Analysis of Connexin43 phosphorylated at S325, S328 and S330 in normoxic and ischemic heart

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Summary

The functional consequences of Connexin43 (Cx43) phosphorylation remain largely unexplored. Using an antibody that specifically recognizes Cx43 phosphorylated at serine residues 325, 328 and/or 330 (pS325/328/330-Cx43), we show that labeling of this form of Cx43 as well as of total Cx43 is restricted to the intercalated disk region of normal ventricular tissue. In ischemic heart, significant relocation of total Cx43 to the lateral edges of myocytes was evident; however pS325/328/330-Cx43 remained predominately at the intercalated disk. Western blots indicated a eightfold decrease in pS325/328/330-Cx43 in ischemic tissue. Peptide-binding- and competition-experiments indicated that our antibody mainly detected Cx43 phosphorylated at S328 and/or S330 in heart tissue. To evaluate how this change in Cx43 phosphorylation contributes to ischemia-induced downregulation of intercellular communication, we stably transfected Cx43−/− cells with a Cx43 construct in which serine residues 325, 328 and 330 had been mutated to alanine (Cx43-TM). Cx43-TM was not efficiently processed to isoforms that have been correlated with gap junction assembly. Nevertheless, Cx43-TM cells were electrically coupled, although development of coupling was delayed. Fully opened channels were only rarely observed in Cx43-TM cells, and Lucifer-Yellow-dye-coupling was significantly reduced compared with wild-type cells. These data suggest that phosphorylation of Cx43 at serine residues 325, 328 and/or 330 influences channel permselectivity and regulates the efficiency of gap junction assembly.

Key words: Connexin43, Gap junction, Heart, Ischemia, Phosphorylation

Introduction

Gap-junction-mediated intercellular communication facilitates direct communication among adjacent cells by allowing passage of ions and small metabolites (White and Paul, 1999; Saez et al., 2003; Sohl and Willecke, 2004). Vertebrate gap junctions, composed of integral membrane proteins from the Connexin gene family, are crucially important in regulating embryonic development, coordinated contraction of excitable cells, tissue homeostasis, normal cell growth and differentiation (Saez et al., 2003; Sohl and Willecke, 2004). Furthermore, connexin mutations have been linked to several diseases (Bergoffen et al., 1993; Gong et al., 1997; Kelsell et al., 1997) including oculodentodigital dysplasia, a disease linked to Connexin43 (Cx43) mutations that can cause atrioseptal defects and arrhythmias (Paznekas et al., 2003). Twenty-one connexin genes have been identified in humans (Sohl and Willecke, 2004). During intercellular channel formation, six connexin proteins oligomerize into a hemichannel or connexon; connexons are then transported to the plasma membrane by as yet unknown mechanisms. The intact channel is formed when one hemi-channel docks with a second in an opposing cell. Once assembled, groups of these intercellular channels (termed gap junctional plaques) mediate the diffusion of ions, amino acids, second messengers and other metabolites between the cytoplasmic spaces of the two cells (White and Paul, 1999; Sohl and Willecke, 2004). The channels can be gated in response to various stimuli, including changes in voltage, pH and connexin phosphorylation. Regulation of gap junctional communication could occur by controlling any one of the steps mentioned above, however, many of the regulatory mechanisms underlying these events remain elusive.

Cx43, the most ubiquitously expressed connexin, is differentially phosphorylated at a dozen or more serine residues throughout its life cycle (Lampe and Lau, 2004). Cx43 from cultured cells commonly demonstrates multiple electrophoretic isoforms when analyzed by SDS-PAGE: a fast-migrating form (sometimes referred to as P0 or NP) that includes the non-phosphorylated isoform, and multiple slow-migrating forms (sometimes termed P1 and P2) (Musil and Goodenough, 1991). Following alkaline phosphatase treatment, the phosphorylated species collapse to the fastest migrating form, suggesting that phosphorylation is the primary covalent modification detected in SDS-PAGE analysis, although no assignment of specific phosphorylation sites to a change in Cx43 migration has been made. Pulse-chase studies with Brefeldin A indicate some Cx43 phosphorylation occurs prior to reaching the plasma membrane (Laird et al., 1995). This phosphorylation event might be necessary for maintaining hemi-channels in their closed state until docking occurs (Bao et al., 2004). In addition, studies investigating phosphorylation in normal rat kidney (NRK) cells show that, Cx43 acquires resistance to Triton X-100 once it has been phosphorylated to...
the slower migrating isoforms and assembled into gap junction plaques (Musil and Goodenough, 1991). Thus, uncharacterized phosphorylation events have been correlated with changes in assembly, acquisition of Triton-X-100-insolubility and, potentially, degradation of Cx43 gap junction channels.

In the normally functioning ventricle, Cx43 is localized to intercalated disks where it supports the longitudinal and transverse spread of the action potential resulting in coordinated contractile activation. Myocardial ischemia leads to Cx43 dephosphorylation and loss of localization to the intercalated disk, which probably contributes to contractile failure and arrhythmias (Beardslee et al., 2000; Schulz et al., 2003). Casein kinase 1 (CK1) mediates phosphorylation of Cx43 at serine residues 325, 328 and/or 330 in vitro. In cultured cells these sites are routinely phosphorylated; inhibition of CK1 reduces phosphorylation at these sites and reduces gap junction assembly (Cooper and Lampe, 2002). We sought to determine here whether phosphorylation at these sites occurs in heart tissue and whether this phosphorylation event is affected during ischemia. Using an antibody specific for Cx43 when phosphorylated at serine residues 325, 328 and/or 330, we show that Cx43 localized to intercalated disks is phosphorylated at one or more of these residues (probably S328 and S330) and that ischemia leads to loss of this phosphorylation and relocalization of the protein. Furthermore, we show that mutation of these sites increases the migration of the phosphorylated Cx43 isoforms on SDS-PAGE, reduces the extent of gap junction formation and reduces gap junctional communication.

Results
Ischemia-induced dephosphorylation of Cx43
Previous studies have shown that the electrophoretic mobility of Cx43 isolated from ischemic heart is increased, such that most of the protein migrates at a position that includes dephosphorylated Cx43 so the assumption has been made that these changes in electrophoretic mobility represent a reduction in phosphorylation at undefined sites (Beardslee et al., 2000; Jain et al., 2003). In addition, inhibition of CK1 has been shown to reduce gap junction assembly and to phosphorylate Cx43 in vitro at one or more of serines 325/328/330 (Cooper and Lampe, 2002). We hypothesized that phosphorylation at sites necessary for efficient assembly of channels might be compromised in the ischemic setting. To evaluate whether these phosphorylation events occur in control tissue and ischemic heart tissue, we generated a polyclonal antibody that uniquely detects Cx43 phosphorylated at serine residues 325, 328 and/or 330 (pS325/328/330-Cx43) – the specificity of which is described in the next section.

In agreement with previous studies, the electrophoretic mobility of total Cx43 from ischemic heart was increased compared with control tissue, possibly indicating that unspecified dephosphorylation event(s) had occurred (Fig. 1, each lane represents a distinct heart). When detected with the phosphorylation-specific antibody, we found that the pS325/328/330-Cx43 content decreased to about 12% of control (P<0.03) in ischemic hearts (to 19% of control when the loss of total Cx43 was taken into account).

The cellular distribution of Cx43 detected with the phosphorylation-specific antibody and one against total Cx43 varied in ischemic vs normal tissue. Fig. 2 shows that in normal heart virtually all Cx43 and pS325/328/330-Cx43 were localized to intercalated disks. As shown previously (Severs et al., 2004), ischemia caused a redistribution of total Cx43 from the intercalated disk to the lateral borders of myocytes. The pS325/328/330-Cx43 signal was reduced at the intercalated disk of ischemic heart tissue, consistent with the reduction of total Cx43 but, surprisingly, pS325/328/330-Cx43 was undetectable at the lateral borders. If phosphorylation at these sites promotes gap junctional functionality, then transverse conduction in the ischemic heart would not be expected to increase (dramatically) despite increased Cx43 at lateral borders because of the reduced phosphorylation of residues S325, S328 and S330 and the consequent poor gap junction functionality.

Analysis of the specificity of the pS325/328/330-Cx43 antibody
To test the specificity of the pS325/328/330-Cx43 antibody and to determine whether phosphorylation at residues S325, S328 and S330 directly affects protein localization and gap junction formation, we transfected constructs encoding either wild-type Cx43 (WT) or Cx43 containing S325A, S328A and S330A mutations (Cx43-TM) into fibroblasts derived from Cx43<sup>−/−</sup> mice, and examined the properties of the gap junctions formed by these cells. We first validated the phosphorylation-specific nature of the antibody by examining its binding to extracts from cells expressing Cx43-WT or Cx43-TM both before and after treatment with alkaline phosphatase. As shown in Fig. 3A, the antibody staining against total Cx43 resulted in bands of 41-44 kDa in all cases and revealed the classic increase in mobility to the fastest migrating position upon alkaline phosphatase treatment. By contrast, pS325/328/330-Cx43-specific antibody bound to Cx43-WT only in the absence of alkaline phosphatase treatment and did not bind to the Cx43-TM at all. Furthermore, pS325/328/330-Cx43 antibody appeared to bind only to the slowest migrating isoform of Cx43; the Cx43-TM-expressing cells also did not efficiently convert Cx43 to the slow-migrating phosphorylated isoforms.

To examine the specificity of the pS325/328/330-Cx43

Fig. 1. Increased electrophoretic mobility of Cx43 in the ischemic heart involves dephosphorylation at serine residues 325, 328 and/or 330. Western blot of total protein isolated from three control (lanes 1,3,5, respectively) or three globally ischemic hearts (30’ at 37°C; lanes 2,4,6, respectively) simultaneously probed for total Cx43 (Cx43NT-1) and Cx43 phosphorylated at S325, S328 and/or S330 followed by GAPDH and vinculin antibodies. Notice the increased mobility of Cx43 and decreased pS325/328/330-Cx43 content in the ischemic samples. Molecular mass standards are 28, 49, 62 and 98 kDa (bottom to top); P, position of the Cx43 P<sup>0</sup> isoform; P<sub>0</sub>, position of the Cx43 P<sub>0</sub> isoform.
antibody and determine which sites are likely to be phosphorylated in vivo, we synthesized three peptides that were phosphorylated at one serine residue each, S325, S328 or S330 (pS325, pS328 or pS330, respectively), two peptides that were phosphorylated at two serine residues each, S325 and S328, as well as S328 and S330 (pS325/328, as well as pS328/330, respectively), and an immunogen peptide that was phosphorylated at all three serine residues (pS325/328/330) and covalently bound these peptides to an activated 96-well plate. The pS325/328/330-Cx43 antibody was incubated in the well and a standard enzyme-linked immunoabsorbent assay (ELISA) was performed (Fig. 3B, open bars). The pS328/330 peptide bound almost as much total antibody as the immunogen pS325/328/330 peptide, indicating that most of the antibody bound an epitope centered on these sites. The singly phosphorylated peptides bound only a small amount of antibody. To further answer the question of the specificity of this antibody preparation, we removed the antibody that bound to the pS328/330 peptide and purified the one specific for pS325/328. Analysis of the binding of this preparation to the six peptides in the ELISA format indicated that the pS325/328- purified antibody indeed mostly bound to the pS325/328 peptide, followed by the triply phosphorylated and the pS325 singly phosphorylated peptides (data not shown). These results indicate that the antibody preparation is phosphorylation-specific although polyclonal with reactivity to all three sites. To examine the ability of the six peptides to block antibody binding to Cx43, we preincubated these peptides with the pS325/328/330-Cx43 antibody prior to western blot analysis of heart lysates. Neither pS325 nor pS328 of the singly phosphorylated peptides interfered significantly with antibody binding to intact Cx43, leaving most of the signal as before, whereas the singly phosphorylated pS330 peptide and all of the more extensively phosphorylated peptides did reduce the signal on the western blot close to background levels (Fig. 3B, filled bars). Further, we tested the ability of the pS325/328-specific antibody and found that it could bind to Cx43 prepared from heart in an immunoblot – albeit much less extensively than the antibody with the pS328/330 reactivity (data not shown).
Combined, these antibody-binding and -competition results indicate that this polyclonal antibody can bind to Cx43 phosphorylated at residues S325, S328 and S330, and also that Cx43 is phosphorylated at S328 and/or S330, and to a lesser extent at S325 in heart tissue.

The role of phosphorylation at S325, S328 and S330 in gap junction assembly

To be more quantitative about the extent of gap junction formation, we assayed the ability of three separate Cx43-TM clones to phosphorylate Cx43 to the P2 isoform and incorporate it into a Triton X-100-insoluble fraction, a hallmark of conventional gap junctions (Musil and Goodenough, 1991). Like most cell lines, fibroblasts expressing wild-type Cx43 showed several isoforms of Cx43 in the whole-cell lysate and a dramatic enrichment of the P2 isoform in the Triton-insoluble fraction (Fig. 4, WT lanes).

Transfectants expressing Cx43 with the TM mutations (clones A-C) showed multiple differences in their Cx43 migration. In the whole-cell fraction, Cx43-TM migrated primarily at P0 with a second band between P0 and P1 and no apparent P2. There was a significant reduction in the total amount of Triton-insoluble Cx43 in clone B and C, and essentially no bands that co-migrated with P2 were observed in any of the clones. Variation in the levels of total Cx43 protein most probably represents clonal heterogeneity among individual Cx43-expressing clones. Since cells expressing Cx43-TM clone A showed similar levels of total Cx43 compared with cells expressing Cx43-WT, we used clone A for most of the subsequent studies. We conclude that phosphorylation of Cx43 at S325, S328 and/or S330 is necessary for Cx43 to migrate as the P2 isoform.

The localization of Cx43 in the clones that express Cx43-TM was also examined by immunofluorescence. In Fig. 5, we show that fibroblasts transfected with Cx43-WT showed both intracellular and punctate, gap junctional plaque-associated Cx43 [compare to the lack of signal in the knockout (KO) panel]. By contrast, the TM-mutant cell lines showed predominately intracellular Cx43, although occasional, apparently appositional, labeling was observed (TM-A clone is shown).

We next determined whether cells expressing Cx43-TM supported intercellular communication. Parental Cx43-knockout (KO) fibroblasts, Cx43-WT-expressing fibroblasts and the three individual Cx43-TM-expressing clones A-C were microinjected with Lucifer Yellow dye and evaluated for dye transfer 3 minutes later. Digital images were taken and the number of recipient cells was quantified. As shown in Fig. 6, the number of cells receiving dye from donor cells was 60-85% lower in the Cx43-TM clones than in Cx43-WT-expressing cells and was comparable to the coupling observed in the parental fibroblasts that lacked Cx43. Although Cx43 protein expression levels varied between the clones (see Fig. 4), all three Cx43-TM clones, including TM-A – the one that expressed similar levels of Cx43, showed significant reductions in communication compared with the wild-type control (P<0.02). These data suggest that phosphorylation at S325, S328 and/or S330 is crucial for efficient dye transfer in mouse fibroblasts.

We next evaluated whether junctional conductance and channel behavior differed as a consequence of the Ser to Ala mutations at serine residues 325, 328 and 330. Based on the dye-coupling data presented above, we expected junctional conductance to be significantly reduced in the Cx43-TM cell clones. This was indeed the case for Cx43-TM cells plated for

Fig. 4. Cx43-TM-expressing cells do not phosphorylate Cx43 to yield the P2 isoform and contain less Triton-insoluble (TI) Cx43. Whole-cell (WC) and TI cellular lysates from cells expressing wild-type Cx43 (WT) or three Cx43-TM clones (TM-A, TM-B, and TM-C) were western blotted and probed for total Cx43 with the Cx43NT1 antibody. A darker exposure of lysates of the TM-C clone is shown. Migration positions of a 50- and 36-kDa marker are indicated with an asterisk and the vinculin loading controls are indicated with a V. The Triton-insoluble vinculin could be associated with the cytoskeleton or adhesive junctions.

Fig. 5. Comparison of Cx43 localization in parental knockout (KO) cells, wild-type Cx43 (WT) cells and Cx43-TM (TM-A)-expressing cells. Apparent gap junctions are marked by arrowheads in the WT panel. Bar, 10 μm.

Fig. 6. Cells expressing Cx43 S325/328/330A are inefficient at dye transfer. Wild-type (WT)-, parental knockout (KO)- and Cx43-TM (clones TM-A, TM-B and TM-C)-expressing cells were microinjected with Lucifer Yellow dye. After 3 minutes of transfer, digital images were taken and the number of recipient cells was determined (n=the number of injected cells). Bars show the mean and error bars ± s.e.m. The extent of dye transfer was significantly different (**P<0.02) for WT compared with all of the other cell types.
Fig. 7. Histogram of channel events observed in Cx43-TM (gray) and Cx43-WT-expressing cells. (A) Event frequency as percent of total events in each bin. (B) Difference induced by mutation of serine residues 325, 328 and 330. Notice the reduced incidence of events corresponding to fully opened channels (106±1 pS) and increased incidence of 75±2 pS. The 54±1 pS population was not different from wild-type and mutant data sets. Difference data were fit using Origin software yielding Chi2 and R2 for fit are 5.677 and 0.94, respectively (~WT, 1851 events, n=13; TM, 418 events, n=7).

Discussion

Previous work has established that myocardial ischemia rapidly induces uncharacterized changes in the phosphorylation and localization of Cx43, however, no mechanistic connections between specific phosphorylation events and changes in conduction or localization have been identified. Here, we show that phosphorylation of Cx43 at S325, S328, and/or S330 was drastically reduced in the ischemic heart. Further, we demonstrate that Cx43 that localized to the lateral borders of myocytes was not phosphorylated at these sites, whereas Cx43 that remained at the intercalated disk retained these modifications. Finally, we show that, when these sites were not phosphorylated, the event frequency for fully opened channels, the overall junctional conductance and the extent of coupling to Lucifer Yellow dye were all significantly reduced.

Although direct Cx43 phosphorylation on serine residues has long been correlated with the formation and function of gap junctions under basal conditions (Musil and Goodenough, 1991), few mechanistic connections to specific phosphorylation events have been drawn. For instance, phosphorylation at S364 was shown to be important for cAMP induced upregulation of gap junction assembly. However, cells expressing Cx43 with a S364A mutation assembled Cx43 into junctions almost as well as cells expressing wild-type Cx43, suggesting S364 is not necessary for assembly in homeostatic cells (TenBroek et al., 2001). Similarly, several serines are known to be involved in mitogen-induced downregulation of communication, including serine residues 255, 279 and 282 (MAPK) and serine residues 262 and 368 (protein kinase C), but these phosphorylation events have been reported to negatively affect channel-gating properties (Lampe et al., 2000; Cottrell et al., 2003). S325, S328 and/or S330 were previously shown to be substrates for CK1 phosphorylation, a kinase implicated in the regulation of gap junction assembly (Cooper and Lampe, 2002). Our phosphorylation-specific antibody could react with Cx43 phosphorylated at all three sites in heart tissue. We probed heart lysates with seven other Cx43 phosphorylation-specific antibodies, that we had either created or are available from commercial sources, and that we find to be specific for phosphorylation at S255, S262, S279-S282, S364, S368 or S372. S325/328/330-Cx43 antibody was the only one that gave a significant decrease in phosphorylation (results not shown). The antibody against pS368 showed increased binding upon hypoxia that we have investigated separately (Ek-Vitorin et al., 2006). Similar to a previous report (Jeyaraman et al., 2003), we found (data not shown) that ischemia led to increased binding of an anti-Cx43 antibody we prepared against a nonphosphorylated peptide corresponding to residues 360-382 of Cx43 that appears to be identical to Zymed 13800, an antibody prepared to a similar epitope that has been reported to bind dephosphorylated Cx43 (Nagy et al., 1997).

We probed the functional consequences of these phosphorylation events by expressing Cx43 with Ser to Ala conversions at all three sites (i.e. the Cx43-TM mutant). This mutant, rather than double- or single-site mutants, was selected for study for two reasons: First, all three sites appear to be phosphorylated because phosphorylation is sequential with the incorporated serines and the remaining sites are not phosphorylated. Second, the functional consequences of phosphorylation at clusters of (phosphorylated) serines can be misinterpreted from single-site mutants because even the remaining sites partially or fully compensate for the missing site or the remaining sites are not phosphorylated, thereby diminishing the ability of these sites to compensate for the missing site or the remaining sites.

We found that a S328A/S330A mutant in Cx43 was not phosphorylated, and this mutant displayed gap junctional conductance and the extent of coupling to Lucifer Yellow dye were all significantly reduced. This mutant, rather than double- or single-site mutants, was selected for study for two reasons: First, all three sites appear to be phosphorylated because phosphorylation is sequential with the incorporated serines and the remaining sites are not phosphorylated. Second, the functional consequences of phosphorylation at clusters of (phosphorylated) serines can be misinterpreted from single-site mutants because even the remaining sites partially or fully compensate for the missing site or the remaining sites are not phosphorylated, thereby diminishing the ability of these sites to compensate for the missing site or the remaining sites.

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24-30 hours after plating. This coupling was mediated by channels with conductances of 70-90 pS, a value intermediate to the fully opened channel (~110 pS) and the previously reported, phosphorylation-induced 50-60–pS substrate (Lampe et al., 2000). These results are consistent with at least two explanations. First, phosphorylation at residues S325, S328 and/or S330 is necessary for Cx43 channels to open to their fully opened state. If correct, the intermediate conductance substate of Cx43 observed in the Cx43-TM cells would have to be poorly permeated by Lucifer Yellow to accommodate the low levels of dye-coupling observed for these cells. Interestingly, this intermediate conductance state is commonly observed in rat insulinoma cells expressing Cx43-WT, which normally do not express Cx45 (Ek-Vitorin et al., 2006), and the junctions formed by these cells are poorly permeated by dye. Second, because Cx43 is expressed at low levels in the parental fibroblasts (Martyn et al., 1997), these intermediate channels could reflect the activity of heteromeric Cx43-Cx45 channels (Martinez et al., 2002). This seems unlikely for two reasons. First, both single-channel data sets (Cx43-TM and Cx43-WT) were obtained from cells whose macroscopic levels of coupling were comparable and, second, heteromeric Cx43-Cx45 channels are not permeated by Lucifer Yellow (Martinez et al., 2002). Regardless of which of these possibilities is correct, the data indicate that loss of the capacity for phosphorylation at S325, S328 and/or S330 renders the junctions inefficient at dye transfer and less conductive.

Which mechanisms might control the level of phosphorylation at S328 and/or S330 during hypoxia? Several groups have shown that phosphatase inhibitors block the change in electrophoretic mobility of Cx43 (Duthe et al., 2001; Jeyaraman et al., 2003; Turner et al., 2004). PK1 has been reported to be the main phosphatase that is active against Cx43 in cardiomyocytes and it appears to be constitutively active (Duthe et al., 2001; Jeyaraman et al., 2003; Turner et al., 2004). We have shown that CK1 can phosphorylate Cx43 at residues S325, S328, and/or S330, and CK1 inhibition leads to loss of phosphorylated Cx43, which supports the role of this kinase in the phosphorylation of these sites (Cooper and Lampe, 2002). However, a constant kinase-phosphatase cycling process has been reported to occur during hypoxia, with Cx43 dephosphorylation resulting from reduced cellular ATP levels without kinase or phosphatase activity changes or Cx43 synthesis (Duthe et al., 2001; Turner et al., 2004). Given that kinase or phosphatase localization could also be crucial, the roles that kinase activity, phosphatase activity and ATP concentration play on the control of the level of phosphorylation at residues S325, S328 and S330 during hypoxia remain to be determined.

In summary, data presented here indicate that Cx43 present in cardiac tissue is phosphorylated at serine residues 325, 328 and/or 330. During ischemia, the amount of total Cx43 is reduced at the intercalated disk and Cx43 relocates to the lateral edges of the myocytes. Cx43 at the intercalated disk retained phosphorylation at S325, S328 and/or S330, whereas Cx43 present at the lateral edges did not. We have shown that phosphorylation at these residues was necessary to form the P2 isoform, and these residues are important in gap junction assembly and stability. Thus, the expected net effect of ischemia would be a measured reduction in longitudinal conduction velocity, due to decreased localization at intercalated disks (Poelzing and Rosenbaum, 2004), without a concomitant increase in transverse conduction velocity, despite increased localization at lateral borders. Whether reduced phosphorylation at S325, S328 and/or S330 triggers the relocation of Cx43 to the lateral edges is not known. However, the spatially specific retention of phosphorylation at these sites at the intercalated disk and loss at the lateral edges would be expected to help retain the anisotropy of conduction in the heart during ischemia.

Materials and Methods

Antibodies and reagents

All general chemicals, unless otherwise noted, were purchased from Fisher Scientific. Mouse anti-Cx43 antibodies, Cx43CT1, Cx43CT2, and Cx343F1 were prepared against amino acid residues 360-382 of Cx43 and antibody Cx43NT1 against amino acids residues 1-20 of Cx43NT1 of Cx43 at the Fred Hutchinson Cancer Research Center Hybridoma Development Facility (Seattle, WA). A rabbit antibody against Cx43 was purchased from Sigma (St Louis, MO, C6219). We made a rabbit anti-pS325/328/330-Cx43 antibody by custom commercial preparation (ProSci Inc., Poway, CA; 13-week schedule) against a synthetic peptide that was phosphorylated at all three serines (pQAGS(p)TIS(p)NS(p)HAQ-amide) that had been linked via the N-terminal cysteine to maleimide-activated KLH (Pierce Biotechnology, Rockford, IL) according to manufacturer’s instructions. Phosphorylation-specific antibody was affinity purified by linking to the phosphorylated peptide and the corresponding nonphosphorylated version to SulfoLink (Pierce Biotechnology, Rockford, IL) according to the manufacturer’s instructions. After washing, phosphorylation-specific antibody was eluted from the column with 5% acetic acid and fractions were neutralized with ammonium acetate. Antibody specific for phosphorylation at S325 and S328 was affinity purified by linking peptides phosphorylated at S325-S328 and S328-S330 (CQQAGS(p)TIS(p)NSH-amide and CGSTIS(p)NS(p)HAQ-amide) to SulfoLink beads to make 1.5-mL columns of each. The polyclonal antibody was then run first over the pS328/330 column and the unbound antibody was run over the pS325/328 column. The pS325/328-specific antibody that bound was then used as described above. All antibodies were aliquoted and stored frozen at −80°C.

Cell culture

Cx43 knockout fibroblasts lacking Cx43 (clone 23-3) or exogenously expressing wild-type Cx43 (Cx43-WT; clone 22C-3) were provided by Erica TenBroek (University of Minnesota). Normal rat kidney cells (ESI) were obtained from American Tissue Type Collection (Manassas, VA). Cells were cultured in Dulbecco’s Modified Essential Medium (DMEM, Fisher Scientific, Pittsburgh PA) supplemented with 5-10% fetal calf serum and antibiotics (100 U/ml penicillin G and 100 μg/ml streptomycin) in a humidified 5% CO2 environment. Cx43 with serine residues 325, 328 and 330 mutated to alanine (Cx43-TM), kindly provided in pmDNA.1 by Steven Taffet (SUNY Health Science Center, Syracuse NY), was subcloned into pIREShygro to make Cx43-knockout fibroblasts, pIREShygro Cx43-TM was transfected into Cx43-knockout fibroblasts (clone 23-3) using Lipofectamine Plus transfection reagent (GibcoBRL/Invitrogen, Carlsbad, CA) according to the manufacturer’s instructions. Cells were selected at 500 μg/ml Hygromycin B and were dilution-cloned in DMEM supplemented with hygromycin and endothelial cell growth supplement (ECGS, 5 mg/ml) to facilitate selection and growth of cells. Three separate clones that stably expressed Cx43-TM were isolated (TM-A, TM-B, TM-C).

Immunodetection of Cx43 in heart

Mouse studies were conducted under FHRC Institutional Animal Care and Use Committee approval. Inbred mice (11 months of age in a FVB/N:C57BL6 background) were anaesthetized (avertin, 0.1 mL per 3 g body weight) and hearts were excised and placed either in ice-cold PBS (Ca2+ and glucose-free) for 30 seconds (control group) or incubated without coronary perfusion in non-oxygenated, glucose-free PBS for 30 minutes at 37°C. After 30 seconds (control) or 30 minutes (ischemic) of incubation, hearts were longitudinally bisected and either sonicated in Laemmli sample buffer (for western analysis) or fixed overnight at 4°C in 10% neutral-buffered formalin (for immunohistochemistry). Although not thoroughly characterized, the 30-minute ischemic period in the presence or absence of 1 mM Ca2+ reproduces the effects of ischemia on Cx43 electrophoretic mobility and gap junction remodeling (see Results) demonstrated in better characterized models of ischemia.

Formalin-fixed tissue was processed, sectioned, immunostained and microscopically analyzed as previously described (King and Lampe, 2004). Briefly, tissue sections were de-paraffinized, antigen retrieved, blocked and detected with rabbit anti-Cx43 (1:250, Sigma) or rabbit anti-pS325/330-Cx43 antibodies.
Approximately the same amount of cardiac tissue for each treatment was sonicated in sample buffer supplemented with 50 mM NaF, 1 mM Na3VO4, 5% mercaptoethanol, 1 mM PMFS and 1/10 complete protease inhibitors (Roche Molecular Biochemicals, Alameda, CA). Insoluble material was removed by centrifugation, and the soluble fraction was separated on 10% SDS-PAGE. After immunoblotting, protein was detected with a rabbit antibody against pS325/328/330-Cx43 and a mouse anti-Cx43 antibody (Cx43NT). The blots were also probed with anti-GAPDH (Ambion, Austin, TX) and anti-vinculin (Sigma) to confirm consistent loading. Primary antibody binding was determined as described above by LI-COR infrared imaging system (Lincoln, NE) and associated software (inverted images are shown).

In each group, we treated with alkaline phosphatase or left untreated equal amounts of protein lysates from fibroblasts expressing Cx43-WT and Cx43-TM (clone A) as previously reported (Lampe et al., 1998), ran the lysates in SDS-PAGE and performed a western blot with primary antibodies were visualized with fluorescent dye-labeled secondary antibodies ([Alexa Fluor-680 goat anti-rabbit, Molecular Probes, Eugene, OR] and IRDye800-conjugated donkey anti-mouse IgG (Rockland Immunocchemicals)] and directly quantified using the LI-COR Biosciences Odyssey infrared imaging system (see Materials and Methods).

Junctional conductance was determined using dual whole-cell voltage-clamp techniques as previously described (Cooper et al., 2002). The plates were equipped with a 40×1.3 n.a. oil objective and a Princeton Instruments digital monochrome camera (Roper Scientific, Inc., Hudson, NH). Event amplitudes were measured by hand for each cell pair and then binned into 6.25-pS bins; the binned data from all pairs were pooled, and the relative frequencies of each bin calculated and plotted. Data were fit using Origin software as previously described (Cooper et al., 2003).

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References


Dye injection

Cx43 phosphorylation at S328 and S330

Cx43 phosphorylation at S328 and S330


