Characterization of Regulatory Volume Decrease in the THP-1 and HL-60 Human Myelocytic Cell Lines

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Exposure to hypotonic stress produces a transient increase in cell volume followed by a regulatory volume decrease (RVD) in both THP-1 and HL-60 cells. In contrast, cells exposed to hypotonic stress in a high K/low Na Hanks' solution not only failed to volume regulate, but displayed a secondary swelling. Thus, while an outward K gradient was required for RVD, the secondary swelling indicated that hypotonic stress increased permeability in the absence of a negative membrane potential. The K channel blocker quinine (1–4 mM) blocked RVD in both cell types. Gramicidin's ability to overcome the quinine block of RVD indicated that RVD is mediated by a quinine-sensitive cation transport mechanism that is independent of the swelling-induced anion transport mechanism. Barium (1–4 mM), another K channel blocker, slowed the rate of RVD, while 4-aminopyridine, charybdotoxin, tetroxalammonium chloride, tetrabutylammonium chloride, and gadolinium had no effect on RVD. Furthermore, RVD was not mediated by calcium-activated conductances, since it occurred normally in Ca-free medium, in medium containing cadmium, and in BAPTA-loaded cells. Gramicidin produced little or no volume change in isotonic medium, suggesting that basal C1 permeability of both THP-1 and HL-60 cells is low. However, swelling induced an anion efflux pathway that is permeable to both chloride and bromide, but is impermeable to methanesulfonate and glutamate. The anion channel blocker 3,5-diiodosalicylic acid (DISA) antagonized RVD in both cell types. In conclusion, RVD in THP-1 and HL-60 cells is mediated by independent anion and cation transport mechanisms that involve both a DISA-sensitive anion pathway and a quinine-inhibitable K efflux pathway, neither of which requires increases in intracellular calcium to be activated.

The presence of impermeable intracellular charged particles results in a tendency for all cells to swell. Animal cells counterbalance this tendency by extruding ions through either conductive pathways or coupled transporters (for review, Sarkadi and Parker, 1991). Volume regulation also occurs when cells, such as kidney epithelial cells, are exposed to anisotonic environments (Lewis and Donaldson, 1990). The pathways involved in homeostatic volume regulation have been extensively studied in vitro by placing cells in anisotonic medium. Exposure to hypotonic medium produces a rapid swelling followed by a volume decrease referred to as a regulatory volume decrease (RVD) in many cell types. In some cells, including erythrocytes, RVD is associated with an increase in the activity of electroneutral transporter(s) (Sarkadi and Parker, 1991), while in other cells, such as lymphocytes, K and Cl conductive pathways underlie RVD (Grinstein and Foskett, 1990).

While volume regulatory responses in lymphocytes and erythrocytes have been well characterized, only two studies have examined RVD in cells of the macrophage lineage (monocytes, macrophages, or promonocytic cells). Novak et al. (1988) demonstrated that, in rabbit alveolar macrophages, RVD depends on the K gradient and is due to a ouabain-insensitive loss of K and Cl, which precedes the swelling-induced changes in surface receptor numbers associated with inhibition of receptor-mediated endocytosis. In the promyelocytic cell line HL-60, RVD is associated with a change in actin polymerization, but the shift in actin polymerization is not required for the transduction of the volume regulatory signal (Hallows et al., 1992). While these two studies demonstrated that myeloid cells, like lymphocytes, exhibit RVD, the ionic mechanisms underlying RVD were not investigated.

It is possible that one or more of the K and Cl conductances, which have been characterized in primary cells of the monocytic-macrophage lineage or in macrophage-like cell lines using electrophysiological tech-

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niques (for review see Gallin, 1991), mediate RVD in these cells. As a first step towards exploring this possibility, this study investigated RVD in two human leukemic cell lines; HL-60 cells, which can be induced to differentiate into either macrophage-like cells or neutrophil-like cells, and THP-1 cells, monocytic-like cells that can be differentiated into mature macrophages (Auwertz, 1991). We demonstrate that while RVD in both these cell lines involves the activation of independent cation and anion permeability pathways, the calcium-activated conductances described in macrophages are unlikely to mediate RVD. Furthermore, the cation permeability pathway activated in these two cell lines has a different sensitivity to pharmacological agents than the n-type K channel that mediates RVD in lymphocytes (Grinstein and Foskett, 1990; Deutsch and Lee, 1988).

**MATERIALS AND METHODS**

**Cells**

HL-60 cells and THP-1 cells were obtained from American Type Culture Collection (Rockville, MD). HL-60 cells and THP-1 cells were maintained in RPMI-1640 supplemented with 10% fetal bovine serum (FBS; HyClone Laboratories, Logan, UT), penicillin (20 units/ml), streptomycin (20 ng/ml), and glutamine (0.3%). Cells were grown in a 5% CO2-air incubator at 37°C and split 3:1 twice a week. Cells were maintained for approximately 3 months, at which time new frozen stocks were thawed.

**Volume measurements**

Volume measurements were carried out electronically using a model ZM Coulter Counter equipped with a 100 μm diameter orifice. The counter was coupled to an IBM compatible personal computer equipped with a Series II Personal Computer Analyzer card (PCAII; Nucleus Inc., Oak Ridge, TN). The PCAII card sorted incoming digitized signals into channels binned to provide histograms in which the channel number was proportional to cell volume. Median cell volume was computed from histograms of approximately 10,000 cells. Polystyrene beads 9–15 μm in diameter (Coulter) in solutions with the same toxicity were used to calibrate the instrument.

Cells were resuspended at approximately 10^6 cells/ml in NaCl Hanks' solution containing 10 mM glucose and 0.5% bovine serum albumin (BSA; normal Hanks') and used for volume analysis within 30 minutes. Volume measurements were performed at room temperature (21–23°C) with control cell volume determined in normal Hanks'. RVD was assessed following dilution with 41% water (0.59x Hanks'). This degree of osmotic stress was chosen because it produced a significant swelling and regulatory response without affecting cell viability. Volume changes were measured over a 30-minute time period and are expressed relative to control cell volume. For later time points where volume changed more slowly, an averaged volume was determined from two to three separate measurements taken over 30 seconds.

**Cell viability**

Cell viability was assessed with a fluorescent microscope using ethidium bromide and acridine orange. Exposure to 41% hypotonic medium for 30 minutes produced no significant change in cell viability. Addition of quinine (4 mM) in either isotonic or hypotonic medium had no effect on the number of viable cells.

**Solutions**

Normal Hanks' referred to as Hanks' contained in mM: 150 NaCl, 1.2 MgCl2, 1.6 CaCl2, 4.6 KCl, 10 HEPES, 10 glucose, and 0.5% BSA, titrated with NaOH to pH 7.3. KCl Hanks' contained in mM: 150 KCl, 1.6 MgCl2, 1.2 CaCl2, 5 NaCl, 10 HEPES, 10 glucose, 0.5% BSA, titrated with KOH to pH 7.3. Low sodium Hanks' was made either with N-methylglucamine chloride (NMDGCl) or tetramethylammonium chloride (TMACl) to replace all but 4 mM NaCl. For studies in which anion selectivity was examined, 150 mM chloride was replaced by other anions. For example, K-methanesulfonate Hanks' contained in mM 150 Kmethanesulfonate, 4.6 KCl, 1.6 CaCl2, 1.2 MgCl2, 10 HEPES, 10 glucose, and 0.5% BSA.

**Calcium measurements**

Cells were loaded with fura in normal Hanks' containing 2.5 μM fura-2-AM and 0.2 mg/ml of pluronic acid with or without 15 μM BAPTA-AM for 1 hour at room temperature. Cells were centrifuged, and loading medium was replaced with normal Hanks'. Following a 15-minute incubation at room temperature fluorescence ratios (excitation 340 and 380; emission 510) were measured at room temperature using an SLM spectrofluorimeter. After obtaining baseline fluorescence measurements, 41% water was added (3 ml for final cuvette volume) and fluorescence changes were monitored for 5–10 minutes. Calibration was accomplished for each cuvette by adding 15 μl of 10% Triton-X 100 to the cuvette to obtain Fmax, followed by the addition of 50 μl of 0.75 M EGTA/TRIS, pH 8, to obtain Fmin. Complete hydrolysis of the fura-AM was confirmed by comparing the fluorescence ratio at 340/380 of lysed fura-loaded cells in high calcium to the fluorescence ratio of free fura recorded under the same conditions. Intracellular calcium ([Ca2+]i) was estimated as described by Grynkiewicz et al. (1985) using a Kd of 135 mM for fura measured in 100 mM KCl, pH 7.2, at 20°C.

**Chemicals**

Gramicidin, 4-acetamido-4'-isothiocyanostilbene-2,2'-disulfonic acid (SITS), 3,5-diiodosalicylic acid (DISA), ouabain, quinine, and tetrabutylammonium chloride (TBACl) were obtained from Sigma (St. Louis, MO), tetraethylammonium chloride from Aldrich (Milwaukee, WI), and charybdotoxin (CTX) from Latoxan (Rosans, France). Gramicidin and DISA were made up as concentrated stocks (100X) in dimethylsulfoxide (DMSO). CTX was made up in Hanks' containing 0.5% BSA and stored in glass vials. CTX's activity was confirmed by demonstrating that it blocked an ionomycin-stimulated K efflux from guinea pig epithelium (Dr. Pamela Gunter-Smith, personal communication). Fura-2-AM, pluronic acid, and BAPTA-AM were obtained from Molecular Probes (Eugene, OR) and made up as 1 mM stock solutions in DMSO.
VOLUME REGULATION IN MYELOCYTIC CELL LINES

A. 140 150
E 130 E 140
120.
110
100
a)
0
110
100
4 8 12 16 20 24 28 32
Time (min)
4 8 12 16 20 24 28 32
B. 140
Time (minutes)
130
10
E E 140
120
0
130
11 (D 120
C 2
110
100
10 L•o 100
0 4 12 16 20 24 28 32
Time (min)
90
90
0 4 8 12 16 20 24 28 32

Fig. 1. Time course of regulatory volume changes in (A) HL-60 cells and (B) THP-1 cells following addition of 41% water at time 0. For this figure and other figures, data are expressed as the percentage volume change compared to control cell volume determined at the beginning of the experiment. Averaged data from three separate runs are shown for each cell type.

Statistics

All data are expressed as mean ± standard error of mean.

RESULTS

RVD occurs in both THP-1 and HL-60 cells

HL-60 and THP-1 cell populations exhibit unimodal volume distributions under isosmotic and hypotonic conditions. Their mean peak volume under isosmotic conditions was 970 ± 4.1 fL and 580 ± 2.4 fL for THP-1 cells and HL-60 cells, respectively. As reported previously by Hallows et al. (1991), HL-60 cells exhibit a transient swelling followed by a decrease in volume when exposed to hypotonic solution (Fig. 1A). A similar RVD response is also present in the THP-1 cell line following exposure to 0.59× Hanks' (Fig. 1B). Peak swelling occurred between 2 and 4 minutes in both cell types, reaching 125–145% of control cell volume in 0.59× solutions. Cell volume returned to within 10% of control levels within 30 minutes at 21–23°C. RVD was characterized by an initial rapid decrease followed by a slower change in volume. The initial rate of RVD measured during the first 5–6 minutes was -3.31%/min ± 0.12 (n = 17) and - 5.48%/min ± 0.7 (n = 11) for THP-1 and HL-60 cells, respectively.

Role of cation permeability

Potassium channel blockers. Since K conductances have been implicated in RVD responses of other cells (Grinstein and Foskett, 1990; Okada and Hazama, 1989), the effects of several different K channel blockers were tested on HL-60 and THP-1 cells exposed to 0.59× Hanks'. In HL-60 cells 1 mM quinine slowed but did not completely block RVD; cell volume at 30 minutes was 120% ± 4.9 (n = 7) of initial control volume compared to a 30-minute volume of 107% ± 3.6 for control measurements in the absence of 1 mM quinine (Fig. 2A). Increasing the quinine concentration to 4 mM blocked RVD completely in HL-60 cells (peak volume vs. 30-minute volume was 142% ± 3.9 vs 139 ± 5.6, n = 3). In contrast, in THP-1 cells, 1 mM quinine completely blocked RVD (volume at 30 minutes was 131% ± 2 of initial control volume, n = 4; Fig. 2B).

Since in some cell types quinine has been reported to affect C1 conductances as well as K conductances (Gogeine and Capek, 1990), experiments were performed using both the cation pore former gramicidin and the K transporter valinomycin to determine whether quinine's block of RVD was due to a block of a cation permeability. Cell volume was monitored in hypotonically stressed cells bathed in medium that contained quinine to which gramicidin was added. Gramicidin, which forms channels permeable to K and Na, enables cells to volume regulate in the presence of a K channel blocker, as long as their anion permeability is unaffected by the K channel blocker. In 0.59× Hanks' containing quinine, addition of gramicidin to THP-1 cells produced a secondary swelling presumably due to the Na influx through gramicidin channels. In contrast, HL-60 cells exposed to gramicidin in 0.59× Hanks' plus
Gramicidin 1 - 6

A.

Fig. 3. Gramicidin induces RVD in the presence of 4 mM quinine. Time course of representative volume responses from (A) HL-60 cells in 0.59x NMDG Hanks' and (B) THP-1 cells in 0.59x TMA-Hanks'.

B.

Fig. 4. Effect of barium on RVD response of (A) HL-60 cells and (B) THP-1 cells. Time course of representative volume responses of cells in 0.59x NaCl Hanks' ( ), 0.59x NaCl Hanks' containing either 1 ( ) or 4 mM BaCl2 ( ).

Fig. 5. Effect of barium on RVD response of (A) HL-60 cells and (B) THP-1 cells. Time course of representative volume responses of cells in 0.59x NaCl Hanks' ( ), 0.59x NaCl Hanks' containing either 1 ( ) or 4 mM BaCl2 ( ).

TABLE 1. Ability of agents to block RVD

<table>
<thead>
<tr>
<th>Agent</th>
<th>Concentration</th>
<th>Block</th>
</tr>
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<tbody>
<tr>
<td>Quinine</td>
<td>0.5-1 mM</td>
<td>++</td>
</tr>
<tr>
<td>Barium</td>
<td>2-4 mM</td>
<td>++</td>
</tr>
<tr>
<td>4-AP</td>
<td>5 mM</td>
<td>+</td>
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<tr>
<td>TEA</td>
<td>10 mM</td>
<td>+</td>
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<tr>
<td>TBA</td>
<td>10 mM</td>
<td>+</td>
</tr>
<tr>
<td>CTX</td>
<td>25 mM</td>
<td>+</td>
</tr>
<tr>
<td>Cadmium</td>
<td>1 mM</td>
<td>+</td>
</tr>
<tr>
<td>DISA</td>
<td>0.3 mM</td>
<td>+</td>
</tr>
<tr>
<td>SITS</td>
<td>0.5-1 mM</td>
<td>+</td>
</tr>
<tr>
<td>Ouabain</td>
<td>1 mM</td>
<td>+</td>
</tr>
<tr>
<td>Gdolinium</td>
<td>10-100 mM</td>
<td>++</td>
</tr>
</tbody>
</table>

1. Agents were added immediately before cells were exposed to 0.59x Hanks' and volume was monitored for the next 30 minutes.
2. +, complete block; +, partial block.
3. Cells were incubated in SITS for 30 minutes prior to hypotonic stress.

Increasing the barium concentration from 1 to 4 mM did not result in further inhibition of RVD (Fig. 4A), while decreasing the barium concentration to 0.5 mM produced a smaller inhibition of the RVD response (data not shown). In THP-1 cells, 4 mM barium was required to significantly slow RVD (Fig. 4B). The K channel blockers, CTX (25 nM), TEACl (10 mM), TBACI (10 mM), and 4-AP (5 mM) had little or no effect on RVD in either cell type (Table 1).

Is the K gradient required for RVD? To determine if the K gradient is required for RVD, both THP-1 and HL-60 cells were placed in a 0.59x KCl Hanks' and cell volume was monitored. Neither cell type exhibited RVD in 0.59x KCl Hanks'. Furthermore, 0.59x KCl Hanks' produced a secondary swelling in both cell types. This is evident in Figure 5 in which representative responses of THP-1 and HL-60 cells to 0.59x KCl
Hanks' are depicted along with RVD responses in 0.59× NaCl Hanks' The secondary swelling response for HL-60 cells was considerably greater (188% ± 2.1 at 30 minutes [n = 5]) than that for THP-1 cells (159% ± 1.7 at 30 minutes [n = 7]). To determine if the diminished secondary swelling in THP-1 cells compared to HL-60 cells reflects lower cation permeability in THP-1 cells compared to HL-60 cells, gramicidin was added to the bath 10-12 minutes after the hypotonic stress and volume were monitored. Gramicidin had little or no effect on the final volume of THP-1 cells after 30 minutes in 0.59× KCl Hanks' (gramicidin-treated volume was 159%, n = 4), suggesting that the anion permeability or gradient of THP-1 cells accounted for the difference in the magnitude of the secondary swelling.

**Anion permeability**

Gramicidin produced little or no change in the mean volume of THP-1 or HL-60 cells bathed in isotonic Hanks', indicating that the basal Cl permeability of THP-1 and HL-60 cells is low. In contrast, Cl permeability increased in response to hypotonic stress since, as noted above (Fig. 3) the addition of gramicidin induced RVD when HL-60 and THP-1 cells were bathed in 0.59× Hanks' plus quinine. Furthermore, gramicidin's ability to overcome the quinine-induced block of RVD indicates that the swelling-induced anion and cation permeabilities are independent.

**Anion channel blockers.** RVD was blocked in both cell types by the anion channel blocker DISA, which has been shown to block RVD in neutrophils (Stoddard et al., 1993). In 0.59× Hanks' the 30-minute volume in DISA-treated cells (0.2 mM) was 136 ± 10 (N = 3) and 136 ± 8 (n = 3) in HL-60 cells and THP-1 cells, respectively. The DISA-induced block of RVD could not be overcome by addition of gramicidin (1 μM). In contrast, SITS (0.5–1 mM) produced only a partial block in THP-1 cells (112% ± 3 of control volume at 30 minutes) and in HL-60 cells (114% ± 4 of control volume at 30 minutes), even when cells were preincubated with 1 mM SITS for 30 minutes before exposure to hypotonic medium (Fig. 6).

**Anion selectivity.** As noted earlier (Fig. 5), HL-60 cells in 0.59× KCl Hanks' failed to volume regulate and showed a secondary volume increase which was presumably due to an influx of both K and Cl. In order to investigate the selectivity of the anion efflux pathway, the magnitude of the secondary volume increase was monitored in HL-60 cells bathed in 0.59× high K Hanks' containing different anions. As shown in Figure 7, bathing HL-60 cells in Kmetanesulfonate Hanks' or Kglutamate Hanks' abolished the secondary volume increase evident in 0.59× KCl Hanks'. In contrast, cells in Kbromide Hanks' displayed a secondary volume increase similar to that of the cells in KCl Hanks'. Thus, the volume-induced anion efflux mechanism is permeable to both Cl and Br and is impermeable to methanesulfonate and glutamate.

**Role of calcium**

To determine if a calcium influx was required for the RVD response, volume was monitored in THP-1 and HL-60 cells exposed to 0.59× Hanks' which contained 1.0 mM EGTA and no calcium. RVD occurred normally in both cell types in this solution. Furthermore, cadmium (1 mM), a calcium channel antagonist in other cells, had no effect on RVD in either cell type (Fig. 8).

Despite the absence of a requirement for calcium influx during RVD, it is possible that swelling causes release of Ca2+ stores that in turn can activate calcium-gated channels. To explore this possibility, THP-1 and HL-60 cells were loaded with either fura, or with
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Fig. 7. Time course of representative volume responses of (A) HL-60 cells and (B) THP-1 cells in 0.59× KCl Hanks' (●), 0.59× KBr Hanks' (○), and in 0.59× K methanesulfonate Hanks' (△).

Fig. 8. Time course of representative volume responses of (A) HL-60 cells and (B) THP-1 cells in 0.59× NaCl Hanks' (●), 0.59× NaCl Hanks' containing either 0 calcium/1 mM EGTA (□) or 1 mM CdCl (○).

BAPTA and fura, and both [Ca] changes and cell volume were monitored during hypotonic stress. Since volume measurements were performed at room temperature, [Ca] measurements also were carried out at room temperature. As shown in Figure 9, swelling induced increases in [Ca] in both THP-1 and HL-60 cells, although the response of THP-1 cells was quite small; the percentage increase in [Ca] compared to baseline levels obtained before addition of 41% water was 273% ± 26 (n = 6) for HL-60 cells and 154% ± 16 (n = 4) for THP-1 cells. BAPTA loading both cell types abolished the induced [Ca] rise (Fig. 9). However, BAPTA loading did not block RVD in either cell type in response to either 0.59× NaCl Hanks' or 0.59× Ca-free, EGTA-containing NaCl Hanks' (shown in Fig. 10).

Effect of swelling on [Ca]. A: B: HL-60 cells and (C, D) THP-1 cells were loaded with either fura-2 am (A, C) or fura-2-AM plus BAPTA-AM, (B, D). At time indicated by arrows, 41% water was added.

Fig. 9. Effect of swelling on [Ca]. A: B: HL-60 cells and (C, D) THP-1 cells were loaded with either fura-2 am (A, C) or fura-2-AM plus BAPTA-AM, (B, D). At time indicated by arrows, 41% water was added.

DISCUSSION

This study demonstrates that RVD in THP-1 cells, a monocytic cell line, and in HL-60 cells, a promonocytic cell line, involves efflux through independent anion and cationic transport mechanisms. While little data are available about the ionic conductances in HL-60 and THP-1 cells (Wieland et al., 1987, 1990), electrophysiological studies on other myeloid cells have characterized several different K and Cl conductances that may be activated during RVD (Gallin, 1991). In many cases, pharmacological blockers of these ionic conductances have been described. Thus we reasoned that examining the ability of these agents to block RVD in THP-1 and HL-60 cells would provide information about the involvement of these ionic conductances in RVD, and therefore, their presence in these cell lines.

Cation permeability pathway

Surprisingly, quinine was the only K channel antagonist that completely blocked RVD in both cell types: 1 mM quinine blocked RVD in THP-1 cells while 4 mM

Effect of gadolinium

Stretch-activated channels, which are inhibited by gadolinium (Yang and Sachs, 1989), have been implicated in RVD in astrocytes and other cells (Medrano and Gruenstein, 1993). However, in the presence of 1-100 μM gadolinium chloride, THP-1 and HL-60 cells exhibited normal RVD (data not shown).
Barium, the other K channel blocker tested that affected RVD in THP-1 and HL-60 cells, only partially blocked RVD. Perhaps the best characterized K conductance in macrophages is a barium-inhibitable voltage-dependent inwardly rectifying K conductance (Kᵢ). The Kᵢ conductance has been cloned recently (Kubo et al., 1993) and, when present, sets the macrophage resting membrane potential near to the K equilibrium potential (Gallin, 1981). This conductance also enables cells to flip between two stable states of resting membrane potential (Gallin, 1981). Barium block of Kᵢ occurs in 300–500 μM extracellular barium (McKinney and Gallin, 1988). In contrast, 1–4 mM barium was required to slow RVD, but failed to completely inhibit it. Thus, the Kᵢ conductance is unlikely to mediate RVD in HL-60 cells or in THP-1 cells. Partial block of RVD by barium could be due either to a partial block by barium of a single cation conductance or to the involvement of more than one cation conductance in RVD, a barium-sensitive conductance and a barium-insensitive conductance in RVD. In the human colon adenocarcinoma HT-29 cell line, where noise analysis has implicated only one type of K channel in volume regulation, the volume-sensitive K channel is completely blocked by quinine but only partially blocked by barium (Illek et al., 1992). Volume regulation in macrophages may involve a similar K channel.

None of the other K channel blockers tested in THP-1 and HL-60 cells inhibited RVD. Consequently, unlike lymphocytes where a CTX-sensitive "delayed-rectifier type" K conductance has been implicated in RVD (Grinstein and Smith, 1990; Grinstein and Foskett, 1990) or cultured proximal tubule cells where a Ca-activated K conductance (Dube et al., 1990) has been implicated in RVD, volume regulatory responses in myeloid cells do not appear to involve either the Ca-activated K conductances or delayed-rectifier type K conductances that have been described in myeloid cells.

The secondary swelling evident in both THP-1 and HL-60 cells bathed in 0.59× KCl Hanks' indicates that opening of the swelling-induced cation permeability does not depend on the K gradient, nor is it inactivated by depolarization. Our experiments with gadolinium, an antagonist of stretch-activated cation channels, also suggest that these channels are not involved in RVD of HL-60 and THP-1 cells. Further studies using both patch clamp techniques and volume measurements are needed to determine the cation permeability pathway(s) that are involved in RVD in macrophages.

**Anion permeability pathway**

Addition of gramicidin to either cell type produced little change in cell volume unless cells were exposed to hypotonic medium, indicating that the basal anion permeability of both THP-1 cells and HL-60 cells is low and that it increases following swelling. As in lymphocytes, our studies with high K hypotonic medium containing different anions indicated that the swelling-induced anionic transport pathway is permeable to both Cl⁻ and bromide ions but relatively impermeable to methanesulfonate and glutamate (Grinstein et al., 1982). While SITS completely blocks RVD in neutrophils (Stoddard et al., 1993) and other cells, it only produced a partial block of the RVD response in THP-1...
and HL-60 cells. On the other hand, RVD was blocked in both cell types by the anion transport blocker DISA and this block could not be overcome by gramicidin. DISA, unlike SITS, is lipophilic and probably crosses the membrane through nonionic diffusion (Simchowitz et al., 1993). In human neutrophils DISA blocked both RVD and the swelling-induced increases in $^{36}$Cl efflux and CI currents (Stoddard et al., 1993; Simchowitz et al., 1993). In addition to blocking RVD in several cell types, DISA is a noncompetitive inhibitor of Cl-CI exchange and Na independent CI/HCO$_3^-$ exchange (Restrepo et al., 1992). Thus, while we can conclude that under our experimental conditions (in HEPES-buffered media without bicarbonate) swelling increases a DISA-inhibitable anionic transport mechanism(s), DISA may block RVD by acting on several different anionic transport mechanisms. In this context, a preliminary report by Hallows et al. (1992) showed that for HL-60 cells suspended in bicarbonate-containing hypotonic medium, the CI efflux was significantly less than the K efflux, suggesting that additional osmolytes are involved in the RVD response in HL-60 cells. In astrocytes, cell swelling induces a rapid release of taurine, which is markedly reduced by the Cl channel inhibitor DIDS (Pasantes-Morales et al., 1990). If the loss of additional osmolytes is also involved in the RVD response of HL-60 cells maintained in bicarbonate-free medium, then DISA must block both the Cl permeability change and the efflux of the additional osmolytes. Alternatively, it is possible that under the conditions of our experiment, the magnitude of the CI efflux is similar to that of K efflux and DISA is acting to block only a Cl efflux. Chloride efflux studies will be needed to differentiate between these possibilities.

Delineating the ionic basis of RVD provides information about the ionic transport mechanisms present in myeloid cells, and may even be useful in assessing changes in transport that occur during activation and/or differentiation. Blood monocytes will be exposed to large changes in tonicity during passage through the kidney. On the other hand, tissue macrophages are probably exposed to only small changes in tonicity at sites of inflammation or following changes in metabolism. Volume regulation following swelling induced by enhanced amino acid uptake has been demonstrated in a number of cells including intestinal enterocytes (MacLeod et al., 1992). It is possible that stimulation of nitrous oxide formation and the subsequent increase in arginine uptake may lead to cell swelling in macrophages.

In conclusion, this study demonstrates that RVD in two myeloid cell lines is mediated by independent quinine-inhibitable cation and DISA-inhibitable anion transport pathways that are not gated by increases in [Ca$^2+$] and that the cation permeability pathway mediating RVD in these cells is different from the pathway mediating RVD in lymphocytes.

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LITERATURE CITED


sium current in differentiating human leukemic (HL-60) cells. J.
stimulating factor (CSF-1) modulates a differentiation-specific in-
ward-rectifying potassium current in human leukemic (HL-60)
Yang, X.C., and Sachs, F. (1989) Block of stretch activated ion chan-
nels in Xenopus oocytes by gadolinium and calcium ions. Science,
243:1068–1071.