults Treated with Hydroxyurea in the Multicenter Study of Hydroxyurea Blood, 2009, 114, 1543-1543.	1.4	3
108	Impact of Hydroxyurea On Employment Among Patients with Sickle Cell Anemia Blood, 2009, 114, 2485-2485.	1.4	3

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109	Patterns of Analgesic Utilization in the Multicenter Study of Hydroxyurea (MSH) Blood, 2009, 114, 2577-2577.	1.4	3
110	Sickle Cell Genetic Markers: Geographic Distribution and Relation to Pain Outcomes in Multicenter Study of Hydroxyurea in Sickle Cell Anemia Blood, 2009, 114, 2582-2582.	1.4	3
111	Stroke in Sickle Cell Anemia After Excellent Response to Hydroxyurea Blood, 2009, 114, 4622-4622.	1.4	0
112	Footprints of Response to Hydroxyurea Are in the Hemogram Blood, 2009, 114, 4619-4619.	1.4	0
113	Meperidine for Acute Sickle Cell Pain in the Emergency Department: Revisited Controversy. Annals of Emergency Medicine, 2008, 51, 217.	0.6	13
114	Disparity in the management of iron overload between patients with sickle cell disease and thalassemia who received transfusions. Transfusion, 2008, 48, 1971-1980.	1.6	28
115	Current Issues in Sickle Cell Pain and Its Management. Hematology American Society of Hematology Education Program, 2007, 2007, 97-105.	2.5	97
116	Morbidity and mortality in chronically transfused subjects with thalassemia and sickle cell disease: A report from the multiâ€center study of iron overload. American Journal of Hematology, 2007, 82, 255-265.	4.1	149
117	Secretory Phospholipase A2Levels in Patients with Sickle Cell Disease and Acute Chest Syndrome. Hemoglobin, 2006, 30, 165-170.	0.8	27
118	Hydroxyurea and sickle cell anemia: effect on quality of life. Health and Quality of Life Outcomes, 2006, 4, 59.	2.4	124
119	Hyperhemolysis during the evolution of uncomplicated acute painful episodes in patients with sickle cell anemia. Transfusion, 2006, 46, 105-110.	1.6	89
120	Folate supplementation and twinning in patients with sickle cell disease. American Journal of Hematology, 2006, 81, 296-297.	4.1	5
121	The Role of Nitric Oxide in Regulation of Red Blood Cell Deformability Blood, 2006, 108, 3730-3730.	1.4	26
122	Utilization of the office, hospital and emergency department for adult sickle cell patients: a five-year study. Journal of the National Medical Association, 2006, 98, 1109-13.	0.8	38
123	Hospital readmission for adult acute sickle cell painful episodes: frequency, etiology, and prognostic significance. American Journal of Hematology, 2005, 79, 17-25.	4.1	231
124	Pain Management of Sickle Cell Disease. Hematology/Oncology Clinics of North America, 2005, 19, 785-802.	2.2	134
125	Non-pharmacologic Management of Sickle Cell Pain. Hematology, 2004, 9, 235-237.	1.5	33
126	Reply to â€~Gender-specific disease modification by NOS3'. British Journal of Haematology, 2004, 126, 161-161.	2.5	0

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127	Sickle Red Cell Microrheology and Sickle Blood Rheology. Microcirculation, 2004, 11, 209-225.	1.8	96
128	Management of acute chest wall sickle cell pain with nebulized morphine. American Journal of Hematology, 2004, 76, 190-191.	4.1	20
129	Safety of Purified Poloxamer 188 in Sickle Cell Disease: Phase I Study of a Nonâ€ionic Surfactant in the Management of Acute Chest Syndrome. Hemoglobin, 2004, 28, 85-102.	0.8	49
130	Sickle Red Cell Microrheology and Sickle Blood Rheology. Microcirculation, 2004, 11, 209-225.	1.8	1
131	Arginase Activity Is a Determinant of Nitric Oxide Levels in Adult Vasoocclusive Sickle Cell Crisis Blood, 2004, 104, 1670-1670.	1.4	1
132	Outcome of Transitioning Pediatric Patients with Sickle Cell Disease to Adult Programs Blood, 2004, 104, 3743-3743.	1.4	5
133	Management of Severe Intractable Sickle Cell Pain with Intrathecal Opioid Analgesia Blood, 2004, 104, 3742-3742.	1.4	Ο
134	Folic Acid Supplementation and Twin Pregnancy in Patients with Sickle Cell Disease Blood, 2004, 104, 3744-3744.	1.4	0
135	l -arginine levels are diminished in adult acute vaso-occlusive sickle cell crisis in the emergency department. British Journal of Haematology, 2003, 120, 532-534.	2.5	55
136	Effect of Hydroxyurea on Mortality and Morbidity in Adult Sickle Cell Anemia. JAMA - Journal of the American Medical Association, 2003, 289, 1645.	7.4	741
137	Treatment of painful sickle cell leg ulcers with topical opioids. Blood, 2002, 99, 1096-1096.	1.4	48
138	Sickle Cell Anaemia. Drugs, 2002, 62, 1143-1172.	10.9	84
139	Sickle cell disease: Current clinical management. Seminars in Hematology, 2001, 38, 307-314.	3.4	25
140	Iron overload is a determinant of morbidity and mortality in adult patients with sickle cell disease. Seminars in Hematology, 2001, 38, 30-36.	3.4	121
141	In vitro exposure to hydroxyurea reduces sickle red blood cell deformability. American Journal of Hematology, 2001, 67, 151-156.	4.1	22
142	Molecular characteristics of pediatric patients with sickle cell anemia and stroke. American Journal of Hematology, 2001, 67, 179-182.	4.1	67
143	Ethical issues in the management of sickle cell pain. American Journal of Hematology, 2001, 68, 127-132.	4.1	13
144	Effect of α-Globin Genotype on the Pathophysiology of Sickle Cell Disease. Fetal and Pediatric Pathology, 2001, 20, 107-121.	0.3	11

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145	Purified Poloxamer 188 for Treatment of Acute Vaso-occlusive Crisis of Sickle Cell Disease. JAMA - Journal of the American Medical Association, 2001, 286, 2099.	7.4	173
146	Iron overload is a determinant of morbidity and mortality in adult patients with sickle cell disease. Seminars in Hematology, 2001, 38, 30-36.	3.4	86
147	Sequential nitric oxide measurements during the emergency department treatment of acute vasoocclusive sickle cell crisis. , 2000, 64, 15-19.		44
148	Cost-effectiveness of hydroxyurea in sickle cell anemia. American Journal of Hematology, 2000, 64, 26-31.	4.1	77
149	Hydration of sickle erythrocytes using a herbal extract ( <i>Pfaffia paniculata</i> ) <i>in vitro</i> . British Journal of Haematology, 2000, 111, 359-362.	2.5	2
150	Elevated serum and bronchoalveolar lavage fluid levels of interleukin 8 and granulocyte colonyâ€stimulating factor associated with the acute chest syndrome in patients with sickle cell disease. British Journal of Haematology, 2000, 111, 482-490.	2.5	1
151	Acute Chest Syndrome in Sickle Cell Anemia. Journal of Intensive Care Medicine, 2000, 15, 123-125.	2.8	2
152	Causes and Outcomes of the Acute Chest Syndrome in Sickle Cell Disease. New England Journal of Medicine, 2000, 342, 1855-1865.	27.0	1,062
153	Elevated serum and bronchoalveolar lavage fluid levels of interleukin 8 and granulocyte colony-stimulating factor associated with the acute chest syndrome in patients with sickle cell disease. British Journal of Haematology, 2000, 111, 482-490.	2.5	34
154	Greater erythrocyte deformability in world-class endurance athletes. American Journal of Physiology - Heart and Circulatory Physiology, 1999, 276, H2188-H2193.	3.2	36
155	Erythropoietic activity in patients with sickle cell anaemia before and after treatment with hydroxyurea. British Journal of Haematology, 1999, 105, 491-496.	2.5	65
156	Erythropoietic activity in patients with sickle cell anaemia before and after treatment with hydroxyurea. British Journal of Haematology, 1999, 105, 491-496.	2.5	1
157	7 Sickle cell disease: clinical management. Best Practice and Research: Clinical Haematology, 1998, 11, 185-214.	1.1	61
158	Viral Burden and Disease Progression in HIV-1–Infected Patients with Sickle Cell Anemia. American Journal of Hematology, 1998, 59, 199-207.	4.1	35
159	Reticulocyte Hemoglobin: <i>An Integrated Parameter for Evaluation of Erythropoietic Activity</i> . American Journal of Clinical Pathology, 1997, 108, 133-142.	0.7	66
160	Molecular basis of asymptomatic β-thalassemia major in an African American individual. , 1997, 69, 196-199.		4
161	BFU-E colony growth in response to hydroxyurea: Correlation between in vitro and in vivo fetal hemoglobin induction. , 1997, 56, 252-258.		34
162	Hydroxyurea and Sickle Cell Anemia Clinical Utility of a Myelosuppressive "Switching―Agent. Medicine (United States), 1996, 75, 300-326.	1.0	294

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163	Factitious sickle cell acute painful episodes: A secondary type of Munchausen syndrome. , 1996, 53, 254-258.		14
164	Neonatal Screening for Sickle Cell Disease in a Metropolitan University Hospital: Efficacy and Problems. Journal of Medical Screening, 1994, 1, 229-232.	2.3	10
165	Hepatitis C in Sickle Cell Anemia. Journal of Clinical Gastroenterology, 1994, 18, 206-209.	2.2	56
166	Evaluation of Serum Immunoglobulins by Protein Electrophoresis and Rate Nephelometry Before and After Therapeutic Plasma Exchange. American Journal of Clinical Pathology, 1992, 98, 243-248.	0.7	14
167	Lack of Effect of Pentoxifylline on Red Blood Cell Deformability. Journal of Clinical Pharmacology, 1992, 32, 1050-1053.	2.0	14
168	Sickle cell anemia with few painful crises is characterized by decreased red cell deformability and increased number of dense cells. American Journal of Hematology, 1991, 36, 122-130.	4.1	81
169	Erythrocyte glycophorin B deficiency may occur by two distinct gene alterations. American Journal of Hematology, 1991, 37, 57-58.	4.1	23
170	Treatment of pain in adults with sickle cell disease. American Journal of Hematology, 1990, 34, 49-54.	4.1	60
171	Modified method of exchange transfusion in sickle cell disease. Journal of Clinical Apheresis, 1990, 5, 183-187.	1.3	18
172	The Prevalence of Avascular Necrosis in Sickle Cell Anemia: Correlation with α-Thalassemia. Hemoglobin, 1989, 13, 649-655.	0.8	51
173	Effect of hydroxyurea on the rheological properties of sickle erythrocytes in vivo. American Journal of Hematology, 1989, 32, 104-111.	4.1	134
174	Erythrocytes in Hb SC disease are microcytic and hyperchromic. American Journal of Hematology, 1988, 28, 37-39.	4.1	19
175	Erythrocyte Rh antigens increase with red cell age. American Journal of Hematology, 1986, 23, 19-24.	4.1	12
176	Biosynthetic Evidence for Stability of Hb N-Baltimore. Hemoglobin, 1985, 9, 489-494.	0.8	1
177	Serum Immunoglobulin Levels in Patients Having Sickle Cell Syndromes. American Journal of Clinical Pathology, 1980, 73, 394-396.	0.7	21
178	FAILURE TO DEMONSTRATE RED CELL MEMBRANE PROTEIN ABNORMALITIES IN SICKLE CELL ANAEMIA. British Journal of Haematology, 1980, 46, 627-629.	2.5	7
179	Delayed hemolytic transfusion reaction in sickle cell anemia. Transfusion, 1980, 20, 477-477.	1.6	5
180	The diagnosis of pulmonary thromboembolism in sickle cell disease. American Journal of Hematology, 1979, 7, 219-232.	4.1	47

#	Article	lF	CITATIONS
181	Leukapheresis for hyperviscosity. Transfusion, 1979, 19, 787-787.	1.6	9
182	Hemoglobin Potomac: Clinical Picture, Biosynths and Stability. Hemoglobin, 1978, 2, 447-451.	0.8	3
183	Stimulation of Lipid Synthesis in Reticulocytes as a Response to Membrane Damage. Blood, 1974, 44, 263-273.	1.4	7
184	Sickle Cell Pain: Biology, Etiology, and Treatment. , 0, , 497-524.		5
185	Pharmacological interventions for painful sickle cell vaso-occlusive crises in adults. The Cochrane Library, 0, , .	2.8	5