Stochastic Nature and Red Cell Population Distribution of the Sickling-induced Ca\(^{2+}\) Permeability

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Abstract

To explore basic properties of the sickling-induced cation permeability pathway, the Ca\(^{2+}\) component (P\(_{\text{sickle-Ca}^2}\)) was studied in density-fractionated sickle cell anemia (SS) discocytes through its effects on the activity of the cells’ Ca\(^{2+}\)-sensitive K\(^+\)-channels (K\(_{\text{Ca}}\)). The instant state of K\(_{\text{Ca}}\) channel activation was monitored during continuous or cyclic deoxygenation of the cells using a novel thiocyanate-dense-cell formation method. Each deoxy pulse caused a reversible, sustained P\(_{\text{sickle-Ca}^2}\) which activated K\(_{\text{Ca}}\) channels in only 10–45% of cells at physiological [Ca\(^{2+}\)]\(_{\text{o}}\) (“activated cells”). After removal of cells activated by each previous deoxy pulse, subsequent pulses generated similar activated cell fractions, indicating a random determination rather than the response of a specific vulnerable subpopulation. The fraction of activated cells rose monotonically with [Ca\(^{2+}\)]\(_{\text{o}}\) along a curve reflecting the cells’ distribution of P\(_{\text{sickle-Ca}^2}\), with values high enough in a small cell fraction to trigger near-maximal K\(_{\text{Ca}}\) channels. Consistent with the stochastic nature of P\(_{\text{sickle-Ca}^2}\), repeated deoxygenated-oxygenated pulsed led to progressive dense cell formation, whereas single long pulses caused one early density shift. Thus deoxygenation-induced Ca\(^{2+}\)-permeabilization in SS cells is a probabilistic event with large cumulative dehydrating potential. The possible molecular nature of P\(_{\text{sickle-Ca}^2}\) is discussed. (J. Clin. Invest. 1997; 99:2727–2735.) Key words: Ca\(^{2+}\)-activated K-channels • sickle cell anemia • red cells • volume control • Ca\(^{2+}\) permeability

Introduction

The prime trigger of sickle cell dehydration is the reversible formation of a permeability pathway, P\(_{\text{sickle}}\). The pathway is somehow activated or generated by interaction of deoxygenated (deoxy)-Hb\(^S\) polymers with red cell membranes, but its nature and precise properties are unknown. P\(_{\text{sickle}}\) is poorly selective for mono and divalent cations (1–6), and is partially inhabitable by stilbene derivatives (7). The magnitude of its dissipative fluxes is apparently similar, at least for divalent cations, in sickle cell anemia (SS) reticulocytes and mature cells of all densities despite large differences in Hb S concentration, extent of polymerization, and sickling morphology (8). The key link between P\(_{\text{sickle}}\) and cell dehydration is calcium. During deoxygenation and sickling of SS red cells, the increase in cell membrane Ca\(^{2+}\) permeability alters the Ca\(^{2+}\)-pump-leak steady state, transiently raising [Ca\(^{2+}\)], to levels presumably sufficient to activate Ca\(^{2+}\)-sensitive K-channels (K\(_{\text{Ca}}\)) in some SS cells and trigger their dehydration (9, 10). In vivo, the sickling-induced increase in mean Ca\(^{2+}\) permeability is documented by the accumulation of cell calcium in endocytic plasma membrane vesicles, mediated by their inwardly directed Ca\(^{2+}\) pumps (11). But in the absence of information about possible heterogeneity of the sickling-induced Ca\(^{2+}\) permeability (P\(_{\text{sickle-Ca}^2}\)), or the effects of elevated [Ca\(^{2+}\)], on the activity of K\(_{\text{Ca}}\) channels, it was not possible to assess the uniformity of cell responses to sickling events or to predict the extent of activation of the K\(_{\text{Ca}}\) channels in the sickled state.

We have attempted here to address some fundamental questions on the properties and possible nature of P\(_{\text{sickle}}\)-Can K\(_{\text{Ca}}\) channel activation be directly demonstrated during a single sickling pulse? If so, are K\(_{\text{Ca}}\) channels activated in all the cells or only in a fraction of cells in each sickling pulse? If only in a fraction, does this response indicate the existence of a vulnerable subpopulation of cells or a probabilistic event randomly affecting different groups of cells in each deoxygenation pulse? Does the response arise from cell to cell differences in P\(_{\text{sickle-Ca}^2}\) or in other factors controlling [Ca\(^{2+}\)]? If P\(_{\text{sickle-Ca}^2}\) is not uniform, how is it distributed? Is P\(_{\text{sickle-Ca}^2}\) sustained for the duration of the sickling pulse, for part of it, or only during oxy-deoxy/deoxy-oxy transitions? To answer these questions we applied an experimental approach originally developed to explore the uniformity of ionophore-induced Ca\(^{2+}\) distributions in normal red cells (12). The method allows detection and separation of subpopulations of red cells with active or inactive K\(_{\text{Ca}}\) channels, with minimal alteration in the Ca\(^{2+}\) distribution which existed within the cells at the time of sampling.

Methods

Experimental design. The general design was to suspend the selected density fraction of SS cells in their own plasma or other specific media, expose the suspension to alternate deoxy-oxygenated (oxy) pulses in a tonometer, sample cells from oxy and deoxy states, and process the samples in conditions designed to preserve both their original

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1. Abbreviations used in this paper: deoxy, deoxygenated; Hb, hemoglobin; Hct, hematocrit; ISC, irreversibly sickled cell; K\(_{\text{Ca}}\) channel, Ca\(^{2+}\)-sensitive K-channel; oxy, oxygenated; SCN, thiocyanate; SS, sickle cell anemia, or homozygous hemoglobin S disease.
Ca\(^{2+}\) content and state of K\(_c\) channel activation, while segregating the cells with active or inactive K\(_c\) channels. The SS cell fraction best suited for the present experiments was the normal density fraction, mostly discocytes, because (a) it is light enough to allow cell dehydration to be detected as density shifts; (b) K\(_c\) channel-mediated dehydration could be studied with minimal interference from the KCl cotransport-mediated activity prevalent in the reticulocyte fraction; and (c) it is the most homogeneous and abundant SS cell fraction.

The design of sample processing was based on earlier studies (12) showing that (a) the diffusional permeability of thiocyanate (SCN) anions was at least 20-fold that of Cl anions; therefore K-permeabilized cells could dehydrate rapidly in low-K SCN media without rate-limiting effects of the anion; (b) the Q\(_0\) of the plasma membrane Ca\(^{2+}\) pump increased sharply from \(\sim 4-4.5\) at temperatures between 5 and 37°C, to \(\sim 7-9\) between 0 and 5°C, so that low temperature could inhibit the Ca-pump and prevent loss of cell Ca\(^{2+}\) in Ca\(^{2+}\)-free media; and (c) the temperature sensitivity of K\(_c\) channel activation remained low between 0 and 37°C; therefore cells with active K\(_c\) channels, suspended in low-K SCN media at 0-5°C, would dehydrate to a level similar to the densest sickle cells (\(\delta > 1.118\)) in 10–30 min. These properties permitted detection of SS cells with active K\(_c\) channels by facilitating their selective shrinkage in conditions which minimized post sampling changes in their Ca\(^{2+}\) content and state of K\(_c\) channel activation.

Based on this background information, aliquots of the cell suspension were transferred from the tonometer at specified times into a large volume of an ice-cold low-K SCN solution containing sufficient Ca chelator EGTA to ensure its excess over [Ca\(^{2+}\)], in the sample. This preserved the Ca\(^{2+}\) content and K\(_c\) channel activation state of the cells at the time of sampling by immediately preventing further passive gain or active loss of Ca\(^{2+}\) (see confirmatory evidence in Results). In the ice bath, cells with active K\(_c\) channels would shrink and could easily be separated from those with inactive channels by density fractionation. Even if the [Ca\(^{2+}\)] in some of the sampled cells had been just high enough for initial K\(_c\) channel activation, as the cells started shrinking in the SCN medium their [Ca\(^{2+}\)] would increase and accelerate further shrinkage. This sequence is favored by the minute Ca\(^{2+}\) binding capacity of cytoplasmic Ca\(^{2+}\) buffers in the submicromolar [Ca\(^{2+}\)] range (13–15). Thus even those cells with minimally active K\(_c\) channels in the original incubation conditions should be detected by this method. Sampling into SCN media per se promotes cell dehydration by its immediate binding to cell hemoglobin with the consequent increase in impermeant anion concentration, loss of cell Cl, and loss of water (16, 17). This effect is relatively small with the consequent increase in impermeant anion concentration, loss of cell Cl, and loss of water (16, 17). This effect is relatively small with the consequent increase in impermeant anion concentration, loss of cell Cl, and loss of water (16, 17). This effect is relatively small with the consequent increase in impermeant anion concentration, loss of cell Cl, and loss of water (16, 17). This effect is relatively small with the consequent increase in impermeant anion concentration, loss of cell Cl, and loss of water (16, 17). This effect is relatively small with the consequent increase in impermeant anion concentration, loss of cell Cl, and loss of water (16, 17). This effect is relatively small with the consequent increase in impermeant anion concentration, loss of cell Cl, and loss of water (16, 17). This effect is relatively small with the consequent increase in impermeant anion concentration, loss of cell Cl, and loss of water (16, 17). This effect is relatively small with the consequent increase in impermeant anion concentration, loss of cell Cl, and loss of water (16, 17). This effect is relatively small with the consequent increase in impermeant anion concentration, loss of cell Cl, and loss of water (16, 17). This effect is relatively small with the consequent increase in impermeant anion concentration, loss of cell Cl, and loss of water (16, 17). This effect is relatively small with the consequent increase in impermeant anion concentration, loss of cell Cl, and loss of water (16, 17). This effect is relatively small with the consequent increase in impermeant anion concentration, loss of cell Cl, and loss of water (16, 17). This effect is relatively small with the consequent increase in impermeant anion concentration, loss of cell Cl, and loss of water (16, 17). This effect is relatively small with the consequent increase in impermeant anion concentration, loss of cell Cl, and loss of water (16, 17). This effect is relatively small with the consequent increase in impermeant anion concentration, loss of cell Cl, and loss of water (16, 17). This effect is relatively small with the consequent increase in impermeant anion concentration, loss of cell Cl, and loss of water (16, 17). This effect is relatively small with the consequent increase in impermeant anion concentration, loss of cell Cl, and loss of water (16, 17). This effect is relatively small with the consequent increase in impermeant anion concentration, loss of cell Cl, and loss of water (16, 17). This effect is relatively small with the consequent increase in impermeant anion concentration, loss of cell Cl, and loss of water (16, 17). This effect is relatively small with the consequent increase in impermeant anion concentration, loss of cell Cl, and loss of water (16, 17).

**Experimental procedure.** Heparinized venous blood was obtained from sickle cell anemia patients and normal controls after informed consent. The red cells were washed three times with solution A containing (mM): 135 NaCl, 5 KCl, 10 Na-phosphate buffer (pH 7.4), 11 glucose, osmolality = 291 mosmol/liter. Theuffy coat of white blood cells and the top layer of light, reticulocyte-rich red cells were removed with each wash. Preliminary tests indicated that this procedure eliminated most cells with \(\delta < 1.095\). The remaining cells were suspended at about 10% hematocrit (Hct) in solution A and layered over arabinogalactan media of relative density 1.106, prepared as described previously (18). After centrifugation for 15 min at 17,000 rpm in a Sorvall RC2B (DuPont Company, Wilmington, DE) with a fixed angle rotor, the dense cell pellet (containing mostly irreversibly sickled cells) was discarded. The remaining cells (initially \(\delta \sim 1.106\), mostly discocytes, were washed twice more in solution A containing, in addition, 50 \(\mu\)M EGTA, and then either used immediately or kept overnight at 4°C, supplemented with 3 mM adenine and 10 mM inosine. After two additional washes (for cells stored overnight), the cells were suspended at between 8 and 40% Hct in plasma or in one of the following media: solution B (mM): NaCl, 135; KCl, 5; Hepes-Na, pH 7.4 at 37°C, 10; solution C: NaSCN, 135; KCl, 5; Hepes-Na, pH 7.40 at 37°C, 10. When indicated in figure legends, these solutions were supplemented with 3 mM adenine, 10 mM inosine, 10 mM glucose, and EGTA or CaCl\(_2\). The cells suspensions were equilibrated in a tonometer (model 237; Instrumentation Laboratory, Inc., Lexington, MA) at 37°C with either air or argon saturated with water vapor at 37°C (for oxy and deoxy conditions, respectively). When the cells were suspended in autologous plasma, the gases also contained 5.6% CO\(_2\). The suspension pH was monitored using a PHM 71 MK2 meter (Radiometer, Copenhagen, Denmark) and a glass microcapillary electrode unit, type E5021, and kept within the range of 7.36–7.48 by addition, when required, of small volumes of 0.15 M HCl or NaHCO\(_3\).

At the times shown in the figures, 0.1-ml samples of the cell suspension were delivered to microfuge tubes containing 0.9 ml of ice-cold SCN medium containing (mM): 138 NaSCN, 2 KCl, 10 Hepes-Na, pH 7.4, and 1 Na-EGTA. After 10–30 min of incubation in the ice bath, 0.3–0.4 ml of arabinogalactan medium, \(\delta = 1.118\), was gently delivered underneath the sample, using a syringe and long needle, and the tube was spun in an Eppendorf microfuge for 5 min (12,000 rpm) at room temperature. Any departures from these procedures are specifically noted in the figure legends. Hemoglobin concentrations were measured after conversion to cyanmethemoglobin using Drabkin’s solution. The fraction of dense cells formed in all conditions was calculated from the ratio of hemoglobin content in dense cell pellets to total hemoglobin in the original samples, corrected for any hemolysis by subtracting the supernatant hemoglobin concentrations of original samples (usually < 2%).

Loading of red cells with the Ca\(^{2+}\) chelator benz2 and measurements of their Ca\(^{2+}\)(6Ca) uptake were performed as previously described (9, 14, 19).

To measure \(^{86}\)Rb influx, the tracer was added to the cells in the tonometer. The cell \(^{86}\)Rb content was measured before and after SCN-density separation, and expressed in mmol per 340 g of Hb, using the specific activity of the tracer relative to the K concentration in the medium in which the cells were suspended, as described (18).

**Results**

**Effects of deoxygenation and calcium on generation of SCN-dense cells.** The experiment of Fig. 1A examined the fraction of SS discocytes which become dense when they are suspended in autologous plasma, exposed to oxy-deoxy periods shown in the figure, and sampled at 10-min intervals into cold SCN medium (see Methods). We see that these SCN-dense cells were formed only from samplings during deoxygenation periods, that their proportion increased during the first 20–30 min of deoxygenation, and that their formation was fully prevented by either reoxygenation or by addition of excess EGTA before sampling the deoxy genated cells. In the experiment of Fig. 1B, cells from the same batch were suspended in a Ca\(^{2+}\)-free plasma-like Cl-buffer. Again, neither deoxygenation in the absence of Ca\(^{2+}\) nor presence of Ca\(^{2+}\) in oxy states generated SCN-dense cells, but addition of Ca\(^{2+}\) to deoxy-cell suspension states promptly triggered the formation of SCN-dense cells. The results in Fig. 1 thus indicate that SCN-dense cell formation occurs only with cells sampled in the deoxygenated state, is strictly dependent on the presence of external Ca\(^{2+}\), and is abolished on reoxygenation. Furthermore, P\(_{\text{K,sickle-Ca}}\) persisted throughout deoxygenation, not just during oxy-deoxy transitions. The strict Ca\(^{2+}\) dependence of SCN-dense
cell formation shows that neither the well-characterized dehydrating effects of SCN (16) nor any residual K:Cl cotransport activity in the discocytes was sufficient to generate dehydration to the levels determined by this method (δ > 1.118).

The continuing increase in SCN-dense cells during the first 30 min of deoxygenation in the presence of [Ca$^{2+}$], (Fig. 1, A and B) raised the question of whether all the cells would eventually be similarly activated. Fig. 2 A shows that with a single long deoxy pulse, ~20% SCN-dense cells were generated within 20 min of deoxygenation, with no further increase. Fig. 2 B shows that each of two consecutive deoxy pulses (separated by an oxy period) generated ~40% SCN-dense cells within 30 min, with no further increase during the rest of each deoxy pulse. In other similar experiments where SS discocytes were suspended either in plasma or in plasmalike Cl-buffers, the plateau fractions of SCN-dense cells were between 10 and 46%, with similar consistency in consecutive pulses on the same suspension. Thus, with physiological [Ca$^{2+}$]o (1.0–1.2...
mM), stable fractions of SCN-dense cells were generated within the first 20–30 min of each deoxy pulse, with plateau levels (<50%) which differed between experiments but were similar for consecutive deoxy pulses with the same cell suspension.

Next, we considered why only a minority of SS discocytes became SCN-dense during each deoxy pulse, and why the plateau levels occurred with consecutive pulses. Were these cells a particularly vulnerable subpopulation, with the same cells affected with each pulse, or was each pulse generating different SCN-dense cells at random, with a similar probability? To address this question, we removed the cells that had become dense during one deoxy pulse and then tested whether subsequent pulses would generate additional SCN-dense cells (Fig. 3). Separation of dense cells from sequential deoxy pulses required suspension of the cells in the tonometer in the SCN buffer. It is important to point out that preliminary experiments under these conditions (not shown) demonstrated that SCN-dense cells were generated from SS discocytes only on deoxygenation in the presence of Ca²⁺, and that normal Hb red cells exposed to the same experimental maneuvers did not form SCN-dense cells, just as when the cells in the tonometer were suspended in plasma or Cl media.

We see that, despite incomplete removal of dense cells from previous pulses, successive sickling pulses recruited new cohorts of dense cells, each similar in size to the initial cohort (Fig. 3). Thus, most of the SS discocytes, when sickled in the presence of Ca²⁺, could become dense in SCN media. The similar proportion of new SCN-dense cells in successive pulses indicates that their generation is a random probabilistic event for each cell each time it is deoxygenated. A negative response to an earlier deoxy pulse does not influence its probability of becoming SCN-dense in a subsequent pulse, with the exception of F-cells. The percentage of Hb F in SCN-dense cells was 12–30% less than that in the original sample, reflecting their expected lower probability of becoming SCN-dense than non-F discocytes.

What determines these probabilities? The clear Ca²⁺ dependence of SCN-dense cell formation points to factors controlling the K⁺ channel activity of each cell, including [Ca²⁺]o (as determined by P₄₅₄Ca and Ca²⁺ pump activity), the number of K⁺ channels, and their Ca²⁺-sensitivity. The similar fraction of SCN-dense cells formed in successive pulses (Fig. 3), together with the knowledge that those cells have varied levels of K⁺ channel activation in the tonometer (see Methods), suggests that cell-to-cell differences in the number and Ca²⁺-sensitivity of the K⁺ channels have little or no influence on the fraction of SCN-dense cells. The pump-leak steady state [Ca²⁺]o level in each deoxy cell reflects the balance between P₄₅₄Ca and Ca²⁺ pump-mediated fluxes. Earlier studies showed that P₄₅₄Ca elicits the elevated [Ca²⁺]o of deoxy-SS cells. Therefore, the main stochastic event determining the fraction of SCN-dense cells with each deoxy pulse must be the extent of the cation-permeabilizing polymer–membrane interaction, which determines P₄₅₄Ca. According to this analysis, each time a cohort of SS cells is deoxygenated, their P₄₅₄Ca varies randomly, following a certain probability distribution among most of the cells.

The distribution of P₄₅₄Ca could be explored further by varying [Ca²⁺]o levels, to test whether more cells become SCN-dense as [Ca²⁺]o is increased. For low [Ca²⁺]o, those cells with the highest P₄₅₄Ca would gain enough Ca²⁺ for K⁺ channel activation, whereas at sufficiently high [Ca²⁺]o, even those cells with low P₄₅₄Ca could become SCN-dense. We therefore explored the increase in SCN-dense cells over a wide range of increasing [Ca²⁺]o.

Two similar experiments (Fig. 4, A and C) show that increasing [Ca²⁺]o generated progressively higher plateau fractions of SCN-dense cells. These plateaus are plotted as a function of [Ca²⁺]o in Fig. 4, B and D, and show a steep, nearly linear dependence on [Ca²⁺]o at [Ca²⁺]o < 5 mM, followed by a leveling off at [Ca²⁺]o > 5 mM. Thus each sickle pulse appears to generate a wide range of P₄₅₄Ca among the SS discocytes, with a continuous distribution and a slope which rises most steeply over physiological [Ca²⁺]o values (~1.0–1.2 mM). At those physiological [Ca²⁺]o levels, the P₄₅₄Ca of most SS discocytes (55–90% in different experiments) were too low to activate K⁺ channels. But at least 70% of SS discocytes exhibited some increased Ca²⁺ permeability, since they became SCN-dense at high [Ca²⁺]o.

Test for saturation of Ca²⁺ influx through P₄₅₄Ca. The leveling off of SCN-dense cells at [Ca²⁺]o > 5 mM (Fig. 4) could reflect either little further increase in P₄₅₄Ca in 20–30% of cells (the tail end of the P₄₅₄Ca distribution) or saturation of Ca²⁺ influx through P₄₅₄Ca. To distinguish between these two possibilities, the mean initial Ca²⁺ influx into chelator-containing SS discocytes was measured as a function of [Ca²⁺]o in oxy and deoxy conditions (Fig. 5). Since Ca²⁺ is distributed unevenly among the cells because of their different P₄₅₄Ca, the interpretation of Ca²⁺ influx results from measured mean cell calcium uptake is quite complex (see Discussion). But the question of whether the leveling off of the fraction of SCN-dense cells at [Ca²⁺]o > 5 mM results from saturation of Ca²⁺ uptake through P₄₅₄Ca can be addressed by observing how Ca²⁺ influx varies with [Ca²⁺]o at high [Ca²⁺]o. Fig. 5 confirms (20, 21) that deoxygenation causes a marked increase in P₄₅₄Ca of SS discocytes, and shows in addition that their Ca²⁺ influx continues to increase at [Ca²⁺]o > 5 mM, without saturating at 16 mM, the highest level tested here. Thus, the leveling off of SCN-dense cell formation at high [Ca²⁺]o (Fig. 4, B and D) ap-

Figure 3. Effect of removing the SCN-dense cells formed in one deoxy pulse on the fraction formed in the next pulse. SS discocytes were suspended at 30% Hct in solution C (with SCN anion) supplemented with adenine, inosine, and 1.6 mM CaCl₂. Samples were taken every 10 min to measure the fraction of cells that became dense (n = 1.118). After 30 min of deoxy, the whole suspension was layered on arabinoxylaminomaltose (n = 1.118) in several microfuge tubes, and after a 5-min spin at 12,000 rpm, the cells with n < 1.118 were washed and resuspended in solution C at 30% Hct in the tonometer, and the deoxygenation cycle was repeated. The third deoxy pulse was for 40 min. Similar results were obtained in another experiment.
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Apparently results from the \(P_{\text{sickle-Ca}}\) distribution among the cells, rather than from limiting Ca\(^{2+}\) influx through \(P_{\text{sickle-Ca}}\). The results in Fig. 4 reflect the distribution of \(P_{\text{sickle-Ca}}\) values generated by full deoxygenation of SS discocytes, and suggest that a substantial fraction, albeit a minority, of SS discocytes deoxygenated at physiological (and even lower) levels of \([\text{Ca}^{2+}]_{o}\) develop sufficiently high \(P_{\text{sickle-Ca}}\) (and therefore \([\text{Ca}^{2+}]_{i}\) elevations) to activate their \(K_{\text{Ca}}\) channels, as demonstrated by the generation of SCN-dense cells. Sickling-induced Ca\(^{2+}\) influx is therefore extremely heterogeneous among the cells. If the mean Ca\(^{2+}\) influx values reported in Fig. 5 are attributed to the small fractions of cells that become SCN-dense at low or physiological \([\text{Ca}^{2+}]_{o}\), fluxes which have been shown to generate dense normal-Hb cells within a few hours of Ca\(^{2+}\) permeabilization with the ionophore A23187 (10). Thus the results in Figs. 4–5 demonstrate that each sickling episode randomly generates small populations of cells with high Ca\(^{2+}\) permeabilities, and sufficient Ca\(^{2+}\) influx to raise \([\text{Ca}^{2+}]_{i}\) to levels producing substantial \(K_{\text{Ca}}\) channel activation. In some cells the activation of \(K_{\text{Ca}}\) channels may be near-maximal, so that cell dehydration becomes limited by the diffusional Cl permeability (\(P_{\text{Cl}}\)).

The mean Ca\(^{2+}\) influx was also measured in dense SS cells fractions (rich in irreversibly sickled cells [ISC]) with the incorporated Ca\(^{2+}\)-chelator benZ. The results of two such experiments were similar, and one is included in Fig. 5. No significant differences in mean Ca\(^{2+}\) uptakes were seen between fully deoxygenated ISC cells, and discocytes, suggesting that the morphological differences in polymer structure and concentration between these two cell fractions (22, 23) may not be relevant determinants of \(P_{\text{sickle-Ca}}\).

Sickling-induced dehydration of SS discocytes in Cl media, without SCN processing. It was important to ascertain whether the properties of \(P_{\text{sickle-Ca}}\) inferred from analysis of SCN-dense cell formation directly reflected those of the original cell suspensions in the tonometer. The observed Ca\(^{2+}\) dependence, distribution, and stochastic nature of \(P_{\text{sickle-Ca}}\) predict that cell
deoxy-oxy pulses produced cumulative density shifts, as expected from high \( P_{\text{sickle-Ca}} \) generation in cells newly activated with each pulse. These results therefore demonstrate that the effects predicted from analysis of the properties of SCN-dense cell formation are indeed obtained for SS cells suspended in physiological saline and exposed to deoxy-oxy pulses in the tonometer.

**Effect of deoxygenation on \(^{86}\text{Rb} \) influx in SCN-dense cells.** Another experimental approach was used to test more directly for the presence of a small fraction of deoxy SS cells with near-maximal \( K_{\text{Ca}} \) channel activation, with rates of dehydration limited by \( P_{\text{Cl}} \). If sickling in \( \text{Ca}^{2+} \)-containing plasma or plasma-like Cl media increased their K permeability so that \( P_{\text{K}} \) substantially exceeded \( P_{\text{Cl}} \), those cells should be hyperpolarized. This hyperpolarization could be further increased by reducing the external K levels below normal. \(^{86}\text{Rb} \) tracer added to this low K medium should accumulate preferentially in the most K-permeabilized, highly hyperpolarized cells, due to the strong inward driving force on the \(^{86}\text{Rb} \) influx (described as a countertransport-like effect in such K-permeabilized cells [24]). Thus a \( \text{Ca}^{2+} \)-dependent, sickling-induced increase in \(^{86}\text{Rb} \)-tracer influx might be evident only in the cells which become dense after SCN treatment. Even though a large fraction of the combined (\( K_{\text{4.4}} \)) content of these cells may be lost during SCN-induced shrinkage, the \(^{86}\text{Rb} \)/K specific activity would be highest in these cells; and since \(^{86}\text{Rb} \) influx through \( P_{\text{sickle}} \) is minimized by the low external K concentration, the higher countertransport-driven influx might exceed the “noise level” of the \(^{86}\text{Rb} \)-influx into all other nonhyperpolarized cells.

Indeed, such an experiment (Fig. 7) showed that a sickling-induced increase in \(^{86}\text{Rb} \) uptake was detectable only in the presence of \([\text{Ca}^{2+}]_o\), and limited to the cells which subsequently became dense in SCN media. Thus a fraction of the cells that became dense in SCN must have had their \( K_{\text{Ca}} \) channels almost fully activated during incubation in a Cl medium in the tonometer, when they were sickled in the presence of \([\text{Ca}^{2+}]_o\).

**Discussion**

To characterize the intermediate steps leading to SS discocyte dehydration in vivo, we investigated their volume responses to deoxygenation. The results indicated that (a) each deoxy pulse caused a reversible increase in \( \text{Ca}^{2+} \) permeability, \( P_{\text{sickle-Ca}} \), which lasted for the duration of the pulse; (b) the presence of external \( \text{Ca}^{2+} \) caused \( K_{\text{Ca}} \) channel activation in a fraction of cells (activated cells); (c) the fraction of activated cells increased within \( \sim 30 \) min to plateau values between 10 and 45\% in different experiments at physiological external \( \text{Ca}^{2+} \) concentrations; (d) the plateau was constant for at least 3 h, and of similar value in consecutive pulses; (e) the fraction of activated cells was determined randomly in each pulse and did not reflect the response of a particularly vulnerable subpopulation of cells; (f) the plateau fraction of activated cells increased monotonically with \([\text{Ca}^{2+}]_o\), along a curve which reflected the distribution of \( P_{\text{sickle-Ca}} \) in the cell population; this distribution indicated that the \( P_{\text{sickle-Ca}} \) generated in each deoxy pulse varied widely among the cells, with the majority of cells having a value too low for \( K_{\text{Ca}} \) channel activation at physiological \([\text{Ca}^{2+}]_o\); (g) as expected from the stochastic nature of \( P_{\text{sickle-Ca}} \), frequent deoxy-oxy pulsing was more efficient in generating larger fractions of dense cells than single deoxygenation pulses; and (h) in the fraction of cells with the highest \( P_{\text{sickle-Ca}} \)
values, $K_{cs}$ channel activation occurred with evidence of marked hyperpolarization, and was thus near maximal, a condition in which cell dehydration becomes rate-limited by $P_C$, which is fairly uniform in the cell population (25).

We consider now the relevance of these results to the process of SS cell dehydration in vivo. In this investigation, the new methods and experimental designs, together with the selection of SS discocytes, allowed a focused study of the dehydration process mediated by $K_{cs}$ channels, with no interference from other mechanisms which may contribute to dehydration in vivo, that is, the K:Cl cotransport and the Na pump. In SS reticulocytes and young red cells the K:Cl cotransport accelerates dehydration according to its level of expression and activity (fast track cells) (21). On the other hand, $K_{cs}$ channel-mediated dehydration operates in deoxygenated SS cells of all ages and densities. Therefore, although this study...
centered on SS discocytes, the conclusions apply to $K_{\text{Ca}}$ channel–mediated dehydration in all SS cells.

An important new finding in this study is that deoxygenation-induced $\text{Ca}^{2+}$ permeabilization of SS cells is a probabilistic event, with a distribution that includes a fraction of cells with $P_{\text{sickle-Ca}}$ high enough for near-maximal $K_{\text{Ca}}$ channel activation in each deoxygen pulse. In a population of deoxygenated SS cells, the probability is expressed as a distribution of $P_{\text{sickle-Ca}}$ values. For an individual circulating SS cell, however, the probabilistic nature of $P_{\text{sickle-Ca}}$ will be reflected in its different $P_{\text{sickle-Ca}}$ value each time it becomes deoxygenated, with higher probabilities for low values. Although the dehydrating potential of each deoxy episode may be small, the cumulative effect of multiple episodes should lead to substantial cell dehydration, as indicated by the results in Fig. 6. However, except for cells trapped in stagnant capillary beds, the fraction of polymer-containing cells in the circulation at any time would be much smaller than that in the fully deoxygenated population in which the $P_{\text{sickle-Ca}}$ distribution was measured here. Therefore, the true dynamic distribution of $P_{\text{sickle-Ca}}$ among circulating SS cells would be a fraction of that inferred from the plateau distribution in the tonometer (Fig. 5). The dehydration profiles obtained within 3–4 h in the tonometer (Fig. 6) may thus take much longer to develop in vivo, although the mechanism would be similar for those cells whose dehydration is mediated primarily by $K_{\text{Ca}}$ channels.

The possible nature of $P_{\text{sickle}}$. With the limited state of present knowledge, these considerations must be highly speculative, but they may help formulate questions and working hypotheses for future experiments. Although it has yet to be established, we will assume for now that $P_{\text{sickle}}$ reflects the general properties of $P_{\text{sickle}}$. The conditions in which $P_{\text{sickle}}$ is induced suggest that it somehow arises from interactions between deoxy-Hb S polymers and membrane components, which either generate a new transport pathway or modify a preexisting one. These polymer–cell membrane interactions may involve known or unidentified contacts. Let us consider what we can explain with known interactions to assess the need of unknown alternatives. One clearly identified configuration for the interaction between polymers and cytosolic membrane domains without cytoskeletal barriers is the protruding polymer spicule; in this configuration fiber surfaces and growing spikes could interact directly with the cytosolic aspect of freely diffusing membrane-embedded proteins (26).

Another documented interaction, between the red cell membrane and soluble hemoglobin, involves the NH$_4$ terminal of band 3 protein and the central cavity of the deoxy-Hb tetramer (27). F.A. Ferrone suggested that this restraining reaction might provide a site energetically favorable for nucleation (personal communication). If deoxy-Hb S polymers participate in this link, bundles of adjacent fibers might generate local areas of high density links with band 3 protein, forming oligomeric aggregates of the band 3 dimers, which could function as a large ion channel. The inhibitory effects of stilbenes on $P_{\text{sickle}}$ (7), though not specific, would be consistent with such a participation of band 3 protein in the structure of the $P_{\text{sickle}}$ pathway. In this context it is of interest to note that although $P_{\text{sickle}}$ is recognized as a poorly selective cation channel, its lack of selectivity may extend to anions as well. Its structure might then resemble that recently suggested for anion-channel–forming trout band 3 protein when expressed in *xenopus* oocytes (28).

With these possibilities in mind, we now ask what insights the present experiments may provide on the properties of $P_{\text{sickle}}$. Initiation of sickle polymer formation by nucleation is a random process with a skewed time-dependent distribution and a 50th order reaction mechanism (29). Once polymerization is initiated within a particular cell, progression to polymer solubility equilibrium ($C_{eq}$) with soluble deoxy-Hb S should be quite rapid. The kinetics of cell recruitment to the fully polymerized equilibrium condition (equivalent to the kinetics of sickling) in the present experiments could therefore be inferred from the time course with which the fraction of SCN-dense cells approached its plateau level (Figs. 1, 2, and 4, A and C). This kinetics was apparently somewhat variable, which could reflect (a) the intrinsic random distribution of nucleation times (Ferrone); (b) cell to cell differences in Hb concentrations of up to 4 g/dl within the discocyte density fraction used here; (c) the presence of fetal hemoglobin-containing cells (F-cells), likely to be either the slowest recruits observed here at high Ca$^{2+}$ levels (Fig. 4, A and C), or cells generating no polymer and no $P_{\text{sickle}}$, and (d) differences in deoxygenation rates in the tonometer, due to different hematocrits, total volumes, or minor pH differences. As previously observed with
the same tonometry system (30), extensive polymerization/sickling appears to have been attained within ~30 min in most conditions (see delays to plateau in Figs. 2, 4, A and C), evoking the participation of at least 60–70% of cells in SCN-dense cell formation at sufficiently high Ca²⁺. At equilibrium, as in the plateau phase of SCN-dense cell formation, all the permeabilizing contacts between polymers and the relevant membrane domains must have been established. If we assume that each effective contact generates a quantum of permeabilization, and therefore that cell to cell differences in P_{sickle} may be due simply to differences in the number of permeabilizing contacts, the results suggest that (a) the number of permeabilizing contacts established in each deoxy pulse is random and variable from pulse to pulse in each cell; (b) for the duration of the deoxy pulse, the permeabilizing contacts must either become established during the brief polymer formation phase and remain structurally stable, or they may result from reversible interactions, with different equilibrium levels in each cell during the formation phase and reversibility in either the formation of the stable polymer contacts, or in the concentration of available interactive polymer contacts (surfaces, ends, or spicules) which form in each pulse; for the concentration and accessibility of membrane targets, whether immobilized by cytoskeletal links or in free lateral diffusion, should not exhibit random variation between pulses.

Several of the speculative assumptions in this account are amenable to experimental testing, using systems such as cytoskeleton-free, Hb-S-containing vesicles derived from isolated spicules, known to contain the proposed reactive components of the permeabilizing contacts (26). Investigation of cation transport and structural changes in vesicles in oxy and deoxy conditions is currently in progress.

2. The kinetics of red cell deoxygenation and oxygenation in the IL tonometer in vitro must be less efficient than that of red cells squeezing through capillaries, where gas equilibration times and diffusion through unixed layers surrounding the cells are minimized. Thus the time required for polymer equilibration and maximal activation of P_{sickles} in vivo should be considerably shorter than in this in vitro system.

Acknowledgments

We wish to thank the Wellcome Trust (United Kingdom), The British Heart Foundation, and the National Institutes of Health (grants HL-28018 and HL-20985) for funds.

References