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3 **Rare double heterozygous of HbD/HbG in a**
4 **Nigerian: A case report**
5

6 **Abstract**

7 **Aim:** To advocate the use of newer and improved methods towards accurate diagnosis of
8 haemoglobinopathies

9 **Case presentation:** A rare case of double heterozygous of HbD/G in a pregnant female
10 Nigerian who had present to the antenatal clinic for routine Haemoglobin electrophoresis.
11 She had previously being diagnosed as HbAS using capillary electrophoresis and HPLC
12 techniques.

13 **Discussion:** Capillary zone electrophoretograms showed the presence of peaks in zone Hb
14 A, Hb D, C and a small peak in Z1 zone. Bio-Rad D10 chromatogram also indicated the
15 presence of four peaks which are identified as Hb A, Hb D, Hb G, and hybrid of HbD/HbG. A
16 peak in Hb D zone of capillary electrophoresis was due to co-migration of Hb D and Hb G
17 variants. Small peak in Z1 zone indicated the presence of alpha chain variant of HbG.

18 **Conclusion:** The case exemplifies the need to use more advanced methods, including DNA
19 analysis in order to accurately diagnose haemoglobinopathies in the nation with largest
20 burden of sickle cell disease.

21 **Key words:** Haemoglobinopathies, haemoglobin electrophoresis, heterozygous, high
22 performance liquid chromatography

23
24 **Introduction**

25 Haemoglobinopathies are a group of genetic disorders that lead to quantitative or qualitative
26 abnormalities in haemoglobin (Hb) variants. Thalassaemias are due to reduced or absent
27 production of structurally normal globin chains while sickle cell disease occurs because of
28 substitution of one amino acid by another. The change in amino acid sequence results in
29 haemoglobin variants with abnormal structures. Many of the haemoglobin variants do not
30 cause symptoms in heterozygous condition but can lead to varying degrees of anaemia and
31 other symptoms in homozygous states or when they coexist with thalassaemias.
32 Double heterozygosity is described when there is a change in the amino acid sequence in
33 both α and β chains of the same individual and it is very rare. Though, sporadic cases of
34 hybrid haemoglobins have been reported in other regions of the world, here we report the
35 very first case of double heterozygosity of an alpha-chain variant hemoglobin G and a beta
36 chain hemoglobin D in a Nigerian.

38 **Case Report**

39 A 28 year old female, Yoruba **by tribe from the Southwestern region of Nigeria** presented to
40 the laboratory for Haemoglobin electrophoresis as part of the routine antenatal investigation.
41 She has no significant past medical history. Her previous Haemoglobin electrophoresis
42 (verbal report from patient) by cellulose acetate at alkaline pH was reported as HbAS. About
43 3mls of venous blood was collected into an ethylenediamine_tetraacetic acid (EDTA) bottle.
44 Complete blood count and solubility tests were also carried out on the sample.

45 Capillary zone electrophoresis (CE) for the sample was carried out using automated Sebia
46 Minicap analyser (Sebia, France) according to the manufacturer's instructions and was
47 repeated with BIORAD D10 high performance liquid chromatography (HPLC) to further
48 identify and confirm the results. The electrophoretogram by Sebia Minicap analyser showed
49 four main peaks as follows: HbA zone (53.5%), HbD zone (22.4%), HbA2 zone (5.1%) which
50 slightly overlaps with an unknown peak in C zone (15.3%). There was also a small peak in
51 the Z1 (2.5%) indicating a variant of alpha chain (fig 1).

52 The BIORAD-D10 high performance liquid chromatography also showed four major peaks
53 HbA (32.5%) with retention time of 1.69 minutes, HbD (30.4%) with a retention time of
54 3.91minutes in an unknown window, HbG (17.1%) with retention time 4.09 minutes in Hb S-
55 window, and an hybrid of HbD/G (12.7%) at retention time of 4.41 minutes in an unknown
56 window (fig 2). A small peak of HbA2 (2%) is also noted on the electrophoretogram. The full
57 blood count showed essentially normal parameters: RBC= $4.3 \times 10^6/\mu\text{L}$, Heamoglobin=
58 12.5g/dl, mean cell corpuscular volume (MCV)=88FL, mean corpuscular haemoglobin
59 concentration (MCHC)=33g/dl, mean corpuscular haemoglobin (MCH)= 29pg The sickling
60 test, **a procedure in which red blood cells sickle in the presence of sodium metabisulphite (a**
61 **reducing agent)** was negative, however, solubility test was not done. The requesting
62 physician was advised on the need for DNA analysis to confirm the diagnosis. However
63 patient was lost to follow up.

64 **Figure 1:** Capillary zone electrophoresis pattern indicates peaks in HbA, Hb D, Hb C, HbA2
65 zones, and additional small peak in Z1 zone indicating alpha chain variant

Sample num.: 19

Name

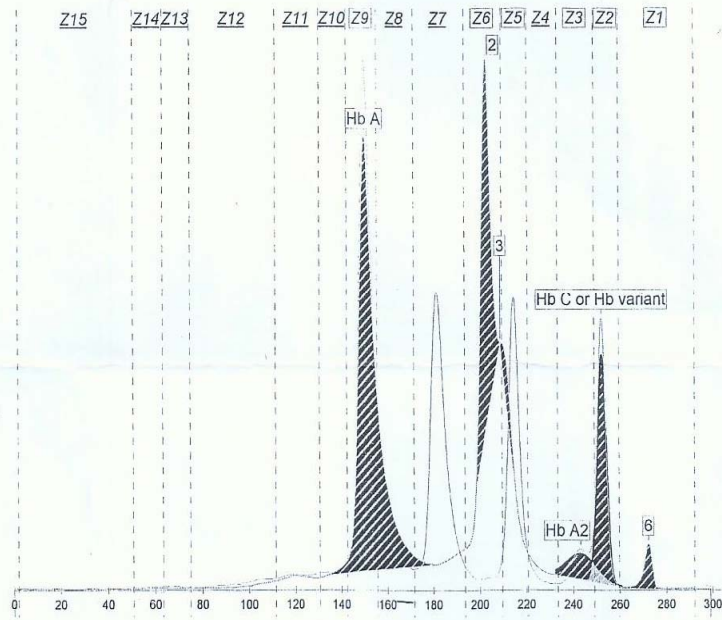
Date :

Date of birth

ID :

Sex

Sample date



Fractions	%	Ref. %	Ref. g/dl
Hb A	53.5		
2	22.4		
3	1.2		
Hb A2	5.1		
Hb C or Hb variant	15.8		
6	2.5		

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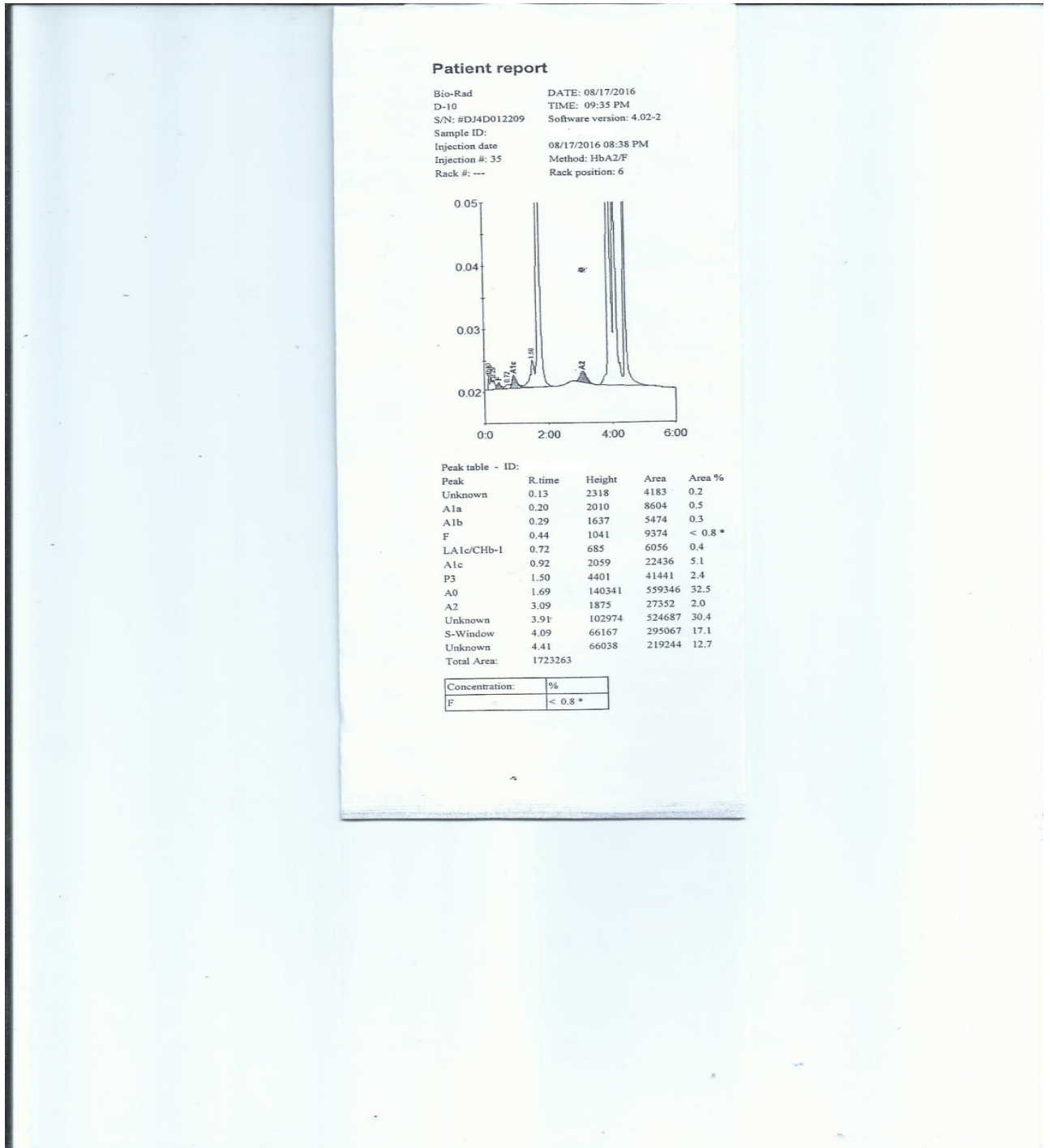
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72 Figure 2: High performance liquid chromatography chromatogram obtained in BIORAD D10
 73 showing 4 major peaks at retention times in minutes 1.69 (HbA), 3.91 (HbD), 4.09 (HbG),
 74 4.41 (hybrid of HbD/G)



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79 **Discussion**

80 There are a few variants of haemoglobin D. Hb D Punjab or Hb D Los Angeles is a type of
81 beta globin gene mutation at 121 codon resulting in replacement of glutamic acid with
82 glutamine (Glu->Gln).¹ The highest prevalence of HbD Punjab is among Sikhs in Punjab,
83 India where it is reported to be around 2%. Heterozygous HbD is a clinically silent
84 condition.¹ There is also HbD Ibadan which was discovered at the University College
85 Hospital Ibadan, Nigeria.² The prevalence of this is currently unknown. Hb D Ibadan results
86 from the replacement of Threonine with Lysine in position 87 of beta chain (Threonine->
87 Lysine).² Haemoglobin G Philadelphia is the most common alpha chain variant and is due to
88 replacement of asparagine with Lysine ($\alpha^{68 \text{ Asn} \rightarrow \text{Lys}}$).³ It occurs in less than 1% of the
89 population of West Africa.⁴

90 Presumptive identification of Hb variants was done by comparing the two methods used.
91 Both techniques clearly indicated that the predominant haemoglobin in this subject was HbA
92 (figs 1 & 2). The second peak on Capillary zone electrophoresis (fig 1) could be either
93 haemoglobin D or G or both travelling within zone 6.⁵ Following review of the literature we
94 confirmed that HbD elutes at approximately 3.91 minutes in D window similar to what was
95 obtained on our HPLC (fig.2).^{7,8} Furthermore, an unknown haemoglobin eluted in the S-
96 window on HPLC at retention time of 4.09 minutes but there was no corresponding pattern in
97 zone 5 (S zone) of CE, suggesting it is unlikely to be HbS but rather HbG which co-eluted
98 with HbD in zone 6 of capillary electrophoresis. Haemoglobins S, D and G also have the
99 same mobility on cellulose acetate paper at alkaline pH which explains the initial diagnosis of
100 HbAS as claimed by the patient. Cellulose acetate method is the routine technique in most
101 laboratories in Nigeria. The unknown pattern in the C window on Capillary electrophoresis is
102 the hybrid of HbD/G (fig 1). This correlates with the unknown peak eluted at 4.41 minutes in
103 the HPLC (fig 2). The inheritance of alpha-chain defects such as HbG-Philadelphia usually
104 results in formation of hybrid haemoglobins.⁹ The small peak of 2.1% in Z1 zone highly
105 suggests the presence of alpha variant of HbG. The haematological parameters and indices
106 were normal in this patient. The patient had no clinical symptoms and had only presented to
107 the hospital for antenatal booking. Therefore, the clinical implications of this inheritance
108 cannot be determined at the moment. The limitation of this study is non-availability of
109 facilities to further confirm the identity of the various haemoglobins in this patient. DNA
110 sequencing of alpha and beta globin genes which is the confirmatory diagnostic method is
111 not readily available in Nigeria. The traditional method of haemoglobin electrophoresis using
112 cellulose acetate in alkaline pH will most probably misdiagnose this patient.

113 The fully automated methods such as HPLC and CE have replaced the cellulose acetate
114 electrophoresis as first-line in the diagnosis of haemoglobinopathies. Apart from the
115 advantages of resolution and automation both allow processing of large batches and require
116 very small samples volumes.^{10,11,12} Quantification and identification of larger proportion of
117 variant haemoglobins can be made. The major disadvantage of HPLC is that it separates
118 glycosylated and other derivatives of haemoglobin making its interpretation complex,
119 however this does not occur with CE. CE has also been found to be more accurate and
120 sensitive for detecting Hb variants than cellulose acetate paper.^{10,13}

121 **Conclusion:** This case exemplifies the relevance of newer techniques in diagnosis of
122 haemoglobinopathies. Quantitative haemoglobin electrophoreses such as Capillary
123 electrophoresis and HPLC have been recently available in some diagnostic laboratories in

124 Nigeria and currently not accessible or affordable to the general population. More cases may
125 be found in the near future as these diagnostic facilities become readily available. It will also
126 enable investigations into the prevalence of Haemoglobin D and G among Nigerians.

127 **Consent: Consent form obtained from the patient.**

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129 **References**

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