

Hemoglobin EE Disease: A Case Report

Awad M. Al-Qahtani¹, Mohamed S. M. Khalil² and Essam M. Ahmed³

¹ Najran University

Received: 12 December 2018 Accepted: 2 January 2019 Published: 15 January 2019

5

Abstract

Background: Hemoglobin E variant results from a G?A substitution resulting consequently in abnormal processing for messenger mRNA. Its interactions with various forms of ? and ? thalassemia produce a very wide range of clinical syndromes. Methods: A Consent has been taken from a 26-year-old male. CBC, Glucose, Vitamin B 12 , C-peptide, estradiol (E 2), follicle stimulating hormone (FSH), free triiodothyronine (FT 3), free thyroxine (FT 4), luteinizing hormone (LH), prolactin, parathyroid hormone (PTH), testosterone, thyroid stimulating hormone (TSH) and vitamin D 3 (25-OH) and HPLC for hemoglobin separation were performed. Results: There was a history of hemolytic anemia due to infection with malaria and just one blood transfusion. There were no significant clinical findings such as organomegaly, icterus, or thalassemic bone changes. C-peptide, E 2 and TSH results were slightly above the normal range. Vit D was slightly insufficient. No Helicobacter pylori Antigen is stool and no clinical abnormalities. All the Hb were abnormal. The patient has low HDL-C which could not be explained. Also the slightly increased hormones of E2 and TSH, the slightly increased C-peptide could not be explained and this requires further investigations. Conclusion: The case was reported as abnormal hemoglobin (Hb E) masking the Hb A2 on HPLC.

23

Index terms—

G?A substitution resulting consequently in abnormal processing for messenger mRNA. Its interactions with various forms of ? and ? thalassemia produce a very wide range of clinical syndromes. Methods: A Consent has been taken from a 26-year-old male. CBC, Glucose, Vitamin B 12 , C-peptide, estradiol (E 2), follicle stimulating hormone (FSH), free triiodothyronine (FT 3), free thyroxine (FT 4), luteinizing hormone (LH), prolactin, parathyroid hormone (PTH), testosterone, thyroid stimulating hormone (TSH) and vitamin D 3 (25-OH) and HPLC for hemoglobin separation were performed.

Results: There was a history of hemolytic anemia due to infection with malaria and just one blood transfusion. There were no significant clinical findings such as organomegaly, icterus, or thalassemic bone changes. C-peptide, E 2 and TSH results were slightly above the normal range. Vit D was slightly insufficient. No Helicobacter pylori Antigen is stool and no clinical abnormalities. All the Hb were abnormal. The patient has low HDL-C which could not be explained. Also the slightly increased hormones of E2 and TSH, the slightly increased C-peptide could not be explained and this requires further investigations.

1 Conclusion:

The case was reported as abnormal hemoglobin (Hb E) masking the Hb A2 on HPLC.

2 I.

Background round 7% of the world's population comprises hemoglobinopathy gene carriers. Almost a total of 1317 Hb variants have been identified (HbVar database) 1 , the four most common worldwide are Hb S, Hb E,

6 DISCUSSION

42 Hb C, and Hb D, in the order of decreasing prevalence 2 . Hb E is the most prevalent variant in Southeast Asia
43 (Thailand, Myanmar, Cambodia, Laos, Vietnam), where its prevalence is 30-60% 3,4,5,6 . The prevalence of Hb
44 E in India is about 0-3.5% with an increased clustering in Kolkata (22%) and Assam (50-80%) 7 .

45 Hemoglobin E variant results from a G?A substitution in codon 26 of the ? globin gene, this produces an
46 abnormal hemoglobin (glutamate is replaced by lysine) and activates a cryptic splice site at codon 25-27 of the
47 ?-globin gene, resulting consequently in abnormal processing for messenger RNA (mRNA). The level of normally
48 spliced mRNA become reduced and because a new stop codon is generated, the abnormally spliced mRNA
49 become nonfunctional 7,8 . Fortunately, only a minor activation of the alternative splicing pathway the mutation
50 is associated with this mutation. Hence there is only a moderate reduction of the normally spliced ? E globin
51 mRNA ??2 .

52 Hb E trait and Hb EE disease are mild disorders. Although Hb E alone does not cause any significant clinical
53 problems, its interactions with various forms of ? and ? thalassemia produce a very wide range of clinical
54 syndromes of varying severity 8 , 9 .

55 Hb E has several compound heterozygotes with common and uncommon ?-globin or ?-globin gene mutations,
56 the most serious Hb E syndrome is Hb E? 0 -thalassemia. Different phenotypes could be noticed with the
57 compound heterozygote state of Hb E?thalassemia ranging from a complete lack of symptoms to transfusion
58 dependency 7,8,3,10 .

59 Experiments were carried out in vitro at temperatures ranging from 38 to 41°C showed that there was mild
60 instability of Hb E but there is no evidence that this is the case in vivo ??5 9 . It is noticeable that, E allele
61 causes mild thalassemia, while ? E ? 0 thalassemia shows a severe phenotype, this marked paradox in phenotype
62 could not be explained up till now. It is reported that Hb E is sensitive to oxidative stress {HbVar database}.
63 Does this or other properties of Hb E can contribute to the severity of the disease? This question is still waiting
64 for an answer 11 .

65 The aim of this report is to present a case of Hb EE discovered accidentally during a routine work trying to
66 cast shadow on some parameters.

67 3 II.

68 4 Case Report

69 A Consent has been taken from a 26-year-old male, from Kolkata, India, came to Najran University Hospital,
70 Saudi Arabia for routine investigation. He did not complain from anemia or receive treatment. He gave a history
71 of hemolytic anemia because of infection with malaria and only one blood transfusion. On examination, there
72 were no significant clinical findings such as organomegaly, icterus, or thalassemic bone changes.

73 CBC were carried out using Sysmex XS 500i (Sysmex, <https://www.sysmex.com/>). The results showed Hb
74 of 13 g/dL, red blood cell (RBC) count of 6.8 x 10 12 /L, mean corpuscular volume (MCV) of 55.6 fl, mean
75 corpuscular hemoglobin (MCH) of 19.6 pg, MCHC concentration of 36 g/dL, and RBC distribution width (RDW-
76 CV) of 20.7%. Peripheral blood smear A showed frequent target cells, and spherocytes as shown in Fig. (1).
77 Serum biochemical analysis were carried out using COBAS C311 (Roche, <https://www.roche.com/>). Results
78 were normal for liver and kidney functions except for mild increase in bilirubin (1.39 mg/dL, mostly indirect
79 of 0.98 mg/dL). Lipid profile was normal except for low high density lipoprotein cholesterol (HDL-C) of 20.6
80 mg/dL...Iron, UIBC and ferritin were found to be 73.3 ug/dL, 250 ug/dL and 149.6 ug/L respectively. Glucose
81 was within normal ranges. Vitamin B 12 , C-peptide, estradiol (E 2), follicle stimulating hormone (FSH), free
82 triiodothyronine (FT 3), free thyroxine (FT 4), luteinizing hormone (LH), prolactin, parathyroid hormone
83 (PTH), testosterone, thyroid stimulating hormone (TSH) and vitamin D 3 (25-OH) were done. C-peptide, E
84 2 and TSH results were slightly above the normal range. Vit D was slightly insufficient. Some parameters
85 were indicated in table (1). Hb A1c did not give results on D-10 HPLC that is why we thought of Hb variant.
86 Hemoglobin (Hb) separation by high performance liquid chromatography (HPLC) using the D-10 instrument
87 (Bio-Rad Laboratories Hercules, California, USA) as in Fig. (2). Nearly all the Hb were abnormal and it was
88 eluted at the Hb A2 window with retention time (rt) of 3.17 minutes. When repeated on Variant II HPLC system
89 ((Bio-Rad) with use of the Variant II Thalassemia Short Program, it showed 86.1 % abnormal Hb with rt of
90 3.8 min along with 2.28 % adult Hb (Hb A) and Hb F around 3% of total Hb. Stool examination was negative
91 for parasites. Helicobacter pylori Antigen is stool and Abs in serum were negative by One Step H. pylori test
92 device (ABON Biopharm, China). Malaria Ag in blood was negative by malaria P/F/Pan rapid test device from
93 ABON, China. Clinical examination showed no abnormality.

94 5 III.

95 6 Discussion

96 The majority of hemoglobinopathy present in the western and eastern provinces of Saudi Arabia. Abuzenadah
97 et al. 7,3 , reported a great heterogeneity at the molecular level in the western province and attributed this to
98 the large population of immigrants there. Hb E was one of the seven common ?-thalassemia alleles reported.

99 Haemoglobinopathy is not common in Najran city. Prevalence reported by Memish et al., 12 of 14.7 % mostly
100 sickle mutations of 14.1 % 8 .

101 Considering his history, clinical findings and laboratory findings, the diagnosis in this case was homozygous
102 Hb EE disease. The patient showed very mild, clinically asymptomatic, hemolytic microcytic hypochromic
103 anemia with many target cells on peripheral blood smear characteristic of Hb EE disease which consistent with
104 the classical presentation of the disease 9 . The presence of mild increase in Hb F indicates the mildness of
105 the pathophysiology of the disease. The presence of a minute quantity of Hb A on HPLC was explained by
106 post-translational modification of Hb E 13 .

107 The patient gave a history of high fever due to malaria infection, with a hemolytic attack and he received a
108 blood transfusion which could match the published of the instability of Hb E in high fever 14 . We noticed the
109 presence of many spherocytes in the peripheral blood smear with an increase of MCHC. Similar findings occur
110 in Hb C due an increase in the activity of K: Clcotransport that induces the loss of K + and subsequently of
111 intracellular water 15 . The patient has low HDL-C which could not be explained. Also the slightly increased
112 hormones of E2 and TSH, the slightly increased C-peptide could not be explained and recommended for further
113 investigation.

114 IV.

115 7 Conclusion

116 Thus we report a case of abnormal hemoglobin (Hb E) masking the Hb A2 on HPLC. Knowledge of such a
117 condition would help in prevention of misdiagnosis. Also we focused on some abnormal findings and recommended
them for further research. ^{1 2}

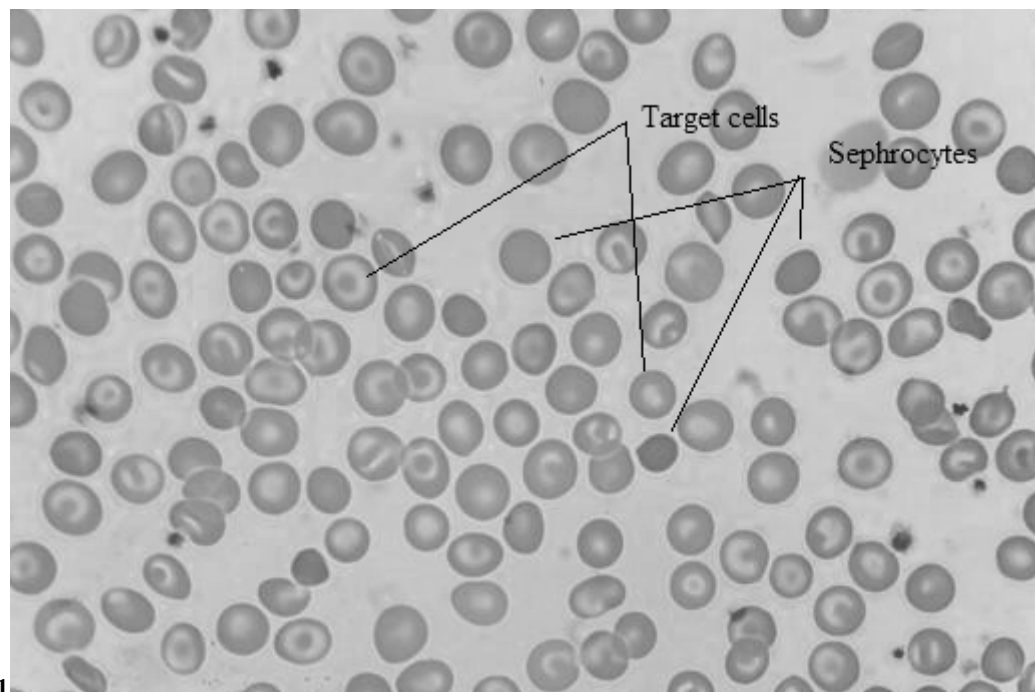
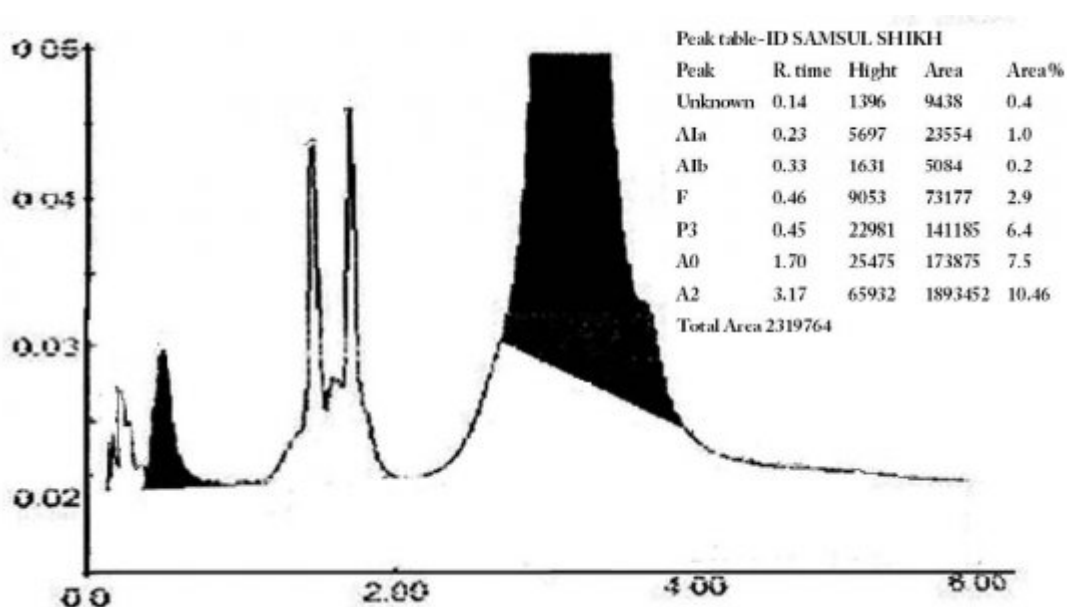


Figure 1: Fig. 1 :

118

¹© 2019 Global Journals

²Mr./Muhammad Shoib Aslam, Medical laboratory technologist Najran University Hospital Ksa, for his cooperation and helps during different stages of this work.



210

Figure 2: Fig. 2 : 10 Hemoglobin

Patient Data
 Sample ID: 80
 Patient ID:
 Name:
 Physician:
 Sex:
 DOB:
 Comments:

Analysis Data
 Analysis Performed: 27/11/2017
 Injection Number: 16300
 Run Number: 48
 Rack ID: 0003
 Tube Number: 2
 Report Generated: 27/11/2017
 Operator ID:

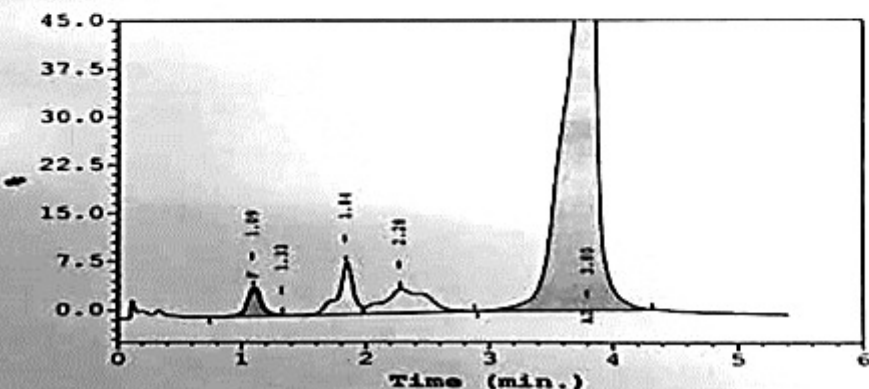
Peak Name	Calibrated Area %	Area %	Retention Time (min)	Peak Area
F	2.2*	---	1.09	54982
P2	---	0.1	1.33	1756
P3	---	4.8	1.84	120225
Ao	---	6.3	2.28	159039
A2	86.1*	---	3.80	2184319

Total Area: 2,520,320

F Concentration = 2.2*%
A2 Concentration = 86.1*%

*Values outside of expected ranges

Analysis comments:



3

Figure 3: Fig. 3 :

1

Investigation	Values	Investigation	Values
Hb	13 g/dl	WBC	11x10 ⁹ /L
RBC	6.8x10 ¹² /L	Eosinophilia	1.9 x 10 ⁹ /L
MCV	55.6 Fl	HDL-C	20.6 mg/dl
MCH	19.6 pg	C-Peptide	6.97 ng/ml
MCHC	36.3 g/dl	E2	46.88 pg/ml
Total bilirubin	1.39 mg/dl	TSH	6.8 pIU/ml
Indirect bilirubin	0.98 mg/dl	Vit-D (25-OH)	27 ng/ml
HDL	20.6 mg/dl		
UIBC	250 µg/dl		
Ferritin	149 µg/dl		
Iron	73.3 µg/dl		

Figure 4: Table 1 :

7 CONCLUSION

119 .1 Acknowledgement

120 The team of this case report acknowledge

121 [Hemoglobin and Disease] , E E Hemoglobin , Disease . (A Case Report)

122 [Oxford and Kingdom ()] , United Oxford , Kingdom . *Blackwell Science* 2001. p. .

123 [Little and Roberts ()] 'A review of variant hemoglobins interfering with hemoglobin A1c measurement'. R R
124 Little , W L Roberts . *J Diabetes Sci Technol* 2009. 3 p. .

125 [Fucharoen et al. ()] 'Clinical manifestation of betathalassemia/hemoglobin E disease'. S Fucharoen , P Ketvichit
126 , P Pootrakul , N Siritanaratkul , A Piankijagum , P Wasi . *J Pediatr Hematol Oncol* 2000. 22 p. .

127 [Olivieri et al. ()] 'Hb E/beta-thalassaemia: a common & clinically diverse disorder'. N F Olivieri , Z Pakbaz ,
128 E Vichinsky , Indian . *J Med Res* 2011. 134 p. .

129 [Yedla et al. ()] 'Hemoglobin E disease and glycosylated hemoglobin'. N Yedla , M S Kuchay , A Mithal . *Indian*
130 *J Endocr Metab* 2015. 19 p. .

131 [Katsanis et al. ()] 'Hemoglobin E: a common hemoglobinopathy among children of Southeast Asian origin'. E
132 Katsanis , K H Luke , E Hsu , J R Yates . *CMAJ* 1987. 137 p. .

133 [Weatherall ()] 'Hemoglobinopathies worldwide: present and future'. D Weatherall . *Curr Mol Med* 2008. 8 p. .

134 [Kohne ()] 'Hemoglobinopathies: clinical manifestations, diagnosis, and treatment'. E Kohne . *Dtsch Arztebl Int*
135 2011. 108 p. .

136 [Weatherall and Clegg ()] 'Inherited haemoglobin disorders: an increasing global health problem'. D J Weatherall
137 , J Clegg . *Bull World Health Organ* 2001. 79 p. .

138 [Rees et al. ()] 'Is hemoglobin instability important in the interaction between hemoglobin E and b thalassemia?'.
139 D C Rees , J B Clegg , D J Weatherall . *Blood* 1998. 92 p. .

140 [Acquaye et al. ()] 'Non-benign sickle cell anaemia in western Saudi Arabia'. J K Acquaye , Omer A Ganeshaguru
141 , K Sejeny , S A Hoffbrand , AV . *Br J Haematol* 1985. 60 p. .

142 [Memish and Saeedi ()] 'Six-year outcome of the national premarital screening and genetic counseling program
143 for sickle cell disease and betathalassemia in Saudi Arabia'. Z A Memish , M Y Saeedi . *Ann Saudi Med* 2011.
144 31 p. .

145 [Gibbons et al.] 'The ? and ?? thalassaeemias in association with structural haemoglobin variants'. R Gibbons ,
146 D R Higgs , N F Olivieri , W Wood . *The Thalassaemia Syndromes*, D J Weatherall, J B Clegg (ed.) (Fourth
147 ed)

148 [Fucharoen and Weatherall (2012)] *The hemoglobin E thalassemias. Cold Spring Harb Perspect Med*, S1
149 Fucharoen , D J Weatherall . doi: 10.1101/ cshperspect.a011734. 2012 Aug 1. 2.

150 [Nagel et al. (2003)] 'The paradox of hemoglobin SC disease'. R L Nagel , M E Fabry , M Steinberg . *Blood Rev*
151 2003 Sep. 17 (3) p. . (Review)