Hepcidin/Ferritin Quotient Helps to Predict Spontaneous Recovery from Iron Loss following Blood Donation

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Introduction

Determining hemoglobin (Hb) levels prior to blood donation is standard in most countries in order to protect blood donors from becoming critically anemic. Hb levels of prospective blood donors are usually measured in capillary blood obtained by finger prick- ing. Regardless of the fact that the reliability of capillary Hb estimation is discussed controversially [1–4], subclinical iron deficiency (SID) of blood donors cannot completely be excluded by measurement of Hb levels. About 200–250 mg of iron is lost with each blood donation [5]. Individuals with plasma ferritin levels smaller than 15 ng/ml are considered having SID [6]. Individuals with SID do not show clinical signs of iron deficiency while having empty iron stores making them prone to develop clinically apparent iron deficiency syndrome including anemia. SID is present in about 10% of individuals who meet the Hb criteria for blood donation [6]. Oral iron supplementation is generally recommended for subjects with frequent blood donations and for patients donating autologous blood prior to surgical interventions, even though autologous blood donation has become a rare procedure performed in less than 1% of cases. However, the term frequent blood donation is not well defined and the efficacy of such a treatment is rarely monitored. In a study of individuals donating a mean of 15 units of blood over 3.5 years, a beneficial effect of oral iron supplementation on iron homeostasis showed significant inter-individual differences [7]. Thus there is a need to determine the two groups of donors: One group of donors might benefit from oral iron substitution, another group, in contrast, need rather longer interdonation intervals. We aimed to address this issue by trying to define an algorithm which could help to distinguish between the two groups.

Therefore, we analyzed plasma concentration of factors which are involved in erythropoiesis and iron homeostasis, such as erythropoietin, reticulocytes, hemoglobin in reticulocyte, twisted gastrulation protein homolog 1, and growth differentiation factor-15.

Results:

56 days after blood donation, donors’ average ferritin dropped to 55% (range 30–100%) compared to the initial value. Of all tested parameters hepcidin showed the highest and most significant changes beginning 1 day after donation and lasting for the whole period of 56 days. Along with ferritin, there was a high variation in hepcidin levels indicating inter-individual differences in hepcidin response to iron loss. Donors with a hepcidin/ ferritin quotient < 0.3 regained 60% of their initial ferritin after 56 days, while those with a quotient ≥ 0.3 reached less than 50%. Conclusion: As hepcidin appears to integrate erythropoietic and iron-loading signals, clinical measurement of hepcidin (together with the hepcidin-ferritin ratio) may become a useful indicator of erythropoiesis and iron kinetics.

Keywords

Hematology · Red cells · Donors · Health research methodology

Introduction

Determining hemoglobin (Hb) levels prior to blood donation is standard in most countries in order to protect blood donors from becoming critically anemic. Hb levels of prospective blood donors are usually measured in capillary blood obtained by finger pricking. Regardless of the fact that the reliability of capillary Hb estimation is discussed controversially [1–4], subclinical iron deficiency (SID) of blood donors cannot completely be excluded by measurement of Hb levels. About 200–250 mg of iron is lost with each blood donation [5]. Individuals with plasma ferritin levels smaller than 15 ng/ml are considered having SID [6]. Individuals with SID do not show clinical signs of iron deficiency while having empty iron stores making them prone to develop clinically apparent iron deficiency syndrome including anemia. SID is present in about 10% of individuals who meet the Hb criteria for blood donation [6]. Oral iron supplementation is generally recommended for subjects with frequent blood donations and for patients donating autologous blood prior to surgical interventions, even though autologous blood donation has become a rare procedure performed in less than 1% of cases. However, the term frequent blood donation is not well defined and the efficacy of such a treatment is rarely monitored. In a study of individuals donating a mean of 15 units of blood over 3.5 years, a beneficial effect of oral iron supplementation on iron homeostasis showed significant inter-individual differences [7]. Thus there is a need to determine the two groups of donors: One group of donors might benefit from oral iron substitution, another group, in contrast, need rather longer interdonation intervals. We aimed to address this issue by trying to define an algorithm which could help to distinguish between the two groups. Therefore, we analyzed plasma concentration of factors which are involved in erythropoiesis and iron homeostasis, such as erythro-
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Plasma levels of ferritin and hepcidin change during infection and inflammation; thus leukocyte count and the concentration of C-reactive protein (CRP) have to be tested concurrently in order to avoid obtaining biased results for ferritin and hepcidin.

In our study we aimed to provide a tool (a single parameter or an algorithm) helping to better estimate recovery from iron loss following blood donation and to better indicate donors profiting from oral iron supplementation in contrast to those who would rather need longer interdonation intervals in order to cope with iron loss. The hepcidin/ferritin quotient here presented qualifies for such an estimation.

Material and Methods

Human Subjects

The study was conducted according to the principles expressed in the Declaration of Helsinki. The study was approved by the Institutional Review Board (Ethical Committee) of Ulm University. All blood donors provided written informed consent for the blood donation and the collection of samples.

The retrospective part of the study on ferritin levels was conducted on 1,664 male first-time donors (representing general population) and 5,073 male repeat donors with a history of at least 10 blood donations. Donors with plasma ferritin levels below 15 ng/ml were considered having SID.

The prospective part of the study was performed on 27 male non-vegetarian healthy regular blood donors without any bleeding events (i.e. blood donations or surgical treatments) within the previous 4 months. Donors were instructed not to change their dietary habits and, specifically, not to take any oral iron supplements.

With regard to the fact that inflammation affects iron homeostasis including concentrations of hepcidin and ferritin, data from donors with CRP levels higher than 5 mg/ml or leukocyte count higher than 10 x 10³/µl were excluded from the analysis.

Blood Samples

Blood samples (10 ml) were taken at the day of blood donation (day 0) as well as 1, 3, 7, 10, and 56 days thereafter in order to measure the following parameters: red cell count, leukocytes (including subtypes), platelet count, Hb, reticulocytes (Reti), Hb in reticulocytes (Hb-Reti), EPO, ferritin, Trf-R, Trf, hepcidin, GDF-15, TWSG1, GDF-15, hepcidin, CRP, ALT, AST, and creatinine.

Enzyme-linked immunosorbent assays (ELISA) for GDF-15 (R&D Systems, Wiesbaden-Nordenstadt, Germany), TWSG1 (Hölzel Diagnostika, Köln, Germany), and hepcidin 25 (DRG Diagnostics, Marburg, Germany) were performed according to the manufacturers’ instructions and using a microplate reader (POLARstar Omega, BMG Labtech, Ortenberg, Germany) to measure absorbance. Blood cell count was measured using the automated blood analyzer CELL-DYN (Abbott, Wiesbaden, Germany).

CRP, ALT, AST, and Trf levels were assessed on ABX pentra 400 (HORIBA Medical, Axon Lab AG, Germany) using reagents from DiaSys Greiner GmbH (Flacht, Germany). Ferritin was measured using Architect i1000 SR and reagents from Abbott (Wiesbaden, Germany) according to manufacturer’s instructions.

EPO was measured on DXI800i (Beckman Coulter, Krefeld, Germany) using Access EPO Assay from Beckman Coulter according to manufacturer’s instructions. Reti-Hb was assessed on hematology analyzer XE-2100 ( Sysmex Europe GmbH, Norderstedt, Germany).

Statistics

Pearson’s product-moment correlation and Student’s t-test were calculated using R Statistical Software, p values ≤ 0.05 were considered statistically significant.

Hepcidin (EPO), ferritin, iron transport protein transferrin (Trf) and its receptor (Trf-R), hepcidin, growth differentiation factor-15 (GDF-15), and twisted gastrulation protein homolog 1 (TWSG1).

EPO rapidly rises in anemia as a result of anemia-induced hypoxia and falls as Hb is corrected [8].

Ferritin is a ubiquitous intracellular protein that stores iron and releases it in a controlled fashion. Plasma ferritin is also an indirect marker of the total amount of iron stored in the body; hence serum ferritin is used as a diagnostic test for iron deficiency.

Hepcidin is a liver-derived hormone that regulates iron absorption and recycling via its effects on ferroportin, the cellular iron export protein [9]. Low hepcidin preserves ferroportin and permits increased iron recycling via its effects on ferroportin, the cellular iron export protein [9].

Even though EPO significantly decreases serum hepcidin levels in humans, it does not appear to be a direct regulator of hepcidin but acts through erythroferrone (ERFE) which is a hormone produced by erythrobasts.

GDF-15 and TWSG1 have been identified through transcriptome analysis as putative erythroblast-derived factors that modulate hepcidin. Recombinant GDF-15 inhibits expression of hepcidin in hepatic cell lines. Expression of GDF-15 peaks in late erythrobasts and is also associated with erythroblast apoptosis. Based on studies in patients with beta-thalassemia and other iron-loading anemias and supported by related animal models and cell-based in vitro studies, it has been proposed that EPO-stimulated erythroblasts secrete mediators that act on the liver to suppress hepcidin production. GDF-15 and TWSG1 have been proposed as a hepcidin suppressor in beta-thalassemia [21, 23].

Trf-R is expressed by erythroblasts showing an increased expression of Trf-R from early to intermediate stages and a decreasing one with maturation. Thus, Trf-R mass reflects number of immature erythroid cells.

Divergent patterns of EPO, soluble Trf-R, and GDF-15 may reflect differing physiologies of these parameters and provide insights into erythropoiesis and iron homeostasis: EPO closely reflects anemia but may also reflect erythroid suppression. Soluble Trf-R reflects overall erythropoietic activity but not necessarily ineffective erythropoiesis or dyserythropoiesis with apoptosis, which appears best reflected by GDF-15.

Given that liver diseases impact iron homeostasis and erythropoiesis by influencing the expression and plasma concentration of hepcidin, EPO, ferritin, and transferrin, liver enzymes (alanine aminotransferase (ALT), aspartate aminotransferase (AST)) need to be tested when measuring these parameters.

Renal excretion plays a major role in clearance of hepcidin. Consistently, patients with chronic kidney diseases have higher hepcidin plasma levels. EPO is expressed in kidney, thus patient with kidney diseases show inadequate EPO concentrations. For these reasons, it is necessary to assess plasma creatinin concentration when measuring EPO and hepcidin.
Results

In our retrospective study on 6,737 male donors, the average ferritin plasma concentration of 1,664 first-time donors (representing general population) was 107.9 ng/ml (SD = 76.2) with 1.3% of individuals presenting SID (i.e. ferritin levels below 15 ng/ml). On the other hand and consistent with published data [6, 26], 15% of repeat donors (n = 5,073) with a history of at least 10 blood donations presented with SID; the average ferritin level of this population was significantly lower being 49.63 ng/ml (SD = 46.5).

The prospective part of the study was performed on 27 male donors with an average age of 34 years (range 19–55 years, mode 22 years) and an average body mass index of 26 kg/m² (21–32 kg/m², mode 25 kg/m²).

There were no clinical or laboratory signs of infection, inflammation, and liver or kidney diseases with normal values for CRP, and leucocyte count, ALT, AST, and creatinine.

All donors met the Hb criteria for blood donation with an average predonation Hb level of 159 g/l (134–185 g/l, mode 161 g/l). Within this group, predonation SID was present in 4 individuals (15%) with ferritin levels ranging from 3.5–11.8 ng/ml resulting in an average of 7.5 ng/ml.

The average plasma ferritin concentration of donors without SID (n = 23) dropped to 55% compared to initial values (i.e. before blood donation) 56 days after blood donation (fig. 1). These results were consistent with published data [26, 27], indicating that 56 days may not be sufficient to recover from iron depletion due to blood loss, if iron supplementation is not provided.

One day after blood donation (day 1), EPO reached its maximum (190% compared to baseline level on day 0), stayed at significantly higher levels until day 10, and declined to 120% on day 56. Accordingly, reticulocytes increased to 150% on day 7 and normalized until day 56 when Hb levels also reached their initial values (fig. 1), all reflecting normal regulation of erythropoiesis in our study group.

Consistent with the increased number of reticulocytes at day 7, plasma concentration of Trf-R – which is expressed on erythroblasts – raised constantly and reached significance 10 days after blood donation (fig. 1). Thus, Trf-R did not qualify for an early predictor. Similarly, average Trf levels reached significant changes not earlier than 56 days and was just 10% compared to its average initial value (data not shown).

No statistical significant changes could be seen for Hb-Reti, GDF-15, and TWSG1 (fig. 1).

Of all tested parameters hepcidin showed the highest and most significant changes beginning 1 day after blood donation and lasting for the whole period of 56 days. Average ferritin and hepcidin plasma concentrations decreased constantly with ferritin levels reaching statistically significant low levels not earlier than 10 days, while the drop in plasma hepcidin concentrations became statistically significant at day 1, already (fig. 1).

Even though the decrease in donors’ average ferritin was significant (p = 0.02), we assessed high inter-individual differences in recovery from iron loss, when comparing the ferritin quotient day 56 / day 0 (d56/d0) ranging from 30 to 100% (average 55%). Thus, we compared the group of donors with ferritin recovery < 55% with those having reached ≥55% of their initial ferritin values after 56 days. For the mentioned two subgroups, neither hepcidin level on day 3 by itself nor ferritin level at day 3 qualified as an indicator to be able to predict which donors would recover best from iron loss (fig. 2).
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Discussion

Oral iron supplementation is generally recommended to blood donors, based on a randomized, placebo-controlled, double-blind study demonstrating that 20 mg of elemental iron per day can adequately compensate for iron loss in males and females who donate whole blood up to 4 (females) or 6 times per year (males). Nevertheless, there are significant inter-individual differences in intestinal iron uptake [7]. Furthermore, Mast and colleagues [28] identified a subgroup of blood donors who do not develop SID despite repeated and frequent blood donations without any oral iron supplementation. This subpopulation is heterozygous for the H63D mutation in HFE and shows significantly low levels of hepcidin and decreased hepcidin/ferritin quotients compared to other individuals. Thus, we tried to find an algorithm which can help to better distinguish between donors who benefit from iron supplementation and those who rather need longer interdonation intervals.

Testing a broad spectrum of parameters we wanted i) to assure the reliability of our data by confirming that the subjects in our study group did not suffer from (subclinical) infections, inflammations, and kidney or liver diseases interfering with measured parameters, ii) to assess erythropoiesis which also impacts iron homeostasis, and iii) to maximize the chance to define an early factor or algorithm which can help to distinguish between donor subpopulations.

Not being sure about the width of the diagnostic window for such an early factor/algorithm, we decided to ask our study subjects not to take any iron supplements which could have narrowed or even closed the possible time window to detect any inter-individual differences in tested parameters. Nevertheless, a prospective study comparing donors with and without oral iron supplementation needs to be performed in order to calculate the predictive value of the hepcidin/ferritin quotient.

Our study was performed on male donors only in order to exclude biases such as menstrual bleeding or different relative blood loss after a donation due to discrepancies in average body weight or dietary habits between the two genders. However, iron homeostasis seems to differ between men and women [7] emphasizing the need for further studies including both genders.

The logarithm of the ratio of the soluble Trf-R to ferritin concentration (log[Trf/R]), which was shown to have a highly linear correlation to body storage iron, is currently the most precise measure of body storage iron available. Thus we also checked for this parameter and confirmed the inability of donors to spontaneously recover from iron loss following blood donation (fig. 1).

Our observation of a weak correlation between GDF-15 and hepcidin levels was consistent with published data [26, 29], indicating that GDF-15 may not directly influence hepcidin levels [30, 31].

As hepcidin appears to integrate erythropoietic and iron-loading signals, clinical measurement of hepcidin (together with the hepcidin-ferritin ratio) may become a useful indicator of erythropoiesis and iron kinetics. Detection of SID in blood donors is important not only to prevent donors from becoming anemic after subsequent blood donations but also because of the fact that iron deficiency affects various
other cellular processes, given that iron is an important constituent of a series of enzymes. For example, previous studies suggested that iron deficiency affects DNA synthesis [32, 33], the immune system, and energy metabolism through impaired mitochondrial electron transport [35, 36].

Furthermore, donors with SID are more prone to develop donation-induced Hb decrease possibly leading to subsequent experience of deferral from blood donation due to non-eligible Hb values. Considering that about 40% of repeat blood donors do not reappear when experiencing a deferral it is of great importance to recognize SID early enough to avoid subsequent Hb-based deferral and also to establish an algorithm to make more individualized recommendations for oral iron uptake which – by itself – would also increase blood donors’ satisfaction and trust in medical surveillance performed by blood donation centers.

We therefore suggest ferritin measurements at the time of first blood donation and once a year for donors with more than 4 donations within 12 months along with a general recommendation for oral iron uptake based on data from Radtke et al. [27] who have already demonstrated that most (but not all) blood donors benefit from a daily oral iron supplementation of 20 mg. Nevertheless, individualized recommendations (particularly for repeat donors) are advisable as a long-term strategy.

Based on our data, we believe that the hepcidin/ferritin quotient may provide an algorithm helping to decide if a donor may benefit from oral iron supplementation or rather from a longer interdonation interval. Assuming that individual hepcidin responsiveness to iron loss does not change, we suggest calculating hepcidin/ferritin quotient for repeat donors (just once) at the time of their 4th donation. This would be a reasonable strategy considering also economic issues.

**Disclosure Statement**

The authors certify that they have no affiliation with or financial involvement in any organization or entity with a direct financial interest in the subject matter or materials discussed in this manuscript.

**References**


