Abstract. Physiological properties of isolated pairs of rat hepatocytes were examined within 5 h after dissociation. These cells become round when separated, but cell pairs still display membrane specializations. Most notably, canaliculi are often present at appositional membranes which are flanked by abundant gap and tight junctions. These cell pairs are strongly dye-coupled; Lucifer Yellow CH injected into one cell rapidly diffuses to the other. Pairs of hepatocytes are closely coupled electrically. Conductance of the junctional membrane is not voltage sensitive: voltage clamp studies demonstrate that \( g_j \) is constant in response to long (5 s) transjunctional voltage steps of either polarity (to \( \pm 40 \) mV from rest). Junctional conductance \( (g_j) \) between hepatocyte pairs is reduced by exposure to octanol (0.1 mM) and by intracellular acidification. Normal intracellular pH \( (pH_i) \), measured with a liquid ion exchange microelectrode, was generally 7.1-7.4, and superfusion with saline equilibrated with 100% CO\(_2\) reduced pH to 6.0-6.5. In the pH range 7.5-6.6, \( g_j \) was constant. Below pH 6.6, \( g_j \) steeply decreased and at 6.1 coupling was undetectable. \( pH_i \) recovered when cells were rinsed with normal saline; in most cases \( g_j \) recovered in parallel so that \( g_j \) values were similar for pHs obtained during acidification or recovery. The low apparent pK and very steep pH--\( g_j \) relation of the liver gap junction contrast with higher pKs and more gradually rising curves in other tissues. If \( H^+ \) ions act directly on the junctional molecules, the channels that are presumably homologous in different tissues must differ with respect to reactive sites or their environment.

LIVER is a tissue rich in gap junctions and it has therefore been a useful source of material for morphological, immunological, and biochemical studies of gap junction protein. The first detailed structural studies on gap junctional membranes included those in sections of intact liver stained with lanthanum (26), and the correlation of images of the gap junction in thin section with freeze-fracture replicas was aided by the abundance of this structure in liver (for review, see reference 2). Most recently, isolated hepatic gap junction membranes have provided material for high resolution structural studies using x-ray diffraction and low dose electron microscopy (18, 42, 47). The relatively large amount of junctional protein in liver has also allowed isolation for immunological and biochemical studies (for example, reference II). Among exciting recent developments are antibodies that block junctional conductance (12, 43) and a partial sequence of amino acid residues comprising the major junctional protein (see reference 21) that has yielded a complete cDNA clone (23). Despite this progress, electrophysiological properties of hepatocyte gap junctions have remained poorly known, in part because of the small size of the cells and the opacity of the intact tissue (see references 9 and 24). Moreover, the component cells are complexly interconnected: coupled cells are arrayed in cords (6) so that models of current flow assuming three-dimensional uniformity cannot be used for spatial analysis (20).
Materials and Methods

Cell Preparation

Cells were dissociated according to the procedure of Berry and Friend (3), using the perfusion medium of Leffert et al. (17). Briefly, rat liver was perfused with collagenase, minced, and incubated for 30 min. Cells were plated in RPMI 1640 (Gibco, Grand Island, NY) or in a hormonally defined medium (25); this medium consisted of RPMI 1640 (Gibco) supplemented with 100 U/ml penicillin, 100 μg/ml streptomycin, 50 ng/ml epidermal growth factor [ Collaborative Research, Inc., Waltham, MA], 30 μg/ml insulin, 10 μg/ml glucagon, 5 μg/ml linoleic acid/BSA [Miles Scientific, Naperville, IL], 20 μM/ml prolactin, 10 μM/ml growth hormone, 0.1 mM zinc, 0.1 μM copper, and 0.3 mM selenium [trace elements from Johnson Matthey, Inc., Malvern, PA] at a density of 5 × 10^6 cells/35-mm diameter plastic petri dish (Falcon Labware, Becton, Dickinson & Co., Oxnard, CA). Hepatocytes were maintained in a 37°C, 5% CO2 incubator and used within 5 h of plating for the studies reported here. Under these conditions hepatocytes survive for at least 1 wk in culture and after 24-48 h in culture can display near normal levels of tissue-specific mRNAs (25). Over the period of time considered here, no difference in electrophysiological properties of the cells (resting potential, input resistance, and gj) were detected, in contrast to the changes that occur a few hours later (see reference 37). We therefore do not distinguish among these cells in this study, except to specify the latency from time of isolation to fixation for electron microscopy.

For studies with multiple microelectrodes the wall of the culture dish was cut with a heated nichrome wire at a height just above the fluid meniscus. Studies were generally conducted at 21°C after rinsing the cells with phosphate-buffered saline (PBS; Gibco) to which Hepes (5 mM, pH 7.4) was added.

Solutions

Electrophysiological experiments were generally performed on cells pairs continuously rinsed (0.5 μl/min in a 1 ml vol dish) with a simple saline solution (Dulbecco's PBS; Gibco) containing (in millimoles) NaCl (137), KCl (2.7), CaCl2 (1.5), glucose (5.0), MgCl2 (0.50), to which was added 5 mM Hepes (pH 7.4). Solutions of octanol (0.2 M) were freshly prepared in this medium. Lucifer Yellow CH (5% wt/vol) was prepared for injection in 150 mM LiCl and microelectrodes were backfilled with this solution. All chemicals were obtained from Sigma Chemical Co. (St. Louis, MO), unless otherwise noted.

Fluorescent Microscopy

Fluorescence micrographs of plated cells used for physiology were taken with xenon arc lamp illumination using a Nikon B filter block (excitation filter 490-510 nm, emission filter 525 nm). Specimens were then frozen sandwiched between flat gold disks in liquid nitrogen–cooled Freon 22 and transferred to a double replica device (Balzers, Hudson, NH). Fracturing was carried out using a Balzers model 301 freeze fracture machine equipped with an electron beam gun for platinum shadowing and a quartz crystal monitor for standardizing replica thickness. Specimens were fractured and replicated at ~185°C; replicas were cleaned overnight in bleach, rinsed in water several times, and examined with a Philips 300 electron microscope.

Electrophysiology

Cell pairs (which comprised a sizable fraction of the cultured cells) were either impaled with microelectrodes (20-30 MΩ, filled with 3 M KCl or KCitrate) connected to homemade electrometers with active bridge circuits or were penetrated with low resistance unpolished patch type electrodes (∼5-10 MΩ, filled with 150 mM KCl, 10 mM Hepes, 10 mM EGTA [pH 7.4]) connected to voltage clamps (see reference 45 for circuit diagrams and further procedural details). With bridge electrometers, electrode resistance was balanced before cell entry and was adjusted as necessary after penetration by neutralization of the fast initial resistive phase in the voltage trace in response to a 30-ms current pulse. For whole cell voltage clamp recordings, cells were slightly indented by the patch type electrode and gentle suction was applied to the rear of the electrode while monitoring in the voltage clamp mode the current produced by 1 mV, 30-ms pulses applied to the pipette. After a high resistance seal (∼4 MΩ) was formed, access to the cell interior was obtained with brief strong suction. Patch clamp circuits were then switched to the current clamp mode and series resistance was compensated using the criteria for bridge balance mentioned above. Finally, circuits were switched back to the voltage clamp mode for recording cell responses. Under current clamp conditions, current pulses were applied alternately to the two cells and conductances of junctional and nonjunctional membranes were calculated by applying the pi-tie transform to measured input and transfer resistances (32). Under whole cell voltage clamp, junctional current is passed by the clamp on one cell in response to a voltage command in the other; gj is the junctional current divided by the transjunctional voltage (see reference 32).

pH-selective Microelectrodes

Microelectrodes were pulled from fiber-containing glass, and, if filled with 3 M KCl, would have had resistances of 20-30 MΩ. Tips were silanized by introducing 1 μl 10% dichlorodimethylsilane (vol/vol in CCl4) into the back of the electrode and then baking tip up in a vacuum oven at 200°C for 30 min. Ion exchange resin (neutral carrier proton exchanger from Pluka AG, Buchs, Switzerland) was then introduced into the electrode shank and the tip quickly filled. pH electrodes were backfilled with 150 mM potassium citrate buffer with citric acid to pH 6.5 and were calibrated before and after each experiment by measuring voltage response to immersion in buffered 150 mM KCl solutions in the range pH 6.0-7.5. Electrode response to a 1 pH-unit change was 90% of the final value within a few seconds; slope was between 55 and 58 mV/pH unit. pH was generally recorded differentially with respect to the voltage electrode in the same cell. Adequacy of intracellular placement of the ion selective electrode was established by comparison of steady-state responses of the voltage and ion selective electrodes in the same cell to a current pulse or voltage command (31). Reproducibility of pH measurements, judged from constancy of gj-pH curves, was within 0.02-0.04 pH unit.

Results

General Appearance of Dissociated Cells

Soon after dissociation, hepatocytes lose the prismatic appearance that characterizes them in reconstructions from thick sections (6) and in scanning electron micrographs of the tissue (20); freshly dissociated cells are round and quickly adhere to the bottom of ordinary tissue culture dishes. In our preparation, cell pairs (Fig. 1 a) and small clusters were common; since cells were in most cases used soon after isolation, the cell pairs we studied were most likely ones in which gap junctions survived the dissociation procedure. Alternatively or in addition, some gap junctions may be newly formed.
When Lucifer Yellow CH was injected into cells soon after dissociation (Fig. 1 b), the dye rapidly spread between or among coupled cells. Often dye spread nonhomogenously when injected into one cell of a group (not illustrated). In some cases, nonuniform dye passage was attributable to the larger sink provided by the extra cell in that direction; in other cases, coupled cells filled at different rates, suggesting that not all cells were coupled equally well. Nevertheless, rapid dye transfer was always (hundreds of injections) observed between pairs of freshly dissociated cells (for example, in one series of experiments comparing rates of dye transfer at 21° and 37°C, there was no failure of dye transfer in 20 cell pairs in each of three dishes at either temperature).

Electron Microscopy

In material fixed shortly after dissociation (1.5 h), pairs or small groups (up to about six cells) of hepatocytes are frequently observed (Fig. 1 c). Cells of such pairs contain one, and occasionally more, prominent euchromatic nuclei. The cytoplasm is dense and filled with mitochondria and rough endoplasmic reticulum with occasional lipid vesicles. Some of the free cell surface has numerous microvilli and presumably represents the area facing the space of Disse in vivo. The other free surface has cytoplasmic projections but fewer microvilli, and may have been appositional regions in vivo (note smooth area to the right of the right-hand cell in Fig. 1 c).

The appositional surfaces between freshly dissociated hepatocytes maintain most of the characteristics found in vivo, including intact bile canaliculi (upper central regions, Fig. 2, a and b). In both fractured (Fig. 2 a) and thin-sectioned (Fig. 2 b) specimens, microvillar projections into the canaliculi are flanked by tight junction networks (labeled T in Fig. 2 a, open arrow in Fig. 2 b). Small gap junctions are scattered within this network (arrowheads, Fig. 2 a); larger ones occur outside it (Fig. 2, c and d), merging with those beneath the tight junction network at the free surface. Where they are not filled with gap junctions, the interstices of tight junction networks are notably particle-poor. Gap junctions are less regularly surrounded by such particle-poor membrane (Fig. 2 c). Aside from variability in their size, the gap junctions are unremarkable. In P faces, gap junctions are composed of ≈7-nm particles irregularly arranged in plaques (Fig. 2 c),
and corresponding arrays of pits are found in E faces (not shown). The space or "gap" between membranes at thinned sectioned gap junctions is 2-4 nm and often is spanned by repeating densities (Fig. 2 d).

**Electrophysiological Properties**

Resting potentials of dissociated hepatocytes averaged \(-40\) mV (in one study using moderately high resistance \([30 \text{ M}\Omega]\) electrodes, resting potentials averaged \(-39.8 \pm 1.5 \text{ mV SD}; n = 38\), similar to values reported from liver cells in vivo \((5, 9, 20)\). Injection of a current pulse into one cell of a pair showed that the cells are electrotonically coupled (Fig. 3 a), also similar to the situation in vivo \((4, 20, 24)\). The voltage in the injected cell appeared with a time constant \(\sim 10-20 \text{ ms}\) and a resulting voltage deflection in the second cell that was generally 20-60% or more of that observed in the first. Typically, input conductances measured with ordinary microelectrodes were in the range \(20-50 \text{ nS}\) and \(g_i\) was \(0.1-1 \mu\text{S}\) (in one extensive series of experiments, \(g_i\) measured from cell pairs in sister cultures within 5 h of dissociation averaged \(0.9 \pm 0.1 \mu\text{S SD}; n = 45\)).

Junctional current is directly measured in dual voltage clamp experiments (Fig. 3 b) as the current that appears in one cell \((I_2\) in these records) in response to a voltage command in the other cell (see Materials and Methods). Voltage

Figure 2. Ultrastructure of ap positional areas. Freeze-fracture (a) and transmission (b) electron micrographs of equivalent canalicular and appositional regions between hepatocytes. Microvilli (M) extend into canalicular space. Tight junction ridges (T in a, arrow in b) and interspersed gap junction particle plaques (arrowheads) stand out in the P face of the appositional membrane in a, and a large gap junction (G) joins the appositional membranes of hepatocytes in b. Enlargements of hepatocyte gap junctions showing aggregated 7-nm P-face particles in fracture (c) and parallel unit membranes with 2-4 nm intercellular gap spanned by subunits in thin section (d). Bars, 0.1 \(\mu\text{m}\).
dependence of \( g_j \) was evaluated by passing long \( V_1 \) pulses of varying amplitudes (to 50 mV) and both polarities. During such voltage pulses (in more than 50 experiments on freshly dissociated hepatocytes from 12 rats), \( I_2 \) was constant and its magnitude was a linear function of \( V_1 \) (Fig. 3 b); \( g_j \) in hepatocytes is thus not appreciably affected by trans junctional voltage. There is also no dependence of \( g_j \) over the normal range of holding potentials (generally 0 to -40 mV) as is demonstrated in the current-voltage curves showing junctional and nonjunctional conductances for a pair of coupled cells (Fig. 3 c). The linearity of the current-voltage curve of the injected cell indicates that the nonjunctional membrane is not appreciably voltage dependent. Linearity of junctional and nonjunctional membranes is also suggested from in vivo studies (20), although the measurements are complicated by the geometry of the tissue.

Electrotonic coupling in several systems is reduced by extracellularly applied octanol (see references 15, 39 and 45). To determine whether this responsiveness is shared by hepatocyte gap junctions, 0.1 or 0.2 mM octanol dissolved in PBS was superfused over the cell pairs while coupling was measured (Fig. 4). Calculated \( g_j \) (plotted as connected dots, in the graph in Fig. 4) precipitously declined (by 80% over 15 s), while nonjunctional conductances (triangles) were not appreciably affected. Rinsing the preparation with PBS slowly restored \( g_j \) toward normal values (recovery to 75% within 3 min, Fig. 4; recovery to 95% within an additional 5 min, not shown here). The experiment illustrated here is typical of 24 responses of cell pairs from six dishes to superfusion with 0.1-0.2 mM octanol; in all cases \( g_j \) was abruptly reduced to <5% its initial value and in each case the effect was largely reversed by rinsing.

These data demonstrate that hepatocyte gap junctions are affected by a reagent that acts in other systems. Two other agents widely reported to affect \( g_j \) are cytoplasmic levels of H and Ca ions (see references 27, 33, 38, and 30 for review). To determine the extent to which \( g_j \) in hepatocytes is pH-dependent, \( g_j \) was measured in pairs of cells using either current injection (Fig. 5) or voltage clamp (Fig. 6), and pH was simultaneously monitored with a pH-sensitive microelectrode in one cell of the pair. Normal pH was 7.1-7.4 (mean 7.2 in 45 measurements) on cells in 18 dishes in Hepes-buffered medium (pH 7.4) but was generally considerably higher (7.8-8.1; measured in 18 cells in three cultures) in bicarbonate-buffered medium at ambient pCO2 levels (extracellular pH, 8.2); this finding is consistent with others in intact tissue (5, 16).

To acidify the cells, medium equilibrated with 100% CO2 was washed over the cells at various flow rates, allowing control of the rate and extent of acidification. In one experiment (Fig. 5) performed with current injection so that the coupling is directly observed, \( \text{pH} \) was reduced from 6.9 to 6.2 by exposure to CO2-equilibrated saline (between the arrows). Coupling between cells was markedly reduced; application of the pi-tee transform to input and transfer voltages showed that \( g_j \) was reduced by 95% while nonjunctional conductances were little affected (plot beneath graph). In the experiment shown in Fig. 6, performed under voltage clamp so that junctional current is directly observed and possible effects of potential are circumvented, the cell was initially at pH 7.8 in a bicarbonate-buffered medium. Exposure to 100% CO2 at moderate flow rate reduced \( \text{pH} \) to ~6.9 with no change in \( g_j \), and very little effect on conductance of the nonjunctional membrane. Flow rate was then increased and the cell
Figure 4. Reduction of coupling by octanol. (Top) Addition of 0.2 mM octanol to the medium (time of application indicated by bar at top of figure) reduced electrical coupling. Currents (I) are pairs of pulses applied alternately to cells 1 and 2. Edges of dark regions of the voltage responses (V1 and V2), transfer voltages associated with pulses in the other cell. Uncoupling in V1 and V2, increase in input voltage and decrease in transfer voltage. (Bottom) Conductances of junctional (●) and nonjunctional membranes (▲, ▼) show that gi is rapidly reduced by octanol and slowly recovers as the drug is washed away; nonjunctional conductances are not strongly affected.

Figure 5. Uncoupling of hepatocytes by acidification: current clamp recording. Display as in Fig. 4 except voltage of pH electrode is also shown. Application of solution equilibrated with 100% CO2 (between the arrows at the top of the figure) increased cytoplasmic acidity (VpH, increased acidity upwards) and decreased electrical coupling (the transfer voltages in traces V1 and V2 decrease and the input voltages increase). Plot of junctional conductances (gj, ●) and pH, (H, scale on right) show that both decrease upon CO2 exposure and recover on rinsing. Nonjunctional conductances (▲, ▼) are stable. The low pH at the beginning of this experiment is due to incomplete recovery from a prior acidification by CO2.
acidified further, reaching a final pHi value of ~6.3. Junctional currents were constant until pHi reached ~6.5 and then decreased by ~90% between pH 6.5 and 6.3. The non-junctional conductance of cell 2 also decreased over this pH range but there was little change in the nonjunctional membrane of cell 1. Decreasing the flow rate of the CO2-containing medium resulted in rapid recovery of pHi to ~6.8 and recovery of gJ to control values. Rapidly flowing CO2 reduced gJ by 90% in each of the more than 40 experiments on 16 cell pairs in which it was tested; gJ was largely restored in each case by rinsing in normal saline.

The relationship between gJ and pHi during acidification and recovery was similar in eight experiments (each representing several acidification–recovery cycles) in which it was quantified (three of which are shown in Fig. 7). As was evident in the experimental data of Figs. 5 and 6, gJ is constant above pH 6.6 but between 6.6 and 6.3 it steeply declines so that for pH values more acidic than 6.2 no detectable gJ remains. These values of gJ at pHi values obtained during acidification are fit by a form of the Hill equation (solid line) with an apparent pK of ~6.45 and a Hill coefficient (n) of ~8 (see Discussion). This value for n is the smallest that would satisfactorily encompass points on the upper and lower phases of the decline in gJ; higher values (to 11) also seem to adequately describe the pHi–gJ relation for hepatocytes.

Measurements of gJ during acidification were generally the same for given pHi values as seen during recovery (Figs. 5, 6, and 7). In two experiments, where pHi changes were rapid and brief, the recovery of gJ became slower and in successive CO2 exposures recovery of gJ lagged the recovery of pHi.

Discussion

Morphological Comments

Pairs and groups of hepatocytes examined soon after the dissociation procedure maintain many of the differentiated structures characteristic of the adult liver. Presumably these cells have been only partly dissociated: some of the appositional surfaces are disrupted, as indicated by the presence of junctional remnants, while other surfaces maintain junctional contacts and specializations seen in vivo (20, 46). The identifiable microvillar free surfaces reminiscent of the intact hepatocyte lining the space of Disse, the correspondence of small microvilli-lined bile canaliculi, and the elaborate junctional network surrounding both support this interpretation.
Figure 7. Correlation of \( g_j \) with \( pH_i \). The relation between normalized junctional conductance \( (G_j) \) and decreasing \( pH_i \) in two voltage clamp experiments (\( O, \bullet \)) and two current clamp experiments (\( \Delta, \Delta \)); points are averages from several acidification–recovery cycles obtained by \( CO_2 \) exposure followed by rinsing with normal saline. The solid curve is a plot of the Hill equation \( (G_j = K^*/(K^* + H^*)) \) where \( G_j \) is normalized \( g_j \), \( H \) is concentration of \( H^+ \) ions, \( K \) is the apparent dissociation constant, and \( n \) is the Hill coefficient. In this plot, fit by eye to the data, \( n = 8 \) and \( K \) corresponds to a \( pK \) of 6.45. The dashed curve is a plot of the Hill equation where \( n = 6 \) and \( K \) corresponds to an apparent \( pK \) of 6.38. Inset, \( G_j \) values during acidification (\( \nabla \)) and recovery (\( \Delta \)).

and suggest that, at early times after dissociation, neither gap junction disappearance associated with regeneration (46) nor random re-association between hepatocytes (7) has yet occurred.

Comparative Considerations of Physiological Properties

The lack of voltage dependence of \( g_j \) in hepatocytes is similar to that found between a number of other vertebrate and invertebrate cell pairs (13, 39, other data reviewed in reference 30). Gap junctions in these tissues thus differ from those in which \( g_j \) is dependent on transjunctional voltage or the potential between the inside and outside the gap (22, 32, 34, 39, other data reviewed in reference 30).

Hepatocyte gap junctions are sensitive to extracellularly applied octanol, as are gap junctions between crayfish septate axons (15), pairs of cardiac myocytes (45), and squid, amphibian, and fish embryonic cells (39). The mechanism of this action is unclear, but is specific for octanol and heptanol (ethanol at 20 mM has no effect in any of these systems).

The resting \( pH_i \) reported here is consistent with that obtained by others applying nuclear magnetic resonance and \( pH \)-sensitive microelectrodes to intact perfused liver (e.g., 5, 16). As was previously pointed out, high resting \( pH_i \) in the face of the very high \( pCO_2 \) of the portal circulation implies that buffer capacity of the cells must be enormous and that active \( H^+ \) extrusion could also play an important role in tissue homeostasis (5, 16). One candidate for such an extrusion mechanism is the Na/H antiport; indeed, these cells acidify profoundly when exposed to amiloride, which inhibits Na/H exchange (2).

The \( pH \) dependence of \( g_j \) in hepatocytes is markedly different from that in other tissues, where small changes in \( pH_i \) from the resting level may alter \( g_j \) (33; sensitivities of various gap junctions to \( pH \) are reviewed in reference 41). For liver, the \( pK \) is so low that \( pH \) gating is unlikely to be relevant to this gap junction under normal conditions. This low apparent \( pK \) possibly explains the previous report of lack of sensitivity of coupling within the intact liver exposed to \( CO_2 \) (19), a treatment likely to acidify cells to a lesser extent in the whole tissue than in culture.

In other tissues, a role for Ca ions has been suggested in control of junctional conductance, but in no case has the sensitivity of \( g_j \) to Ca been shown to be within a physiological range (27, 38, 40). Lack of sensitivity of hepatocyte gap junctions to Ca has recently been demonstrated in an elegant
study of the role of intracellular Ca and cytoplasmic contractile elements in the control of canalicular movement (44). In these experiments Ca injection sufficient to cause massive canalicular contractions did not block dye transfer through gap junctions located nearby. Our experimental results with Ca-selective microelectrodes indicate that Ca levels at least as high as 10 μM (obtained with Ca ionophore and vasopressin application) do not reduce gj (Spray, D. C., and J. C. Saez, unpublished observations) and are consistent with reported insensitivity in other systems.

**Implications for Biochemical and Biophysical Studies**

It seems reasonable that the pK and Hill coefficient contain information that is relevant for gap junction structure, since the closing of the channel at low pH presumably involves the binding of n or more protons to a site characterized by the apparent pK (33; while it seems likely to us that the binding site is an intrinsic portion of the channel molecule, we acknowledge that in most tissues action of a channel-associated modulatory molecule has not been excluded; see for example references 14 and 30 but see also 36).

The Hill coefficient may indicate the degree of cooperativity among subunits that comprise the channel. Presumably all gap junctions are dodecamers of subunits, hexamers being contributed by each cell. If there is one H+-binding site per subunit and occupation of all sites are required for channel closure, with maximum cooperativity n should be 12 (or 8 [the square of the Hill plot, see reference 33] if there are two independent hexamers in series, either of which can gate the channel). For n, n > 8, consistent with a model where hexamers are arranged in series, conformational change of either hexamer causing channel closure. Alternatively, binding to 12 sites may be required for closure but with lower cooperativity (low cooperativity is apparently the case for other gap junctions regardless of closure mechanism, where values of n range from 1 to ~5 [see references 33, 45, and 41 for a review]).

Conductance of all gap junctions examined except lens is pH dependent although the slope and pK of the pH–gj relations vary considerably from one tissue to another (see reference 41 for a review). In three tissues (lens, liver, and heart) partial sequences of the apparent junctional proteins are

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