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Cell volume regulation: osmolytes, osmolyte transport, and signal transduction

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Abstract In recent years, it has become evident that the volume of a given cell is an important factor not only in defining its intracellular osmolality and its shape, but also in defining other cellular functions, such as transepithelial transport, cell migration, cell growth, cell death, and the regulation of intracellular metabolism. In addition, besides inorganic osmolytes, the existence of organic osmolytes in cells has been discovered. Osmolyte transport systems—channels and carriers alike—have been identified and characterized at a molecular level and also, to a certain extent, the intracellular signals regulating osmolyte movements across the plasma membrane. The current review reflects these developments and focuses on the contributions of inorganic and organic osmolytes and their transport systems in regulatory volume increase (RVI) and regulatory volume decrease (RVD) in a variety of cells. Furthermore, the current knowledge on signal transduction in volume regulation is compiled, revealing an astonishing diversity in transport systems, as well as of regulatory signals. The information available indicates the existence of intricate spatial and temporal networks that control cell volume and that we are just beginning to be able to investigate and to understand.

General introduction

For a number of years, cell volume regulation in mammalian cells has been considered of minor physiological importance since overall osmolality of the plasma is one of the very tightly regulated parameters of the body. In the meantime, however, it has been realized

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that quite a variety of tissues such as renal medullary cells and chondrocytes are exposed to anisotonic extracellular media and thus require volume regulatory mechanisms. Anisotonic conditions can also arise in pathological conditions such as hypo- or hypernatraemia when homeostatic functions of the body are insufficient (Law 1999; Law 1998; Verbalis et al. 1989; Verbalis 1994). Similarly, cells encased in a rigid surrounding, such as brain cells, depend on cell volume regulation for their proper function. Furthermore, even at normal extracellular osmolalities cells can generate transmembrane osmotic differences due to uptake of (organic) osmolytes or during transepithelial transport of solutes (McCarty and O'Neil 1991).

Cell volume changes also occur globally during maturation of erythrocytes, cell growth, differentiation, hypertrophy, and apoptosis. Locally, cell migration and shape changes require volume adaptations (Lang et al. 1998b). Finally, cell volume has been identified as a mechanism that regulates cell metabolism (Lang et al. 1989). This effect is particularly evident in the liver, where cell swelling increases protein and glycogen synthesis and cell shrinkage increases protein and glycogen breakdown (Häussinger 1996; Häussinger 1998). Thus, cell volume and its importance during the whole life cycle of a cell has become more and more evident and this area of physiology and pathophysiology has been attracting an increasing number of investigators.

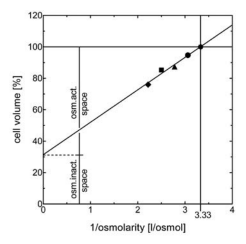
During recent years, it also became more and more accepted that not only inorganic osmolytes such as K⁺ and Cl⁻ are employed to restore osmotic equilibrium across the cell membranes but that also organic osmolytes play a significant role in cell volume homeostasis.

In most reviews, inorganic and organic osmolytes and their respective volume regulatory transporters are dealt with separately. The current review combines the two and discusses the question of a putative identity of some systems. This contribution also takes into account the progress made in identifying the molecular entities of the transporters by molecular biology techniques. Furthermore, the current state of knowledge with regard to the regulation and coordination of the various osmolyte transporters is reviewed in order to shed some light on the intriguing signal transduction networks used in cell volume regulation.

The osmometric behavior of cells

Any changes in cell volume of animal cells under anisotonic conditions are, in principle, based on a distinct permeability of the plasma membrane to water. In many instances, this permeability is increased by specialized membrane proteins, which mediate the facilitated diffusion of water, so-called water channels or aquaporins (Maunsbach et al. 1997; Marinelli and LaRusso 1997; Ma and Verkman 1999). Given a sizeable water permeability at its outer membrane, a cell then exhibits an "osmometric" behavior as its first passive response to anisotonicity, i.e., a movement of water will occur, which changes cell volume until the difference in osmotic pressure is equalized and a new chemical equilibrium of water across the plasma membrane is achieved. From a physicochemical point of view, however, a cell is not a *perfect* osmometer, which means that its volume will not change to the same extent as is predicted by the law of Boyle/van't Hoff. This means that apparently some 20%–40% of total cell volume usually comprises a phase that is not "cytosolic," which means that it is not osmotically active (Lucke and McCutcheon 1932). In practice, the osmotically active/inactive space of a living cell can be easily determined graphi-

Fig. 1 With all Na⁺ importers of RVI blocked (i.e., with 10⁻³ mol/ l amiloride plus 10⁻⁴ mol/l furosemid; see text for details), the passive changes of rat hepatocyte volumes after 10 min exposure to hypertonicity (327, 360, 400, and 450 mOsm/l) are a linear function of the reciprocal of extracellular osmolarity. The value of 3.33 is equivalent to 300 mOsm/l where cell volumes equal 100% (control, ★). The intercept of the regression line with the volume axis is equivalent to the "osmotically inactive space" that amounts to 31.3% of total cell volumes (from Wehner and Tinel 2000)



cally by plotting its relative volume versus the reciprocal of extracellular osmolarity, which is equivalent to a Boyle/van't Hoff-Plot (see Fig. 1).

The intercept of the regression line with the Y-axis then gives the osmotically inactive space; its difference from the 100% mark is equivalent to the osmotically active part. This type of analysis, which deals with the passive osmotic behavior of a cell, however, is only valid if the active compensatory processes by which cells readjust their volume are not activated and/or are inhibited efficiently. In rat hepatocytes, for instance, this was achieved by blocking all Na⁺ import mechanisms mediating regulatory volume increase (RVI) (namely Na⁺ conductance, Na⁺/H⁺ antiport, and Na⁺-K⁺-2Cl⁻ symport) by 10⁻³ Mol/l amiloride and 10⁻⁴ Mol/l furosemide and by then determining the actual changes of cell volume after 10 min of cell shrinkage, under four different hypertonic conditions. From these measurements, an osmotically inactive space could be determined that amounts to some 31% of total cell volume (Wehner and Tinel 2000).

Inorganic and organic osmolytes

For a number of years, research on volume regulation in mammalian cells has focused on the movement of the main intracellular inorganic osmolytes such as potassium and chloride. Compensatory changes in these osmolytes form indeed the backbone for volume regulation when only small perturbations of the intra- or extracellular osmolarity occur.

In recent years it has become, however, more and more evident that—similar to other phyla in nature—in mammalian cells also organic osmolytes are employed in volume regulation (Yancey et al. 1982; Hoffmann et al. 1988; Kinne 1993; Burg 1995; Burg 1996; Junankar and Kirk 2000).

Organic osmolytes comprise polyols such as sorbitol and myo-inositol, methylamines such as glycerophosphorylcholine and betaine, and amino acids such as taurine, glutamic acid, and \(\mathcal{B}\)-alanine amongst others (Yancey and Burg 1989; Yancey et al. 1982; Beck et al. 1985; Beck et al. 1998).

There are several hypotheses as to why in some cells organic solutes are used in addition to inorganic solutes. One hypothesis is the "compatible osmolyte hypothesis" which

is based on the observation that high concentrations of inorganic salts such as NaCl or KCl perturb the function of enzymes or other proteins (Noulin et al. 1999), whereas organic solutes do not (Yancey et al. 1982). Another hypothesis reflects the "counteracting osmolyte principle," which refers to the finding that methylamines attenuate the destabilizing effect of high concentrations of urea on protein function (Burg 1996). A third aspect is that several organic osmolytes are electroneutral and can replace inorganic osmolytes which, when released across the cell membrane, may change the membrane potential and thereby neuronal excitability in the brain (Iwasa et al. 1980) or driving forces for electrogenic sodium-cotransport systems. Again, the brain is particularly interesting because of the existence of reuptake systems for excitatory amino acids that are transported by sodium-cotransport systems (Curtis and Johnston 1974).

One of the major places for organic osmolyte accumulation is the renal medulla, where accumulation occurs because of a broad range of extracellular osmolalities exceeding normal osmolality (for review see Beck et al. 1985; Bagnasco et al. 1986; Yancey and Burg 1989; Garcia-Perez and Burg 1991; Kinne et al. 1993; Kinne 1993; Kinne et al. 1995; Kinne 1998; Grunewald and Kinne 1999; Kinne et al. 2001) in particular in the direction of hyperosmolality (Grunewald et al. 1993a; Grunewald et al. 1994; Handler and Kwon 2001). Also chondrocytes (de Angelis et al. 1999; Hall and Bush 2001) encounter hyperosmolality in the extracellular space because of the high concentration of fixed charges in the mucopolysaccharides in which they are embedded. Limited expandability in the brain (Strange 1992; Pasantes-Morales et al. 1994a; Pasantes-Morales et al. 1994b; Pasantes-Morales et al. 2000b) and regulation of cell transparency in the lens (lens fiber cells; Burg and Kador 1988; Cammarata et al. 2002), also require organic osmolytes.

Intracellular accumulation of organic osmolytes involves mainly two processes. The first represents uptake across the cell membrane by specific transport systems, the second intracellular generation of the osmolyte by metabolic reactions. Examples of the former are sodium-cotransport systems for myo-inositol, taurine, and betaine which use the sodium gradient across the plasma membrane as driving force. The general and specific properties of these cotransport systems are summarized in the section entitled "Organic osmolytes in RVI."

Metabolic reactions are primarily responsible for the generation of sorbitol from D-glucose and glycerophosphorylcholine (GPC). D-glucose taken up by the cells via uniport or symport systems can be readily converted into sorbitol by the enzyme aldose reductase using NADPH as cofactor (Gabbay 1973; Ohta et al. 1990). In an extensive study on renal inner medullary aldose reductase, the affinity for D-glucose was found to be about 370 mMol/l, the affinity for NADPH was 7.5 µMol/l. Thus, in the intact cell with a NADPH concentration of about 0.4 mMol/l the enzyme is saturated with NADPH, but only to a small extent with D-glucose. Due to the high $K_{\rm m}$ of the enzyme, in the intact cell sorbitol synthesis is characterized by the affinity of the cellular glucose uptake systems. For example, a value of about 60 mMol/l is obtained in intact IMCD cells, which reflects the $K_{\rm m}$ usually observed for glucose uniporters GLUT 1 or GLUT 2 (Grunewald et al. 1993b). Regulation of the synthesis occurs mainly by altering the $V_{\rm max}$ of the enzyme; a strong increase is observed in a variety of cells when they are exposed to hypertonic conditions in vivo or in vitro (Bedford et al. 1987; Bagnasco et al. 1988; Grunewald et al. 1998; Lang et al. 1998a). However, in disease states such as diabetes, the NADPH/NADP ratio is another important regulator of the rate of sorbitol synthesis (Grunewald et al. 1993b).

Sorbitol breakdown yielding fructose can occur via the sorbitol dehydrogenase pathway with NAD as the hydrogen acceptor. In some tissues that use sorbitol as organic osmolyte, these pathways are restricted to different, sometimes closely associated cells (Grunewald et al. 1995; Kinne et al. 1997). In general, the activity of sorbitol dehydrogenase is much lower than that of aldose reductase, thereby limiting the importance of this enzyme in osmotic adaptation (Grunewald et al. 1995; Kinne et al. 1997; Grunewald et al. 1998).

The major pathways involved in the metabolism of GPC in IMCD cells, for example, are the following. The precursor of GPC, choline, is taken up by sodium-independent transport systems. Choline is phosphorylated and then incorporated into the phospholipid phosphatidylcholine (PC). By a stepwise removal of the fatty acid residues (involving phospholipase A2), first lysophosphatidylcholine and then GPC are generated. Interestingly, this reaction seems to use a pool of PC different from that present in the plasma membrane. Breakdown of GPC is mediated by the GPC: choline diesterase reaction yielding choline and phosphoglycerol. The latter two reactions seem to be slow compared to the synthetic pathway. The predominant "osmosensitive" enzyme is the GPC: choline diesterase since studies in vivo and in vitro indicate that after longterm exposure of cells to hypertonic conditions, the rate of enzymatic breakdown is reduced and thus the overall concentration of the organic osmolyte is increased (Zablocki et al. 1991; Bauernschmitt and Kinne 1993).

The release of organic osmolytes from the cells involves channel-like proteins, in some instances organic and inorganic osmolytes share the same transporter. Thus, a complex picture emerges in which inorganic and organic osmolyte levels have to be controlled in a well-coordinated manner (Burg 1996). This coordination requires feedback systems that have to be elucidated and taken into account when considering volume regulatory processes in the cells.

Regulatory volume increase

Role of inorganic osmolytes in RVI

In order to achieve RVI, the intracellular osmolyte content has to be augmented rapidly. To this end, transport systems for inorganic osmolytes are activated as the first cellular response. The main osmolyte taken up by the cells is sodium since favorable driving forces for this cation exist across the plasma membrane. Sodium is subsequently exchanged against potassium by the action of the Na⁺,K⁺-ATPase, to restore the original sodium gradients and its electrochemical potential.

There are also mechanisms set in place that reduce loss of intracellular potassium and sometimes also chloride, depending on its electro-chemical equilibrium.

Na⁺/H⁺ antiport

Na⁺/H⁺ antiporters (NHEs) catalyze the secondary active and electroneutral exchange of H⁺ against Na⁺. With one exception, they are most important for the regulation of cell pH, but some NHEs are effective mediators of RVI. Six NHE isoforms have been cloned to date (NHE1–NHE6) which exhibit a common molecular organization (see for review

Orlowski and Grinstein 1997; Counillon and Pouyssegur 2000; Ritter et al. 2001). They consist of two main functional domains with amino acid identities of 45%-60% and 25%-35%, respectively. These are an N-terminal portion of 10-12 transmembrane regions (depending on the hydropathy algorithm used) and a large intracellular C-terminal part that is equipped with regulatory sites (see below). NHE1 to NHE5 are located in the plasma membrane, while NHE6 is sorted to the mitochondrial inner membrane. NHE1 and NHE6 are ubiquitously expressed whereas NHE2-NHE5 are restricted to specific tissues. In epithelia, NHE1 appears to be largely confined to the basolateral membrane (Coupaye-Gerard et al. 1996) but, in cell lines like OK (opossum kidney) or MDCK (Madin-Darby canine kidney), it may also occur apically (Noel et al. 1996). NHE2, NHE3, and NHE4 are found preferentially in the gastrointestinal tract and kidney where they appear to reside mainly in the apical membrane (Soleimani et al. 1994; Dudeja et al. 1996; Noel et al. 1996; Hoogerwerf et al. 1996; Sun et al. 1997; Bookstein et al. 1997). The presence of NHE2 in the kidney and NHE4 in the intestine, however, is discussed controversially (Bookstein et al. 1997). NHE5 was detected in nonepithelial tissues, preferentially in the brain, but also in spleen, testis, and skeletal muscle (Klanke et al. 1995). In many cells, several NHE isoforms are expressed and assigned to specific membrane domains so that (together with other transporters) they achieve a concerted cross-talk in the vectorial movement of ions; this especially holds for epithelia (Ritter et al. 2001).

Some NHEs are inhibited by submillimolar concentrations of amiloride and, more selectively, by its derivatives dimethyl-amiloride and ethyl-isopropyl-amiloride (Tse et al. 1994; Counillon and Pouyssegur 2000). There are significant differences in sensitivity to these blockers among the different isoforms, which (for amiloride) amount to almost four orders of magnitude. The amiloride sensitivity generally follows the order NHE1 > NHE2 >> NHE3 > NHE4 (Orlowski and Grinstein 1997; the latter two isoforms are therefore sometimes referred to as being amiloride-insensitive (Counillon and Pouyssegur 2000). The mitochondrial isoform NHE6 exhibits a rather low sensitivity to amiloride, but it is efficiently blocked by the analog, benzamil (Brierley et al. 1989).

With respect to their role in cell volume regulation, it was found that NHE1, NHE2, and NHE4 are activated under hypertonic conditions, whereas NHE3 is inhibited (Orlowski and Grinstein 1997; Hoffmann and Mills 1999; Ritter et al. 2001). NHE1 is clearly the Na⁺/H⁺ antiport most commonly employed in RVI, which is not surprising in regard to the ubiquitous expression of the protein. As a matter of fact, the shrinkage-induced activation of Na⁺/H⁺ antiport that had been reported in a variety of preparations prior to the molecular definition of NHE isoforms can now in many instances be attributed to NHE1 (Lang et al. 1998a; Lang et al. 1998b; Hoffmann and Mills 1999).

Concerning regulation, hypertonicity appears to shift the pH_i sensitivity of NHE1 to more alkaline values (Grinstein et al. 1985). Interestingly, although the C-terminal domain of NHE1 exhibits a number of phosphorylation sites (that are actually used for the activation by hormones and mitogens), no direct phosphorylation of the protein occurs upon cell shrinkage (Sardet et al. 1990; Sardet et al. 1991; Grinstein et al. 1992). Rather, there appears to be a significant hypertonicity-induced cross-talk between NHE1, stress fibers and the small GTPase Rho so that osmotically-induced changes in the actin texture (also involving integrins) are very likely to be part of the signaling cascade-mediating activation (Watson et al. 1992; Grinstein et al. 1993; Vexler et al. 1996; Hooley et al. 1996; Tominaga et al. 1998; Tominaga and Barber 1998). In addition, Ca²⁺ may participate in the stimulation because the C-terminal domain of NHE1 contains two calmodulin binding sites and deletion or mutation of the high affinity binding site significantly reduces the

osmotic sensitivity of the transporter (Bertrand et al. 1994). It is worth mentioning that NHE1 is also subject to transcriptional osmotic regulation (Ritter et al. 2001).

The hypertonic activation patterns of NHE2 are less well-defined. Proline-rich regions in the C-terminal portion of the protein appear to be involved which resemble SH3 binding domains and might reflect sites of interaction with the cytoskeleton and/or signaling molecules. Truncation experiments, however, revealed that these regions are probably involved in the proper targeting of NHE2 rather than its regulation (Chow et al. 1999). Besides this, NHE2 is subject to transcriptional control (Soleimani et al. 1994; Bai et al. 1999).

NHE3 is inhibited under hypertonic conditions. This effect is due to a reduction of the maximal transport velocity rather than to changes in substrate affinities (Nath et al. 1996). NHE3 is highly sensitive to the actin texture of cells (Kurashima et al. 1999; Szaszi et al. 2000). The protein associates with two regulatory factors, namely NHERF1 and NHERF2, and NHERF1 could be shown to bind to ezrin, an actin-binding protein of the ezrin-radix-in-moesin (ERM) family that links membrane proteins to the cytoskeleton. Based on deletion experiments, it could also be shown that the same C-terminal region of NHE3 mediates actin sensitivity and NHERF binding (Kurashima et al. 1999).

Moreover, among the small GTPases of the Rho family controlling actin assembly, the inhibitory form of RhoA (but not Rac1 and Cdc42) greatly depressed NHE3 activity and comparable effects were observed upon a specific block of Rho kinase (ROK), a downstream effector of RhoA; furthermore, inhibition of ROK reduced the phosphorylation of myosin light chain (MLC). These data strongly suggest that the RhoA-ROK signalling pathway is a mechanism for the control of NHE3 activity, which is, at least in part, achieved by controlling the phosphorylation of MLC and, consequently, the organization of the actin cytoskeleton (Szászi et al. 2000).

Interestingly, considerable amounts of NHE3 are present in recycling endosomes (Ritter et al. 2001). This predominantly intracellular location may contribute to its paradoxical behavior during RVI.

NHE4 is clearly activated under hypertonic conditions; it exhibits a bell-shaped profile with maximal functionality close to 500 mOsm/l (Bookstein et al. 1994). This relatively low osmotic sensitivity is readily explained in terms of NHE4 tissue distribution with maximal expression found in the renal medulla (Bookstein et al. 1994; Ritter et al. 2001) where a high extracellular osmolality prevails.

Na^+ - K^+ - $2Cl^-$ symport

NKCC1 and NKCC2 are mediators of electroneutral Na⁺, K⁺, Cl⁻ cotransport across cell membranes at a stoichiometry of 1:1:2 (with very few exceptions; see Haas and Forbush, III 2000; Russell 2000 for reviews). On the molecular level, they share many similarities with other members of Cl⁻-dependent cation transporters, namely the Na⁺-Cl⁻ symporter (NCC) and the four isoforms of K⁺-Cl⁻ symporters (KCC1 to KCC4) cloned so far. These similarities include molecular weights in the range of 110 kD to 130 kD (deglycosylated), 12 predicted transmembrane regions, and large hydrophilic intracellular N- and C-terminal domains. The most conserved regions of these transporters are the transmembrane domains, as well as the putative intracellular loops connecting them, particularly the one between TM2 and TM3. NKCCs are blocked by the "loop" diuretics bumetanide and furosemide at micromolar concentrations; KCCs exhibit a lower affinity to these compounds, whereas a possible inhibition of NCC remains ambiguous (Russell 2000). NKCC1 exhibits

a broad tissue distribution and is found in many secretory epithelia (where it resides in the basolateral membrane), as well as in a variety of nonepithelial cells. In contrast, NKCC2 is only present (apically) in the thick ascending limb of Henle's loop and in the macula densa of the kidney (Haas and Forbush, III 2000; Russell 2000).

NKCC1 is one of the most routinely employed transporters of cell volume regulation (Russell 2000) during RVI (Lang et al. 1998a; Lang et al. 1998b; Hoffmann and Mills 1999; Haas and Forbush, III 2000). Activation of NKCC1 appears to involve phosphorylation at serine/threonine residues and cell shrinkage results in NKCC1 phosphorylation in a number of cells. In most instances, however, the actual kinase in charge remains to be identified (Hoffmann and Mills 1999; Haas and Forbush, III 2000). Recently, the phosphorylation of NKCC1 by c-Jun NH₂-terminal kinase (JNK), a member of the mitogen-activated protein kinase (MAPK) family, was reported from bovine aortic endothelial cells that were shrunken under hypertonic or isotonic conditions (Klein et al. 1999). In rat hepatocytes, the hypertonic activation of Na⁺-K⁺-2Cl⁻ symport (as well as that of Na⁺ conductance) was inhibited by staurosporine, as well as by the PKC specific blocker bis-indolylmaleimide I (BIM; Heinzinger et al. 2001). In human tracheal epithelial cells, the hypertonic activation of NKCC1 appeared to be mediated by PKC- δ ; this process is also likely to involve the (extracellular signal-regulated) kinase ERK (Liedtke and Cole 2002). In addition to phosphorylation and dephosphorylation, the activity of NKCC1 appears to be regulated by the state of the actin network, as well as by accessory proteins that remain to be characterized (Haas and Forbush, III 2000; Russell 2000).

Cation channels

Compared to the symport system discussed above, the rates of ion transport by channels are some 4–5 orders of magnitude higher. Accordingly, any modulation of channel activity in response to changes of cell volume will serve as a fast and very efficient regulatory mechanism.

If one considers the electrochemical driving forces for Na⁺, K⁺, and Cl⁻ transport across most cell membranes, the activation of Na⁺-selective channels and conductive Na⁺ entry would be a highly efficient mediator of RVI. The resultant depolarization of membrane voltage would favor a parallel conductive entry of Cl⁻. Na⁺ accumulating inside the cell would then be extruded via Na⁺, K⁺-ATPase so that constant driving forces for Na⁺-coupled cotransporters are ensured. In sum, a net intracellular gain of K⁺ and Cl⁻ and thus a rapid increase of cell volume would be achieved by these mechanisms.

There is an increasing number of systems from which the hypertonic activation of cation channels is reported. Two main classes of channels can be distinguished based on their sensitivity to amiloride. In the following, first the amiloride sensitive channels will be discussed.

The activation of conductive Na⁺ entry as a mechanism of RVI was originally proposed by Hoffmann (Hoffmann 1978) and Okada (Okada and Hazama 1989) for Ehrlich ascites tumor cells and the Intestine 407 cell line, respectively. In 1995, a hypertonic stimulation of cell membrane Na⁺ conductance was reported from current-clamp recordings on rat hepatocytes in confluent monolayer culture (Wehner et al. 1995; see Fig. 2).

Furthermore, in a quantitative study, it could be shown that the relative contribution of Na⁺ conductance, Na⁺/H⁺ antiport, and Na⁺-K⁺-2Cl⁻ symport to the initial uptake of Na⁺ under hypertonic stress was approximately 4:1:1 (Wehner and Tinel 1998). This clearly

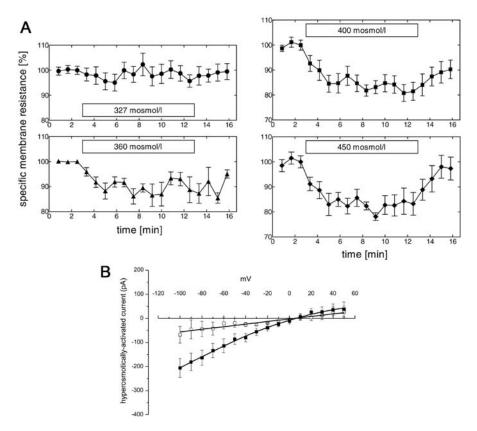


Fig. 2 A Cable analysis of specific membrane resistance, reflecting the hypertonic activation of Na⁺ conductance in rat hepatocytes. Experiments were carried out in the continuous presence of 0.5 mM quinine; extracellular osmolarity was increased from 300 mOsm/l to the osmolarities indicated; means±SEM; n=4–5. **B** Current-voltage relationships of hypertonicity-induced membrane currents in rat hepatocytes obtained with (two-channel microelectrode) voltage-clamp techniques. The differences between currents obtained at 400 mOsm/l and 300 mOsm/l are depicted for cells injected with control-oligo-DNA (\blacksquare) or anti-α-rENaC oligo DNA (\square) (from Wehner and Tinel 2000; Böhmer and Wehner 2001)

renders Na⁺ conductance the prominent mechanism of RVI in this system. The Na⁺ conductance was inhibited by amiloride with an apparent K_i of 6 μ Mol/l and its overall sensitivity profile was EIPA > amiloride > benzamil (Wehner et al. 1997; Böhmer et al. 2000). Hence, at first sight, the hypertonicity-induced Na⁺ conductance of rat hepatocytes may reflect a low (amiloride) affinity type rather than an epithelial Na⁺ channel (ENaC); the latter typically exhibits K_i values in the upper nanomolar range and an inverse pharmacological profile (Garty and Palmer 1997; Fyfe et al. 1998). In voltage-clamp experiments on rat hepatocytes, a relatively low cation selectivity of the channel was detected with a P_{Na}/P_K of 1.4 (Böhmer and Wehner 2001). In patch-clamp experiments in the cell-attached configuration, hypertonicity-induced single channel events with a unitary conductance of 6 pS were recorded (P. Lawonn and F. Wehner, unpublished observation). With respect to intracellular signaling events, the hypertonic activation of Na⁺ conductance (and Na⁺-K⁺-2Cl⁻ symport, but not Na⁺/H⁺ antiport) is mediated by PKC (Heinzinger et al. 2001).

In the human hepatoma cell-line HepG2, hypertonic stress led to the activation of a nonselective cation conductance that was clearly sensitive to amiloride with 65% inhibition at 10⁻⁵ Mol/l (Wehner et al. 2002a). Interestingly, this hypertonicity-induced conductance was also partially inhibited by Gd³⁺ (10⁻⁴ Mol/l) and flufenamate (10⁻⁵ Mol/l) which are typical blockers of nonselective but amiloride-insensitive cation channels in a variety of systems (see below). Benzamil and EIPA (at 10⁻⁵ Mol/l each) had no effect on HepG2 cation conductance (Wehner et al. 2002a). In Ehrlich-Lettre ascites tumor cells, hypertonic conditions activated a cation conductance that did not discriminate between Na+, K+, and Li⁺, but that was impermeable to NMDG and choline. In the range of 10⁻⁶ Mol/l to 10^{-3} Mol/l this cation conductance was blocked by the ion channel inhibitors benzamil > Gd³⁺ > amiloride > EIPA (in order of potency; Lawonn et al. 2003). In immortalized human nonpigmented ciliary epithelial cells, shrinkage upon return from hypo-osmotic to normosmotic conditions was followed by a partial (post-RVD) RVI (Civan et al. 1996) that was mediated by the parallel operation of Na⁺/H⁺ antiport, Na⁺-Cl⁻ symport, Na⁺-K⁺-2Cl⁻ symport, as well as by conductive Na⁺ entry. In the range of 10⁻⁶ Mol/l and 10⁻⁵ Mol/l, this Na⁺ conductance was significantly inhibited by amiloride and—even more efficiently-by benzamil, the most effective blocker of the ENaC (Civan et al. 1997; Garty and Palmer 1997; Fyfe et al. 1998). In principal cells of rat cortical collecting duct hypertonic stress led to a distinct depolarization of membrane voltage coinciding with an increase of cell Na⁺ (Schlatter et al. 1997). These effects were partially inhibited by 10⁻⁵ Mol/l amiloride or Gd³⁺ and the effects were additive. This result was interpreted in terms of a parallel activation of Na⁺ channels and nonselective cation channels.

In human red blood cell ghosts, hypertonic stress induced a cation conductance that appeared to be equally permeable to Na⁺ and K⁺ (but impermeable to NMDG) and that was partially inhibited by 10⁻⁴ Mol/l amiloride (Huber et al. 2001). These findings support earlier flux studies on lamprey erythrocytes in which hypertonic stress led to the activation of an amiloride-sensitive Na⁺ transport that could not be attributed to Na⁺/H⁺ or Na⁺/Na⁺ antiport (Gusev and Sherstobitov 1996).

In U937 macrophages, hypertonic stress led to the activation of an inward current with a distinct selectivity for Na⁺ over K⁺ that was inhibited by amiloride with an apparent K_i close to 1 μ Mol/l. At the single channel level, this cation conductance appeared to be related to a 6 pS channel. Interestingly, these channels are upregulated by pretreatment of the cells with glucocorticoids, which are known to regulate macrophage function (Gamper et al. 2000).

The shrinkage-activated amiloride sensitive cation channel appears to be related to (α) ENaC. In rat hepatocytes, antisense nucleotides attenuated the activation of the channel (Böhmer and Wehner 2001; see Fig. 2B). In human ciliary body, α -ENaC mRNA was identified (Civan et al. 1997). Similarly, mRNA for α , β , and γ subunits was detected in rat hepatocytes and these subunits could also be identified on the protein level (Böhmer and Wehner 2001). Although the nonselectivity of hypertonicity-induced channels is quite in contrast to the high P_{Na}/P_{K} reported for α -, β -, γ -rENaC (which is as high as 40; Fyfe et al. 1998) this is not contradictory per se to a possible contribution of ENaCs because their biophysical properties (also including single-channel conductance) strongly depend on subunit composition (Garty and Palmer 1997) as well as their relative affinity to amiloride and its derivatives (Benos et al. 1997; Garty and Palmer 1997). Furthermore osmo-sensitivity of α -, β -, γ -ENaC was reported when the channel subunits were expressed in oocytes (Awayda and Subramanyam 1998). Since some of the results are, however, conflicting,

further studies are necessary to clarify the relation between ENaC and the amiloride-sensitive cation channels.

The second group of hypertonicity-induced cation channels is clearly amiloride-insensitive up to concentrations of 10⁻⁴ Mol/l. Quite typically, however, these channels are inhibited by the anti-inflammatory drug flufenamate (10⁻⁴ Mol/l), as well as by Gd³⁺ in the range of 10⁻⁵ to 10⁻³ Mol/l. They are expressed in human nasal epithelial cells (Chan and Nelson 1992), in the human colon cell-lines CaCo-2 and HT₂₉ (Nelson et al. 1996; Koch and Korbmacher 1999), in the mouse cortical collecting duct cell line M1 (Volk et al. 1995), as well as in BSC-1 renal epithelial cells (derived from the African green monkey), A 10 vascular smooth muscle cells (established from rat embryonic aorta), and Neuro-2a cells (derived from mouse neuroblastoma; Koch and Korbmacher 1999). In general, these channels are rather nonselective with respect to monovalent cations, although there are certain peculiarities. In human nasal epithelium and in CaCo-2 cells, for instance, channels did not discriminate much between Na+, K+, and Cs+, but exhibited a significantly lower permeability to Li⁺ and, in the latter system, there was also a distinct permeability to NMDG (P_{Na}/P_{NMDG}=0.56; Chan and Nelson 1992; Nelson et al. 1996). In contrast, in the M1 cell-line the channel appeared to be equally permeable to Na⁺, K⁺, Cs⁺, Li⁺, and Rb⁺ but virtually impermeable to NMDG (Korbmacher et al. 1995; Volk et al. 1995). In addition, there appear to be significant differences in the relative permeabilities of amilorideinsensitive channels to Cl⁻ with P_{Na}/P_{Cl} values in the range of 60 (Korbmacher et al. 1995) to 1.7 (Nelson et al. 1996). In some systems, the hypertonic activation of channels is Ca²⁺independent (Chan and Nelson 1992) whereas, in others, channel activity appears to require a minimum $(Ca^{2+})_i$ of 1 μ Mol/I (Korbmacher et al. 1995; Koch and Korbmacher 1999). Nevertheless, these channels do not appear to conduct significant amounts of divalent cations (Korbmacher et al. 1995; Koch and Korbmacher 2000). The unitary conductances of hypertonicity-induced but amiloride-insensitive channels generally is in the range of 15 pS to 27 pS (Korbmacher et al. 1995; Koch and Korbmacher 1999).

Most interestingly, at least some of these amiloride-insensitive nonselective cation channels are typically inhibited by cytosolic ATP concentrations in the millimolar range (Koch and Korbmacher 1999). This raises some concerns as to their actual role in RVI because, under most physiological conditions these channels will remain silent. Also of note in this respect, in a recent study on isolated rat colonic crypt cells, there was no detectable activation of nonselective cation channels under hypertonic stress (Weyand et al. 1998), whereas these cells are known to express Ca²⁺-activated channels which become detectable in excized inside-out patches and which are very similar to those observed in HT29 cells (Bleich et al. 1996). Clearly, whereas nonselective and amiloride-insensitive cation channels appear to be ubiquitously expressed, their actual contribution to cell volume regulation remains to be elucidated.

In principle, the activation of cation conductances—either sensitive or insensitive to amiloride—that do not discriminate much between Na⁺ and K⁺ will lead to both Na⁺ influx as well as K⁺ efflux. Because of the inside negative membrane voltage, however, overall cation gain will significantly exceed overall cation loss equivalent to an increase of the overall intracellular osmotic activity.

K⁺ channels

In most systems, the continuous channel-mediated K^+ leakage out of cells is equivalent to a significant loss of cellular osmolytes. This K^+ loss is commonly compensated for by the activation of Na⁺, K^+ -ATPase. Nevertheless, it is rather obvious that any inhibition of K^+ channels under hypertonic conditions will per se facilitate RVI and, in some cells, this mechanism appears to be an important mediator of volume regulation.

An inhibition of K⁺ conductance was originally reported for the basolateral membrane of toad and rabbit urinary bladder (Lewis et al. 1985; Donaldson et al. 1989), as well as of frog skin (Costa et al. 1987; Leibowich et al. 1988) where cells were shrunken by extracellular Cl⁻ removal (leading to a loss of Cl⁻ because of the lower membrane permeability of the Cl⁻ substitute used when compared to Cl⁻ itself) and/or by increasing extracellular osmolarity. In addition, hypertonic stress reduced basolateral K⁺ conductance in rabbit proximal tubule (Lapointe et al. 1990; Macri et al. 1997) and human nasal epithelium (Willumsen et al. 1994) and it decreased conductive K⁺ loss in MDCK cells (Ritter et al. 1991). In mouse liver, a reduction of cell membrane K⁺ conductance appeared to be the main mechanism of RVI (Graf et al. 1988; Wang and Wondergem 1991) and a transient inhibition of K⁺ channels may contribute to the volume response of rat hepatocytes (Wehner et al. 1995; Wehner and Tinel 1998). Interestingly, whole-cell recordings on freshly isolated hippocampal neurons revealed a decrease of voltage-gated K⁺ currents under hypertonic conditions (Huang and Somjen 1997). In dissociated rabbit corneal epithelial cells, a large conductance K⁺ channel (167 pS in symmetrical 150 mMol/l KCl) could be identified that appeared to mediate the decrease in whole-cell K⁺ conductance to 44% of control upon change to 130% extracellular osmolarity (Farrugia and Rae 1993). In a recent study on isolated rat colonic crypts, it was found that hypertonic stress (+50 mOsm/l and +100 mOsm/l) led to membrane depolarizations by 12 mV and 22 mV, respectively, coinciding with decreases in whole-cell conductance to 70% and 50% of control (Weyand et al. 1998). On the molecular level, these effects appeared to be correlated to a 16 pS K⁺ channel that experienced decreases in activity $(N \cdot P_0)$ to 47% and 44% when compared to isotonic conditions. Of note, this reduction of channel activity was most likely triggered by a hypertonicity-induced decrease of (Ca²⁺)_i. The channel was inhibited by (10⁻⁴ Mol/l) Ba²⁺ and (10⁻⁹ Mol/l) charybdotoxin (Bleich et al. 1996).

Anion channels

Inhibition of anion channels during RVI might directly contribute to the gain in osmolytes if the intracellular anion (Cl^- or HCO_3^-) activity is above electrochemical equilibrium. Indirectly, it restricts the movement of the counterion for K^+ flux across the membrane, thus impeding K^+ losses.

Decreases of Cl⁻ conductance under hypertonic conditions have been reported only from a limited number of preparations. In cultured human nasal epithelium, hypertonic stress reduced apical Cl⁻ conductance (Willumsen et al. 1994) whereas, in rabbit collapsed proximal convoluted tubules, the partial Cl⁻ conductance of the basolateral membrane was inhibited (Macri et al. 1997). In some instances, the decrease of Cl⁻ conductance under hypertonic conditions may reflect an inhibition of hypotonicity-induced Cl⁻ channel activation. In human vas deferens cells in primary culture, for example, with 290 mOsm/l solutions in the experimental bath, as well as in the patch pipette, there was a slowly devel-

oping increase of Cl⁻ conductance once the whole-cell configuration was achieved; this effect could be reversed by an increase of extracellular osmolarity (Winpenny et al. 1996). The increase of Cl⁻ conductance was interpreted to be due to the additional osmotic activity of intracellular macromolecules and the resultant swelling of cells.

Organic osmolytes in RVI

In most cells investigated, RVI after exposure to a hypertonic extracellular medium occurs within minutes and involves the net uptake of inorganic osmolytes. During this short-term regulation of cell volume, changes in organic osmolytes transport are generally not involved, although it has been observed that the plasma membrane permeability for organic osmolytes such as sorbitol (e.g., in IMCD cells grown at 600 mOsm/l and exposed to 900 mOsm/l medium) decreases (Bagnasco et al. 1988; Grunewald and Kinne 1989). Thus, leak pathways for organic osmolytes are downregulated to make the intracellular accumulation of organic osmolytes—by transporters or metabolic synthesis—more effective. Whether these leak pathways are identical to the ones activated during RVD remains to be investigated.

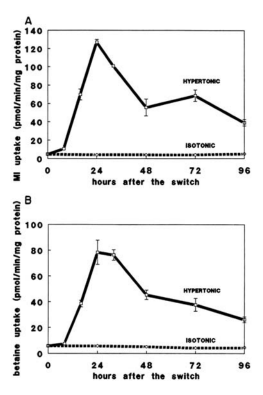
If exposure of cells to hypertonicity is extended to periods of hours or days, adaptive changes take place aimed to replace the augmented intracellular inorganic electrolytes by organic osmolytes. To this end, intracellular synthesis is increased (sorbitol and GPC) or intracellular breakdown is decreased (GPC; for review see Burg 1995; Burg et al. 1997). For taurine, betaine, and myo-inositol, the predominant effect is on the rate of uptake, which increases severalfold in a matter of hours (see Fig. 3).

Thus, strictly speaking, organic osmolytes are not directly involved in RVI; they serve, however, in the long run to maintain cellular electrolyte homeostasis. Therefore, the cellular uptake systems for organic osmolytes and their regulation are described in the following paragraphs.

Na⁺-*Cl*⁻-taurine symport

Taurine, a β -amino acid with a sulfonic acid instead of a carboxylic acid as head group occurs in plasma at concentrations of 40 µMol/l. It has several major functions: as partner in the formation of taurocholate, as neurotransmitter (Jacobsen and Smith 1968; Curtis and Johnston 1974), and as an organic osmolyte. Although several tissues, including liver and astrocytes, can synthesize taurine from cysteine, the high intracellular concentration (e.g., 20 mMol/l in IMCD cells) can only be achieved by an active uptake into the cell (Ruhfus et al. 1998). The transporter mediating this uptake is a secondary active sodiumchloride taurine cotransport system in which up to three sodium ions are translocated with one chloride ion and one taurine molecule. Taurine transport with these characteristics has been found in renal (Chesney et al. 1985a; Kinne et al. 1998; Wolff and Kinne 1988; Zelikovic et al. 1989; Zelikovic and Budreau-Patters 1999) and intestinal brush border (O'Flaherty et al. 1997), in liver cells (Warskulat et al. 1997a, Warskulat et al. 1997b; Peters-Regehr et al. 1999), in H4IIE hepatoma cells (Warskulat et al. 1997b), Ehrlich ascites tumor cells (Hoffmann et al. 1988), astrocytes (Beetsch and Olson 1996), the retinal pigment epithelium (Handler and Kwon 2001), bovine aortic endothelial cells (Qian et al. 2000), renal IMCD cells, and MDCK cells (Uchida et al. 1991). The apparent affinity

Fig. 3 Time course of basolateral myo-inositol (A) and betaine (B) uptake into MDCK cells switched into hypertonic medium. On day 0, cells cultured on filters in defined medium with 10% FBS were switched to same medium made hypertonic (500 mOsm/l) by addition of raffinose. Isotonic cells were maintained in isotonic defined medium with 10% FBS. Uptake was performed at 37°C for 30 min with 10 µMol/l myo-inositol and 10 μMol/l betaine. Results are means±SD to triplicate samples (from Yamauchi et al. 1991)



 (K_m) for taurine is in the range of 10 μM to 100 μM (Uchida et al. 1991). The transporter also transports β -alanine and, with a lower affinity, γ -amino butyric acid (GABA). Molecular cloning from MDCK cells revealed that the transporter corresponds to a protein with 655 amino acids with a relative molecular mass of 74 kDa and probably has twelve transmembrane helices (Uchida et al. 1992). It shows significant amino acid sequence similarity to the MDCK cell betaine/GABA transporter and other Na⁺, Cl⁻-dependent neurotransmitter transporters (Guastella et al. 1990). In dog tissues, the order of mRNA abundance for this transporter (dTAUT) was kidney cortex \approx kidney medulla > ileal mucosa > brain > liver > heart > epididymis. Taurine transporters cloned from other sources such as LLC-PK1 cells from pig kidney, human retinal pigment epithelium, thyroid, placenta, and bovine aortic endothelial cells (Qian et al. 2000) were of similar size (621 and 620 amino acids, respectively) and have a high amino acid sequence similarity.

In a variety of cells and tissues such as MDCK cells (Uchida et al. 1991), human and bovine lens epithelial cells (Cammarata et al. 2002), bovine aortic endothelial cells (Qian et al. 2000), rat liver macrophages (Warskulat et al. 1997a), H4IIE hepatoma cells (Warskulat et al. 1997b), primary cultures of rat hepatocytes (Warskulat et al. 1997a), rat astrocyte cultures (Beetsch and Olson 1996), and human Caco-2 cells (Satsu et al. 1999), exposure to hyperosmotic media results in an increase in $V_{\rm max}$ of taurine transport and a concomitant increase in transporter mRNA (Burg et al. 1997; Handler and Kwon 2001; Cammarata et al. 2002).

High affinity taurine transport is inhibited by phosphorylation via PKC by decreasing affinity for taurine and sodium and reducing maximal velocity (Kulanthaivel et al. 1991;

Brandsch et al. 1993; Nakamura et al. 1996; Mollerup and Lambert 1998; Lima et al. 2000; Qian et al. 2000). Furthermore, taurine uptake adaptively decreases when taurine availability is augmented (Chesney et al. 1985b; Han et al. 1998). Although these regulatory processes appear not to be linked to the response to hypertonicity, they complicate the study of the molecular mechanisms of the latter. However, it can be assumed that the mechanisms involved in osmotic adaptation are similar to the ones described below for the betaine transport system, since both systems respond in a very comparable manner.

Na⁺-Cl⁻-betaine symport

In medullary kidney cells and chondrocytes, which use betaine in volume regulation, intracellular concentrations of up to 50 mMol/l are found. Although cells usually contain choline dehydrogenase that catalyses the synthesis of betaine from choline, most of the betaine is taken up from the extracellular medium. The same holds for MDCK (Yamauchi et al. 1991) and PAP-HT25 cells (Ferraris et al. 1996). The uptake of betaine is dependent on the presence of both sodium and chloride and the transport system (Yamauchi et al. 1992; Moeckel et al. 1997) has been identified to belong to the family of brain GABA and noradrenaline transporters (Guastella et al. 1990).

Initial cloning of the transporter from MDCK cells revealed a protein of 614 amino acids with a molecular weight of 69 kDa. Its proposed membrane topology predicts 12 transmembrane helices and an intracellular NH₂ and COOH terminus (Yamauchi et al. 1992). When expressed in oocytes, the protein showed both betaine and γ-amino-n-butyric acid (GABA) transport activities that were chloride- as well as sodium-dependent (Matskevitch et al. 1999; Forlani et al. 2001). Because of these transport properties and the 43%–49% identity in the amino acid sequence (Rasola et al. 1995) with the rat brain GABA transporter and the human brain noradrenaline transporter, it was named betaine-GABA-transporter (BGT1).

Consecutive studies showed that the gene extends over 28 kilobases and consists of 18 exons. In addition, the 5' end of the gene has three different first exons, thus a complex mixture of mRNAs exists. Three main types of mRNA (A, B, and C) can be distinguished that differ considerably in their 5' untranslated sequences. Each type is expressed in a tissue-specific manner: kidney medulla (and MDCK cells) contain all three types, brain and kidney cortex, type B and C, and liver only type A (Takenaka et al. 1995; Burg et al. 1997).

Betaine transport shows apparent affinities ($K_{\rm m}$) of 0.3 mMol/l–0.5 mMol/l in MDCK cells, and in oocytes that express BGT1, the affinity for GABA is at or below 0.1 mMol/l. At 1 mMol/l substrate, the current induced in BGT 1-expressing oocytes decreased in the following order: betaine > GABA > diaminobutyric acid = β -alanine > proline. At 1 mMol/l betaine, the affinity for sodium was 93.3 mMol/l and the affinity for chloride 76.1 mMol/l (Yamauchi et al. 1992).

In electrophysiological studies, a coupling ratio of Na/Cl/betaine of 3:2:1 was found. Furthermore, these studies revealed significant kinetic differences compared to the neuronal GABA-transporter GAT-1 (Matskevitch et al. 1999).

For the symport process, a transport model of ordered binding was proposed (Matskevitch et al. 1999; Forlani et al. 2001) in which betaine binds first to the extracellular side of a transporter. Sodium binding occurs thereafter, followed by chloride binding. Translocation of betaine and sodium is already observed in the absence of chloride, but chloride

augments the translocation. As for other cotransport systems, this model would predict that betaine facilitates sodium and chloride binding to the transporter; corresponding increases in the transporter affinities at higher betaine concentrations have been observed. The exact mechanism of chloride (and sodium and betaine) translocation are unknown to date. It should be mentioned, however, that in squid motor neurons betaine activates a large Cl⁻ selective current which is sodium-dependent (Petty and Lucero 1999). This observation raises the possibility that the betaine transporter might posses chloride channel properties.

Augmented extracellular osmolality increases betaine uptake into MDCK cells (Nakanishi et al. 1990; Yamauchi et al. 1991; Kempson 1998), rabbit PAP-HT 25 cells (renal papillary epithelium; Ferraris et al. 1996), and porcine chondrocytes (de Angelis et al. 1999) at the level of the transporter by increasing $V_{\rm max}$ but not $K_{\rm m}$ of betaine uptake (see Fig. 3). The increase in uptake is preceded by an enhanced transcription of BGT1 in MDCK cells followed by increases in BGT1-mRNA. As in the case of the sodium-myoinositol transporter a "tonicity-responsive enhancer element" (designated TonE) of the gene is involved that spans -69/-50 of the sequence (Burg et al. 1997; Handler and Kwon 2001). In transgenic mice, the 5'-flanking region which contains TonE also mediates osmotic regulation of transcription in vivo (Handler and Kwon 2001). Osmoregulation of BGT1 mRNA has also been observed in renal medulla in the intact rat (Miyai et al. 1996; Moeckel et al. 1997), and in rat hepatic stellate cells (Peters-Regehr et al. 1999). Although the time course and extent of regulation of BGT-1 and SMIT by hypertonicity are quite similar, the regulatory pathways may be different at one point or another (Atta et al. 1999)

In addition to the relatively slow adaptation of betaine transport in vitro, more rapid changes have also been reported. For example, in mouse isolated perfused straight proximal tubules, basolateral betaine uptake (measured as substrate-dependent depolarization) is enhanced when the extracellular osmolality is increased (Völkl and Lang 2001). The speed of regulation suggests nongenomical processes. A similar rapid activation was observed in squid motor neurons (Petty and Lucero 1999). Whether these responses involve protein phosphatases that, potentially by dephosphorylation, could stimulate betaine transport is not clear.

Hypertonic activation of betaine uptake shows some more peculiarities. Under isotonic conditions betaine uptake into MDCK cells proceeds only across the basolateral membrane; after exposure to hypertonicity basal-lateral uptake is increased, but in addition, significant uptake across the apical membrane is observed (Yamauchi et al. 1991). This finding might be related to the existence of various isoforms of BGT1 (see above) that might differ in their cellular location.

Na⁺-Myo-inositol symport

Physiological plasma concentrations of myo-inositol range in mammals from 4.5 mMol/l to 6.6 mMol/l, whereas intracellular concentrations up to 133 mMol/l can be found in rat glial (Strange et al. 1991) and renal medullary cells (Bagnasco et al. 1986; Nakanishi et al. 1988; Wirthensohn et al. 1989; Yancey and Burg 1989; Garcia-Perez and Burg 1991; Sizeland et al. 1993; Grunewald et al. 1995; Grunewald and Kinne 1999; Handler and Kwon 2001). This large concentration difference suggests active uptake of myo-inositol into these cells. Similarly, active myo-inositol uptake is observed in rat pancreatic islets, bovine lens epithelial cells, hamster small intestine, rat mesanglial cells, rat hepatocytes, crystalline lense, rabbit peripheral nerve, retinal pigment epithelial cells, rabbit ciliary

body, isolated rat Schwann cells, and endothelial cells (for references see Porcellati et al. 1999). This uptake process involves a sodium/myo-inositol symport system (SMIT) which as secondary active transport system employs the transmembrane electrochemical potential difference of sodium for intracellular accumulation of myo-inositol. Apparent K_m values for myo-inositol range from 30 µMol/l in MDCK cells (Kwon et al. 1992) to ~94 µMol/l in rabbit renal brush border (Hammerman et al. 1980) and 104 µMol/l in rabbit TALH cells (Yorek et al. 1999). Interestingly, also D-glucose is a substrate of this transporter (Hammerman et al. 1980; Hager et al. 1995). Phlorizin, the well-known inhibitor of the sodium-D-glucose cotransporter, also inhibits the sodium-myo-inositol transporter (K_i <60 μMol/l; Strange et al. 1991; Hager et al. 1995). Similarities between the two transporters were also detected at the DNA and protein level. There is 46% amino acid identity overall and apparently similar membrane topology of the two transporters; SMIT is a polypeptide of 718 amino acids with a relative molecular mass of 79.5 kDa (Kwon et al. 1992). Tissue abundance of mRNA in dog tissues is kidney medulla > kidney cortex > brain (Kwon et al. 1992). The SMIT cloned recently from bovine lens epithelial cells shows a 92% identity with the MDCK cell transporter (Zhou et al. 1994).

The sodium to myo-inositol stoichiometry of the transporter is 2:1 (Hager et al. 1995). The stoichiometry and electrogenicity of the transporter forms the basis for the extensive intracellular accumulation of myo-inositol. Kinetic and biochemical characteristics vary from tissue to tissue and cell to cell. These variations have recently been explained by the existence of alternate splicing, which generates isoforms that differ in their intracellular protein kinase A and protein kinase C phosphorylation sites at the carboxy terminus (Porcellati et al. 1999). In epithelial cells, the cellular location can also differ; thus, in the renal proximal tubule the Na⁺/myo-inositol symport is present in the luminal brush border membrane, whereas in more distal renal cells the basolateral location prevails (see Grunewald and Kinne 1999).

SMIT is increased by augmentation of the extracellular osmolality in MDCK cells (Nakanishi et al. 1989; Yamauchi et al. 1991), glial cells (Strange et al. 1991; Kwon et al. 1992), neuronal cells (Yamashita et al. 1999), mesothelial cells (Matsuoka et al. 1999), bovine lens epithelial cells (Cammarata et al. 2002), and human retinal epithelial cells (Handler and Kwon 2001). In MDCK cells grown on filters, a 25-fold increase in uptake rate was observed within 24 h after exposure of the cells to hypertonic media (Yamauchi et al. 1991).

After restoring isotonicity, the transport rate returns to normal levels within 1 d. In both instances only the $V_{\rm max}$ of the transport is affected and $K_{\rm m}$ remains constant, suggesting a change in the number of transporters (Nakanishi et al. 1989). In similar studies in rat C6 glioma cells, myo-inositol uptake also increased after exposure of the cells to 440 mOsm/l solutions; however, the time course was slower and maximum uptake was obtained after 48 h (Strange et al. 1991). The increased uptake in response to hypertonicity is preceded by an increased abundance of mRNA for the transporter, which is the direct result of increased transcription of the gene (Yamauchi et al. 1993). Activation of transcription depends on an enhancer element named tonicity responsive enhancer (TonE). The SMITgene is regulated by multiple TonEs in its 5'-flanking region (Burg et al. 1997; Zhou and Cammarata 1999; Handler and Kwon 2001). An increase in SMIT m-RNA is also found in a TALH cell line derived from rabbit renal medulla (Yorek et al. 1999) and in mesothelial cells (Matsuoka et al. 1999). In the latter cell line, the first significant increase in mRNA could be observed already 1 h after exposure of the cells to 490 mOsm. The increase in transporter mRNA has also been observed under in vivo conditions, for example, in rat in-

ner medullary collecting duct cells when d DAVP was administered to chronically diuretic rats, a maneuver that rapidly increases the extracellular osmolality in the medulla (Burger-Kentischer et al. 1999). Injection of NaCl into rats increases SMIT-mRNA significantly within 1 h. Comparing the experiments in intact rats with cultured cells, the transporter mRNA seems to respond more rapidly in vivo than in vitro (Yamauchi et al. 1995).

Amino acid transport system A

Amino acid transport system A (System A), which mediates sodium-dependent uptake of neutral amino acids into mammalian cells, appears to be also subject to regulation by extracellular osmolality. In MDCK cells, as well as chondrocytes, an upregulation within 4–6 h of hypertonic exposure was observed, well before any change in BGT1 activity (Chen et al. 1996a; de Angelis et al. 1999; Horio et al. 1997; Kempson 1998). The increase in System A is blocked by inhibitors of RNA and protein synthesis, suggesting that an increase in the number of transporters is part of the mechanism (Kempson 1998). The adaptation of this system appears to be a relatively early response of cells exposed to hypertonicity; therefore, it deserves further investigation.

Regulatory volume decrease

Inorganic osmolytes in RVD

After cell swelling, transport systems are immediately activated that mediate the release of the major intracellular inorganic osmolytes potassium and chloride. Their transmembrane movement occurs either via separate pathways or directly coupled to each other.

K⁺ channels

Due to the outwardly directed K^+ gradient in most animal cells, any increase of K^+ channel activity will augment the conductive exit of K^+ . In addition, the increase of cell membrane K^+ conductance will hyperpolarize the cell membrane and (even if basal Cl^- conductance is not changing) this hyperpolarization will favor conductive Cl^- efflux. Likewise, if there is an initial increase of Cl^- conductance and if cell Cl^- is above electrochemical equilibrium (as it is in most systems) this will augment Cl^- exit and depolarize the cell membrane. Membrane depolarization in turn will facilitate conductive K^+ efflux. The most effective mechanism of RVD will, of course, be the parallel activation of K^+ and Cl^- channels. Because of the pronounced voltage-mediated coupling between both pathways conductive K^+ and Cl^- release may result in a quasielectroneutral mode of KCl export.

An increase of cell membrane K⁺ conductance under hypotonic conditions and its significance for RVD have been reported from a variety of preparations (see Lang et al. 1998a; Lang et al. 1998b for comprehensive reviews). In many studies, these mechanisms were analyzed by means of ⁸⁶Rb⁺ (or ⁴²K⁺) fluxes, intracellular microelectrode recordings and rapid ion-substitutions, whole-cell patch-clamp measurements, and by use of specific K⁺ channel blockers. Here, we will focus on data at the molecular level that were obtained

by means of single-channel recordings and by the cloning and expression of some of the volume-activated K⁺ channels.

 BK_{Ca} or maxi- K^+ channels exhibit large (big) unitary conductances in the range of some 100 pS to 250 pS and show under symmetrical high K^+ solutions a linear current-to-voltage relation (see Vergara et al. 1998 for review). In most instances, BK_{Ca} channels are inhibited by Ba^{++} , quinine, and TEA (tetraethylammonium), as well as by (the scorpion peptide) charybdotoxin. They are selectively blocked by (the scorpion toxin) iberiotoxin. BK_{Ca} channels are found in neurons, skeletal muscle, smooth muscle, and in epithelial cells, where they reside in the apical membrane. BK_{Ca} channels are activated by membrane depolarization and micromolar concentrations of cell Ca^{2+} . It is this Ca^{2+} sensitivity that might function as a link to a swelling-induced increase in intracellular calcium. Likewise, the voltage-dependence of BK_{Ca} channels may reflect the coupling mechanism to a hypotonicity-induced activation of Cl^- channels (see above).

 BK_{Ca} channels consist of hetero-oligomeric complexes of the pore-forming α -subunits (first cloned from Drosophila as dSlo (Atkinson et al. 1991; Adelman et al. 1992) and the auxiliary β -subunits (Garcia-Calvo et al. 1994; Behrens et al. 2002). The α -subunits are expressed in multiple splice variants generating functional diversity among different cells, but they exhibit a high degree of homology among species. Their membrane topology is related to that of voltage-gated K^+ channels (see below; Toro et al. 1998; Vergara et al. 1998; Jensen et al. 2001). They contain seven transmembrane segments (so that the N-terminus is most likely extracellular), a positively charged TM4-segment (that is probably part of the voltage sensor), and four additional hydrophobic segments forming a unique secondary structure at the C-terminal cytosolic tail that appears to mediate Ca^{2+} binding (Schreiber and Salkoff 1997).

The β -subunit is likely to function as an additional modulator of channel characteristics, e.g., with respect to Ca²⁺ sensitivity, but it does not appear to be an obligatory component of all BK_{Ca} channels. Four β -subunits have been cloned that are mainly expressed in smooth muscle, endocrine cells, epithelia, and the central nervous system (Behrens et al. 2002). β -subunits have a proposed topology of spanning the membrane twice, with N- and C-termini inside the cell (Toro et al. 1998; Vergara et al. 1998).

The activation of BK_{Ca} channels under hypotonic conditions could be demonstrated in *Necturus* and rabbit proximal tubule cells (Dubé et al. 1990; Filipovic and Sackin 1991; Kawahara et al. 1991), in principal cells of rabbit and rat cortical collecting tubule (Ling et al. 1992; Schlatter 1993; Stoner and Morley 1995; Hirsch and Schlatter 1997), in clonal kidney (Vero) cells derived from African green monkey (Hafting and Sand 2000), in acinar cells of rat lacrimal gland (Park et al. 1994), in embryonic chick hepatocytes (Pon and Hill 1997), and in guinea pig jejunal villus enterocytes (MacLeod and Hamilton 1999b). In the A_3 cell line derived from rabbit medullary thick ascending limb, hypotonic stress, as well as negative pressure applied to the patch pipette (suction), significantly increased the open probability of BK_{Ca} channels (Taniguchi and Guggino 1989). Comparable effects were observed in intercalated cells of rat cortical collecting tubule (Pácha et al. 1991), in the human colonic cell line CaCo-2 (Bear 1991), and in vascular smooth muscle cells (Kirber et al. 1992).

These results suggest a direct effect of membrane stretch on the channel itself or on a closely associated component as a mode of channel regulation. Also of note is the distinct sensitivity of BK_{Ca} channels to extracellular ATP as it was observed in Vero cells (Hafting and Sand 2000). Thus, ATP release may well function as an autocrine (or paracrine) mech-

anism of hypotonic channel activation. In addition, a variety of protein kinases and phosphatases appear to have an impact on BK_{Ca} channel activity (Levitan 1994; Vergara et al. 1998; Hafting and Sand 2000) but their precise role in channel regulation remains to be elucidated.

 IK_{Ca} channels are activated by cytosolic Ca^{2+} activities in the upper nanomolar range and exhibit an *intermediate conductance* that (at 0 mV) equals 20 pS–80 pS. They are voltage-independent but weakly inwardly rectifying (under symmetrical high K^+ conditions), (see Vergara et al. 1998; Jensen et al. 2001 for review). The apparently large range of unitary conductances reported for these channels is in part due to differences in experimental design and the very pronounced dependence of these channels on the actual (extracellular) K^+ activity. IK_{Ca} channels are weakly inhibited by quinine but efficiently blocked by charybdotoxin and (more selectively) by clotrimazole. The latter compound was found to have a high affinity to the "Gardós channel" (this channel is the IK_{Ca} of erythrocytes and actually the first Ca^{2+} -dependent K^+ transport described; Gardos 2002) and was therefore considered to be of potential use for the treatment of sickle cell anemia (Jensen et al. 2001). IK_{Ca} channels are virtually insensitive to the bee venom toxin apamin, which discriminates them from most isoforms of SK_{Ca} channels (see below; Jensen et al. 2001). Interestingly, IK_{Ca} channels are selectively activated by the channel modulator 1-EBIO (1-ethyl-benzimidazolinone; Jensen et al. 2001).

The first IK_{Ca} channel was cloned from human tissues (*KCNN4*; Ishii et al. 1997; Joiner et al. 1997; Logsdon et al. 1997; Jensen et al. 1998) and shortly thereafter the orthologues of mouse and rat followed (Warth et al. 1999; Vandorpe et al. 2002). The human IK_{Ca} channel is 88% and 90% identical to the mouse and rat channel, respectively, and 40% to 42% identical to SK_{Ca} channels. The short cytosolic N-terminus is followed by six transmembrane segments and a long intracellular C-terminus (Vergara et al. 1998; Jensen et al. 2001). The channel pore is most probably located in a hydrophobic pocket between TM5 and TM6 and the channel is likely to function as a homotetramer. The Ca^{2+} sensitivity of IK_{Ca} channels appears to be mediated by calmodulin binding to a C-terminal domain. Using Northern blot analyses, IK_{Ca} channels were mainly detected in tissues rich in epithelia and endothelia (Jensen et al. 2001). This is of note because of the known high rates of vectorial transport in such systems and the resultant need for an effective osmotic cell homoiostasis.

There are several reports which identify this type of channel as the mediator of conductive hypotonicity-induced K⁺ release. In the human epithelial cell line Intestine 407, hypotonic swelling led to a significant increase of cell Ca²⁺, thereby activating IK_{Ca} channels (Okada et al. 2001). Human T lymphocytes express a Ca²⁺-activated K⁺ conductance that was activated by hypotonic conditions (Khanna et al. 1999). The K⁺ conductance was slightly inwardly rectifying and blocked by charybdotoxin as well as clotrimazole. In transformed Madin-Darby canine kidney (MDCK-F) cells, hypotonic stress increased cell Ca²⁺ and activated a 53 pS K⁺ channel that was inwardly rectifying (Schwab et al. 1993) and blocked by charybdotoxin, Ba²⁺, and TEA (Schwab and Oberleithner 1996; Schwab et al. 1999; Schneider et al. 2000). Because MDCK-F cells employ cell volume regulatory mechanisms for locomotion, cell migration could be shown to be inhibited with an identical pharmacological profile (Schwab and Oberleithner 1996). Moreover, the channel appeared to be activated by 1-EBIO (Schwab et al. 1999). In the distal nephron cell line A6 derived from *Xenopus* kidney, cell swelling activated a K⁺ channel that (in symmetrical 130 mMol/l K⁺) was slightly inwardly rectifying with a unitary conductance of 29 pS (at

0 mV) and inhibited by quinine (Nilius et al. 1995). It is tempting to speculate that the channel may be Ca²⁺-sensitive because, in A6 cells, hypotonic stress was shown to elicit marked increases of cell Ca²⁺ that paralleled RVD and the activation of a TEA-sensitive K⁺ conductance (Yu and Sokabe 1997; Urbach et al. 1999).

In some systems, the putative role of IK_{Ca} channels in the RVD process is not yet conclusively defined. In MDCK cells for instance, hypotonic stress was reported to transiently activate K⁺ channels that were Ca²⁺-sensitive and in cell-attached patches (with 150 mMol/l K⁺ in the pipette); these channels were inwardly rectifying with a unitary conductance of 29 pS at 0 mV (Weiss and Lang 1992). From the same system, Banderali and Roy (Banderali and Roy 1992a) reported a Ca²⁺-sensitive channel that showed a significant increase of open probability upon hypotonic cell swelling as well and that was blocked by quinine (Roy and Banderali 1994). In symmetrical 145 mMol/l K⁺, however, the channel clearly exhibited a linear current-to-voltage relation with a unitary conductance of 24 pS (Banderali and Roy 1992a; Roy and Banderali 1994). In an early report on Ehrlich ascites tumor cells, RVD was found to involve the activation of an inwardly rectifying K+ channel that (under symmetrical high K⁺ conditions) exhibited unitary conductances of 40 pS and 15 pS in the negative and positive voltage range, respectively. The channel appeared to be Ca²⁺-activated and its open probability did not depend on membrane voltage (Christensen and Hoffmann 1992). In a more recent study on the same preparation, however, Ca²⁺ signaling did not seem to play any significant role in RVD at all, and hypotonicity-induced K⁺ release was found to be inhibited by Ba²⁺ but clearly not by charybdotoxin or clotrimazole (Jorgensen et al. 1997). In some other systems, cell swelling activates K⁺ channels of intermediate conductance that apparently are not related to IK_{Ca}. In Necturus proximal tubule for instance, cell-attached patches on the basolateral membrane revealed the presence of a K+ channel that could be activated by negative pressure as well as by a reduction in bath osmolarity (Sackin 1989). There was no detectable voltage dependence of channel gating and the channel did not appear to be Ca²⁺-sensitive. Under symmetrical high K⁺ conditions (70 mMol/l), the channel exhibited a linear current-to-voltage relation and a conductance of 36 pS (Filipovic and Sackin 1992). In cell-attached patches on Xenopus proximal tubule, a basolateral K+ channel was observed that was activated by negative pressure and blocked by Ba²⁺ and was supposed to be involved in RVD (Kawahara 1990). With 100 mMol/l K⁺ in the pipette, the channel showed little if any inward rectification with a unitary conductance of 28 pS and it did not appear to be Ca²⁺-sensitive.

SK_{Ca} channels under symmetrical high K⁺ conditions exhibit a *small conductance* of some 4 pS–18 pS (at 0 mV) and are activated by nanomolar concentrations of Ca²⁺. They are inwardly rectifying and voltage-independent (see Vergara et al. 1998 for review). In electrically excitable cells, SK_{Ca} channels mediate the slow after-hyperpolarization following action potentials. SK_{Ca} channels have been recently cloned from—or identified in—rat brain and colon (rSK1–3 and rSK4, respectively), as well as human brain (hSK1) and placenta (hSK4; Köhler et al. 1996; Warth et al. 1999; Okada 1997; Joiner et al. 1997). The topology of SK_{Ca} channels is similar to that of voltage-gated K⁺ channels with six transmembrane helices, an intracellular C- and N-terminus, positively charged residues in the TM4 segment, and the pore-forming region between TM5 and TM6. Primary sequences, however, are significantly different. The channels are likely to function as homotetramers, and gating of SK_{Ca} channels involves an interaction of calmodulin with C-terminal domains. SK_{Ca} isoforms differ markedly with respect to their sensitivity to apamin: SK2 is highly sensitive to the bee venom toxin (with a K_d of 60 pMol/l for rSK2) whereas SK1

and SK4 are not significantly inhibited in concentrations up to 100 nMol/l; SK3 exhibits an intermediate sensitivity (Vergara et al. 1998). The sensitivity to apamin is defined by a distinct group of amino acids in the deep pore of the channels, which also determine channel sensitivity to D-tubocurarine, an additional, quite selective blocker. In a recent study, a full-length 2.1 k-bp cDNA (hSK2) highly homologous to rat brain rSK2 was isolated from a human liver cDNA library and identical cDNAs were obtained from human primary hepatocytes, human HuH-7 hepatoma cells, and human Mz-ChA-1 cholangiocarcinoma cells (Roman et al. 2002). Stable transfection of CHO cells with hSK2 resulted in the expression of an apamin-sensitive K⁺ conductance as revealed by whole-cell patch-clamp recordings. Thus far, there is only limited evidence for the involvement of SK_{Ca} in volume regulation. Although in Mz-ChA-1 cells hypotonic stress led to a prominent increase of K⁺ conductance, and this effect as well as RVD was partially inhibited by apamin (at 50 nMol/l and 100 nMol/l, respectively), Ba²⁺ (at 5 mMol/l) exhibited a complete block of RVD, suggesting that additional K⁺ channels contribute to this process (Roman et al. 2002).

Two members of voltage-gated K⁺ channels could be defined as mediators of RVD, namely Kv1.3 and Kv1.5. They belong to the group of delayed rectifier channels that were first cloned from (electrically) excitable tissues (Stühmer et al. 1989; Tamkun et al. 1991) where they stabilize resting potential and mediate the rapid repolarization of action potentials. K⁺ channels of the Kv group typically exhibit six transmembrane helices, an intracellular C- and N-terminus, and a positively charged TM4 segment that most likely is part of the gating machinery. Kv channels were found to be also expressed in electrically nonexcitable cells such as T-lymphocytes (Lewis and Cahalan 1995). There is increasing evidence suggesting that they play a role in cell volume regulation (Felipe et al. 1993; Lewis and Cahalan 1995). It was found, for instance, that a mouse T-lymphocyte cell-line (CTLL-2) that does not express voltage-dependent K+ channels did not exhibit any significant RVD under hypotonic conditions. Transient transfection of these cells with Kv1.3, however, reconstituted their capability of an almost complete volume regulatory response and led to the generation of significant voltage-activated K+ currents (Deutsch and Chen 1993). Moreover, the RVD was blunted in the presence of 50 nMol/l charybdotoxin. In a study on human T-lymphocytes both hSK2 (see above) and Kv1.3 were found to contribute to K⁺ conductance and RVD (Khanna et al. 1999). In this investigation, the contribution of Kv1.3 was quantified in a differential approach by use of the specific blocker (scorpion toxin) margatoxin (applied at 5 nMol/l). Stable transfection of a mouse fibroblast cell line (Ltk⁻) with Kv1.5 prevented isotonic cell swelling (that was elicited by the application of dexamethasone) and elicited sizeable voltage-activated K⁺ currents (Felipe et al. 1993). Both effects of channel expression were blocked by 60 μ Mol/l quinine.

The I_{sK} protein was first cloned from rat kidney by use of the *Xenopus* oocyte expression system (Takumi et al. 1989). Its most remarkable feature is its *small size* of 126–130 amino acids (depending on the species) with a single transmembrane α -helical domain (Busch and Maylie 1993; Busch and Suessbrich 1997). In various systems, expression of I_{sK} elicited a K^+ conductance upon membrane depolarization that was very slowly activating with time constants of 10 s and more (Tai et al. 1997). I_{sK} functions as a regulatory subunit of K_vLQT1 (which exhibits the typical topology of voltage-dependent K^+ channels and a tissue distribution that is very similar to that of I_{sK}) thus forming the functional "minK" channels (Barhanin et al. 1996; Tai et al. 1997). Alternative partners for both K_vLQT1 , as well as I_{sK} , however, may also exist.

In *Xenopus* oocytes that were injected with I_{sK} cRNA (and that were later found to endogenously express a K_vLQT1 analog; Sanguinetti et al. 1996), slowly developing voltage-activated K⁺ currents were detected (Busch et al. 1992). Hypotonic stress led to a pronounced increase of these currents, to an accelerated activation, and to a shift in voltage dependence to more negative membrane voltages. The above effects were completely abolished in Ca^{2+} -free solutions (Busch et al. 1992). A significant contribution of minK channels to the RVD process was also reported from vestibular dark cells of gerbil inner ear (Shiga and Wangemann 1995; Wangemann et al. 1995) and from mouse tracheal epithelium (Lock and Valverde 2000). In the latter system, RVD was insensitive to Ba^{2+} and apamin and only weakly inhibited by TEA, which is to be expected for I_{sK} . In contrast, clofilium (a quaternary ammonium blocker, used at $100~\mu\text{Mol/l}$), which is a rather selective blocker of the K_vLQT/I_{sK} complex, potently inhibited RVD. Moreover, in tracheal epithelial cells, the I_{sK} (–/–) knockout mouse, RVD was no more detectable (Lock and Valverde 2000).

In a recent study, TASK-2 (KCNK5) was identified as a mediator of hypotonicity-induced conductive K^+ release in Ehrlich ascites tumor cells (Niemeyer et al. 2001). TASK, for TWIK (*T*andem of P domains in *Weak Inward* rectifier K^+ channels)-related acid-sensitive K^+ channels, belong to the group of two pore-domain K^+ channels (K_{2P} ; Lesage and Lazdunski 2000). These channels have four transmembrane segments and they operate as dimers. There is a widespread distribution of K_{2P} channels in both (electrically) excitable and nonexcitable tissues. K_{2P} channels appear to define the background K^+ conductive properties of many cell membranes. They are insensitive to most classical K^+ channel blockers (Lesage and Lazdunski 2000). When expressed in HEK293 cells, TASK-2 elicited hypotonicity-induced currents that are very similar to those in Ehrlich cells, e.g., with respect to ion sensitivity ($K^+ > Rb^+ >> Cs^+ > NH_4^+ > Na^+ \cong Li^+$) and inhibition by clofilium, with an IC_{50} of 25 μ Mol/l. In addition, mTASK-2 (but not mTASK-1 and mTASK-3) could actually be identified in Ehrlich cells (Niemeyer et al. 2001).

In conclusion, hypotonicity-induced K^+ channels represent an effective mechanism for the selective release of K^+ during RVD. They differ considerably with respect to unitary conductance, gating mechanism, and molecular organization. On the other hand, the high degree of variability among the numerous proteins actually recruited for conductive K^+ release underlines the general importance of this mechanism for RVD (Wehner 1998).

Cl⁻ channels

The apparently most abundant form of hypotonicity-induced Cl⁻ channels belong to a group that has been named VSOR (volume-sensitive outwardly-rectifying) Cl⁻ channels (Okada 1997), VRAC (for "volume-regulated anion channels" (Nilius and Droogmans 2001), or VSOAC (volume-sensitive organic osmolyte-anion channels; Jackson et al. 1994). These channels are referred to as mediators of I_{Cl,swell} (Nilius et al. 1999) or I_{Cl,vol} (Hoffmann and Mills 1999). They appear to be ubiquitously expressed (see Strange et al. 1996; Okada 1997 for comprehensive reviews; see also Nilius and Droogmans 2001). An example for such a channel is given in Fig. 4.

Their significant outward rectification persists under symmetrical Cl⁻ conditions. An interesting feature of VSOR Cl⁻ channels is their pronounced inactivation when membrane voltage is stepped from very negative values to above +50 mV. The channels exhibit intermediate conductances in the range of 10 pS to 40 pS at negative and 40 pS to 80 pS at

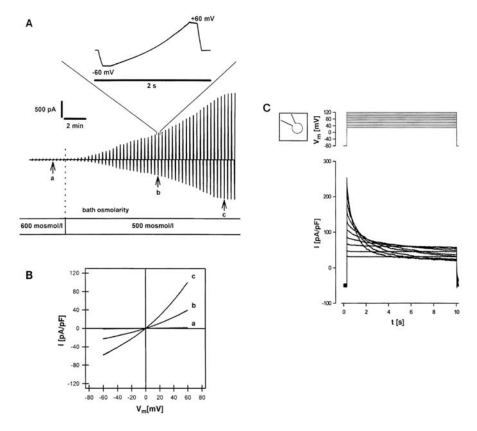


Fig. 4 A Activation of anion conductance in rat inner medullary collecting duct (IMCD) cells under hypotonic conditions. Whole-cell patch-clamp recording with symmetrical CsCl solutions; the osmolarity of the pipette solution was 600 mOsm/l. At the time indicated, osmolarity of the superfusate was reduced from 600 mOsm/l to 500 mOsm/l. Voltage ramps from -60 to +60 mV and 2 s duration were applied every 15 s. **B** Current-voltage relationship before (*line a*), at 6.5 min (*line b*), and at 11 min (*line c*) of hypotonicity; currents were referred to membrane capacitance. **C** Time-dependent inactivation of hypotonicity-induced Cl⁻ conductance at depolarizing voltages. Membrane voltage was stepped from -80 mV to positive values in the range of +30 to +120 mV with 10 mV increment after each pulse. Pulse duration and interval between pulses were 10 s and 15 s, respectively (**A** and **B** from Boese et al. 1996b; **C** from Boese et al. 1996a)

positive membrane voltages. The activation of VSOR Cl⁻ channels strictly depends on the nonhydrolytic binding of ATP at physiological intracellular concentrations (Jackson et al. 1994; Jackson et al. 1996; Meyer and Korbmacher 1996). Extracellular ATP in contrast, induces an open-channel block and it does so in its Mg^{2+} -free form (Nilius et al. 1994a; Jackson and Strange 1995a; Jackson et al. 1996; Tsumura et al. 1996). VSOR Cl⁻ channels exhibit the permeability sequence $SCN^- > I^- > NO_3^- > Br^- > Cl^- > HCO_3^- > F^- > isethionate > gluconate > glycine > taurine > aspartate, glutamate (Boese et al. 1996a; Strange et al. 1996; Okada 1997; Nilius and Droogmans 2001). The halide selectivity corresponds to Eisenman's sequence I, suggesting a relatively weak interaction of permeant ions with cationic sites inside the channel when compared to their dehydration energy (Wright and Diamond 1977). The physiological significance of the permeability of the channels to cer-$

tain amino acids that function as organic osmolytes will be discussed below (see the section entitled "Swelling activated release").

VSOR Cl⁻ channels are inhibited by submillimolar concentrations of SITS and DIDS. This inhibition is reversible and it is voltage-dependent so that outward currents are more sensitive than inward currents (which is supposed to reflect the preferred entrance of these compounds into the channel mouth at positive membrane voltages; Boese et al. 1996b; Okada 1997). In a voltage-independent fashion, the channel is also efficiently inhibited by NPPB and, in some preparations, by DPC and its derivatives flufenamate and niflumate. Interestingly in some systems, VSOR Cl⁻ channels were found to be blocked by inhibitors of p-glycoprotein (the product of the MDR1 gene), like tamoxifen, 1,9-dideoxy-forskolin, verapamil, and quinidine, which led to the hypothesis of a close molecular correlation between these membrane proteins. This notion appeared initially to be strongly supported by experiments in which p-glycoprotein was heterologously expressed in various cell systems (Gill et al. 1992; Valverde et al. 1992). Currently even a possible role of p-glycoprotein as a regulator of VSOR Cl⁻ channels is questioned (Okada 1997; Nilius and Droogmans 2001).

Despite the rather ubiquitous presence of VSOR Cl⁻ channels, their molecular entity still remains a mystery. One of the reasons for this is the absence of any specific inhibitor of VSOR Cl⁻ channels. A possible molecular candidate is I_{Cln}, a protein that was originally cloned from a Madin Darby canine kidney (MDCK) cell library and that elicited a nucleotide-sensitive ("n"), outwardly rectifying chloride current (with rapid inactivation at positive voltages) when expressed in Xenopus oocytes (Paulmichl et al. 1992). Because of the presence of VSOR Cl⁻ channels (or functionally related channels) in almost all cells, however, it was always difficult to decide whether the I_{Cln} protein reflects a channel regulator or the channel itself (Strange 1998; Fürst et al. 2000b). A possible role of I_{Cln} as a regulator has been inferred from the observation that, in most instances, it appears to be localized in the cytoplasm rather than in the plasma membrane (Krapivinsky et al. 1994; Buyse et al. 1997). On the other hand, a hypertonicity-induced translocation of the I_{Cln} protein from the cytosol to the plasma membrane has also been reported (Laich et al. 1997; Musch et al. 1997; Musch et al. 1998). This observation per se could, of course, equally well reflect the insertion of a channel or the movement of a regulatory factor. Hydrophobicity analysis indicated that I_{Cln} lacks transmembrane helices that are believed to form the pore of most vertebrate ion channels (Paulmichl et al. 1992). Accordingly, a putative model of an I_{Cln} channel was proposed consisting of a homodimer with each monomer containing four β strands so that an eight-stranded, antiparallel β -barrel transmembrane pore is formed (Paulmichl et al. 1992; Fürst et al. 2000b). Interestingly, when purified I_{Cln} was reconstituted in lipid bilayers, single channel events could be detected that exhibited a significant rectification; ion substitutions, however, revealed that the channels observed were rather cation- than anion-selective (Li et al. 1998; Fürst et al. 2000a). In heart lipid bilayers, however, chloride selectivity was found (Garavaglia et al. 2002).

CIC-2 and CIC-3 were also considered as molecular candidates for VSOR Cl⁻ channels. The first member of this group of voltage-gated channels (CIC-0) was cloned from the electric ray *Torpedo* (Jentsch et al. 1990). Typically, ClCs exhibit ten to twelve transmembrane helices and the N- and C-termini are located in the cytosol. Most ClCs (except some prokaryotic isoforms) contain two hydrophobic domains close to the C-terminus that may be involved in their cellular sorting (Schmidt-Rose and Jentsch 1997; Jentsch et al. 1999). ClCs were supposed to function as dimers, with each monomer showing a separate pore.

This structure could recently be proven by X-ray analysis on two bacterial ClC isoforms with 3 Å resolution (Dutzler et al. 2002). CIC-2 is a broadly expressed Cl⁻ channel (Jentsch et al. 1999) that is activated by strong (nonphysiological) hyperpolarizations, acidic extracellular pH, and cell swelling (Gründer et al. 1992). However, it clearly differs from VSOR Cl⁻ channels with respect to its voltage dependence (it is *inwardly* rectifying) and its anion selectivity (P_{Cl} > P_I). An actual role of ClC-2 in cell volume regulation was a matter of debate (Jentsch et al. 1999), but recent evidence supports this notion. In Xenopus oocytes heterologously expressing CIC-2, for instance, hypotonic stress elicited a significant increase of cell membrane Cl⁻ conductance (Stegen et al. 2000). Likewise in Sf9 cells, expression of ClC-2 led to swelling-activated chloride currents that could be specifically inhibited by anti-ClC-2 antibodies (Xiong et al. 1999). In mouse atrial and ventricular myocytes, a Cl⁻ current was observed that was activated by hyperpolarization of membrane voltage and current activation was increased by cell swelling (and inhibited by cell shrinkage). The selectivity sequence of currents was Cl⁻ > I⁻ >> aspartate, and RT-PCR and Northern blot analysis confirmed transcriptional expression of ClC-2 in both tissues (Duan et al. 2000). In rat HTC hepatoma cells, hypotonic stress elicited inwardly rectifying Cl⁻ currents that could be inhibited by anti-ClC-2 antibodies. Moreover, a cDNA closely homologous to rat brain ClC-2 was isolated and ClC-2 mRNA and membrane protein expression was demonstrated by in situ hybridization, immuno-cytochemistry, and Western blot analysis (Roman et al. 2001).

CIC-3 exhibits many similarities to VSOR Cl⁻ channels including outward rectification, unitary conductance, anion selectivity, voltage-dependent inactivation, and inhibition by extracellular nucleotides, stilbene derivatives, and tamoxifen (Strange 1998; Jentsch et al. 1999; Hume et al. 2000). There are some significant differences remaining, however. Among these is the enormous basal activity of ClC-3 when compared to VSOR Cl⁻ channels, the latter being undetectable under isotonic conditions. Moreover, ClC-3 appears to be inhibited by PKC, whereas PKC stimulates VSOR Cl⁻ channels in many preparations (Okada et al. 1998; Okada 1998; Strange 1998).

In most systems, hypotonic stress elicits a significant increase in cell Ca²⁺. This in turn may lead to the activation of Ca²⁺-dependent Cl⁻ channels (CaCCs) that may contribute significantly to the overall volume response (Jentsch 1996; Hoffmann and Mills 1999; Nilius and Droogmans 2001). In Ehrlich ascites tumor cells and in the mouse inner medullary collecting duct cell line mIMCD-K2, for instance, the contribution of both VSOR Clchannels and CaCCs to the hypotonicity-induced Cl⁻ release could be determined in a quantitative fashion (Pedersen et al. 1998; Boese et al. 2000). CaCCs activate slowly at positive and inactivate rapidly at negative voltages. They exhibit strong outward rectification with unitary conductances in the range of 3 pS to 10 pS and the ion permeability sequence is I⁻ > Cl⁻ > gluconate (see Nilius and Droogmans 2001 for review). Channel activity is high at positive and low at negative voltages. CaCC currents are blocked by DIDS, NPPB, niflumate, and tamoxifen. The molecular correlate of CaCCs have not yet been unequivocally identified. The first member of a gene family of Ca²⁺ dependent Cl⁻ channels was cloned from bovine trachea (Cunningham et al. 1995) and since then a number of isoforms followed (Fuller and Benos 2000; Fuller et al. 2001). These "CLCA" are heterodimers of some 90 kDa and 35 kDa that are cleaved from a common precursor (at the inner loop between TM3 and TM4) and the most likely topology are five transmembrane regions with an extracellular N- and an intracellular C-terminus. There are a number of similarities between CLCAs and CaCCs but also significant differences, including unitary conductances which appear to be higher in CLCAs. An actual role of CLCAs in cell volume regulation remains to be demonstrated (Nilius and Droogmans 2001; Papassotiriou et al. 2001).

BCl (or maxi Cl⁻) channels that are activated under hypotonic conditions have been reported from cardiac myocytes of new-born rats (Coulombe and Coraboeuf 1992), mouse N1E115 neuroblastoma cells (Falke and Misler 1989), rat cortical astrocytes in primary culture (Jalonen 1993), the rabbit renal cortical collecting duct cell line RCCT-28A (Schwiebert et al. 1994), and primary cultures of bovine pigmented ciliary epithelium (Zhang and Jacob 1997). Typically, BCl channels exhibit unitary conductances in the range of 100 pS to 400 pS and a linear current-to-voltage relationship. In most instances, up to five identical subconductance levels were detected. BCl channels have a bell-shaped voltage dependence with very low activity outside the range of -40 mV to +40 mV. The anion-to-cation selectivity is higher than ten and the channels are inhibited by DIDS, NPPB, and DPC, but apparently not by tamoxifen. BCl channels may be activated by membrane stretch or by patch excision from nonswollen cells. Based on the effects of cytochalasins, a direct regulatory interaction with the actin cytoskeleton is assumed (Strange et al. 1996; Nilius et al. 1997). BCl channels may be related to the voltage-dependent anion channel (VDAC), a porin-like channel found in eukaryotic mitochondria or to the brain-derived voltage-dependent anion channel 1 (BR1-VDAC), which is also expressed in lymphocytes, kidney, heart, and skeletal muscle (Dermietzel et al. 1994; Thinnes et al. 2001).

Hypotonicity-induced SCl (or mini Cl⁻) channels with unitary conductances in the range of 2 pS to 10 pS have been reported from single-channel patch-clamp studies on the ventricular membrane of choroid plexus (Christensen et al. 1989), Ehrlich ascites tumor cells (Christensen and Hoffmann 1992), and pigmented and nonpigmented bovine ciliary epithelium (Zhang and Jacob 1997). It is because of these low conductances that the biophysical properties and the actual contribution of SCl channels to macroscopic conductances were sometimes hard to define. In the human colon cell-line HT₂₉, for instance, the Cl⁻ channels activated by cell swelling appeared to be too small to be detected by use of single-channel patch-clamp techniques (Kubitz et al. 1992). Stationary noise-analysis of whole-cell currents was considered a way out and unitary currents in the range of 0.26 pS to 6 pS were determined by this method in mouse and human T and B lymphocytes (Lewis et al. 1993; Schumacher et al. 1995), human neutrophils (Stoddard et al. 1993), bovine chromaffin cells (Doroshenko and Neher 1992), and various endothelial, epithelial, fibroblast, and blood cells (Nilius et al. 1994b).

As was outlined in detail by Strange and coworkers (Jackson and Strange 1995b; Jackson et al. 1996), however, this approach may lead to a marked underestimation of unitary conductances (and to a considerable overestimation of the actual number of channel molecules involved) because some basic assumptions made—the number of channels remains constant and these channels respond to cell swelling with a gradual increase in open probability—may not be valid (see also Nilius et al. 1997). Instead, nonstationary noise-analysis may be more appropriate as could be shown for (at least some) VSOR Cl⁻ channels which respond to hypotonic stress with an abrupt shift of open probability from virtually zero to a value very close to unity so that the number of channels contributing to the overall membrane response is increasing, i.e., it is not constant (Jackson and Strange 1995b; Boese et al. 1996a). Accordingly, some of the above data obtained by noise-analysis may

need reevaluation, and the apparently low unitary conductances determined may actually be considerably higher.

Taken together, despite the enormous efforts undertaken in recent years to define the biophysical and molecular properties of hypotonicity-induced Cl⁻ channels, the overall image is still elusive and, in some instances, rather conflicting results were obtained. One of the most important steps to be achieved will be towards the molecular understanding of VSOR Cl⁻ channels, which appear to be the most commonly used mechanism of conductive Cl⁻ release during RVD.

K^+ - Cl^- symport

Electroneutral KCl release appears to be one of the major features during RVD in many systems (Lang et al. 1998a; Lauf and Adragna 2000; Hoffmann and Mills 1999; Lang et al. 1998b).

K+-Cl- symport was first defined in erythrocytes as Cl- dependent and ouabain-insensitive K⁺ transport and, thereafter, detected in many other cell types (see Lauf and Adragna 2000 for review). Four isoforms of K⁺-Cl⁻ symporters (KCC1–KCC4) have been cloned to date which exhibit significant molecular similarities to the Na⁺-Cl⁻ symporter (NCC) and to the known mediators of Na⁺-K⁺-2Cl⁻ symport (NKCC1 and NKCC2; see the section entitled "Na+-K+-2Cl- symport"). All together, they form the superfamily of cation-Cl⁻ cotransporters (CCCs) that appears to originate from an ancient short transport protein found in cyanobacteria (Gillen et al. 1996). CCCs contain twelve putative transmembrane domains with large N- and C-terminal regions in the cytosol. Typically, however, KCCs have a large extracellular loop between TM5 and TM6 that is variably N-glycosylated between the different isoforms (Gillen et al. 1996; Lauf and Adragna 2000). TM2 probably reflects the binding/transport site for K⁺, whereas TM4 and TM7 are likely to be involved in Cl⁻ transport (Lauf and Adragna 2000). KCC1 appears to be ubiquitously expressed (Haas and Forbush III 2000; Lauf and Adragna 2000), while KCC2 was found exclusively in the brain (Payne 1997; Jarolimek et al. 1999). KCC3 was most prominent in heart and kidney, but also detectable in skeletal muscle, placenta, liver, lung, pancreas, endothelial cells, and in the central nervous system (Hiki et al. 1999; Mount et al. 1999; Pearson et al. 2001). KCC4 exhibited a rather restricted expression pattern with significant amounts of transcript only in muscle, lung, heart, and kidney (Mount et al. 1999). The likely function of KCC2 in the central nervous system is to decrease cell Cl⁻ below its electrochemical equilibrium and thus to facilitate the hyperpolarizing potency of GABA or glycine-gated Cl⁻ channels.

KCC1 (Gillen et al. 1996; Holtzman et al. 1998; Mount et al. 1999; Su et al. 1999; Mercado et al. 2000; Lauf et al. 2001) and KCC3 (Mount et al. 1999; Race et al. 1999), as well as KCC4 (Mercado et al. 2000), are clearly activated under hypotonic conditions (when heterologously expressed in HEK-293 cells or *Xenopus* oocytes). KCC2 was found to be insensitive to cell swelling (Payne 1997). With respect to the regulation of KCCs, *dephosphorylation* by specific phosphatases (e.g., PP1A) appears to be the trigger for stimulation (Lauf and Adragna 2000). This is an intriguing difference to NKCCs which are activated by *phosphorylation* and which are mediators of the opposite cellular volume response, namely RVI. Also certain protein kinases do participate in KCC regulation. In rat vascular smooth muscle cells, it was found that protein kinase G (PKG)-1 posttranscriptionally led to an acute upregulation of KCC3 mRNA (Di Fulvio et al. 2001b). Moreover, the NO/

cGMP signaling pathway participates in the mRNA regulation of KCC1, which is the major mediator of RVD in these cells (Di Fulvio et al. 2001a).

Cation channels

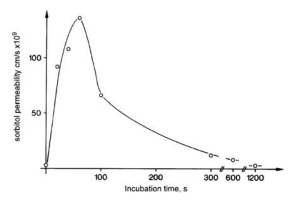
In some systems, hypotonic stress leads to the activation of nonselective cation channels. At first sight this may be surprising because, due to the steeper electrochemical Na^+ than K^+ gradient across the cell membrane, nonselective cation channels are expected to mediate a net uptake rather than a release of osmolytes. It has to be considered, however, that some of these channels exhibit a slightly higher permeability for K^+ when compared to Na^+ . Moreover, certain nonselective cation channels are permeable to divalent cations so that they may play a role in Ca^{2+} signaling. Some nonselective cation channels are sensitive to membrane stretch.

A cation channel with a P_K/P_{Na} of 4.7 was reported from cultured rat mesangial cells (Craelius et al. 1993). In cell-attached patches, this channel could be activated by hypotonic stress and by negative pressure (suction) applied to the patch pipette. The channel exhibited unitary conductances of 76 pS and 40 pS with high K⁺ and high Na⁺ in the pipette, respectively. It was impermeable to Ca^{2+} and Ba^{2+} and its mean open time increased with membrane depolarization. In the rat pheochromocytoma cell-line PC12, cell swelling, as well as negative pressure, activated a cation channel of 46 pS with high K⁺ and 27 pS with high Na⁺ in the (cell-attached) pipette (Cornet et al. 1993). In ventricular myocytes of the rabbit, hypotonicity induced a cation conductance with a P_K/P_{Na} of 5.9 that was blocked by $10 \, \mu \text{Mol/l} \, \text{Gd}^{3+}$ (Clemo and Baumgarten 1997).

A second group of cation channels does not discriminate significantly among small monovalent cations, but these channels exhibit a sizeable permeability to Ca²⁺ (and/or Ba²⁺). Channels of this type were found in Ehrlich ascites tumor cells (Christensen and Hoffmann 1992), rat hepatocytes (Bear 1990), single proximal tubule cells of frog kidney (Robson and Hunter 1994), porcine cerebral capillaries (Popp et al. 1992; Popp et al. 1993), rat atrial cells (Kim and Fu 1993; Kim 1993), guinea-pig gastric smooth muscle cells (Yamamoto and Suzuki 1996), mouse N1E115 neuroblastoma cells (Falke and Misler 1989), as well as in *Rana* and *Xenopus* oocytes (Taglietti and Toselli 1988; Yang and Sachs 1989; Hurst and Hunter 1990; Schütt and Sackin 1997). Unitary conductances were in the range of 15 pS to 40 pS and, in most instances, channels were blocked by micromolar concentrations of Gd³⁺.

Recently, first molecular correlates to hypotonicity-induced, nonselective, and Ca²⁺-permeable cation channels could be identified in human, rat, mouse, and chicken. These are the "osmosensitive, transient receptor potential channel 4" (OTRPC4; Strotmann et al. 2000), TRP 12 (Wissenbach et al. 2000), and the "vanilloid receptor-related, osmotically activated channel" (VR-OAC; Liedtke et al. 2000). They belong to a family of gene products that are structurally related to the light-activated transient receptor potential (Trp) cation channel from *Drosophila melanogaster* (Montell and Rubin 1989). Typically, these osmo-sensitive channels have six putative membrane-spanning domains, a cytosolic N-and C-terminus, and a pore loop between TM5 and TM6; of note, there are three ankyrin-repeat domains close to the N-terminus (Liedtke et al. 2000). The channels appear to be abundantly expressed in kidney but also in lung, liver, spleen, fat tissue, heart, brain, testis, and endothelial cells. When expressed in HEK293 cells, OTRPC4 channels were activated under hypotonic conditions and exhibited outward rectification with unitary conductances

Fig. 5 Time course of activation of sorbitol efflux from IMCD cells during exposure to hypotonic solution. IMCD cells isolated at 600 mOsm/l were exposed at time 0 to a medium of 300 mOsm/l. Membrane permeability was calculated based on the difference between intracellular and extracellular concentration of sorbitol (from Czekay et al. 1994)



of 30 pS for inward and 90 pS for outward currents (Strotmann et al. 2000). P_{Ca}/P_{Na} was 0.8 and P_K : P_{Cs} : P_{Na} : P_{Li} equaled 1.9:1.3:1:0:0.9 (Strotmann et al. 2000; Nilius et al. 2001). The cation channel was insensitive to membrane stretch. It was inhibited by Gd^{3+} (70%) and La^{3+} (30 %) at 100 μ Mol/l each and completely blocked by 10 μ Mol/l of the poly-cation ruthenium red, an effective inhibitor of the vanilloid receptor VR1 and the vanilloid receptor-like protein VRL-1 (Strotmann et al. 2000). Expression of VR-OAC in CHO cells led to hypotonicity-induced whole-cell currents with strong outward rectification (and with dual rectification in the absence of extracellular Ca^{2+} ; Liedtke et al. 2000). In cell-attached measurements (at +80 mV), unitary current events corresponding to a conductance of 310 pS became detectable. VR-OAC channels were blocked by 500 μ Mol/l Gd^{3+} . Transient transfection of CHO and HEK293 cells with TRP 12 led to increased cytosolic Ca^{2+} levels when compared to nontransfected cells and, upon hypotonic stimulation, cell Ca^{2+} further increased (Wissenbach et al. 2000).

Swelling and stretch-activated channels that are nonselective for small monovalent cations and apparently impermeable to Ca²⁺ were found in the basolateral membrane of frog proximal tubule (Hunter 1990), in the mouse kidney cell line mIMCD-3 (Ono et al. 1994), and in human epididymal cells (Chan et al. 1994). Unitary conductances were in the range of 7 pS to 25 pS. In toad urinary bladder and in the A6 cell line derived from *Xenopus* kidney, cell swelling activated nonselective cation channels that were blocked by extracellular Ca²⁺ at millimolar concentrations (van Driessche and Erlij 1994; van Driessche et al. 1994). The actual role of these channels in the RVD process is completely unclear and remains to be elucidated.

Organic osmolytes in RVD

After exposure to hypotonic media, cells begin almost immediately to release organic osmolytes in addition to inorganic osmolytes. This release is not the consequence of a general increase of the membrane permeability to organic solutes, but shows a marked specificity (Grunewald and Kinne 1989; Furlong et al. 1991) and thus involves specific release pathways. These pathways are only briefly activated as exemplified for sorbitol efflux from IMCD cells in Fig. 5 (Kinne 1998).

Some of them have quite low osmotic thresholds. In some cells, organic osmolyte release accounts for about 50% of the total osmolytes released during RVD; thereby, its im-

portance in overall cell volume regulation becomes evident. It also becomes necessary for the cells to coordinate the various channel activities so that they can operate in concert at the proper activity. How this coordination is achieved is one of the enigmas that remain to be solved.

Swelling-activated taurine release

Taurine release induced by cell swelling is a well-studied phenomenon in Ehrlich Ascites tumor cells (Hoffmann et al. 1988), articular chondrocytes (Hall and Bush 2001), HeLa cells (Hall et al. 1996; Stutzin et al. 1999), glial cells (Jackson and Strange 1993; Roy 1995; Roy and Malo 1992), astrocytes (Pasantes-Morales et al. 1994b; Mongin et al. 1999; Olson 1999), liver cells (Warskulat et al. 1997a), endothelial cells (Weik et al. 1998; Nilius and Droogmans 2001), neurons (Pasantes-Morales et al. 1994a), and renal medullary cells (Banderali and Roy 1992b; Ruhfus and Kinne 1996; Sanchez-Olea et al. 1991). During RVD, 30% to 50% of the intracellular taurine leave the cell. Taurine efflux is passive and directed only by the concentration difference (Sanchez et al. 1991; Roy and Malo 1992), lacks saturability and transstimulation. In addition, taurine flux in a variety of cells is inhibited by a range of anion channel blockers (Goldstein and Davis 1994; Boese et al. 1996b; Goldstein et al. 1996; Hall et al. 1996; Kinne et al. 1996; Ruhfus and Kinne 1996; Junankar and Kirk 2000). These properties suggest that taurine efflux is mediated by a transport system which is more similar to a channel than to a transporter (Kirk 1997; Kirk and Strange 1998; Perlman and Goldstein 1999; Shen et al. 2002). The search for such channels has revealed the following candidates thus far (see also Pasantes-Morales et al. 1994b).

VSOR-Cl⁻ channel, VSOAC, or VRAC

As discussed in more detail above (see the section entitled "Cl⁻ channels") these channels are outwardly rectifying (Boese et al. 1996a; Kirk and Strange 1998; Nilius and Droogmans 2001) and have significant permeabilities to anionic amino acids, including those with anionic side chains (glutamate and aspartate), as well as the anionic form of zwitterionic amino acids such as taurine. Attempts to further investigate the substrate specificity with regard to the organic osmolytes by measuring the effect of extracellular amino acids on RVD in glia cells (Pasantes-Morales et al. 1994b) or by electrophysiological studies revealed that the channel has a higher permeability to β -amino acids compared to α -amino acids. The anionic amino acids glutamate and aspartate appear to have a lower permeability than electroneutral amino acids of similar size (Pasantes-Morales et al. 1994b; Kirk 1997).

If compared to the chloride permeability of the channel, P_{taurine}/P_{Cl}— ranges from 0.15 to 0.25 in rat and human glial cells (Jackson and Strange 1993) and rat inner medullary collecting duct cells (Boese et al. 1996a), and to 0.49 in MDCK cells (Banderali and Roy 1992b). The size of the pore has been estimated to be between 5.6 Å and 11 Å (Jackson and Strange 1993; Nilius and Droogmans 2001). The channel therefore, seems to select the substrates both according to size and charge of the molecule.

Some electrophysiological studies also suggest that myo-inositol and sorbitol interact with the channel—since they are electroneutral molecules, no actual proof of translocation across the channel could be provided (Jackson and Strange 1993).

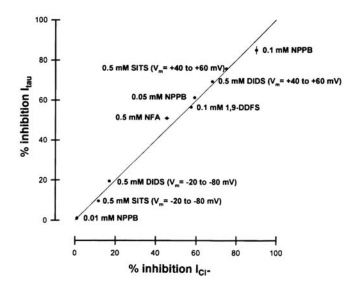


Fig. 6 Effects of various anion channel blockers on swelling-activated anion conductance in rat IMCD cells. Inhibition of volume-activated anion conductance under symmetrical taurine conditions (300 nM; pH 7.8; 31 nM charged; %inhibition I_{Tau}) plotted against inhibition of anion currents in symmetrical 140 mM CsCl conditions (%inhibition I_{Cl}) by various anion channel blockers (unpaired experiments); n=5 for all experimental conditions. For most data points, SE values are smaller than the symbols used. NFA niflumic acid, DDFS 1,9-dideoxyforskolin, NPPB 5-nitro-2-(3-phenylpropylamino)benzoate, DIDS 4,4'-diisothiocyanostilbene-2,2disulfonic acid, SITS 4-acetamino-4'-isothiocyanostilbene-2,2disulfonic acid (from Boese et al. 1996b)

In a variety of cells, VSOR-Cl⁻ channel and the taurine efflux pathways have an almost identical sensitivity to anion channel blockers. Figure 6 shows as an example the findings in rat IMCD cells. Because the rate of activation is also similar, these data suggest that chloride and taurine share the same release path.

Interestingly, VSOAC activity is regulated by the metabolic status of the cell. When intracellular ATP falls, the relative taurine efflux decreases. This has been observed in rat glial cells (Jackson et al. 1994), skate hepatocytes (Jackson et al. 1996), endothelial cells (Oike et al. 1994), and rat IMCD cells (Ruhfus and Kinne 1996). In patch clamp experiments, it was demonstrated that ATP or nonhydrolysable ATP derivatives have to be present at the cytoplasmic side of the membrane for activation of the channel by cell swelling (Jackson et al. 1994; Oiki et al. 1994; Oike et al. 1994; Jackson et al. 1996). These results point against a direct role of phosphorylation in the activation process. However, both phorbol esters and cAMP potentiate swelling induced organic osmolyte flux in C6 glioma cells, involving phosphorylation of VSOAC as a modifier of channel properties (Jackson and Strange 1993).

In agreement with findings of Motais (Motais et al. 1991) in trout red blood cells and in skate erythrocytes (Wittels et al. 2000) on swelling-activated taurine efflux, the activation of VSOAC is suppressed by high intracellular chloride concentrations (Jackson et al. 1996). The regulation by ATP and Cl⁻ has been interpreted to indicate that the cell very sparingly uses the metabolically expensive organic osmolytes only in situations where sufficient intracellular energy remains for a rapid recovery (Jackson et al. 1994; Jackson et al. 1996).

ICl_n protein

The ICl_n protein was cloned first from MDCK cells and exhibits a nucleotide-sensitive outwardly rectifying chloride current when expressed in oocytes (Krapivinsky et al. 1994; Fürst et al. 2000a). These properties are similar to those of swelling-activated chloride channels (see the section entitled "Cl- channels"). When reconstituted in lipid bilayers (Garavaglia et al. 2002) and in oocytes (Stegen et al. 2000), ICl_n shows a significant permeability for taurine. Its proposed molecular structure composed of \(\beta \)-barrels also bears similarities to porins invoked in organic solute transport in bacteria (Schirmer et al. 1995; Saier 2000). Furthermore, in neonatal myocytes, taurine efflux accompanies ICl_n transposition from the cytoplasm to the plasma membrane with a similar time course, suggesting, but not proving, that ICl_n is a regulator of the taurine channel or the channel itself. Knock down studies using antibodies or antisense oligonucleotides showed a significant reduction of RVD-related chloride currents (Gschwentner et al. 1995; Chen et al. 1999a); studies involving overexpression of ICl_n showed the reverse (Hubert et al. 2000). Unfortunately, no data on organic osmolyte fluxes are available from these experiments. Thus, the role of ICln as one of the swelling-activated taurine channels remains to be determined. The potential role of the Cl⁻/HCO₃⁻ antiporter and of phospholemman as taurine channel is discussed in the section entitled "Cl-/HCO₃- antiport" and the section entitled "Phospholemman," respectively.

Finally, it should be noted that in several cells separate pathways for chloride and taurine have been described; thus, additional transporters seem to exist (Shennan et al. 1994; Lambert and Hoffmann 1994; Stutzin et al. 1999).

Myo-inositol release pathways

Flux studies in rat C6 glioma cells (Jackson and Strange 1993) and rat inner medullary collecting duct cells (Ruhfus and Kinne 1996; Ruhfus et al. 1998) revealed striking similarities between swelling-activated myo-inositol and taurine fluxes. In glioma cells and IMCD cells, the rate of activation, the extent of release, and the sensitivity to anion channel inhibitors were virtually identical (Jackson and Strange 1993; Ruhfus and Kinne 1996). In addition, dependence on intracellular ATP levels could be demonstrated in the two cell types for both fluxes. In IMCD cells, the majority of the taurine and myo-inositol occurs across the same membrane at the basal-lateral cell side (Kinne et al. 1996; Ruhfus et al. 1998). Furthermore, the activation pathways (Ca⁺⁺-independent and G-protein independent) are identical (Ruhfus and Kinne 1996). Electrophysiological investigations on the VSOAC in glia cells also showed that myo-inositol probably interacts with the channel (Jackson and Strange 1993).

Thus, it is very likely that myo-inositol efflux occurs via the same channel as taurine. Also, studies in skate erythrocyte suggest that taurine and myoinositol share a swelling-activated transport pathway (Goldstein and Davis 1994; Goldstein et al. 1996; Perlman and Goldstein 1999).

Sorbitol release pathways

Swelling-activated efflux of sorbitol has been observed in C6 glioma cells (Jackson and Strange 1993), a rabbit renal papillary cell line (Siebens and Spring 1989; Garty et al.

1991), HeLa Cells (Hall et al. 1996), rat IMCD cells (Grunewald and Kinne 1989; Ruhfus et al. 1998), rabbit kidney TALH cells (Kinne et al. 2000), and porcine urinary bladder cells (Mahler et al. 1998). In all instances tested, the flux, either measured as unidirectional sorbitol uptake (Jackson and Strange 1993; Hall et al. 1996; Kinne et al. 2000) or as solute efflux (Garty et al. 1991), lacked saturation kinetics. Furthermore, anion channel blockers such as NPPB, DIDS, 1,9-dideoxyforskolin, and cation channel blockers such as quinidine, inhibited sorbitol efflux in most systems, suggesting the involvement of a channel-like protein. With regard to the specificity towards the polyol, the length of the polyol chain and the orientation of the hydroxyl group at C6 appear to affect the permeability, although to a relatively small extent (Napathorn and Spring 1994).

In most cells, except IMCD cells, anions (chloride, iodide), potassium, choline, thymine, and taurine seem to be translocated via the same swelling-activated channel as sorbitol. This conclusion is based on the very similar inhibitory profile of various inhibitors on the fluxes (Hall et al. 1996), a similar lag phase before opening, and a similar dependence on the intracellular ATP level. In addition, in electrophysiological experiments in C6 cells sorbitol affected the volume-activated anion channel in such a way as to suggest an interaction with the anion binding site (Jackson and Strange 1993).

In rat IMCD cells, however, several differences between taurine/myo-inositol efflux and sorbitol efflux have been found. DIDS and SITS stimulate rather than inhibit sorbitol efflux; time of activation and closure of the two efflux pathways differ, as does the osmotic threshold for activation (Kinne 1998). Furthermore, activation of the sorbitol efflux pathway is calcium-dependent, probably involving activation of G-proteins (Ruhfus et al. 1996). Thus, for example, sorbitol efflux is reduced at low extracellular calcium and stimulated by arachidonic acid (Tinel et al. 1997; Kinne et al. 2000) and mastoparan under isotonic conditions (Ruhfus et al. 1996), whereas taurine (and myo-inositol) efflux is not affected by a change in overall calcium or by mastoparan, and inhibited by arachidonic acid (Kinne et al. 1996). However, local changes in the calcium concentration appear to affect swelling-induced Cl⁻ channels (Lemonnier et al. 2002).

It has to be noted, however, that in IMCD cells a considerable (about 50%) fraction of swelling-activated sorbitol efflux is not calcium dependent; thus, the existence of an additional channel—similar to the one proposed for other cells—cannot be excluded.

On the other hand, there is recent evidence for a highly selective sorbitol channel. Studies on malaria-induced pathways in chicken erythrocytes have shown that a solute channel is induced after infection with Plasmodium gallicaneum, which exhibits an about 17-fold higher permeability to sorbitol than to taurine and is relatively insensitive to DIDS and tamoxifen (Staines et al. 2002).

Other systems involved in volume regulation

Cl⁻/HCO₃⁻ antiport

The superfamily of bicarbonate transporters (BTs) consists of two groups, namely, the AE family of anion exchangers (i.e., Cl⁻/HCO₃⁻ antiporters) and the family of HCO₃⁻ transporters that are coupled to Na⁺ (NBCs; see Boron 2001 for review). Whereas no direct contribution of NBCs to cell volume regulation has been reported so far, Cl⁻/HCO₃⁻ antiport is of considerable importance for, both RVI as well as RVD. Four isoforms have been

cloned to date (AE1–AE4) starting with murine AE1 (erythrocyte band 3) as early as 1985 (Kopito and Lodish 1985). The erythrocyte AE1 is a 929 amino acid protein with two functional domains, a hydrophilic one that is related to its role as an anchor for the cytoskeleton, and a hydrophopic one reflecting its function as an anion exchanger with 12 transmembrane regions (Kopito and Lodish 1985). An N-terminal truncated isoform of AE1 is present in the kidney (Brosius III et al. 1989). AE2 is widely expressed in nonerythroid tissues, including the kidney, and AE3 appears to be mainly expressed in electrically excitable tissues. Recently, AE4 was cloned from kidney; it appears to represent the apical CI^-/HCO_3^- antiporter in β -intercalated cells (Tsuganezawa et al. 2001).

One of the prime functions of Cl⁻/HCO₃⁻ antiport in cell volume regulation is the HCO₃⁻-dependent uptake of Cl⁻ that is induced parallel to the hypertonic activation of Na⁺/H⁺ antiport. Both transporters are functionally coupled via cell pH and the net result of their activity is a quasi electroneutral uptake of Na⁺ and Cl⁻. Activation of Cl⁻/HCO₃⁻ antiport parallel to that of Na⁺/H⁺ antiport has been reported from a variety of system, including mammalian hepatocytes (Graf and Häussinger 1996), human retinal pigment epithelial and nonpigmented ciliary epithelial cells (Civan et al. 1994b; Civan et al. 1996), C6 glioma cells (Mountian et al. 1996), flounder red blood cells (Weaver et al. 1999), as well as from mouse medullary thick ascending limb (Hebert and Sun 1988). In addition, a parallel hypertonic stimulation of endogenous NHE1 and of heterologously expressed AE2 was reported from *Xenopus* oocytes (Jiang et al. 1997). A concerted activation of Na⁺/H⁺ antiport and Cl⁻/HCO₃⁻ antiport was also found in renal epithelial (MDCK-F) cells that employ these mechanisms of RVI for the purpose of locomotion (Klein et al. 2000).

Equally well, Cl⁻/HCO₃⁻ antiport may be activated under hypotonic conditions. In skate and trout erythrocytes, and also when heterologously expressed in *Xenopus* oocytes, AE1 appears to mediate the hypotonicity-induced release of Cl⁻ (see the section entitled "Swelling activated taurine release;" Musch et al. 1994; Fievet et al. 1995; Motais et al. 1997; Guizouarn et al. 2001). Of note, it may do so in a channel-like fashion and, at the same time, it also appears to increase cell membrane Na⁺ and K⁺ permeabilities (Guizouam et al. 2001).

AE 1 is one of the most clearly established molecular identities involved in taurine transport from fish red blood cells that are nucleated (Motais et al. 1991; Motais et al. 1997; Guizouarn et al. 2001). In particular, for AE 1 derived from trout, it has been convincingly demonstrated that expression of the protein in oocytes elicits both anion conductance and taurine transport, which show the same inhibitor profile and are clearly distinguishable from the endogenous I_{Cl} swell of the oocytes (Fievet et al. 1995; Fievet et al. 1998; Koomoa et al. 2002).

The fact that in oocytes AE 1—contrary to intact trout red blood cells—cannot be activated by hypotonic stress and is not DIDS-sensitive might reflect different conformations of the protein in the two systems. Different conformations that change transport characteristics and sensitivity to modification have also been reported for ICl_n and phospholemman, and might be a general property of volume-regulated organic osmolyte channels. Another caveat remains: mouse AE 1 does not act as an osmolyte channel. In addition, in eel erythrocytes, significant differences between the AE 1-mediated SO₄ transport and swelling-activated taurine transport were observed (Lewis et al. 1996). Thus, a role of AE 1 in taurine release from mammalian cells remains to be demonstrated.

Phospholemman

Phospholemman (PLM) is a small membrane protein of 15 kDa which was first purified, cloned, and sequenced from dog heart (Palmer et al. 1991). The mature protein consists of 72 amino acids and has one single transmembrane domain. The name PLM denotes its high degree of phosphorylation, which (under most physiological conditions) appears to be mediated by protein kinases A and C (Palmer et al. 1991; Walaas et al. 1994; Mounsey et al. 1999). When expressed in Xenopus oocytes, PLM induces anion currents that are activated by large hyperpolarizations of membrane voltage and most likely it does so by inducing endogenous oocyte channels (Moorman et al. 1992; Mounsey et al. 1999). On the other hand, when reconstituted in lipid bilayers, PLM could be shown to form functional ion channels itself which exhibited rather slow gating kinetics and an enormous unitary conductance that is close to 700 pS (Moorman et al. 1995). It was also found that PLM (again in lipid bilayers) exhibits two different modes of operation, namely a cation- and an anion-selective one, and that the protein appears to spontaneously switch between these modes (Kowdley et al. 1997; Moorman and Jones 1998). PLM also exhibits a remarkably high permeability for taurine with a Ptaurine/PCI as high as 70 (Moorman et al. 1995) and a high selectivity with regard to the NH₂ group at the β -position (Moorman et al. 1995; Kowdley et al. 1997; Moorman and Jones 1998).

Overexpression of PLM in HEK293 cells significantly increased the membrane currents elicited by *hypotonic* stress (Moorman and Jones 1998) as well as the release of Cl⁻ (I⁻) and taurine during RVD (Morales-Mulia et al. 2000); moreover, this osmolyte release could be markedly reduced by use of PLM-specific antisense oligonucleotides (Morán et al. 2001).

In solitary rat hepatocytes in primary culture, which could be shown to exhibit a distinct RVI (Kirschner et al. 1998) and which do express PLM, *hypertonic* conditions activate a channel that very much resembles PLM with respect to its unitary conductance, its gating pattern, and its nonselectivity for Na⁺ over K⁺. Moreover, in *Xenopus* oocytes expressing hepatocyte PLM, hypertonic stress induces a nonselective cation conductance and noise-analysis reveals the activation of a channel with characteristics that appear to be similar to those of PLM (Kirschner et al. 2003).

Taken together these results suggest a role of PLM in both RVD *and* RVI. This could mean that PLM functions as a last line of defense so that a cell under strong anisotonic conditions opens a large osmotic shunt just to survive. As an alternative view, given the regulatory role PLM appears to play in some systems (Moorman et al. 1992; Mounsey et al. 1999) and its high degree of phosphorylation per se (Palmer et al. 1991; Walaas et al. 1994; Mounsey et al. 1999) the protein may well be part of the signaling machinery that actually tunes the ion transporters responsible for cell volume regulation.

Na+, K+-ATPase

Cell volume homeostasis depends on the balance between osmolyte entry into the cell and removal of osmolytes from the cell interior. For cells that possess sodium entry pathways, such as sodium channels, Na⁺/H⁺ antiporter, Na⁺, K⁺, 2Cl⁻ symporter, or other sodium cotransporters, sodium entering the cells is removed by the primary active Na⁺, K⁺-ATPase. The activity of this enzyme, thereby plays an important role in maintaining the cell volume as indicated by the cell swelling observed in isotonic media when ouabain is present

in the incubation medium. This observation reflects the actual stoichiometry of Na⁺, K⁺-ATPase of 3 Na⁺/2 K⁺ which continuously leads to a net loss of cellular inorganic osmolytes. In this context, it is also of note that in some systems an *inhibition* of the enzyme appears to directly contribute to the RVI processes (see below).

Since the first description of the enzymatic activity in 1957 by Skou (Skou 1957), the transport cycle has been described in great detail, the two subunits α and β have been identified, and a cell-specific distribution of their isoforms has been described (for review see Blanco and Mercer 1998).

The α -subunit has a molecular weight of 110 kDa, includes ten predicted transmembrane domains, and is responsible for the catalytic activity of the enzyme. During the pump's catalytic cycle the α -unit is transiently phosphorylated on a residue of the large cytoplasmic loop between transmembrane domains 4 and 5. This protein also contains phosphorylation sites for PKA and PKC. Residues of the fourth transmembrane segments contribute to the cation selectivity of the enzyme (for review see Dunbar and Caplan 2000).

The β -subunit has a molecular mass of approximately 65 kDa and elicits only one transmembrane segment with a cytosolic N-terminus. The extracellular C-terminus is heavily glycosylated (for review, see Dunbar and Caplan 2000). Assembly between the α - and β -subunits is required for the enzyme complex to exit the ER.

In addition, a γ -subunit has been identified; there is still considerable debate as to whether this subunit is required for the enzyme activity and/or whether it represents a regulatory membrane protein, which in some cells is closely associated with the enzyme (for review see Therien and Blostein 2000). The variability of the expression of the different subunits, the composition of which changes the properties of the enzyme considerably, results in a large variation of the response of the enzyme to changes in extracellular osmolality. In the context of this chapter, therefore, only a few examples of typical responses will be discussed.

During RVI following the exposure of cells to shrinkage in hypertonic media, the Na⁺, K⁺-ATPase is particularly important in cells that primarily and predominantly activate regulatory osmolyte transport systems, which enhance the intracellular sodium content, such as the Na⁺/H⁺ antiporter, the Na⁺, K⁺, 2Cl⁻ symporter, or a sodium channel, as discussed above. In a study in which osmolyte and Na⁺ transport balances of rat hepatocytes were quantified as a function of hypertonic stress (see Fig. 7), it could be demonstrated that ouabain-sensitive ⁸⁶Rb uptake, representing the transport activity of the Na⁺, K⁺-ATPase, increased almost fourfold in a saturable fashion with increasing extracellular osmolarity (Wehner and Tinel 2000).

This increase could be completely accounted for by the rise in intracellular sodium concentration and the apparent Michaelis-Menten constant for sodium of the enzyme of 12 mMol/l.

The regulation of transport activity by changes of intracellular sodium represents the most straightforward and well-known response of the Na⁺, K⁺-ATPase to an osmotic challenge (Wojnowski and Oberleithner 1991). There are, however, also instances where either no increase in transport activity or even a decrease during RVI is observed. No change in ouabain-sensitive 86 Rb⁺ uptake was, for example, observed in the human hepatoma cell line Hep G2 (Wehner et al. 2002a). In view of the already relatively high Na⁺, K⁺-ATPase transport activity in these cells under isotonic conditions, the authors argued that the enzyme might normally function at V_{max} level and therefore a further stimulation might not

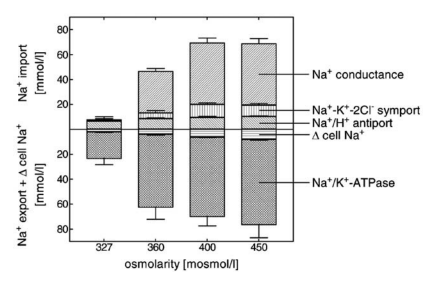


Fig. 7 Balance between Na⁺ import and Na⁺ export by the Na⁺, K⁺-ATPase plus the actual increase of cell Na⁺. Rat hepatocytes in primary culture were exposed to increasing osmolarities. Note the saturability of the Na⁺, K⁺-ATPase pump rate (from Wehner and Tinel 2000)

be possible. In Malpighian tubule, an inhibition of Na⁺, K⁺-ATPase was observed during RVI. Here, the increase in intracellular adenosine, which inhibits the enzyme activity, was supposed to cause this effect (Caruso-Neves and Lopes 2000). In view of the fact that the Na⁺, K⁺-ATPase activity may also be altered by various kinases and phosphatases, as well as eicosanoids, intracellular ATP levels, intracellular pH (Goldstein and Davis 1994; Therien and Blostein 2000), rate of K⁺ recycling across the membrane, and the membrane potential, the response of the enzyme during volume regulation can differ considerably depending on the cellular system being investigated.

It has to be noted that the alterations in Na⁺, K⁺-ATPase transport activity described above included regulation via phosphorylation, changes in transport rate, substrate, or inhibitor concentrations, or modifications in driving forces without changes in mRNA or enzyme content. Changes in the latter two parameters are observed when cells are exposed in vitro to hypertonic media for an extended time period. Thus, in mouse inner medullary collecting duct cells (Ohtaka et al. 1996), MDCK cells (Bowen 1992), human proximal tubule cells (Yordy and Bowen 1993), and vascular smooth muscle cells (Muto et al. 1998), changes in α -mRNA and β -mRNA are observed as early as one h after exposure of cells to a hypertonic medium. A maximum response is found between 8 h and 24 h of anisotonic incubation. The increase in mRNA leads to an augmentation of Na⁺, K⁺-ATPase abundance as demonstrated by a higher enzymatic activity in cell homogenates. An in vivo modulation of Na⁺, K⁺-ATPase expression was also observed in the renal papilla of water-deprived mice versus water-loaded mice, suggesting an osmoregulation of the enzyme in the intact animal (Capasso et al. 2001).

The mechanisms involved in the upregulation are still unclear; changes in the intracellular sodium concentration combined with an increased synthesis of intermediate regulatory proteins have been discussed (Yordy and Bowen 1993; Muto et al. 1998).

The physiological rationale for this long term adaptation is probably the same as for the short-term activation during RVI. In both instances, the cell aims to maintain a suitable sodium gradient across the plasma membrane. This mechanism assures the proper operation of sodium cotransport systems that include sodium osmolyte cotransport systems (see the section entitled "Organic osmolytes in RVI") and also systems regulating important intracellular parameters, such as intracellular pH and intracellular calcium. Thus, the activation of a net osmolyte export system during RVI, a reaction which at first sight appears to be counterproductive, fulfills an important function for general cell activity.

The effect of cell swelling on the Na⁺, K⁺-ATPase is difficult to estimate since a variety of stimulating and inhibiting parameters change simultaneously. Thus, cell swelling reduces intracellular sodium concentration, but activates K⁺ channels (see the section entitled "K⁺ channels") that facilitate K⁺ recycling and thus could increase Na⁺, K⁺-ATPase activity. The increase in membrane potential would tend to decrease the transport rate of the electrogenic pump. The net effect of these—and other—regulatory changes differs from cell to cell. Nevertheless, there is evidence from various tissues that ouabain inhibits RVD, pointing to a significant role of the Na⁺, K⁺-ATPase in removing excess inorganic osmolytes from the cells (Fraser and Swanson 1994; Grunewald et al. 1994; Kanli and Norderhus 1998).

Signal transduction in volume regulation

Sensory systems and set points

Although the search for sensory mechanisms of cell volume changes has been carried out very intensively during recent years, a conclusive model of an osmosensor in mammalian cells has not yet been presented. The main cause for this gap lies in the nature of cellular signalling events. Usually, several signalling mechanisms are activated in parallel and some of them do influence each other mutually. This integrative character of signalling networks makes the assignment of signalling events to distinct pathways difficult. The concept that not one single osmosensor but rather a complex sensory system exists, has to be taken into account. Such a sensory system has to fulfill two major requirements. On the one hand, it has to be very sensitive to perceive even slight alterations of cell volume. On the other hand, the sensor must amplify and transfer the information to trigger signalling cascades leading to the activation of volume regulatory processes.

The mechanisms that have been reported to be implicated in sensing changes of cell hydration can be classified into three major groups, according to their cellular location. They can be (1) associated with the cytoskeleton perceiving changes in the interaction between extracellular matrix and plasma membrane receptors; (2) intracellular solute sensors sensitive to changes in macromolecular crowding and/or ionic strength and (3) membrane-based sensors which record alterations of cell membrane stretch.

Extracellular matrix and cytoskeleton

Cells adhere to the ambient extracellular matrix (ECM) via cellular integrins which are clustered to focal adhesions upon binding of ECM-ligands. Proteins that are associated with these adhesion sites may serve to maintain structural integrity, and equally well, they

may be involved in signal transduction. For instance, the focal adhesion kinase FAK is a tyrosine kinase that is involved in the transduction of integrin-mediated cellular responses (Burridge and Zhong 1997). The activity of both integrins and FAK is altered by focal adhesion clustering.

Focal adhesions and the intracellular actin cytoskeleton effect each other mutually in their structures. Interestingly, evidence has been provided which indicates that integrins could be part of a volume-sensing system. In rat skeletal muscle, for example, the ECM-integrin-mediated cell adhesion and the cytoskeleton are involved in mechanochemical transduction of cell volume changes to chemical signals modulating glutamine transport (Low and Taylor 1998). ECM-integrin binding also transduces changes in cell volume and/or shape into changes in endothelial permeability of cultured bovine aortic endothelia and frog mesenteric capillaries (Kajimura et al. 1997). At the frog neuromuscular junction, integrins contribute to the hypertonic enhancement of transmitter release (Kashani et al. 2001).

The finding that integrins may be involved in sensing cell volume changes is coincidental with the circumstance that in nearly all systems studied so far volume changes cause alterations of the cytoskeletal organization. In the majority of the systems examined, hypotonic cell swelling evokes disorganization of the F-actin cytoskeleton. This has been shown for Ehrlich Ascites tumor cells (Pedersen et al. 2001), rat IMCD cells (Czekay et al. 1994) shark rectal gland and C6 glial cells (reviewed in (Henson 1999; Papakonstanti et al. 2000; Pedersen et al. 2001). In contrast, hypotonic actin polymerization could be detected in rat hepatocytes (Theodoropoulos et al. 1992) and HL-60 cells (Hallows et al. 1991). Increasing extracellular tonicity enhances the relative F-actin content in Ehrlich Ascites tumor cells (Pedersen et al. 2001) and lowers it in HL-60 cells (Hallows et al. 1991). These examples illustrate how diverse the response of the actin cytoskeleton to cell volume changes may be.

The alteration of cell volume evokes changes of cell morphology which are paralleled by the development of mechanical tension. This tension impinges on both the cytoskeleton and the interaction between integrins and their ECM-ligands. In both cases, integrins are altered in their activity by conformational changes. They may be activated and clustered to focal adhesions or inactivated and dispersed from focal adhesions (Burridge and Chrzanowska-Wodnicka 1996; Ingber 1997; Schoenwaelder and Burridge 1999).

The assembly of integrins to focal adhesions can influence the cytoskeleton by two synergistic pathways, both of them involving activation of the small GTPase Rho. The contribution of Rho to the regulation of volume-sensitive transport mechanisms has been reported for several systems, (see Nilius et al. 1999; Okada 1997).

Activation of RhoA by integrin-mediated adhesion (Renshaw et al. 1996; Clark et al. 1998; Ren et al. 1999) and/or by isometric tension of cytoskeletal components (Burridge and Chrzanowska-Wodnicka 1996) induces the formation of stress fibers (Amano et al. 1997). This change is caused by influencing the state of contractile structures or by effecting the conformation of actin-associated proteins (Schoenwaelder and Burridge 1999). Rho activates Rho-kinase, which phosphorylates and thereby inhibits myosin phosphatase. This results in elevated phosphorylation of myosin light chain (MLC; Kimura et al. 1996). The consequence is that myosin filament assembly and actin-activated myosin ATPase activity (Burridge and Chrzanowska-Wodnicka 1996) are enhanced. This leads to the bundling of actin filaments and the assembly of stress fibers. Rho may also phosphorylate MLC directly (Amano et al. 1996).

The alteration of stress fiber assembly by actin-associated proteins occurs via lipid derived messengers. The activation of PI-5 kinase by RhoA elevates the intracellular level of phosphatidylinositol 4,5-bisphosphate (PIP₂; Chong et al. 1994). PIP₂ induces conformational changes of the actin-associated proteins vinculin (Weekes et al. 1996), ezrin-radix-in-moesin (ERM; Niggli et al. 1995), and profilin. Thereby, binding sites for actin and other proteins are exposed. The resulting changes in stress fiber tension are then transduced to the integrins, which thereby are altered in their activity. This bidirectional signalling between the cytoskeleton and integrins allows the creation of a feedback-loop, by which the duration of volume-generated signalling can be determined (Burridge and Chrzanowska-Wodnicka 1996; Schoenwaelder and Burridge 1999).

In this context, another property of the sensor of volume changes, namely, the transduction of the perceived information to effectors, may be realized by three different mechanisms. Firstly, alterations in the cytoskeletal structure may affect the activity of signalling molecules by changing their intracellular localization. Secondly, the cytoskeleton may directly interact with and act upon volume regulatory transporters and, thirdly, vesicular transport, if involved in cell volume regulation, may be influenced, (see for example Czekay et al. 1994).

In many studies cited above, it is not yet clear whether the perturbations of the cytoskeleton observed are the cause or the consequence of volume changes and/or regulatory processes. Furthermore, a direct interaction with the target proteins and the consequences of this interaction have not been recorded in the intact cell. Thus, a higher spatial and temporal resolution of these processes is necessary to be able to conclusively judge upon the role of the cytoskeleton and the ECM in volume sensing.

As was outlined above, the alteration of cell adhesion effects FAK, as well as integrin signalling. For the activation of volume-sensitive Cl⁻ currents, it has been proposed that the stimulation of a tyrosine kinase is one of the early events in osmosensing (Sadoshima and Izumo 1993; Tilly et al. 1993). In most instances, however, tyrosine-phosphorylarions last only for a short period of time, which is mainly due to the presence of tyrosine-specific protein phosphatases. Accordingly, these tyrosine-phosphorylations, in most systems, are translated into longer lasting signalling events, namely, by the activation of serine/threonine phosphorylation cascades, such as the mitogen-activated protein kinase (MAP kinase) pathways. The sequential patterns by which MAP kinases are activated offer the opportunity of an amplification, stabilization, and integration of intracellular signalling events in response to extracellular stimuli at the same time. In fact, MAP kinases could be shown to be involved in the regulation of a variety of fundamental physiological processes such as cytoskeletal organization, nuclear transcription, protein translation, cell growth and differentiation, as well as cell volume homoeostasis.

Activation of ERK1/2 MAP kinases by cell volume changes appears to be a quite common phenomenon. A swelling-induced activation occurs in most mammalian cell types (van Der Wijk et al. 2000). In contrast, activation of ERK1/2 upon hypertonic cell shrinkage was observed, for instance, in MDCK cells (Itoh et al. 1994), H4IIE rat hepatoma cells (Schliess et al. 1996), and in bovine aortic endothelial cells (Duzgun et al. 2000). In these systems, hypertonic activation occurs within 10 min and the maximal induction is reached after 20 min–30 min.

The activity of p38 MAP kinase can be upregulated by hypertonicity as well as by hypotonicity (see Shrode et al. 1998). Hypertonic activation occurs, for example in U 937 and in IMCD-3 cells. In the latter system, the activation of p38 is achieved after approxi-

mately 10 min (Zhang and Cohen 1996; Shrode et al. 1997). Hypotonic activation of p38 has been observed in the perfused rat liver (Vom Dahl et al. 2001), in HTC rat liver hepatoma cells (Feranchak et al. 2001), and in intestine 407 cells (van Der Wijk et al. 1998; van Der Wijk et al. 2000). For the perfused rat liver and intestine 407 cells it could be demonstrated that p38 is activated rapidly and without a detectable lag phase. In IMCD-3 cells, the JNK MAP kinase is activated under hypertonic conditions (van Der Wijk et al. 1998). The swelling-induced activation of JNK is a slow process, occurring with a delay of some 10 min–20 min (van Der Wijk et al. 2000).

Though MAP kinases are activated by the anisotonic alteration of cell volume in many systems, a direct involvement in volume regulatory mechanisms has only been observed in a few systems. In human cervical cancer cells, for example, swelling-activated ERK1/2 is linked to the activation of Cl⁻ currents, K⁺ currents and taurine transport (van Der Wijk et al. 2000; Shen et al. 2001). In contrast, even though Na⁺/H⁺ antiport can be activated in an ERK1/2-dependent manner, the osmotic activation of this antiporter does not involve ERK1/2 (or p38 MAP kinase) in Chinese hamster lung fibroblasts (Aharonovitz and Granot 1996; Bianchini et al. 1997; van Der Wijk et al. 2000). Moreover, Nilius et al. demonstrated that hypotonic activation of MAP kinases are not involved in the activation of volume-sensitive Cl⁻ currents (Nilius et al. 1997).

In most instances, the apparent absence of effects of MAP kinases on the (fast) regulatory movements of inorganic osmolytes can be explained in terms of their slower time course of activation that instead reflects that of organic osmolyte transport and metabolism. Accordingly, osmosensitive MAP kinases may be related to cell volume regulation by activating transcription factors, since the required upregulation of the corresponding metabolizing enzymes and organic osmolyte transporters is realized by enhancing their levels of transcription. An adjusted gene expression thus allows the cells to sustain cell volume disturbances for a prolonged period of osmotic stress.

Macromolecular crowding and ionic strength

Macromolecules occur in a high intracellular concentration; they occupy typically 20%-30% of the total cell volume (Ellis 2001). This noncytosolic and not osmotically active space (see the section entitled "The osmometric behavior of cells") excludes molecules of low molecular weight. This phenomenon, known as macromolecular crowding, has fundamental effects on the rates and equilibria of biochemical reactions (Ellis 2001). Under conditions of increased macromolecular crowding, the reduction of excluded volume causes a decrease of total free energy of the solution. Therefore, the equilibrium constants of binding and association reactions that exclude the least volume to all the other macromolecules present are increased. Macromolecular crowding alters biochemical reaction rates by influencing both molecular diffusion rates and thermodynamic activities. While molecular diffusion rates will be reduced by increased crowding (Luby-Phelps 2000), their thermodynamic activity will be enhanced. The net result of these opposing effects depends on the nature of the rate-limiting step. If a reaction rate is limited by the encounter rate of the initial molecules, the rate will be decreased by crowding due to decreased diffusion rates. If the rate is limited by the activity of the transition complex, it will be increased by crowding due to enhanced thermodynamic activities (Ellis 2001).

Macromolecular crowding is reduced by increasing ionic strength, because proteins interact with the ambient electrolytes. The volume regulatory set points are thereby shifted

to smaller volumes (Parker et al. 1995). Since even slight changes in macromolecular crowding are sufficient to evoke large changes in enzymatic activity, macromolecular crowding has been discussed as a candidate for a cell volume sensory system. Indeed, the volume sensitive transporters Na⁺/H⁺ symporter (Parker et al. 1995), K⁺-Cl⁻ symport (Minton et al. 1992; Minton 1994; Parker et al. 1995), VSOAC channel (Emma et al. 1997), and Cl⁻ channels (Hoffmann and Pedersen 1998; Nilius et al. 1998) are influenced by macromolecular crowding and ionic strength in various systems. A reduction of ionic strength activates some transport systems involved in RVD. An increase in ionic strength is supposed to mediate the induction of sodium organic osmolyte transporters and osmosensitive enzymes during prolonged exposure of cells to hypertonic media (see Burg 1995; and the section entitled "Organic osmolytes in RVI").

Macromolecular crowding and ionic strength are quite global parameters of idealized cells that can not yet be related to various cellular compartments and the place where signal generation and transduction occurs. Thus, their importance in localized reactions remains to be determined.

Cell membrane stretch

Cell swelling causes stretching of the plasma membrane, whereas shrinkage leads to a decrease in cell membrane stretch (Kinnunen 2000). These alterations of membrane tension may generate a signal perceived by stretch-sensitive channels. Among the stretch-sensitive channels, both stretch-activated and stretch-inactivated channels can be found. Stretch-activated channels can either be nonselective cation channels permeable for K⁺, Na⁺ and Ca²⁺, or channels selective for Ca²⁺, K⁺ or Cl⁻ (Hoffmann and Dunham 1995; Lang et al. 1998a).

The activation of *nonselective* stretch-activated cation channels upon cell swelling was demonstrated in several cells (Christensen 1987; Lansman et al. 1987; Falke and Misler 1989; Bear 1990; Christensen and Hoffmann 1992). Activation of these channels contributes to RVD, probably mainly by the resulting Ca²⁺ current (Hoffmann and Dunham 1995; Lang et al. 1998a). The resulting (local) increase in intracellular Ca²⁺ then activates Ca²⁺-sensitive K⁺ channels (Christensen 1987; Ubl et al. 1988). The loss of K⁺ is accompanied by osmotic water movement, leading to volume decrease (see the sections entitled "K⁺ channels" and "Cation channels").

The activation of *selective* stretch-activated ion channels contributes directly to cell volume regulation. Stretch- and volume-sensitive K⁺ channels were demonstrated for *Necturus* proximal tubule cells (Filipovic and Sackin 1992), gall bladder cells (Vanoye and Reuss 1999) and enterocytes (Dubinsky et al. 2000). Similarly, cell membrane stretch may also lead to the activation of stretch-sensitive maxi Cl⁻ channels, which were reported to be activated upon cell swelling in several systems (see the section entitled "Cl⁻ channels").

Stretch-inhibited channels are not as common as stretch-activated channels, and their relevance for cell volume regulation is usually regarded as negligible (Morris 1990; Hoffmann and Dunham 1995). Nevertheless, Suzuki et al. showed, that a cation channel cloned from rat kidney is inactivated by membrane stretch due to cell swelling and activated by cell shrinkage (Suzuki et al. 1999).

The mechanism by which stretch-sensitive channels are regulated by cell volume changes has not yet been fully elucidated. Signals modulating stretch-sensitive channels could be mechanical tension, conveyed by the membrane or the cytoskeleton (Sokabe et al. 1991; Ingber 1997), or the release of stretch-sensitive messengers, such as fatty acids (Kirber et al. 1992), or ATP (Wang et al. 1996).

Though a variety of ion channels were reported to respond to membrane stretch caused by cell volume alterations, the physiological significance of this signal remains questionable, since the activation of these channels requires considerable stretch (Okada 1997). For example, it could be demonstrated that significant cell membrane stretch only arises when cell volume increases by 60%; volume regulatory mechanisms are, however, already activated when cell volume is altered by 5%–10% (Al Habori 2001). Thus, stretch-sensitive channels may only play a role in sensing cell volume changes when excessive cell swelling occurs and other signalling systems have failed (Lang et al. 1998a).

Lipid derived messengers

Phosphoinositides

Lipid signalling through phosphoinositides was reported to be involved in cell survival, cytoskeletal remodeling, metabolic control, and vesicular trafficking (Wymann and Pirola 1998). Since all these processes are related to cell volume regulation (Lang et al. 1998a; Hoffmann and Mills 1999), it seems not surprising that there is ample evidence for the involvement of lipid-derived messengers in processes maintaining osmolyte balance. Phosphoinositides can interfere at different levels with signalling cascades. As signalling intermediates, they can serve as substrate and cofactor for phospholipases or phosphatidylinositol kinases. They are also capable of regulating various enzymes like Rho GTPases and profilin and modulating ion channels and transporters.

For Ehrlich Ascites tumor cells, it could be shown that the level of phosphatidylinositol 4,5-bisphosphate [PI(4,5)P₂] responds to cell volume alterations. A decrease and an increase in cellular PI(4,5)P₂ levels occurs upon cell shrinkage and cell swelling, respectively (Hoffmann and Mills 1999). Similarly, a PI(4,5)P₂ decrease could be shown for rat hepatocytes (Baquet et al. 1991; Vom Dahl et al. 1991) upon cell shrinkage. In Swiss 3T3 mouse fibroblasts, hypertonic stress increases the levels of phosphatidylinositol 3,4-bisphosphate [PI(3,4)P₂] and phosphatidylinositol 3,4,5-trisphosphate [PI(3,4,5)P₃; Van der Kaay et al. 1999].

The effects of cell volume changes on $PI(3,4,5)P_3$ levels, phosphatidylinositol 3-kinase (PI3 K) and phosphatidylinositol 5-phosphatase activity, and the initiated signalling pathways are very variable. In rat skeletal muscle, hypotonicity as well as hypertonicity induce changes of glutamine uptake that are dependent on the activity of PI3 K (Low and Taylor 1998). The authors assume that PI3 K may serve to maintain an integrin-dependent mechanochemical transduction mechanism in an active state. The activity of PI3 K is also required to initiate the hypertonic activation of Na^+ conductance in rat hepatocytes (Wehner et al. 2002b).

In MDCK cells, hyperosmolality activates the serine-threonine-kinase Akt (also known as protein kinase B) in a PI3K-dependent manner (Terada et al. 2001). The product of PI3 K, PI(3,4,5)P₃, interacts with Akt itself as well as with 3-phosphoinositide-dependent kinases (PDK1 and PDK2). The actual activation of Akt occurs via Thr and Ser phosphorylation by PDK1 and PDK2 (Alessi et al. 1997; Stephens et al. 1998).

It was reported that the activation of Akt by hypertonicity prevents hyperosmolality-induced apoptotic changes in MDCK cells (Terada et al. 2001). Together with the finding that the hypotonic activation of PI3 K and Akt in HepG2 cells stimulates proliferation (Kim et al. 2001) ,this suggests that both enzymes influence the osmo-sensitive balance between cell survival and apoptosis.

In Swiss 3T3 mouse fibroblasts, hypertonicity-activated PI3 K elevates $PI(3,4,5)P_3$ levels without enhancing Akt activity, due to negative regulation of Akt by osmotic stress (Van der Kaay et al. 1999). This is coincident with the hypertonic, $PI(3,4,5)P_3$ -dependent inhibition of $PKB\alpha$ in HEK-293 cells caused by inhibition of phosphorylation and activation of dephosphorylation of Akt, probably via PP2A (Meier et al. 1998). The inhibition of Akt seems to be triggered by $PI(3,4,5)P_3$ levels, independent of whether these occur by activation of PI3 K (Van der Kaay et al. 1999) or by inhibition of phosphatidylinositol 5-phosphatase (Meier et al. 1998).

Hypertonic accumulation of PI(3,4,5)P₃ activates p70 S6 kinase in Swiss 3T3 cells (Van der Kaay et al. 1999). This activation of p70 S6 kinase could be related to cell volume regulation on the basis of alterations in glycogen metabolism, since p70 S6 kinase activity is required for osmotically triggered changes in glycogen synthesis in rat skeletal muscle; during RVI, glycogen synthesis is inhibited, whereas it is stimulated during RVD (Low et al. 1996). The effect of cell volume changes on glycogen metabolism has also been demonstrated for rat hepatocytes (Häussinger et al. 1991). In this system, hypotonicity activates both PI3 K and p70 S6 kinase, but only PI3 K is involved in the hypotonic regulation of glycogen synthase (Krause et al. 1996).

The transcription of the serum and glucocorticoid dependent kinase (sgk) is clearly sensitive to cell volume changes. In the human hepatoma cell line HepG2, upregulation of sgk occurs upon cell shrinkage, whereas swelling reduces sgk transcript levels (Waldegger et al. 1997). Furthermore, sgk, like Akt and p70 S6, is a downstream target of PI3 K (Kobayashi and Cohen 1999). Most interestingly, stimulation of sgk results in the activation of ENaC in A6 cells (Chen et al. 1999b), rabbit cortical collecting duct cells (Naray-Fejes-Toth et al. 1999), and in *Xenopus* oocytes (Böhmer et al. 2000). For the latter system, it could also be demonstrated that sgk increases the abundance of the ENaC-protein in the plasma membrane (Alvarez et al. 1999; Wagner et al. 2001). In addition, sgk stimulates the activity of the neuronal K+ channel Kv 1.3 as well as that of Na+-K+-ATPase and may thereby contribute to cell volume regulation (Setiawan et al. 2002; Wärntges et al. 2002).

The phosphorylation sites at which Akt and other protein kinases, such as p70 S6 kinase, PKC δ , and SGK are activated exhibit high similarity. This suggests that PDK1 and PDK2 may contribute equally well to the PI3K-dependent activation of these enzymes (Kobayashi and Cohen 1999).

Moreover, in rat HTC hepatoma cells (Feranchak et al. 1998) and in Mz-Cha-1 human cholangiocarcinoma cells (Feranchak et al. 1999), PI3 K contributes to cell volume regulation by the activation of Cl⁻ secretion via ATP release. The hypotonic activation of Na⁺/H⁺ exchange and HCO₃⁻ absorption in rat renal medullary thick ascending limb (MTAL) requires PI3 K activity as well (Good et al. 2000). The fact, that PI3 K is activated by *hypotonicity* as well as by *hypertonicity* in several systems reflects the general role of PI3 K as a key enzyme in signalling cascades. The actual result of this activation is then defined by its downstream targets that may vary considerably between different cells.

Arachidonic acid and eicosanoids

Arachidonic acid (AA) can be directly released from phosphatidylcholine and phosphatidylchanolamine by the activity of phospholipase A_2 (PLA₂). The cleavage of phosphatidylinositol by phospholipase C yields arachidonic acid out of diacylglycerol (DAG). Phospholipase D produces phosphatic acid, which can be transferred to arachidonic acid by the action of diglyceride lipase. Phosphatic acid may also be hydrolysed by PLA₂ into AA and lysophosphatic acid (LPA).

An increase in the intracellular concentration of arachidonic acid and its derivates upon hypotonic cell swelling has been shown for rat IMCD cells (Tinel et al. 1997), human neuroblastoma cells (Basavappa et al. 1998), and Ehrlich Ascites tumor cells (Thoroed et al. 1997). This release of AA seems to be mainly caused by the activation of PLA₂, as shown for Ehrlich Ascites tumor cells (Thoroed et al. 1997), human platelets (Margalit et al. 1993a), and human neuroblastoma cells (Basavappa et al. 1998).

In agreement with these findings, inhibition of PLA₂ prevents the activation of hypotonically activated events like Cl⁻ current in bovine pigmented ciliary epithelial cells (Mitchell et al. 1997) and iodide efflux in rat brain endothelial cells (von Weikersthal et al. 1997).

Influence on volume regulatory mechanism can be carried out by AA itself or by eicosanoids made from AA. AA is able to directly inhibit volume-sensitive Cl⁻ currents in guinea pig antral gastric myocytes (Xu et al. 1997), rat osteoblast-like cells (Gosling et al. 1996), Ehrlich Ascites tumor cells (Hoffmann and Lambert 1994), human neuroblastoma cells (Basavappa et al. 1998), and rat hepatocytes (Sakai et al. 1996). The activation of C1C-2 Cl⁻channels by AA has been reported for the human lung epithelial cell lines Calu-3, A549, and BEAS-2B (Cuppoletti et al. 2001), and for HEK-293 cells (Tewari et al. 2000). AA is capable of activating K⁺ channels in the colonic secretory cell line T84 (Devor and Frizzell 1998), in rat neuronal cells (Kim et al. 1995), and in rabbit cortical collecting tubule (CCT) cells (Ling et al. 1992).

Via alteration of cytosolic Ca²⁺ (Tinel et al. 1997; Jorgensen et al. 1999), AA can influence Ca²⁺-sensitive Cl⁻ and K⁺ currents, as well as sorbitol efflux in IMCD cells (see the section entitled "Sorbitol release pathways").

The variety of the response of organic osmolyte release pathways in IMCD cells to AA is illustrated in Fig. 8. Here, isotonic taurine and *myo*-inositol efflux is inhibited, whereas AA can mimic the hypotonic response for sorbitol, betaine, and GPC.

AA can be metabolized in several pathways; two of them have been reported to be involved in mediating mechanisms of cell volume regulation. The lipoxygenase pathway leads to formation of leukotrienes and prostanoids, whereas prostaglandins and thromboxanes are formed in the cyclooxygenase pathway by the activity of cyclooxygenase and peroxidase.

Products of the lipoxygenase pathway seem to be involved mainly in the activation of RVD. The 5-LIP metabolite LTD₄ is able to activate hypotonic Cl⁻ and K⁺ currents in Ehrlich Ascites tumor cells (Jorgensen et al. 1996); the isotonic cell shrinkage induced by LTD₄ occurs by the activation of the same mechanisms (Lambert 1989). In addition, LTD₄ acts as a second messenger for the taurine leak pathway and thereby contributes to RVD in this system (Lambert and Hoffmann 1993). An alteration of K⁺ channels by 5-LIP product has also been shown for mudpuppy red blood cells (Light et al. 1997). In human skin fibroblasts, the activity of 5-LIP is required for the activation of swelling-induced Cl⁻

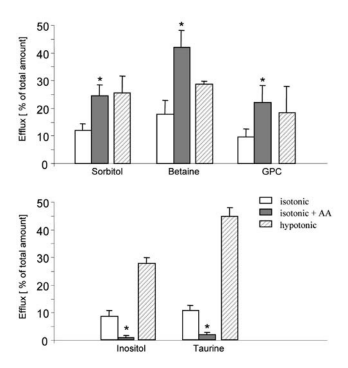


Fig. 8 Effect of arachidonic acid (AA) and hypotonicity on osmolyte efflux in rat inner medullary collecting duct (IMCD) cells. Efflux of osmolytes (in % of total amount; \pm SE/10 min) was determined after 10 min exposure to 10 μ M AA at isotonicity or after a decrease of extracellular osmolarity ($600\rightarrow300$ mOsm/l) by means of HPLC; n=4 for all experimental conditions (B. Ruhfus and R.K. Kinne, unpublished data)

efflux (Mastrocola et al. 1993). The 12-LIP product hepoxilin A₃ is involved in the hypertonic activation of K⁺ channels in human platelets (Margalit et al. 1993b).

In Ehrlich Ascites tumor cells, the stimulation of leukotriene synthesis is concomitant with an inhibition of prostaglandin synthesis (Lambert et al. 1987). The acceleration of RVD by LTD₄ or AA and the deceleration by prostaglandin E₂ (PGE₂) in mouse colonic crypts (Mignen et al. 1999) confirms that the reciprocal alteration of lipoxygenase and cyclooxygenase pathway affect synergistically volume recovery in these systems. In Ehrlich Ascites tumor cells, PGE₂ has no effect on the swelling-induced Cl⁻ permeability or K⁺ permeability (Lambert et al. 1987). However, since PGE₂ is able to activate Na⁺ channels, a decrease in PGE₂ synthesis may contribute to volume decrease by a reduction in the passive membrane conductance for Na⁺ (Hoffmann 1978). In MDCK cells (Steidl et al. 1991) and human erythrocytes (Li et al. 1996), PGE2 has been reported to activate K⁺ channels resulting in a decrease of cell volume. Cl⁻ channels can be activated by cyclooxygenasedependent PGE₂ synthesis in the hepatoma cell line HTC (Kilic and Fitz 2002). For human amnion-derived WISH cells, it could be shown that COX-2 gene expression is upregulated by hypotonic stress; as a consequence thereof, PGE2 release is increased (Lundgren et al. 1997). Of note, there is no evidence for the involvement of PLA₂-signalling in cell volume restoration upon hypertonic stress.

An involvement of phospholipase C (PLC) in cell volume regulation has been reported after hypotonic cell swelling, as well as after hypertonic cell shrinkage. PLC can influence

volume regulatory processes by different mechanisms, depending on the products of the reaction.

Ca²⁺ oscillations can be IP₃– as AA-sensitive as well (Ling et al. 1992; Tinel et al. 1997); both messengers result from PLC activity. Involvement of PLC in hypotonic Ca²⁺ oscillations has been reported for rat IMCD cells (Tinel et al. 1997), rat caudal artery endothelial cells (Shinozuka et al. 2001), rabbit proximale tubule cells (O'Neil and Leng 1997), and bovine aortic endothelial cells (Oike et al. 2000). The relevance of Ca²⁺ signalling for cell volume regulation is discussed in detail in the section entitled "Intracellular calcium."

In pigmented ciliary epithelial cells, volume-sensitive Cl⁻ currents are dependent on the activity of both PLA₂ and PLC. The contribution of PLC may be the synthesis of AA or enhancement of PLA₂ activity via a PKC-dependent phosphorylation of PLA₂ (Mitchell et al. 1997).

Phosphatic acid and lysophosphatic acid

Phospholipase D (PLD) generates the lipid second messengers phosphatic acid (PA) and choline by the cleavage of phosphatidylcholine. Only few examples exist for the involvement of PLD in volume-regulatory signalling. In skate erythrocytes, hypotonic treatment increases the turnover of phosphatidylcholine and the formation of PA due to the activation of PLD (Musch and Goldstein 1990). Sorbitol-induced osmotic stress enhances activity of PLD and PA release in rat adipocytes, leading to activation of aPKCs and translocation of GLUT4 transporters (Sajan et al. 2002). The PIP₂-synthesizing enzyme PIP 5-kinase can be regulated by PA; therefore, PLD may play a signalling role in PIP₂ sensitive events (Cockcroft 2001). Since alterations of PIP₂ levels are related to cell volume regulation, PLD may contribute to volume-dependent phosphoinositide signalling.

PA can be metabolized to AA by the action of PLD, but the relevance of PLD for volume-induced AA metabolism seems to be minor. In Ehrlich Ascites tumor cells, the contribution of PLD to the hypotonic AA release could be excluded (Thorough et al. 1997).

Enzymatic conversion of PA by extracellular PLA₂ yields lysophosphatic acid (LPA); the initialization of intracellular signalling cascades occurs by binding of LPA to G protein coupled receptors (Goetzl and An 1998). For Ehrlich Ascites tumor cells, it could be revealed that extracellular application of LPA caused cell shrinkage by evoking Ca²⁺ oscillations and activation of Ca²⁺-sensitive K⁺ and Cl⁻ currents. As a consequence of this volume decrease, the Na⁺/H⁺ exchanger is activated (Pedersen et al. 2000). On the basis of the similarity between cell signalling after hypertonic cell shrinkage and after LPA treatment, the authors assume an interrelation in the regulation of both events.

Diacylglycerol

In erythrocytes, the intracellular level of diacylglycerol (DAG) is increased by hypotonicity (Musch and Goldstein 1990). DAG is a product of PLC and able to activate protein kinase C (PKC). PKC is a serine/threonine kinase of central importance in intracellular signalling cascades; hence, it is not astonishing that PKC is activated upon hypotonic cell swelling as well as upon hypertonic cell shrinkage in various systems.

Contribution of PLC to RVI after hypertonic stress has been reported for NIH/3T3 mouse fibroblasts (Zhuang et al. 2000) and Ehrlich Ascites tumor cells (Jensen et al. 1993); in both systems, PLC acts via PKC. In NIH/3T3 cells, the hypertonic phosphorylation of ERK 1/2 is due to DAG-dependent activation of classical and novel PKCs (Zhuang et al. 2000). Hypertonic activation of PLC enhances Na⁺/K⁺/2Cl⁻ cotransport activity in Ehrlich Ascites tumor cells via Ca²⁺-dependent PKC (Jensen et al. 1993).

In Ehrlich Ascites tumor cells, hypertonicity-activated PKC contributes to RVI by activating NHE and NKCC, and by inhibiting volume-sensitive Cl⁻ currents (Larsen et al. 1994; Hoffmann and Dunham 1995; O'Donnell et al. 1995; Pedersen et al. 1996). Inhibition of volume-sensitive Cl⁻ currents by PKC has also been reported for human nonpigmented ciliary epithelial cells (Coca-Prados et al. 1996), for cardiac ClC-3 expressed in NIH/3T3 cells (Duan et al. 1997), and HeLa cells (Hardy et al. 1995); however, in other systems, PKC may either stimulate chloride currents or leave them unaffected (reviewed in Pasantes-Morales et al. 2000a). The contribution of PKC to shrinkage-induced activation of NKCC is also cell-specific. While NKCC is downregulated by PKC in sweat glands (Toyomoto et al. 1997), NKCC remains unaffected in bovine endothelial cells (O'Donnell et al. 1995). PKC-dependent activation of NKCC has been demonstrated for rat hepatocytes; in this system, hypertonic Na⁺ currents are activated by PKC, while hypertonicity activated NHE is not regulated by PKC (Heinzinger et al. 2001). In NIH/3T3 cells, hyperosmolality enhances the activity of c and n PKC, leading to activation of ERK1/2 (Zhuang et al. 2000).

After osmotic cell swelling, PKC is activated in the late phase of RVD in Ehrlich Ascites tumor cells; inhibition of VSCC by PKC terminates the volume response (Larsen et al. 1994). In contrast, hypotonic Cl⁻ membrane permeability is enhanced by PKC in HTC rat hepatoma and Mz-ChA-1 human cholangiocarcinoma cells (Roman et al. 1998).

The swelling-induced release of organic osmolytes is also influenced by the hypertonic activation of PKC. While inhibition of PKC prevents the hypertonic taurine release in rat cortical cup model (Estevez et al. 1999), the osmo-sensitive taurine release in cerebellar granule neurons is only prevented after chronic downregulation of PKC (Morales-Mulia et al. 2001). Thus, lipid-derived messengers are certainly involved in the cellular response to osmotic challenges. The complexity of the networks they are part of precludes, however, prediction of the outcome of the experiments aimed at elucidating their physiological role. Apparently, various cells have developed different responses; the reason for this phenomenon may lie in their differentiation into functionally diverse entities.

Cyclic nucleotides

The intracellular amounts of cAMP, as well as cAMP-dependent phosphorylation, have been demonstrated to be unaffected by osmotic cell shrinkage in duck and turkey erythrocytes (Kregenow et al. 1976; Alper et al. 1980). Initiation of the cAMP pathway is more likely to inhibit RVI, since cAMP-dependent activation of PKA can prevent the uptake of the organic osmolytes betaine and myo-inositol, as shown for MDCK cells (Preston et al. 1995). The hypertonic increase of intracellular cAMP suppresses the activation of Erk and p38; both MAP-kinases are known to be activated after hypertonic cell shrinkage in various systems (Orlic et al. 2002). However, cAMP is able to evoke cell swelling in sweat glands (Ohtsuyama et al. 1993). Osmotic cell swelling has been found to increase intracel-

lular cAMP levels in various systems and cAMP leads to cell volume decrease (reviewed in Lang et al. 1998a).

In rectal gland tubular cells from *Squalus acanthias*, cAMP-mediated cell shrinkage occurs by stimulation of NaCl excretion. A PKA-dependent phosphorylation of CFTR-type Cl⁻ channels is followed by activation of Na⁺-K⁺-2Cl⁻ cotransporter (Greger 1996). In the human hematopoietic myeloblastic leukemia cell line ML-1, outward-rectified Cl⁻ channels are activated by PKA (Xu and Lu 1994). The activation of volume-sensitive Cl⁻ currents is inhibited by cAMP in myocardial cells, indicating that activation of these channels requires dephosphorylation of a cAMP-dependent phosphorylation site (Hall et al. 1995; Nagasaki et al. 2000).

Although hypotonicity increases intracellular cAMP levels in S49 mouse lymphoma cells, cAMP is not involved in triggering volume regulatory processes (Watson et al. 1991). As shown for the hypotonic myo-inositol efflux in human NT2-N neurons, PKA does not contribute to the osmotic regulation of this transporter (Novak et al. 2000). The efflux of organic osmolytes can be inhibited by PKA, since inhibition of PKA enhances the speed of the hypotonic activation of anionic amino acid conductance in *Leishmania major* promastigotes (Vieira et al. 1997).

Intracellular calcium

An involvement of calcium in volume regulation has been found in many cellular systems. In particular, the regulation of cell swelling seems to often be accompanied by calcium signals. In the case of hypertonic stress and cell shrinkage, other intracellular signal transduction processes seem to be responsible for the recovery of the cell volume and thus the reports about an involvement of calcium in RVI are very rare (Marchenko and Sage 2000; Erickson et al. 2001). In this overview, we will focus, therefore, on the role of calcium during RVD.

The core process of volume regulation after cell swelling is a release of osmolytes, resulting in reduction of cellular water content. Many different transport processes involved in RVD have been found to be calcium-dependent. Calcium-regulated potassium conductance via BK_{Ca}, IK_{Ca}, and SK_{Ca} have been found in many cells (see the section entitled "K+ channels"). Also chloride conductances induced by volume changes have been observed to be regulated by calcium/calmodulin-dependent processes (see the section entitled "Cl- channels"). In guinea pig jejunal villus epithelial cells, a calcium/calmodulin kinase II-mediated phosphorylation is a critical determinant of the volume regulation (MacLeod and Hamilton 1999a). In some cells, the calcium action on the same transport system seems to be transmitted via different effector proteins; for example, in human cervical cancer cells, myosin light-chain kinase and protein kinase C regulate a volume-sensitive chloride channel (Chou et al. 1998; Shen et al. 2002). In astrocytes, chloride conductance as well as taurine release are processes regulated by calcium and calmodulin (Li et al. 2002). In rat IMCD cells, calcium is involved in the release of sorbitol (Ruhfus et al. 1996). Recently, a coupling between arginine vasopressin-induced intracellular calcium mobilization and apical exocytosis was investigated in isolated perfused rat IMCD segments (Yip 2002). Vasopressin induced a rapid increase of the intracellular calcium followed by sustained calcium oscillations and changes in cell volume. Apical exocytosis accompanied by an increase of osmotic water permeability could be prevented by an inhibition of intracellular calcium release. The study shows that in IMCD cells vasopressin, via

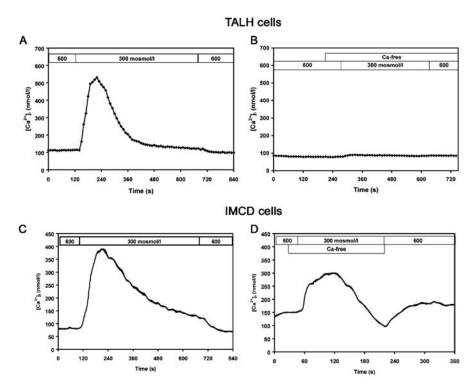


Fig. 9 Changes in the intracellular calcium concentration of TALH cells (**A**, **B**) and IMCD cells (**C**, **D**). At time indicated osmolarity was reduced from 600 mOsm/l to 300 mOsm/l by omission of sucrose. The experiments in calcium-containing solutions show a sustained calcium increase in both TALH (**A**) and IMCD (**C**) cells. In low-calcium solutions there was a calcium increase in the IMCD-cells due to a calcium release from intracellular stores (**D**). In TALH cells (**B**) no changes of the calcium concentration could be observed in calcium-free solutions as a prevention of the calcium influx made the calcium-induced calcium release impossible (**A** and **B** from Tinel et al. 2002; **C** and **D** from Tinel et al. 1994)

the V₂ receptor, triggers calcium oscillations which are required for exocytotic insertion of aquaporin-2 (Yip 2002). Activation of the sorbitol channel in rabbit TALH cells involves phosphorylation by a calcium/calmodulin kinase (Kinne-Saffran and Kinne 1997).

The source of the calcium involved in the volume regulation seems to be cell type-dependent. In several cells, RVD strictly depends on calcium entry across the plasma membrane. In these systems, the volume response can be inhibited by extracellular calcium removal or by addition of calcium channel blockers. A swelling-induced rise in intracellular calcium in rabbit medullary thick ascending limb cells was absent in low-calcium solutions (Montrose-Rafizadeh et al. 1991; see Fig. 9).

Similarly, strict dependence on calcium entry was found in, e.g., osteosarcoma cells (Yamaguchi et al. 1989), rabbit proximal tubule cells (Suzuki et al. 1990), intestine 407 cells (Hazama and Okada 1988), and cultured mouse primary sensory neurons (Viana et al. 2001). The diversity of calcium channels which are responsible for the calcium influx during hypotonic stress in different cell types is very high. Cell swelling induces stretch of the plasma membrane; thus, it is possible that stretch-activated calcium channels are activated by this process. Indeed, there are many reports showing that Gd³⁺, a nonspecific in-

hibitor of stretch-activated channels, can block the calcium increase during hypotonic stress (Chen et al. 1996b; Urbach et al. 1999; Miyauchi et al. 2000; Viana et al. 2001).

Stretch-activated calcium influx might, in some cells, be mediated by voltage-sensitive calcium pathways, as it has been shown in osteocytes (Miyauchi et al. 2000). In these cells, antisense oligodeoxynucleotides against the α 1C subunit of voltage-operated L-type calcium channels abolished the calcium increase during hypotonic stress. The expression of the α 1C subunit of voltage-operated L-type calcium channels has also been shown in UMR-106 cells (Kizer et al. 1999). The existence of a voltage-activated calcium L-type channel which may participate in the volume regulation has also been identified in the apical membrane of cultured rabbit proximal tubule cells using the patch-clamp technique (Zhang and O'Neil 1996). In many other cell types the calcium increase during hypotonic stress was sensitive to blockers of voltage-activated calcium channels (Montrose-Rafizadeh and Guggino 1991; Bender et al. 1994; Mignen et al. 1999). In inner medullary collecting duct cells (IMCD), diltiazem, nifedipine, and verapamil partly reduced the calcium increase (Tinel et al. 2000). However, the concentration of the blocker was quite high (40 μMol/l). Thus, calcium influx seems to occur via calcium channels which possess pharmacological characteristics different from those of classical voltage-gated channel. The fact that an inhibitor of receptor-mediated channels (SKF 96365) also reduced the calcium influx in IMCD cells supports the idea (Tinel et al. 2000). Channels involved in hypotonic calcium influx in thick ascending limb of Henle's loop (TALH) cells could be inhibited by nifedipine, but not diltiazem or verapamil (Tinel et al. 2002). These results can be correlated with the existence of homologous genes for different types of subunits and several types of auxiliary subunits of voltage-dependent calcium channels (Isom et al. 1994). It can also explain the difference compared to the characteristics of the calcium influx observed in TALH cells, which were not adapted to higher osmolarity (Montrose-Rafizadeh and Guggino 1991). In these cells, hypotonicity-induced calcium entry occurred via a nifedipine- and verapamil-sensitive pathway.

A complete inhibition of the calcium increase by calcium channel blockers or lack of the calcium signal in calcium-free solution has often been interpreted as proof that the intracellular calcium stores are not involved in the calcium rise during hypotonic stress. The assumption would be correct only if no ryanodine-sensitive calcium stores exist in the cell. Using caffeine, it has been shown that, e.g., TALH cells possess calcium stores that, although at first sight not involved in the volume regulation, can be mobilized experimentally in isotonic solution (Tinel et al. 2002). Caffeine causes calcium release from ryanodine-sensitive stores. The ryanodine receptor in the membrane of these stores is sensitive to calcium, which in micromolar concentration stimulates the receptor and triggers a calcium release (Tinel et al. 2000). This phenomenon is referred to as calcium-induced calcium release (CICR), a mechanism responsible for, e.g., heart muscle contraction. In TALH cells, the existence of CICR during hypotonic stimulation has been shown using manganese quenching experiments (see Figs. 9, 10).

Calcium release from intracellular stores followed the calcium influx during hypotonic stress (Tinel et al. 2002). The involvement of CICR in volume regulation has already been described in a human intestinal epithelial cell line (Intestine 407; Hazama and Okada 1990). For the volume regulation of these cells, the calcium influx is necessary, but not sufficient, unless it elicits CICR. In rabbit corneal epithelial cell, hypotonic stress has been found to induce a calcium release from ryanodine-sensitive stores followed by activation of a calcium influx (Wu et al. 1997). The authors describe the mechanism as CICR; how-

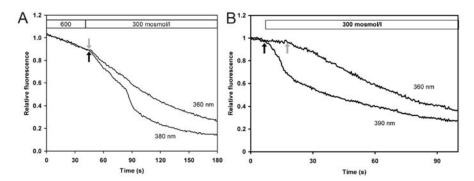


Fig. 10 Quench of fura-2 fluorescence by manganese under hypotonic conditions. The osmolarity was reduced from 600 mOsm/l to 300 mOsm/l by omission of sucrose. A In TALH cells, the hypotonic stress in calcium-containing solutions induced a monoexponential decay of the fluorescence intensity at 360 nm, reflecting a constant calcium influx. At 380 nm, the fluorescence decay was biphasic. During the first 40 s, both manganese and calcium entered the cell and induced a reduction of the fluorescence intensity. The additional fast decay was due to calcium-induced calcium release. B In IMCD cells hypotonic stress in calcium-free solutions induced a fast decay of the fluorescence at 390 nm due to calcium release from intracellular stores followed by an opening of a calcium influx pathway visible as a decrease in the fluorescence at 360 nm. → start of increase in intracellular calcium, ⇒ start of calcium-entry from the extracellular medium (A from Tinel et al. 2002; B from Tinel et al. 1994)

ever, it corresponds more to the store-operated calcium influx. The RVD in these cells was also strictly dependent on the calcium rise. In TALH cells, the volume regulation was only partly calcium-dependent; thus, the calcium release from intracellular stores probably works as an amplifier of the signal. CICR is very important for the generation of calcium oscillations and calcium waves (Tinel et al. 2000). In some cells, there are initiation sites where the signal appears first before spreading through the cell as a wave (Tinel et al. 1999). Such local calcium events during the volume regulation of TALH cells could control membrane recycling, as has been described to occur in IMCD cells (Czekay et al. 1994). During exposure of IMCD cells to hypotonicity, the submembranal actin web rapidly disintegrates, and reserve vesicles, probably containing a sorbitol transporter, move to and fuse with the basolateral plasma membrane. The fusion causes a rapid increase in sorbitol permeability (see Fig. 4 above). These membrane areas are recovered by internalization, and the transport systems for sorbitol are concomitantly retrieved (Czekay et al. 1994).

The majority of studies concerning the role of calcium in volume regulation describe a biphasic calcium rise consisting of a calcium influx followed by calcium release from intracellular stores which are not ryanodine-sensitive. Manoeuvres leading to depletion of the intracellular stores or inhibiting the calcium release abolish the calcium increase in these cells during hypotonic stress (Hoffmann et al. 1984; McCarty and O'Neil 1991; Bender et al. 1994; Civan et al. 1994a; Mignen et al. 1999). Intracellular stores seem to participate in the shaping of the calcium signal when the changes of the cell volume occur in isotonic solution, e.g., in Ehrlich ascites tumor cells after stimulation with lysophosphatidic acid (Pedersen et al. 2000), or in pneumocytes after pretreatment with insulin (Marunaka et al. 1999). In some of the reports, the calcium stores are described as being inositol-trisphosphate (IP₃)-sensitive. An increase of the intracellular concentration of IP₃ has been measured during isotonic swelling of hepatocytes (Baquet et al. 1991), and during volume

regulation of rabbit proximal tubule cells (Suzuki et al. 1990) and astrocytes (Bender et al. 1993).

Although in IMCD cells the calcium rise could be blocked by inhibitors of the calcium release from IP₃-sensitive stores, the signal transduction pathways leading to the calcium mobilization seem to be different (Tinel et al. 1994). In these cells, cell swelling in hypotonic solution is paralleled by an increase in intracellular calcium concentration typically appearing in two distinct phases. Whereas the first phase of the calcium response reflects predominantly calcium release from intracellular stores, the second sustained component is due to an increase in calcium entry from the extracellular compartment (Mooren and Kinne 1994; Tinel et al. 1994; see Figs. 9 and 10). However, measurement of intracellular IP₃ revealed no significant changes in the concentration of this second messenger (Tinel et al. 1997). Instead, inhibition of the arachidonic acid metabolism led to a considerable reduction of calcium release, as well as RVD. Thus, in these cells the calcium mobilization during RVD may be initiated by arachidonic acid or by its metabolites. Under isotonic conditions, an application of arachidonic acid induces a considerable increase in the intracellular calcium concentration in IMCD cells, both in high and in low calcium solutions, indicating that IMCD cells posses arachidonic acid-sensitive calcium stores. Extracellular ATP induces in these cells an increase in the intracellular IP₃ concentration and calcium rise, which reveals the existence of functionally active IP₃-sensitive stores. Thus, IMCDcells possess arachidonic acid-sensitive as well as IP3-sensitive calcium stores. These stores may be physically distinct; alternatively, the arachidonic acid-sensitive stores may be a part of the IP₃-sensitive pool. This is not contradictory to the finding that stimulation of calcium release by IP₃ during volume regulation apparently does not occur in IMCD cells. Arachidonic acid could change the affinity of the IP₃-receptor for IP₃ so that the IP₃ concentration already present in the cytosol of nonstimulated cells might induce a calcium release. It is possible that cooperation of different calcium stores control the volume regulation of these cells.

A combination of two different calcium-releasing mechanisms in the regulation of the cell volume has also been found in astrocytes (Bender and Norenberg 1994). In these cells, intracellular calcium release from CICR-stores and from IP₃-sensitive intracellular calcium stores play a role in the mechanism of potassium release under hyposmotic conditions. In submandibular gland acinar cells, the calcium increase is a result of a calcium release from agonist-sensitive stores and from mitochondria (Park 2002). Mitochondria are known to participate in modulating calcium signals in other cells (Hoffmann et al. 1984), and it is possible that they fulfill this function also during volume regulation, but this question remains to be elucidated.

The signal transduction processes leading to generation of the second messengers and to the calcium release (IP₃ or arachidonic acid) involves probably the action of phospholipases. Inhibitors of the phospholipase A_2 impaired the volume regulation of, e.g., human neuroblastoma cells (Basavappa et al. 1998) and human platelets (Margalit et al. 1993a). In cultured rabbit principal cells, apical application of the phospholipase A_2 agonist increased single-channel open probability of a volume-sensitive potassium channel (Ling et al. 1992). In IMCD cells, the activation of the arachidonic acid cascade during RVD occurs via activation of phospholipase C and probably phospholipase A_2 (Mooren and Kinne 1994; see also the section entitled "Arachidonic acid and eicosanoids").

Usually phospholipases are activated by G-proteins. In IMCD cells, G-proteins are involved in the generation of the calcium signal (Mooren and Kinne 1994). Involvement of

G-proteins in the regulation of RVD has also been reported to occur in human platelets (Margalit et al. 1993a), and inhibition of RVD by pertussis toxin has been described in proximal tubule cells (Suzuki et al. 1990).

How G-proteins are actually activated during osmotic swelling remains obscure. A mechanoreceptor which could serve as a volume sensor has only been described for neurons (Oliet and Bourque 1993). A mechanical-biochemical coupling involving G-proteins and phospholipase A₂ during cell swelling has been postulated for platelets (Margalit et al. 1993a). OTRPC4, a nonselective calcium-permeable channel which belongs to the TRP channel family, has been proposed to be a candidate for a molecular sensor that confers osmosensitivity in mammalian cells (Strotmann et al. 2000). It exhibits spontaneous activity in isotonic media and is rapidly activated by a decrease (and is inhibited by an increase) of the extracellular osmolarity. Changes in osmolarity of as little as 10% result in significant changes in intracellular calcium concentration in HEK293 cells expressing the channel. This increase resulted exclusively from calcium influx from the extracellular space, with no involvement of calcium release from intracellular stores. In contrast to the members of the classical TRP channel family that are activated secondary to the activation of phospholipase C-coupled receptors, the OTRPCs have been proposed to be involved in the transduction of physical and chemical stimuli. However, OTRPC4 activity was unaffected by pressure-induced stretching of the membrane. The authors propose the channel as a universal sensor in mammalian cells, but the results from other studies where calcium signals could be abolished by inhibition of different steps of the signal transduction contradict this assumption.

In some cells, an autocrine (or paracrine) action of ATP has been found to initiate the calcium signalling during the volume regulation. During osmotic swelling, intracellular ATP is released to the extracellular space in a number of cells. In the immediate vicinity of the cell surface, released ATP has been shown to reach a concentration high enough to simulate P₂-purinergic receptors in a human epithelial cell line, intestine 407 (Dezaki et al. 2000). RVD in these cells is facilitated by ATP by augmenting intracellular calcium rise via the stimulation of purinergic P_{2Y2} receptors. Similar observations were made in endothelial cells from the rat caudal artery (Shinozuka et al. 2001). In hepatocytes, the signalling pathways seem to be more complicated. Ferenchak et al. have found a constitutive and volume-dependent ATP release, which was a critical determinant of membrane chloride permeability. P2 receptor antagonist suramin abolished the volume recovery after exposure of hepatocytes to hypotonic buffer (Feranchak et al. 2000). However, Roe et al. have shown that the swelling-induced calcium signal in hepatocytes was unaffected by either extracellular ATP depletion or blockade of P₂ receptors (Roe et al. 2001). The swelling elicited an increase of the intracellular calcium, which was essential for ion channel activation and volume recovery, but this increase did not stem from activation of volumesensitive P2 receptors. Taken together, the results from both groups indicate that in hepatocytes two different pathways initiate the calcium signalling during hypotonic stress: a purinergic-dependent and a purinergic-independent pathway.

The mechanisms guaranteeing the interplay between the calcium release and calcium influx are not very well understood. In IMCD cells, the calcium release from the stores is a prerequisite for the activation of the calcium influx (Tinel et al. 1994). The opening of the plasma membrane ion channels happens possibly to prolong the calcium response beyond the point where intracellular stores are depleted and/or to support the refilling of these stores after calcium depletion. If the calcium release takes place in calcium-free iso-

tonic solutions, a readdition of calcium to the extracellular medium induces a sustained increase of the intracellular calcium concentration. This indicates that the IMCD cells possess a capacitative calcium influx mechanism. The activation of this pathway is not dependent on an increase of the intracellular calcium concentration. Thus, there has to be another signal activating the calcium channels in the plasma membrane. In IMCD cells, the cytoskeleton could forward the message from the empty stores to the plasma membrane as Mooren and Kinne have observed a complete inhibition of the hypotonic calcium increase by cytochalasin B (Mooren and Kinne 1994). The signal transduction from the stores to the plasma membrane takes some 17 s, as could be estimated using the manganese quenching technique (Tinel et al. 1994; see Fig. 10).

In proximal straight tubule, a temporary dependence of the RVD on calcium has been observed (McCarty and O'Neil 1991). There is a short period of time (about 60 s), during which extracellular calcium is required. Outside of this "calcium window," RVD would inactivate and could not be reactivated by subsequent addition of calcium. It was found that the calcium permeability did not inactivate over several minutes, indicating that the temporary dependence of RVD on extracellular calcium is not due to the transient activation of calcium entry pathway (McCarty and O'Neil 1991).

A very interesting new aspect of the interaction between calcium stores and channels has been found by Lemonnier et al. in human prostate cancer epithelial cells (Lemonnier et al. 2002). In these cells, although variations of both intracellular and extracellular calcium concentrations had no visible effects on a volume-regulated anion channel, calcium influx via store-operated channels strongly modulates it. The authors suggest that interaction between the volume-regulated anion channel and calcium occurs in the confined compartments at the inner surface of the membrane that are not accessible to changes of global intracellular calcium. These domains can be readily reached by calcium entering the cell via plasma membrane, especially through the store-operated calcium channels. This preferred access of store-operated calcium channels to the volume-regulated anion channels suggests colocalization of these channels in the cell membrane (Lemonnier et al. 2002).

This finding belongs to the rare results illustrating the importance of the spatio-temporal aspect of calcium signalling. The intensive work during the last few years showed that, e.g., in secretory cells, the global calcium rise observed in many studies is mainly due to experimental manipulations and that local calcium events (sparks, oscillations, waves) reflect what actually happens under physiological conditions (Tinel et al. 1999; Tinel et al 2000). Future work concerning calcium signalling during volume regulation should concentrate much more on the local calcium events. It would be very important to recognise the mechanisms confining the signals in time and space to allow regulation of specific volume-related processes without influencing other cellular processes influenced by calcium.

Conclusions and perspectives

The data presented above document once more the complexity of events that ultimately result in the maintenance of cell volume and its restoration after disturbance by an osmotic difference between intra- and extracellular space. This complexity, although difficult to decipher, provides different cells with various mechanisms that are adapted to protect themselves and at the same time to fulfill their overall role in the organism. Within one

particular cell, the presence of multiple systems might open multiple possibilities to achieve a graded response to osmotic stimuli.

To obtain further insight into this complexity several steps at different levels of resolution are necessary. At the level of the single molecules, proper molecular and pharmacological identification, proper biophysical characterization, and proper analysis of biochemical alterations have to be continued and intensified. At the cellular level, the simultaneous measurement of the activities of the most important osmolyte transporters and the concentration of the various osmolytes at a higher temporal and spatial resolution have to be performed. The same holds for the determination of the localization and activity of the various elements of the signal transduction network.

Only then will it be possible to describe the dynamic nature of cell volume regulation and its underlying mechanisms in a quantitative way based on causal relationships.

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References

- Adelman JP, Shen KZ, Kavanaugh MP, Warren RA, Wu YN, Lagrutta A, Bond CT, North RA (1992) Calcium-activated potassium channels expressed from cloned complementary DNAs. Neuron 9:209–216
- Aharonovitz O, Granot Y (1996) Stimulation of mitogen-activated protein kinase and Na⁺/H⁺ exchanger in human platelets. Differential effect of phorbol ester and vasopressin. J Biol Chem 271:16494–16499
- Al Habori M (2001) Macromolecular crowding and its role as intracellular signalling of cell volume regulation. Int J Biochem Cell Biol 33:844–864
- Alessi DR, Deak M, Casamayor A, Caudwell FB, Morrice N, Norman DG, Gaffney P, Reese CB, Mac-Dougall CN, Harbison D, Ashworth A, Bownes M (1997) 3-Phosphoinositide-dependent protein kinase-1 (PDK1): structural and functional homology with the Drosophila DSTPK61 kinase. Curr Biol 7:776–789
- Alper SL, Palfrey HC, DeRiemer SA, Greengard P (1980) Hormonal control of protein phosphorylation in turkey erythrocytes. Phosphorylation by cAMP-dependent and Ca²⁺-dependent protein kinases of distinct sites in goblin, a high molecular weight protein of the plasma membrane. J Biol Chem 255:11029–11039
- Alvarez de la Rosa D, Zhang P, Naray-Fejes-Toth A, Fejes-Toth G, Canessa CM (1999) The serum and glucocorticoid kinase sgk increases the abundance of epithelial sodium channels in the plasma membrane of Xenopus oocytes. J Biol Chem 274:37834–37839
- Amano M, Chihara K, Kimura K, Fukata Y, Nakamura N, Matsuura Y, Kaibuchi K (1997) Formation of actin stress fibers and focal adhesions enhanced by Rho-kinase. Science 275:1308–1311
- Amano M, Ito M, Kimura K, Fukata Y, Chihara K, Nakano T, Matsuura Y, Kaibuchi K (1996) Phosphorylation and activation of myosin by Rho-associated kinase (Rho- kinase) J Biol Chem 271:20246– 20249
- Atkinson NS, Robertson GA, Ganetzky B (1991) A component of calcium-activated potassium channels encoded by the Drosophila slo locus. Science 253:551–555
- Atta MG, Dahl SC, Kwon HM, Handler JS (1999) Tyrosine kinase inhibitors and immunosuppressants perturb the myo-inositol but not the betaine cotransporter in isotonic and hypertonic MDCK cells. Kidney Int 55:956–962
- Awayda MS, Subramanyam M (1998) Regulation of the epithelial Na⁺ channel by membrane tension. J Gen Physiol 112:97–111

- Bagnasco S, Balaban R, Fales HM, Yang YM, Burg M (1986) Predominant osmotically active organic solutes in rat and rabbit renal medullas. J Biol Chem 261:5872–5877
- Bagnasco SM, Murphy HR, Bedford JJ, Burg MB (1988) Osmoregulation by slow changes in aldose reductase and rapid changes in sorbitol flux. Am J Physiol 254:C788–C792
- Bai L, Collins JF, Muller YL, Xu H, Kiela PR, Ghishan FK (1999) Characterization of cis-elements required for osmotic response of rat Na⁺/H⁺ exchanger-2 (NHE-2) gene. Am J Physiol 277:R1112–R1119
- Banderali U, Roy G (1992a) Activation of K⁺ and Cl⁻ channels in MDCK cells during volume regulation in hypotonic media. J Membr Biol 126:219–234
- Banderali U, Roy G (1992b) Anion channels for amino acids in MDCK cells. Am J Physiol 263:C1200–C1207.
- Baquet A, Meijer AJ, Hue L (1991) Hepatocyte swelling increases inositol 1,4,5-trisphosphate, calcium and cyclic AMP concentration but antagonizes phosphorylase activation by Ca²⁺-dependent hormones. FEBS Lett 278:103–106
- Barhanin J, Lesage F, Guillemare E, Fink M, Lazdunski M, Romey G (1996) KvLQT1 and IsK (minK) proteins associate to form the IKs cardiac potassium current. Nature 384:78–80
- Basavappa S, Pedersen SF, Jorgensen NK, Ellory JC, Hoffmann EK (1998) Swelling-induced arachidonic acid release via the 85-kDa cPLA2 in human neuroblastoma cells. J Neurophysiol 79:1441–1449
- Bauernschmitt HG, Kinne RKH (1993) Metabolism of the "organic osmolyte" glycerophosphorylcholine in isolated rat inner medullary collecting duct cells. II. Regulation by extracellular osmolality. Biochim Biophys Acta 1150:25–34
- Bear CE (1990) A nonselective cation channel in rat liver cells is activated by membrane stretch. Am J Physiol 258:C421–C428
- Bear CE (1991) A K⁺-selective channel in the colonic carcinoma cell line: CaCo-2 is activated with membrane stretch. Biochim Biophys Acta 1069:267–272
- Beck F, Dörge A, Rick R, Thurau K (1985) Osmoregulation of renal papillary cells. Pflugers Arch 405 Suppl 1, S28–S32
- Beck FX, Burger-Kentischer A, Müller E (1998) Cellular response to osmotic stress in the renal medulla. Pflugers Arch 436:814–827
- Bedford JJ, Bagnasco SM, Kador PF, Harris HW Jr., Burg MB (1987) Characterization and purification of a mammalian osmoregulatory protein, aldose reductase, induced in renal medullary cells by high extracellular NaCl. J Biol Chem 262:14255–14259
- Beetsch JW, Olson JE (1996) Hyperosmotic exposure alters total taurine quantity and cellular transport in rat astrocyte cultures. Biochim Biophys Acta 1290:141–148
- Behrens R, Nolting A, Reimann F, Schwarz M, Waldschütz R, Pongs O (2002) hKCNMB3 and hKCNMB4, cloning and characterization of two members of the large-conductance calcium-activated potassium β subunit family. FEBS Lett 474:99–106
- Bender AS, Norenberg MD (1994) Calcium dependence of hypo-osmotically induced potassium release in cultured astrocytes. J Neurosci 14:4237–4243
- Bender AS, Mantelle LL, Norenberg MD (1994) Stimulation of calcium uptake in cultured astrocytes by hypo-osmotic stress-effect of cyclic AMP. Brain Res 645:27–35
- Bender AS, Neary JT, Norenberg MD (1993) Role of phosphoinositide hydrolysis in astrocyte volume regulation. J Neurochem 61:1506–1514
- Benos DJ, Fuller CM, Shlyonsky V Gh, Berdiev BK, Ismailov II (1997) Amiloride-sensitive Na⁺ channels: Insights and outlooks. News Physiol Sci 12:55–61
- Bertrand B, Wakabayashi S, Ikeda T, Pouysségur J, Shigekawa M (1994) The Na⁺/H⁺ exchanger isoform 1 (NHE1) is a novel member of the calmodulin-binding proteins. J Biol Chem 269:13703–13709
- Bianchini L, L'Allemain G, Pouyssegur J (1997) The p42/p44 mitogen-activated protein kinase cascade is determinant in mediating activation of the Na⁺/H⁺ exchanger (NHE1 isoform) in response to growth factors. J Biol Chem 272:271–279
- Blanco G, Mercer RW (1998) Isozymes of the Na-K-ATPase: heterogeneity in structure, diversity in function. Am J Physiol 275:F633–F650
- Bleich M, Riedemann N, Warth R, Kerstan D, Leipziger J, Hör M, van Driessche W, Greger R (1996) Ca²⁺ regulated K⁺ and nonselective cation channels in the basolateral membrane of rat colonic crypt base cells. Pflugers Arch 432:1011–1022
- Boese SH, Glanville M, Gray MA, Simmons NL (2000) The swelling-activated anion conductance in the mouse renal inner medullary collecting duct cell line mIMCD-K2. J Membr Biol 177:51–64
- Boese SH, Kinne RKH, Wehner F (1996a) Single-channel properties of swelling-activated anion conductance in rat inner medullary collecting duct cells. Am J Physiol 271:F1224–F1233

- Boese SH, Wehner F, Kinne RKH (1996b) Taurine permeation through swelling-activated anion conductance in rat IMCD cells in primary culture. Am J Physiol 271:F498–F507
- Böhmer C, Wagner CA, Beck S, Moschen I, Melzig J, Werner A, Lin J-T, Lang F, Wehner F (2000) The shrinkage-activated Na⁺ conductance of rat hepatocytes and its possible correlation to rENaC. Cell Physiol Biochem 10:187–194
- Böhmer C, Wehner F (2001) The epithelial Na⁺ channel (ENaC) is related to the hypertonicity-induced Na⁺ conductance in rat hepatocytes. FEBS Lett 494:125–128
- Bookstein C, Musch MW, DePaoli A, Xie Y, Villereal M, Rao MC, Chang EB (1994) A unique sodiumhydrogen exchange isoform (NHE-4) of the inner medulla of the rat kidney is induced by hyperosmolarity. J Biol Chem 269:29704-29709
- Bookstein C, Xie Y, Rabenau K, Musch MW, McSwine RL, Rao MC, Chang EB (1997) Tissue distribution of Na⁺/H⁺ exchanger isoforms NHE2 and NHE4 in rat intestine and kidney. Am J Physiol 273:C1496–C1505
- Boron WF (2001) Sodium-coupled bicarbonate transporters. JOP 2:176-181
- Bowen JW (1992) Regulation of Na⁺-K⁺-ATPase expression in cultured renal cells by incubation in hypertonic medium. Am J Physiol 262:C845–C853
- Brandsch M, Miyamoto Y, Ganapathy V, Leibach FH (1993) Regulation of taurine transport in human colon carcinoma cell lines (HT- 29 and Caco-2) by protein kinase C. Am J Physiol 264:G939–G946
- Brierley GP, Davis MH, Cragoe EJ Jr., Jung DW (1989) Kinetic properties of the Na⁺/H⁺ antiport of heart mitochondria. Biochemistry (Mosc) 28:4347–4354
- Brosius FC III, Alper SL, Garcia AM, Lodish HF (1989) The major kidney band 3 gene transcript predicts an amino-terminal truncated band 3 polypeptide. J Biol Chem 264:7784–7787
- Burg MB (1995) Molecular basis of osmotic regulation. Am J Physiol 268:F983-F996
- Burg MB (1996) Coordinate regulation of organic osmolytes in renal cells. Kidney Int 49:1684-1685
- Burg MB, Kador PF (1988) Sorbitol, osmoregulation, and the complications of diabetes. J Clin Invest 81:635-640
- Burg MB, Kwon ED, Kültz D (1997) Regulation of gene expression by hypertonicity. Annu Rev Physiol 59:437–455
- Burger-Kentischer A, Müller E, Neuhofer W, März J, Thurau K, Beck F (1999) Expression of aldose reductase, sorbitol dehydrogenase and Na⁺/myo- inositol and Na⁺/Cl⁻-/betaine transporter mRNAs in individual cells of the kidney during changes in the diuretic state. Pflugers Arch 437:248–254
- Burridge K, Chrzanowska-Wodnicka M (1996) Focal adhesions, contractility, and signaling. Annu Rev Cell Dev Biol 12:463–518
- Burridge K, Zhong C (1997) Focal adhesion assembly. Trends Cell Biol 7:342–347
- Busch AE, Maylie J (1993) MinK channels: A minimal channel protein with a maximal impact. Cell Physiol Biochem 3:270–276
- Busch AE, Suessbrich H (1997) Role of the ISK protein in the IminK channel complex. Trends Pharmacol Sci 18:26–29
- Busch AE, Varnum M, Adelman JP, North RA (1992) Hypotonic solution increases the slowly activating potassium current ISK expressed in *Xenopus* oocytes. Biochem Biophys Res Commun 184:804–810
- Buyse G, Voets T, Tytgat J, De Greef C, Droogmans G, Nilius B, Eggermont J (1997) Expression of human pI_{Cln} and ClC-6 in Xenopus oocytes induces an identical endogenous chloride conductance. J Biol Chem 272:3615–3621
- Cammarata PR, Schafer G, Chen SW, Guo Z, Reeves RE (2002) Osmoregulatory alterations in taurine uptake by cultured human and bovine lens epithelial cells. Invest Ophthalmol Vis Sci 43:425–433
- Capasso JM, Rivard CJ, Berl T (2001) Long-term adaptation of renal cells to hypertonicity: role of MAP kinases and Na-K-ATPase. Am J Physiol Renal Physiol 280, F768-F776
- Caruso-Neves C, Lopes AG (2000) Sodium pumps in the Malpighian tubule of Rhodnius sp. An Acad Bras Cienc 72:407–412
- Chan HC, Fu WO, Chung YW, Huang SJ, Chan PSF, Wong PYD (1994) Swelling-induced anion and cation conductances in human epididymal cells. J Physiol 478:449–460
- Chan HC, Nelson DJ (1992) Chloride-dependent cation conductance activated during cellular shrinkage. Science 257:669–671
- Chen JG, Coe M, McAteer JA, Kempson SA (1996a) Hypertonic activation and recovery of system A amino acid transport in renal MDCK cells. Am J Physiol 270:F419–F424
- Chen Y, Simasko SM, Niggel J, Sigurdson WJ, Sachs F (1996b) Ca²⁺ uptake in GH3 cells during hypotonic swelling: the sensory role of stretch-activated ion channels. Am J Physiol 270:C1790–C1798
- Chen L, Wang L, Jacob TJ (1999a) Association of intrinsic pICln with volume-activated Cl⁻ current and volume regulation in a native epithelial cell. Am J Physiol 276:C182–C192

- Chen SY, Bhargava A, Mastroberardino L, Meijer OC, Wang J, Buse P, Firestone GL, Verrey F, Pearce D (1999b) Epithelial sodium channel regulated by aldosterone-induced protein sgk. Proc Natl Acad Sci USA 96:2514–2519
- Chesney RW, Gusowski N, Dabbagh S (1985a) Renal cortex taurine content regulates renal adaptive response to altered dietary intake of sulfur amino acids. J Clin Invest 76:2213–2221
- Chesney RW, Gusowski N, Dabbagh S, Theissen M, Padilla M, Diehl A (1985b) Factors affecting the transport of β -amino acids in rat renal brush-border membrane vesicles. The role of external chloride. Biochim Biophys Acta 812:702–712
- Chong LD, Traynor-Kaplan A, Bokoch GM, Schwartz MA (1994) The small GTP-binding protein Rho regulates a phosphatidylinositol 4-phosphate 5-kinase in mammalian cells. Cell 79:507–513
- Chou CY, Shen MR, Hsu KS, Huang HY, Lin HC (1998) Involvement of PKC-α in regulatory volume decrease responses and activation of volume-sensitive chloride channels in human cervical cancer HT-3 cells. J Physiol 512:435–448
- Chow CW, Woodside M, Demaurex N, Yu FH, Plant P, Rotin D, Grinstein S, Orlowski J (1999) Prolinerich motifs of the Na⁺/H⁺ exchanger 2 isoform. Binding of Src homology domain 3 and role in apical targeting in epithelia. J Biol Chem 274:10481–10488
- Christensen O (1987) Mediation of cell volume regulation by Ca²⁺ influx through stretch-activated channels. Nature 330:66–68
- Christensen O, Hoffmann EK (1992) Cell swelling activates K⁺ and Cl⁻ channels as well as nonselective, stretch-activated cation channels in Ehrlich ascites tumor cells. J Membr Biol 129:13–36
- Christensen O, Simon M, Randlev T (1989) Anion channels in a leaky epithelium. A patch-clamp study of choroid plexus. Pflugers Arch 415:37–46
- Civan MM, Coca-Prados M, Peterson-Yantorno K (1994a) Pathways signaling the regulatory volume decrease of cultured nonpigmented ciliary epithelial cells. Invest Ophthalmol Vis Sci 35:2876–2886
- Civan MM, Marano CW, Matschinsky FW, Peterson-Yantorno K (1994b) Prolonged incubation with elevated glucose inhibits the regulatory response to shrinkage of cultured human retinal pigment epithelial cells. J Membr Biol 139:1–13
- Civan MM, Coca-Prados M, Peterson-Yantorno K (1996) Regulatory volume increase of human nonpigmented ciliary epithelial cells. Exp Eye Res 62:627–639
- Civan MM, Peterson-Yantorno K, Sánchez-Torres J, Coca-Prados M (1997) Potential contribution of epithelial Na⁺ channel to net secretion of aqueous humor. J Exp Zool 279:498–503
- Clark EA, King WG, Brugge JS, Symons M, Hynes RO (1998) Integrin-mediated signals regulated by members of the rho family of GTPases. J Cell Biol 142:573–586
- Clemo HF, Baumgarten CM (1997) Swelling-activated Gd³⁺-sensitive cation current and cell volume regulation in rabbit ventricular myocytes. J Gen Physiol 110:297–312
- Coca-Prados M, Sanchez-Torres J, Peterson-Yantorno K, Civan MM (1996) Association of ClC-3 channel with Cl⁻ transport by human nonpigmented ciliary epithelial cells. J Membr Biol 150:197–208
- Cockcroft S (2001) Signalling roles of mammalian phospholipase D1 and D2. Cell Mol Life Sci 58:1674–1687
- Cornet M, Ubl J, Kolb H-A (1993) Cytoskeleton and ion movements during volume regulation in cultured PC12 cells. J Membr Biol 133:161–170
- Costa PMF, Fernandes PL, Ferreira HG, Ferreira KTG, Giraldez F (1987) Effects of cell volume changes on membrane ionic permeabilities and sodium transport in frog skin (*Rana ridibunda*). J Physiol (Lond) 393:1–17
- Coulombe A, Coraboeuf E (1992) Large-conductance chloride channels of new-born rat cardiac myocytes are activated by hypotonic media. Pflugers Arch 422:143–150
- Counillon L, Pouyssegur J (2000) The expanding family of eukaryotic Na⁺/H⁺ exchangers. J Biol Chem 275:1-4
- Coupaye-Gerard B, Bookstein C, Duncan P, Chen XY, Smith PR, Musch M, Ernst SA, Chang EB, Kleyman TR (1996) Biosynthesis and cell surface delivery of the NHE1 isoform of Na⁺/H⁺ exchanger in A6 cells. Am J Physiol 271:C1639–C1645
- Craelius W, Ross MJ, Harris DR, Chen VK, Palant CE (1993) Membrane currents controlled by physical forces in cultured mesangial cells. Kidney Int 43:535–543
- Cunningham SA, Awayda MS, Bubien JK, Ismailov II, Arrate MP, Berdiev BK, Benos DJ, Fuller CM (1995) Cloning of an epithelial chloride channel from bovine trachea. J Biol Chem 270:31016–31026
- Cuppoletti J, Tewari KP, Sherry AM, Kupert EY, Malinowska DH (2001) ClC-2 Cl- channels in human lung epithelia: activation by arachidonic acid, amidation, and acid-activated omeprazole. Am J Physiol Cell Physiol 281:C46–C54
- Curtis DR, Johnston GA (1974) Amino acid transmitters in the mammalian central nervous system. Ergeb Physiol 69:97–188

- Czekay RP, Kinne-Saffran E, Kinne RKH (1994) Membrane traffic and sorbitol release during osmo- and volume regulation in isolated rat renal inner medullary collecting duct cells. Eur J Cell Biol 63:20–31
- de Angelis E, Petronini PG, Borghetti P, Borghetti AF, Wheeler KP (1999) Induction of betaine-gammaaminobutyric acid transport activity in porcine chondrocytes exposed to hypertonicity. J Physiol 518 (Pt 1), 187–194
- Dermietzel R, Hwang TK, Buettner R, Hofer A, Dotzler E, Kremer M, Deutzmann R, Thinnes FP, Fishman GI, Spray DC, (1994) Cloning and in situ localization of a brain-derived porin that constitutes a large-conductance anion channel in astrocytic plasma membranes. Proc Natl Acad Sci USA 91:499–503
- Deutsch C, Chen L-Q (1993) Heterologous expression of specific K⁺ channels in T lymphocytes: Functional consequences for volume regulation. Proc Natl Acad Sci USA 90:10036–10040
- Devor DC, Frizzell RA (1998) Modulation of K⁺ channels by arachidonic acid in T84 cells. II. Activation of a Ca²⁺-independent K⁺ channel. Am J Physiol 274:C149–C160
- Dezaki K, Tsumura T, Maeno E, Okada Y (2000) Receptor-mediated facilitation of cell volume regulation by swelling-induced ATP release in human epithelial cells. Jpn J Physiol 50:235–241
- Di Fulvio M, Lauf PK, Adragna NC (2001a) Nitric oxide signaling pathway regulates potassium chloride cotransporter-1 mRNA expression in vascular smooth muscle cells. J Biol Chem 276:44534–44540
- Di Fulvio M, Lincoln TM, Lauf PK, Adragna NC (2001b) Protein kinase G regulates potassium chloride cotransporter-3 expression in primary cultures of rat vascular smooth muscle cells. J Biol Chem 276:21046–21052
- Donaldson PJ, Chen LK, Lewis SA (1989) Effects of serosal anion composition on the permeability properties of rabbit urinary bladder. Am J Physiol 256:F1125–F1134
- Doroshenko P, Neher E (1992) Volume-sensitive chloride conductance in bovine chromaffin cell membrane. J Physiol (Lond) 449:197–218
- Duan D, Winter C, Cowley S, Hume JR, Horowitz B (1997) Molecular identification of a volume-regulated chloride channel. Nature 390:417–421
- Duan D, Ye L, Britton F, Horowitz B, Hume JR (2000) A novel anionic inward rectifier in native cardiac myocytes. Circ Res 86:E63–E71
- Dubé L, Parent L, Sauvé R (1990) Hypotonic shock activates a maxi K⁺ channel in primary cultured proximal tubule cells. Am J Physiol 259:F348–F356
- Dubinsky WP, Mayorga-Wark O, Schultz SG (2000) Potassium channels in basolateral membrane vesicles from necturus enterocytes: stretch and ATP sensitivity. Am J Physiol Cell Physiol 279:C634–C638
- Dudeja PK, Rao DD, Syed I, Joshi V, Dahdal RY, Gardner C, Risk MC, Schmidt L, Bavishi D, Kim KE, Harig JM, Goldstein JL, Layden TJ, Ramaswamy K (1996) Intestinal distribution of human Na⁺/H⁺ exchanger isoforms NHE-1, NHE- 2, and NHE-3 mRNA. Am J Physiol 271:G483–G493
- Dunbar LA, Caplan MJ (2000) The cell biology of ion pumps: sorting and regulation. Eur J Cell Biol 79:557–563
- Dutzler R, Campbell EB, Cadene M, Chait BT, MacKinnon R (2002) X-ray structure of a CIC chloride channel at 3.0 A reveals the molecular basis of anion selectivity. Nature 415:287–294
- Duzgun SA, Rasque H, Kito H, Azuma N, Li W, Basson MD, Gahtan V, Dudrick SJ, Sumpio BE (2000) Mitogen-activated protein phosphorylation in endothelial cells exposed to hyperosmolar conditions. J Cell Biochem 76:567–571
- Ellis RJ (2001) Macromolecular crowding: obvious but underappreciated. Trends Biochem Sci 26:597–604 Emma F, McManus M, Strange K (1997) Intracellular electrolytes regulate the volume set point of the organic osmolyte/anion channel VSOAC. Am J Physiol 272:C1766–C1775
- Erickson GR, Alexopoulos LG, Guilak F (2001) Hyperosmotic stress induces volume change and calcium transients in chondrocytes by transmembrane, phospholipid, and G-protein pathways. J Biomech 34:1527–1535
- Estevez AY, O'Regan MH, Song D, Phillis JW (1999) Hypo-osmotically induced amino acid release from the rat cerebral cortex: role of phospholipases and protein kinases. Brain Res 844:1–9
- Falke LC, Misler S (1989) Activity of ion channels during volume regulation by clonal N1E115 neuroblastoma cells. Proc Natl Acad Sci USA 86:3919–3923
- Farrugia G, Rae J (1993) Effect of volume changes on a potassium current in rabbit corneal epithelial cells. Am J Physiol 264:C1238–C1245
- Felipe A, Snyders DJ, Deal KK, Tamkun MM (1993) Influence of cloned voltage-gated K⁺ channel expression on alanine transport, Rb⁺ uptake, and cell volume. Am J Physiol 265:C1230–C1238
- Feranchak AP, Berl T, Capasso J, Wojtaszek PA, Han J, Fitz JG (2001) p38 MAP kinase modulates liver cell volume through inhibition of membrane Na⁺ permeability. J Clin Invest 108:1495–1504
- Feranchak AP, Fitz JG, Roman RM (2000) Volume-sensitive purinergic signaling in human hepatocytes. J Hepatol 33:174–182

- Feranchak AP, Roman RM, Doctor RB, Salter KD, Toker A, Fitz JG (1999) The lipid products of phosphoinositide 3-kinase contribute to regulation of cholangiocyte ATP and chloride transport. J Biol Chem 274:30979–30986
- Feranchak AP, Roman RM, Schwiebert EM, Fitz JG (1998) Phosphatidylinositol 3-kinase contributes to cell volume regulation through effects on ATP release. J Biol Chem 273:14906–14911
- Ferraris JD, Burg MB, Williams CK, Peters EM, Garcia-Perez A (1996) Betaine transporter cDNA cloning and effect of osmolytes on its mRNA induction. Am J Physiol 270:C650–C654
- Fievet B, Gabillat N, Borgese F, Motais R (1995) Expression of band 3 anion exchanger induces chloride current and taurine transport: structure-function analysis. EMBO J 14:5158–5169
- Fievet B, Perset F, Gabillat N, Guizouarn H, Borgese F, Ripoche P, Motais R (1998) Transport of uncharged organic solutes in Xenopus oocytes expressing red cell anion exchangers (AE1 s) Proc Natl Acad Sci USA 95:10996–11001
- Filipovic D, Sackin H (1991) A calcium-permeable stretch-activated cation channel in renal proximal tubule. Am J Physiol 260:F119–F129
- Filipovic D, Sackin H (1992) Stretch- and volume-activated channels in isolated proximal tubule cells. Am J Physiol 262:F857–F870
- Forlani G, Bossi E, Perego C, Giovannardi S, Peres A (2001) Three kinds of currents in the canine betaine-GABA transporter BGT-1 expressed in *Xenopus laevis* oocytes. Biochim Biophys Acta 1538:172–180
- Fraser CL, Swanson RA (1994) Female sex hormones inhibit volume regulation in rat brain astrocyte culture. Am J Physiol 267:C909–C914
- Fuller CM, Benos DJ (2000) Ca²⁺-activated Cl⁻ channels: a newly emerging anion transport family. News Physiol Sci 15:165–171
- Fuller CM, Ji HL, Tousson AM, Elble RC, Pauli BU, Benos DJ (2001) Ca²⁺-activated Cl⁻ channels: a newly emerging anion transport family. Pflugers Arch 443 Suppl 1, S107–S110
- Furlong TJ, Moriyama T, Spring KR (1991) Activation of osmolyte efflux from cultured renal papillary epithelial cells. J Membr Biol 123:269–277
- Fürst J, Bazzini C, Jakab M, Meyer G, König M, Gschwentner M, Ritter M, Schmarda A, Bottà G, Benz R, Deetjen P, Paulmichl M (2000a) Functional reconstitution of ICln in lipid bilayers. Pflugers Arch 440:100–115
- Fürst J, Jakab M, Konig M, Ritter M, Gschwentner M, Rudzki J, Danzl J, Mayer M, Burtscher CM, Schirmer J, Maier B, Nairz M, Chwatal S, Paulmichl M (2000b) Structure and function of the ion channel ICln. Cell Physiol Biochem 10:329–334
- Fyfe GK, Quinn A, Canessa CM (1998) Structure and function of the Mec-ENaC family of ion channels. Semin Nephrol 18:138–151
- Gabbay KH (1973) The sorbitol pathway and the complications of diabetes. N Engl J Med 288:831–836
- Gamper N, Huber SM, Badawi K, Lang F (2000) Cell volume-sensitive sodium channels upregulated by glucocorticoids in U937 macrophages. Pflugers Arch 441:281–286
- Garavaglia L, Rodighiero S, Bertocchi C, Manfredi R, Fürst J, Gschwentner M, Ritter M, Bazzini C, Botta G, Jakab M, Meyer G, Paulmichl M (2002) ICln channels reconstituted in heart-lipid bilayer are selective to chloride. Pflugers Arch 443:748–753
- Garcia-Calvo M, Knaus HG, McManus OB, Giangiacomo KM, Kaczorowski GJ, Garcia ML (1994) Purification and reconstitution of the high-conductance, calcium-activated potassium channel from tracheal smooth muscle. J Biol Chem 269:676–682
- Garcia-Perez A, Burg MB (1991) Renal medullary organic osmolytes. Physiol Rev 71:1081-1115
- Gardos G (2002) The function of calcium in the potassium permeability of human erythrocytes. Biochim Biophys Acta 30:653–654
- Garty H, Furlong TJ, Ellis DE, Spring KR (1991) Sorbitol permease: an apical membrane transporter in cultured renal papillary epithelial cells. Am J Physiol 260:F650–F656
- Garty H, Palmer LG (1997) Epithelial sodium channels: Function, structure, and regulation. Physiol Rev 77:359–396
- Gill DR, Hyde SC, Higgins CF, Valverde MA, Mintenig GM, Sepúlveda FV (1992) Separation of drug transport and chloride channel functions of the human multidrug resistance P-glycoprotein. Cell 71:23–32
- Gillen CM, Brill S, Payne JA, Forbush B, III (1996) Molecular cloning and functional expression of the K-Cl cotransporter from rabbit, rat, and human—A new member of the cation-chloride cotransporter family. J Biol Chem 271:16237–16244
- Goetzl EJ, An S (1998) Diversity of cellular receptors and functions for the lysophospholipid growth factors lysophosphatidic acid and sphingosine 1-phosphate. FASEB J 12:1589–1598
- Goldstein L, Davis EM (1994) Taurine, betaine, and inositol share a volume-sensitive transporter in skate erythrocyte cell membrane. Am J Physiol 267:R426–R431

- Goldstein L, Davis-Amaral EM, Musch MW (1996) Organic osmolyte channels: transport characteristics and regulation. Kidney Int 49:1690–1694
- Good DW, Di Mari JF, Watts BA, III (2000) Hypo-osmolality stimulates Na⁺/H⁺ exchange and HCO₃⁻ absorption in thick ascending limb via PI 3-kinase. Am J Physiol Cell Physiol 279:C1443–C1454
- Gosling M, Poyner DR, Smith JW (1996) Effects of arachidonic acid upon the volume-sensitive chloride current in rat osteoblast-like (ROS 17/2.8) cells. J Physiol 493:613–623
- Graf J, Haddad P, Häussinger D, Lang F (1988) Cell volume regulation in liver. Renal Physiol Biochem 11:202–220
- Graf J, Häussinger D (1996) Ion transport in hepatocytes: Mechanisms and correlations to cell volume, hormone actions and metabolism. J Hepatol 24 Suppl. 1:53–77
- Greger R (1996) The membrane transporters regulating epithelial NaCl secretion. Pflugers Arch 432:579–588
- Grinstein S, Rothstein A, Cohen S (1985) Mechanism of osmotic activation of Na⁺/H⁺ exchange in rat thymic lymphocytes. J Gen Physiol 85:765–787
- Grinstein S, Woodside M, Sardet C, Pouyssegur J, Rotin D (1992) Activation of the Na⁺/H⁺ antiporter during cell volume regulation. Evidence for a phosphorylation-independent mechanism. J Biol Chem 267:23823–23828
- Grinstein S, Woodside M, Waddell TK, Downey GP, Orlowski J, Pouyssegur J, Wong DC, Foskett JK (1993) Focal localization of the NHE-1 isoform of the Na⁺/H⁺ antiport: assessment of effects on intracellular pH. EMBO J 12:5209–5218
- Gründer S, Thiemann A, Pusch M, Jentsch TJ (1992) Regions involved in the opening of CIC-2 chloride channel by voltage and cell volume. Nature 360:759–762
- Grunewald RW, Kinne RKH (1989) Intracellular sorbitol content in isolated rat inner medullary collecting duct cells. Regulation by extracellular osmolarity. Pflugers Arch 414:178–184
- Grunewald RW, Kinne RKH (1999) Osmoregulation in the mammalian kidney: the role of organic osmolytes. J Exp Zool 283:708–724
- Grunewald JM, Grunewald RW, Kinne RKH (1993a) Ion content and cell volume in isolated collecting duct cells: effect of hypotonicity. Kidney Int 44:509–517
- Grunewald RW, Weber II, Kinne-Saffran E, Kinne RKH (1993b) Control of sorbitol metabolism in renal inner medulla of diabetic rats: regulation by substrate, cosubstrate and products of the aldose reductase reaction. Biochim Biophys Acta 1225:39–47
- Grunewald JM, Grunewald RW, Kinne RKH (1994) Regulation of ion content and cell volume in isolated rat renal IMCD cells under hypertonic conditions. Am J Physiol 267:F13–F19
- Grunewald RW, Weber II, Kinne RKH (1995) Renal inner medullary sorbitol metabolism. Am J Physiol 269:F696–F701
- Grunewald RW, Wagner M, Schubert I, Franz HE, Muller GA, Steffgen J (1998) Rat renal expression of mRNA coding for aldose reductase and sorbitol dehydrogenase and its osmotic regulation in inner medullary collecting duct cells. Cell Physiol Biochem 8:293–303
- Gschwentner M, Nagl UO, Woll E, Schmarda A, Ritter M, Paulmichl M (1995) Antisense oligonucleotides suppress cell-volume-induced activation of chloride channels. Pflugers Arch 430:464–470
- Guastella J, Nelson N, Nelson H, Czyzyk L, Keynan S, Miedel MC, Davidson N, Lester HA, Kanner BI (1990) Cloning and expression of a rat brain GABA transporter. Science 249:1303–1306
- Guizouarn H, Gabillat N, Motais R, Borgese F (2001) Multiple transport functions of a red blood cell anion exchanger, tAE1: its role in cell volume regulation. J Physiol (Lond) 535:497–506
- Gusev GP, Sherstobitov AO (1996) An amiloride-sensitive, volume-dependent Na⁺ transport across the Lamprey (*Lampetra fluviatilis*) erythrocyte membrane. Gen Physiol Biophys 15:129–143
- Haas M, Forbush B, III (2000) The Na-K-Cl cotransporter of secretory epithelia. Annu Rev Physiol 62:515-534
- Hafting T, Sand O (2000) Purinergic activation of BK channels in clonal kidney cells (Vero cells) Acta Physiol Scand 170:99–109
- Hager K, Hazama A, Kwon HM, Loo DD, Handler JS, Wright EM (1995) Kinetics and specificity of the renal Na⁺/myo-inositol cotransporter expressed in *Xenopus* oocytes. J Membr Biol 143:103–113
- Hall AC, Bush PG (2001) The role of a swelling-activated taurine transport pathway in the regulation of articular chondrocyte volume. Pflugers Arch 442:771–781
- Hall JA, Kirk J, Potts JR, Rae C, Kirk K (1996) Anion channel blockers inhibit swelling-activated anion, cation, and nonelectrolyte transport in HeLa cells. Am J Physiol 271:C579–C588
- Hall SK, Zhang J, Lieberman M (1995) Cyclic AMP prevents activation of a swelling-induced chloridesensitive conductance in chick heart cells. J Physiol 488:359–369
- Hallows KR, Packman CH, Knauf PA (1991) Acute cell volume changes in anisotonic media affect F-actin content of HL-60 cells. Am J Physiol 261:C1154–C1161

- Hammerman MR, Sacktor B, Daughaday WH (1980) myo-Inositol transport in renal brush border vesicles and it inhibition by D-glucose. Am J Physiol 239:F113–F120
- Han X, Budreau AM, Chesney RW (1998) Molecular cloning and functional expression of an LLC-PK1 cell taurine transporter that is adaptively regulated by taurine. Adv Exp Med Biol 442:261–268
- Handler JS, Kwon HM (2001) Cell and molecular biology of organic osmolyte accumulation in hypertonic renal cells. Nephron 87:106–110
- Hardy SP, Goodfellow HR, Valverde MA, Gill DR, Sepulveda V, Higgins CF (1995) Protein kinase C-mediated phosphorylation of the human multidrug resistance P-glycoprotein regulates cell volume-activated chloride channels. EMBO J 14:68–75
- Häussinger D (1996) The role of cellular hydration in the regulation of cell function. Biochem J 313:697–710
- Häussinger D (1998) Osmoregulation of liver cell function: signalling, osmolytes and cell heterogeneity. Contrib Nephrol 123:185–204
- Häussinger D, Hallbrucker C, Vom Dahl S, Decker S, Schweizer U, Lang F, Gerok W (1991) Cell volume is a major determinant of proteolysis control in liver. FEBS Lett 283:70–72
- Hazama A, Okada Y (1988) Ca²⁺ sensitivity of volume-regulatory K⁺ and Cl⁻ channels in cultured human epithelial cells. J Physiol 402:687–702
- Hazama A, Okada Y (1990) Involvement of Ca²⁺-induced Ca²⁺ release in the volume regulation of human epithelial cells exposed to a hypotonic medium. Biochem Biophys Res Commun 167:287–293
- Hebert SC, Sun A (1988) Hypotonic cell volume regulation in mouse medullary thick ascending limb: effects of ADH. Am J Physiol 255:F962–F969
- Heinzinger H, van den Boom F, Tinel H, Wehner F (2001) In rat hepatocytes, the hypertonic activation of Na⁺ conductance and Na⁺-K⁺-2Cl⁻ symport—but not Na⁺-H⁺ antiport—is mediated by protein kinase C. J Physiol (Lond) 536:703–715
- Henson JH (1999) Relationships between the actin cytoskeleton and cell volume regulation. Microsc Res Tech 47:155–162
- Hiki K, D'Andrea RJ, Furze J, Crawford J, Woollatt E, Sutherland GR, Vadas MA, Gamble JR (1999) Cloning, characterization, and chromosomal location of a novel human K+-Cl⁻ cotransporter. J Biol Chem 274:10661–10667
- Hirsch JR, Schlatter E (1997) Ca²⁺-dependent K⁺ channels in the cortical collecting duct of rat. Wien Klin Wochenschr 109:485–488
- Hoffmann EK (1978) Regulation of cell volume by selective changes in the leak permeabilities of Ehrlich ascites tumor cells. In Jorgensen CB, Skadhauge E (eds) Osmotic and volume regulation. Alfred Benzon Symposium XI., Munksgaard, Copenhagen, pp. 397–417
- Hoffmann EK, Dunham PB (1995) Membrane mechanisms and intracellular signalling in cell volume regulation. Int Rev Cytol 161:173–262
- Hoffmann EK, Lambert IH (1994) On the similarity between the small Cl⁻ channel and the taurine channel activated after cell swelling in Ehrlich ascites tumor cells. Jpn J Physiol 44 Suppl 2:S49–S53
- Hoffmann EK, Lambert IH, Simonsen LO (1988) Mechanisms in volume regulation in Ehrlich ascites tumor cells. Renal Physiol Biochem 11:221–247
- Hoffmann EK, Mills JW (1999) Membrane events involved in volume regulation. Curr Top Membr 48:123-196
- Hoffmann EK, Pedersen SF (1998) Sensors and signal transduction in the activation of cell volume regulatory ion transport systems. Contrib Nephrol 123:50–78
- Hoffmann EK, Simonsen LO, Lambert IH (1984) Volume-induced increase of K⁺ and Cl[−] permeabilities in Ehrlich ascites tumor cells. Role of internal Ca²⁺. J Membr Biol 78:211–222
- Holtzman EJ, Kumar S, Faaland CA, Warner F, Logue PJ, Erickson SJ, Ricken G, Waldman J, Kumar S, Dunham PB (1998) Cloning, characterization, and gene organization of K-Cl cotransporter from pig and human kidney and C. elegans. Am J Physiol 275:F550–F564
- Hoogerwerf WA, Tsao SC, Devuyst O, Levine SA, Yun CH, Yip JW, Cohen ME, Wilson PD, Lazenby AJ, Tse CM, Donowitz M (1996) NHE2 and NHE3 are human and rabbit intestinal brush-border proteins. Am J Physiol 270:G29–G41
- Hooley R, Yu CY, Symons M, Barber DL (1996) G α 13 stimulates Na⁺-H⁺ exchange through distinct Cdc42-dependent and RhoA-dependent pathways. J Biol Chem 271:6152–6158
- Horio M, Yamauchi A, Moriyama T, Imai E, Orita Y (1997) Osmotic regulation of amino acids and system A transport in Madin-Darby canine kidney cells. Am J Physiol 272:C804–C809
- Huang R, Somjen GG (1997) Effects of hypertonia on voltage-gated ion currents in freshly isolated hippocampal neurons, and on synaptic currents in neurons in hippocampal slices. Brain Res 748:157–167
- Huber SM, Gamper N, Lang F (2001) Chloride conductance and volume-regulatory nonselective cation conductance in human red blood cell ghosts. Pflugers Arch 441:551–558

- Hubert MD, Levitan I, Hoffman MM, Zraggen M, Hofreiter ME, Garber SS (2000) Modulation of volume regulated anion current by I(Cln) Biochim Biophys Acta 1466:105–114
- Hume JR, Duan D, Collier ML, Yamazaki J, Horowitz B (2000) Anion transport in heart. Physiol Rev 80:31–81
- Hunter M (1990) Stretch-activated channels in the basolateral membrane of single proximal cells of frog kidney. Pflugers Arch 416:448–453
- Hurst AM, Hunter M (1990) Stretch-activated channels in single early distal tubule cells of the frog. J Physiol (Lond) 430:13–24
- Ingber DE (1997) Tensegrity: the architectural basis of cellular mechanotransduction. Annu Rev Physiol 59:575–599
- Ishii TM, Silvia C, Hirschberg B, Bond CT, Adelman JP, Maylie J (1997) A human intermediate conductance calcium-activated potassium channel. Proc Natl Acad Sci USA 94:11651–11656
- Isom LL, De Jongh KS, Catterall WA (1994) Auxiliary subunits of voltage-gated ion channels. Neuron 12:1183–1194
- Itoh T, Yamauchi A, Miyai A, Yokoyama K, Kamada T, Ueda N, Fujiwara Y (1994) Mitogen-activated protein kinase and its activator are regulated by hypertonic stress in Madin-Darby canine kidney cells. J Clin Invest 93:2387–2392
- Iwasa K, Tasaki I, Gibbons RC (1980) Swelling of nerve fibers associated with action potentials. Science 210:338–339
- Jackson PS, Churchwell K, Ballatori N, Boyer JL, Strange K (1996) Swelling-activated anion conductance in skate hepatocytes: Regulation by cell Cl⁻ and ATP. Am J Physiol 270:C57–C66
- Jackson PS, Morrison R, Strange K (1994) The volume-sensitive organic osmolyte-anion channel VSOAC is regulated by nonhydrolytic ATP binding. Am J Physiol 267:C1203–C1209
- Jackson PS, Strange K (1993) Volume-sensitive anion channels mediate swelling-activated inositol and taurine efflux. Am J Physiol 265:C1489–C1500
- Jackson PS, Strange K (1995a) Characterization of the voltage-dependent properties of a volume-sensitive anion conductance. J Gen Physiol 105:661–677
- Jackson PS, Strange K (1995b) Single-channel properties of a volume-sensitive anion conductance. Current activation occurs by abrupt switching of closed channels to an open state. J Gen Physiol 105:643-660
- Jacobsen JG, Smith LH (1968) Biochemistry and physiology of taurine and taurine derivatives. Physiol Rev 48:424–511
- Jalonen T (1993) Single-channel characteristics of the large-conductance anion channel in rat cortical astrocytes in primary culture. Glia 9:227–237
- Jarolimek W, Lewen A, Misgeld U (1999) A furosemide-sensitive K+-Cl⁻ cotransporter counteracts intracellular Cl⁻ accumulation and depletion in cultured rat midbrain neurons. J Neurosci 19:4695–4704
- Jensen BS, Jessen F, Hoffmann EK (1993) Na⁺, K⁺, Cl⁻ cotransport and its regulation in Ehrlich ascites tumor cells. Ca²⁺/calmodulin and protein kinase C dependent pathways. J Membr Biol 131:161–178
- Jensen BS, Strobaek D, Christophersen P, Jorgensen TD, Hansen C, Silahtaroglu A, Olesen SP, Ahring PK (1998) Characterization of the cloned human intermediate-conductance Ca²⁺-activated K⁺ channel. Am J Physiol 275:C848–C856
- Jensen BS, Strobaek D, Olesen SP, Christophersen P (2001) The Ca²⁺-activated K⁺ channel of intermediate conductance: a molecular target for novel treatments? Curr Drug Targets 2:401–422
- Jentsch TJ (1996) Chloride channels: A molecular perspective. Curr Opin Neurobiol 6:303-310
- Jentsch TJ, Friedrich T, Schriever A, Yamada H (1999) The CLC chloride channel family. Pflugers Arch 437:783–795
- Jentsch TJ, Steinmeyer K, Schwarz G (1990) Primary structure of Torpeto marmorata chloride channel isolated by expression cloning in Xenopus oocytes. Nature 348:510–514
- Jiang L, Chernova MN, Alper SL (1997) Secondary regulatory volume increase conferred on Xenopus oocytes by expression of AE2 anion exchanger. Am J Physiol 272, C191-C202
- Joiner WJ, Wang LY, Tang MD, Kaczmarek LK (1997) hSK4 a member of a novel subfamily of calcium-activated potassium channels. Proc Natl Acad Sci USA 94:11013–11018
- Jorgensen NK, Christensen S, Harbak H, Brown AM, Lambert IH, Hoffmann EK, Simonsen LO (1997) On the role of calcium in the regulatory volume decrease (RVD) response in Ehrlich mouse ascites tumor cells. J Membr Biol 157:281–299
- Jorgensen NK, Lambert IH, Hoffmann EK (1996) Role of LTD4 in the regulatory volume decrease response in Ehrlich ascites tumor cells. J Membr Biol 151:159–173
- Jorgensen NK, Petersen SF, Hoffmann EK (1999) Thrombin-, bradykinin-, and arachidonic acid-induced Ca²⁺ signaling in Ehrlich ascites tumor cells. Am J Physiol 276:C26–C37
- Junankar PR, Kirk K (2000) Organic osmolyte channels: a comparative view. Cell Physiol Biochem 10:355–360

- Kajimura M, O'Donnell ME, Curry FE (1997) Effect of cell shrinkage on permeability of cultured bovine aortic endothelia and frog mesenteric capillaries. J Physiol 503:413–425
- Kanli H, Norderhus E (1998) Cell volume regulation in proximal renal tubules from trout (Salmo trutta) J Exp Biol 201 (Pt 9):1405–1419
- Kashani AH, Chen BM, Grinnell AD (2001) Hypertonic enhancement of transmitter release from frog motor nerve terminals: Ca²⁺ independence and role of integrins. J Physiol 530:243–252
- Kawahara K (1990) A stretch-activated K⁺ channel in the basolateral membrane of Xenopus kidney proximal tubule cells. Pflugers Arch 415:624–629
- Kawahara K, Ogawa A, Suzuki M (1991) Hypo-osmotic activation of Ca-activated K channels in cultured rabbit kidney proximal tubule cells. Am J Physiol 260:F27–F33
- Kempson SA (1998) Differential activation of system A and betaine/GABA transport in MDCK cell membranes by hypertonic stress. Biochim Biophys Acta 1372:117–123
- Khanna R, Chang MC, Joiner WJ, Kaczmarek LK, Schlichter LC (1999) hSK4/hIK1 a calmodulin-binding KCa channel in human T lymphocytes. Roles in proliferation and volume regulation. J Biol Chem 274:14838–14849
- Kilic G, Fitz JG (2002) Heterotrimeric G-proteins activate Cl- channels through stimulation of a cyclooxygenase-dependent pathway in a model liver cell line. J Biol Chem 277:11721–11727
- Kim D (1993) Novel cation-selective mechanosensitive ion channel in the atrial cell membrane. Circ Res 72:225–231
- Kim D, Fu C (1993) Activation of a nonselective cation channel by swelling in atrial cells. J Membr Biol 135:27–37
- Kim D, Sladek CD, Aguado-Velasco C, Mathiasen JR (1995) Arachidonic acid activation of a new family of K⁺ channels in cultured rat neuronal cells. J Physiol 484 (Pt 3):643–660
- Kim RD, Roth TP, Darling CE, Ricciardi R, Schaffer BK, Chari RS (2001) Hypo-osmotic stress stimulates growth in HepG2 cells via protein kinase B-dependent activation of activator protein-1. J Gastrointest Surg 5:546–555
- Kimura K, Ito M, Amano M, Chihara K, Fukata Y, Nakafuku M, Yamamori B, Feng J, Nakano T, Okawa K, Iwamatsu A, Kaibuchi K (1996) Regulation of myosin phosphatase by Rho and Rho-associated kinase (Rho-kinase) Science 273:245–248
- Kinne RKH (1993) The role of organic osmolytes in osmoregulation: from bacteria to mammals. J Exp Zool 265:346–355
- Kinne RKH (1998) Mechanisms of osmolyte release. Contrib Nephrol 123:34-49
- Kinne RKH, Czekay RP, Grunewald JM, Mooren FC, Kinne-Saffran E (1993) Hypotonicity-evoked release of organic osmolytes from distal renal cells: systems, signals, and sidedness. Renal Physiol Biochem 16:66–78
- Kinne RKH, Boese SH, Kinne-Saffran E, Ruhfus B, Tinel H, Wehner F (1996) Osmoregulation in the renal papilla: membranes, messengers and molecules. Kidney Int 49:1686–1689
- Kinne RKH, Grunewald RW, Ruhfus B, Kinne-Saffran E (1997) Biochemistry and physiology of carbohydrates in the renal collecting duct. J Exp Zool 279:436–442
- Kinne RKH, Tinel H, Kipp H, Kinne-Saffran E (2000) Regulation of sorbitol efflux in different renal medullary cells: similarities and diversities. Cell Physiol Biochem 10:371–378
- Kinne RKH, Kipp H, Ruhfus B, Wehner F, Boese SH, Kinne-Saffran E (2001) Organic osmolyte channels in the renal medulla: Their properties and regulation. Am Zool 41:728–733
- Kinne RKH, Ruhfus B, Tinel H, Boese SH, Wehner F, Kinne–Saffran E (1995) Renal organic osmolytes: signal transduction pathways and release mechanisms. In de Santo NG, Capasso G (eds.) Acid-base and electrolyte balance. Molecular, cellular and clinical aspects. Istituto Italiano per gli Studi Filosofici, Cosenza, pp. 237–242
- Kinne-Saffran E, Kinne RKH (1997) Sorbitol uptake in plasma membrane vesicles isolated from immortalized rabbit TALH cells: activation by a Ca²⁺/calmodulin-dependent protein kinase. J Membr Biol 159:231–238
- Kinnunen PK (2000) Lipid bilayers as osmotic response elements. Cell Physiol Biochem 10:243-250
- Kirber MT, Ordway RW, Clapp LH, Walsh Jr. JV, Singer JJ (1992) Both membrane stretch and fatty acids directly activate large conductance Ca²⁺-activated K⁺ channels in vascular smooth muscle cells. FEBS Lett 297:24–28
- Kirk K (1997) Swelling-activated organic osmolyte channels. J. Membr. Biol. 158:1–16
- Kirk K, Strange K (1998) Functional properties and physiological roles of organic solute channels. Annu Rev Physiol 60:719–739
- Kirschner U, Tinel H, Rosin-Steiner S, Giffey A, Kinne RKH, Wehner F (1998) Single rat hepatocytes in primary culture as a model system for the study of regulatory volume increase (RVI) in liver. Nova Acta Leopold 306:299–303

- Kirschner U, Van Driessche W, Werner A, Wehner F (2003) Hypertonic activation of phospholemman in solitary rat hepatocytes in primary culture. FEBS Lett 537:151-156
- Kizer N, Harter L, Hruska K, Alvarez U, Duncan R (1999) Volume regulatory decrease in UMR-106.01 cells is mediated by specific α1 subunits of L-type calcium channels. Cell Biochem Biophys 31:65–79
- Klanke CA, Su YR, Callen DF, Wang Z, Meneton P, Baird N, Kandasamy RA, Orlowski J, Otterud BE, Leppert M, (1995) Molecular cloning and physical and genetic mapping of a novel human Na⁺/H⁺ exchanger (NHE5/SLC9A5) to chromosome 16q22.1. Genomics 25:615–622
- Klein JD, Lamitina ST, O'Neill WC (1999) JNK is a volume-sensitive kinase that phosphorylates the Na-K-2Cl cotransporter in vitro. Am J Physiol 277:C425–C431
- Klein M, Seeger P, Schuricht B, Alper SL, Schwab A (2000) Polarization of Na⁺/H⁺ and Cl⁻/HCO₃⁻ exchangers in migrating renal epithelial cells. J Gen Physiol 115:599–608
- Kobayashi T, Cohen P (1999) Activation of serum- and glucocorticoid-regulated protein kinase by agonists that activate phosphatidylinositide 3-kinase is mediated by 3-phosphoinositide-dependent protein kinase-1 (PDK1) and PDK2. Biochem J 339:319–328
- Koch JP, Korbmacher C (1999) Osmotic shrinkage activates nonselective cation (NSC) channels in various cell types. J Membr Biol 168:131–139
- Koch JP, Korbmacher C (2000) Mechanism of shrinkage activation of nonselective cation channels in M-1 mouse cortical collecting duct cells. J Membr Biol 177:231–242
- Köhler M, Hirschberg B, Bond CT, Kinzie JM, Marrion NV, Maylie J, Adelman JP (1996) Small-conductance, calcium-activated potassium channels from mammalian brain. Science 273:1709–1714
- Koomoa DL, Musch MW, Goldstein L (2002) Comparison of the osmolyte transport properties induced by trAE1 versus IClswell in *Xenopus* oocytes. J Membr Biol 185:57–63
- Kopito RR, Lodish HF (1985) Primary structure and transmembrane orientation of the murine anion exchange protein. Nature 316:234–238
- Korbmacher C, Volk T, Segal AS, Boulpaep EL, Frömter E (1995) A calcium-activated and nucleotide-sensitive nonselective cation channel in M-1 mouse cortical collecting duct cells. J Membr Biol 146:29–45
- Kowdley GC, Ackerman SJ, Chen Z, Szabo G, Jones LR, Moorman JR (1997) Anion, cation, and zwitterion selectivity of phospholemman channel molecules. Biophys J 72:141–145
- Krapivinsky GB, Ackerman MJ, Gordon EA, Krapivinsky LD, Clapham DE (1994) Molecular characterization of a swelling-induced chloride conductance regulatory protein, pl_{Cln}. Cell 76:439–448
- Krause U, Rider MH, Hue L (1996) Protein kinase signaling pathway triggered by cell swelling and involved in the activation of glycogen synthase and acetyl-CoA carboxylase in isolated rat hepatocytes. J Biol Chem 271:16668–16673
- Kregenow FM, Robbie DE, Orloff J (1976) Effect of norepinephrine and hypertonicity on K influx and cyclic AMP in duck erythrocytes. Am J Physiol 231:306–311
- Kubitz R, Warth R, Allert N, Kunzelmann K, Greger R (1992) Small-conductance chloride channels induced by cAMP, Ca²⁺, and hypotonicity in HT29 cells: ion selectivity, additivity and stilbene sensitivity. Pflugers Arch 421:447–454
- Kulanthaivel P, Cool DR, Ramamoorthy S, Mahesh VB, Leibach FH, Ganapathy V (1991) Transport of taurine and its regulation by protein kinase C in the JAR human placental choriocarcinoma cell line. Biochem J 277 (Pt 1), 53–58
- Kurashima K, D'Souza S, Szászi K, Ramjeesingh R, Orlowski J, Grinstein S (1999) The apical Na⁺/H⁺ exchanger isoform NHE3 is regulated by the actin cytoskeleton. J Biol Chem 274:29843–29849
- Kwon HM, Yamauchi A, Uchida S, Preston AS, Garcia-Perez A, Burg MB, Handler JS (1992) Cloning of the cDNa for a Na+/myo-inositol cotransporter a hypertonicity stress protein. J Biol Chem 267:6297– 6301
- Laich A, Gschwentner M, Krick W, Nagl UO, Fürst J, Hofer S, Susanna A, Schmarda A, Deetjen P, Burckhardt G, Paulmichl M (1997) I_{Cln} a chloride channel cloned from kidney cells, is activated during regulatory volume decrease. Kidney Int 51:477–478
- Lambert IH (1989) Leukotriene-D4 induced cell shrinkage in Ehrlich ascites tumor cells. J Membr Biol 108:165–176
- Lambert IH, Hoffmann EK (1993) Regulation of taurine transport in Ehrlich ascites tumor cells. J Membr Biol 131:67–79
- Lambert IH, Hoffmann EK (1994) Cell swelling activates separate taurine and chloride channels in Ehrlich mouse ascites tumor cells. J Membr Biol 142:289–298
- Lambert IH, Hoffmann EK, Christensen P (1987) Role of prostaglandins and leukotrienes in volume regulation by Ehrlich ascites tumor cells. J Membr Biol 98:247–256
- Lang F, Busch GL, Ritter M, Völkl H, Waldegger S, Gulbins E, Häussinger D (1998a) Functional significance of cell volume regulatory mechanisms. Physiol Rev 78:247–306

- Lang F, Busch GL, Völkl H (1998b) The diversity of volume regulatory mechanisms. Cell Physiol Biochem 8:1–45
- Lang F, Stehle T, Häussinger D (1989) Water, K⁺, H⁺, lactate and glucose fluxes during cell volume regulation in perfused rat liver. Pflugers Arch 413:209–216
- Lansman JB, Hallam TJ, Rink TJ (1987) Single stretch-activated ion channels in vascular endothelial cells as mechanotransducers? Nature 325:811–813
- Lapointe J-Y, Garneau L, Bell PD, Cardinal J (1990) Membrane crosstalk in the mammalian proximal tubule during alterations in transepithelial sodium transport. Am J Physiol 258:F339–F345
- Larsen AK, Jensen BS, Hoffmann EK (1994) Activation of protein kinase C during cell volume regulation in Ehrlich mouse ascites tumor cells. Biochim Biophys Acta 1222:477–482
- Lauf PK, Adragna NC (2000) K-Cl cotransport: properties and molecular mechanism. Cell Physiol Biochem 10:341–354
- Lauf PK, Zhang J, Gagnon KB, Delpire E, Fyffe RE, Adragna NC (2001) K-Cl cotransport: immunohistochemical and ion flux studies in human embryonic kidney (HEK293) cells transfected with full-length and C- terminal-domain-truncated KCCl cDNAs. Cell Physiol Biochem 11:143–160
- Law RO (1998) The role of taurine in the regulation of brain cell volume in chronically hyponatraemic rats. Neurochem Int 33:467–472
- Law RO (1999) Amino acid efflux and cell volume regulation in cerebrocortical minislices prepared from chronically hyponatraemic and hypernatraemic rats. Neurochem Int 35:423–430
- Lawonn P, Hoffmann EK, Hougaard C, Wehner F (2003) A cell shrinkage-induced cation conductance with a novel pharmacology in Ehrlich-Lettre-ascites tumour cells. FEBS Lett (in press)
- Leibowich S, DeLong J, Civan MM (1988) Apical Na⁺ permeability of frog skin during serosal Cl⁻ replacement. J Membr Biol 102:121–130
- Lemonnier L, Prevarskaya N, Shuba Y, Vanden Abeele F, Nilius B, Mazurier J, Skryma R (2002) Ca²⁺ modulation of volume-regulated anion channels: evidence for colocalization with store-operated channels. FASEB J 16:222–224
- Lesage F, Lazdunski M (2000) Molecular and functional properties of two-pore-domain potassium channels. Am J Physiol 279:F793–F801
- Levitan IB (1994) Modulation of ion channels by protein phosphorylation and dephosphorylation. Annu Rev Physiol 56:193–212
- Lewis RA, Bursell JD, Kirk K (1996) Anion-selectivity of the swelling-activated osmolyte channel in eel erythrocytes. J Membr Biol 149:103–111
- Lewis RS, Cahalan MD (1995) Potassium and calcium channels in lymphocytes. Annu Rev Immunol 13:623-653
- Lewis RS, Ross PE, Cahalan MD (1993) Chloride channels activated by osmotic stress in T lymphocytes. J Gen Physiol 101:801–826
- Lewis SA, Butt AG, Bowler MJ, Leader JP, Macknight ADC (1985) Effects of anions on cellular volume and transepithelial Na⁺ transport across toad urinary bladder. J Membr Biol 83:119–137
- Li CH, Breton S, Morrison R, Cannon CL, Emma F, Sanchez-Olea R, Bear C, Strange K (1998) Recombinant pI_{Cln} forms highly cation-selective channels when reconstituted into artificial and biological membranes. J Gen Physiol 112:727–736
- Li G, Liu Y, Olson JE (2002) Calcium/calmodulin-modulated chloride and taurine conductances in cultured rat astrocytes. Brain Res 925:1–8
- Li Q, Jungmann V, Kiyatkin A, Low PS (1996) Prostaglandin E2 stimulates a Ca²⁺-dependent K⁺ channel in human erythrocytes and alters cell volume and filterability. J Biol Chem 271:18651–18656
- Liedtke CM, Cole TS (2002) Activation of NKCC1 by hyperosmotic stress in human tracheal epithelial cells involves PKC-delta and ERK. Biochim Biophys Acta 1589:77–88
- Liedtke W, Choe Y, Marti-Renom MA, Bell AM, Denis CS, Sali A, Hudspeth AJ, Friedman JM, Heller S (2000) Vanilloid receptor-related osmotically activated channel (VR-OAC) a candidate vertebrate osmoreceptor. Cell 103:525–535
- Light DB, Mertins TM, Belongia JA, Witt CA (1997) 5-Lipoxygenase metabolites of arachidonic acid regulate volume decrease by mudpuppy red blood cells. J Membr Biol 158:229–239
- Lima L, Cubillos S, Guerra A (2000) Regulation of high affinity taurine transport in goldfish and rat retinal cells. Adv Exp Med Biol 483:431–440
- Ling BN, Webster CL, Eaton DC (1992) Eicosanoids modulate apical Ca²⁺-dependent K⁺ channels in cultured rabbit principal cells. Am J Physiol 263:F116–F126
- Lock H, Valverde MA (2000) Contribution of the IsK (MinK) potassium channel subunit to regulatory volume decrease in murine tracheal epithelial cells. J Biol Chem 275:34849–34852
- Logsdon NJ, Kang J, Togo JA, Christian EP, Aiyar J (1997) A novel gene, hKCa4, encodes the calcium-activated potassium channel in human T lymphocytes. J Biol Chem 272:32723–32726

- Low SY, Rennie MJ, Taylor PM (1996) Modulation of glycogen synthesis in rat skeletal muscle by changes in cell volume. J Physiol 495:299–303
- Low SY, Taylor PM (1998) Integrin and cytoskeletal involvement in signalling cell volume changes to glutamine transport in rat skeletal muscle. J Physiol 512:481–485
- Luby-Phelps K (2000) Cytoarchitecture and physical properties of cytoplasm: volume, viscosity, diffusion, intracellular surface area. Int Rev Cytol 192:189–221
- Lucke B, McCutcheon M (1932) The living cell as an osmotic system and its permeability to water. Physiol Rev 12:68–139
- Lundgren DW, Moore RM, Collins PL, Moore JJ (1997) Hypotonic stress increases cyclooxygenase-2 expression and prostaglandin release from amnion-derived WISH cells. J Biol Chem 272:20118–20124
- Ma TH, Verkman AS (1999) Aquaporin water channels in gastrointestinal physiology. J Physiol (Lond) 517:317–326
- MacLeod RJ, Hamilton JR (1999a) Ca²⁺/Calmodulin kinase II and decreases in intracellular pH are required to activate K⁺ channels after substantial swelling in villus epithelial cells. J Membr Biol 172:59–66
- MacLeod RJ, Hamilton JR (1999b) Increases in intracellular pH and Ca²⁺ are essential for K⁺ channel activation after modest "physiological" swelling in villus epithelial cells. J Membr Biol 172:47–58
- Macri P, Breton S, Marsolais M, Lapointe JY, Laprade R (1997) Hypertonicity decreases basolateral K⁺ and Cl⁻ conductances in rabbit proximal convoluted tubule. J Membr Biol 155:229–237
- Mähler S, Kinne-Saffran E, Fujisue H, Kinne RKH, Föllmann W (1998) Regulation of sorbitol content in cultured porcine urinary bladder epithelial cells. Am J Physiol 274:F342–F347
- Marchenko SM, Sage SO (2000) Hyperosmotic but not hypo-osmotic stress evokes a rise in cytosolic Ca²⁺ concentration in endothelium of intact rat aorta. Exp Physiol 85:151–157
- Margalit A, Livne AA, Funder J, Granot Y (1993a) Initiation of RVD response in human platelets: mechanical-biochemical transduction involves pertussis-toxin-sensitive G protein and phospholipase A2. J Membr Biol 136:303–311
- Margalit A, Sofer Y, Grossman S, Reynaud D, Pace-Asciak CR, Livne AA (1993b) Hepoxilin A3 is the endogenous lipid mediator opposing hypotonic swelling of intact human platelets. Proc Natl Acad Sci USA 90:2589–2592
- Marinelli RA, LaRusso NF (1997) Aquaporin water channels in liver: Their significance in bile formation. Hepatology 26:1081–1084
- Marunaka Y, Niisato N, O'Brodovich H, Post M, Tanswell AK (1999) Roles of Ca²⁺ and protein tyrosine kinase in insulin action on cell volume via Na⁺ and K⁺ channels and Na⁺/K⁺/2Cl⁻ cotransporter in fetal rat alveolar type II pneumocyte. J Membr Biol 168:91–101
- Mastrocola T, Lambert IH, Kramhoft B, Rugolo M, Hoffmann EK (1993) Volume regulation in human fibroblasts: role of Ca²⁺ and 5-lipoxygenase products in the activation of the Cl⁻ efflux. J Membr Biol 136:55–62
- Matskevitch I, Wagner CA, Stegen C, Broer S, Noll B, Risler T, Kwon HM, Handler JS, Waldegger S, Busch AE, Lang F (1999) Functional characterization of the Betaine/gamma-aminobutyric acid transporter BGT-1 expressed in *Xenopus* oocytes. J Biol Chem 274:16709–16716
- Matsuoka Y, Yamauchi A, Nakanishi T, Sugiura T, Kitamura H, Horio M, Takamitsu Y, Ando A, Imai E, Hori M (1999) Response to hypertonicity in mesothelial cells: role of Na⁺/myo-inositol cotransporter. Nephrol Dial Transplant 14:1217–1223
- Maunsbach AB, Marples D, Chin E, Ning G, Bondy C, Agre P, Nielsen S (1997) Aquaporin-1 water channel expression in human kidney. J Am Soc Nephrol 8:1–14
- McCarty NA, O'Neil RG (1991) Calcium-dependent control of volume regulation in renal proximal tubule cells: I. Swelling-activated Ca²⁺ entry and release. J Membr Biol 123:149–160
- Meier R, Thelen M, Hemmings BA (1998) Inactivation and dephosphorylation of protein kinase Balpha (PKBalpha) promoted by hyperosmotic stress. EMBO J 17:7294–7303
- Mercado A, Song LY, Vázquez N, Mount DB, Gamba G (2000) Functional comparison of the K+-Cl⁻ cotransporters KCC1 and KCC4. J Biol Chem 275:30326–30334
- Meyer K, Korbmacher C (1996) Cell swelling activates ATP-dependent voltage-gated chloride channels in M-1 mouse cortical collecting duct cells. J Gen Physiol 108:177–193
- Mignen O, Le Gall C, Harvey BJ, Thomas S (1999) Volume regulation following hypotonic shock in isolated crypts of mouse distal colon. J Physiol 515:501–510
- Minton AP (1994) Influence of macromolecular crowding on intracellular association reactions: possible role in volume regulation. In Strange K (ed) Cellular and molecular physiology of cell volume regulation. CRC Press, Boca Raton, pp 181–190
- Minton AP, Colclasure GC, Parker JC (1992) Model for the role of macromolecular crowding in regulation of cellular volume. Proc Natl Acad Sci USA 89:10504–10506

- Mitchell CH, Zhang JJ, Wang L, Jacob TJ (1997) Volume-sensitive chloride current in pigmented ciliary epithelial cells: role of phospholipases. Am J Physiol 272:C212–C222
- Miyai A, Yamauchi A, Moriyama T, Kaneko T, Takenaka M, Sugiura T, Kitamura H, Ando A, Tohyama M, Shimada S, Imai E, Kamada T (1996) Expression of betaine transporter mRNA: its unique localization and rapid regulation in rat kidney. Kidney Int 50:819–827
- Miyauchi A, Notoya K, Mikuni-Takagaki Y, Takagi Y, Goto M, Miki Y, Takano-Yamamoto T, Jinnai K, Takahashi K, Kumegawa M, Chihara K, Fujita T (2000) Parathyroid hormone-activated volume-sensitive calcium influx pathways in mechanically loaded osteocytes. J Biol Chem 275:3335–3342
- Moeckel GW, Lai LW, Guder WG, Kwon HM, Lien YH (1997) Kinetics and osmoregulation of Na⁺- and Cl⁻-dependent betaine transporter in rat renal medulla. Am J Physiol 272:F100–F106
- Mollerup J, Lambert IH (1998) Calyculin A modulates the kinetic constants for the Na⁺-coupled taurine transport in Ehrlich ascites tumor cells. Biochim Biophys Acta 1371:335–344
- Mongin AA, Cai Z, Kimelberg HK (1999) Volume-dependent taurine release from cultured astrocytes requires permissive [Ca²⁺]_i and calmodulin. Am J Physiol 277:C823–C832
- Montell C, Rubin GM (1989) Molecular characterization of the *Drosophila* trp locus: a putative integral membrane protein required for phototransduction. Neuron 2:1313–1323
- Montrose-Rafizadeh C, Guggino WB (1991) Role of intracellular calcium in volume regulation by rabbit medullary thick ascending limb cells. Am J Physiol 260:F402–F409
- Montrose-Rafizadeh C, Guggino WB, Montrose MH (1991) Cellular differentiation regulates expression of Cl⁻ transport and cystic fibrosis transmembrane conductance regulator mRNA in human intestinal cells. J Biol Chem 266:4495–4499
- Mooren FC, Kinne RKH (1994) Intracellular calcium in primary cultures of rat renal inner medullary collecting duct cells during variations of extracellular osmolality. Pflugers Arch 427:463–472
- Moorman JR, Ackerman SJ, Kowdley GC, Griffin MP, Mounsey JP, Chen ZH, Cala SE, O'Brian JJ, Szabo G, Jones LR (1995) Unitary anion currents through phospholemman channel molecules. Nature 377:737–740
- Moorman JR, Jones LR (1998) Phospholemman: A cardiac taurine channel involved in regulation of cell volume. Adv Exp Med Biol 442:219–228
- Moorman JR, Palmer CJ, John III JE, Durieux ME Jones LR (1992) Phospholemman expression induces a hyperpolarization-activated chloride current in Xenopus oocytes. J Biol Chem 267:14551–14554
- Morales-Mulia M, Pasantes-Morales H, Morán J (2000) Volume sensitive efflux of taurine in HEK293 cells overexpressing phospholemman. Biochim Biophys Acta 1496:252–260
- Morales-Mulia S, Cardin V, Torres-Marquez ME Crevenna A, Pasantes-Morales H (2001) Influence of protein kinases on the osmosensitive release of taurine from cerebellar granule neurons. Neurochem Int 38:153–161
- Morán J, Morales-Mulia M, Pasantes-Morales H (2001) Reduction of phospholemman expression decreases osmosensitive taurine efflux in astrocytes. Biochim Biophys Acta 1538:313–320
- Morris CE (1990) Mechanosensitive ion channels. J Membr Biol 113:93-107
- Motais R, Guizouarn H, Garcia-Romeu F (1991) Red cell volume regulation: the pivotal role of ionic strength in controlling swelling-dependent transport systems. Biochim Biophys Acta 1075:169–180
- Motais R, Fievet B, Borgese F, Garcia-Romeu F (1997) Association of the band 3 protein with a volume-activated, anion and amino acid channel: a molecular approach. J Exp Biol 200:361–367
- Mounsey JP, Lu KP, Patel MK, Chen ZH, Horne LT, John JE, III, Means AR, Jones LR, Moorman JR (1999) Modulation of Xenopus oocyte-expressed phospholemman-induced ion currents by co-expression of protein kinases. Biochim Biophys Acta 1451:305–318
- Mount DB, Mercado A, Song L, Xu J, George AL, Jr., Delpire E, Gamba G (1999) Cloning and characterization of KCC3 and KCC4, new members of the cation-chloride cotransporter gene family. J Biol Chem 274:16355–16362
- Mountian I, Chou KY, van Driessche W (1996) Electrolyte transport mechanisms involved in regulatory volume increase in C6 glioma cells. Am J Physiol 271:C1041–C1048
- Musch MW, Goldstein L (1990) Hypotonicity stimulates phosphatidylcholine hydrolysis and generates diacylglycerol in erythrocytes. J Biol Chem 265:13055–13059
- Musch MW, Leffingwell TR, Goldstein L (1994) Band 3 modulation and hypotonic-stimulated taurine efflux in skate erythrocytes. Am J Physiol 266:R65–R74
- Musch MW, Luer CA, Davis-Amaral EM, Goldstein L (1997) Hypotonic stress induces translocation of the osmolyte channel protein pICln in embryonic skate (Raja eglanteria) heart. J Exp Zool 277:460–463
- Musch MW, Davis-Amaral EM, Vandenburgh HH, Goldstein L (1998) Hypotonicity stimulates translocation of ICln in neonatal rat cardiac myocytes. Pflugers Arch 436:415–422
- Muto S, Ohtaka A, Nemoto J, Kawakami K, Asano Y (1998) Effects of hyperosmolality on Na, K-ATPase gene expression in vascular smooth muscle cells. J Membr Biol 162:233–245

- Nagasaki M, Ye L, Duan D, Horowitz B, Hume JR (2000) Intracellular cyclic AMP inhibits native and recombinant volume-regulated chloride channels from mammalian heart. J Physiol 523:705–717
- Nakamura H, Huang SH, Takakura K (1996) High-affinity taurine uptake and its regulation by protein kinase C in human glioma cells. Adv Exp Med Biol 403:377–384
- Nakanishi T, Balaban RS, Burg MB (1988) Survey of osmolytes in renal cell lines. Am J Physiol 255:C181-C191
- Nakanishi T, Turner RJ, Burg MB (1989) Osmoregulatory changes in myo-inositol transport by renal cells. Proc Natl Acad Sci USA 86:6002–6006
- Nakanishi T, Turner RJ, Burg MB (1990) Osmoregulation of betaine transport in mammalian renal medullary cells. Am J Physiol 258:F1061–F1067
- Napathorn S, Spring KR (1994) Further characterization of the sorbitol permease in PAP-HT25 cells. Am J Physiol 267:C514–C519
- Naray-Fejes-Toth A, Canessa C, Cleaveland ES, Aldrich G, Fejes-Toth G (1999) sgk is an aldosterone-induced kinase in the renal collecting duct. Effects on epithelial Na⁺ channels. J Biol Chem 274:16973– 16978
- Nath SK, Hang CY, Levine SA, Yun CH, C., Montrose MH, Donowitz M, Tse CM (1996) Hyperosmolarity inhibits the Na⁺/H⁺ exchanger isoforms NHE2 and NHE3: An effect opposite to that on NHE1. Am J Physiol 270:G431–G441
- Nelson DJ, Tien XY, Xie WW, Brasitus TA, Kaetzel MA, Dedman JR (1996) Shrinkage activates a nonselective conductance: involvement of a Walker-motif protein and PKC. Am J Physiol 270:C179–C191
- Niemeyer MI, Cid LP, Barros LF, Sepulveda FV (2001) Modulation of the two-pore domain acid-sensitive K⁺ channel TASK-2 (KCNK5) by changes in cell volume. J Biol Chem 276:43166–43174
- Niggli V, Andreoli C, Roy C, Mangeat P (1995) Identification of a phosphatidylinositol-4,5-bis-phosphate-binding domain in the N-terminal region of ezrin. FEBS Lett 376:172–176
- Nilius B, Droogmans G (2001) Ion channels and their functional role in vascular endothelium. Physiol Rev 81:1415-1459
- Nilius B, Sehrer J, Droogmans G (1994a) Permeation properties and modulation of volume-activated Clcurrents in human endothelial cells. Br J Pharmacol 112:1049–1056
- Nilius B, Sehrer J, Viana F, De Greef C, Raeymaekers L, Eggermont J, Droogmans G (1994b) Volume-activated Cl⁻ currents in different mammalian nonexcitable cell types. Pflugers Arch 428:364–371
- Nilius B, Sehrer J, De Smet P, van Driessche W, Droogmans G (1995) Volume regulation in a toad epithelial cell line: Role of coactivation of K⁺ and Cl⁻ channels. J Physiol (Lond) 487:367–378
- Nilius B, Eggermont J, Voets T, Buyse G, Manolopoulos V, Droogmans G (1997) Properties of volumeregulated anion channels in mammalian cells. Prog Biophys Mol Biol 68:69–119
- Nilius B, Prenen J, Voets T, Eggermont J, Droogmans G (1998) Activation of volume-regulated chloride currents by reduction of intracellular ionic strength in bovine endothelial cells. J Physiol 506:353–361
- Nilius B, Voets T, Prenen J, Barth H, Aktories K, Kaibuchi K, Droogmans G, Eggermont J (1999) Role of Rho and Rho kinase in the activation of volume-regulated anion channels in bovine endothelial cells. J Physiol (Lond) 516:67–74
- Nilius B, Prenen J, Wissenbach U, Bodding M, Droogmans G (2001) Differential activation of the volumesensitive cation channel TRP12 (OTRPC4) and volume-regulated anion currents in HEK-293 cells. Pflugers Arch 443:227–233
- Noel J, Roux D, Pouysségur J (1996) Differential localization of Na⁺/H⁺ exchanger isoforms (NHE1 and NHE3) in polarized epithelial cell lines. J Cell Sci 109:929–939
- Noulin JF, Brochiero E, Lapointe JY, Laprade R (1999) Two types of K⁺ channels at the basolateral membrane of proximal tubule: inhibitory effect of taurine. Am J Physiol 277:F290–F297
- Novak JE, Agranoff BW, Fisher SK (2000) Regulation of Myo-inositol homeostasis in differentiated human NT2-N neurons. Neurochem Res 25:561–566
- O'Donnell ME, Martinez A, Sun D (1995) Endothelial Na-K-Cl cotransport regulation by tonicity and hormones: phosphorylation of cotransport protein. Am J Physiol 269:C1513-C1523
- O'Flaherty L, Stapleton PP, Redmond HP, Bouchier-Hayes DJ (1997) Intestinal taurine transport: a review. Eur J Clin Invest 27:873–880
- O'Neil RG, Leng L (1997) Osmo-mechanically sensitive phosphatidylinositol signaling regulates a Ca^{2+} influx channel in renal epithelial cells. Am J Physiol 273:F120–F128
- Ohta M, Tanimoto T, Tanaka A (1990) Characterization of aldose reductase and aldehyde reductase from the medulla of rat kidney. Chem Pharm Bull (Tokyo) 38:1639–1643
- Ohtaka A, Muto S, Nemoto J, Kawakami K, Nagano K, Asano Y (1996) Hyperosmolality stimulates Na-K-ATPase gene expression in inner medullary collecting duct cells. Am J Physiol 270:F728–F738
- Ohtsuyama M, Suzuki Y, Samman G, Sato F, Sato K (1993) Cell volume analysis of gramicidin-treated eccrine clear cells to study regulation of Cl channels. Am J Physiol 265:C1090–C1099

- Oike M, Droogmans G, Nilius B (1994) The volume-activated chloride current in human endothelial cells depends on intracellular ATP. Pflugers Arch 427:184–186
- Oike M, Kimura C, Koyama T, Yoshikawa M, Ito Y (2000) Hypotonic stress-induced dual Ca²⁺ responses in bovine aortic endothelial cells. Am J Physiol Heart Circ Physiol 279:H630–H638
- Oiki S, Kubo M, Okada Y (1994) Mg²⁺ and ATP-dependence of volume-sensitive Cl⁻ channels in human epithelial cells. Jpn J Physiol 44: S77–S79
- Okada Y (1997) Volume expansion-sensing outward-rectifier Cl⁻ channel: fresh start to the molecular identity and volume sensor. Am J Physiol 273:C755–C789
- Okada Y (1998) Cell volume-sensitive chloride channels. Contrib Nephrol 123:21–33
- Okada Y, Hazama A (1989) Volume-regulatory ion channels in epithelial cells. News Physiol Sci 4:238–242
- Okada Y, Oiki S, Hazama A, Morishima S (1998) Criteria for the molecular identification of the volumesensitive outwardly rectifying Cl⁻ channel. J Gen Physiol 112:365–367
- Okada Y, Maeno E, Shimizu T, Dezaki K, Wang J, Morishima S (2001) Receptor-mediated control of regulatory volume decrease (RVD) and apoptotic volume decrease (AVD) J Physiol (Lond) 532:3–16
- Oliet SH, Bourque CW (1993) Mechanosensitive channels transduce osmosensitivity in supraoptic neurons. Nature 364:341–343
- Olson JE (1999) Osmolyte contents of cultured astrocytes grown in hypo-osmotic medium. Biophys Acta 1453:175–179
- Ono S, Mougouris T, DuBose TD, Jr., Sansom SC (1994) ATP and calcium modulation of nonselective cation channels in IMCD cells. Am J Physiol 267:F558–F565
- Orlic T, Loomis WH, Shreve A, Namiki S, Junger WG (2002) Hypertonicity increases cAMP in PMN and blocks oxidative burst by PKA-dependent and -independent mechanisms. Am J Physiol Cell Physiol 282:C1261–C1269
- Orlowski J, Grinstein S (1997) Na+/H+ exchangers of mammalian cells. J Biol Chem 272:22373-22376
- Pácha J, Frindt G, Sackin H, Palmer LG (1991) Apical maxi K channels in intercalated cells of CCT. Am J Physiol 261:F696–F705
- Palmer CJ, Scott BT, Jones LR (1991) Purification and complete sequence determination of the major plasma membrane substrate for cAMP-dependent protein kinase and protein kinase C in myocardium. J Biol Chem 266:11126–11130
- Papakonstanti EA, Vardaki EA, Stournaras C (2000) Actin cytoskeleton: a signaling sensor in cell volume regulation. Cell Physiol Biochem 10:257–264
- Papassotiriou J, Eggermont J, Droogmans G, Nilius B (2001) Ca²⁺-activated Cl⁻ channels in Ehrlich ascites tumor cells are distinct from mCLCA1:2 and 3. Pflugers Arch 442:273–279
- Park K, Lee S, Elliott AC, Kim JS, Lee JH (2002) Swelling-induced Ca²⁺ release from intracellular calcium stores in rat submandibular gland acinar cells. J Membr Biol 186:165–176
- Park K-P, Beck JS, Douglas IJ, Brown PD (1994) Ca²⁺-activated K⁺ channels are involved in regulatory volume decrease in acinar cells isolated from the rat lacrimal gland. J Membr Biol 141:193–201
- Parker JC, Dunham PB, Minton AP (1995) Effects of ionic strength on the regulation of Na/H exchange and K-Cl cotransport in dog red blood cells. J Gen Physiol 105:677–699
- Pasantes-Morales H, Chacon E, Sanchez-Olea R, Moran J (1994a) Volume regulation in cultured neurons: pivotal role of taurine. Adv Exp Med Biol 359:317–323
- Pasantes-Morales H, Murray RA, Sanchez-Olea R, Moran J (1994b) Regulatory volume decrease in cultured astrocytes. II. Permeability pathway to amino acids and polyols. Am J Physiol 266:C172–C178
- Pasantes-Morales H, Cardin V, Tuz K (2000a) Signaling events during swelling and regulatory volume decrease. Neurochem Res 25:1301–1314
- Pasantes-Morales H, Franco R, Torres-Marquez ME, Hernandez-Fonseca K, Ortega A (2000b) Amino acid osmolytes in regulatory volume decrease and isovolumetric regulation in brain cells: contribution and mechanisms. Cell Physiol Biochem 10:361–370
- Paulmichl M, Li Y, Wickman K, Ackerman M, Peralta E, Clapham D (1992) New mammalian chloride channel identified by expression cloning. Nature 356:238–241
- Payne JA (1997) Functional characterization of the neuronal-specific K-Cl cotransporter: implications for [K⁺]_o regulation. Am J Physiol 273:C1516–C1525
- Pearson MM, Lu J, Mount DB, Delpire E (2001) Localization of the K+-Cl⁻ cotransporter, KCC3, in the central and peripheral nervous systems: expression in the choroid plexus, large neurons and white matter tracts. Neuroscience 103:481–491
- Pedersen S, Hoffmann EK, Hougaard C, Lambert IH (2000) Cell shrinkage is essential in lysophosphatidic acid signaling in Ehrlich ascites tumor cells. J Membr Biol 173:19-29
- Pedersen SF, Hoffmann EK, Mills JW (2001) The cytoskeleton and cell volume regulation. Comp Biochem Physiol A Mol Integr Physiol 130:385–399

- Pedersen SF, Kramhoft B, Jorgensen NK, Hoffmann EK (1996) Shrinkage-induced activation of the Na⁺/ H⁺ exchanger in Ehrlich ascites tumor cells: mechanisms involved in the activation and a role for the exchanger in cell volume regulation. J Membr Biol 149:141–159
- Pedersen SF, Prenen J, Droogmans G, Hoffmann EK, Nilius B (1998) Separate swelling- and Ca²⁺-activated anion currents in Ehrlich ascites tumor cells. J Membr Biol 163:97–110
- Perlman DF, Goldstein L (1999) Organic osmolyte channels in cell volume regulation in vertebrates. J Exp Zool 283:725–733
- Peters-Regehr T, Bode JG, Kubitz R, Häussinger D (1999) Organic osmolyte transport in quiescent and activated rat hepatic stellate cells (Ito cells) Hepatology 29:173–180
- Petty CN, Lucero MT (1999) Characterization of a Na⁺-dependent betaine transporter with Cl⁻ channel properties in squid motor neurons. J Neurophysiol 81:1567–1574
- Pon DC, Hill CE (1997) Existence, properties, and functional expression of "Maxi- K"-type, Ca²⁺-activated K⁺ channels in short-term cultured hepatocytes. J Cell Physiol 171:87–94
- Popp R, Hoyer J, Gögelein H (1993) Mechanosensitive nonselective cation channels in the antiluminal membrane of cerebral capillaries (blood-brain barrier) EXS 66:101–105
- Popp R, Hoyer J, Meyer J, Galla H-J, Gögelein H (1992) Stretch-activated nonselective cation channels in the antiluminal membrane of porcine cerebral capillaries. J Physiol (Lond) 454:435–449
- Porcellati F, Hosaka Y, Hlaing T, Togawa M, Larkin DD, Karihaloo A, Stevens MJ, Killen PD, Greene DA (1999) Alternate splicing in human Na⁺-MI cotransporter gene yields differentially regulated transport isoforms. Am J Physiol 276:C1325–C1337
- Preston AS, Yamauchi A, Kwon HM, Handler JS (1995) Activators of protein kinase A and of protein kinase C inhibit MDCK cell myo-inositol and betaine uptake. J Am Soc Nephrol 6:1559–1564
- Qian X, Vinnakota S, Edwards C, Sarkar HK (2000) Molecular characterization of taurine transport in bovine aortic endothelial cells. Biochim Biophys Acta 1509:324–334
- Race JE, Makhlouf FN, Logue PJ, Wilson FH, Dunham PB, Holtzman EJ (1999) Molecular cloning and functional characterization of KCC3 a new K-Cl cotransporter. Am J Physiol 277:C1210–C1219
- Rasola A, Galietta LJ, Barone V, Romeo G, Bagnasco S (1995) Molecular cloning and functional characterization of a GABA/betaine transporter from human kidney. FEBS Lett 373:229–233
- Ren XD, Kiosses WB, Schwartz MA (1999) Regulation of the small GTP-binding protein Rho by cell adhesion and the cytoskeleton. EMBO J 18:578–585
- Renshaw MW, Toksoz D, Schwartz MA (1996) Involvement of the small GTPase rho in integrin-mediated activation of mitogen-activated protein kinase. J Biol Chem 271:21691–21694
- Ritter M, Fürst J, Wöll E, Chwatal S, Gschwentner M, Lang F, Deetjen P, Paulmichl M (2001) Na⁺/H⁺ exchangers: Linking osmotic dysequilibrium to modified cell function. Cell Physiol Biochem 11:1–18
- Ritter M, Steidl M, Lang F (1991) Inhibition of ion conductances by osmotic shrinkage of Madin-Darby canine kidney cells. Am J Physiol 261:C602–C607
- Robson L, Hunter M (1994) Volume-activated, gadolinium-sensitive whole-cell currents in single proximal cells of frog kidney. Pflugers Arch 429:98–106
- Roe MW, Moore AL, Lidofsky SD (2001) Purinergic-independent calcium signaling mediates recovery from hepatocellular swelling: implications for volume regulation. J Biol Chem 276:30871–30877
- Roman R, Feranchak AP, Troetsch M, Dunkelberg JC, Kilic G, Schlenker T, Schaack J, Fitz JG (2002) Molecular characterization of volume-sensitive SK_{Ca} channels in human liver cell lines. Am J Physiol 282:G116–G122
- Roman RM, Bodily KO, Wang Y, Raymond JR, Fitz JG (1998) Activation of protein kinase C alpha couples cell volume to membrane Cl⁻ permeability in HTC hepatoma and Mz-ChA-1 cholangiocarcinoma cells. Hepatology 28:1073–1080
- Roman RM, Smith RL, Feranchak AP, Clayton GH, Doctor RB, Fitz JG (2001) ClC-2 chloride channels contribute to HTC cell volume homeostasis. Am J Physiol 280:G344–G353
- Roy G (1995) Amino acid current through anion channels in cultured human glial cells. J Membr Biol 147:35–44
- Roy G, Banderali U (1994) Channels for ions and amino acids in kidney cultured cells (MDCK) during volume regulation. J Exp Zool 268:121–126
- Roy G, Malo C (1992) Activation of amino acid diffusion by a volume increase in cultured kidney (MDCK) cells. J Membr Biol 130:83–90
- Ruhfus B, Kinne RKH (1996) Hypotonicity-activated efflux of taurine and myo-inositol in rat inner medullary collecting duct cells: evidence for a major common pathway. Kidney Blood Press Res 19:317–324
- Ruhfus B, Tinel H, Kinne RKH (1996) Role of G-proteins in the regulation of organic osmolyte efflux from isolated rat renal inner medullary collecting duct cells. Pflugers Arch 433:35–41
- Ruhfus B, Bauernschmitt HG, Kinne RKH (1998) Properties of a polarized primary culture from rat renal inner medullary collecting duct (IMCD) cells. In Vitro Cell Dev Biol Anim 34:227–231

- Russell JM (2000) Sodium-potassium-chloride cotransport. Physiol Rev 80:211–276
- Sackin H (1989) A stretch-activated K+ channel sensitive to cell volume. Proc Natl Acad Sci USA 86:1731–1735
- Sadoshima J, Izumo S (1993) Mechanical stretch rapidly activates multiple signal transduction pathways in cardiac myocytes: potential involvement of an autocrine/paracrine mechanism. EMBO J 12:1681–1692
- Saier MH, Jr (2000) Families of proteins forming transmembrane channels. J Membr Biol 175:165-180
- Sajan MP, Bandyopadhyay G, Kanoh Y, Standaert ML, Quon MJ, Reed BC, Dikic I, Farese RV (2002) Sorbitol activates atypical protein kinase C and GLUT4 glucose transporter translocation/glucose transport through proline-rich tyrosine kinase-2, the extracellular signal-regulated kinase pathway and phospholipase D. Biochem J 362:665–674
- Sakai H, Kakinoki B, Diener M, Takeguchi N (1996) Endogenous arachidonic acid inhibits hypotonicallyactivated Cl⁻ channels in isolated rat hepatocytes. Jpn J Physiol 46:311–318
- Sanchez OR, Pasantes-Morales H, Lazaro A, Cereijido M (1991) Osmolarity-sensitive release of free amino acids from cultured kidney cells (MDCK) J Membr Biol 121:1–9
- Sanchez-Olea R, Moran J, Schousboe A, Pasantes-Morales H (1991) Hypo-osmolarity-activated fluxes of taurine in astrocytes are mediated by diffusion. Neurosci Lett 130:233–236
- Sanguinetti MC, Curran ME, Zou A, Shen J, Spector PS, Atkinson DL, Keating MT (1996) Coassembly of KvLQT1 and minK (IsK) proteins to form cardiac IKs potassium channel. Nature 384:80–83
- Sardet C, Counillon L, Franchi A, Pouyssegur J (1990) Growth factors induce phosphorylation of the Na⁺/ H⁺ antiporter, glycoprotein of 110 kD. Science 247:723–726
- Sardet C, Fafournoux P, Pouyssegur J (1991) Alpha-thrombin, epidermal growth factor, and okadaic acid activate the Na⁺/H⁺ exchanger, NHE-1, by phosphorylating a set of common sites. J Biol Chem 266:19166–19171
- Satsu H, Miyamoto Y, Shimizu M (1999) Hypertonicity stimulates taurine uptake and transporter gene expression in Caco-2 cells. Biochim Biophys Acta 1419:89–96
- Schirmer T, Keller TA, Wang YF, Rosenbusch JP (1995) Structural basis for sugar translocation through maltoporin channels at 3.1 A resolution. Science 267:512–514
- Schlatter E (1993) Regulation of ion channels in the cortical collecting duct. Renal Physiol Biochem 16:21–36
- Schlatter E, Ankorina-Stark I, Cermak R, Haxelmans S, Kleta R, Hirsch JR (1997) Cell shrinkage activates a cation conductance in principal cells of rat cortical collecting duct. Cell Physiol Biochem 7:321-332
- Schliess F, Sinning R, Fischer R, Schmalenbach C, Häussinger D (1996) Calcium-dependent activation of Erk-1 and Erk-2 after hypo-osmotic astrocyte swelling. Biochem J 320:167–171
- Schmidt-Rose T, Jentsch TJ (1997) Transmembrane topology of a CLC chloride channel. Proc Natl Acad Sci USA 94:7633–7638
- Schneider SW, Pagel P, Rotsch C, Danker T, Oberleithner H, Radmacher M, Schwab A (2000) Volume dynamics in migrating epithelial cells measured with atomic force microscopy. Pflugers Arch 439:297–303
- Schoenwaelder SM, Burridge K (1999) Bidirectional signaling between the cytoskeleton and integrins. Curr Opin Cell Biol 11:274–286
- Schreiber M, Salkoff L (1997) A novel calcium-sensing domain in the BK channel. Biophys J 73:1355–1363
- Schumacher PA, Sakellaropoulos G, Phipps DJ, Schlichter LC (1995) Small-conductance chloride channels in human peripheral T lymphocytes. J Membr Biol 145:217–232
- Schütt W, Sackin H (1997) A new technique for evaluating volume sensitivity of ion channels. Pflugers Arch 433:368–375
- Schwab A, Oberleithner H (1996) Plasticity of renal epithelial cells: The way a potassium channel supports migration. Pflugers Arch 432:R87-R93
- Schwab A, Westphale H-J, Wojnowski L, Wünsch S, Oberleithner H (1993) Spontaneously oscillating K⁺ channel activity in transformed Madin-Darby canine kidney cells. J Clin Invest 92:218–223
- Schwab A, Schuricht B, Seeger P, Reinhardt J, Dartsch PC (1999) Migration of transformed renal epithelial cells is regulated by K⁺ channel modulation of actin cytoskeleton and cell volume. Pflugers Arch 438:330–337
- Schwiebert EM, Mills JW, Stanton BA (1994) Actin-based cytoskeleton regulates a chloride channel and cell volume in a renal cortical collecting duct cell line. J Biol Chem 269:7081–7089
- Setiawan I, Henke G, Feng Y, Böhmer C, Vasilets LA, Schwarz W, Lang F (2002) Stimulation of *Xenopus* oocyte Na⁺,K⁺ ATPase by the serum and glucocorticoid-dependent kinase sgk1. Pflugers Arch 444:426–431

- Shen MR, Chou CY, Browning JA, Wilkins RJ, Ellory JC (2001) Human cervical cancer cells use Ca²⁺ signalling, protein tyrosine phosphorylation and MAP kinase in regulatory volume decrease. J Physiol 537:347–362
- Shen MR, Furla P, Chou CY, Ellory JC (2002) Myosin light chain kinase modulates hypotonicity-induced Ca²⁺ entry and Cl⁻ channel activity in human cervical cancer cells. Pflugers Arch 444:276–285
- Shennan D, McNeillie S, Curran D (1994) The effect of a hypo-osmotic shock on amino acid efflux from lactating rat mammary tissue: stimulation of taurine and glycine efflux via a pathway distinct from anion exchange and volume-activated anion channels. Exp Physiol 79:797–808.
- Shiga N, Wangemann P (1995) Ion selectivity of volume regulatory mechanisms present during a hypo-osmotic challenge in vestibular dark cells. Biochim Biophys Acta 1240:48–54
- Shinozuka K, Tanaka N, Kawasaki K, Mizuno H, Kubota Y, Nakamura K, Hashimoto M, Kunitomo M (2001) Participation of ATP in cell volume regulation in the endothelium after hypotonic stress. Clin Exp Pharmacol Physiol 28:799–803
- Shrode LD, Krump E, Grinstein S (1998) Activation of protein kinases upon volume changes: role in cellular homeostasis. Contrib Nephrol 123:79–93
- Shrode LD, Rubie EA, Woodgett JR, Grinstein S (1997) Cytosolic alkalinization increases stress-activated protein kinase/c-Jun NH2-terminal kinase (SAPK/JNK) activity and p38 mitogen-activated protein kinase activity by a calcium-independent mechanism. J Biol Chem 272:13653–13659
- Siebens AW, Spring KR (1989) A novel sorbitol transport mechanism in cultured renal papillary epithelial cells. Am J Physiol 257:F937–F946
- Sizeland PC, Chambers ST, Lever M, Bason LM, Robson RA (1993) Organic osmolytes in human and other mammalian kidneys. Kidney Int 43:448–453
- Skou JC (1957) The influence of some cations on an adenosine triphosphatase from peripheral nerves. Biochim Biophys Acta 23:394–401
- Sokabe M, Sachs F, Jing ZQ (1991) Quantitative video microscopy of patch clamped membranes stress, strain, capacitance, and stretch channel activation. Biophys J 59:722–728
- Soleimani M, Singh G, Bizal GL, Gullans SR, McAteer JA (1994) Na⁺/H⁺ exchanger isoforms NHE-2 and NHE-1 in inner medullary collecting duct cells. Expression, functional localization, and differential regulation. J Biol Chem 269:27973–27978
- Staines HM, Godfrey EM, Lapaix F, Egee S, Thomas S, Ellory JC (2002) Two functionally distinct organic osmolyte pathways in Plasmodium gallinaceum-infected chicken red blood cells. Biochim Biophys Acta 1561:98–108
- Stegen C, Matskevich I, Wagner CA, Paulmichl M, Lang F, Broer S (2000) Swelling-induced taurine release without chloride channel activity in Xenopus laevis oocytes expressing anion channels and transporters. Biochim Biophys Acta 1467:91–100
- Steidl M, Ritter M, Lang F (1991) Regulation of potassium conductance by prostaglandins in cultured renal epitheloid (Madin-Darby canine kidney) cells. Pflugers Arch 418:431–436
- Stephens L, Anderson K, Stokoe D, Erdjument-Bromage H, Painter GF, Holmes AB, Gaffney PR, Reese CB, McCormick F, Tempst P, Coadwell J, Hawkins PT (1998) Protein kinase B kinases that mediate phosphatidylinositol 3,4,5-trisphosphate-dependent activation of protein kinase B. Science 279:710–714
- Stoddard JS, Steinbach JH, Simchowitz L (1993) Whole cell Cl⁻ currents in human neutrophils induced by cell swelling. Am J Physiol 265:C156–C165
- Stoner LC, Morley GE (1995) Effect of basolateral or apical hypo-osmolarity on apical maxi K channels of everted rat collecting tubule. Am J Physiol 268:F569–F580
- Strange K (1992) Regulation of solute and water balance and cell volume in the central nervous system. J Am Soc Nephrol 3:12–27
- Strange K (1998) Molecular identity of the outwardly rectifying, swelling-activated anion channel: Time to reevaluate pICln. J Gen Physiol 111:617–622
- Strange K, Emma F, Jackson PS (1996) Cellular and molecular physiology of volume-sensitive anion channels. Am J Physiol 270:C711–C730
- Strange K, Morrison R, Heilig CW, DiPietro S, Gullans SR (1991) Upregulation of inositol transport mediates inositol accumulation in hyperosmolar brain cells. Am J Physiol 260:C784–C790
- Strotmann R, Harteneck C, Nunnenmacher K, Schultz G, Plant TD (2000) OTRPC4 a nonselective cation channel that confers sensitivity to extracellular osmolarity. Nat Cell Biol 2:695–702
- Stühmer W, Ruppersberg JP, Schroter KH, Sakmann B, Stocker M, Giese KP, Perschke A, Baumann A, Pongs O (1989) Molecular basis of functional diversity of voltage-gated potassium channels in mammalian brain. EMBO J 8:3235–3244

- Stutzin A, Torres R, Oporto M, Pacheco P, Eguiguren AL, Cid LP, Sepulveda FV (1999) Separate taurine and chloride efflux pathways activated during regulatory volume decrease. Am J Physiol 277:C392– C402
- Su W, Shmukler BE, Chernova MN, Stuart-Tilley AK, De Franceschi L, Brugnara C, Alper SL (1999) Mouse K-Cl cotransporter KCC1: cloning, mapping, pathological expression, and functional regulation. Am J Physiol 277:C899–C912
- Sun AM, Liu Y, Dworkin LD, Tse CM, Donowitz M, Yip KP (1997) Na⁺/H⁺ exchanger isoform 2 (NHE2) is expressed in the apical membrane of the medullary thick ascending limb. J Membr Biol 160:85–90
- Suzuki M, Kawahara K, Ogawa A, Morita T, Kawaguchi Y, Kurihara S, Sakai O (1990) [Ca²⁺]_i rises via G protein during regulatory volume decrease in rabbit proximal tubule cells. Am J Physiol 258:F690–F696
- Suzuki M, Sato J, Kutsuwada K, Ooki G, Imai M (1999) Cloning of a stretch-inhibitable nonselective cation channel. J Biol Chem 274:6330–6335
- Szaszi K, Grinstein S, Orlowski J, Kapus A (2000) Regulation of the epithelial Na⁺ /H⁺ exchanger isoform by the cytoskeleton. Cell Physiol Biochem 10:265–272
- Szászi K, Kurashima K, Kapus A, Paulsen A, Kaibuchi K, Grinstein S, Orlowski J (2000) RhoA and rho kinase regulate the epithelial Na⁺/H⁺ exchanger NHE3 Role of myosin light chain phosphorylation. J Biol Chem 275:28599–28606
- Taglietti V, Toselli M (1988) A study of stretch-activated channels in the membrane of frog oocytes: Interaction with Ca²⁺ ions. J Physiol (Lond) 407:311–328
- Tai KK, Wang KW, Goldstein SA, N (1997) MinK potassium channels are heteromultimeric complexes. J Biol Chem 272:1654–1658
- Takenaka M, Bagnasco SM, Preston AS, Uchida S, Yamauchi A, Kwon HM, Handler JS (1995) The canine betaine gamma-amino-n-butyric acid transporter gene: diverse mRNA isoforms are regulated by hypertonicity and are expressed in a tissue-specific manner. Proc Natl Acad Sci USA 92:1072–1076
- Takumi T, Ohkubo H, Nakanishi S (1989) Cloning of a membrane protein that induces a slow voltage-gated potassium current. Science 242:1042–1045
- Tamkun MM, Knoth KM, Walbridge JA, Kroemer H, Roden DM, Glover DM (1991) Molecular cloning and characterization of two voltage-gated K⁺ channel cDNAs from human ventricle. FASEB J 5:331–337
- Taniguchi J, Guggino WB (1989) Membrane stretch: A physiological stimulator of Ca²⁺-activated K⁺ channels in thick ascending limb. Am J Physiol 257:F347–F352
- Terada Y, Inoshita S, Hanada S, Shimamura H, Kuwahara M, Ogawa W, Kasuga M, Sasaki S, Marumo F (2001) Hyperosmolality activates Akt and regulates apoptosis in renal tubular cells. Kidney Int 60:553–567
- Tewari KP, Malinowska DH, Sherry AM, Cuppoletti J (2000) PKA and arachidonic acid activation of human recombinant ClC-2 chloride channels. Am J Physiol Cell Physiol 279:C40–C50
- Theodoropoulos PA, Stournaras C, Stoll B, Markogiannakis E, Lang F, Gravanis A, Häussinger D (1992) Hepatocyte swelling leads to rapid decrease of the G-/total actin ratio and increases actin mRNA levels. FEBS Lett 311:241–245
- Therien AG, Blostein R (2000) Mechanisms of sodium pump regulation. Am J Physiol Cell Physiol 279:C541–C566
- Thinnes FP, Walter G, Hellmann KP, Hellmann T, Merker R, Kiafard Z, Eben-Brunnen J, Schwarzer C, Gotz H, Hilschmann N (2001) Gadolinium as an opener of the outwardly rectifying Cl⁻ channel (ORCC) Is there relevance for cystic fibrosis therapy? Pflugers Arch 443:S111-S116
- Thoroed SM, Lauritzen L, Lambert IH, Hansen HS, Hoffmann EK (1997) Cell swelling activates phospholipase A2 in Ehrlich ascites tumor cells. J Membr Biol 160:47–58
- Tilly BC, van den Berghe N, Tertoolen LG, Edixhoven MJ, de Jonge HR (1993) Protein tyrosine phosphorylation is involved in osmoregulation of ionic conductances. J Biol Chem 268:19919–19922
- Tinel H, Petersen OH (2000) Transport across intracellular organelles. In Seldin DW, Giebisch G (eds) The kidney. Physiology and pathophysiology. Lippincott Williams and Wilkins, Philadelphia, pp 294–319
- Tinel H, Wehner F, Sauer H (1994) Intracellular Ca²⁺ release and Ca²⁺ influx during regulatory volume decrease in IMCD cells. Am J Physiol 267:F130–F138
- Tinel H, Wehner F, Kinne RKH (1997) Arachidonic acid as a second messenger for hypotonicity-induced calcium transients in rat IMCD cells. Pflugers Arch 433:245–253
- Tinel H, Cancela JM, Mogami H, Gerasimenko JV, Gerasimenko OV, Tepikin AV, Petersen OH (1999) Active mitochondria surrounding the pancreatic acinar granule region prevent spreading of inositol trisphosphate-evoked local cytosolic Ca²⁺ signals. EMBO J 18:4999–5008
- Tinel H, Kinne-Saffran E, Kinne RKH (2000) Calcium signalling during RVD of kidney cells. Cell Physiol Biochem 10:297–302

- Tinel H, Kinne-Saffran E, Kinne RKH (2002) Calcium-induced calcium release participates in cell volume regulation of rabbit TALH cells. Pflugers Arch 443:754–761
- Tominaga T, Barber DL (1998) Na-H exchange acts downstream of RhoA to regulate integrin-induced cell adhesion and spreading. Mol Biol Cell 9:2287–2303
- Tominaga T, Ishizaki T, Narumiya S, Barber DL (1998) p160ROCK mediates RhoA activation of Na-H exchange. EMBO J 17:4712–4722
- Toro L, Wallner M, Meera P, Tanaka Y (1998) Maxi-K_{Ca} a unique member of the voltage-gated K channel superfamily. News Physiol Sci 13:112–117
- Toyomoto T, Knutsen D, Soos G, Sato K (1997) Na-K-2Cl cotransporters are present and regulated in simian eccrine clear cells. Am J Physiol 273:R270–R277
- Tse C-M, Levine SA, Yun CHC, Brant SR, Nath S, Pouysségur J, Donowitz M (1994) Molecular properties, kinetics and regulation of mammalian Na⁺/H⁺ exchangers. Cell Physiol Biochem 4:282–300
- Tsuganezawa H, Kobayashi K, Iyori M, Araki T, Koizumi A, Watanabe S, Kaneko A, Fukao T, Monkawa T, Yoshida T, Kim DK, Kanai Y, Endou H, Hayashi M, Saruta T (2001) A new member of the HCO₃⁻ transporter superfamily is an apical anion exchanger of α-intercalated cells in the kidney. J Biol Chem 276:8180–8189
- Tsumura T, Oiki S, Ueda S, Okuma M, Okada Y (1996) Sensitivity of volume-sensitive Cl⁻ conductance in human epithelial cells to extracellular nucleotides. Am J Physiol 271:C1872–C1878
- Ubl J, Murer H, Kolb HA (1988) Hypotonic shock evokes opening of Ca²⁺-activated K channels in opossum kidney cells. Pflugers Arch 412:551–553
- Uchida S, Nakanishi T, Kwon HM, Preston AS, Handler JS (1991) Taurine behaves as an osmolyte in Madin-Darby canine kidney cells. Protection by polarized, regulated transport of taurine. J Clin Invest 88:656–662
- Uchida S, Kwon HM, Yamauchi A, Preston AS, Marumo F, Handler JS (1992) Molecular cloning of the cDNA for an MDCK cell Na⁺- and Cl⁻-dependent taurine transporter that is regulated by hypertonicity. Proc Natl Acad Sci USA 89:8230–8234
- Urbach V, Leguen I, O'Kelly I, Harvey BJ (1999) Mechanosensitive calcium entry and mobilization in renal A6 cells. J Membr Biol 168:29–37
- Valverde MA, Díaz M, Sepúlveda FV, Gill DR, Hyde SC, Higgins CF (1992) Volume-regulated chloride channels associated with the human multidrug-resistance P-glycoprotein. Nature 355:830–833
- Van der Kaay J, Beck M, Gray A, Downes CP (1999) Distinct phosphatidylinositol 3-kinase lipid products accumulate upon oxidative and osmotic stress and lead to different cellular responses. J Biol Chem 274:35963–35968
- van Der Wijk T, Dorrestijn J, Narumiya S, Maassen JA, de Jonge HR, Tilly BC (1998) Osmotic swellinginduced activation of the extracellular-signal-regulated protein kinases Erk-1 and Erk-2 in intestine 407 cells involves the Ras/Raf-signalling pathway. Biochem J 331:863–869
- van Der Wijk T, Tomassen SF, de Jonge HR, Tilly BC (2000) Signalling mechanisms involved in volume regulation of intestinal epithelial cells. Cell Physiol Biochem 10:289–296
- van Driessche W, Erlij D (1994) Cell swelling activates a poorly selective monovalent cation channel in the apical membrane of toad urinary bladder. Pflugers Arch 428:1–8
- van Driessche W, De Smet P, De Smedt H (1994) Poorly selective cation channels in the apical membrane of A6 cells. Pflugers Arch 426:387–395
- Vandorpe DH, Shmukler BE, Jiang L, Lim B, Maylie J, Adelman JP, De Franceschi L, Capellini MD, Brugnara C, Alper SL (2002) cDNA cloning and functional characterization of the moese Ca²⁺-gated K⁺ channel, mIK1. J Biol Chem 273:21542–21553
- Vanoye CG, Reuss L (1999) Stretch-activated single K⁺ channels account for whole-cell currents elicited by swelling. Proc Natl Acad Sci USA 96:6511–6516
- Verbalis JG (1994) Pathogenesis of hyponatremia in an experimental model of the syndrome of inappropriate antidiuresis. Am J Physiol 267:R1617–R1625
- Verbalis JG, Drutarosky MD, Ertel RJ, Vollmer RR (1989) Adaptive responses to sustained volume expansion in hyponatraemic rats. J Endocrinol 122:421–431
- Vergara C, Latorre R, Marrion NV, Adelman JP (1998) Calcium-activated potassium channels. Curr Opin Neurobiol 8:321–329
- Vexler ZS, Symons M, Barber DL (1996) Activation of Na⁺-H⁺ exchange is necessary for RhoA-induced stress fiber formation. J Biol Chem 271:22281–22284
- Viana F, de la PE, Pecson B, Schmidt RF, Belmonte C (2001) Swelling-activated calcium signalling in cultured mouse primary sensory neurons. Eur J Neurosci 13:722–734
- Vieira LL, Lafuente E, Blum J, Cabantchik ZI (1997) Modulation of the swelling-activated amino acid channel of Leishmania major promastigotes by protein kinases. Mol Biochem Parasitol 90:449–461

- Volk T, Frömter E, Korbmacher C (1995) Hypertonicity activates nonselective cation channels in mouse cortical collecting duct cells. Proc Natl Acad Sci USA 92:8478–8482
- Völkl H, Lang F (2001) Electrophysiology of betaine transport in isolated perfused straight proximal tubule. Pflugers Arch 442:136–140
- Vom Dahl S, Hallbrucker C, Lang F, Häussinger D (1991) Role of eicosanoids, inositol phosphates and extracellular Ca²⁺ in cell-volume regulation of rat liver. Eur J Biochem 198:73–83
- Vom Dahl S, Dombrowski F, Schmitt M, Schliess F, Pfeifer U, Häussinger D (2001) Cell hydration controls autophagosome formation in rat liver in a microtubule-dependent way downstream from p38MAPK activation. Biochem J 354:31–36
- von Weikersthal SF, Hickman ME, Hladky SB, Barrand MA (1997) Hypotonicity-induced changes in anion permeability of cultured rat brain endothelial cells. Biochim Biophys Acta 1325:99–107
- Wagner CA, Ott M, Klingel K, Beck S, Melzig J, Friedrich B, Wild KN, Bröer S, Moschen I, Albers A, Waldegger S, Tummler B, Egan ME, Geibel JP, Kandolf R, Lang F (2001) Effects of the serine/threonine kinase SGK1 on the epithelial Na⁺ channel (ENaC) and CFTR: implications for cystic fibrosis. Cell Physiol Biochem 11 209–218
- Walaas SI, Czernik AJ, Olstad OK, Sletten K, Walaas O (1994) Protein kinase C and cyclic AMP-dependent protein kinase phosphorylate phospholemman, an insulin and adrenaline- regulated membrane phosphoprotein, at specific sites in the carboxy terminal domain. Biochem J 304:635–640
- Waldegger S, Barth P, Raber G, Lang F (1997) Cloning and characterization of a putative human serine/ threonine protein kinase transcriptionally modified during anisotonic and isotonic alterations of cell volume. Proc Natl Acad Sci USA 94:4440–4445
- Wang K, Wondergem R (1991) Effects of hyperosmotic medium on hepatocyte volume, transmembrane potential and intracellular K⁺ activity. Biochim Biophys Acta 1069:187–196
- Wang Y, Roman R, Lidofsky SD, Fitz JG (1996) Autocrine signaling through ATP release represents a novel mechanism for cell volume regulation. Proc Natl Acad Sci USA 93:12020–12025
- Wangemann P, Liu J, Shen Z, Shipley A, Marcus DC (1995) Hypo-osmotic challenge stimulates transepithelial K⁺ secretion and activates apical IsK channel in vestibular dark cells. J Membr Biol 147:263–273
- Wärntges S, Friedrich B, Henke G, Duranton C, Lang PA, Waldegger S, Meyermann R, Kuhl D, Speckmann EJ, Obermüller N, Witzgall R, Mack AF, Wagner HJ, Wagner A, Bröer S, Lang F (2002) Cerebral localization and regulation of the cell volume-sensitive serum- and glucocorticoid-dependent kinase SGK1. Pflugers Arch 443:617–624
- Warskulat U, Wettstein M, Häussinger D (1997a) Osmo-regulated taurine transport in H4IIE hepatoma cells and perfused rat liver. Biochem J 321:83–690
- Warskulat U, Zhang F, Häussinger D (1997b) Taurine is an osmolyte in rat liver macrophages (Kupffer cells) J Hepatol 26:1340–1347
- Warth R, Hamm K, Bleich M, Kunzelmann E, Von Hahn T, Schreiber R, Ullrich E, Mengel M, Trautmann N, Kindle P, Schwab A, Greger R (1999) Molecular and functional characterization of the small Ca²⁺-regulated K⁺ channel (rSK4) of colonic crypts. Pflugers Arch 438:437–444
- Watson AJ, Levine S, Donowitz M, Montrose MH (1992) Serum regulates Na⁺/H⁺ exchange in Caco-2 cells by a mechanism which is dependent on F-actin. J Biol Chem 267:956–962
- Watson PA, Giger KE, Frankenfield CM (1991) Activation of adenylate cyclase during swelling of S49 cells in hypotonic medium is not involved in subsequent volume regulation. Mol Cell Biochem 104:51–56
- Weaver YR, Kiessling K, Cossins AR (1999) Responses of the Na⁺/H⁺ exchanger of European flounder red blood cells to hypertonic, β -adrenergic and acidotic stimuli. J Exp Biol 202:21–32
- Weekes J, Barry ST, Critchley DR (1996) Acidic phospholipids inhibit the intramolecular association between the N- and C-terminal regions of vinculin, exposing actin-binding and protein kinase C phosphorylation sites. Biochem J 314:827–832
- Wehner F (1998) Cell volume-regulated cation channels. Contrib Nephrol 123:8–20
- Wehner F, Tinel H (1998) Role of Na⁺ conductance, Na⁺-H⁺ exchange, and Na⁺- K⁺-2Cl⁻ symport in the regulatory volume increase of rat hepatocytes. J Physiol (Lond) 506:127–142
- Wehner F, Tinel H (2000) Osmolyte and Na⁺ transport balances of rat hepatocytes as a function of hypertonic stress. Pflugers Arch 441:12–24
- Wehner F, Sauer H, Kinne RKH (1995) Hypertonic stress increases the $\mathrm{Na^+}$ conductance of rat hepatocytes in primary culture. J Gen Physiol 105:507–535
- Wehner F, Tinel H, Kinne RKH (1997) Pharmacology of volume activated Na⁺ conductance in rat hepatocytes. Physiologist 40:A-4
- Wehner F, Lawonn P, Tinel H (2002a) Ionic mechanisms of regulatory volume increase (RVI) in the human hepatoma cell-line HepG2. Pflugers Arch 443:779–790

- Wehner F, Olsen H, Bierhals K, Lin C-T, Waldmann H (2002b) Signal transduction in the hypertonic activation of rat hepatocyte Na⁺ conductance. FASEB J A56
- Weik C, Warskulat U, Bode J, Peters-Regehr T, Häussinger D (1998) Compatible organic osmolytes in rat liver sinusoidal endothelial cells. Hepatology 27:569–575
- Weiss H, Lang F (1992) Ion channels activated by swelling of Madin Darby canine kidney (MDCK) cells. J Membr Biol 126:109–114
- Weyand B, Warth R, Bleich M, Kerstan D, Nitschke R, Greger R (1998) Hypertonic cell shrinkage reduces the K⁺ conductance of rat colonic crypts. Pflugers Arch 436:227–232
- Willumsen NJ, Davis CW, Boucher RC (1994) Selective response of human airway epithelia to luminal but not serosal solution hypertonicity. Possible role for proximal airway epithelia as an osmolality transducer. J Clin Invest 94:779–787
- Winpenny JP, Mathews CJ, Verdon B, Wardle CJ, C., Chambers JA, Harris A, Argent BE, Gray MA (1996) Volume-sensitive chloride currents in primary cultures of human fetal vas deferens epithelial cells. Pflugers Arch 432:644–654
- Wirthensohn G, Lefrank S, Schmolke M, Guder WG (1989) Regulation of organic osmolyte concentrations in tubules from rat renal inner medulla. Am J Physiol 256:F128–F135
- Wissenbach U, Bodding M, Freichel M, Flockerzi V (2000) Trp12 a novel Trp related protein from kidney. FEBS Lett 485:127–134
- Wittels KA, Hubert EM, Musch MW, Goldstein L (2000) Osmolyte channel regulation by ionic strength in skate RBC. Am J Physiol Regul Integr Comp Physiol 279:R69–R76
- Wojnowski L, Oberleithner H (1991) Hypertonicity in fused Madin-Darby canine kidney cells: transient rise in NaHCO₃ followed by sustained KCl accumulation. Pflugers Arch 419:43–50
- Wolff NA, Kinne R (1988) Taurine transport by rabbit kidney brush-border membranes: coupling to sodium, chloride, and the membrane potential. J Membr Biol 102:131–139
- Wright EM, Diamond JM (1977) Anion selectivity in biological systems. Physiol Rev 57:109-156
- Wu X, Yang H, Iserovich P, Fischbarg J, Reinach PS (1997) Regulatory volume decrease by SV40-transformed rabbit corneal epithelial cells requires ryanodine-sensitive Ca²⁺-induced Ca²⁺ release. J Membr Biol 158:127–136
- Wymann MP, Pirola L (1998) Structure and function of phosphoinositide 3-kinases. Biochim Biophys Acta 1436:127–150
- Xiong H, Li C, Garami E, Wang Y, Ramjeesingh M, Galley K, Bear CE (1999) ClC-2 activation modulates regulatory volume decrease. J Membr Biol 167:215–221
- Xu B, Lu L (1994) Protein kinase A-regulated Cl⁻ channel in ML-1 human hematopoietic myeloblasts. J Membr Biol 142:65–75
- Xu WX, Kim SJ, So I, Kang TM, Rhee JC, Kim KW (1997) Volume-sensitive chloride current activated by hypo-osmotic swelling in antral gastric myocytes of the guinea-pig. Pflugers Arch 435:9–19
- Yamaguchi DT, Green J, Kleeman CR, Muallem S (1989) Characterization of volume-sensitive, calciumpermeating pathways in the osteosarcoma cell line UMR-106-01. J Biol Chem 264:4383-4390
- Yamamoto Y, Suzuki H (1996) Two types of stretch-activated channel activities in guinea-pig gastric smooth muscle cells. Jpn J Physiol 46:337–345
- Yamashita T, Yamauchi A, Miyai A, Taniguchi M, Yoshimine T, Tohyama M (1999) Neuroprotective role of Na⁺/myo-inositol cotransporter against veratridine cytotoxicity. J Neurochem 72:1864–1870
- Yamauchi A, Kwon HM, Uchida S, Preston AS, Handler JS (1991) Myo-inositol and betaine transporters regulated by tonicity are basolateral in MDCK cells. Am J Physiol 261:F197–F202
- Yamauchi A, Uchida S, Kwon HM, Preston AS, Robey RB, Garcia-Perez A, Burg MB, Handler JS (1992) Cloning of a Na⁺- and Cl⁻-dependent betaine transporter that is regulated by hypertonicity. J Biol Chem 267:649–652
- Yamauchi A, Uchida S, Preston AS, Kwon HM, Handler JS (1993) Hypertonicity stimulates transcription of gene for Na⁺-myo-inositol cotransporter in MDCK cells. Am J Physiol 264:F20–F23
- Yamauchi A, Miyai A, Shimada S, Minami Y, Tohyama M, Imai E, Kamada T, Ueda N (1995) Localization and rapid regulation of Na[†]/myo-inositol cotransporter in rat kidney. J Clin Invest 96:1195–1201
- Yancey PH, Burg MB (1989) Distribution of major organic osmolytes in rabbit kidneys in diuresis and antidiuresis. Am J Physiol 257:F602–F607
- Yancey PH, Clark ME, Hand SC, Bowlus RD, Somero GN (1982) Living with water stress: evolution of osmolyte systems. Science 217:1214–1222
- Yang X-C, Sachs F (1989) Block of stretch-activated ion channels in Xenopus oocytes by gadolinium and calcium ions. Science 243:1068–1071
- Yip KP (2002) Coupling of vasopressin-induced intracellular Ca²⁺ mobilization and apical exocytosis in perfused rat kidney collecting duct. J Physiol 538:891–899

- Yordy MR, Bowen JW (1993) Na,K-ATPase expression and cell volume during hypertonic stress in human renal cells. Kidney Int 43:940–948
- Yorek MA, Dunlap JA, Lowe WL, Jr (1999) Osmotic regulation of the Na⁺/myo-inositol cotransporter and postinduction normalization. Kidney Int 55:215–224
- Yu W-G, Sokabe M (1997) Hypotonically induced whole-cell currents in A6 cells: relationship with cell volume and cytoplasmic Ca²⁺. Jpn J Physiol 47:553–565
- Zablocki K, Miller SP, Garcia-Perez A, Burg MB (1991) Accumulation of glycerophosphocholine (GPC) by renal cells: osmotic regulation of GPC:choline phosphodiesterase. Proc Natl Acad Sci USA 88:7820–7824
- Zelikovic I, Stejskal-Lorenz E, Lohstroh P, Budreau A, Chesney RW (1989) Anion dependence of taurine transport by rat renal brush-border membrane vesicles. Am J Physiol 256:F646–F655
- Zelikovic I, Budreau-Patters A (1999) Cl⁻ and membrane potential dependence of amino acid transport across the rat renal brush border membrane. Mol Genet Metab 67:236–247
- Zhang JJ, Jacob TJ, C (1997) Three different Cl⁻ channels in the bovine ciliary epithelium activated by hypotonic stress. J Physiol (Lond) 499:379–389
- Zhang MI, O'Neil RG (1996) An L-type calcium channel in renal epithelial cells. J Membr Biol 154:259– 266
- Zhang Z, Cohen DM (1996) NaCl but not urea activates p38 and jun kinase in mIMCD3 murine inner medullary cells. Am J Physiol 271:F1234–F1238
- Zhou C, Agarwal N, Cammarata PR (1994) Osmoregulatory alterations in myo-inositol uptake by bovine lens epithelial cells. Part 2: Cloning of a 626 bp cDNA portion of a Na⁺/myo-inositol cotransporter, an osmotic shock protein. Invest Ophthalmol Vis Sci 35:1236–1242
- Zhou C, Cammarata PR (1999) Characterization of the minimal osmotic response element of the bovine Na⁺/myo-inositol cotransporter gene. Exp Eye Res 68:137–142
- Zhuang S, Hirai SI, Ohno S (2000) Hyperosmolality induces activation of cPKC and nPKC, a requirement for ERK1/2 activation in NIH/3T3 cells. Am J Physiol Cell Physiol 278:C102–C109