Activation, Permeability, and Inhibition of Astrocytic and Neuronal Large Pore (Hemi)channels**

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Background: The permeability and physiological role of several large pore (hemi)channels are unresolved.
Results: Large pore (hemi)channels, when heterologously expressed, display isoform-specific permeability and gating for ions and fluorescent dyes.
Conclusion: Large pore channels have isoform-specific transport characteristics that can be used for their identification.
Significance: Although large pore channels have characteristic properties in overexpression systems, these properties may be undetectable in native cells.

Astrocytes and neurons express several large pore (hemi)channels that may open in response to various stimuli, allowing fluorescent dyes, ions, and cytoplasmic molecules such as ATP and glutamate to permeate. Several of these large pore (hemi)channels have similar characteristics with regard to activation, permeability, and inhibitor sensitivity. Consequently, their behaviors and roles in astrocytic and neuronal (patho)physiology remain undefined. We took advantage of the Xenopus laevis expression system to determine the individual characteristics of several large pore channels in isolation. Expression of connexins Cx26, Cx30, Cx36, or Cx43, the pannexins Px1 or Px2, or the purinergic receptor P2X7 yielded functional (hemi)channels with isoform-specific characteristics. Connexin hemichannels had distinct sensitivity to alterations of extracellular Ca2+ and their permeability to dyes and small atomic ions (conductance) were not proportional. Px1 and Px2 exhibited conductance at positive membrane potentials, but only Px1 displayed detectable fluorescent dye uptake. P2X7, in the absence of Px1, was permeable to fluorescent dyes in an agonist-dependent manner. The large pore channels displayed overlapping sensitivity to the inhibitors Brilliant Blue, gadolinium, and carbonoxolone. These results demonstrated isoform-specific characteristics among the large pore membrane channels; an open (hemi)channel is not a nonselective channel. With these isoform-specific properties in mind, we characterized the divalent cation-sensitive permeation pathway in primary cultured astrocytes. We observed no activation of membrane conductance or Cx43-mediated dye uptake in astrocytes nor in Cx43-expressing C6 cells. Our data underscore that although Cx43-mediated transport is observed in overexpressing cell systems, such transport may not be detectable in native cells under comparable experimental conditions.

Neurons and glia cells express a variety of large pore channels, including connexins, pannexins, and purinergic ionotropic receptors (1–3). Connexins are the molecular building blocks of gap junctions, but select connexin isoforms have in recent years been demonstrated to function as transmembrane transport routes in their hemichannel configuration (4–8). A range of large pore (hemi)channels appears to open to the extracellular space in response to a variety of stimuli, such as removal of divalent cations from the extracellular solution, addition of receptor agonist, and/or at positive membrane potentials (4, 6, 8–10). Channel opening is commonly measured with various fluorescent dyes of molecular sizes ranging from ~300 to ~700 Da (9, 11, 12). Proportionality between dye uptake and permeability to biologically relevant molecules has, on occasion, been implied. However, the exact permeability profile for the majority of these channel types has not been determined.

Experimental determination of channel-specific characteristics may be obscured by the parallel expression of a range of large pore channels in the cell types of interest, their overlapping permeability profile, and their cross-sensitivity to commonly used inhibitors (13–17). Knock-out and knockdown strategies for several of the implicated large pore channels have been successfully applied to address this issue (9, 16, 18–20). However, interpretation of the findings is complicated by many uncontrolled factors, including protein-protein interactions, alternative splice variants, altered transcriptomes, and/or altered basal ATP concentrations, among others (9, 21–24), thus rendering the exact contribution of the different (hemi)-channels to dye uptake, ion conductance, and ATP/glutamate release unresolved.
Although several of these large pore channels serve as dye/ATP conduits as well as ion channels, some channels exhibit a noticeable disconnect between the ability to pass dyes/ATP and current; although dye uptake and ATP glutamate/cAMP permeability can be detected at physiologically relevant membrane potentials in the negative range (4, 9, 27–29), membrane conductance for Px1 and Cx34 appears predominantly at positive membrane potentials (1, 4, 9, 25). Prolonged exposure to positive membrane potentials may even be required to detect Cx43-mediated membrane conductance (26). This seeming paradox may indicate that large pore channels do not act as nonselective freely diffusible pores but may, rather, possess the ability to selectively gate their permeability pathway (30).

Of the 21 different mammalian connexin isoforms (31), astrocytes express primarily Cx30 and Cx43 (and possibly Cx26), whereas Cx36 is the major neuronal connexin (3, 32–35). Connexins create gap junctional channels at cell to cell interfaces, allowing intercellular communication by transfer of current and small molecules (31). The permeability profile of gap junctional channels varies among isoforms (36); for example, Cx32 and Cx43 gap junctions have similar anionic/cationic current selectivity but orders of magnitude difference in their ATP/adenosine permeability (37). The molecular basis for these differences is unclear, although affinity between pore and permeant has been suggested for fluorescent dyes (38). Although connexins 26, 30, 36, and 43 are all deemed functional (36), Cx32 and Cx43 gap junctions have similar anionic/cationic current selectivity (39). Cx38 antisense has therefore been suggested that the endogenous Xenopus Cx38 does not contribute to DCFS-induced dye uptake and membrane current in our hands (30). Cx38 antisense has therefore not been co-injected in this study.

**Cell Cultures**—Cortical astrocytes were prepared from neonatal or P7 rat pups (Sprague-Dawley) or neonatal mouse pups (Cx43<sup>−/−</sup> /mice) and subcultured as detected by immunohistochemistry and Western blotting, and no compensatory changes in Cx30 expression were detected (16). Briefly, meningeal tissue was removed and the neocortex minced by pipetting, and the dissociated cells were plated on glass coverslips with culture medium consisting of Earle’s minimum essential media (Invitrogen) supplemented with 10% fetal bovine serum (HyClone, Logan, UT), 20 mM glucose, 10 units/ml penicillin, and 10 μg/ml streptomycin. Culture medium was changed every 3–4 days. Cells were used after 1–4 weeks in culture.

Native C6 glioma cells and C6 glioma cells transduced with Cx43 (C6 Cx43) were kindly provided by Zeeland Pharma, Denmark. Cells were plated at low density on glass coverslips with culture medium consisting of Dulbecco’s modified Eagle’s medium (DMEM) containing 10% fetal bovine serum (FBS), 100 units/ml penicillin, and 100 μg/ml streptomycin at 37 °C with 5% CO2 and used after 1–3 days in culture.

**Electrophysiology**—X. laevis oocytes were continuously superfused with NaCl solution (containing in mM: 100 NaCl, 2 KCl, 1 CaCl2, 1 MgCl2, 10 HEPS, 4 Tris ((HOCH2)2CNH2), pH 7.4, at room temperature) or NaCl solution without divalent cations (DCFS) in which Ca<sup>2+</sup> and Mg<sup>2+</sup> salts were substituted with equiosmolar NaCl with or without 30 μM BzATP (see under “Chemicals”) as indicated in the figure legends. The solution change took place with a custom-made continuous superfusion system (Drummond Scientific, PA) and kept in a custom-made continuous superfusion system (Drummond Scientific, PA) and kept in a

**EXPERIMENTAL PROCEDURES**

**Heterologous Expression in X. laevis Oocytes**—Oocytes were surgically removed from X. laevis frogs (Nasco or National Center for Scientific Research, France) in accordance with The Danish National Committee for Animal Studies under the Ministry of Justice. Stage V–VI oocytes were defolliculated and isolated as described previously (45) by incubation in Klorodi medium containing (in mM) 90 NaCl, 1 KCl, 1 MgCl2, 1 CaCl2, 5 HEPS, pH 7.4, at 18 °C for up to 24 h prior to cRNA injection. cDNA encoding hCx26 (obtained from Thomas W. White, Stony Brook University, New York), mCx30 and mCx36 (obtained from Klaus Willecke, Bonn University), rCx43 (obtained from Zealand Pharma, Denmark), hP2X<sub>7</sub> (obtained from Gunter Schmalzing, University of Kaachen via Rainer Schreiber, University of Regensburg, Germany), and rPx1 and rPx2 (obtained from Robert Bruzzone via Gina Sosinsky, University of California at San Diego School of Medicine (46)) were subcloned into the expression vector pXOOM optimized for oocyte expression by addition of untranslated RNA-stabilizing elements (47). The coding sequence of the cDNA was verified by sequencing prior to linearization downstream from the poly(A) segments. The cDNA was in vitro transcribed to cRNA with mMessage mMachine T7 (Ambion, Denmark) followed by transcript purification with MEGAglo (Ambion, Denmark). Oocytes were injected with 10 ng of cRNA using a Nanoject microinjection (Drummond Scientific, PA) and kept in Klorodi medium at 18 °C for 3–6 days prior to experiments. We have previously demonstrated that the endogenous Xenopus Cx38 does not contribute to DCFS-induced dye uptake and membrane current in our hands (30). Cx38 antisense has therefore not been co-injected in this study.

**Experimental Procedures**

2 The abbreviations used are: DCFS, divalent cation-free solution; ANOVA, analysis of variance; BzATP, 2’(3’)-O-(4-benzoylbenzoyl)ATP; a.u., arbitrary unit; Cx, connexin; h, human; r, rat; m, mouse; pF, picofarad.
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flow system. Although trace amounts of divalent cations are likely to be present in the DCFS solution, we estimate this concentration to be below 3 μM during the experimental condition based on trace amounts in the chemicals present in the test solution, glassware, and the weak Ca\(^{2+}\)-chelating properties of Tris((HOC\(_2\)\(_2\))\(_3\)CNH\(_2\)). Borosilicate glass capillary electrodes were pulled to a resistance of 1–2 megohms when filled with 1 M KCl. Currents were recorded from a holding potential of −50 mV by application of 100-ms voltage steps from −140 to +60 mV or 10-s voltage steps from −60 to +60 mV in intervals of 20 mV following 1 min of exposure to experimental solution, and I/V relations were obtained from the steady-state current levels. The recordings were performed with a Dagan Clampator interfaced to a PC with a Digidata 1320 A/D converter and pClamp 9.2 (Both Axon Instruments, CA). Currents were filtered at 3 kHz and sampled at 2 kHz. Dye uptake was measured using a monochromator (Polychrome V, TILL Photonics, Germany) exciting at 450 nm for ethidium and 490 nm for Yo-Pro, and emission was measured with filters allowing passage of wavelengths >510 nm (Leica Microsystems, Wetzlar, Germany).

Cultured astrocytes were seeded on coverslips that were mounted on the stage of an inverted microscope (Zeiss Axiovert S10). The cells were superfused at 1 ml/min with NaCl solution (containing in mM: 124 NaCl, 3 KCl, 1 MgCl\(_2\), 1.8 CaCl\(_2\), 5 d-glucose, 20 HEPES, pH 7.4, 37 °C) or similar solution without divalent cations (DCFS, equiosmolar substitution with NaCl). The solution change took place with continuous superfusion. Patch pipettes were pulled from borosilicate capillaries (GC150F, Harvard Apparatus, Edenbridge, UK) to a resistance of 4–6 megohms. The pipettes were front-loaded with pipette solution (containing in mM: 70 K\(_2\)SO\(_4\), 10 NaCl, 1 CaCl\(_2\), 1 MgCl\(_2\), 10 HEPES, pH 7.2) and subsequently back-filled with the same pipette solution including 0.26 mM amphotericin B (Sigma) for perforated patch clamping (49). A gigaohm seal was created on single cells, and the reduction in access resistance, due to amphotericin B generating monovalent ion-permeable pores in the membrane, was monitored as changes in the capacitive transients. A plateau in access resistance was obtained following removal of divalent cations, and regions of interest were placed randomly over single cells, with a minimum of five cells chosen per experiment. Fluorescence was sampled at 10-s intervals and recorded with MetaFluor software (Metaphor 7.6.2.0, Molecular Devices). Ethidium uptake was measured using a monochromator (Polychrome V, TILL Photonics) exciting at 340 nm and a filter allowing passage of wavelengths >520 nm for emission. Cells were superfused at 0.7 ml/min with NaCl solution (containing in mM: 124 NaCl, 3 KCl, 1.8 CaCl\(_2\), 1 MgCl\(_2\), 5 d-glucose, 20 HEPES, and 50 μM ethidium bromide, pH 7.4, room temperature), followed by a solution change via continuous superfusion to DCFS (Ca\(^{2+}\) and Mg\(^{2+}\) substituted by equiosmolar NaCl). Linear regression analysis was performed on the uptake during the last 5 min (out of 10 min) in each solution, and slopes were normalized to those obtained in control solution.

C6 cells were plated at low density on glass coverslips. The coverslips were mounted in a chamber on the stage of an inverted microscope (Zeiss Axiovert S100) with a 40×1.3 NA FLUAR oil objective (Carl Zeiss Micro Imaging GmbH, Jena, Germany). No morphological changes in C6 cells were observed following removal of divalent cations, and regions of interest were placed randomly over single cells, with a minimum of five cells chosen per experiment. Fluorescence was sampled at 10-s intervals and recorded with MetaFluor software (Metafluor 7.6.2.0, Molecular Devices). Ethidium uptake was measured using a monochromator (Polychrome V, TILL Photonics) exciting at 340 nm and a filter allowing passage of wavelengths >520 nm for emission. Cells were superfused at 0.7 ml/min with NaCl solution (containing in mM: 124 NaCl, 3 KCl, 1.8 CaCl\(_2\), 1 MgCl\(_2\), 5 d-glucose, 20 HEPES, and 50 μM ethidium bromide, pH 7.4, room temperature), followed by a solution change via continuous superfusion to DCFS and finally in NaCl-EDTA (Ca\(^{2+}\) and Mg\(^{2+}\) substituted by 2 mM EDTA).

Statistics and Data Analysis—All oocyte uptake experiments were repeated at least three times, each with oocytes from dif-
ficient animal donors and each with five oocytes per condition. Dye uptake in cultured rat and mouse primary culture of astrocytes was carried out with three different astrocytic preparations. All experiments on wild-type and Cx43 knock-out mouse astrocytes were performed in parallel (same days, same solutions, and same number of days in culture). Dye uptake in C6 cells was carried out on three distinct batches. n for uptake in oocytes, astrocytes, or C6 indicates the number of experiments, and n for electrophysiology indicates the number of individual oocytes/astrocytes.

Best fit was determined by the extra sum-of-squares F test, and three- or four-parameter logistic nonlinear regression analysis was performed with Prism 6.02 (GraphPad Software) on the Ca\(^{2+}\)- or inhibitor-sensitive component of ethidium uptake or current using Equations 1 and 2.

\[
Y = \min + \frac{(100\% - \min)}{1 + 10^\alpha(X - \log IC_{50})}
\]  

(Eq. 1)

\[
Y = \min + \frac{(100\% - \min)}{1 + 10^\alpha((\log IC_{50} - X) \cdot \text{Hill slope})}
\]  

(Eq. 2)

Astrocytic and C6 dye uptake was analyzed with linear regression and normalized to uptake obtained in control solution. Fluorescence is given as arbitrary units (a.u.). Statistical analysis was performed with Prism 6.02, using t test or ANOVA with Holm-Šidák multiple comparison post hoc test. Because equal variances are a prerequisite for using ANOVA, we used Kruskal-Wallis with Dunn’s post hoc test in cases where between groups variance differed significantly as determined by the Brown-Forsythe test. Regression analysis and statistical tests are indicated in legends. p < 0.05 was considered significant. Values in text are given as average ± S.E.

**Chemicals**—Inhibitors and fluorescent dyes from Sigma were prepared as follows: GdCl\(_3\) (H\(_2\)O, 100 mM), BzATP (H\(_2\)O, 3 mM, prepared freshly every 2nd week and stored at 5°C), Brilliant Blue G (H\(_2\)O, 50 mM), carbenoxolone (H\(_2\)O, 100 mM, prepared freshly every 2nd week and stored at 5°C), ethidium bromide (H\(_2\)O, 25 mM), Yo-Pro (DMSO, 1 mM), propidium iodide (H\(_2\)O, 1 mM), carboxyfluorescein (H\(_2\)O, 10 mM), DMEM, FBS, penicillin, and streptomycin were from Invitrogen.

**RESULTS**

**Dye Permeability and Conductance of Connexin-expressing Oocytes**—To determine the isoform-specific permeability characteristics of connexins Cx26, Cx30, Cx36, and Cx43, these constructs were expressed in *X. laevis* oocytes. As an initial measure of connexin hemichannel opening, we determined the uptake of the fluorescent molecule ethidium (314 Da) into individual connexin-expressing oocytes.

In control solution containing divalent cations (1 mM of each), the ethidium uptake observed in Cx26- and Cx30-expressing oocytes was not significantly different from that of the uninjected oocytes (Fig. 1a, white bars), indicating no basal hemichannel activity by these connexin isoforms. In contrast, Cx36- and Cx43-expressing oocytes displayed a significantly higher ethidium uptake than the uninjected oocytes (341 ± 45 and 207 ± 20 *versus* 111 ± 12 a.u., n = 5–7, p < 0.05) in a gadolinium-sensitive manner (50 μM, Fig. 1a, hatched bars). Removal of divalent cations (*i.e.* Mg\(^{2+}\) and Ca\(^{2+}\)) can induce connexin hemichannel opening (6, 8, 11, 29), and we, accordingly, exposed oocytes to DCFS (Fig. 1a, gray bars). The ethidium uptake of uninjected oocytes was insensitive to removal of divalent cations (111 ± 20 *versus* 101 ± 5 a.u., n = 7), whereas DCFS induced a significant increase in ethidium uptake in Cx30- and Cx43-expressing oocytes (557 ± 108 *versus* 142 ± 5 and 432 ± 57 *versus* 207 ± 20 a.u., respectively, n = 5–6, p < 0.001 and 0.05, gray bars) in a gadolinium-sensitive manner (Fig. 1a, gray hatched bars). Exposure to DCFS had no significant effect on ethidium uptake in Cx26- or Cx36-expressing oocytes compared with control solution (p > 0.4).

The Ca\(^{2+}\) sensitivity of the DCFS-activated Cx30 hemichannel was determined in the absence of Mg\(^{2+}\). Cx30-expressing oocytes were exposed to increasing concentrations of extracellular Ca\(^{2+}\), and the hemichannel activity was determined as the Cx-mediated ethidium uptake following subtraction of the contribution from the uninjected oocyte. The Cx30 IC\(_{50}\) value for Ca\(^{2+}\) was similar to that of Cx43 (44 ± 6 μM for Cx30 and 30 ± 9 μM for Cx43 (30), n = 5–6), although the Hill coefficient (slope) was steeper for Cx30 than for Cx43 (the Ca\(^{2+}\) sensitivity for Cx43-mediated dye uptake (Fig. 1b, dashed line) was adapted from Ref. 30)). Cx30-mediated hemichannel activity was fully inhibited by 0.5 mM Ca\(^{2+}\) (Fig. 1b), whereas Cx43-mediated hemichannel activity reached its minimum at 2 mM Ca\(^{2+}\) (30).

To further probe for connexin isoform-specific differences, we determined the hemichannel-mediated permeability toward another commonly used fluorescent probe, Yo-Pro (375 Da). The experiments were carried out in DCFS. Cx-mediated Gd\(^{3+}\)-sensitive Yo-Pro uptake was observed exclusively in Cx30-expressing oocytes (13,815 ± 5759 *versus* 3820 ± 996 a.u. in uninjected oocytes, n = 7 and 12, p < 0.01, Fig. 1c), whereas no significant uptake was observed in oocytes expressing the other tested isoforms. A similar pattern was observed with propidium iodide (668 Da) (data not shown). These data illustrate that dye uptake via Cx hemichannels displays isoform-specific characteristics regarding both Ca\(^{2+}\) sensitivity and dye selectivity.

To determine whether hemichannel-mediated dye permeability was paralleled by ionic conductance, we applied two-electrode voltage clamp to connexin-expressing oocytes (Fig. 2). In control solution, the membrane current for Cx26- and Cx30-expressing oocytes was slightly increased compared with that of uninjected oocytes (250 ± 40 and 316 ± 33 nA *versus* 143 ± 8 nA at +60 mV, n = 15, 10, and 19, p < 0.05), whereas the membrane current for Cx36- and Cx43-expressing oocytes was comparable with the uninjected control (137 ± 8, 168 ± 9, and 143 ± 8 nA at +60 mV, n = 8–19). Exposure to DCFS (1 min) induced large Cx-mediated membrane currents in oocytes expressing Cx26 and Cx30 (2041 ± 415 and 2390 ± 631 nA at +60 mV, n = 15 and 10, p < 0.01 compared with uninjected oocytes; 317 ± 32 nA, n = 19), whereas the DCFS-induced membrane current for Cx36- and Cx43-expressing oocytes was similar to that of the uninjected control oocytes. Applying a longer pulse of 10 s to Cx36-expressing oocytes also failed to...
activate a conductance, and the current in DCFS was similar to that of the uninjectected oocytes (455 ± 155 versus 385 ± 150 nA at +60 mV, n = 7, p > 0.7). The permeation pathway through connexin hemichannels, when expressed in *Xenopus* oocytes, appears to be highly selective toward fluorescent dyes and/or ions. The different connexins thus display distinct isoform-specific behavior in their hemichannel configuration.

**Dye Permeability and Conductance of Pannexin-expressing Oocytes**—Pannexins constitute another group of large pore channels and may have some functionality in common with the connexin hemichannels. We determined the dye permeability and membrane conductance in oocytes expressing two pannexin isoforms, Px1 or Px2. Px1- and Px2-expressing oocytes displayed an outwardly rectifying membrane conductance at positive membrane potentials although smaller for Px2-expressing oocytes than for Px1-expressing oocytes (1130 ± 61, 365 ± 39, and 133 ± 10 nA at +60 mV for Px1, Px2, and uninjectected, respectively, p < 0.001, n = 9 – 17) (Fig. 3a).

In control solution, Px1-expressing oocytes displayed greater uptake of ethidium and Yo-Pro than the uninjectected oocytes (Fig. 3, b and c, white bars). The ethidium and Yo-Pro uptake of Px2-expressing oocytes was similar to those obtained in the uninjectected oocytes, illustrating no detectable ethidium and Yo-Pro permeability through Px2 (Fig. 3, b and c). Exposure to DCFS had no effect on ethidium uptake in Px1- or Px2-expressing oocytes (Fig. 3b, gray bars). We found a tendency, although it did not reach statistical significance, toward carbboxyfluorescein permeability through Px1 (766 ± 122 a.u., n = 6, p > 0.27) and Px2 (981 ± 304 a.u., n = 5, p > 0.15) as compared with that of the uninjectected oocytes (469 ± 130 a.u., n = 6) (Fig. 3d). Exposure to the P2X7 agonist BzATP (10 μM, see below) did not affect dye uptake in Px1- or Px2-expressing oocytes (data not shown). Thus, Px1 displayed permeability to ethidium and Yo-Pro in a nongated manner at resting membrane potential, whereas Px1-mediated ion conductance was observed only at positive membrane potentials.

**Dye Permeability and Conductance of P2X7-expressing Oocytes**—To determine whether the purinergic receptor P2X7 possesses the inherent ability to allow dye permeation (versus the proposed requirement for Px1 (42, 44)), we determined the permeability and activation profile for P2X7-expressing oocytes. In control solution, as well as in DCFS, P2X7-expressing oocytes displayed membrane conductance similar to that of the uninjectected oocytes (compare Fig. 4a (P2X7) and 3a (Uninj)). Addition of the P2X7 agonist BzATP (30 μM) to the test solution induced a large, nonrectifying, and fully reversible membrane conductance (Fig. 4a, inset). Note that the current pattern induced by BzATP was biphasic, with an initial "shoulder" (Fig. 4a, inset). BzATP-induced current was not observed in uninjectected oocytes (data not shown).

In control solution, the dye uptake of P2X7-expressing oocytes was similar to that of the uninjectected oocytes regardless of connexin expression in DCFS in the absence (gray bars) or presence of 50 μM Gd3+ (hatched bars). Values are given in arbitrary units. n = 3–12 experiments, and statistical significance was determined using one-way ANOVA with Holm–Šidák post hoc test and illustrated as in a. Error bars indicate S.E. ns, not significant, *, p < 0.05; **, p < 0.01; ***, p < 0.001.

![FIGURE 1. Differences in hemichannel-mediated ethidium and Yo-Pro uptake following removal of divalent cations.](image-url)
of the dye used (i.e. ethidium, Yo-Pro, or carboxyfluorescein; Fig. 4, b–d, white bars). Addition of BzATP (10 μM) did not affect dye uptake in the uninjected oocytes (Fig. 4, b–d, black bars) but promoted dye uptake in P2X7-expressing oocytes as follows: ethidium (184 ± 29 versus 56 ± 8 a.u., n = 7, p < 0.01, b), Yo-Pro (4598 ± 1273 versus 1250 ± 167 a.u., n = 6, p < 0.01, c), and carboxyfluorescein (3479 ± 1155 versus 497 ± 114 a.u., n = 6, p < 0.05, d). Ethidium uptake in P2X7-expressing oocytes was unaffected by removal of divalent cations from control solution (Fig. 4b, gray bar), whereas the combination of BzATP and DCFS resulted in a larger P2X7-mediated ethidium uptake than seen with BzATP alone (compare Fig. 4, b and e). P2X7 therefore mediated dye permeation and biphasic current in the absence of Px1, its proposed interaction partner for large pore formation.

**Pxl and P2X7 Co-expression**—To test for a potential functional interaction between P2X7 and Pxl, these constructs were co-expressed in oocytes. In control solution, membrane conductance of Pxl/P2X7-expressing oocytes was comparable with that of Pxl-expressing oocytes in size and rectification (compare Fig. 3a with Fig. 5a). Upon addition of BzATP to the test solution, the conductance increased and resembled that of non-rectifying P2X7-expressing oocytes (compare Figs. 4a and 5a).

In control solution, the Pxl/P2X7-expressing oocytes were permeable to ethidium (369 ± 80 versus 60 ± 12 a.u., n = 5–6, p < 0.01, Fig. 3b) and Yo-Pro (5348 ± 984 versus 1214 ± 238 a.u., n = 5, p < 0.01, Fig. 3c) but not significantly to carboxyfluorescein (1137 ± 382 versus 503 ± 134 a.u., n = 5–6, Fig. 3d). These levels of dye uptake were similar to the observations with oocytes expressing only Pxl (Fig. 3, b–d). Exposure to DCFS did not increase ethidium uptake in Pxl/P2X7-expressing oocytes (Fig. 5b), and exposure to the P2X7 agonist BzATP did not further increase the uptake of ethidium, Yo-Pro, or carboxyfluorescein (Fig. 5, b–d). Therefore, co-expression of Pxl and P2X7 did not confer novel membrane conductance or dye uptake.

**Potency of Brilliant Blue, Gadolinium, and Carbenoxolone**—Several inhibitors are commonly used to identify different large pore channels in complex cell systems, although their potency and selectivity for the different (hemi)channels have not been fully characterized, limiting their utility (15, 16). We determined the potency of three commonly used inhibitors, Brilliant Blue, gadolinium (Gd3+), and carbenoxolone in blocking ethidium uptake via connexin hemichannels, Pxl, and P2X7 expressed in oocytes. Cx26-expressing oocytes were not tested because they lack measurable ethidium permeability (Fig. 1a). The contribution from the native oocyte membrane (the uninjected oocytes) was subtracted to generate the inhibition curves. Brilliant Blue had the greatest potency against P2X7 with an IC50 13.6 ± 2.3 μM, n = 4 (Fig. 6a) although an effect on Pxl and the Cx hemichannels was apparent at higher concentrations (IC50: Cx30, 82 ± 23, 169 ± 22, and 37 ± 6 μM for Cx30, Cx43, and Pxl, respectively, n = 3–4). Gd3+ inhibited Cx30-, Cx36-, and P2X7-mediated ethidium uptake with similar potency (IC50: Cx30, 2.4 ± 0.4, 2.0 ± 0.4, and 2.8 ± 1.0 μM for Cx30, Cx36, and P2X7, respectively, n = 3–6, Fig. 6b), which is comparable with the IC50 value previously described for Cx43 (0.4 ± 0.1 μM (30)). Notably, the Gd3+ sensitivity of Cx30-mediated currents was higher than that obtained for dye uptake (0.3 ± 0.1 versus 2.4 ± 0.4 μM, n = 6, p < 0.01). In contrast,
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Curiously, Cx36 displayed carbenoxolone-induced increase in dye uptake, to 348 ± 45% of control at 200 μM, n = 5 (Fig. 6d). These inhibitor profiles demonstrate isoform-specific differences among the expressed (hemi)channels.

Hemichannel Activity in Cultured Rat Astrocytes—Having established key characteristics of possible astrocytic large pore channels in an isolated setting, we attempted to identify the molecular identity of the channel underlying DCFS-induced permeability in a native setting. First, we determined the DCFS-activated membrane current and dye permeability in primary cultures of rat cortical astrocytes. In conventional whole cell patch clamping, the intracellular content may be partially replaced with that of the pipette solution which may interfere with intracellular regulation of connexin hemichannels. The electrophysiological recordings were thus carried out using the perforated patch clamp technique to avoid replacing the cytosol with the pipette solution. The astrocytic membrane current was determined in control solution containing Ca²⁺ and Mg²⁺ and subsequently in DCFS (Fig. 7a, top panel, and summarized in b). The membrane current was not significantly increased in the absence of divalent cations (27 ± 7 versus 23 ± 8 pA/pF at +60 mV, n = 11 cells, p > 0.9) and was not significantly decreased upon addition of 50 μM Gd³⁺ to the DCFS (24 ± 8 pA/pF at +60 mV, n = 10, p > 0.9) (Fig. 7a and b). In cultured rat astrocytes, no DCFS-induced, Gd³⁺-sensitive membrane current above background was detected at the tested membrane potentials.

Dye permeability to ethidium and Yo-Pro was determined by adding dye to an astrocytic culture following determination of baseline background fluorescence. The dye uptake was subsequently monitored continuously. Cultured astrocytes were exposed to control solution (containing Ca²⁺ and Mg²⁺) for 10 min followed by exposure to DCFS (see Fig. 7c for representative traces). The rate of dye uptake was quantified for the last 5 min in control solution and in DCFS, and the data were summarized and plotted as the “fold increase” in dye uptake rate upon removal of divalent cations. DCFS induced a significant increase in uptake of both fluorescent dyes (3.1 ± 0.3-fold for Yo-Pro uptake, n = 5, p < 0.01, and 1.4 ± 0.1-fold for ethidium uptake, n = 9, p < 0.05) (Fig. 7d, gray bars). The DCFS-induced uptake of Yo-Pro and ethidium were, however, insensitive to addition of 50 μM Gd³⁺ to the DCFS, n = 5–9 (Fig. 7d, hatched bars and inset). Thus, removal of divalent cations from the extracellular solution did promote dye uptake into cultured astrocytes, but the lack of Gd³⁺ sensitivity raised questions about the molecular mechanism underlying the dye uptake.

Dye Permeability of Cultured Astrocytes from WT and Cx43 cKO Mice—To resolve a putative role of Cx43 in astrocytic dye uptake, DCFS-induced ethidium and Yo-Pro uptake were determined, in parallel, in cultured astrocytes from wild-type (WT) and Cx43 cKO mice. Ethidium and Yo-Pro were added simultaneously, and the dye uptake was monitored as described above. Similar to the observations in rat astrocytes, a basal dye uptake was observed in control solution containing Ca²⁺ and Mg²⁺ (Fig. 8a). Following exposure to DCFS, the dye uptake of WT astrocytes increased to 1.12 ± 0.04-fold of the control for ethidium and 1.39 ± 0.09-fold for Yo-Pro (n = 35 experiments,
rates of ethidium uptake (Fig. 9). Cellular fluorescence was monitored continuously to increase the possibility for detection of Cx43-mediated dye uptake. p > 0.05 (Fig. 8b). However, we detected no significant difference between the DCFS-induced uptake in WT and Cx43 cKO astrocytes with ethidium (1.12 ± 0.04, n = 35 for WT versus 0.95 ± 0.06, n = 18 for Cx43 cKO, p > 0.35) and Yo-Pro (1.39 ± 0.09, n = 35 for WT versus 1.29 ± 0.16, n = 18 for Cx43 cKO, p > 0.45) (Fig. 8b). It is important to note that cells exposed to control solution for time periods similar to those in which we tested DCFS revealed no difference in dye uptake as a function of time (n = 5, p > 0.75, data not shown). In conclusion, DCFS-induced uptake of ethidium and Yo-Pro in mouse astrocytes was minor and independent of Cx43 expression.

**Cx43 in Rat C6 Glioma Cells**—With the genetic approach above providing evidence against a significant contribution from Cx43 in DCFS-induced dye uptake in cultured astrocytes, we next tested the C6 glioma cell line, with negligible endogenous expression of connexins and pannexins (50, 51), against Cx43. With the genetic approach above providing evidence against a significant contribution from Cx43 in DCFS-induced dye uptake in cultured astrocytes, we next tested the C6 glioma cell line, with negligible endogenous expression of connexins and pannexins (50, 51), against Cx43. The cDNA encoding Cx43 was transduced into the genome of the native C6 cells. Cx43 was not detectable in WT C6 cells but was clearly seen in cells transduced with Cx43 (Fig. 9a). Wild-type C6 cells and Cx43-expressing C6 cells were exposed to ethidium in a sequence of solutions as follows: control solution followed by DCFS and subsequently by DCFS plus EDTA (to ensure complete chelation of any residual divalent cations and therefore increase the possibility for detection of Cx43-mediated dye uptake). Cellular fluorescence was monitored continuously (Fig. 9b). The native and transduced cells displayed comparable rates of ethidium uptake (Fig. 9b), as summarized in Fig. 9c (n = 3 matched experiments with a total of 52 and 49 cells for WT and Cx43, respectively). The rates of ethidium uptake during these experiments never varied from those seen in C6 Cx43 cells continuously exposed to control solution containing 1.8 mM Ca\(^{2+}\) for the same period of time (n = 3 experiments with 28 cells, data not shown). In summary, no increase in dye uptake of C6 cells was produced by DCFS regardless of whether Cx43 was expressed or not.

**DISCUSSION**

We have in this study demonstrated that large pore channels of different molecular identities have overlapping, although isoform-specific, channel characteristics when expressed heterologously in *Xenopus* oocytes. Curiously, we observed no direct link between atomic ion conductance and dye permeability for the different pores.

Removal of divalent cations from the extracellular solution induces hemichannel opening in a variety of connexins (6, 8, 11, 29), and accordingly, we observed hemichannel activity in oocytes expressing Cx26, Cx30, and Cx43 upon exposure to DCFS. Cx36, however, was constitutively permeable to ethidium and unaffected by the concentration of divalent cations in the extracellular solution. Cx43 displayed a slight ethidium uptake in the presence of 1 mM Ca\(^{2+}\), which is in accordance with our previous findings where Cx43 is only partially closed at this Ca\(^{2+}\) concentration (30). The sensitivity to extracellular Ca\(^{2+}\) for dye uptake through Cx30 was distinct from that of Cx43, with a steeper slope and near-complete inhibition for Cx30-mediated dye uptake at 0.5 mM Ca\(^{2+}\), although the Cx30 and Cx43 IC\(_{50}\) values for Ca\(^{2+}\) were similar (this study and see...
Ref. 30). It must be noted, however, that removal of divalent cations may have a variety of secondary cellular effects such as (i) membrane depolarization which may (52) or may not (30, 53) promote Cx43 hemichannel opening, and (ii) changes in intracellular Ca\(^{2+}\)/H\(^{+}\) concentration that may, in itself, affect hemichannel opening (54, 55). Once the opening was induced in the respective hemichannels, the permeation pattern differed among the connexins: Cx30-expressing oocytes displayed Gd\(^{3+}\)/H\(^{+}\)-sensitive ion conductance and permeability to both ethidium, Yo-Pro, and propidium, whereas Cx36- and Cx43-expressing oocytes displayed only ethidium permeability (in a Gd\(^{3+}\)/H\(^{+}\)-sensitive fashion), and Cx26-expressing oocytes solely exhibited Gd\(^{3+}\)/H\(^{+}\)-sensitive membrane conductance and no dye permeability. However, it should be noted that oocytes expressing Cx36 and Cx43 had a tendency to reduced Yo-Pro and propidium uptake, although not to a statistically significant degree. This could indicate that another minor Yo-Pro/propidium-permeable transport system, native to the oocyte membrane, may be outcompeted by the experimental overexpression of connexins. Additionally, the Yo-Pro/propidium uptake and the gadolinium-sensitive component appeared greater for Cx26-expressing than for Cx36- and Cx43-expressing oocytes. Thus, we cannot exclude slight Cx26-mediated dye permeability in the present setting. These data may align with previously published data on limited DCFS-induced Cx26-mediated dye uptake in Cx-expressing HeLa cells: the low number (21%) of cells permitting DCFS-induced dye uptake (56) and the limited (~2-fold) DCFS-induced increase in dye uptake (57).

Activation of hemichannel-mediated atomic ion conductance by removal of extracellular divalent cations has previously been demonstrated at physiologically relevant membrane potentials for Cx26 and Cx30 expressed in Xenopus oocytes or mammalian cell lines (6, 8, 30), whereas it is often reported that highly positive membrane potentials (>40 mV) or addition of cytokines is required to observe Cx43-mediated conductance in mammalian cell systems (4, 25, 26, 58). Lack of Cx36 hemichannel-mediated conductance has previously been demonstrated (59, 60). Interestingly, although Cx36, in its gap-junction configuration, has an extremely low single channel conductance (10–15 pS), Cx36 gap junctions are nevertheless permeable to the fluorescent dyes Lucifer Yellow and ethidium (61, 62).

Our data indicate that the connexin hemichannels have distinct extracellular Ca\(^{2+}\)-mediated activation regimes and display distinct permeability toward ions as well as toward dyes of different molecular composition. Although the difference may seem more subtle, cell to cell gap junctional channels do share peculiar relations between their single channel conductance and selectivity. A thorough comparison by Harris and Locke (63) showed no correlation between single channel conductance and the selectivity for atomic cations versus anions. When dealing with molecules, the complexity only increases, and some connexins, such as Cx30, are in their gap-junction configuration, have an extremely low single channel conductance (10–15 pS), Cx36 gap junctions are nevertheless permeable to the fluorescent dyes Lucifer Yellow and ethidium (61, 62).

A disconnect between hemichannel-mediated dye permeability and conductance, such as that observed in this study for Cx26, Cx36, and Cx43, seems counterintuitive due to the ~15 Å large pore reported in the crystal structures of connexins 26 and 43 (66, 67). However, inclusion of amino acid side chains lining the pore will reduce the pore size to ~5 Å (66). In addition, the C and N termini may create an intracellular plug with potential participation in channel gating (68). Connexin hemichannels in their intact and functional form may therefore not operate with an open ~15 Å large pore but may, via the amino

![Figure 5. No additive effect of co-expressing Px1 and P2X, on membrane conductance and dye permeability.](image-url)
acid side chains lining the pore, their termini, and extracellular loops, generate a form of isoform-specific pore selectivity filters as well as permeant-specific gating properties. It must be noted that molecular mechanisms allowing permeability of large molecules, while concomitantly excluding ions, are observed in an unrelated class of large pore membrane channels, the aquaglyceroporins such as AQP9 (69, 70).

Pannexins belong to another class of large pore channels that appear not to form intercellular junctional pores (71). Px1 has been suggested as an alternative to the proposed Cx43-mediated astrocytic hemichannel activity (9, 72) as it shares key properties with the connexin hemichannels, including dye permeability, ATP release, and propagation of Ca$^{2+}$/H$^{+}$ waves (14, 15, 71). Oocytes expressing either Px1 or Px2 displayed an outwardly rectifying conductance at positive membrane potentials, which is characteristic for pannexins whether heterologously expressed in mammalian cells or *Xenopus* oocytes (9, 44, 71, 73). The Px2-expressing oocytes displayed smaller membrane conductance than that of the Px1-expressing oocytes, but without the knowledge of unit expression level, we cannot, at present, exclude that the difference in membrane conductance originates from lower expression levels of Px2. No dye uptake was detectable in the Px2-expressing oocytes, although Px1-expressing oocytes displayed a constitutively large uptake of ethidium and Yo-Pro. Notably, the large dye uptake was readily observed at basal negative membrane potentials, whereas the membrane conductance appeared exclusively at positive membrane potentials in a strictly voltage-gated manner. In analogy with the connexin hemichannels, pannexins therefore possess the ability to distinguish between fluorescent dyes and atomic ions with a distinct gating paradigm for each permeant. The Px1-mediated ethidium uptake was not affected by removal of divalent cations, and with this lack of DCFS-induced opening, it appears that Px1 and Cx43 may well open under distinct physiological conditions, and therefore each serve a different physiological purpose in native cells.

P2X purinergic receptors are agonist-induced ion channels, some of which (e.g., P2X$_7$) undergo pore dilation upon prolonged activation, allowing for permeation of large molecules and/or increased current (40, 41, 74). Px1 has been proposed to enter into a functional complex with P2X$_7$ and act as the molecular origin of the large pore (42–44). BzATP-induced nonrectifying membrane conductance was observed in P2X$_7$-expressing oocytes with the previously reported characteristics of P2X$_7$-mediated conductance (10, 75). Expression of P2X$_7$ allowed for agonist-induced permeation of three different fluorescent dyes (ethidium, Yo-Pro, and carboxyfluorescein). Although a few reports suggest inherent lack of dye permeability in P2X$_7$ (42–44), P2X$_7$-mediated large molecule and fluorescent dye permeability (in the absence of Px1) have, in analogy with the present data, been documented in mammalian cell lines (40, 76), thus arguing against the proposed requirement for Px1 to act as the large pore channel during P2X$_7$-mediated dye uptake.

**FIGURE 6.** Dye uptake in the presence of Brilliant Blue, gadolinium, and carbenoxolone. Ethidium uptake or membrane conductance (Cx30, gadolinium; I at $-60$ mV) was determined in Cx30-, Cx36-, Cx43-, Px1-, and P2X$_7$-expressing oocytes exposed to either Brilliant Blue (BB), gadolinium (Gd$^{3+}$), or carbenoxolone (CBX). Connexin-mediated uptake was determined in DCFS; Px1-mediated uptake was obtained in control solution, and BzATP (10 $\mu$M) was employed to obtain the P2X$_7$-mediated uptake. Following subtraction of the contribution from the uninjected oocytes, ethidium uptake as a function of inhibitor concentration was fitted using linear regression (Gd$^{3+}$, Px1) or a three (Brilliant Blue and carbenoxolone; Cx30, Gd$^{3+}$; Cx30, Cx36, and P2X$_7$) or four (Brilliant Blue; Cx43, Px1, and P2X$_7$) parameter logistic equation. $a$, uptake with Brilliant Blue concentrations from 0 to 1000 $\mu$M, $n = 3–4$ experiments with 5–6 points. $b$, uptake with Gd$^{3+}$ from 0 to 1000 $\mu$M, $n = 3–6$ experiments with 6–7 points (2–4 for Px1). Included is the Cx30-mediated current, obtained by a two-electrode voltage clamp as in Fig. 2 and fitted using a four-parameter logistic equation. $c$ and $d$, uptake with carbenoxolone from 0 to 200 $\mu$M, $n = 3–5$ experiments with 5 points. Each data point averages $n$ experiments; line indicates best fit, and IC$_{50}$ was determined as the average of independent regression analyses.
The I/V relationship of the membrane conductance observed in Px1/P2X7-expressing oocytes was indistinguishable from that of the Px1-expressing oocytes and in the presence of BzATP indistinguishable from that of the P2X7-expressing oocytes, thus indicating that both large pore channels are functionally expressed and capable of conducting ions, independently of each other, in a nonsynergistic way upon co-expression in Xenopus oocytes. The BzATP-induced current traces obtained in P2X7-expressing oocytes consistently presented with a shoulder prior to full channel opening. This particular pattern has previously been observed in mammalian cells and was suggested to originate from initial BzATP-induced activation of P2X7 (the shoulder) and a subsequent P2X7-mediated activation of Px1 (the full current activation) (42). Observation of such a shoulder in oocytes exclusively expressing P2X7 suggests that this current pattern is inherent to P2X7 and thus argues against the requirement of Px1 for full activation. Co-expression of Px1 and P2X7 did not lead to synergistic effects on the dye uptake. In the absence of P2X7 agonist, Px1/P2X7-expressing oocytes displayed the large constitutive dye uptake also observed in oocytes expressing Px1 alone. Curiously, for reasons presently unknown, BzATP failed to induce further dye uptake in Px1/P2X7-expressing oocytes. One possible explanation for this discrepancy could be the previously reported BzATP-mediated inhibition of Px1 (77) that we, however, did not observe in this study. Taken together, it appears that P2X7 possesses the ability to undergo pore dilation and thus, in the absence of Px1, allow for permeation of fluorescent dyes. Although P2X7 receptors have the capability to function independently of Px1, our data by no means preclude that Px1 may be activated by other receptors, as has been demonstrated for α-adrenergic and NMDA receptors (78, 79).

A range of studies has been carried out on complex cell systems and molecular identity to a certain transport pathway assigned partly by the aid of pharmacological inhibitors (42, 43, 60, 80, 81). Although several of these agents act in a potent fashion, their specificity is either nonexisting or awaits determination, which complicates finite interpretation of the individual role of the different large pore channels (15, 16, 82). A recent approach that may offer more specificity has been the mimetic peptide Gap19, which inhibited Cx43 hemichannels without affecting Cx40 or Px1 (83). In this study, Brilliant Blue was a potent inhibitor of P2X7-mediated dye uptake in accordance with Ref. 84, but it also affected connexin hemichannels and Px1 albeit at slightly higher concentrations (the latter previously observed by (77)). Gadolinium, however, inhibited dye uptake of all tested connexin hemichannels and P2X7 to a

**FIGURE 7.** Dye permeability but not membrane conductance of rat cortical astrocytes is increased following exposure to DCFS. a, membrane conductance obtained with the perforated patch clamp technique was recorded from a holding potential of −80 mV by application of 1-s voltage steps from −100 to +60 mV in intervals of 20 mV in control solution with 1.8 mM Ca2+ and 1 mM Mg2+. In panels b and c, cumulative tracings are presented as in a and determined at the end of pulses, with currents in control solution (white symbols), DCFS (gray symbols), and DCFS + 50 μM Gd3+ (black-bordered gray symbols). n = 10–11, currents were normalized to membrane capacitance (pA/pF). c, dye uptake visualized by fluorescence microscopy with representative traces of ethidium and Yo-Pro uptake in cells in control solution followed by exposure to DCFS. d, summarized Yo-Pro and ethidium uptake rates in DCFS in the absence (gray bars) or presence (hatched bars) of 50 μM Gd3+ from experiments as in c (see inset) with uptake rates normalized to uptake in control solution, n = 5 and 9 experiments for Yo-Pro and ethidium, respectively. Statistical significance was determined using one-way (c and d) or two-way (b) ANOVA with Holm-Sidak post hoc test, and one sample t test against the value 1.0 for DCFS (d). Statistical significance is indicated over bars for difference compared with uptake in control solution and over lines for the given in-group comparisons. Error bars indicate S.E. *, p < 0.05; **, p < 0.01; ns, not significant.
similar extent while displaying a higher potency toward Cx30-mediated current versus dye uptake. Px1 was completely insensitive to gadolinium even at very high concentrations, as previously observed (44, 85). Carbenoxolone, curiously, increased Cx36-mediated dye uptake, inhibited Cx30-mediated dye uptake, and displayed a relatively low potency toward Cx43, Px1, and P2X7. The increase in Cx36-mediated uptake may reflect relief from inhibition by intramembrane arachidonic acid as reported for short chain alcohols on Cx36 but not Cx43 gap junctional channels (86). Carbenoxolone has previously been reported to display a partial effect on Yo-Pro uptake in Px1-expressing HEK293 cells (87), whereas much higher potency is found in electrophysiological experiments in both oocytes and complex cell systems expressing Px1 (73, 85). Based on the higher Gd$^{3+}$ sensitivity observed with Cx30-mediated current versus dye uptake (this study), it is possible that Px1-mediated currents and dye uptake are differentially affected by carbenoxolone. Isoform-specific inhibitor profiles thus exist among the connexins in their hemichannel configuration, and the inhibitor potency varies with transport mode (dye versus current). These data now add, to the already extensive list of promiscuous large pore channel inhibitors (15, 82), Gd$^{3+}$ as a potent P2X$_7$-inhibitor, Brilliant Blue as a connexin hemichannel inhibitor, and carbenoxolone as more potent toward Cx30 than Cx43. However, according to the present data, gadolinium provides a useful tool with which to distinguish between pannexin- and connexin-hemichannel-mediated dye permeability (while bearing in mind the plethora of other Gd$^{3+}$-sensitive channels expressed in native cells).
Large Pore (Hemi)channels

Heterologous expression systems such as Xenopus oocytes and mammalian cell lines are designed to overexpress a given protein to an extent where the background permeabilities become negligible. The advantage thereof is to provide a setting in which the biophysical properties of a transport protein of interest can be determined. It must, however, be emphasized that in expression systems and cultured cells, one cannot be assured of correct protein processing and/or the presence of any required cofactor, etc. Observations obtained with gross overexpression of a given protein could, however, not be detectable in a native setting where expression may be much lower and a range of other transport proteins may contribute to the measured transport. We therefore aimed to determine the large pore channel activated by DCFS in astrocytes using the characteristics identified in this study. In cultured rat cortical astrocytes, however, we detected no evidence of a DCFS-induced current at negative membrane potentials (4, 28, 30). In rat astrocytes, we detected a DCFS-induced increase in membrane potentials in Cx43-expressing HeLa cells (4, 25). It is, at present, not clear why differences in electrophysiological approaches may play a role versus fast whole cell patch versus perforated whole cell patch). The astrocytic data of this study are in agreement with data from Cx43-expressing HeLa cells and Xenopus oocytes, thus showing a lack of significant Cx43-mediated current at negative membrane potentials (4, 28, 30). In rat astrocytes, we detected a DCFS-induced increase in uptake of ethidium and Yo-Pro which was, however, insensitive to Gd3+ and therefore unlikely to be Cx43-mediated. A few research groups have previously reported DCFS-induced Lucifer Yellow uptake in astrocytic cultures (29, 89), whereas other groups failed to detect DCFS-induced permeability to ethidium or calcein when astrocytes were grown in conventional medium (53, 90). With the lack of a specific Cx43 inhibitor, we took advantage of the conditional Cx43 knock-out mice but detected no significant difference between the DCFS-induced uptake of ethidium or Yo-Pro in a comprehensive series of parallel experiments on astrocytes obtained from these animals in comparison with those obtained from WT mice. Cx43-expressing C6 cells (transduced into the genome rather than transfected and hence presumably not overexpressing Cx43 to the same level as that obtained by transient transfections) have previously been shown to support gap-junctional coupling (91). Nevertheless, these cells did not present with DCFS-induced dye uptake under the employed experimental conditions. Although Cx43 hemichannels are indeed able to transport small quantities of ethidium upon exposure to DCFS when overexpressed in a cellular system (this study and Refs. 4, 11), in a complex system with parallel dye/ion transport routes their contribution to dye/ion transport may simply not be detectable upon removal of divalent cations. However, a range of experimental factors may well promote experimentally detectable Cx43-mediated dye uptake and ion conductance, e.g. species-specific channel function, culture conditions, choice of fluorescent dye, presence of cytokines, the metabolic state of the cells, excessively positive membrane potentials, and/or the mechanism of solution exchange. In relation to the latter, removal of control solution by suction and subsequent addition of test solution may well induce a mechanical stimulus, which in itself has been shown to affect Cx43 hemichannel opening under calcium-free conditions (88).

In conclusion, we have demonstrated that a range of neuronal and astrocytic channels, when heterologously expressed, display type- and isoform-specific extracellular Ca2+ sensitivity and permeation. Although the central pore of these channels is of relatively large diameter, molecular mechanisms appear to be in place to prevent nonselective free diffusion of molecules upon channel opening. Several of the tested (hemi)channels possess the ability to gate conductance and fluorescent dye permeability independently. We detected no obvious DCFS-induced Cx43-dependent dye uptake in cultured astrocytes and conclude, under our experimental conditions, that although Cx43 may indeed be able to allow dye permeation and conductance under certain experimental conditions, their contribution to these transport parameters may, upon removal of divalent cations, be negligible in complex native cells such as astrocytes. It seems probable, however, that future studies may reveal additional variables that impact the function and gating of Cx43 hemichannels in astrocytes.

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