Activation of Akt, Not Connexin 43 Protein Ubiquitination, Regulates Gap Junction Stability*

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The pore-forming gap junctional protein connexin 43 (Cx43) has a short (1–3 h) half-life, and both the proteasome and lysosome have been implicated in Cx43 turnover.

Results: Use of lysine to arginine mutants and kinase inhibitors showed that Akt activation stabilizes gap junctions.

Conclusion: Akt phosphorylation of Cx43 stabilizes gap junctions.

Significance: Linkage of Akt to gap junction stability may mechanistically explain their cell survival/growth roles.

The passage of ions and small metabolites between adjacent cells occurs via gap junction-mediated intercellular communication (1–3). Vertebrate gap junctions, composed of integral membrane proteins from the connexin gene family, are critically important in regulating development, coordinated contraction of excitable cells, tissue homeostasis, and controlled cell growth and differentiation (2, 3). Connexin mutations have been linked to several diseases (4–6) including oculodentodigital dysplasia, a disease caused by connexin 43 (Cx43)™ mutations that can lead to craniofacial abnormalities, atrioseptal defects, and arrhythmias (7). Cx43, the most ubiquitous connexin, oligomerizes into a hexamer or “connexon” in the trans-Golgi network (8) and, after transport to the plasma membrane and association with other connexons in neighboring cell membranes, can form a gap junction.

Cx43 is highly phosphorylated, and many of the modifications appear to increase gap junction formation whereas others inhibit formation or decrease gap junction channel open time (9–11). Given the unusually short half-life (1–3 h) of Cx43 in cell culture and tissues, Cx43 turnover also appears to be highly regulated (12–16). Cx43 degradation, mostly based on the use of inhibitors, has been reported to involve both the proteasome and lysosome (17–21). Several reports have shown that a ubiquitin ladder can be observed upon immunoprecipitation of Cx43 (17, 22–25). Furthermore, interaction of Cx43 with the ubiquitin ligase Nedd4 and the endocytic adapter protein Eps15 also indicates that Cx43 might be ubiquitinated and degraded via the proteasomal or endocytic pathway (22, 26). Reports indicate that endoplasmic reticulum-associated degradation is involved in the turnover of misfolded Cx43 and is sensitive to heat shock and cytotoxic stress (27). However, a recent report using a mutant version of Cx43 with all lysines converted to arginines indicated that this process does not involve ubiquitination of Cx43 (28). Several other observations are not consistent with polyubiquitination of Cx43 leading to proteasomal degradation. Treatment with proteasomal inhibitors often leads to increased levels of Cx43 within gap junctions (i.e. they become larger) with increased phosphorylation, but little change in total Cx43, whereas treatment with lysosomal inhibitors leads to increased levels of the protein (≥95% of cell surface Cx43 was retained for up to 6 h after lysosomal inhibition) (29). A clear polyubiquitin ladder co-labeled with Cx43 and ubiquitin antibodies has not been shown nor has a specific lysine acceptor for ubiquitin been identified in Cx43 that when

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2 The abbreviations used are: Cx43, connexin 43; MDCK, Madin-Darby canine kidney; NRK, normal rat kidney; TPA, 12-O-tetradecanoylphorbol-13-acetate.

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mutated to arginine prevents ubiquitination and subsequent internalization and degradation.

Our laboratories have spent significant time looking for Cx43 ubiquitination and the possible lysine targets for this process. Among several other mutations, we created a construct representing full-length Cx43 with all of the lysines converted to arginines that maintains the same net charge but that could not be ubiquitinated at lysine residues. When expressed in cells that did not express wild-type Cx43, this mutant version trafficked to the plasma membrane, formed gap junctions, and responded to proteasomal inhibitors in a manner similar to wild-type Cx43, i.e. junctions became larger in immunofluorescence studies, and slower migrating Cx43 was observed in immunoblots, essentially demonstrating that direct Cx43 ubiquitination was not necessary to observe the effects of proteasomal inhibition on gap junction size. We then turned our search to other proteins that might be regulated by ubiquitination that could in turn regulate Cx43 localization within the plasma membrane.

We found that Akt (protein kinase B) is the most likely candidate for the following reasons: Akt becomes ubiquitinated and phosphorylated (activated) to translocate to the plasma membrane and phosphorylate membrane proteins (30). Proteasomal inhibition led to increased phosphorylation of Akt substrates including Cx43. Inhibition of Akt with specific Akt inhibitors or with a dominant negative version of Akt (either of which dramatically reduce Akt activity) resulted in loss of the proteasomal inhibitor effect, i.e. junctions remained smaller, and less phosphorylated Cx43 was observed. Our data support a model where ubiquitination of Akt leads to increased Akt activity and direct phosphorylation of Cx43, resulting in increased junctional size.

EXPERIMENTAL PROCEDURES

Antibodies and Other Reagents—All general chemicals, unless otherwise noted, were purchased from Fisher Scientific. 12-O-tetradecanoylphorbol-13-acetate (TPA, 75 nm final concentration) was purchased from Sigma Co. Akt VIII and MG132 were purchased from EMD Biosciences (Gibbstown, NJ) and both used at 10 μM. Mouse anti-Cx43 antibodies, Cx43CT1 and Cx43IF1, prepared against amino acids 360–382 of Cx43 and antibody Cx43NT1 against amino acids 1–20 (Cx43NT1) of Cx43 (described in Refs. 31–33) were made at the Fred Hutchinson Cancer Research Center Hybridoma Development Facility (Seattle, WA). Anti-hemagglutinin (HA)-tagged monoclonal antibody was from Covance (Princeton, NJ). A rabbit anti-Cx43 (PNRF) was obtained from A. Boynton (34). Akt, phospho-Akt (Ser-473), and phospho-(Ser/Thr) Akt substrate antibodies were purchased from Cell Signaling (Danvers, MA) or Santa Cruz Biotechnology, Inc. (Santa Cruz, CA).

Cell Line Maintenance, Treatment, cDNA Transfection, and Microinjection—Madin-Darby canine kidney (MDCK) cells that do not express Cx43 (35) and normal rat kidney (NRK)-52E cells (ATTC) were cultured in DMEM (Mediatech, Pittsburgh PA) supplemented with 10% FCS and antibiotics (Invitrogen) except where indicated. Where MG132 treatment was followed by TPA incubation, cells were pretreated with MG132 for 30 min followed by TPA for 1 h. For experiments where cells were treated with Akt VIII followed by MG132, cells were pretreated with 10 μM Akt VIII inhibitor for 30 min followed by 10 μM MG132 for 3 h. Cells were microinjected with the gap junction-permeable dye Alexa Fluor 488 (Invitrogen) as described previously (37) and are reported as the number of recipient cells receiving dye in 3 min.

Immunoblotting—Cells were lysed in sample buffer containing 50 mM NaF, 500 μM Na3VO4, 2 mM PMSF, and 1× Complete Protease inhibitor (Roche Applied Science), and cellular proteins were separated by SDS-PAGE (10% polyacrylamide gel). After electrophoresis, protein was transferred to nitrocellulose, the membrane was blocked, and antibodies were incubated as indicated previously (32). IRDye800 donkey anti-mouse (Rockland Immunochemicals, Gilbertsville, PA) and Alexa Fluor 680 goat anti-rabbit (Invitrogen) secondary antibodies were used to visualize rabbit and mouse primary antibodies, respectively, simultaneously and were quantified directly using the Li-Cor Biosciences Odyssey infrared imaging system and associated software (Lincoln, NE). Statistical differences were calculated via a paired two-tailed t test.

Immunofluorescence—Cells were washed twice in PBS, and fixed in cold methanol/acetone (50:50) for 1 min followed by a 1-h block in 1% BSA in PBS. Cells were incubated with a mouse anti-Cx43 antibody (Cx43IF1) or rabbit anti-Cx43 in blocking solution for 1 h. Following several PBS washes, the cultures were incubated with Alexa Fluor 546-conjugated goat anti-rabbit antibody and/or Alexa Fluor 488-conjugated goat anti-mouse antibody for 30–60 min and counterstained with DAPI (Molecular Probes), followed by several washes in PBS. The coverslips were mounted onto slides with DABCO anti-fade medium (25 mg/ml of 1,4-diazobicyclo-(2,2,2)octane (Sigma) diluted in 90% glycerol and 10% PBS, pH 8.6) and viewed with a Zeiss LSM 510 laser scanning fluorescence microscope.

Immunoprecipitation—Anti-HA-tagged antibody inked to agarose (Syd Laboratories, Malden, MA) or the PNRF anti-Cx43 antibody was used in immunoprecipitation reactions as described previously (38). Briefly, cells were lysed in radioimmunoprecipitation assay buffer (0.5% deoxycholate, 0.5% Triton X-100, 100 mM NaCl, 10 mM EDTA, 50 mM NaF, 500 μM Na3VO4, 2 mM PMSF, and 1× Complete Protease inhibitor, 25 mM Tris-HCl, pH 7.2), precleared, incubated with the HA-antibody-coated agarose or PNRF and protein A beads, washed three times with cold radioimmunoprecipitation assay buffer, and eluted with sample buffer prior to SDS-PAGE and immunoblotting.
RESULTS

Several studies have shown that treatment of Cx43-expressing cells with proteasomal inhibitors increases the level of Cx43 present in gap junctions often without increasing the total level of Cx43 in the cells. On the other hand, treatment with lysosomal inhibitors leads to an increase in the P0 or fastest migrating form of Cx43. Results consistent with these observations are illustrated in Fig. 1 for NRK and MDCK cells expressing Cx43. NRK cells endogenously express Cx43 and have been a common cell line to examine Cx43 trafficking and gap junction assembly (e.g. 39). In these cells, treatment with the lysosomal degradation inhibitor ammonium chloride (NH₄Cl) led to a 33% statistically significant ($p < 0.001, n = 9$) increase in the fastest migrating Cx43 species compared with control in SDS-PAGE whereas the proteasomal inhibitor MG132 led to a 53% statistically significant increase ($p < 0.001, n = 9$) in the slower migrating isoforms at the expense of the faster migrating bands (Fig. 1A). The MDCK cells used in this study do not express endogenous Cx43 (35) (Fig. 1A, middle panel) and hence were transfected with a construct containing wild-type Cx43 (MDCK43). These cells showed similar responses to these drugs as the NRK cells with increases of 36 and 45% in the fastest migrating and slower migrating Cx43 species in response to NH₄Cl and MG132 treatment, respectively, compared with control ($p < 0.001, n = 9$).

Immunofluorescence studies to localize Cx43 in control, NH₄Cl-, and MG132-treated NRK (Fig. 1B, top) and MDCK43 (Fig. 1B, bottom) cells showed little change in junctional Cx43 upon NH₄Cl treatment and a readily apparent increase in punctate and continuous junctional staining in the presence of MG132.

We were interested in determining whether and at which lysine residue Cx43 is ubiquitinated and created several lysine to arginine mutants including a construct of Cx43 where all of the 27 lysine residues in Cx43 are converted to arginines. We stably expressed this K/R mutant version of Cx43 in MDCK cells (termed MDCK43K/R) and examined the protein localization and ability to dye transfer. When MDCK43K/R cells were examined in immunoblots after treatment with proteasomal and lysosomal inhibitors (Fig. 2A), we were surprised to find that the mutant Cx43 reacted very similarly to wild-type Cx43 (MDCK43 cells). Significant increases of 23 and 68% in the amount of the fastest migrating and slower migrating Cx43 species compared with control were observed in SDS-PAGE in response to NH₄Cl and MG132 treatment, respectively (both $p < 0.001, n = 9$). Similarly, Cx43 immunolocalization in MDCK43K/R cells appeared generally similar. Fig. 2B shows the expression of the K/R mutant version of Cx43 at apparent ER/Golgi and gap junctional membranes (CON) via immunofluorescence. NH₄Cl led to few overt changes whereas MG132 caused an obvious increase in the amount of junctional Cx43 (Fig. 2B, D). To examine whether gap junctions formed by the K/R mutant were functional, Alexa Fluor 488 dye transfer was measured 3 min after microinjection. MDCK43K/R cells transferred dye to 3.3 ± 1.7 (mean ± S.D.) cells in contrast to 6.0 ± 3.6 in MDCK cells expressing wild-type Cx43. Because there could be differences in Cx43 expression between the clones, we cannot calculate reliable differences in efficiency due to the mutations but can conclude that the mutated version is capable of forming functional channels.

Reagents that activate MAPK and PKC such as EGF and TPA have been reported to lead to increased turnover of gap junctions and increased Cx43 ubiquitination (24) and could possibly represent an alternative pathway for Cx43 ubiquitination and turnover. Therefore, we decided to see whether TPA treatment alone or in combination with proteasomal inhibition affected MDCK43 and MDCK43K/R cells differently. As expected, either TPA or MG132 treatment decreased the mobility of Cx43 in SDS-PAGE (Fig. 3A), although each reagent probably resulted in phosphorylation at different serines in Cx43 (24, 40). The mobility of Cx43 from TPA/MG132-treated cells did not appear to vary from MG132 treatment alone for either MDCK43 or MDCK43K/R cells. This “elimination” of the effect of TPA was more apparent in immunofluorescence studies. TPA treatment led to a near complete loss of gap junctional structures whereas a combined treatment of TPA and MG132 not only retained gap junctions but did not appear to vary from MG132 treatment alone (compare Fig. 3B, right column, with...
Because Cx43 in the MDCK43K/R cells could not be ubiquitinated, these results led us to question the relevance of Cx43 ubiquitination for gap junctional stability and whether proteasomal inhibition was actually affecting the activity of a different protein that regulated gap junctional localization of Cx43.

Given that Cx43 localization to junctions is regulated via phosphorylation, we tested the ability of a variety of inhibitors of kinases known to phosphorylate Cx43 to see whether they affected its localization during proteasomal inhibition. Akt has been previously shown to phosphorylate Cx43 (41), and an inhibitor of Akt named Akt VIII (42) completely eliminated the MG132-dependent shift in Cx43 migration found in SDS-PAGE of NRK, MDCK43, and MDCK43K/R cells (Fig. 4A). Furthermore, 3 h of Akt VIII treatment caused Cx43 to localize to cytoplasmic membranes and almost completely eliminated Cx43 in junctional structures (Fig. 4B).

Although Akt VIII is reported to be highly selective (42), we wanted to decrease Akt activity by an alternate method and chose to use a dominant negative Akt-K197A construct (36). When Akt-K197A was expressed in either NRK or MDCK43 cells, the cells no longer responded to MG132 with a shift to the slower migrating Cx43 species in SDS-PAGE compared with controls (Fig. 5A). Similar to Akt VIII treatment, Cx43 immunofluorescence in Akt-K197A-transfected MDCK43 cells treated with MG132 showed mostly cytoplasmic localization (Fig. 5C).

Ubiquitination of Akt has been reported to cause translocation of Akt to the plasma membrane and increase its activity via phosphorylation at Ser-473 (43, 44). We wanted to determine whether MG132 treatment led to increased Akt activity, phosphorylation/activation at Ser-473, increased Akt substrate phosphorylation in general, and increased Cx43 phosphorylation in our cellular systems, thereby making the mechanistic connection between drug treatment and effect. When MDCK43 cells were treated with Akt VIII or MG132, Akt phosphorylation at Ser-473 was dramatically decreased or increased, respectively, as detected with the phospho-Akt (Ser-473) phosphospecific antibody (Fig. 6A; Akt is ~60 kDa).

Akt functional activity (i.e., the ability to phosphorylate substrates at Akt consensus sequences) can be assayed via an antibody (45) that recognizes the Akt substrate consensus sequence K/RXK/RXXS/T where S/T is phosphorylated (named phospho-(Ser/Thr) Akt substrate antibody). Akt activity was 52% higher ($p = 0.004, n = 6$) upon MG132 treatment when densitometry is performed on all bands in an immunoblot of cell lysates probed with the phospho-(Ser/Thr) Akt substrate antibody (i.e., compared with control) (Fig. 6B). Akt VIII treatment showed a nonsignificant 10% decrease in Akt activity as defined
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Figure 4. Cx43 in NRK, MDCK43, and MDCK43K/R cells with or without lysosomal, proteasomal, and Akt activity inhibitors. A, representative immunoblot for Cx43 in cells treated with NH₄Cl, MG132, or Akt VIII inhibitor. Similar banding patterns were observed in nine independent experiments. Positions of prestained standards of the designated molecular mass in kDa are indicated on the left. B, immunofluorescence microscopy for Cx43 in cells treated with NH₄Cl, MG132, or Akt VIII inhibitor. Scale bar, 25 μm.

Akt could be isolated with HA-antibody-coated beads. In Fig. 6D, we show that MG132 treatment resulted in increased total HA-ubiquitin signal (left panel) and the appearance of a band that overlays exactly with a faint band observed when probed with an anti-Akt antibody in the input cell lysate (marked with an asterisk; this is the same lane blotted with the two different antibodies and detected in different optical channels). When the HA-bead immunoprecipitate was probed with the Akt antibody, MG132 increased the quantity of Akt immunoprecipitated whereas Akt VIII treatment decreased the amount. These results and the molecular size of the apparent ubiquitinated Akt band are essentially the same as reported by Adachi et al. (46).

Discussion

We sought to determine how Cx43 phosphorylation and ubiquitination were linked. However, after trying and failing to discover either sites of Cx43 ubiquitination or a functional need for ubiquitination in the cell types we analyzed, we hypothesized that another ubiquitinated protein that was degraded by the proteasome might in fact regulate gap junctional Cx43 retention and/or removal. We now believe that Akt is the protein that regulates Cx43 gap junctional retention for the following reasons: Akt membrane localization, activation, and phosphorylation of membrane protein substrates are regulated by Akt ubiquitination/activity abrogated the effect of the proteasomal inhibitor on Cx43 in gap junctions and Cx43 phosphorylation and dramatically reduced the amount of larger stable junctions. Cx43 phosphorylation at Akt sites was increased by proteasome to limit signaling, a process that is inhibited by blocking proteasomal degradation (46). Our Cx43 K/R mutant protein that contains no lysine acceptors for ubiquitination behaved very similarly to wild-type Cx43 in terms of the effects of TPA treatment and proteasomal inhibition, indicating that ubiquitination is not necessary for the enhanced degradation or stabilization of gap junctional Cx43 caused by these reagents, respectively. Moreover, Akt VIII treatment or transfection with a dominant negative Akt that dramatically reduced Akt ubiquitination/activity abrogated the effect of the proteasomal inhibitor on Cx43 in gap junctions and Cx43 phosphorylation and dramatically reduced the amount of larger stable junctions. Cx43 phosphorylation at Akt sites was increased by proteasomal inhibition and decreased by Akt VIII treatment. Akt has been shown previously to phosphorylate and potentially interact with Cx43 particularly at the edges of gap junction plaques (41). We hypothesize that Akt-dependent Cx43 phosphorylation could result in gap junction stabilization, and we are currently testing this linkage.

Cx43 has a very short half-life (1–3 h) compared with most integral membrane proteins. Many factors apparently influence gap junction assembly and degradation, including cell-cell adhesion and protein kinase activation. For example, protein kinase A (47, 48) and casein kinase 1 (31) activation have been shown to increase the amount of Cx43 within gap junctions whereas MAPK (49) activity decreases junctional Cx43. Treatment of cells with proteasomal inhibitors increases the level of phosphorylated Cx43 and the amount of Cx43 within gap junctions (50, 51). Early results with inhibitors led to speculation that the proteasome might actively degrade ubiquitinated Cx43 that had been in the membrane (17). Because subsequent results more clearly implicated the lysosome in Cx43 degradation (18, 51, 52), Cx43 monoubiquitination has been proposed...
that Cx43 ubiquitination cannot occur, particularly in cells that overexpress components of the ubiquitin conjugation pathway, rather that it is not necessary for the regulation of Cx43 gap junctional stability that is affected negatively and positively by TPA or proteasomal inhibitor treatment, respectively. In addition, we do not discount the role that ubiquitin modification may have in the interactions of Nedd4, Hrs, and Tsg101 with Cx43 (20, 22, 26) because these proteins have also been reported to affect Akt regulation (53–55). Importantly, in our cellular systems, we could not find Cx43 ubiquitination despite significant efforts. On the other hand, Akt ubiquitination and activation, as had been shown previously in other cellular systems (46), was readily apparent and changed in the manner expected for gap junctional stabilization. We did observe an ~50-kDa band resulting from Cx43 phosphorylation that could explain a similar band observed in other studies in the range for ubiquitinated Cx43. We found that Akt and Cx43 could co-immunoprecipitate (data not shown), also possibly explaining ubiquitin-positive bands.

We have shown that Akt activity is necessary for MG132-dependent stabilization of gap junctions using two molecularly distinct methods for reducing Akt activity. Thus, although Cx43 ubiquitination is not necessary, Akt activity is required for gap junction stabilization induced by proteasomal inhibition. We conclude that inhibition of proteasomal degradation leads to ubiquitination and phosphorylation/activation of Akt, direct Akt phosphorylation of Cx43 and other proteins at Akt substrate sites, and the stabilization of gap junction structures in the plasma membrane. We hypothesize that some of the proposed roles for gap junctional communication may be partially explained by the PI3K/Akt/mTOR signaling pathways because gap junctional communication and Akt both play roles in the control of apoptosis suppression, inhibition of stress-regulated kinase cascades, and cell cycle regulation (56). This interplay between a proto-oncogene and a tumor suppressor protein has some interesting biological implications and could help explain why gap junctions appear to act as tumor suppressors and yet can promote growth under other conditions (57).

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