Measurement of Red Cell Lifespan and Aging

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Introduction

Human red blood cells (RBC), after differentiating from erythroblasts in the bone marrow, are released into the blood and survive in the circulation for approximately 115 days. In humans and some other species, RBC normally survive in a nonrandom manner. This means that all of the RBC in an age cohort are removed by the reticuloendothelial system at about the same time. In practice there is considerable variation in the lifespan of human RBC. In a normal individual with a mean RBC lifespan of 115 days, this value may vary between 70 and 140 days [1, 2]. Among individuals the mean lifespan varies by approximately ±15% [1]. As discussed below, there are circumstances in which even this relatively small normal variation can be important.

An accurate method to determine RBC lifespan that took advantage of emerging knowledge of the ABO system was first published by Winifred Ashby in 1919 [3, 4]. The differ-
tial agglutination technique used anti-A and anti-B antiserum to measure the lifespan of type O RBC that were given to type A or type B recipients. This technique was difficult to perform since it included a quantitative agglutination of the subjects RBC in posttransfusion blood samples that was subject to many variables, followed by manual counting of the non-agglutinated RBC to quantitate the remaining transfused RBC. Furthermore, this method could not be applied to autologous RBC. Nevertheless, the Ashby technique was accurate, and remained the standard for measurement of allogeneic RBC lifespan for almost 40 years. It is an example of methods in which a representative sample of circulating RBC from the donor that can be distinguished from the recipient’s RBC is infused into the recipient and identified in the circulation for (ideally) the lifetime of the RBC. Typically, the population of RBC that are infused represent the entire range of RBC ages in the donor, from RBC that were just released from the bone marrow to RBC that are senescent and due for removal from the circulation. In this type of ‘population’ study normal RBC are lost from the circulation immediately after reinfusion, and are removed in a linear manner for the greater part of the lifespan. Today, RBC population studies are performed with a label that is placed on the RBC ex vivo, making it possible to study both donor and autologous RBC.

The survival of labeled RBC can be analyzed in a number of ways to characterize the lifespan and help to identify mechanisms of RBC removal from the circulation. There are 3 basic time periods that must be considered. The first is the time it takes for the infused, labeled RBC to mix with the circulating unlabeled RBC. This is normally about 5 min [5], but may be longer in some patients, especially those with splenomegaly. During this time the concentration of labeled RBC cannot be determined, and any severely damaged RBC that are lost from the circulation are not accounted for unless there is an independent determination of the expected postinfusion concentration that can be compared with the measured value immediately after mixing occurs. For example, when measuring the survival characteristics of stored RBC a sample of fresh RBC (which are assumed to survive during the mixing period) with a different label may be co-infused. The second time period is the first 24 h after infusion of the labeled RBC. This period is critically important since RBC that have been damaged by storage or labeling will typically be removed from the circulation during this time. RBC that make it through this window will usually have good long-term survival [6]. In the evaluation of cell storage media the U.S. Food and Drug Administration has relied mostly on 24-hour recovery, setting a minimum acceptable mean value of 75% with additional requirements for the range of values [7]. The third time period is long-term survival which is about 4 months. There has been added interest in the long-term RBC survival after storage and transfusion in recent years, as studies have become available that imply detrimental effects during this time period in patients who received blood that was stored for longer periods [8]. Accurate measurement of long-term survival requires determination of the amount of remaining labeled RBC for all or most of the RBC lifespan. The current standard label, $^{51}$Cr, is not well-suited for this task as explained below. Optimal determination of long-term survival also requires a steady state situation, with the important variable depending on the label used. For example, $^{51}$Cr and other radioactive labels require a stable blood volume, while biotin and other flow cytometry-based labels require a stable number of total circulating RBC.

Another type of RBC lifespan study uses a synchronized age cohort of RBC that is labeled metabolically in vivo during a short period of time with either a radioactive or stable isotope. The isotope is typically delivered via orally administered labeled glycine, which is incorporated into both the heme and globin components of hemoglobin. This type of analysis does not average the survival of RBC generated for 4 months, but instead yields the survival of RBC that were all produced at about the same time. In a ‘cohort’ study it is possible to follow the various stages in the RBC lifespan, including release from the bone marrow during a time period that reflects normoblast differentiation kinetics, a long plateau period during which the only normal loss of label is related to presumably denatured and aggregated hemoglobin that has been removed from RBC that continue to circulate [9, 10], and finally the removal of RBC during a time period that represents the variability in RBC lifespan. An advantage of cohort studies is that all of the RBC coming out of the marrow are included in the analysis. This may not be true for population studies, since short-lived RBC will be underrepresented.

### Methods to Determine RBC Survival Characteristics

#### Methods for Labeling a Representative Population of Circulating RBC

$^{51}$Cr

$^{51}$Cr is a radioisotopic noncovalent population label. In the form of Na$_2$$^{51}$CrO$_4$, it is incubated with packed or washed RBC [11–13] and incorporated into RBC as chromate VI through anion channels. Once inside the RBC it binds noncovalently to hemoglobin, and over 90% of the label is incorporated into RBC. After reinfusion, Cr elutes from the labeled RBC at a relatively constant rate of about 1% per day [12]. Advantages of this label are that it has been used extensively to measure human RBC survival characteristics and is available in approved formulations for this purpose. Labeling is convenient and standardized, and the counters required for quantitation are widely available. Its half-life (27.8 days) is well-suited for a 30-day RBC survival analysis which is adequate for many purposes. A disadvantage is that in many countries $^{51}$Cr cannot be used due to radioactivity. Even where it can be used, such as the USA, it is avoided in chil-
dren and pregnant women. In research studies blood samples may be considered to be radioactive waste and must be handled accordingly, making parallel assays difficult. While standard tables, derived by assuming a linear survival of 115 days [14] or by comparison with diisopropylphosphofluoridate (DF32P)-labeled RBC [15], are available to correct for elution up to 30 days, these are average values and elution for individuals and RBC subpopulations [16] may vary significantly. The combination of elution and decay makes it difficult to follow the labeled RBC accurately for the entire RBC lifespan.

**Biotin**

Biotin is a nonradioactive covalent population label. Washed RBC are reacted with N-hydroxysuccinimide(NHS)-biotin or sulfo-NHS-biotin to covalently label membrane proteins with biotin [1, 2]. RBC from postinfusion blood samples are reacted with fluorescently conjugated streptavidin and the biotin-labeled RBC quantitated as a percentage of the total RBC by flow cytometry. The nonradioactive biotin label has been used in newborns to determine RBC volume and lifespan [17] and has been shown to correlate with 51Cr-labeled RBC [18, 19]. There is some loss of label from the RBC as they age, but all the labeled RBC remain in the positive flow cytometric region, and thus the loss of label has no effect on quantitation. This, together with the very high number of RBC that can be analyzed in each postinfusion sample, and the very low false-positive number in the cytometer, makes it possible to accurately determine the entire RBC lifespan using microliter-sized blood samples. Since it is possible to place different levels of biotin on separate cell populations and distinguish them in the flow cytometer, RBC lifespan can be determined for up 3 populations of RBC concurrently [2]. The biotin-labeled RBC can be examined with multicolor flow cytometry to determine how variables change as the RBC age. Examples include fetal hemoglobin (HbF)-containing RBC and externalized phosphatidylserine [19–22]. The labeled RBC can be isolated by means of streptavidin-coated magnetic beads and assayed by high-performance liquid chromatography (HPLC) or other methods [1]. Examples of this type of analysis include HbF and HbA1c quantitation [1, 19–22]. The major potential disadvantage of the biotin label is the development of antibodies against biotinylated RBC. In one report [23] transient antibodies were detected using an agglutination technique in 3 of 20 subjects that were given biotin-labeled RBC. None of these antibodies appeared to have an effect on RBC lifespan, and none could be detected after 1 year.

**Methods Utilizing Metabolic Precursors or Products**

**Carbon Monoxide**

Carbon monoxide (CO) can be used for the quantitation of heme turnover. This method is based on the principle that the catabolism of heme is the only endogenous source of CO, with 1 molecule of CO resulting from each heme. Therefore, CO production is directly related to heme turnover [24–28]. However, hemoglobin accounts for only about 80% of heme turnover, and this may vary in disease states. Production of CO may be quantitated over several hours by means of a rebreathing system [24–28], or an estimate made by measurement of alveolar CO [29–31]. This method has the advantage that measurement of CO production can be used to measure heme turnover serially over a relatively short time period. Disadvantages include that while heme turnover can be reliably determined with the aid of a rebreathing system, there is uncertainty concerning the contribution of other heme-containing proteins. Alveolar sampling, while convenient, is subject to environmental influences.

**Labeled Glycine**

This is a stable isotope metabolic cohort label. Glycine is a precursor of heme and globin, and may be given orally to label the hemoglobin in RBC. Glycine is incorporated into RBC precursors over the entire range of normoblasts that synthesize hemoglobin, and thus the amount of label in the blood increases for up to 25 days after oral administration [32]. It has the advantages that no ex vivo labeling is required, and all RBC released from the bone marrow are included in the analysis. A disadvantage is that this method requires mass spectroscopic analysis.

**Applications**

RBC survival determinations are rarely performed for clinical purposes, but there are a number of investigational circumstances in which they are useful, including: i) evaluation of RBC storage and pathogen inactivation [33–35]; ii) studies of the pathophysiology of sickle cell disease [19–22]; iii) the importance of RBC lifespan in diabetes as an explanation of mismatches between blood glucose and HbA1c [1]; iv) studies of the optimum RBC transfusion product for premature newborns [17]; v) poorly understood anemias including those associated with chronic inflammation [36] and the elderly; vi) investigations of RBC aging and senescence. Studies of transfusion products require a population-type, ex vivo label. However, the metabolic glycine-based cohort label may be ideal for studies of disease pathophysiology.

**The Biotin Label in RBC Aging Studies**

The most useful label for RBC aging studies is biotin. This label not only allows precise quantitation of RBC survival for the full lifespan, but also makes possible detailed evaluation of changes that occur in the circulation as the RBC age.
The shape of the curve. This is useful in plotting changes that occur as the RBC age. However, for many purposes a simpler analysis is adequate, including a linear extrapolation to the X-axis or a time to 50% removal. While it is possible to study a relatively pure population of old RBC using this technique by allowing them to age in vivo, it is not possible to study young RBC without the presence of older RBC. This limitation can be partially overcome by isolating for study very young unla-

**Assays of Aging RBC Based on Multicolor Flow Cytometry**

Using the biotin label and 2 fluorescent colors it is possible to measure any property that can be analyzed by flow cytometry as RBC age. The first color (e.g. FITC) is used to identify the biotin-labeled RBC, and the second color (e.g. phycoerythrin) to evaluate the property. Since a large number of RBC are counted this analysis can be continued for essentially the entire RBC lifespan. In animal studies with in vivo biotinylation, this technique has been used to determine cell age-related changes in membrane immunoglobulin [43, 46], aminophospholipid translocase (flippase) activity [47], enzyme activities [40], and many other properties [38, 39, 42] In human studies with ex vivo biotinylation, measured variables include HbF-containing RBC [19, 20, 22], RBC with external phosphatidylserine [21], and the time for disappearance of reticulocytes [19].

**Assays of Aging RBC Based on Magnetic Isolation**

Another powerful analytical approach, which can be applied with either in vivo or ex vivo labeling, is the isolation of bi-

**Time-Dependent Changes in the Age of Biotin-Labeled RBC after Reinfusion**

In animal studies it is possible to label all the RBC in vivo with biotin with an intravenous injection of sulfo-NHS-biotin [37–43]. At subsequent times the biotin-negative RBC are all younger than the time since administration of the labeling reagent, and the biotin-positive RBC are all older than that time. The negative RBC include reticulocytes, which may bias any assays performed on those RBC. It is also possible in animals to combine in vivo biotinylation with another label to follow a true age cohort as the RBC age in the circulation [44]. In human studies RBC are labeled ex vivo, and in a typical adult reinfusion of 10 ml of labeled RBC will give about 0.5% positive RBC in the flow cyrometer immediately after reinfusion and mixing in the circulation. Initially, the labeled and unla-

**Fig. 1.** Survival of normal RBC labeled with biotin. Six hematologically normal subjects (NDM1-NDM6) are shown. The 1-day recovery, compared to 10–20 min after reinfusion, was 96.8 ± 1.4 (1 standard deviation) %. The value at 1 day was taken as 100% for evaluation of long-term lifespan. This research was originally published in Blood in 2008 [1].

**Fig. 2.** RBC were labeled with biotin and reinfused into a normal subject. At the times indicated labeled cells were isolated with streptavidin-coated magnetic beads and the fraction HbA1c determined by HPLC. The mean RBC age was calculated for each time point (inverted triangles) according to [45]. The open square represents the value for TfR+ reticulocytes isolated with anti-TfR-coated magnetic beads and assayed the same way. These data indicate that the amount of HbA1c increases linearly as the RBC age. This research was originally published in Blood in 2008 [1].
otin-labeled RBC with the aid of streptavidin-coated magnetic labeled beads [1]. For human studies in which 10 ml of packed labeled RBC are reinused, there are initially about 0.5% labeled RBC in the circulation. This corresponds to 5 μl of labeled RBC per ml of packed RBC. After 90% of the labeled RBC are removed from the circulation, and the remaining labeled RBC in a normal subject are at least about 110 days old, there are about 0.5 μl of labeled RBC per ml of packed RBC. Any assay that can be performed on these volumes of RBC can therefore be applied to aging RBC. For example, hemoglobin composition of aging sickle RBC and the rate of formation of HbA1c have been determined by HPLC [1] (fig. 2).

Conclusion

It is nearly 100 hundred years since the first accurate measurements of red cell lifespan and the first clues of their age-dependent removal from the circulation. During this time there has been great interest in mechanisms involved in determining the proper time for removal and marking the cell for uptake by the reticuloendothelial system. Information concerning the changes that take place as a red cell ages is central to these questions, and the early differential agglutination techniques used for measuring lifespan were well suited for these studies since they allowed the isolation of aged RBC. However, they were not applied in this way, and their complete replacement by radioisotope labels in the mid 20th century made such studies impossible. Only recently with the introduction of the biotin label has a method become available that allows the detection, analysis, and isolation of aging RBC, and thus detailed studies of their properties.

Disclosure Statement

The author declared no conflict of interest.

References


