Potential Protective Effect of Sickle Cell Gene Allele on HIV Infection

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Abstract

Both Sickle Cell Disease and HIV Infection have overlapping worldwide distributions, with Africa being the epicenter for these two conditions [Figure 1 & 2] [1,2]. Sickle Cell Disease is a genetic condition affecting chromosome 11 in which homozygous recessive individuals will have significant negative consequences secondary to sickling phenomenon of their red blood cells, while those that are heterozygous have a survival advantage based on a well-described resistance to malaria [3]. The Human Immunodeficiency Virus takes advantage of surface proteins on white cells, resulting in lytic destruction and consequent CD4+ lymphopenia and immunosuppression [4]. Both diseases have significant morbidity and mortality, with limited treatment options. Nevertheless, observations in our pediatric HIV program in Newark, NJ, one of the first epicenters for Perinatal HIV Infection (PHIV), suggested that children with sickle cell disease infrequently were infected with HIV despite being from a high-risk population5 (Table 1) [5]. Current understanding of the interplay of genetic impact on surface membranes supports the idea that genetic variations might impact susceptibility to infectious diseases. Similar to its protective effect against malaria, sickle cell trait may provide protection against acquisition of HIV infection through changes in CXCR4 and CCR5 receptors on CD4+ lymphocytes, possibly blocking viral entry and infection [6]. Potential mechanisms for this disparity might provide better understanding and approaches to HIV infection based on sickle cell status.

Table 1: HIV Status by Hemoglobinopathy Status among Perinatal HIV-Exposed Infants (Newark, NJ 1993-2003) [5]. *P=0.48 by Fisher’s Exact (odds ratio=0.62; 95% Confidence interval=0.21 to 1.8).

<table>
<thead>
<tr>
<th>HIV/Hemoglobin Status</th>
<th>HIV+</th>
<th>HIV-</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hemoglobinopathy (AS)</td>
<td>4</td>
<td>44</td>
<td>48</td>
</tr>
<tr>
<td>No Hemoglobinopathy (AA)</td>
<td>40</td>
<td>236</td>
<td>276</td>
</tr>
<tr>
<td>Total</td>
<td>44</td>
<td>280</td>
<td>324</td>
</tr>
</tbody>
</table>

The authors have had a long standing hypothesis that the Sickle Cell Gene Allele (SCGA) may confer a protective effect against acquisition and/or progression of HIV infection. Rutgers NJMS in Newark, NJ has long been a center of PHIV care and research since the onset of the HIV pandemic, developing domestic and global clinical, educational and research expertise in PHIV infection.
leading to the control and subsequent prevention of PHIV in the US. Among our large population of PHIV African American children, we noted a lower than expected occurrence of Sickle Cell Hemoglobinopathy (SCH) suggesting relative protection against PHIV acquisition, and by inference, progression. However, though the data from the Newark PHIV program is suggestive of the potential protective effect of the SCGA, it lacks the patient base needed to achieve sufficient statistical power to confirm this potential association [5]. Sub-Saharan Africa has approximately 64% of the world’s population living with HIV, 91% of children under 15 years of age living with HIV, and approximately 10-40% of the world’s SCD population [1,7]. The population distribution of individuals in Africa with Hbs allele and HIV are seemingly non-overlapping, supporting preliminary data from our pediatric cohort [1,2,5]. This non-overlapping distribution is unexpected, given the high prevalence of both Sickle Cell Disease and HIV in Africa. South Africa, having a low distribution of SCGA has one of the higher concentrations of HIV infectivity in Africa, confirming the significance of these observations. These findings should stimulate further investigation into the molecular basis for this protective effect, which could influence the direction of new therapies (drug, genetic, and immune-based approaches) for the treatment and prevention of HIV.

There are three lines of evidence that suggest Sickle Cell Hemoglobin (SCH) has a protective effect against HIV acquisition/progression. First, anecdotal observations by investigators at Rutgers NJMS noted the above, which suggested lower transmission rates of PHIV among HIV exposed neonates with SCGA compared with those without [5]. Second, an epidemiological study by Nouria et al. [8] documented a two-fold lower risk of HIV infection and comorbidity in patients with SCD among African-American adult patient hospital discharges. This effect was observed over a 13-year period in over 400,000 patient records. Further, the authors of this article offered several possible mechanisms for the apparent impact of SCGA on the course of HIV disease including up-regulation of heme oxygenase-1, hypoxia related to the anemia, higher expression of inflammatory cytokines, inhibition of HIV transcription in the presence of reduced iron, as well as use of hydroxyurea in SCD patients. Third, SCGA is already documented to have protective effects against other infectious diseases, such as Malaria. Although this conferred protection occurs by conformational changes in the hemoglobin molecule by the SCGA inhibiting the penetration and proliferation of the Malaria Parasite in RBCs, there may be other characteristics of this gene allele product, expressed on lymphocytes and other white blood cells that may also confer protection against HIV attachment, penetration and/or proliferation [9].

A possible mechanism of protection could exist within the genome of patients with SCGA. Past literature has already elucidated the mechanism by which certain Caucasians of Northern European descent have conferred resistance to HIV [10]. These persons possess a 32-base-pair deletion on the CCR5 white blood cell surface protein, one of two chemoreceptors utilized by HIV-1 to enter the leukocyte. The CCR5Δ32 mutation found in these individuals results in a dysfunctional surface protein, preventing the virus from entering the host [11]. In the case of individuals with SCGA, the authors further propose that the TRIM5α protein could play a role in the protective effect. This protein disassembles and degrades viral capsid binding, preventing the pre-integration complex from arriving at the nucleus, thus interfering with reverse transcription [12]. A study in 2004 conducted by Stremlau et al. [13] identified TRIM5α as the blocking factor responsible for blocking HIV-1 strains from infecting Old World Monkeys13. Subsequent research has shown that rhesus TRIM5α (TRIM5α<sub>hu</sub>) appears to be more stable, and thus more potent, than human TRIM5α (TRIM5α<sub>sa</sub>), lending a reason as to why the majority of humans are immune to HIV-1 infection. Examination into the effect of various polymorphisms in TRIM5α<sub>sa</sub> revealed differing frequencies of HIV infection among African American cohorts. The TRIM5α 136Q polymorphism—indicating a glutamate at residue 136 in the TRIM5α protein – was identified as possessing a potential protective effect, as it was found at a greater frequency in the High-Risk Exposed Uninfected African American population [14]. Furthermore, the TRIM5α gene locus is found in close proximity to the HBB gene locus on human chromosome 11, lending the possibility that linkage disequilibrium interaction between the two could exist. Because a large proportion of African American individuals possess the SCGA, the authors further propose that a potential link between SCGA and TRIM5α 136Q may explain the lower-than-expected proportion of HIV-infected individuals with the SCGA.

While a recent systematic review concluded that SCD slows the progression of HIV into AIDS, and conversely HIV infection complicates the course of patients with SCD [15], no well-controlled study of a relationship between the presence of SCGA and the acquisition or progression of HIV infection has been conducted. Such a study would identify the sickle cell status (SS vs. SC vs. SA vs. AA) in HIV-infected cohorts and, conversely, the HIV status in identified sickle cell cohorts. The population of HIV-infected patients who carry the SCGA would be the target population of a prospective study to determine the impact associated with the co-morbidity of HIV infection and SCGA status. The high prevalence of SCGA and HIV infection in the Sub-Saharan African population and the preliminary observations of a possible impact of SCGA status and HIV incidence as well as the progression of HIV-1 disease, justifies further study of the possible interaction between these two significant global health problems. The results of such a study would have a major impact on the management and follow-up of individuals followed in these countries as well as encourage collaboration with and within these resource-constrained areas of the world to address the global health challenges experienced by much of the world’s population.

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Author Contributions

JO made original observation that SCD appeared in his population of PHIV infants to be underrepresented. HS assisted in recruiting international support for this hypothesis. JM consulted on presence of PHIV in her large cohort of SCD patients. AS provided technical support in data evaluation. KS, GG, JK, RM, PP, AD, OF, and FD provided technical support in data analysis and manuscript preparation.

References


