

Original Research Article

Association between Non-Secretion of ABH Antigens and Sickle Cell Anaemia

1 ABSTRACT

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Aim: To determine whether non-secretion of ABH blood group antigens was associated with Sickle Cell Anaemia.

Materials and Methods: Haemagglutination inhibition test was carried out on saliva samples from 300 individuals; 100 of whom had haemoglobin (Hb) genotype AA, 100 HbAS, 50 HbAC and 50 HbSS. ABO blood grouping was carried out by standard methods and Haemoglobin genotype test was performed by cellulose acetate electrophoresis technique.

Results: Eighteen percent (18%) of HbAA, 23% of HbAS, 18% of HbAC and 42% of HbSS individuals were non-secretors of ABH antigens ($p = 0.007$). Non-secretion of ABH substances was more associated with HbSS persons than HbAA ($p = 0.002$), HbAS ($p = 0.016$) and HbAC ($p = 0.009$) individuals.

Conclusion: Non-secretion of ABH blood group substances is associated with Sickle Cell Anaemia.

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Keywords: Haemoglobin genotype, ABO blood group, Sickle cell anaemia, ABH antigens, Secretor status

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1. INTRODUCTION

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In Southwestern Nigeria, in addition to normal haemoglobin A, haemoglobins S and C exist bringing about variants HbAA, HbAS, HbAC, HbSS, HbCC and HbSC among the people in the region [1]. Haemoglobinopathies especially sickle cell anaemia poses a lot of health challenges in Nigeria [2]. Sickle cell anaemia (SCA) is an inherited disorder caused by mutation resulting in replacement of amino-acid glutamic acid with valine at the 6th base position of the beta globin chain. It is a genetic blood disorder characterized by the presence of 2 alleles of the abnormal haemoglobin S (HbSS) in the red cell instead of HbAA with high morbidity and mortality rates [2]. It is recognized by the United Nations as a global public health concern and the World Health Organisation has recommended that by 2020, half of

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20 its members should have set up Sickle Cell Anaemia (SCA) control programmes [3].
21 Worldwide, SCA is estimated to affect 20-25 million people and annually about 300,000
22 children are born with the disorder [4]; approximately 250,000 of whom are in sub-Saharan
23 Africa [5] with 50-80% of affected children dying before the age of 5 years [6].
24 The ABO blood group and secretor status of a person are inherited independently. While the
25 ABH (FUT 1) gene codes for the ABO blood group, the secretor (FUT 2) gene interacts with
26 the ABH (FUT 1) gene to determine the secretor status of an individual [7]. Individuals can
27 be homozygous (SeSe) or heterozygous (Sese) secretors or non-secretors (sese). Non-
28 secretion of ABH antigens has been associated with a number of non-communicable
29 diseases and disorders such as autoimmune diseases [8, 9], blood clotting and thrombotic
30 diseases [10, 11], immunological disorders [12], myocardial infarction [13, 14], rheumatic
31 heart disease [15, 16], duodenal ulcers [17]. Apart from one study [18] which investigated
32 frequency distribution of secretors and non-secretors in HbAA and HbSS individuals in Zaria,
33 Northwestern Nigeria, we are not aware of any other investigation that has related secretor
34 status with haemoglobin variants. In this study, we hypothesized that secretor status varied
35 significantly with haemoglobin variants and that non-secretion of ABH antigens was
36 associated with sickle cell anaemia.

37 38 **2. METHODOLOGY**

39 40 **2.1 Study Area and Population**

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42 This study was carried out in Osogbo, Southwestern Nigeria. It is the capital of Osun State.
43 Osogbo city seats the Headquarters of both Osogbo Local Government Area (situated at
44 Oke Baale Area of the city) and Olorunda Local Government Area (situated at Igbonna Area
45 of the city). It is some 88 kilometers by road northeast of Ibadan with coordinates Latitude
46 7.767-7.770°N and Longitude 4.557-4.567°E. A total of 300 participants were screened for
47 this study: 100 HbAA individuals, 100 HbAS, 50 HbAC and 50 HbSS. They were drawn from
48 apparently healthy staff, students and patients of LAUTECH Teaching Hospital visiting the

49 General Out Patient Department for routine examination. Informed consent was obtained
50 from all the participants. Ethical approval for this study was obtained from the Ethical
51 Committee of the College of Health Sciences, Ladoke Akintola University of Technology,
52 Osogbo, Osun State, Nigeria.

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54 **2.2 Collection of Blood Samples**

55 Blood samples were collected for Haemoglobin genotype test and ABO blood grouping. A
56 sample of 3 ml of venous blood was collected from each participant into
57 ethylenediaminetetraacetic acid (EDTA) bottle. Haemoglobin genotype test was performed
58 using cellulose acetate electrophoresis method [1]. In an alkaline pH (8.2-8.6), Hb is a
59 negatively charged molecule and will migrate towards the anode. Different Hbs move at
60 different rates depending on their net charge which is controlled by the amino acid
61 composition of their globin chain. The ABO grouping system is based on agglutination
62 reaction. When a red blood cell carrying an antigen is exposed to its corresponding antibody,
63 they react with each other to form agglutination or clumping. ABO blood group tests were
64 performed by standard techniques [19].

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66 **2.3 Collection of Saliva Samples**

67 Saliva samples were collected from participants for the determination of their secretor status;
68 2 ml of saliva was collected from each participant for determination of secretor status using
69 haemagglutination inhibition test [20]. If the saliva is from a secretor, the soluble blood
70 antigens in it react and neutralize the antibodies in the antiserum. So when red blood cells of
71 appropriate blood group are added to the test mixture of the saliva and antiserum, there will
72 be no free antibody to agglutinate them because the antibodies have already been
73 neutralized by the antigens in the saliva. Therefore the reaction will be negative for
74 agglutination. However, if the saliva is from a non-secretor, there will be no blood group
75 antigens in it and so the antibodies in the antiserum will not be neutralized but free to react

76 with appropriate test cells when added to produced agglutination. Laboratory investigations
77 were carried out on samples collected in the Research Laboratory of the Department of
78 Medical Laboratory Science, College of Health Sciences, Ladoke Akintola University of
79 Technology, Osogbo, Nigeria.

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81 **2.4 Data Analysis**

82 The statistical package for social sciences (SPSS version 16) was used for statistical
83 analysis. Differences in proportions or percentages were tested by Chi-square test. A p-
84 value of < 0.05 was considered significant.

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91 **3. RESULTS**

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93 A total of 300 persons comprising 100 HbAA, 100 HbAS, 50 HbAC and 50 HbSS individuals
94 participated in this study. Table 1 shows the age and sex distributions of the study
95 population. There were no significant differences in the age ($p = 0.998$) and sex ($p = 0.718$)
96 distributions among the four groups of haemoglobin variants.

97 The distributions of the haemoglobin variants of the study participants in relation to secretor
98 status are given in Table 2. Of the 100 individuals with HbAA, 18% were non-secretors;

99 23%, 18% and 42% of the HbAS, HbAC and HbSS individuals respectively were non-

100 secretors. Non-secretion of ABH antigens varied significantly with haemoglobin variants (χ^2
101 = 11.99, $df = 3$, $p = 0.007$). Further Chi-square tests showed that non-secretion of ABH

102 antigens was more associated with HbSS individuals than HbAA individuals ($\chi^2 = 9.978$, $df =$

103 1, $p = 0.002$), HbAS individuals ($\chi^2 = 5.805$, $df = 1$, $p = 0.016$) and HbAC individuals ($\chi^2 =$

104 6.857, $df = 1$, $p = 0.009$). There was no significant variation in secretion of ABH antigens

105 among HbAA, HbAS and HbAC individuals ($\chi^2 = 0.938$, $df = 2$, $p = 0.626$). Altogether, non-

106 secretors in the HbSS group (42.0%) were significantly higher than those in the non-SS
 107 (HbAA, AS and AC) group (20.0%) ($\chi^2 = 11.163$, $df = 1$, $p < 0.001$).

108 The distributions of the haemoglobin variants of the study participants in relation to ABO
 109 blood group are given in Table 2. Group AB was excluded from the analysis due to its small
 110 number across the haemoglobin variants. There was no significant association in the
 111 distributions of haemoglobin variants in relation to ABO blood group ($\chi^2 = 5.69$, $df = 6$, $p =$
 112 0.458).

113 The distributions of the non-secretors of the study participants with respect to haemoglobin
 114 variants and ABO blood group is given in Table 3. Of the 100 AA individuals, 18 were non-
 115 secretors (10 non-group O and 8 group O); of the 100 AS individuals, 23 were non-secretors
 116 (14 non-group O and 9 group O); of the 50 AC individuals, 9 were non-secretors (6 non-
 117 group O and 3 group O) while 21 of the 50 SS individuals were non-secretors (14 non-group
 118 O and 7 group O). Altogether, of the 150 non-O blood group participants, 44 (29.3%) were
 119 non-secretors while 27 of the 150 (18.0%) group O participants were non-secretors ($\chi^2 =$
 120 5.332 , $df = 1$, $p = 0.021$).

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123 **Table 1: Age and Sex distributions among the Study Participants**

	Haemoglobin Variants				Total 300	p-value
	HbAA n=100(%)	HbAS n=100(%)	HbAC n=50(%)	HbSS n=50(%)		
Age group(years)						0.998
16 - 25	32(32.0)	35(35.0)	17(34.0)	15(30.0)	99(33.0)	
26 - 35	28(28.0)	27(27.0)	14(28.0)	15(30.0)	84(28.0)	
>36	40(40.0)	38(38.0)	19(38.0)	20(40.0)	117(39.0)	
Sex						0.718
Male	45(45.0)	48(48.0)	26(52.0)	27(54.0)	146(48.7)	
Female	55 (55.0)	52(52.0)	24(48.0)	23(46.0)	154(51.3)	

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Table 2: Distribution of the Haemoglobin Variants of the Study Participants in Relation to Secretor Status and ABO Blood Group

	Haemoglobin Variants				Total 300	p-value
	HbAA n=100(%)	HbAS n=100(%)	HbAC n=50(%)	HbSS n=50(%)		
Secretor status						0.007
Secretor	82(82.0)	77(77.0)	41(82.0)	29(58.0)	229(76.3)	
Non-secretor	18(18.0)	23(23.0)	09(18.0)	21(42.0)	71(23.7)	
ABO Blood Group						0.458
A	22(22.0)	23(23.0)	11(22.0)	16(32.0)	72(24.0)	
B	19(19.0)	26(26.0)	13(26.0)	14(28.0)	72(24.0)	
AB	02(2.0)	02(2.0)	01(2.0)	01(2.0)	06(2.0)	
O	57(57.0)	49(49.0)	25(50.0)	19(38.0)	150(50.0)	

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Table 3: Distribution of the Non-Secretors of the Study Participants with Respect to Haemoglobin Variants and ABO Blood Group

	Haemoglobin Variants				Total(NS)
	HbAA(NS)	HbAS(NS)	HbAC(NS)	HbSS(NS)	
ABO Blood Group					
Non-O	43(10)	51(14)	25(06)	31(14)	150(44)
O	57(08)	49(09)	25(03)	19(07)	150(27)
Total	100(18)	100(23)	50(09)	50(21)	300(71)

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NS: Non-Secretor

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4. DISCUSSION

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Previous studies in this study area had shown that secretor status was independent of sex

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[20]. A similar finding was reported in Calabar, South south Nigeria[21]. Similarly, in this

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study locality, the distribution of ABO blood group and haemoglobin variants had been

152 reported to be sex independent [1, 22] which were in line with ABO studies carried out in the
153 same region by other researchers [23, 24].

154 In this study, we tested the hypothesis that non-secretors were more associated with HbSS
155 compared to the other haemoglobin variants. The frequency of non-secretors in SCA (HbSS)
156 individuals was significantly higher than the frequency of non-secretors in the other
157 haemoglobin variants (HbAA, HbAS and HbAC) showing that secretor status varied
158 significantly with haemoglobin variants. A study in Northwestern Nigeria reported a higher
159 frequency of non-secretor in HbSS individuals compared to HbAA individuals [18].

160 Also in this study, non-secretion of ABH substances was more associated with persons of
161 non-O group compared to those of O group. Previous studies in the study area and
162 elsewhere had reported lower proportion of group O non-secretors compared to non-O
163 group non-secretors [7, 20]. Another study in the area showed that malaria was less
164 associated with group O secretors than non-group O secretors [22]. These studies showed
165 that with respect to ABO blood group system, more group O persons were secretors
166 compared to the other groups. The protective effect offered by group O individuals had been
167 linked to higher incidence of secretor compared to non-O group [7].

168 Non-secretion of ABH antigens has been associated with many non-communicable diseases
169 and disorders [8 - 17]. Similarly, sickle cell anaemia individuals are known to have several
170 complications including chronic pain, intermittent painful episodes, musculoskeletal
171 problems, stroke, pulmonary hypertension and septicaemia [3, 25]. It is not unlikely that the
172 complications exhibited by majority of the persons with sickle cell disorder might largely be
173 due to their inability to secrete ABH substances. Also, the observed association might be
174 linked with the Le^a antigens which are present in greater amounts on the epithelial surface of
175 non-secretors [26]. The positive interaction observed between HbSS and inability to secrete
176 ABH antigens could be suggestive of the fact that the sickle cell gene and the secretor gene
177 might directly or indirectly interact to confer susceptibility on persons with sickle cell
178 anaemia. We opine that the severity of symptoms and complications observed in HbSS

179 patients could be due to their inability to secrete ABH antigens. Further studies can be
180 carried out to confirm or disprove this view.

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182 **4. CONCLUSION**

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184 Secretor status varies significantly with haemoglobin variants and inability to secrete ABH
185 antigens is associated sickle cell anaemia.

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189 **COMPETING INTERESTS**

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191 Authors have declared that no competing interests exist.

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194 **REFERENCES**

195

196 1. Igbeneghu C, Olisekodiaka MJ, Akinola FFS, Odaibo AB. Impact of haemoglobin variants
197 AS and AC on Asymptomatic falciparum malaria among adults in Iwo, Southwestern Nigeria.
198 SJAMS. 2015a;3(1A):17-20.

199

200 2. Emechebe GO, Onyire NB, Orji ML, Achigbu KI. Sickle cell disease in Nigeria - A review.
201 IOSR-JDMS. 2017;16(1):87-94.

202

203 3. Mulumba LL, Wilson L. Sickle cell disease among children in Africa: An integrative
204 literature review and global recommendations. Intern J Afr Nursing Sci. 2015;3:56-64.

205

206 4. Azar S, Wong TE. Sickle cell disease: a brief update. Med Clin North Am. 2017;
207 101(2):375-393.

208

209 5. Macharia AW, Mochamah G, Uyoga S, Ndila CM, Nyutu G, Makale J, Tendwa M, Nyatichi
210 E, Ojal J, Shebe M, Awuondo KO, Mturi N, Peshu N, Tsofa B, Scott JAG, Maitland K and
211 Williams TN. The clinical epidemiology of sickle cell anaemia in Africa. Am J Haematol.
212 2018;93(3):363-370.

213

214 6. Aygun B, Odame I. A global perspective on Sickle cell disease. Pediatr Bld Cancer. 2012;
215 59(2): 386-390.

216

217 7. Jaff MS. Higher frequency of secretor phenotype in O blood group-its benefits in
218 prevention and /or treatment of some diseases. Intern J Nanomed. 2010;5:901-905.

219

220 8. Shinebaum R, Blackwell CC, Forster PJ, Hurst NP, Weir DM, Nuki G. Non-secretion of
221 ABO blood group antigens as host susceptibility factor in the spondyloarthropathies. BMJ
(Clin Res Ed). 1987;294(6566):208-210.

222

223 9. Shinebaum R. ABO blood group and secretor status in the spondyloarthropathies. FEMS
224 Microbiol Immunol. 1989;1(6-7):389-395.

225

226 10. Orstavik, KH, Kornstad L, Reisner H, Berg K. Possible effect of secretor locus on plasma
227 concentration of factor VIII and von Willebrand factor. Blood. 1989;73(4):990-993.

- 228
229 11. Orstavik KH. Genetics of plasma concentration of von Willebrand factor. *Folia Haematol*
230 *Int Mag Klin morphol Blutforsch.* 1990;117(4):527-531.
231
- 232 12. Al-Agidi SK, Shukri SM. Association between immunoglobulin levels and known genetic
233 markers in an Iraqi population. *Ann Hum Biol.* 1982;9(6): 565-569.
234
- 235 13. Hein HO, Sorensen H, Suadican P, Gyntelberg F. The Lewis blood group - a new
236 genetic marker of ischaemic heart disease. *J Intern med.* 1992;232(6):481-487.
237
- 238 14. Ellison RC, Zhang Y, Myers RH, Swanson JL, Higgins M, Eckfeldt J. Lewis blood group
239 phenotype as an independent risk factor for coronary heart disease (the NHLBI Family Heart
240 Study). *Am J Cardiol.* 1999;83(3):345-348.
241
- 242 15. Robinson WM, Salzano FM, Achutti AC, Franco MH. Blood groups, salivary secretion
243 and other immunologic variables in rheumatic fever and rheumatic heart disease. *Acta*
244 *Anthropogenet.* 1984;8(3-4):217-221.
245
- 246 16. Jhingam B, Mehra NK, Reddy KS, Taema V, Valdya MC, Bhatia ML. HLA, Blood groups
247 and secretor status in patients with established rheumatic fever and rheumatic heart
248 disease. *Tissue Ag.* 1986;3:172-178.
249
- 250 17. Dickey W, Collins JSA, Watson RGP, Sloan JM, Porter KG. Secretor status and
251 helicobacter pylori infection are independent risk factors for gastroduodenal disease. *Gut.*
252 1994;34(3):351-353.
253
- 254 18. Olorunshola KV, Audu I. ABO (H) secretor status of sickle cell disease patients in Zaria,
255 Kaduna State, Nigeria. *Niger J Physiol Sci.* 2013;28:29-34.
256
- 257 19. Igbeneghu C, Odaibo GN, Olaleye DO, Odaibo AB. Malaria Infection and ABO blood
258 grouping in Iwo community, Southwestern Nigeria. *Res J Med Sci.* 2012;6(5):247-250.
259
- 260 20. Igbeneghu C, Olisekodiaka JM, Alabi T, Onuegbu JA, Oseni BA, Odaibo A. ABH
261 secretors status in Osogbo, Southwestern Nigeria. *Indian J Fund Appl Life Sci.*
262 2015b;5(3):42-47.
263
- 264 21. Emeribe AO, Igweagu CA and Ossim EE. ABH secretor status in saliva of Calabar
265 Municipal residents. *East Afr Med J.* 1992;69(1):27-30.
266
- 267 22. Igbeneghu C, Olisekodiaka MJ, Okanlawon BM, Onuegbu JA, Odaibo AB. Non-
268 Secretors of ABH Antigens are susceptible to falciparum malaria. *SJAMS.*
269 2015c;3(5A):1838-1841.
270
- 271 23. Falusi AG, Ademowo CA, Latunji CA, Okeke AC, Olatunji PO, Onyekwere TO, Jimmy
272 EO; Raji Y. Distribution of ABO and Rh genes in Nigeria. *Afr J Med Med Sci.* 2000;29:23-26.
273
- 274 24. Bakare AA, Azeez MA, Agbolade JO. Gene frequencies of ABO and rhesus blood
275 groups and haemoglobin variants in Ogbomoso, Southwest Nigeria. *Afr J Biotech.*
276 2006;5(3):224-229.
277
- 278 25. Kapoor S, Little JA, Pecker LH. Advances in the treatment of Sickle cell disease. *Mayo*
279 *Clin Proc.* 2018;93(12):1810-1824.
280

281 26. Raza MW, Backwell CC, James VS, Ogilvie MM, Weir DM, Molyneaux P et al.
282 Association between secretor status and respiratory viral illness. BMJ. 1991;303:815-818.
283
284

UNDER PEER REVIEW