Little is known about age-related changes in red blood cell (RBC) membrane transport and homeostasis. We investigated first whether the known large variation in plasma membrane calcium (PMCA) pump activity was correlated with RBC age. Glycated hemoglobin, Hb A1c, was used as a reliable age marker for normal RBCs. We found an inverse correlation between PMCA strength and Hb A1c content, indicating that PMCA activity declines monotonically with RBC age. The previously described subpopulation of high-Na\(^+\), low-density RBCs had the highest Hb A1c levels, suggesting it represents a late homeostatic condition of senescent RBCs. Thus, the normal densification process of RBCs with age must undergo late reversal, requiring a membrane permeability increase with net NaCl gain exceeding KCl loss. Activation of a nonselective cation channel, Pcat, was considered the key link in this density reversal. Investigation of Pcat properties showed that its most powerful activator was increased intracellular Ca\(^{2+}\). Pcat was comparably selective to Na\(^+\), K\(^+\), choline, and N-methyl-D-glucamine, indicating a fairly large, poorly selective cation permeability pathway. Based on these observations, a working hypothesis is proposed to explain the mechanism of progressive RBC densification with age and of the late reversal to a low-density condition with altered ionic gradients.

Introduction

Our recent findings of a large variation in plasma membrane Ca\(^{2+}\) pump (PMCA) activity of normal human red blood cells (RBCs) prompted the question of whether this variation was related to cell age and, if so, whether an age-related decline in PMCA activity might account for some observed changes in aging RBCs. The PMCA is the only active Ca\(^{2+}\) transporter in human RBCs. At saturating [Ca\(^{2+}\)], levels, its mean maximal Ca\(^{2+}\) extrusion rate, \(V_{\text{max}}\), is about 10 to 15 mmol/(340 g Hb)\(^{-1}\)h\(^{-1}\), whereas the normal mean pump-leak Ca\(^{2+}\) turnover is only about 50 mmol/(340 g Hb)\(^{-1}\)h\(^{-1}\). Thus, in physiological conditions the mean pump activity is a small fraction of its mean \(V_{\text{max}}\). This balance, however, is far from uniform among the RBCs, whose PMCA activity showed a broad unimodal distribution pattern, with a coefficient of variation of about 50\%. The cells with the weaker Ca\(^{2+}\) pumps would have higher [Ca\(^{2+}\)], levels than those with stronger pumps. Because RBC [Ca\(^{2+}\)], levels control several transport and biochemical processes affecting volume regulation, metabolism, and rheology, the resulting variations in [Ca\(^{2+}\)], may generate marked differences in RBC composition. There were early indications that PMCA activity decreased with RBC age, but there has been no systematic study of the age-activity relation over the full range of pump variation observed.

Here we devised a method to segregate RBCs according to their PMCA strength to be able to measure directly the age of the segregated RBCs. We used the fraction of glycated hemoglobin, Hb A1c, as a reliable measure of RBC age. Because protein glycosylation is a nonenzymic process linear with time and glucose concentration, the Hb A1c fraction is a reliable age marker for normal human RBCs, which, unlike those of diabetic patients, are exposed to relatively stable glucose levels in vivo. In the course of investigating the age-activity relation of the PMCA we observed an unexpected reversal of the main pattern of RBC behavior, as detailed below, which could be explained by the presence of a small fraction of aged RBCs unable to dehydrate by K\(^+\) permeabilization. Such RBCs had been described before, but their age had not been measured. We now demonstrate that they have a high Hb A1c content, suggesting that they represent a normal terminal condition of senescent RBCs.

For a dense, low-Na\(^+\), aging RBC to change to a lesser dense, higher-Na\(^+\), terminal condition in the circulation, it must undergo a change in membrane permeability whereby net NaCl gain exceeds net KCl loss, leading to progressive net salt and water gain. A likely candidate for such a process was the nonselective cation permeability pathway originally described by Crespo et al\(^{17}\) as "prolytic" (following a nomenclature originally used by Ponder to describe effects of lysins on RBC homeostasis\(^{18,19}\)). The prolytic pathway exhibited a sufficiently high Na\(^+\) permeability to exceed the extrusion capacity of the Na pump, leading to the swelling and eventual lysis of the affected RBCs. We undertook a detailed investigation of the main properties of the "prolytic" pathway, which we designate here "Pcat," applying methods that examine the responses of the entire RBC population to the factors that activate this pathway.

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Materials and methods

Approval for these studies was obtained from the institutional review boards of the University of Cambridge, United Kingdom, and from the Albert Einstein College of Medicine, Bronx, NY.

Solutions

The main solutions used (to which specific additions were made, as noted) were (all concentrations in mM) as follows: (1) “90K”: KCl, 90; NaCl, 55; HEPES-Na, pH 7.5 at 37°C, 10, and MgCl₂, 0.15. SCN was added with special care to minimize aspiration of lighter RBCs. The washed RBCs buffy coat (platelets and white cells) was removed after each wash, with added EGTA, and washed twice more with solution 90K alone. The isoelectric point of impermeant cell anions and RBC hydration that are specified otherwise in the legends of figures, the final concentrations of the most frequently added solutes were (in mM): NaOH-neutralized EGTA (ethylene glycol-bis(2-aminoethylether)-N,N,N',N'-tetraacetic acid), 0.1; CaCl₂, 0.12; inosine, 5; ionophore A23187 (from 2 mM stock in ethanol), 0.02%; giving a coefficient of variation of 0.35%. Thus, the Ca²⁺-free RBCs above the oil and the pelleted dense cells still containing Ca²⁺ were harvested into microtube vials by gentle resuspension in 0K medium and sample for Hb to estimate their yields. After centrifugation the RBC pellets were refrigerated for 24 to 48 hours or frozen for later determination of Hb A1c.

This method provided a cumulative separation of RBCs with a PMCA Vₘₐₓ above or below a given value at each time point. In the earliest samples nearly all the RBCs would be in the pellets but, with time, increasing proportions of RBCs would have extruded their Ca²⁺ and would remain on top of the oil. Hb A₁c measurements required at least 2.5 to 3 μL of RBCs, and when the yields (from the top and bottom of the oil layer in the initial and final samples, respectively) were too small to measure Hb A₁c, the RBCs harvested over 2 consecutive sampling periods were pooled.

Preparation of RBCs for the age-PMCA correlation study

After obtaining informed written consent in accordance with the Declaration of Helsinki, venous blood of healthy volunteers was drawn into tubes with EDTA, washed twice in ice-cold 90K solution with added EDTA, and washed twice more with solution 90K alone. Theuffy coat (platelets and white cells) was removed after each wash, with special care to minimize aspiration of lighter RBCs. The washed RBCs were suspended at 10% hematocrit (Hct) in 90K supplemented with inosine and Ca²⁺-lactate and temperature-equilibrated for 10 minutes at 37°C before the Ca²⁺-load protocol.

Preparation of RBCs for the study of Pcat properties

RBCs obtained as above were washed twice in ice-cold 90K solution with added EGTA and twice more with solution 90K alone. Theuffy coat (platelets and white cells) was carefully aspirated after each wash to ensure near complete white cell removal. The washed RBCs were suspended at 5% Hct in 90K supplemented with inosine (5 mM), gentamicin, and Ca²⁺, as indicated, and temperature-equilibrated for 10 minutes at 37°C before addition of ionophore. Other additions, ion replacements, as well as the specific concentrations of Ca²⁺, ionophores, and inhibitors used in the experiments are indicated in the figure legends.

Protocol to segregate RBCs according to PMCA strength

We designed the following method to separate RBCs according to their maximal Ca²⁺ extrusion capacity (PMCA Vₘₐₓ). The RBCs were rapidly and uniformly loaded with about 500 μmol(340 g Hb)⁻¹ of total Ca²⁺, [Ca⁵⁺], by addition of 10 μM ionophore A23187 to the 10% Hct suspension containing 120 μM external Ca²⁺, [Ca²⁺]. The 90K medium prevented net K⁺ fluxes via Ca²⁺-activated Gardos channels, thus preserving the original RBC volumes.22 After 2 minutes to allow Ca²⁺ equilibration, the ionophore was extracted to restore the original low-membrane Ca²⁺ permeability by transferring the suspension into 10 volumes of stirred, ice-cold 90K medium with 1.5% albumin. The low temperature inhibited the PMCA, trapping the original RBC Ca²⁺ load.23 The suspension was spun at 2500g for 5 minutes at 0°C. The RBC pellets were pooled and washed twice more at 0°C in solution 90K with albumin and three in 90K alone.

The washed, ionophore-free packed RBCs (about 75% Hct) were resuspended (final Hct, about 14%) in 90K medium supplemented with inosine in a magnetically stirred glass vial in a water bath at 37°C; this immediately reactivated Ca²⁺ extrusion22 (t₀ = 0) at the pump Vₘₐₓ rate of each RBC. At 15- to 30-second intervals, 0.5 mL samples of this suspension were delivered into tubes containing 12 mL of 0K medium at 0°C to 4°C to instantly block further RBC Ca²⁺ extrusion and trap their residual Ca²⁺ content. Because Gardos channels have a relatively low temperature dependence,26 dilution into 0K medium triggered rapid dehydration of any RBCs containing sufficient Ca²⁺ to activate the channels. After 40 minutes in the ice bath to allow maximal dehydration of the Ca²⁺-containing RBCs, the tubes were centrifuged and the RBCs were resuspended in 1 mL of 0K medium at 20°C, layered onto 0.4 mL of DEP oil (1.117 g/mL) in 1.5 mL microtube vials, and spun 1 minute at 10 000 g. The Ca²⁺-free RBCs above the oil and the pelleted dense cells still containing Ca²⁺ were harvested into microtube vials by gentle resuspension in 0K medium and sample for Hb to estimate their yields. After centrifugation the RBC pellets were refrigerated for 24 to 48 hours or frozen for later determination of Hb A₁c.

Reproducibility of the Hb A₁c measurements was estimated on 20 aliquots of a single blood sample, whose mean ± standard deviation was 5.71% ± 0.02%, giving a coefficient of variation of 0.35%. Thus, the intrinsic measurement error of the reported Hb A₁c values is within the size of the points in the figure below. Preliminary experiments (not shown) using Co²⁺ instead of albumin washes to terminate ionophore-mediated Ca²⁺ fluxes showed that Co²⁺ alters the size and shape of the Hb A₁c peaks, suggesting that Ca²⁺ might crosslink Hb with some other cell or lysate component that coeluted with genuine glycosylated Hb A₁c. This precluded Ca²⁺ use when measuring Hb A₁c. Similar tests with variable Ca²⁺ concentrations showed no such effects.

Separation of the small subpopulation of high-Na⁺ RBCs: measurement of yield, Hb A₁c, and Na⁺ contents

Because segregation of RBCs according to their PMCA Vₘₐₓ involved cell separation across a density boundary after K⁺ permeabilization, it was important to distinguish any fraction of K⁺-permeabilized RBCs unable to
dehydrate to levels exceeding that boundary for reasons unrelated to their Ca\(^{2+}\) pump activity (ie, the low-density, high-Na\(^+\) RBCs).\(^{16}\) These low-density RBCs were found in blood from healthy subjects and in much higher proportions in blood from sickle cell anemia (SS) and thalassemia patients.\(^{16,31-33}\) Depending on whether the observed resistance to dehydration followed K\(^+\) permeabilization induced by valinomycin, or via Gardos channels\(^{34}\) following Ca\(^{2+}\) loads, they were named “valres” or “calres” cells, respectively. To separate calres cells, LK-washed RBCs were suspended at 10% Hct in LK medium containing 3 mM KCl and 50 \(\mu\)M CaCl\(_2\) and incubated at 37\(^\circ\)C. After temperature equilibration, the ionophore A23187 was added to a final suspension concentration of 10 \(\mu\)M. After 10 minutes to allow maximal dehydration, 1.4 mL aliquots of this suspension were layered over 0.5 mL of DEP oil in 2 mL microfuge tubes, spun for 1 minutes at 10000g, and the tiny fractions of calres RBCs on top of the oil harvested and pooled using LK medium. Yields of calres RBCs and their Hb A1c fraction were measured directly in this suspension as described above for PMCA-separated samples. For Na\(^+\)-K\(^+\) measurements the cells were washed 3 times in 10 volumes of isotonic MgCl\(_2\), lysed, and diluted in distilled water to the required optimal dilution to activate Ca\(^{2+}\) channels\(^{24}\) following Ca\(^{2+}\) loads, they were named “valres” or “calres” cells, respectively. To separate calres cells, LK-washed RBCs were suspended at 10% Hct in LK medium containing 3 mM KCl and 50 \(\mu\)M CaCl\(_2\) and incubated at 37\(^\circ\)C. After temperature equilibration, the ionophore A23187 was added to a final suspension concentration of 10 \(\mu\)M. After 10 minutes to allow maximal dehydration, 1.4 mL aliquots of this suspension were layered over 0.5 mL of DEP oil in 2 mL microfuge tubes, spun for 1 minutes at 10000g, and the tiny fractions of calres RBCs on top of the oil harvested and pooled using LK medium. Yields of calres RBCs and their Hb A1c fraction were measured directly in this suspension as described above for PMCA-separated samples. For Na\(^+\)-K\(^+\) measurements the cells were washed 3 times in 10 volumes of isotonic MgCl\(_2\), lysed, and diluted in distilled water to the required optimal reading range and measured by flame photometry (Corning model 455, Surrey, United Kingdom).

**Experimental protocol for the study of Pcat properties**

RBC dehydration was previously described as the main trigger of Pcat.\(^{17}\) Accordingly, the experimental design was first to uniformly dehydrate all RBCs of any given sample and then follow their pattern and rate of rehydration from the timed changes in the profile of migrating osmotic fragility curves. The LK( + Ca\(^{2+}\))-suspended RBCs (5% Hct) were K\(^-\)permeabilized by the addition of valinomycin or A23187 at time 0 and were maximally dehydrated in 10 minutes. Samples of 0.5 mL were taken at varying time intervals up to 22 hours and processed for Na\(^+\)-K\(^+\) contents, as described above, and for osmotic fragility. Variations of the basic protocol are described in “Results” and in each of the figures. Osmotic fragility was measured by the profile migration method, as previously described.\(^{19,34,35}\) The osmotic fragility curves represent the integrals of population distributions\(^{19}\) and are shown in standard format, plotting the percent hemolysis as a function of relative toxicity.

**Results**

**Relation between PMCA activity and RBC age**

Figure 1 shows the patterns of yield recovery for RBCs that had fully extruded their Ca\(^{2+}\) load (\(\delta\) less than 1.117, harvested from the top of the DEP oil), or had not yet done so (\(\delta\) more than 1.117, harvested from pellets), at each time interval after temperature activation of PMCA-mediated Ca\(^{2+}\) extrusion. The results are typical of 9 experiments with blood from 6 donors. The selected results (Figure 1) exemplify the 2 patterns observed, differing mainly in the rate of increase in the yield of light RBCs (filled symbols). This rate reflects differences in the fraction of fast-pumping RBCs in samples from different donors, with faster-climbing yields of light RBCs indicating a greater fraction of high-V\(_{\text{max}}\) cells. The time required for full Ca\(^{2+}\) extrusion varied from 2 to 4.5 minutes in these experiments. Even at the earliest sampling time (30 seconds), the fraction of cells with \(\delta\) less than 1.117 was never 0 but varied between 1.5% and 5%. With an initial Ca\(^{2+}\) load of about 500 \(\mu\)mol(340 g Hb)\(^{-1}\), the RBCs would require pumping rates of over 60 mmol(340 g Hb)\(^{-1}\)h\(^{-1}\) to fully extrude this load in less than 30 seconds. We see below that such high-pumping cells account for only part of the RBCs in these early light fractions.

The relation between PMCA activity and RBC age is shown in Figure 2. The percent Hb A1c is plotted as a function of the yields of light RBCs. The ordinate reports the mean yield of light cells at the lowest yields of dense RBCs, which is analyzed in the text. The dotted lines in panels B and C represent extrapolated Hb A1c values at vanishingly small yields, pointing to the values expected among the youngest RBC fractions (lowest Hb A1c) and oldest RBC fractions (calres cells, highest Hb A1c).
those with the weakest pumps, represented by the points approaching the lowest yields (δ more than 1.117) with the highest Hb Alc values, indicating their older age. The quasilinear trend of increasing Hb Alc values of dense cells with decreasing yields shows that the decline in PMCA activity is an approximate linear function of RBC age. The mirror image variation in Hb Alc with yield of the light RBCs (δ less than 1.117) demonstrates the reciprocal trend, a progressive increase in PMCA activity with increasing RBC youth.

The reversal of the downward trend in the Hb Alc values observed at the lowest yields of light RBCs (δ less than 1.117) was seen in all experiments, varying only in magnitude (compare Figure 2A-C). A “contaminant” presence of high-Na⁺ calres RBCs, unable to dehydrate to densities above 1.117 g/mL by K⁺ permeabilization, was an obvious candidate to explain the observed reversal, if those RBCs had high Hb Alc levels. Measurements in 14 blood samples from 7 different donors showed Hb Alc levels (mean ± SEM) of 8.19% ± 0.9% and yields of calres RBCs of 3.2% ± 1.1%. The mean Hb Alc in the unfracti nated controls was 5.77% ± 0.06%. The Na⁺ content of calres cells was measured in 5 samples from a single donor from blood drawn on different days; the Na⁺ content of the calres cells was (mean ± SEM) 51.6 ± 4.3 mmol(340 g Hb)⁻¹, whereas that of the unfracti nated cells was 10.0 ± 0.1. Thus, calres RBCs from healthy donors had the high Hb Alc and Na⁺ contents, which could explain the observed reversal as a variable mix of low and high Hb Alc cells. Together with the early observations, the present results show that senescent RBCs reverse their progressive densification process toward a low-density, low-K⁺, high-Na⁺ condition, requiring a change in their normal cation permeability properties.

The properties of Pcat

In preliminary experiments with the profile migration method, we confirmed many of the results of Crespo et al: the elevated permeability of Pcat to Na⁺ and K⁺ and the lack of effect of many transport inhibitors (ouabain, amiloride, bumetanide, furosemide, tetrodotoxin, chlorpromazine, trifluoperazine, oligomycin, mepaca rine, and tetracaine). We also found that quinine was partially inhibitory (50% inhibition at 1 mM), that clotrimazole and vanadate were slightly stimulatory, and that iodoacetic acid and N-ethyl maleimide (NEM), known inhibitors of a nonspecific, voltage-dependent channel of RBC membranes (NSVDC), had no effect. These results are not detailed here. We focus rather on results that provide additional new information about Pcat properties.

Figure 3A illustrates the screening test applied to expose Pcat using the profile migration protocol. A sudden Ca²⁺ load produced...
uniform dehydration of the RBCs, seen as a rapid, marked, and largely parallel left shift in the osmotic fragility curve; after 10 minutes, the RBCs have attained near-maximal dehydration. If they retained their normal Na\(^+\) permeability, they would remain in this dehydrated state indefinitely, as seen in Figure 3D, in which 120 mM external NaCl was isosmotically replaced by 240 mM sucrose. In high-Na\(^+\) media (Figure 3A), however, the osmotic fragility curves showed a slow and persistent right shift, eventually leading to the lysis of many RBCs. Unlike the largely uniform initial left shift, the right shifts occurred with progressively lower slopes of the osmotic fragility curves; thus, all the RBCs underwent rehydration, but their rehydration rates showed a continuous variation, with no sharp discontinuities. Thus, the extent of Pcat activation was variable among the RBCs, causing some to lyse in 1 to 3 hours and others after 20 hours (Figures 3A-C,E-G). The clear distinction between dehydration and rehydration processes shown here in the presence of SCN\(^-\) could not be seen so distinctly in the absence of SCN\(^-\), when dehydration, rate-limited by the Cl\(^-\) permeability, was much slower. However, the long-term rehydration pattern was identical with and without SCN\(^-\) (not shown), indicating that prolonged exposure to 10 mM SCN\(^-\) had no effect on the rehydration response.

To investigate whether the observed variations in Pcat were related to RBC age, Hb A1c was measured in RBCs exposed to the dehydration-rehydration protocol of Figure 3A before and after prolonged incubations (6 to 22 hours). If RBC age correlated with the rehydration rate, the Hb A1c levels of the unlysed cells recovered after prolonged incubations would have differed from the original mean level. In 4 experiments with blood from 3 donors, the differences between mean Hb A1c percents in controls and unlysed cells recovered at late times, after lysis of 25% to 42% of the cells, were 0, 0.1, 0, and -0.1, averaging 0.00% ± 0.04% (SEM). Thus, at least for most of the RBCs, age was not a significant determinant of the observed differences in hydration rate and hence of Pcat activation. These results do not exclude the possibility that small subpopulations of aged or young RBCs were among those lysing first due to a high Pcat response.

Comparison of Figure 3A and B shows that after similar dehydration the rate of rehydration was much faster when intracellular Ca\(^{2+}\) was increased. Rapid extraction of the RBCs’ Ca\(^{2+}\) load by addition of Na-EGTA in large excess over [Ca\(^{2+}\)], at t = 10 minutes, after the cells had dehydrated, markedly reduced the rate of rehydration in both high-Na\(^+\) (Figure 3B) and high-K\(^+\) media (Figure 3C) but did not inhibit it completely. Figure 3B also illustrates the response of RBCs dehydrated with valinomycin in the absence of Ca\(^{2+}\) loads. Thus, the minor level of Pcat activation observed after Ca\(^{2+}\) extraction cannot be attributed to a residual irreversible effect of a previous Ca\(^{2+}\) load. The similarity of dehydration-induced rehydration rates of Ca\(^{2+}\)-free cells in high-Na (Figure 3B) and high-K (Figure 3C) media suggests that the Pcat-mediated permeability for these 2 cations is similar. Microscopic observation on timed samples of profoundly dehydrated RBCs incubated in culture media at 37°C after ionophore and Ca\(^{2+}\) extraction showed slow and uneven rehydration, with many initially crenated RBCs regaining biconcave or spheroidal shapes.80

In all these experiments, rehydration occurred in the absence of contaminant white cells and platelets.

In the experiments of Figure 3E-H, the RBCs were suspended in 90K-SCN medium, which prevented their dehydration following Ca\(^{2+}\) loads (Figure 3E-G) or valinomycin treatment (Figure 3H). All Ca\(^{2+}\)-loaded RBCs showed a hydration response that was not significantly modified by charybdotoxin (Figure 3F) or ouabain (Figure 3G), indicating that inhibition of Gardos channels or Na pumps had minimal effects on this response. On the other hand, the valinomycin-treated RBCs showed a barely detectable hydration response (Figure 3H). The results in Figure 3E-H thus show that Pcat can be activated by increased [Ca\(^{2+}\)]\(_i\), alone, without dehydration, and that K\(^+\) permeabilization itself has no effect on Pcat. In the conditions of Figure 3A, rehydration started with the cells initially profoundly dehydrated. When cells dehydrate by the net loss of KCl, the concentration of Cl\(^-\) in the isotonic effluent (about 150 mM) is higher than in the cells (about 100 mM). The resulting dilution of the intracellular Cl\(^-\) concentration increases the [Cl\(^-\)]/[Cl\(^-\)]\(_o\) ratio, and the anion exchanger rapidly restores the equality [Cl\(^-\)]/[Cl\(^-\)]\(_o\) = [H\(^+\)]/[H\(^+\)]\(_o\), with consequent intracellular acidification.41,42 In addition, the increase in impermeant anion concentration in the dehydrated RBCs hyperpolarizes the cells.41,43 Therefore, rehydration in the conditions of Figure 3A starts with the cells dehydrated, acidified, and hyperpolarized. On the other hand, in the initial conditions of Figure 3E, cell volume, pH, and membrane potential were all at physiological levels at the start of hydration. Because hydration proceeded comparably in both conditions, Ca\(^{2+}\) activation of Pcat appears not to be much affected by dehydration, pH, or membrane potential, although subtle effects cannot be ruled out from these data.

The measured changes in mean RBC Na\(^+\) and K\(^+\) contents during the dehydration-rehydration protocol of Figure 3A are shown in Figure 4A. The RBC K\(^+\) content fell sharply in the first 10 minutes and leveled off by 90 minutes, whereas the rate of Na\(^+\) gain was fairly constant throughout rehydration. These are average values for the RBC population as a whole and do not reflect the

![Figure 4](http://www.bloodjournal.org/blood/issue-files/200708/15aug/0000001338.jpg)
heterogeneity of the response, clearly discernible from the changing slope of the right-shifting osmotic fragility curves during rehydration (Figure 3A). From the data in Figure 4A it is possible to obtain a rough estimate of the mean value of the Na\(^+\) permeability, \(P_{Na}\), through Pcat, an important parameter for comparison with other permeation pathways in the RBC. Computation of the mean \(P_{Na}\) through Pcat can be done by simulating the experimental conditions with a model of proven reliability and by searching for the \(P_{Na}\)-Pcat values that would fit best the experimental data in Figure 4A. The computations were performed with the Lew–Bookchin model,\(^{41,43}\) and the results are shown in Figure 4B. The \(P_{Na}\)-Pcat value required to generate the agreement apparent in the comparison of Figure 4A and B was 0.015 h\(^{-1}\).

Pcat activation. The range of \([Ca^{2+}]_{i}\) values over which Pcat activation increases in all the RBCs was far above physiological \([Ca^{2+}]_{i}\) levels; at the \([Ca^{2+}]_{i}\) levels of 50 \(\mu\)M used in most of the experiments reported here, the Pcat response was below its maximal \(Ca^{2+}\)-saturated value.

The cation selectivity of Pcat was investigated by isosmotic replacement of NaCl (Figure 6A, LK) with choline chloride (Figure 6B, choline) or with N-methyl-D-glucamine chloride (NMDG) (Figure 6C). Rehydration occurred in all conditions, indicating that choline and NMDG were permeable through Pcat. In 1 experiment we investigated the Mg\(^{2+}\) permeability through Pcat. After Ca\(^{2+}\) loading, the ionophore A23187 was extracted with albumin at 4°C and the RBCs resuspended in iso-osmotic MgCl\(_2\) with 10 mM NaSCN and 1 mM vanadate to inhibit the PMCA and retain the Ca\(^{2+}\) load. The results showed an almost identical rehydration pattern as that of Figure 6C, indicating that Pcat is permeable to Mg\(^{2+}\). These results suggest that Pcat has a wide ionophoric path in the open state with poor cation selectivity.

**Discussion**

The present results demonstrate that the activity of the plasma membrane Ca\(^{2+}\) pump declines monotonically with RBC age, that the subpopulation of high-Na\(\text{+}\) “calres” RBCs represents old RBCs near the end of their circulating life span, and that RBCs have a poorly selective cation permeability pathway, Pcat, activated primarily by elevated \([Ca^{2+}]_{i}\). These results have important implications for understanding the changes in RBCs during physiological senescence and also for the interpretation of the statistical distributions of some classic RBC indices widely used in clinical laboratory tests.

The mechanism of programmed RBC senescence has been the subject of intense research for a few decades.\(^{46}\) Experimental evidence strongly supports terminal macrophage recognition and removal of RBCs following well-characterized antigenic changes in outer surface protein domains.\(^{47,53}\) There is also abundant evidence of gradual age-related RBC changes, including reduced enzymatic activities;\(^{54-56}\) cross-linking of cytoskeletal components;\(^{57,58}\) cumulative oxidative damage;\(^{59,60}\) and loss of lipid asymmetry.\(^{61,62}\)

The generally increasing density of aging RBCs has been recognized,\(^{46,63,66}\) and density fractionation has been used extensively to separate age cohorts. Yet the mechanism of densification has seldom been addressed, and little is known about the composition of aging RBCs. The condition documented here for the high-Hb A1c calres RBCs is one of increased Na\(^+\) permeability, not the often-suggested terminal condition of profoundly dehydrated RBCs.\(^{66,67}\) The present

![Figure 5. Effect of Ca\(^{2+}\) on the magnitude of the rehydration response.](image-url)
findings of a monotonic decline in PMCA activity with RBC age and of the advanced age of calres RBCs suggest a working hypothesis for the mechanism of the homeostatic changes in aging RBCs: the decline in PMCA activity would drive the pump-leak Ca\(^{2+}\) balance toward higher [Ca\(^{2+}\)], levels, which when exceeding the threshold activation of Gardos channels would induce a gradual net loss of KCl and water, leading to progressive RBC densification. With PMCA \(V_{\text{max}}\) variations of, for example, 60 and 2 mmol/(340 g Hb)\(^{-1}\) h\(^{-1}\), estimated pump-leak steady-state [Ca\(^{2+}\)] levels may vary from about 15 to 80 nM in young and old RBCs, respectively, leading to low-level Gardos channel activation in aging RBCs.\(^{45,69}\) Toward the final days in the circulation, the progressively increased [Ca\(^{2+}\)], levels, together with RBC dehydra-
tion, would activate Pcat, allowing the influx of Na\(^+\) at a rate exceeding the balancing capacity of the Na pump, eventually leading to net NaCl gain in excess of KCl loss and osmotic cell swelling toward a low-density terminal state (similar to the density of reticulocytes).\(^{16}\)

Valres and calres RBCs, with elevated Na\(^+\) contents, were originally detected in density-fractionated samples of sickle cell anemia and normal blood, sharing the same light-density fraction as reticulocytes\(^{16}\) of 1.070 to 1.090 g/mL. This is well beyond the prelytic density range of 1.050 to 1.060 g/mL, suggesting that intravascular lysis may not play a significant role in the normal circulatory removal of RBCs. However, the actual density spread of circulating calres RBCs deserves further investigation.

Elevated [Ca\(^{2+}\)], was shown to be a much more powerful activator of Pcat than RBC dehydra-
tion, which, without elevated [Ca\(^{2+}\)], was only a weak stimulus of Pcat. Permeation of very large cations such as NMDG (Figure 6) suggests that Pcat presents a large-conductance, wide-ionophoric path in the open state. At uniformly high Ca\(^{2+}\) loads, Pcat was activated in all the RBCs of each blood sample but to different extents, with a unimodal pattern of variation. This variation was not age-related for the bulk of the RBC population, but the precision limits of the measurements did not exclude enhanced Pcat sensitivity in up to 5% to 6% of RBCs. Pcat activation was observed in the absence of contaminant white cells or platelets and is therefore strictly a RBC response. Although Pcat has all the attributes required to explain the density reversal process leading to calres-valres cell formation, there is no conclusive evidence for its activation in vivo. The conditions that activate Pcat experimentally, elevated [Ca\(^{2+}\)], and RBC dehydra-
tion, do occur in vivo to an extent that would appear sufficient to elicit a very slow net sodium gain and density reversal, with the oldest RBCs reaching a light terminal density similar to that of reticulocytes.

Finally, let us consider the implications of RBC densification with age to the statistical distribution of the hemoglobin concentration (HC) in normal RBCs. The distributions of Hb contents (CH) and volumes (V) of human RBCs have coefficients of variation of about 13%,\(^{34,70,71}\) If these 2 distributions were independent, the coefficient of variation (CV) of the HC, representing the ratio of Hb content to volume in each RBC, would be much larger than 13%. But the measured CV of the HC is only about 7%, indicating the biologic priority of uniformity of HC during RBC production. The measured CH distribution probably changes little after RBC release from the bone marrow, with perhaps minor Hb loss by exovesicula-
tion.\(^{12}\) On the other hand, the measured V distribution is a composite of its original distribution at birth and its progressive reduction as the RBCs become increasingly dense with age. Therefore, the V and HC distributions of day-1 RBC cohorts must be much narrower than those measured for the whole RBC population comprising about 120 different day cohorts. This conclusion is supported by early flow cytometric measurements of HC distributions on density-separated RBCs,\(^{34}\) in which the CV of each one of 8 density fractions was about 3% (Table 2, column 24). Because each fraction represented a roughly 15-day age cohort, this result suggests that the coefficient of variation of the HC distribution of RBCs released from the bone marrow must be much less than 3%, an amazing degree of uniformity for a process delivering 10 billion RBCs per hour to a healthy adult’s circulation.

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Authorship

Contribution: V.L.L. designed the experiments, analyzed the data, and drafted the manuscript; N.D. and Z.E. performed experiments and contributed to data analysis; A.M. and L.V. performed experiments; T.T. contributed to data analysis and manuscript revision;
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Effects of age-dependent membrane transport changes on the homeostasis of senescent human red blood cells

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