Ionic imbalance, in addition to molecular crowding, abates cytoskeletal dynamics and vesicle motility during hypertonic stress

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Cell volume homeostasis is vital for the maintenance of optimal protein density and cellular function. Numerous mammalian cell types are routinely exposed to acute hypertonic challenge and shrink. Molecular crowding modifies biochemical reaction rates and decreases macromolecule diffusion. Cell volume is restored rapidly by ion influx but at the expense of elevated intracellular sodium and chloride levels that persist long after challenge. Although recent studies have highlighted the role of molecular crowding on the effects of hypertonicity, the effects of ionic imbalance on cellular trafficking dynamics in living cells are largely unexplored. By tracking distinct fluorescently labeled endosome/vesicle populations by live-cell imaging, we show that vesicle motility is reduced dramatically in a variety of cell types at the onset of hypertonic challenge. Live-cell imaging of actin and tubulin revealed similar arrested microfilament motility upon challenge. Vesicle motility recovered long after cell volume, a process that required functional regulatory volume increase and was accelerated by a return of extracellular osmolality to isosmotic levels. This delay suggests that, although volume-induced molecular crowding contributes to trafficking defects, it alone cannot explain the observed effects. Using fluorescent indicators and FRET-based probes, we found that intracellular ATP abundance and mitochondrial potential were reduced by hypertonicity and recovered after longer periods of time. Similar to the effects of osmotic challenge, isovolumetric elevation of intracellular chloride concentration by ionophores transiently decreased ATP production by mitochondria and abated microfilament and vesicle motility. These data illustrate how perturbed ionic balance, in addition to molecular crowding, affects membrane trafficking.

A basic challenge for all animal cells is to maintain constant volume and intracellular electrolyte composition (1, 2). K⁺ is the principal intracellular cation, and Na⁺ is extruded from the cell interior. Cellular concentration of these ions is controlled by energy-consuming transport mechanisms, primarily the Na⁺/K⁺-ATPase. Cl⁻ is a major anion in the extracellular milieu that is maintained at relatively low concentrations (5–40 mM) within the cytosol by a variety of plasma membrane transporters, including the CFTR and members of the chloride channel, voltage-sensitive (CIC) family of Cl⁻ channels (3). Together, these ions not only maintain cell volume and a biologically compatible intracellular milieu but also allow passive diffusional movement of ions down their electrochemical gradients that is central to cell function, from general cell division to the more specialized processes of secretion and contractibility. Various types of environmental challenge lead to electrolyte imbalance, as illustrated by variations of environmental toxicity that immediately alter cell volume and intracellular ion composition (4–6). Cells shrink rapidly in hypertonic environments and swell in hypotonic environments. Cell volume is reestablished quickly—within minutes—in mammalian cells by salt (Na⁺ and Cl⁻) influx (in response to hypertonic stress) or K⁺ efflux (in response to hypotonic stress), but such changes occur at the expense of electrolyte imbalance. Prechallenge intracellular ion composition is reestablished gradually over time (hours to days) by readjusting the intracellular levels of nonionic solutes, principally amino acids and other organic compounds (2, 6) that have far fewer effects on enzyme activity.

Numerous organisms across the animal kingdom are routinely exposed to fast variations of environmental osmolality. In mammals, many cell types must deal with hypertonic challenge as part of their daily function. These include blood cells passing through the renal medulla and intestinal epithelial cells after food ingestion (7). Moreover, hypertonic fluids are routinely administered to treat disorders such as hyponatremia, cystic fibrosis, and hemorrhagic shock (8–10). Hypertonicity affects a wide range of cellular processes. For example, it causes DNA damage (11) and protein damage and aggregation (5, 12), promotes autophagy (13), and leads to apoptosis when challenge is overwhelming (4). Membrane trafficking is a major process affected by hypertonicity that probably contributes significantly to the change incurred upon challenge. Indeed, we and others have observed that endocytosis, exocytosis, and vesicular trafficking are reduced from the onset of challenge, and that the reduction can persist hours after challenge (14–19). However, how immediate changes in the cell’s biophysical parameters influence these events is unclear. By inducing cell shrinkage, hypertonicity increases intracellular component density and molecular crowding (20). Increased crowding

Significance

Maintenance of cell volume and ionic homeostasis are fundamental to cell function. Cells shrink when extracellular osmolality is increased, and although mechanisms controlling cell volume recovery are well documented, the mechanisms underlying changes in membrane trafficking that occur during adaptation to high osmolality are not well understood. Molecular crowding arising from cell shrinkage can alter protein structure and has been proposed to provoke changes in trafficking events. However, cell volume recovery involves a rapid influx of ions, and the effects of ionic imbalance on trafficking dynamics in living cells remain largely unexplored. We found that high levels of chloride and loss of ATP, in addition to molecular crowding, contribute to altered cytoskeletal and vesicular dynamics during hypertonic stress.

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of macromolecules and organelles by water efflux results in cytoplasm stiffening (21), protein misfolding and aggregation (12), and reduces protein diffusion (22). Therefore it is reasonable to suppose that molecular crowding may interfere with at least some trafficking events. On the other hand, because osmotically induced cell shrinkage is invariably accompanied by increased intracellular salt concentration, the relative contribution of each occurrence on membrane trafficking is unclear.

High levels of intracellular salt after hypertonic challenge may perturb a number of parameters that collectively affect membrane trafficking. Microtubules (MT) and actin filaments play paramount roles in membrane trafficking. Their dynamic nature creates forces that help move cargo, and they supply “tracks” for long-range delivery of cargo (23). A key feature of MT biology is dynamic instability, so that tubules continuously switch between episodes of steady growth and rapid shrinkage (24), allowing the cell to probe the cytoplasm constantly and adapt quickly to environmental change. A hallmark feature of hypertonicity is that it induces actin cytoskeleton reorganization (4), and we recently have shown that it also induces fast MT remodeling (13). Because the dynamics of microfilament assembly/disassembly are sensitive to ionic conditions (25, 26), a sudden increase in intracellular salt could alter their dynamics and impinge on membrane trafficking. High levels of intracellular salt also could affect membrane trafficking by modulating available energy. Increased ion concentration decreases mitochondrial matrix volume and disrupts key parameters of the respiratory chain, as shown in isolated mitochondria (27, 28) and as predicted by mathematical models of mitochondrial bioenergetics (29). Thus, hypertonicity conceivably could affect mitochondrial function by decreasing ATP production by mitochondria. However, the influence of hypertonicity on mitochondrial function in intact cells has been investigated mostly in the context of apoptosis induced by severe hypertonic stress, which disrupts mitochondrial potential (30–32).

In the present study, we examined the effects of molecular crowding, high intracellular Cl− concentration [Cl−]i, microfilament remodeling, and altered mitochondrial function by hypertonicity on vesicle motility. We show that strong but sustainable hypertonic challenge to mammalian cells immediately halts both microfilament motility and the motility of a wide variety of endosome-vesicle subpopulations. We demonstrate that vesicle motility recovers significantly more slowly than cell volume and that an isovolumetric increase in [Cl−], by ionophores triggers mitochondrial depolarization and decreases ATP levels, microfilament dynamics, and vesicle motility. Thus, these data illustrate how intracellular ionic imbalance, in addition to molecular crowding, affects membrane trafficking and cell function.

Results
Hypertonicity Induces a Transient and Ubiquitous Decrease in Vesicle Motility. We have shown previously that hypertonicity reduces endocytosis and exocytosis and affects vesicle distribution concomitant with autophagy in porcine LLC-PK1 renal proximal tubule-like cells (13, 18). We also showed that the great majority of cells survive when medium osmolality is increased from 300 to 500 mOsmol/kg but not to 700 mOsmol/kg (13). To explore further the mechanisms underlying these changes in intracellular trafficking, we analyzed the motility of FITC-dextran–loaded endosomes in live LLC-PK1 cells by spinning-disk confocal imaging (Fig. 1). About 90% of FITC-dextran–loaded endosomes were mobile under isosmotic conditions, indicating that the dextran was incorporated into functional endosomes efficiently. Consistent with expected patterns (33), movement consisted of either quick, short trajectories that frequently changed direction or longer, more linear trajectories that sometimes spanned the entire length of cells. Endosome velocity was broadly distributed, ranging from the fast motility characteristic of MT-based movement (30–60 µm/min [34, 35]) to the slower motility distinctive of movement mediated by myosin motor proteins and actin treadmilling [1.2–12 µm/min (36–40)]. Because of their large size, movement of dextran-loaded endosomes can be tracked easily by live-cell imaging (Fig. 1D). We quantified their motility by measuring the mean endosome displacement over 2 min at 12-s intervals. We considered only mobile endosomes that remained in the focal plane during the entire 2-min time interval, thus limiting our analysis to slower-moving endosomes. The distribution of the average velocity (displacement/time) of individual endosomes during a 2-min time interval under isotonic conditions is depicted in Fig. 1B. Despite the high variability (SD) in the velocity of individual endosomes (10.6 ± 5.2 µm/min), the average velocity of tracked endosomes (30 for each 2 min time interval) was highly reproducible among experiments (10.6 ± 1.1 µm/min) (Fig. SL4). The motility of at least 95% of all labeled endosomes was abolished almost completely immediately after NaCl challenge.
Reduced vesicle motility by hypertonicity is a general phenomenon affecting numerous endosome/vesicle subpopulations in various cell types. (A) Endosome/vesicle motility before and after NaCl challenge (500 mOsm/kg) in LLC-PK1 cells transfected with various GFP-tagged Rab isoforms (Rab7, -10, and -11) or ssYFP, each used as a marker for specific endosome/vesicle subpopulations as shown in the table at right. The motility of all subpopulations was abolished immediately after challenge, whereas recovery rates varied, as shown in the table at right. (B) Kymographs illustrate different rates of recovery in cells expressing GFP-Rab10, ssYFP, or GFP-Rab11. (C–E) The effects of NaCl challenge on the motility of dextran-loaded endosomes in primary hepatocytes (C), primary pancreatic β cells (D), and cultured pancreatic MIN6 cells (E). (Also see Figs. S1 and S2 and Movies S2 and S3.) Error bars show mean ± SEM from four independent experiments.

Fig. 2. Reduced vesicle motility by hypertonicity is a general phenomenon affecting numerous endosome/vesicle subpopulations in various cell types. (A) Endosome/vesicle motility before and after NaCl challenge (500 mOsm/kg) in LLC-PK1 cells transfected with various GFP-tagged Rab isoforms (Rab7, -10, -11, and ssYFP), each used as a marker for specific endosome/vesicle subpopulations as shown in the table at right. The motility of all subpopulations was abolished immediately after challenge, whereas recovery rates varied, as shown in the table at right. (B) Kymographs illustrate different rates of recovery in cells expressing GFP-Rab10, ssYFP, or GFP-Rab11. (C–E) The effects of NaCl challenge on the motility of dextran-loaded endosomes in primary hepatocytes (C), primary pancreatic β cells (D), and cultured pancreatic MIN6 cells (E). (Also see Figs. S1 and S2 and Movies S2 and S3.) Error bars show mean ± SEM from four independent experiments.
MT motility generally was decreased by urea, actin motility increased above baseline levels shortly after challenge and then returned to baseline levels.

To provide a framework within which shifts of microfilament motility caused by hypertonic challenge and their effects on vesicle motility can be compared, we next investigated how sudden shifts of microfilament polymerization by NaCl challenge affect vesicle motility. We previously observed that decreased MT motility by NaCl is associated with immediate MT depolymerization, with repolymerization occurring within minutes of challenge (13). Cell-permeable urea also induced strong MT depolymerization upon challenge (Fig. S4A). Unlike MTs, actin polymerizes quickly after challenge by either NaCl or urea and remains polymerized for long periods of time, as revealed by Western blot analysis of fractionated G- vs. F-actin and by TRITC-phalloidin fluorescence analysis (Fig. S5). Then, we compared the effects of NaCl with those of chemicals that alter microfilament polymerization or polymer stability (Fig. 3 B and C). We previously showed that taxol, an agent that stabilizes MT polymers, and nocodazole, which interferes with tubulin polymerization, immediately increase and decrease tubulin polymerization, respectively (13). Taxol also increased actin polymerization (Fig. S5 B and C) but had no effect on either actin (Fig. S6 B and C) or vesicle motility (Fig. 3C). Similar to taxol, nocodazole increased actin polymerization (Fig. S5 B and C) but strongly decreased both actin (Fig. S6 B and C) and vesicle motility (Fig. 3C). It also restored Rmax after NaCl challenge (Fig. 3C). Colchicine, another chemical agent that interferes with tubulin polymerization, also decreased vesicle motility (Fig. 3C). Jasplakinolide, an agent that stabilizes actin filaments by binding to F-actin, immediately increased actin polymerization (Fig. S5) and decreased actin filament motility (Fig. S6 B and C) but had no effect on MT motility (Fig. S6 A and C). It immediately reduced, but did not abolish, vesicle motility and, interestingly, had no effect on Rmax when applied after NaCl challenge (Fig. 3B). Latrunculin B, an agent that complexes with G-actin, thereby inhibiting actin polymerization, immediately induced actin depolymerization (Fig. S5) and had no effect on either MT (Fig. S6 A and C) or vesicle motility (Fig. 3B).

These data show that chemical agents that decrease microfilament motility reduce vesicle motility (Fig. 3D) and indicate that a decrease in microfilament dynamics, rather than sudden shifts in their polymerization states per se that conceivably could interfere with membrane trafficking events, may be implicated in the reduction of vesicle motility by NaCl challenge. Our data also suggest that neither jasplakinolide nor latrunculin B alters MT remodeling, at least not to the levels achieved by hypertonic stress. This finding is supported by our observation that neither chemical agent affects end-binding protein 1 (EB1) comet-like structure or motility, which can be used as a means to follow MT growth (Movies S5 and S6; the effects of NaCl on EB1 motility are described in ref. 13). On the other hand, the stabilization or destabilization of MT networks by taxol and nocodazole, respectively, induces actin polymerization, as previously observed (48-53).

**Full Recovery of Vesicle Motility Following NaCl Hypertonic Challenge Occurs Long After Cell Volume Recovery.** We next examined how changes in cell volume caused by either NaCl or cell-permeable urea affect vesicle motility by measuring the fluorescence intensity of calcine-AM. NaCl spontaneously reduced LLC-PK1 cell volume in a dose-dependent manner (Fig. 4A). Equiosmolar urea produced a similar loss of cell volume (Fig. 4A), indicating that water efflux upon urea challenge precedes influx of urea in LLC-PK1 cells. As expected, cell volume recovered significantly faster after urea challenge than after NaCl challenge, with R1/2 occurring at 1.0 ± 0.03 min for urea challenge and at 2.5 ± 0.03 min for NaCl challenge. Cell volume recovered almost completely (96 ± 0.1%) after NaCl challenge and recovered com-
Fig. 4. Vesicle motility following NaCl-hyperotic challenge recovers well after cell volume is restored. (A, Left) LLC-PK₁ cell volume was measured using calcein-AM under isotonic conditions and after challenge (arrow) with NaCl (350 or 500 mM) or urea (500 mM). (Right) LLC-PK₁ cell volume (solid lines) and FITC-dextran–loaded endosome motility (vesicle mvt, dotted lines) after challenge with NaCl or urea (500 mM) normalized to maximal variations for comparison of their recovery kinetics. (B) Projected FITC-dextran–loaded endosome motility in LLC-PK₁ cells as a function of cell volume. (C) LLC-PK₁ cells were challenged or not with an RVI inhibitor mixture (red arrow), and cell volume (Left) and FITC-dextran–loaded endosome motility (Right) were measured before and after (arrow) NaCl challenge (500 mM). (D) LLC-PK₁ cell volume (Left) and FITC-dextran–loaded endosome motility (Right) before and after NaCl challenge (500 mM) and after osmolality returned to basal, isotonic conditions, either immediately after challenge or after a longer period (12 min), when near-maximal volume recovery was reached (blue and red arrows, respectively). Error bars show mean ± SEM from four independent experiments.

Macromolecule Dynamics Are Reduced by an Isovolumetric Increase in [Cl⁻]. Loss of intracellular water upon hypertonic challenge increases Na⁺ and Cl⁻ levels within cells. These levels remain elevated during RVI as the driving force for water reentry into cells. Consequently, hyperosmotic water influx induces an electrolyte imbalance and depolarizes the membrane of various cell types, including proximal tubule cells (54–56). We first tested whether membrane depolarization by an acute isotonic increase in extracellular K⁺ concentration (72 mM) or K⁺ channel blockade by tetraethylammonium chloride (5 µM) had an effect on vesicle motility in LLC-PK₁ cells. No differences were found, indicating that cell depolarization is not responsible for the observed defect in vesicle trafficking. As a model of ionic imbalance, we examined the effects of high levels of [Cl⁻] on macromolecule motility under isovolumetric conditions by challenging cells with a previously described (57) combination of tributyltin and nigericin that already had been validated in LLC-PK₁ cells (58). The Cl⁻/OH⁻ antiporter tributyltin forms pores in the cell membrane that allow extracellular Cl⁻ to equilibrate with intracellular Cl⁻. The H⁺/K⁺ ionophore nigericin is added to clamp intracellular pH. We confirmed that [Cl⁻] was increased by tributyltin/nigericin ionophores (TBTN) by loading LLC-PK₁ cells with N-(ethoxycarbonylmethyl)-6-methoxyquinolinium bromide (MQAE), a fluorescent dye that quenches in the presence of Cl⁻ ions (59). To account for fluorescence changes induced by differences in cell volume, cells were loaded simultaneously with calcein-AM, and the ratio of MQAE/calcein fluorescence was recorded. MQAE fluorescence was quenched by TBTN, NaCl, and also by glucose challenge (500 mM) (Fig. 5A and B and Fig. S7A). Similar to the effect of osmolyte challenge, fluorescence remained low for long periods of time in the presence of TBTN, which did not affect cell volume (Fig. 5A). As expected, contrary to the effects of TBTN and NaCl challenge, MQAE fluorescence was quenched only transiently by urea (Fig. 5B). We conclude that TBTN at least partly mimics ionic imbalance by NaCl challenge without the changes in cell volume that invariably accompany hypertonic challenge. TBTN rapidly decreased but did not abolish the motility of FITC-dextran–loaded endosomes (Fig. 5C). Interestingly, it decreased motility to a lesser extent than the steady-state motility observed following strong (500 mM) NaCl challenge. We estimated actual [Cl⁻] values in LLC-PK₁ cells using a previously described calibration method performed in this same cell line (59). [Cl⁻] was estimated to be 16.9 ± 0.6 mM under resting, isotonic conditions and increased strongly after NaCl, urea, or TBTN challenge (Fig. S7C and D). The relationship among cell volume, [Cl⁻], and FITC-dextran–loaded endosome motility is depicted in Fig. 5D and E. Endosome motility decreased steadily in response to increased [Cl⁻] induced by TBTN, which does not alter cell volume, and then fell dramatically when [Cl⁻] was increased by ≥20 mM. Endosome motility similarly was reduced at
50 mM after long periods of NaCl challenge (i.e., at a time when cell volume had recovered almost fully). Endosome motility then fell dramatically to near immobilization when [Cl−] was increased above 50 mM immediately after either NaCl or urea challenge, likely reflecting the combined effects of increased [Cl−] and reduced cell volume. These figures also indicate that, although urea influx allows fast recovery of cell volume and [Cl−], the intracellular accumulation of urea actually decreases endosome motility, although to a lesser extent than in response to elevated [Cl−]. This finding may help explain the residual loss of endosome motility by hypertonicity could affect microfilament dynamics and vesicle motility. We examined this possibility by monitoring real-time changes in intracellular ATP concentration ([ATP]) in LLC-PK1 cells using the recently developed FRET-based ATP indicator ATeam (60). We compared the effects of NaCl, urea, and the ATP synthase inhibitor oligomycin on YFP/CFP emission ratios of wild-type and mutant (AT1.03R122K/R128K) ATeam probes (Fig. 6A). As expected, the basal emission ratio of the mutant probe, which does not respond to [ATP], (60), was lower than that of the wild-type probe (0.42 mutant FRET ratio compared with 1.3 wild-type FRET ratio). Unexpectedly, the emission ratio of the mutant probe was increased immediately by NaCl challenge (Fig. 6A). Equiosmolar urea produced a similar effect (Fig. 6A). Because equiosmolar NaCl and urea reduce cell volume to similar extents (Fig. 4A), this ATP-independent increase in the FRET emission ratio probably is caused by a conformational change in the probe’s sensor domain that results from cell shrinkage-induced molecular crowding and/or high ionic strength. This effect increased dramatically with the magnitude of osmotic challenge and prevented us from conducting experiments with a challenge above 400 mOsmol/kg. Despite this drawback, the emission ratio of the wild-type probe was partly quenched immediately following NaCl challenge (400 mOsmol/kg) and, to a lesser extent, after urea

High Levels of [Cl−] Decrease Mitochondrial Function and Intracellular ATP Content. Membrane trafficking is a tightly regulated process that relies on available energy. Perturbations of ATP production by hypertonicity could affect microfilament dynamics and vesicle motility. We examined this possibility by monitoring real-time changes in intracellular ATP concentration ([ATP]) in LLC-PK1 cells using the recently developed FRET-based ATP indicator

Fig. 5. An isovolumetric increase in [Cl−], decreases vesicle and microfilament motility. (A and B) LLC-PK1 cells were challenged (arrows) with the ionophores nigericin (5 μM)/tributyltin chloride (10 μM) (TBTN) (A), NaCl (500 mOsmol/kg), or urea (500 mOsmol/kg) (B). Error bars show mean ± SEM from three independent experiments. Relative changes of MQAE fluorescence were measured as the normalized emission ratio of MQAE (360 nm) to calcine (488 nm). MQAE quenching by KSCN (150 mM) confirms the probe was operating within its dynamic range. Inset in A shows changes in cell volume measured with calcine alone. (C) Decreased FITC-dextran-loaded endosome motility (E) in LLC-PK1 cells. (D) Time-lapse pseudocolored images of GFP-α-tubulin and GFP-β-actin in LLC-PK1 cells before and after arrow) TBTN challenge illustrating decreased but not abolished motility in both microfilaments after challenge. (Scale bar: 10 μm.) Representative images are shown on the left, and motility quantification is shown on the right. Error bars show mean ± SEM of three independent experiments. *P < 0.05 vs. control.
challenge (400 mOsmol/kg), indicating that each osmolyte decreases [ATP], but does so to different extents. Contrary to the high levels of [Cl\textsuperscript{−}], that are sustained by NaCl challenge (Fig. 5B), [ATP], recovered almost completely after long periods of challenge (Fig. 6A). [ATP] also decreased at first but then recovered completely after glucose challenge (Fig. 5B). As expected, a combination of oligomycin and the glycolysis inhibitor 2-deoxy-glucose (2-DOG) decreased [ATP] (Fig. 6A). Similar to our FRET data, analysis of [ATP] by a fluorometric assay revealed that [ATP] was decreased by NaCl challenge and, to a lesser extent, by urea challenge (500 mOsmol/kg), returned to basal levels after longer periods of challenge, and was decreased by oligomycin/2-DOG (Fig. 6B). FRET analysis of LLC-PK\textsubscript{i} cells additionally revealed that [ATP] was decreased initially by an isovolumetric increase in [Cl\textsuperscript{−}], induced by TBTN and recovered only partly after longer periods of challenge (Fig. 6C), possibly reflecting increased ATP consumption and/or loss.

Our data indicate that increased [Cl\textsuperscript{−}], induced by NaCl and TBTN might contribute to decreased vesicle and microfilament motility partly because of decreased [ATP], at least during the early stages of challenge. This notion led us to examine the effects of ATP depletion by oligomycin/2-DOG on macromolecular dynamics in LLC-PK\textsubscript{i} cells. Similar to the effect of TBTN, oligomycin/2-DOG decreased but did not abolish the motility of FITC-dextran-loaded endosomes (Fig. 6D) and reduced MT and actin motility (Fig. 6E). These observations indicate that a sudden increase in [Cl\textsuperscript{−}], transiently decreases [ATP], which in turn affects macromolecular dynamics.

The decreased [ATP], induced by NaCl challenge could arise from perturbed mitochondrial function. Ratiometric imaging of the mitochondrial membrane potential (ΔΨ) dye JC-1 in LLC-PK\textsubscript{i} cells revealed an F488/F568 signal ratio that increased significantly after NaCl challenge (Fig. 6F). Similar to recovered levels of [ATP], after long periods of challenge (Fig. 6 A and B), ΔΨ gradually returned toward basal levels after longer periods of time (Fig. 6F). Cells challenged with TBTN displayed similar, transient mitochondrial dysfunction despite sustained elevated levels of [Cl\textsuperscript{−}], (Fig. 6F). Mitochondria exist as a network that is continuously shaped by fusion and fission events, and their excessive fragmentation is a common consequence of mitochondrial dysfunction (61). Both NaCl and TBTN challenge increased mitochondrial fission, and mitochondrial remodelling was sustained even after glucose challenge (Fig. 6C). Therefore, a sudden increase in [Cl\textsuperscript{−}], transiently decreases [ATP], at least partly as a consequence of mitochondrial dysfunction.

Discussion

The motility of various endosome-vesicle subpopulations was nearly abolished immediately after NaCl-hypertonic challenge, even when osmolality was increased only moderately (by 50 mOsmol/kg). This observation made in various mammalian cell types is similar to the decreased vesicle trafficking observed in yeast exposed to severe osmotic stress (3 M sorbitol; i.e., 3,000 mOsmol/kg) (14). Osmotic compression per se and accompanying molecular crowding undoubtedly play a large part in the vesicle immobilization that accompanies water efflux, because the intracellular space is filled to near capacity with macromolecules and organelles under basal conditions (62). An optimal protein density that maximizes biochemical reaction rates may exist in cells (22). By increasing macromolecule density, cell shrinkage hinders protein diffusion and biochemical reactions (14, 22). This idea is supported by the finding that eukaryotic cells under compressive stress behave as strong glass because of crowding-induced stiffening of the cytoplasm (21). Although vesicle motility certainly is affected by molecular crowding, our data suggest that other factors that arise as part of the hypertonic stress response may be involved, because volume recovery did not immediately relieve the depression in vesicular dynamics. Because RVI relies on electrolyte import, we hypothesized that electrolyte imbalance could be one of these additional factors. Although both the intracellular Na\textsuperscript{+} concentration [Na\textsuperscript{+}], and [Cl\textsuperscript{−}], increase in response to hypertonicity, our study focused on investigating how high levels of [Cl\textsuperscript{−}], may affect vesicle motility in conjunction with molecular crowding, taking advantage of tools for measuring and manipulating [Cl\textsuperscript{−}], already optimized in LLC-PK\textsubscript{i} cells (58, 59). Similar to the effect of NaCl challenge, vesicle motility was abolished when cells were exposed to high concentrations (corresponding to 500 mOsmol/kg) of cell-permeable osmolytes (urea in LLC-PK\textsubscript{i} cells and glucose in pancreatic β cells) but, unlike NaCl challenge, recovered quickly thereafter. Water is drawn back into the cell rapidly as cell-permeable osmolytes accumulate intracellularly. As a result, cell volume is restored quickly, and the accumulation of inorganic electrolytes inside the cell is minimized. The situation is very different in cells challenged with cell-impermeable osmolytes, such as NaCl and mannitol. In this case, cell volume is restored by RVI mechanisms that increase intracellular concentrations of inorganic electrolytes. After NaCl challenge, but not after urea challenge, vesicle motility recovers significantly more slowly than cell volume and remains low well after cell volume is restored. Motility is restored quickly by re-exposing cells to isotonic conditions, which restore intracellular electrolyte composition, but recovery is reduced by an isovolumetric increase of [Cl\textsuperscript{−}], by ionophores. These data demonstrate that vesicle motility is abated by high, but physiological, levels of [Cl\textsuperscript{−}], and after cell volume was recovered. In addition, along with molecular crowding, high levels of [Cl\textsuperscript{−}], probably contribute to reduced motility at the onset of hypertonic challenge, because water loss increases intracellular electrolyte concentration.

We show that a sudden increase of [Cl\textsuperscript{−}], transiently decreases ATP abundance and that chemical block of ATP synthesis reduces vesicle and microfilament motility. These observations indicate that attenuated levels of available ATP by high levels of intracellular salt may contribute to decreased macromolecule motility. ATP is consumed at approximately the same rate as it is synthesized. Our data indicate that high concentrations of [Cl\textsuperscript{−}], may impair mitochondrial function, thereby decreasing ATP synthesis, probably because protein crowding is even higher in mitochondria than in the cytosol (63) and because alterations in matrix volume strongly affect mitochondrial function (64, 65). Accordingly, hypertonicity initially decreases the volume of isolated mitochondria (64), and mitochondrial volume increases at a slower rate of time than cell volume (Fig. 6C). Mitochondria were reported to dissipate ΔΨ in cultured Vero cells within minutes of challenge (30). We show that ΔΨ is decreased by both hypertonic and TBTN challenge. We also show that, reflecting the recovery of cellular ATP abundance, ΔΨ returns to basal levels after long periods of challenge by either stimulus, although [Cl\textsuperscript{−}], levels remain elevated. Although our study focused on the effects of high [Cl\textsuperscript{−}], imbalance of other ions (particularly Na\textsuperscript{+} but potentially others as well) is likely to be involved. Our data argue that an initial ionic imbalance temporarily impairs mitochondrial function but does not preclude mitochondria from adapting to their new milieu. We interpret our data as evidence that mitochondria are able to adapt and restore ATP production despite elevated intracellular ionic strength, but that this adaptation takes a few minutes. This adaptation may involve restored K\textsuperscript{+} equilibrium across the mitochondrial inner membrane, because matrix volume is controlled by the kinetic equilibrium between K\textsuperscript{+} entry and efflux (64, 65). Other mitochondrial ion transporters also may be involved in restoring mitochondrial ΔΨ. [ATP], was decreased almost instantaneously upon NaCl challenge, faster than after oligomycin/2-DOG challenge, probably reflecting fast ΔΨ dissipation by the quasi-instantaneous cell shrinkage induced by NaCl as opposed to the slower effects of chemicals that need time to cross cell membranes. Increased ATP consumption after hypertonic challenge also might participate in decreasing [ATP]. Altered protein structure by hypertonicity might lead to widespread and immediate hydrolysis of protein-bound ATP and/or to rapid ATP/ADP
cycling, analogous to the so-called “catastrophe” events of MT depolymerization in which bound GTP is hydrolyzed immediately. Increased activity of ATP-consuming transporters that reestablish ionic balance and, notably, elevated Na\textsuperscript{+}/K\textsuperscript{−}-ATPase activity by high [K\textsuperscript{−}]), also may contribute to decreased [ATP]. Interestingly, our data show that although [ATP], is reduced similarly immediately after NaCl or glucose challenge, it recovers completely after glucose challenge but not after NaCl challenge. This recovery after glucose challenge suggests that ATP synthesis is initially reduced by hypertonicity-induced cell shrinkage but that increased glucose availability helps restore [ATP]. The combined effects of fast [Cl\textsuperscript{−}]\textsuperscript{−} recovery and increased ATP availability in glucose-challenged cells may help explain the higher steady-state levels of vesicle motility achieved in some cell types (LLC-PK\textsubscript{1} and Huh7) and the fast recovery of motility in pancreatic β cells.

The recovery of ATP abundance and ΔΨ recovery precede the recovery of vesicle and microfilament motility, indicating that the effects of [Cl\textsuperscript{−}] do not result from depleted ATP synthesis alone. Perturbed ionic balance within the vesicles themselves, which could alter vesicular shape and/or recruitment of microfilament tethering complexes, may contribute to the effects of [Cl\textsuperscript{−}]\textsuperscript{−}. Indeed, recent data examining defects in endosomal CIC-family channel mutants suggest that luminal Cl\textsuperscript{−} content plays a role in endosomal trafficking beyond V-ATPase charge compensation (which facilitates endosomal acidiﬁcation), which may include fission/fusion as well as endosomal volume regulation (3). We have shown recently that autophagy is induced by hypertonic stress (13). In this regard, it would be revealing to examine how vesicle motility that is reduced by ionic imbalance affects autophagy, because this process relies on autophagosome trafﬁcking and fusion events with lysosomes/vacuoles (66). Vacuoles also contribute to protecting osmotically challenged cells by sequestrating Na\textsuperscript{+} and Cl\textsuperscript{−}. This sequestration lowers intracellular osmolality and thereby helps avert the deleterious effects of cytoplasmic salt accumulation in plants and yeast (67). It remains to be determined whether a similar accumulation of salt in the vacuoles of at least some mammalian cell types may occur following hypertonic challenge and whether this mechan-ism participates in the mammalian cellular adaptation process.

Monovalent salts such as NaCl modify the ionic strength of solutions, such as the cellular cytoplasm, and can exert stabilizing or destabilizing effects on proteins, with the net effect depending on the speciﬁc charge distribution of the protein and on the microscopic charge interactions within the protein (68–70). Electrolyte concentration also is central to the screening of surface charge–charge interactions that influence protein stability, interaction, and function. Along with molecular crowding, increased ionic strength probably explains protein misfolding by hypertonicity (5, 12, 13, 71). Similarly, it might be expected to alter the functionality of at least a subset of proteins by affecting biomolecular electrostatic interactions. It has long been known that MT polymerization is decreased at high ionic strength in vitro (26, 72). This effect would help explain our observation that hypertonicity induces transient MT depolymerization in living cells. Actin ﬁlament assembly also is affected by hypertonicity, which increases their polymerization (4). Salt modulates actin assembly and its mechanical properties through nonspeciﬁc electrostatic effects (25). Occupancy of a distinct site by cations at concentrations similar to those occurring after strong hypertonic challenge recently was found also to stiffen actin ﬁlaments (73). Our data show that [Cl\textsuperscript{−}] decreases the motility of both actin ﬁlaments and MTs and that chemical agents that decrease microﬁlament motility also reduce vesicle motility. These data link microfilament dynamics reduced by [Cl\textsuperscript{−}], to decreased vesicle motility, which might arise from a combination of effects of salt bound to microﬁlaments, ATP loss, and/or modulated activities of any of dozens of proteins that bind to MTs and actin and that regulate microﬁlament assembly/disassembly dynamics.

Analogous to cell volume that is constrained within limits that maximize biochemical reaction rates (14, 22), intracellular elec-
the Image Correlation Spectroscopy (ICS) Image plug-in suite, developed by Jay Urru (Stower’s Institute for Medical Research, Kansas City, MO) (74) was used. Two to three subtractions (80 × 90 pixels that avoided nuclei and areas of obvious whole-cell movements) per movie from three independent movies were analyzed per condition. Movie segments encompassing 10 frames (2-min timespans) were analyzed for each time point, starting 2 min before manipulation or at the indicated times. For each region and time point, 16 flow-velocity vectors were calculated by the program. To obtain the average movement of each subregion, the absolute value of the velocity vectors, defined as \( \text{velocity} = \sqrt{\text{velocity}_x^2 + \text{velocity}_y^2} \), was calculated for each vector, and the average of the 16 values was reported.

**Cell Volume Analysis.** Cells were loaded with calcein-AM (5 μM) for 15 min at 37 °C. Regions of interest were drawn around the areas within cells that remained in focus. Photobleaching/dye leak correction was performed by extrapolating baseline fluorescence loss, estimated by fitting values from the first 6 min with a one-phase exponential decay function (Y = Span × exp(-X/Time Constant) + Plateau). Background-subtracted and photobleaching-corrected calcein fluorescence, which is inversely proportional to cell volume (75), was converted to cell-volume changes. Recovery phases were fitted with an exponential association function \( Y = (R_{\text{max}} - \text{Bottom}) \times (1 - \exp(-X/\text{In} (2/R_{\text{max}} + \text{Bottom})). \) Results are expressed normalized to baseline. RVI was delayed by treating cells with benzamil (10 μM), bumetanide (50 μM), and gadolinium (100 μM).

**Measurement of Relative Changes in \([Cl^{-}]_i\) by MQAE Analysis.** Cells were loaded with 5 mM MQAE for 60 min at 37 °C, and 5 μM calcein-AM was added during the last 15 min. Cells were alternately excited at 360 nm/488 nm, and emission was collected at 530 nm to visualize MQAE or calcine, respectively. Photobleaching correction was performed as described above on MQAE and calcein signals separately, before the \( F_{\text{calc}}/F_{\text{ref}} \) ratio was calculated. To transform ratio values into actual \([Cl^{-}]_i\) values, three independent calibration curves were constructed as described previously (59) and averaged. Briefly, cells loaded with MQAE/calcein-AM were sequentially exposed to solutions containing TBTN and 0, 30, 50, 80, 110, or 130 mM Cl\(^-\), in which clorurate was used to replace Cl\(^-\) as the counterion to maintain osmolarity, for 5 min each. The background-subtracted and photobleaching-corrected average ratio values within the middle of the 5-min exposure time frame were plotted against the known Cl\(^-\) concentration values of each calibration solution. The resulting curve then was fitted with a one-phase exponential decay function (which gave the best fit), and the equation was used to transform ratio values into mM (Cl\(^-\)) values.

**Analysis of Cellular Glucose, ATP, and Mitochondrial Function.** To compare glucose-influx kinetics of LLC-PK\(_1\) and MIN6 cells, cells transfected with the FRET-based glucose sensor FLI12Pglu-700Delta6 (76) were incubated in glucose-free Hank’s medium for 10 min before imaging. Images were collected using 440 nm excitation and alternate 485/535 nm emission as the solution was changed to an iso-osmolar Hanks’ solution containing 5 mM glucose, a change previously shown to lie within the middle of the dynamic range of the probe (76). Changes in cellular ATP levels were measured using a similar excitational emission protocol, using the FRET-based ATeam probe (60) and by fluorometric analysis using the ATP Assay Kit (Abcam), in which the excitation/emission fluorescence (535/587 nm) was measured using a SpectraMax Paradigm microplate reader (Molecular Devices). Mitochondrial depolarization was measured as an increase in the background-subtracted 488/568 nm JC-1 ratio. Mitochondrial morphology was quantified using Mitotracker Red CMXRos as described (77). Data from at least 100 cells per condition were collected in each experiment. For experiments performed with oligomycin, glucose was replaced by 2-DG (10 mM).

**Western Blot Analysis and F-Actin/G-Actin Assay.** Cells lysates were prepared and quantified as described previously (78). Actin polymerization was quantified by determining the Triton X-100-soluble (G-actin)/Triton X-100-insoluble (F-actin) ratio. (See SI Materials and Methods for details) or by a TRITC-phalloidin methanol extraction method (79) adapted to 96-well format. F-actin content values are expressed as background-subtracted fluorescence values normalized to nontreated controls.

**Statistics.** Results are given as the mean ± SEM from \( n \) independent experiments. Each experiment was performed on cells from the same passage. All experiments were performed at least three times. Statistical differences were assessed by one-way ANOVA, and curve fitting was performed using GraphPad Prism software. \( *p \leq 0.05 \).

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