The effects of old and recent migration waves in the distribution of HBB*S globin gene haplotypes

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Abstract

Sickle cell hemoglobin is the result of a mutation at the sixth amino acid position of the beta (β) globin chain. The HBB*S gene is in linkage disequilibrium with five main haplotypes in the β-globin-like gene cluster named according to their ethnic and geographic origins: Bantu (CAR), Benin (BEN), Senegal (SEN), Cameroon (CAM) and Arabian-Indian (ARAB). These haplotypes demonstrated that the sickle cell mutation arose independently at least five times in human history. The distribution of βS haplotypes among Brazilian populations showed a predominance of the CAR haplotype. American populations were clustered in two groups defined by CAR or BEN haplotype frequencies. This scenario is compatible with historical records about the slave trade in the Americas. When all world populations where the sickle cell gene occurs were analyzed, three clusters were disclosed based on CAR, BEN or ARAB haplotype predominance. These patterns may change in the next decades due to recent migrations waves. Since these haplotypes show different clinical characteristics, these recent migrations events raise the necessity to develop optimized public health programs for sickle cell disease screening and management.

Keywords: βS globin haplotypes, sickle cell disease, Hemoglobin S, migration.

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Introduction

Sickle cell hemoglobin is the result of a single nucleotide mutation (GAG→GTG) at the sixth amino acid position of the beta (β) globin gene (HBB). Sickle cell anemia (SCA) is caused by HBB*S homozygosity. This gene has a worldwide distribution (Piel et al., 2010). The disease is a severe chronic hemolytic anemia, but its clinical course is highly variable. Although not completely understood, many factors have been suggested to be modulators of this variability, such as coinheritance with Hb C, α and β thalassemias, as well as high fetal hemoglobin (HB F) levels (Higgs et al., 1982; Frenette and Atweh, 2007).

The HBB*S gene is in linkage disequilibrium with five main haplotypes defined by single nucleotide polymorphisms (SNPs) in the β-globin-like gene cluster. These haplotypes are named according to their ethnic and geographic origins: Bantu (CAR, originated in South-Central and East Africa), Benin (BEN, in Midwest Africa), Senegal (SEN, in Atlantic West Africa), Cameroon (CAM, along the west coast of Africa), and Arabian-Indian (ARAB, from the Indian subcontinent and the eastern Arabian peninsula). Based on this haplotype distribution it has been demonstrated that the HBB*S mutation arose at least five times in human history (Pagnier et al., 1984; Kulozik et al., 1986; Lapouméroulie et al., 1992). Moreover these haplotypes have also been investigated in association with clinical features of the disease in order to disclose if some characteristics associated with disease severity such as HB F levels were also associated with a specific haplotype (Steinberg, 2009). It is essential to know about the old and recent dispersions of these haplotypes considering their clinical heterogeneities and their implications to public health programs for sickle cell disease screening and management.

HBB*S haplotypes have been studied in different Brazilian populations (Table 1), as tools to clarify population origins, since the sickle cell mutation is absent among Native Americans and it was introduced into the American continent basically by gene flow from Africa during the slave trade from the 16th to the 19th century (Zago et al., 1995; Salzano and Bortolini, 2002). In this study, we compared the HBB*S haplotypes frequencies in sickle cell disease patients from several world populations, in order to disclose the effects of old and recent wave migrations in the distribution of HBB*S haplotypes.
Material and Methods

A systematic review was performed to find studies that describe sickle cell haplotypes in different world populations. When more than one study from the same population was available, mean haplotype frequencies were calculated. A Wright’s FST (Weir and Hill, 2002) analysis was performed using ARLEQUIN 3.0 (Excoffier et al., 2005) to determine the differentiation among populations based on haplotype frequencies. Principal component analysis (PCA) was performed to summarize the distribution of populations based on the pairwise FST using SPSS v.18 software.

This study also included information about 110 non-consanguineous SCD patients from Rio Grande do Sul, southern region of Brazil, screened using isoelectric focusing (IEF) and/or cation exchange high performance liquid chromatography (HPLC) and confirmed by a PCR-RFLP approach with Ddel enzyme (Wagner et al., 2010). All patients were ascertained by the Neonatal Screening Reference Service or health care centers. The Ethics Committee of the Federal University of Rio Grande do Sul approved the study protocol.

Genomic DNA was isolated from peripheral blood samples using a salting out procedure (Lahiri and Nurnberger Jr, 1991). Haplotype analysis was performed by PCR-RFLP for the following polymorphic sites in the β globin gene cluster: HindIII-Gγ, HindIII-Aγ, HincII-ψβ, HincII, 3'ψβ, HinfI- S'β as previously described (Sutton et al., 1989). Haplotypes were inferred using the Multiple Locus Haplotype Analysis program (Long, 1999).

Results and Discussion

HBB*S haplotypes identified in several Brazilian populations are shown in Table 1. The CAR haplotype was the most frequent one, followed by the BEN haplotype. These results are in accordance with historical reports on slave traffic to Brazil. It is estimated that during the period between 1701 and 1816, 68% of the imported slaves came from Angola and the remainder from the Benin region. From 1843 to 1871, 90% of slaves came from Congo, Angola and Mozambique (Curtain, 1969). The SEN haplotype has its higher frequency in Brazil in Belém, in the northern region (Cardoso and Guerreiro, 2006). This is in accordance on what was expected based on the slave trade historical data of Atlantic West African populations to northern Brazil (10%), considering the high frequency of this haplotype in Senegal (Currat et al., 2002). The CAM haplotype was always in lower frequencies, with 0.9% in Rio Grande do Sul and 0.9-1.3% in other Brazilian regions, probably due to domestic slave trade and later internal migrations from regions supplied with slaves from Central West Africa, where this haplotype has been found (Oner et al., 1992). These results confirmed the diversity of the African influence in Brazilian regions.

PCA (Figure 1) demonstrated that two components explained 98.9% of the variance observed among Brazilians. The first component showed a group composed by Rio Grande do Sul (RS), Pará (PA), Pernambuco (PE), São Paulo (SP) and Rio Grande do Norte (RN) populations, where the CAR haplotype has a high frequency (from 66 to 81%). The other group was composed by Rio de Janeiro...
(RJ), Bahia (BA) and Ceará (CE) populations, where the CAR and BEN haplotypes have similar frequencies.

The Brazilian populations were then compared to other American populations. The PCA (Figure 2) showed the American populations distributed in different clusters. In this analysis, three groups explained 98.9% of the variance observed. Populations with higher frequencies of CAR are clustered together (Uruguay, Brazil, Panama and Mexico), whereas populations with higher BEN frequencies formed another cluster (USA, Canada, Trinidad, Guadeloupe and Jamaica). The other populations present similar BEN and CAR haplotype frequencies and formed a third cluster comprising Venezuela, Suriname, Colombia and Cuba. This cluster pattern appears to reflect geographical data, since a North, Central and South America separation can be observed, except for Mexico. This distribution could also be explained by historical reports of colonial power in these countries: Spain, France and Great Britain (Curtain, 1969). The British and French bought slaves from Midwestern African regions, where the BEN haplotype was prevalent, while slaves imported by the Spanish and Portuguese colonizers were mainly from Atlantic Central Africa, where CAR haplotype was the most prevalent.

Table 2 and the PCA of world populations (Figure 3) showed the distribution expected according to the haplotypes’ distribution and origin. Three different components could be observed with ARAB, CAR or BEN haplotype predominance. The first group was composed by Kuwait, Bahrain, Iran, India, United Arab Emirates and Senegal. All of them have a predominance of the Arabian-Indian (ARAB) haplotype, except Senegal. The second group was composed by Madagascar, Mexico, Angola, Tanzania, Kenya, Congo, Uganda, Brazil, Uruguay and Panama. All of them have a predominance of the Bantu (CAR) haplotype. The third group was composed by USA, Jordan, Tunisia, Guadeloupe, Canada, Jamaica, Suriname, Greece, Cameroon, Oman, Palestine, Algeria, Venezuela, Egypt, Syria, Cuba, Saudi Arabia, Turkey, Nigeria, Colombia, Sudan, Portugal and Italy. These populations have a predominance of the Benin (BEN) haplotype. The trade slave to the Americas and migration routes to the Mediterranean areas and the Middle East from West Africa determines the BEN haplotype predominance in these regions. Finally, the ARAB haplotype predominated in areas where it was originally derived.

This clear pattern of origin and dispersal of HBB*S haplotypes can suffer radical changes in the next decades due to global migrations. At present, the mobility of humans has reached unimaginable levels. This mobility can affect the epidemiology of several diseases, with an increase in the risk of a local disease spreading globally and the introduction of deleterious alleles into populations in
which they were previously absent. Information about the number of international migrants in the last decades showed a noticeable difference between migrants with and without HB S. Whereas the number of migrants without HB S increased from 92.6 million in 1960 to 165.2 million in 2000, the number of migrants with this hemoglobin increased faster (from 1.6 million in 1960 to 3.6 million in 2000) (Piel et al., 2014). The estimated number of migrants from African countries, India and Middle East with HB S moving to North America, Western Europe and Australia increased (Piel et al., 2014). An increase in the Arab-Indian haplotype frequency in several countries in the next decades could potentially be expected due migration processes that are occurring from the Middle East to Europe (Figure 4).

A similar process can also be observed in Brazil, where the number of migrants from Bolivia, Haiti and Senegal increased in the last years. The dispersal of these migrants is still uneven, but Bolivians tend to remain in São Paulo state while Senegalese individuals tend to move to Rio Grande do Sul (Figure 4). Therefore, an increase in the contribution of the Senegal haplotype is expected in southern Brazil, reflecting this new migration process. No studies about HBB*S haplotypes in Haiti population are available. This country does not have any national newborn screening program to measure the prevalence of SCD. Nevertheless, a study with infants born in Port-au-Prince showed that the prevalence of SCD in Haitian newborns appears to be more than twice higher than that found among African Americans in the United States (Rotz et al., 2013). This study showed a prevalence of the SCD genotypes Hb SS and HbSC of 1:173 newborns. The authors discuss the importance to consider these results carefully, since many children are born outside hospitals in Haiti, and therefore this prevalence may probably be an underestimate (Rotz et al., 2013). Since Haiti was colonized by French the most probable frequent haplotype would be BEN, as observed in Guadeloupe (Kéclard et al., 1997). Considering this information, independent from the HBB*S haplotype that predominates in these migrants, an increase in HB S prevalence in Brazil is expected in the next years. It is important to consider that the effect of migration cannot be assessed only by the number of migrants, but also by their behavior and habits. In this context, it is essential to consider that a higher intermarriage rate is likely among migrants from the same group, leading to an increase in sickle cell disease

Figure 2 - PCA based on FST distances calculated using haplotype frequencies showing clustering patterns for different American populations according to HBB*S haplotypes.
| Continents | Population | N     | CAR | BEN | SEN | CAM | ARAB | Atypical | Reference                  |
|------------|------------|-------|-----|-----|-----|-----|------|----------|----------------------------|
| Africa     | Algeria    | 20    | -   | 100.0 | -   | -   | -   | -       | Pagnier et al., 1984      |
|            | Angola     | 44    | 95.5 | 4.5  | -   | -   | -   | -       | Lavinha et al., 1992      |
|            | Cameroon   | 1082  | 0.5 | 73.8 | 0.2 | 19.1 | 0.3 | 6.1     | Bitoungui et al., 2015    |
|            | Congo      | 232   | 90.9 | 9.1   | -   | -   | -   | -       | Mouélé et al., 1999       |
|            | Egypt      | 28    | -   | 100.0 | -   | -   | -   | -       | El-Hazmi et al., 1999     |
|            | Guinea     | 40    | 22.5 | -    | -   | 77.5 | -   | -       | Sow et al., 1995          |
|            | Kenya      | 111   | 98.2 | 1.8   | -   | -   | -   | -       | Ojwang et al., 1987       |
|            | Mauritania | 90    | 4.4  | 8.9   | 77.8 | -   | -   | 5.6     | Veten et al., 2012        |
|            | Nigeria    | 669   | 0.9  | 93.3  | -   | 3.4 | -   | 2.4     | Adekile et al., 1992      |
|            | Senegal    | 90    | -    | -    | 100.0 | -   | -   | -       | Currat et al., 2002       |
|            | Sudan      | 143   | 2.8  | 29.4  | 18.2 | 35.0 | -14.7 | 5.7     | Hewitt et al., 1996       |
|            | Tanzania   | 41    | 100.0 | -   | -   | -   | -   | -       | Oner et al., 1992         |
|            | Tunisia    | 332   | 2.7  | 60.5  | -   | -   | -   | 36.7    | Moumni et al., 2011       |
|            | Uganda     | 208   | 99.5 | -    | 0.5 | -   | -   | -       | Mpalampa et al., 2012     |
| America    | Brazil     | 1176  | 65.0 | 31.5 | 3.0 | 0.5 | -   | -       | *                          |
|            | Canada     | 61    | 11.5 | 49.2  | 13.1 | 13.1 | -   | 13.1    | Oner et al., 1992         |
|            | Colombia   | 229   | 29.7 | 33.2  | 4.4 | 4.4 | 0.4 | 27.9    | Fong et al., 2013         |
|            | Cuba       | 198   | 40.9 | 51.0  | 8.1 | -   | -   | -       | Muniz et al., 1995        |
|            | Guadeloupe | 830   | 11.1 | 74.9  | 6.1 | 2.3 | 0.7 | 5.1     | Kéclard et al., 1997      |
|            | Jamaica    | 446   | 8.3  | 76.0  | 5.2 | -   | -   | 10.5    | Mpalampa et al., 2012     |
|            | Mexico     | 33    | 78.8 | 18.2  | -   | -   | -   | 3.0     | Magaña et al., 2002       |
|            | Panama     | 200   | 51.0 | 30.0  | 8.5 | 4.0 | 1.0 | 5.5     | Rusanova et al., 2011     |
|            | Surinam    | 77    | 29.9 | 53.2  | 2.6 | 2.6 | -   | 11.7    | Oner et al., 1992         |
|            | Trinidad   | 283   | 17.3 | 61.8  | 8.5 | 3.5 | 3.2 | 5.6     | Jones-Lecointe et al., 2008 |
|            | USA        | 806   | 16.0 | 62.4  | 9.4 | 4.7 | 1.5 | 6.0     | Crawford et al., 2002     |
|            | Uruguay    | 10    | 60.0 | 20.0  | -   | -   | -   | 20.0    | Luz et al., 2006          |
|            | Venezuela  | 359   | 36.4 | 51.5  | 10.6 | 1.5 | -   | -       | **                         |
| Asia       | Bahrain    | 37    | 5.4  | 2.7   | -   | -   | 89.2 | 2.7     | Al-Arrayed and Haltes, 1995 |
|            | India      | 140   | -    | -    | -   | -   | 91.4 | 8.6     | Mukherjee et al., 2004    |
|            | Iraq       | 128   | 7.8  | 69.5  | -   | -   | 12.5 | 10.2    | Al-Allawi et al., 2012    |
|            | Iran       | 162   | 3.1  | 11.7  | 3.7 | 2.5 | 53.7 | 25.3    | Rahimi et al., 2003       |
|            | Jordan     | 20    | -    | 80.0  | -   | -   | 20.0 | -       | El-Hazmi et al., 1999     |
|            | Kuwait     | 125   | 5.6  | 11.2  | -   | -   | 80.8 | 2.4     | Adekile and Haider, 1996  |
|            | Lebanon    | 100   | 15.0 | 73.0  | -   | -   | 10.0 | 2.0     | Inati et al., 2003        |
|            | Oman       | 117   | 21.4 | 52.1  | -   | -   | 26.5 | -       | Daar et al., 2000         |
|            | Palestine  | 118   | 5.1  | 88.1  | -   | -   | -   | 6.8     | Samarah et al., 2009      |
|            | Saudi-Arabia | 124 | -   | 98.4  | -   | -   | 1.6  | -       | El-Hazmi et al., 1999     |
|            | Syria      | 18    | -    | 66.7  | -   | -   | 33.3 | -       | El-Hazmi et al., 1999     |
|            | United Arab Emirates | 94 | 25.5 | 22.3 | -   | -   | 52.1 | -       | El-Kalla and Baysal, 1998 |
| Europe     | Greece     | 14    | -    | 92.9  | 7.1 | -   | -   | -       | Oner et al., 1992         |
|            | Italy      | 64    | -    | 100.0 | -   | -   | -   | -       | Schilirò et al., 1992     |
|            | Portugal   | 33    | 42.4 | 36.4  | 21.2 | -   | -   | -       | Lavinha et al., 1992      |
|            | Turkey     | 214   | -    | 96.3  | -   | -   | 0.5  | 3.3     | Oner et al., 1992         |

N: number of chromosomes; *mean frequency for Brazilian populations showed in Table 1; **mean frequency for Arends et al., 2000; Moreno et al., 2002.
prevalence. Some religious or cultural beliefs could be also a factor complicating an effective genetic counseling. The public health system agents should be prepared to address these problems in the best way possible.

Several chromosomes were identified as atypical (chromosomes with less common haplotypes) in all populations. Some of these atypical haplotypes were previously studied and diverse genetic mechanisms were inferred as
involved in their origin, such as recombination, point substitutions, or nonreciprocal sequence transfer (conversion) in the pre-existing common haplotypes instead of recurrent de novo HBB*S mutations (Zago et al., 2000). Subsequently, it was demonstrated that these events can be observed in typical HBB*S haplotypes in a way similar to those that generate atypical haplotypes (Zago et al., 2001). An extended haplotype within the HBB gene cluster is composed by three elements: a four repeats sequences configuration (AT)\(\times\)N12(AT)\(\gamma\) motif within the 5' HS2 region of \(\beta\)-LCR site, (TG)n (CG)n motif within IVSII region of fetal globin gene (\(\delta\)\(\gamma\) and \(\delta\)\(\gamma\)), and (AT)\(\times\)Ty motif within 5' region of \(\beta\)-globin gene region. Molecular investigations of this extended haplotype confirmed that the atypical haplotypes are obtained through recombination among the classical SNPs in the \(\beta\)-globin-like gene cluster and these sites in the extended haplotype region (Moumni et al., 2014).

In addition to population origin effects, these waves of migration can have important effects on public health. It was well established that there is a substantial phenotypic heterogeneity among patients with sickle cell anemia. In general, carriers of the CAR haplotype have the most severe clinical course, while carriers of the Senegal or Arab-Indian haplotypes have the best clinical course. Carriers of the BEN haplotype are intermediate in this respect. As HBB*S presence alone cannot explain this heterogeneity among patients, environmental influences and variations in others genes are likely to modulate the sickle cell anemia phenotype. The main pathophysiological factor determining disease severity is the Hb F concentration, leading to a reduced severity in patients with higher concentrations of this hemoglobin. In addition to Hb F concentration, \(\alpha\)-thalassemia can also affect the disease phenotype because both decrease Hb S polymerization. Several genetic and epigenetic factors modulate Hb F levels, such as the locus control region (LCR), the Hb F-related quantitative trait locus (QTL) and secretion-associated and RAS-related gene (SARI). In addition, several SNPs in candidate genes have been associated with subphenotypes of sickle cell anemia. For example, nonhemorrhagic stroke has been associated with variation in \(\text{VCAM1}, \text{TNFA}, \text{ADRB2}, \text{IL4R}, \text{LDLR}, \text{HLA}, \text{ANXA2}, \text{SELP}\) and \(\text{TGF-}\beta\text{/BMP}\) genes (a complete review about this topic could be found in Steinberg, 2009).

Considering the possible increase in Hb S frequency in Brazil due the recent wave migrations, it should be important to consider a more appropriate public health policy, including screening, adequate care and counseling, not only to Brazilians but also to migrants. Sometimes it could be difficult for migrants to have full access to public health services due to linguistic, cultural, religious, and social barriers but the government’s role is to provide the best opportunities to everyone.

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References

Adekile AD and Haider MZ (1996) Morbidity, \(\beta^\text{s}\) haplotype and \(\alpha\)-globin gene patterns among sickle cell anemia patients in Kuwait. Acta Haematol 96:150-154.

Adekile AD, Kitundu MN, Gu LH, Lancelos KD, Adeodu OO and Huisman TH (1992) Haplotypes in SS patients from Nigeria; characterization of one atypical beta S haplotype no. 19 (Benin) associated with elevated HB F and high G gamma levels. Ann Hematol 65:41-45.

Adorno EV, Zanette A, Lyra I, Seixas MO, Reis MG and Gonçalves MS (2008) Clinical and molecular characteristics of sickle cell anemia in northeast of Brazil. Genet Mol Biol 31:621-625.

Al-Allawi NA, Jalal SD, Nerwey FF, Al-Sayan GO, Al-Zebari SS, Alshingaly AA, Markous RD, Jubrael JM and Hamamy H (2012) Sickle cell disease in the Kurdistan population of northern Iraq. Hemoglobin 36:333-342.

Al-Arrayed SS and Haltes N (1995) Features of sickle-cell disease in Bahrain. East Mediterr Health J 1:112-118.

Arends A, Alvarez M, Velázquez D, Bravo M, Salazar R, Guevara JM and Castillo O (2000) Determination of beta-globin gene cluster haplotypes and prevalence of alpha-thalassemia in sickle cell anemia patients in Venezuela. Am J Hematol 64:87-90.

Bezerra MAC, Santos MNN, Araújo AS, Gomes YM, Abath FG and Bandeira MGC (2007) Molecular variations linked to the grouping of \(\beta\) and \(\alpha\)-globin genes in neonatal patients with sickle cell disease in the state of Pernambuco, Brazil. Hemoglobin 31:1-6.

Bitoungui VJN, Pule GD, Hanchard N, Ngogang J and Wonkam A (2015) Beta-globin gene haplotypes among Cameroonian and review of the global distribution: Is there a case for a single sickle mutation origin in Africa? OMICS 19:171-179.

Cabral CHK, Serafim ESS, de Medeiros WRDB, Fernandes TAAM, Kimura EM, Costa FF, Sonati MF, Rebecchi IMM and de Medeiros TMD (2011) Determination of \(\beta^\text{s}\) haplotypes in patients with sickle-cell anemia in the state of Rio Grande do Norte, Brazil. Genet Mol Biol 34:421-424.

Cardoso GL and Guerreiro JF (2006) African gene flow to North Brazil as revealed by HBB*S gene haplotype analysis. Am J Hum Biol 18:93-98.

Costa FF, Arruda VR, Gonçalves MS, Miranda SRP, Carvalho MH, Sonati MF, Saad SOT, Gesteira F, Fernandes D, Nascimento ML, et al. (1984) \(\beta^\text{s}\)-gene cluster haplotypes in sickle cell anemia patients from two regions of Brazil. Am J Hematol 46:96-97.

Crawford DC, Caggana M, Harris KB, Lorey F, Nash C, Pass KA, Tempelis C and Olney RS (2002) Characterization of \(\beta\)-globin haplotypes using blood spots from a population-based cohort of newborns with homozygous HbS. Genet Med 4:328-335.

Currat M, Trabuchet G, Rees D, Perrin P, Harding RM, Clegg JB, Langaney A and Excoffier L (2002) Molecular analysis of the \(\beta\)-globin gene cluster in the Niokholo Mandenka popula-
tion reveals a recent origin of the \(\beta\)-S Senegal mutation. Am J Hum Genet 70:207-223.

Curtain PD (1969) The Atlantic Slave Trade: A census. University of Wisconsin Press, Milwaukee, 338 p.

Daar S, Hussain HM, Gravel D, Nagel RL and Krishnamoorthy R (2000) Genetic epidemiology of \(\alpha\)S in Oman: Multicentric origin for the \(\beta\)S gene. Am J Hum Genet 64:39-46.

Elderbery AY, Mills J, Mohamed BA, Cooper AJ, Mohammed AO, Eltieb N and Old J (2012) Molecular analysis of the \(\beta\)-globin gene cluster haplotypes in a Sudanese population with sickle cell anemia. Int J Lab Hematol 34:262-266.

El-Hazmi MA, Warsy AS, Bashir N, Beshlawi A, Hussain IR, Tantamy S and Qubaili F (1999) Haplotypes of the \(\beta\)-globin gene as prognostic factors in sickle-cell disease. East Mediterr Health J 5:1154-1158.

El-Kalla S and Baysal E (1998) Genotype-phenotype correlation in sickle cell disease in the United Arab Emirates. J Pediatr Hematol Oncol 15:237-242.

Excoffier L, Laval G and Schneider S (2005) Arlequin (version 3.5): An integrated software package for population genetics data analysis. Evol Bioinform Online 1:47-50.

Fleury MK (2007) Haplotipos do cluster da globina beta em pacientes com anemia falciforme no Rio de Janeiro: Aspectos clínicos e laboratoriais. Rev Bras Anál Clín 39:89-93.

Fong C, Lizaralde-Iragorri MA, Rojas-Gallardo D and Barreto G (2013) Frequency and origin of haplotypes associated with the \(\beta\)-globin gene cluster in individuals with trait and sickle cell anemia in the Atlantic and Pacific coastal regions of Colombia. Genet Mol Biol 36:494-497.

Frenette PS and Atweh GF (2007) Sickle cell disease: Old discoveries, new concepts, and future promise. J Clin Invest 117:850-858.

Galiza Neto GC, Pitombeira MS, Vieira HF, Vieira MLC and Farias DAB (2005) Analysis of \(\beta\)-globin gene haplotypes in Ceará, Brazil. J Bras Patol Med Lab 41:315-321.

Gonçalves MS, Nechtmann JF, Figueiredo MS, Kerbauy J, Arruda VR, Sonati MF, Saad SOT, Costa FF and Stoming TA (1994) Sickle cell disease in a Brazilian population from São Paulo: A study of the \(\beta\)S haplotypes. Hum Hered 44:322-327.

Gonçalves MS, Bomfim GC, Maciel E, Cerqueira I, Lyra I, Zanette A, Bomfim G, Adorno EV, Albuquerque AL, Pontes A, et al. (2003) \(\beta\)-haplotypes in sickle cell anemia patients from Salvador, Bahia, Northeastern Brazil. Braz J Med Biol Res 36:1283-1288.

Hewitt R, Krause A, Goldman A, Campbell G and Jenkins T (1996) Beta-globin haplotype analysis suggests that a major source of Malagasy ancestry is derived from Bantu-speaking populations. J Hum Genet 58:1303.

Higgs DR, Aldridge BE, Lamb J, Clegg JB, Weatherall DJ, Hayes RJ, Grandison Y, Lowrie Y, Mason KP, Serjeant GE, et al. (1982) The interaction of alpha-thalassemia and homozygous sickle-cell disease. N Engl J Med 306:1441-1446.

Inati A, Taher A, Bou Alawi W, Koussa S, Kaspar H, Shbaklo H and Zalloua PA (2003) Beta-globin gene cluster haplotypes and \(\text{Hb}F\) levels are not the only modulators of sickle cell disease in Lebanon. Eur J Haematol 70:79-83.

Jones-Lecointe A, Smith E, Romana M, Gilbert MG, Charles WP, Saint-Martin C and Kéclard L (2008) Beta-globin gene cluster haplotypes and \(\alpha\)-thalassemia in sickle cell disease patients from Trinidad. Am J Hum Biol 20:342-344.

Kéclard L, Romana M, Lavocat E, Saint-Martin C, Berchel C and Mérault G (1997) Sickle cell disorder, beta-globin gene cluster haplotypes and \(\alpha\)-thalassemia in neonates and adults from Guadeloupe. Am J Hematol 55:24-27.

Kulozok AE, Wainscoat JS, Serjeant GR, Kar BC, Al-Awamy B, Essan GJ, Falusi AG, Haque SK, Hilali AM, Kate S, et al. (1986) Geographical survey of beta S-globin gene haplotypes: Evidence for an independent Asian origin of the sickle-cell mutation. Am J Hum Genet 39:239-244.

Lahiri DK and Nurnberger Jr JI (1991) A rapid non-enzymatic method for the preparation of HMW DNA from blood for RFLP studies. Nucleic Acids Res 19:5444.

Lapouméroulie C, Dunda O, Ducrocq R, Trabuchet G, Mony-Lobé M, Bodo JM, Carnevale P, Labie D, Elion J and Krishnamoorthy R (1992) A novel sickle cell mutation of yet another origin in Africa: The Cameroon type. Hum Genet 89:333-337.

Lavinha J, Gonçalves J, Faustino P, Romião L, Osório-Almeida L, Peres MJ, Picanço I, Martins MC, Ducrocq R, Labie D, et al. (1992) Importation route of the sickle cell trait into Portugal; Contribution of molecular epidemiology. Hum Biol 64:891-901.

Long JC (1999) Multiple Locus Haplotype Analysis. Software and documentation distributed by the author. Laboratory of Neurogenetics, National Institute on Alcohol Abuse and Alcoholism, National Institutes of Health, Bethesda.

Luz JÁ, Sans M, Kimura EM, Albuquerque DM, Sonati MF and Costa FF (2006) \(\alpha\)-thalassemia, \(\text{Hb}S\) and beta-globin gene cluster haplotypes in two Afro-Uruguayan sub-populations from northern and southern Uruguay. Genet Mol Biol 29:595-600.

Magaña MT, Onguy Z, Tagle J, Bentura G, Cobian JG, Perea FJ, Casas-Castañeda M, Sánchez-López YJ and Ibarra B (2002) Analysis of beta S and beta A genes in a Mexican population with African roots. Blood Cells Mol Dis 28:121-126.

Moreno N, Martínez JA, Blanco Z, Osorio L and Hackshaw P (2002) Beta-globin gene cluster haplotypes in Venezuela sickle cell patients from the state of Aragua. Genet Mol Biol 25:21-24.

Moulé R, Pambou O, Feingold J and Galactéros F (1999) \(\alpha\)-thalassemia in Bantu population from Congo-Brazzaville: Its interaction with sickle cell anemia. Hum Genet 105:118-125.

Moumni I, Ben Mustapha M, Sassi S, Zorai A, Ben Mansour I, Douzi K, Chouachi D, Melloulfi F, Bejaoui M and Abbas S (2014) Haplotype map of sickle cell anemia in Tunisia. Dis Markers 2014:938301.

Moumni I, Ilkbel BMM, Leila C, Fethi M, Amine Z, Mohamed B and Salem A (2011) Restriction mapping of \(\beta^+\) locus among Tunisian sickle cell patients. Am J Hum Biol 23:815-819.

Mpampe L, Nduguwa CM, Ddungu H and Idrro R (2012) Foetal haemoglobin and disease severity in sickle cell anaemia patients in Kampala, Uganda. BMC Hematol 12:e11.

Mukherjee MB, Surve RR, Gangakhedkar RR, Ghosh K, Colah RB and Mohanty D (2004) Beta-globin gene cluster haplotypes linked to the beta S gene in western India. Hemoglobin 28:157-161.

Muniz A, Corral L, Alaez C, Svrach E, Espinosa E, Carbonell N, di Leo R, Felicetti L, Nagel RL and Martinez G (1995) Sickle cell anemia and beta-gene cluster haplotypes in Cuba. Am J Hematol 49:163-164.
Ojwang PJ, Ogada T, Beris P, Hattori Y, Lanclos KD, Kutlar A, Kutlar F and Huismann TH (1987) Haplotypes and a globin gene analyses in sickle cell anemia patients from Kenya. Br J Haematol 65:211-215.

Oner C, Dimovski AJ, Olivieri NF, Schiliro G, Codrington JF, Fattoum S, Adekile AD, Oner R, Yüregir GT, Altay C, et al. (1992) Beta S haplotypes in various world populations. Hum Genet 89:99-104.

Pagnier J, Mears JG, Dunda-Belkhodja O, Schaefer-Rego KE, Beldjord C, Nagel RL and Labie D (1984) Evidence for the multicentric origin of the sickle cell hemoglobin gene in Africa. Proc Natl Acad Sci USA 81:1771-1773.

Pante-de-Sousa G, Mousinho-Ribeiro RC, dos Santos EJM, Zago MA and Guerreiro JF (1998) Origin of the hemoglobin S gene in a northern Brazilian population: The combined effects of slave trade and internal migrations. Genet Mol Biol 21:365-373.

Piel FB, Patil AP, Howes RE, Nyangiri OA, Gething PW, Williams TN, Weatherall DJ and Hay SI (2010) Global distribution of the sickle cell gene and geographical confirmation of the malaria hypothesis. Nat Commun 1:e104.

Piel FB, Tatem AJ, Huang Z, Gupta S, Williams TN and Weatherall DJ (2014) Global migration and the changing distribution of sickle haemoglobin: A quantitative study of temporal trends between 1960 and 2000. Lancet Glob Health 2:e80-e89.

Rahimi Z, Karimi M, Haghshenass M and Merat A (2003) Beta-globin gene cluster haplotypes in sickle cell patients from southwest Iran. Am J Hematol 74:156-160.

Rotz S, Arty G, Dall’Amico R, de Zen L, Zanoli F and Bodas P (2013) Prevalence of sickle cell disease, hemoglobin S, and hemoglobin C among Haitian newborns. Am J Hematol 88:827-828.

Rusanova I, Cossio G, Moreno B, Javier Perea F, De Borace RG, Perea M, Escames G and Acuña-Castroviejo D (2011) β-globin gene cluster haplotypes in sickle cell patients from Panama. Am J Hum Biol 23:377-380.

Salzano FM and Bortolini MC (2002) The evolution and genetics of Latin American populations. Cambridge University Press, Cambridge, 512 p.

Samarah F, Ayesh S, Athanasiou M, Christakis J and Vavatsi N (2009) BetaS-globin gene cluster haplotypes in the West Bank of Palestine. Hemoglobin 33:143-149.

Schiliro G, Samperi P, Consalvo C, Gangarossa S, Testa R, Miraglia V and Lo Negro L (1992) Clinical, hematological, and molecular features in Sicilians with sickle cell disease. Hemoglobin 16:469-480.

Steinberg MH (2009) Genetic etiologies for phenotypic diversity in sickle cell anemia. Sci World J 9:46-67.

Silva LB, Gonçalves RP and Rabenhorst SHB (2009) Analysis of sickle cell anemia haplotypes in Fortaleza reveals the ethnic origins of Ceará state population. J Bras Patol Med Lab 45:115-118.

Sow A, Peterson E, Josifovska O, Fabry ME, Krishnamoorthy R and Nagel RL (1995) Linkage disequilibrium of the Senegal haplotype with the β⁶ gene in the Republic of Guinea. Am J Hematol 50:301-303.

Sutton M, Bouhassira EE and Nagel RL (1989) Polymerase chain reaction amplification applied to the determination of beta-like globin gene cluster haplotypes. Am J Hematol 32:66-69.

Veten FM, Abdelhamid IO, Meiloud GM, Ghaber SM, Salem ML, Abbes S and Houmeida AO (2012) Hb S [β⁶ (A3) Glu/Val, GAG > GTG] and β-globin gene cluster haplotype distribution in Mauritania. Hemoglobin 36:311-315.

Wagner SC, de Castro SM, Gonzalez TP, Santin AP, Zaleski CF, Azevedo LA, Dreau H, Henderson S, Old J and Hutz MH (2010) Neonatal screening for hemoglobinopathies: Results of a public health system in South Brazil. Genet Test Mol Biomarkers 14:565-569.

Weir BS and Hill WG (2002) Estimating F-statistics. Annu Rev Genet 36:721-750.

Zago MA, Figueiredo MS and Ogo SH (1992) Bantu βS cluster haplotype predominates among Brazilian blacks. Am J Phys Anthropol 88:295-298.

Zago MA, Melo Santos EJ, Clegg JB, Guerreiro JF, Martinson JJ, Norwich J and Figueiredo MS (1995) Alpha-globin gene haplotypes in South American Indians. Hum Biol 67:535-546.

Zago MA, Silva Jr WA, Dalle B, Gualandro S, Hutz MH, Lapoumeroulie C, Tavella MH, Araujo AG, Krieger JE, Elion J, et al. (2000) Atypical beta(s) haplotypes are generated by diverse genetic mechanisms. Am J Hematol 63:79-84.

Zago MA, Silva WA Jr, Gualandro S, Yokomizu IK, Araujo AG, Tavela MH, Gerard N, Krishnamoorthy R and Elion J (2001) Rearrangements of the beta-globin gene cluster in apparently typical beta S haplotypes. Haematologica 86:142-145.

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