A new insight into apheresis platelet donation by sickle cell trait carriers: evidences of safety and quality

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Introduction

In countries with a high prevalence of the sickle cell trait (SCT), which is often determined by neonatal screening programs, a significant proportion of blood donors may be SCT carriers [1]. In Brazil, for example, where SCT prevalence ranges from 1.1% to 9.8% in the overall population [2], the trait is found in up to 2.48% of blood donors [3-7]. Because individuals with SCT are usually asymptomatic, many of them are unaware of their condition at the time of donation [1]. Considering the potential risks associated with SCT blood donation for both donor and recipient, particularly those with sickle cell disease [8], routine HbS screening in first-time blood donations is generally recommended [9].

After the implementation of universal leukoreduction in several countries, the management and use of SCT blood products became a challenge for blood banks. The trait represents the most common cause of filter failure [10], and the units that manage to pass through the filter may have prolonged filtration time and high residual white blood cell (WBC) counts, despite the use of high performance filters [11]. These results appear to be a consequence of mechanical differences in SCT red blood cells (RBCs), which are inherently stiffer and more viscous than those from healthy donors [12].

As a result, some countries have issued specific recommendations regarding the processing and use of SCT RBCs products. In Brazil, RBCs products with the SCT cannot undergo leukoreduction and their use is prohibited in some clinical scenarios, such as patients with hemoglobinopathies, severe acidosis or hyperthermia [13]. The American Association of Blood Banks (AABB) [14] and the European Committee on Blood Transfusion (ECBT) [15] have similar recommendations. Therefore, SCT donors should preferentially donate non-RBC products [16]. In this context, platelethropheresis may be a promising alternative to maximize SCT donors. However, there is a blank regarding safety of such donations. For instance, ECBT prohibits SCT carriers to donate RBC products by apheresis, but there is no mention to apheresis platelet donation [15], which is often allowed or prohibited based on Blood Banks’ experience only.

Current standards have defined cut-off values of the main parameters used in the assessment of in vitro apheresis platelet's quality. Actual platelet yield, pH and residual WBC count are specified in virtually all major standards for apheresis platelets quality, but with some slight differences in their threshold values. The Brazilian Ministry of Health’s (BMH) technical regulation for hemotherapy procedures and the AABB Standards for Blood Banks and Transfusion Services agree in recommendations for two of these parameters: a minimum platelet yield of $3 \times 10^{11}$ platelets in 90% of sampled units and a residual WBC count below $5 \times 10^6$ per unit [13,14]. On the other hand, the European Committee on Blood Transfusion is less demanding, requiring a platelet yield of at least $2 \times 10^{11}$ per unit and a residual leukocyte count below $3 \times 10^6$ per unit [15]. The AABB indicates that at least 90% of apheresis platelets should have a pH ≥ 6.2 at the end of the storage time [14], while the European and the Brazilian regulations specify a pH greater than 6.4 at the end of shelf-life, with additional recommendation for volume and, in the Brazilian requirements, a negative culture result [13,15].

This study sought to assess the frequency of apheresis platelet donation by SCT carriers, the safety of the procedure to this population of donors and the in vitro quality of their apheresis platelet concentrates.

Methods

This is a retrospective study which included all apheresis platelet donations by SCT carriers between January 2013 and December 2016 at Hematology and Hemotherapy Center of Ceará, Brazil. Apheresis platelet units were obtained using Trima Accel® Automated Blood Collection System (Terumo BCT; Denver, USA). Hemoglobin status was screened by electrophoresis and the result was confirmed by high performance liquid chromatography. All data were obtained using the blood bank software.

Information on in vitro quality parameters of platelethpheresis was only available for the years of 2015 and 2016. AABB, ECBT and the BMH recommendations were used as standards for apheresis platelets quality. Parameters of SCT donors were compared to those of a control group composed of non-SCT carriers matched for sex who donated in the same period. For each SCT donor, two non-SCT donors were included in the control group. There was one exception: only one female non-SCT donor was included to pair with one of the female SCT donors in that period, since no other female match was available, because of the transfusión-related acute lung injury (TRALI) prevention protocol.

The data were analyzed using SPSS 20.0 for Windows (SPSS Inc. Chicago, IL, USA) and expressed as mean ± standard deviation (SD). The Kolgomorov-Smirnov test was used to sort out variables which

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follow a Gaussian curve from those which do not. In order to compare variable means between groups, Student’s t or Mann-Whitney tests were performed accordingly.

Results

A total of 2,434 platelet donors were included, accounting for 5930 donations. Hemoglobin electrophoresis was performed in 1921 donors (78.93%). Of those, 27 (1.4%) had the SCT, 1885 were homozygous for HbA (98.1%) and 9 were heterozygous for HbC (0.46%). The only variables that followed a normal distribution were concentrates’ platelet yield and donors’ pre-donation platelet count.

The overall mean number of donations per individual was 2.44 ± 4.38, with median of 1 and maximum of 83 donations, as shown in Table 1. A total of 1,431 donors (62.9%) donated only once in the period of the study.

The SCT group was responsible for 104 donations (1.75%), with a mean, median and maximum number of donations per individual of 3.85 ± 6.86, 2 and 35 respectively. There was no statistically significant difference between the SCT and non-SCT donor’s number of donations (3.85 ± 6.86 vs. 2.44 ± 4.38, p = 0.096). One-time donors accounted for 48.1% (13 individuals) of SCT participants. None of the SCT donors presented any side-effect or complication that may have arisen due to apheresis donation, whilst 24 HbAA donors presented 25 episodes of adverse reactions to the donation, mainly hypocalcemic symptoms.

Platelet quality parameters were available for 13 SCT donors, eleven of which were males (86.61%). Mean age in this group was 33.69 ± 12.47 years. Twenty-five donors were included in the control group, with 22 males (88%) and mean age of 37.84 ± 11.50 years. There was no statistically significant difference in age (37.84 ± 11.50 vs. 33.69 ± 12.47 years, p = 0.312) or pre-donation platelet count (254.84 ± 56.17 vs. 254.31 ± 66.11 thousand platelets/mm³, p = 0.979) between the two groups, as showed in Table 2.

Table 3 displays the cut-off values recommended by the ECBT, AABB and the BMH followed by a comparison between in vitro parameters for quality assessment of the apheresis platelet units collected from the SCT group and the control group.

Although SCT causes no lifespan shortening or clinical repercussions in the majority of cases, some specific situations, such

Table 1. Comparison of number of donations and episodes of adverse reaction in all participants and in the SCT group

<table>
<thead>
<tr>
<th></th>
<th>Number of donations</th>
<th>Adverse Reactions</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean</td>
<td>Median</td>
</tr>
<tr>
<td>All donors (n = 2434)</td>
<td>2.44 ± 4.38</td>
<td>1</td>
</tr>
<tr>
<td>SCT donors (n = 27)</td>
<td>3.85 ± 6.86</td>
<td>2</td>
</tr>
<tr>
<td>Non-SCT donors (n = 2407)</td>
<td>2.42 ± 4.34</td>
<td>1</td>
</tr>
<tr>
<td>p-value</td>
<td>0.096*</td>
<td>---</td>
</tr>
</tbody>
</table>

*Mann-Whitney test. P-values ≤ 0.05 were considered statistically significant.

Table 2. Comparison of donor’s features between the SCT and the control group

<table>
<thead>
<tr>
<th></th>
<th>Control group (n=26)</th>
<th>SCT group (n=13)</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pre-donation platelet count (10³/mm³)</td>
<td>254.84 ± 56.17</td>
<td>254.31 ± 66.11</td>
<td>0.979*</td>
</tr>
<tr>
<td>Age (years)</td>
<td>37.84 ± 11.50</td>
<td>33.69 ± 12.47</td>
<td>0.312*</td>
</tr>
</tbody>
</table>

*Student’s t test. P-values ≤ 0.05 were considered statistically significant.

Table 3. Cut-off values of quality parameters recommended by ECBT, AABB and BMH and comparison between mean values of quality control parameters of SCT and control groups

<table>
<thead>
<tr>
<th></th>
<th>ECBT’s recommendation</th>
<th>AABB’s recommendation</th>
<th>BMH’s recommendation</th>
<th>Mean values</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Control group (n=26)</td>
<td>SCT group (n=13)</td>
<td>p-value</td>
<td></td>
</tr>
<tr>
<td>Platelet yield</td>
<td>≥ 2</td>
<td>≥ 3</td>
<td>≥ 3</td>
<td>5.03 ± 1.52</td>
</tr>
<tr>
<td>Residual WBC count</td>
<td>&lt; 300</td>
<td>&lt; 5</td>
<td>&lt; 5</td>
<td>0.07 ± 0.02</td>
</tr>
<tr>
<td>pH</td>
<td>&gt; 6.4</td>
<td>&gt; 6.2</td>
<td>&gt; 6.4</td>
<td>7.63 ± 0.26</td>
</tr>
<tr>
<td>Volume</td>
<td>&gt; 40³</td>
<td>---</td>
<td>&gt;200³</td>
<td>339.76 ± 98.71</td>
</tr>
<tr>
<td>Negative culture</td>
<td>---</td>
<td>Negative</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>Swirling</td>
<td>Positive</td>
<td>Positive</td>
<td>Positive</td>
<td>---</td>
</tr>
</tbody>
</table>

ECBT, European Committee on Blood Transfusion. AABB, American Association of Blood Banks. BMH, Brazilian Ministry of Health. SCT, Sickle cell trait. WBC, White blood cell count.

*Student’s t test. # Mann-Whitney test. P-values ≤ 0.05 were considered statistically significant.
as severe dehydration and exercise, may trigger kidney lesions, rhabdomyolysis or even death in SCT carriers [17]. This is, to our knowledge, the first study to objectively assess the safety of platelet apheresis donation by SCT carriers and the in vitro quality parameters of their platelet concentrates.

Considering all types of apheresis procedures, platelethropheresis is the second cause of donors’ adverse reactions, behind only plasmapheresis [18]. Possible adverse reactions include blood access injuries, vaso-vagal symptoms and alkalosis caused by citrate overload [18]. In the present study, no SCT donor presented any adverse reaction to the apheresis procedure. Furthermore, the mean number of donations from SCT individuals was similar to that from the overall population and there was no statistically significant difference from the non-SCT group. Such findings suggest that there is no adverse reaction specific to SCT donors that could force this group to discontinue apheresis donation over time.

Whilst buffy-coat platelet concentrates must undergo leukoreduction after collection in order to avoid adverse reactions, platelethropheresis returns leukocytes back to the donor’s circulation during collection [19]. It is considered a very efficient procedure, at least in non-SCT blood. Nevertheless, whether the apheresis technique is as efficient in depleting leukocytes from SCT blood was an unanswered question. The present study showed a similar residual WBC count in both SCT and control groups, as showed in Table 3.

Previous studies have analyzed the in vitro quality of apheresis platelets, particularly the pH of the concentrates [20,21]. In one of these studies, a strong correlation of platelet pH values was found within donors. Furthermore, an association was found between certain donor characteristics, namely age and sex, and a low platelet yield [21]. Both findings suggest a role of donor characteristics in the quality of their platelet products. In none of these studies, however, hemoglobin status has been specified as one of these characteristics.

The quality of a blood component is also influenced by storage lesion, which is an umbrella term that refers to the various mechanisms of deterioration of blood components during storage. Platelets are especially sensitive to bacterial contamination and storage lesion caused by exposure to foreign surfaces, shear stress during centrifugation, acidic pH, trauma, amongst other factors [22]. As a result, platelet units present a short shelf-life of 5 days in average if stored at 22ºC (71.6°F), with the possibility of being extended to 7 days if stored at 18ºC (64.4°F) or if additive solutions are used [23].

Measurement of pH is an important tool for evaluating storage lesion. Values of pH below 6.0 cause irreversible damage to platelets, which become improper for use. The fall in pH is ascribed to the production of CO₂ due to platelet anaerobic glucose metabolism [24]. The platelet yield is a parameter primarily linked to the purpose of the platelet transfusion. In other words, if a platelet unit does not gather enough platelets, its quality is obviously poor, once the clinical end-result will be compromised. However, a recent study has demonstrated that if the platelet count is too high (above 5 x 10¹¹ platelets per unit) in platelet units that underwent Intercept pathogen inactivation, an increase in storage lesion rates was observed, especially in apheresis platelets [25]. In the present work, there was no statistically significant difference of pH and platelet yield between the SCT and the control group.

Although it is not mandatory, the swirling test is another useful way of assessing the quality of platelets, more precisely, their morphological integrity. It consists of a qualitative test, in which light is cast through the platelet concentrate. It is said to be positive when the light is scattered by the unit content, which is an indirect sign that the platelets are morphologically preserved [15]. In our sample, all concentrates presented positive swirling, regardless of belonging to the SCT group or not.

Conclusion

Apheresis platelet concentrates from SCT donors have similar quality parameters to those from non-SCT donors. No increase in the rates of adverse reactions was observed in the SCT group. Therefore, SCT individuals seem to be as good source of apheresis platelets as the general population and blood banks which struggle with scarcity of platelet donations may greatly benefit from routinely allowing SCT individuals to donate by platelethropheresis, especially where there is a high SCT prevalence. Further studies with a greater number of SCT participants and laboratory evaluation of extracorporeal circulation effects may help to better elucidate this issue.

Acknowledgement

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