Serge Pissard

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/3141133/publications.pdf

Version: 2024-02-01

1937685 1372567 17 95 4 10 citations h-index g-index papers 17 17 17 208 docs citations times ranked citing authors all docs

#	Article	IF	CITATIONS
1	Haematological determinants of cardiac involvement in adults with sickle cell disease. European Heart Journal, 2016, 37, 1158-1167.	2.2	45
2	A new gene associated with a \hat{l}^2 -thalassemia phenotype: the observation of variants in SUPT5H. Blood, 2020, 136, 1789-1793.	1.4	13
3	Maternal consumption of quinine-containing sodas may induce G6PD crises in breastfed children. European Journal of Pediatrics, 2017, 176, 1415-1418.	2.7	9
4	Inherited or acquired modifiers of iron status may dramatically affect the phenotype in dehydrated hereditary stomatocytosis. European Journal of Haematology, 2018, 101, 566-569.	2.2	9
5	Improvements in phenotype studies of hemoglobin disorders brought by advances in reversedâ€phase chromatography of globin chains. International Journal of Laboratory Hematology, 2015, 37, 279-286.	1.3	5
6	HFE Genotyping by Amplification Refractory Mutation System–Denaturing HPLC. Clinical Chemistry, 2002, 48, 769-772.	3.2	3
7	A Second Observation of the Rare Frameshift Mutation in the \hat{l}^2 -Globin Gene: Codon 46 (+A) (Hbb:c.138_139insA). Hemoglobin, 2011, 35, 157-161.	0.8	2
8	Hb Savaria [α49(CE7)Ser→Arg; <i>HBA2</i> : c.150C > A]: A New Case and Complete Description. Hemoglobin, 2016, 40, 267-269.	0.8	2
9	Evidence for a gene conversion in a Hb Arya Carrier [α codon 47 Asp>Asn, Hb A1(or Hb A2):c.142 G>A]. International Journal of Laboratory Hematology, 2017, 39, e55-e59.	1.3	2
10	Analysis of Rare deletional Thalassemia Using Custom CGH Array DNA Chip. Blood, 2010, 116, 4278-4278.	1.4	2
11	Hb Olivet (<i>HBA1</i> : C.40G > A; p.Ala14Thr), a Novel Silent Hemoglobin Variant in Two Families of Distinct Origin. Hemoglobin, 2016, 40, 349-352.	0.8	1
12	Short in-Frame Insertions/Deletions in the Coding Sequence of the α-Globin Gene. Consequences of the 3D Structure and Resulting Phenotypes: Hb Choisy as an Example. Hemoglobin, 2018, 42, 287-293.	0.8	1
13	HFE genotyping by amplification refractory mutation system-denaturing HPLC. Clinical Chemistry, 2002, 48, 769-72.	3.2	1
14	Red Blood Cells Free α Hemoglobin Pool: A Biomarker to Monitor Imbalanced α/Non α-Globin Chain Synthesis in β-Thalassemia Intermedia: The ALPHAPOOL Study. Hemoglobin, 2019, 43, 344-344.	0.8	0
15	Molecular Approach to Prenatal Diagnosis of Hemoglobinopathies and Thalassemias. Hemoglobin, 2019, 43, 335-335.	0.8	0
16	Hb Angers: A new α2â€globin variant [α2 (140)(HC2) TyrÂ→ÂSer; <i>HBA2</i> : C.422 A>C] with increased oxygen affinity leading to erythrocytosis. International Journal of Laboratory Hematology, 2021, 43, e114-e117.	1.3	0
17	Le déficit en G6PD. Journal De Pediatrie Et De Puericulture, 2021, 34, 244-250.	0.0	0