Ouabain Modulates the Distribution of Connexin 43 in Epithelial Cells

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Key Words
Ouabain • Cnx43 • Gap junctions • MDCK cells • c-Src and ERK1/2

Abstract
Background/Aims: The fact that ouabain has been identified as an endogenous substance, led us to inquire its physiological role in epithelial cells. Based on previous observations, we hypothesized that it influences processes related to cell contacts. Previously we have shown that nanomolar concentrations of ouabain up-regulate tight junctions, accelerate ciliogenesis, and increase gap junctional intercellular communication (GJIC). Given that silencing assays indicated that connexin 43 (Cnx43) is involved in the GJIC response, in the present work we study whether ouabain affects Cnx43 expression and distribution. Methods: We seeded confluent monolayers of epithelial renal MDCK cells and incubated them with 10 nM ouabain during 1 h. Then we measured, by densitometric analysis of Western blot assays, the amount of Cnx43 in cells and in fractions enriched of plasma membrane. We also studied its localization with immunofluorescence and confocal microscopy. Results: Cnx43 is remarkably displayed, outlining the borders of cells gathered in clusters, randomly scattered throughout the monolayer. Ouabain increases the density of such clusters, as well as the average number of cells per cluster, without inducing the synthesis of new Cnx43. It also promotes relocation towards the membrane, of subunits already available. The fact that such changes are inhibited by PP2 and PD98059 indicates that a signaling pathway, that includes c-Src and ERK1/2, is involved in this response. Conclusion: Ouabain induces the translocation of Cnx43 from the cytoplasm to the plasma membrane. These findings support our hypothesis that one of the physiological roles of ouabain is the modulation of physiological processes that depend on cell to cell contacts.

Introduction

Ouabain is a highly toxic compound of remarkable interest on cellular physiology and medicine. Originally obtained from Acokanthera ouabaio and Strophanthus gratus, both A. Ponce and I. Larre shared first authorship.
plants native to eastern Africa, it was traditionally used as an arrow poison for hunting and warfare [1]. Also known as g-strophanthin, ouabain belongs to a group of compounds known as cardiotonic steroids, because of its chemical structure, and its influence on heart function. When administered at toxic dosages it produces heart failure and death, nonetheless, at controlled dosages has been used therapeutically to treat heart arrhythmias and hypotension [2, 3]. A substance identical to the plant-derived ouabain was reported to be endogenously present in the plasma of some mammalian species, including humans [4]. Further studies have shown endogenous ouabain to be produced in the adrenal cortex and hypothalamus [5, 6]. These findings have led to regard ouabain as a hormone [7, 8], whose physiological role is still not fully understood.

Ouabain binds to the Na+/K+-ATPase, blocking its pumping function. This leads to an increase of intracellular sodium, which in turn reduces the activity of the Na+/Ca2+-exchanger, producing an increase of intracellular calcium. This scheme accounts for the effect of ouabain on the heart because it results in higher cardiac contractility and an increase in cardiac vagal tone [9, 10]. Recently it has been found that Na+/K+-ATPase also acts as a receptor of ouabain [11, 12]. Binding of ouabain to Na+/K+-ATPase unleash a cascade of interrelated signal transduction pathways that involves inositol (3, 4, 5)-tris-phosphate receptor; Src kinase, tyrosine phosphorylation, epidermal growth factor receptor, Ras and the Ras/Raf/MEK/ MAPK signaling pathway [13-15]. Following this scheme, ouabain has shown to influence a wide variety of physiological processes, including proliferation [16, 17], growth [18], apoptosis [19-21] and cell mobility [22] in different tissues and organs. It also has been shown to influence the regulation of salt-sensitive blood pressure [23], salt handling in the kidney [24], vascular tone, Na+ homeostasis [25] and embryonic kidney cell’s survival during malnutrition [26].

We have studied the effect that ouabain, in a nanomolar dose (10 nM), has on the physiology of epithelial cells. Using MDCK epithelial cells as a model, we have shown that it influences several processes related to cell-cell contact [27]: (1) The hermeticity and the molecular composition of tight junctions [28], a fundamental component of epithelial transporting phenotype [29]; (2) It accelerates ciliogenesis, a feature requiring close cell-cell contacts and that is a visible sign of polarity [30]. More recently we have focused on its effect on Gap Junctional Intercellular Communication (GJIC).

Gap junctions are molecular assemblies between neighboring cells, which by this mean exchange small molecules, such as ions, second messengers and metabolites [31, 32]. Gap junctions are found in cells of almost every organ and tissue, but are particularly abundant in tissues where a coordinated response is required, as for instance in cardiac and smooth muscle or in endocrine glands. Gap junctions are displayed as plate-shaped structures between cells. Such plates are clusters of intercellular channels, formed by head-to-head docking of two hemichannels or connections, contributed by each of two neighboring cells. Connexons are in turn formed by six tetraspan membrane proteins known as connexins (Cnx) [33, 34]. There exist a variety of distinct types of connexins, which are differentially expressed among tissues of vertebrate species to form homomeric or heteromeric connexons. The human genome contains 21 connexin isoforms, whereas the mouse has 20 [35]. Connexins are named by its molecular weight, for instance Cnx32 represents the connxin protein of 32 kDa. From among the different types Cnx43 is of striking interest because it is one of the most abundantly expressed in a variety of organs and tissues in mammals, besides that its expression has been described to be related to a variety of cellular processes such as cellular proliferation and apoptosis [36-39].

To date little is known about the effect of ouabain on GJIC. It has been reported that 100 μM ouabain produces no effect on GJIC of primary cultures of rat Sertoli cells and cardiac myocytes [40], whereas it caused a significant degree of junctional uncoupling in isolated rabbit corneal endothelium [41]. Ouabain (1 mM) has been shown to inhibit GJIC in rat aortic A7r5 smooth muscle cells, monkey COS-1 fibroblasts and human HeLa epithelial cells [42]. An effect that has been shown to depend on localized changes in [Ca2+] (i) through modulation of Na+/Ca2+-exchanger activity [43].
In a previous work we showed, by dye transfer assays and electrical capacitance measurements, that ouabain 10 nM enhances GJIC up to 475% within 1 hour in MDCK cells [44]. The fact that such response occurs quickly (after 10 minutes of addition of ouabain), and that it is not affected by cycloheximide or Actinomycin D suggest that ouabain does not induce the synthesis of new connexins in order to enhance GJIC but rather promotes the relocation of subunits already synthesized. Using silencing assays, we also demonstrate that this ouabain-induced enhancement of GJC involves Cnx43.

Therefore, in the present work we resorted to immunohistochemical methods to elucidate how ouabain influences the expression and distribution of Cnx43.

Materials and Methods

Cell culture

Starter MDCK-II cell cultures (Canine renal; American Type Culture Collection, CCL-34). The cells were grown at 36.5°C in a 5% CO₂ atmosphere in Dulbecco’s modified Eagle medium (DMEM; life technologies, Carlsbad, CA, USA) supplemented with penicillin-streptomycin 10,000 U/µg/ml (In Vitro, Acayucan, México) and 10% fetal bovine serum (GIBCO). This medium referred to as CDMEM. The cells were harvested with trypsin-EDTA (In Vitro) and seeded on glass coverslips placed at 24-well multi-dishes (3524; Costar Corning, NY, USA) for immunofluorescence and Western blot analysis.

Ouabain challenge protocol

Cells were cultured at a saturating density of ~70% and maintained for one day in CDMEM, then kept for 24 h in DMEM containing 1% fetal bovine serum before treatment with or without 10 nM ouabain (O-3125; Sigma-Aldrich, St Louis MO, USA). For signaling studies, monolayers were exposed to either 25 µM PD98059 (513000; Merck Millipore, Darmstadt GE), an inhibitor of MEK1/2 kinase that impairs the activation of the Extracellular Regulated Kinase 1/2 (ERK1/2), or 10 µM PP2 (529573; Merck Millipore), an inhibitor of c-Src kinase. In both cases, those inhibitors were added 1 h before ouabain challenge.

Immunofluorescence

Monolayers on coverslips were washed three times with ice-cold PBS containing Ca²⁺, fixed and permeabilized with methanol for 8 min at -20°C, washed three times with PBS, blocked for 1 h with 0.5% bovine serum albumin and incubated overnight at 4°C with rabbit anti-Cnx43 antibody (71-0700; Invitrogen) or mouse anti-α1-Na⁺/K⁺-ATPase (MA-3-929; Thermo Fisher Scientific, Waltham, MA, USA). The samples were washed, incubated with secondary antibody (goat anti-rabbit-FITC, 65-6111, Thermo Fisher Scientific), or goat-anti-mouse-TRICT (115-123003; Jackson Immunoresearch, PA, USA), rinsed, incubated with 1 µg/ml DAPI (Sigma-Aldrich, St Luis, MO, USA) in PBS for five minutes, mounted in vectashield (H-1000; Vector Laboratories, Burlingame, CA, USA), and examined by confocal microscopy (SP8, equipped with Plan-NeoFluar 63x NA 1.4 objective, Leica Microsystems, Wetzlar, Ger). Sampled images were analyzed with the software Las AF (Leica) and FIJI (FIJI is Just Image J), imported into GIMP (GNU is not Unix Image Manipulation Program) to adjust brightness and contrast and to construct Figures.

Immunoblot

Cell monolayers were washed three times with ice-cold PBS with Ca²⁺ and then incubated at 4°C for 10 min with lysis buffer (24948; Santa Cruz catalog, CA, USA), supplemented with a protease inhibitor cocktail (Complete™ 469311600; Sigma-Aldrich), for protein extraction. These protein extracts were then centrifuged for 10 min at 17,000 g and the supernatant recovered to measure total protein content with the Micro BCA™ assay (23235; Thermo Fisher Scientific). An equal volume of 2x Laemmli sample buffer (161-0737; Bio-Rad, Hercules, CA, USA) was then added to the protein extracts, that were subsequently resolved by SDS-polyacrylamide gel electrophoresis (PAGE), and transferred to PVDF sheets (Hybond-P, Sigma-Aldrich) with the help of a Transblot Turbo (Bio-Rad), following the instructions of the manufacturer. These sheets were blocked for 10 min with 0.05 % bovine serum albumin in Tris Buffer Saline (TBS, 10 mM Tris-HCl, 100 mM NaCl) supplemented with 0.1 % Tween-20 (TBS-T), incubated with a rabbit-anti-Cnx43 antibody in TBS-T, supplemented with BSA 0.5 %, for 20 min at room temperature, washed with...
TBS-T and then incubated with a HRP-goat-anti-rabbit (G-21234; Sigma-Aldrich). The blotting procedure was performed with the help of SNAP id 1.0 (Millipore, Billerica, MA, USA) following the instructions provided by the manufacturer. Bands were resolved by chemiluminescence (RPN2232 ELC, GE Healthcare Life Sciences, Malborough, MA, USA) using Hyperfilm (GE Healthcare Life Sciences). Then membranes were stripped by incubation (30 min at 37 °C with gentle shaking) in 3% sodium hydrosulfite (157953; Sigma-Aldrich) in TBS-T, and incubated with mouse anti-actin (a kind gift from Dr. José Manuel Hernández from the Department of Cell Biology, Cinvestav) or rabbit-anti-gp135 (a generous gift from Raya A., Department of Physiology, Cinvestav México City), depending on the experiment, and processed as described above. Densitometric analysis was performed with Kodak 1D 3.5.4 software (Eastman Kodak; Rochester, NY, USA). Cnx43 densitometric measurements were related to those from actin or gp135. For the purpose of comparison, the corrected optical density of monolayers under control condition was arbitrarily standardized with a value of 1.

Plasma membrane protein extraction
Plasma membrane proteins were extracted using a kit obtained from Abcam (Plasma membrane extraction kit ab65400, Abcam, Cambridge, UK). Briefly: all procedures were developed at 4°C. Monolayers were scraped from 100 mm Petri dishes (four for each condition), resuspended in 8 ml of PBS, then centrifuged (1057 g, 5 min). Precipitates were re-suspended in 2 ml of lysis buffer (provided by the manufacturer), then homogenized with 150 strokes in a Dounce Homogenizer (cat. D8938, Sigma-Aldrich) with ice and centrifuged at 700 g for 10 min at 4°C. The supernatant was then centrifuged in a new vial for 30 min at 10000 g and the precipitate was further processed to perform a step gradient plasma membrane extraction. The precipitate is resuspended in an upper phase solution (provided by the manufacturer) and an equal volume of the lower phase solution of higher density (provided by the manufacturer) is then added. After gently shaken, the extract is centrifuged 1000g for 5 min, the superior phase is recovered and the remaining lower phase is mixed again with an upper phase solution to repeat the extraction. Both superior phases are combined and the step gradient plasma membrane extraction repeated once more. The final upper phase was diluted with 5 volumes of water and centrifuged at 17,000 g for 10 min. The pellet is re-suspended in TX-100 0.1 % in water. These plasma membrane extracts were then processed by immunoblot.

Analysis of Cnx43 and Na+/K+-ATPase colocalization
To estimate the co-localization of Cnx43 with the Na+/K+-ATPase of the plasma membrane, we used the LAS-AF software (v 2.30, Leica). We first captured 10 images for monolayers cultured 1 hour in control conditions or with 10 nM ouabain. Then we selected a region of interest in each image, choosing the membrane labeling, identified by the “chicken fence” like pattern, and avoiding the cytoplasmic label as much as possible. The signals of both channels were overlaid to reveal co-localized pixels threshold (30 %) and background (20 %) level, based on the “Red-Green” or “Red-blue” scattergram offered by the LAS-AF software (Leica). The co-localization rate (%) was calculated with the same image analysis software (Leica, % co-localization rate = co-localization area/area foreground; area foreground = area image - area background). The average of data obtained from 10 images was considered a value of a single independent experiment.

Statistical analyses
Statistical analyses were performed using Prism 5 (GraphPad software). The results are expressed as the means ± standard error. Statistical significance was estimated via one-way analysis of variance (ANOVA) followed by Bonferroni’s multiple comparison or Student’s t-test and was denoted as follows: * P<0.05, ** P<0.005 and *** P<0.001 and n is the number of independent experiments.

Results
Ouabain does not promote synthesis of Cnx43
Our previous findings, that 10 nM ouabain triggers a quick enhancement of GJIC, plus the fact that this response is not impaired by inhibitors of RNA or protein synthesis [44] suggest that no synthesis of connexins are required to achieve this effect. To test this
**Fig. 1.** Ouabain does not increase the expression level of Cnx43. (A) Cell lysates were prepared from MDCK monolayers, either control or incubated with 10 nM ouabain for 1 hour, then equal amounts of total cell protein were subjected to SDS-PAGE and western blotting with anti-Cnx43 antibodies. In both cases a multi-band pattern was obtained. The blots were then stripped and re-probed with anti-actin antibodies. (B) The Cnx43 signal density of gels, as shown in (A), was measured and normalized with to actin label. The statistical analysis shows non-significant difference between control and treatment. “n” stands for the number of assays analysed.

**Fig. 2.** Ouabain changes the distribution pattern of Cnx43. (A, B) Representative images showing the pattern of distribution of Cnx43 in MDCK monolayers, under (A) control conditions, or (B) after one hour incubation with 10 nM ouabain. (A) Under control conditions, many cells express Cnx43 signal (green) as a faint borderline, suggesting basolateral localization (white arrowheads), although intracellular staining is also observed (yellow arrows). Clusters of cells expressing a more intense Cnx43 signal are clearly noticed (red arrow). (B) Ouabain increases the density of these clusters, as well as the number of cells that form them. (C) Statistical analysis of the density of the clusters of cells expressing Cnx43, as a function of time, after ouabain challenge. The error bar in the 0 time point is smaller than the symbol. (D) Comparison of the number of cells per group in control and ouabain-treated monolayers. (*) indicates a statistically significant difference (p<0.05) between control and treatment groups. “n” stands for the number of images observed.

In hypothesis, we compared, by densitometry analysis of Western blots, the protein level of Cnx43 from total protein extracts obtained from MDCK monolayers, without (control) and after 1 hour of treatment with 10 nM ouabain. Figure 1A shows representative blots of both cases. The two of them exhibit a typical pattern, consisting in several bands, due to distinct phosphorylated states [45-48]. Figure 1B shows the corresponding statistical analysis of densitometric measurements, normalized with actin. The fact that no significant difference in the density of control and treatment batches supports further the hypothesis that ouabain per se does not increase the expression level of Cnx43, at least within one hour of treatment.

**Ouabain induces changes in the pattern of distribution of Cnx43**

Next, we investigated whether ouabain enhances GJIC by promoting changes in the distribution of Cnx43. Figure 2 shows the pattern exhibited by Cnx43 in the absence (control, 2A) and in the presence of 10 nM ouabain (Fig. 2B). In the former case, Cnx43 is expressed all over the monolayer, where cells tend to express Cnx43 as a faint chicken fence-like pattern, characteristic of a basolateral distribution (white arrows), although abundant intracellular staining is also observed (yellow arrows). It is strikingly noticeable the presence of clusters of cells, randomly scattered throughout the monolayer, where Cnx43 is more intensely expressed (red arrow). Treatment with 10 nM ouabain produces an increase in the density of clusters of cells expressing Cnx43, as shown in Figure 2B. This increase is statistically significant (p<0.05), as shown in Figure 2C. The number of clusters per group in control and ouabain-treated monolayers was compared, and the results are shown in Figure 2D. The presence of clusters of cells expressing Cnx43 in the ouabain-treated monolayer is statistically significant (p<0.05) compared to the control group.

**Fig. 3.** Ouabain induces changes in the pattern of distribution of Cnx43. (A) Cell lysates were prepared from MDCK monolayers, either control or incubated with 10 nM ouabain for 1 hour, then equal amounts of total cell protein were subjected to SDS-PAGE and western blotting with anti-Cnx43 antibodies. In both cases a multi-band pattern was obtained. The blots were then stripped and re-probed with anti-actin antibodies. (B) The Cnx43 signal density of gels, as shown in (A), was measured and normalized with to actin label. The statistical analysis shows non-significant difference between control and treatment. “n” stands for the number of assays analysed.
those clusters of cells (Fig. 2C) which is significant from as early as 5 minutes after addition of ouabain 10 nM to external media. Ouabain also causes a significant increase of the average number of cells per cluster (Fig. 2D).

Ouabain promotes relocation of Cnx43 to the membrane. We noticed that the Cnx43 borderline pattern exhibited by cells, within clusters of monolayers treated with ouabain, appears more intense and delineated as compared to those from untreated monolayers. This suggests that ouabain promotes recruitment of Cnx43 subunits towards the basolateral membrane. To test such possibility, we estimated how close Cnx43 co-localizes with Na+/K+-ATPase, which was taken as a basolateral membrane marker. Figure 3 shows two representative images of the co-staining of Na+/K+-ATPase (red) and Cnx43 (green) in a control monolayer (Fig. 3A), and in a monolayer treated for 1 h with 10

Fig. 3. Ouabain increases the colocalization of Cnx43 with α-Na+/K+-ATPase. Monolayers contained with antibodies against Cnx43 (green) and α-Na+/K+-ATPase (red) cultured in control conditions (A) and with 10 nM ouabain added to the media for 1 hour (B). (C) Percentage of colocalization rate in images obtained from monolayers cultured in control (white bar) and 10 nM ouabain (red bar). (**) indicates a statistically significant difference (p<0.005) between control and treatment groups. "n" stands for the number of images analyzed.

Fig. 4. Ouabain increases Cnx-43 in plasma membrane. Protein extracts from plasma membranes were obtained from MDCK monolayers, either under control or 1 hour of 10 nM ouabain treatment. After SDS-PAGE, immunoblots were made to reveal Cnx43. (A) representative immunoblots. (B) densitometric analysis. (**) indicates a statistically significant difference (p<0.005) between control and treatment groups. "n" stands for the number of blots analyzed.

Fig. 5. Ouabain induces membrane relocation of Cnx43 through the activation of c-Src and ERK1/2. (A) (Above) Representative examples of western blot assays (Below) statistical analyses of the comparison of relative density of Cnx43/gp135 labels under control (white bar), ouabain (red bar) and ouabain plus PP2 (blue bar) treatments. The (*) and (**) signs indicate a statistically significant difference between the bars that indicate the horizontal lines. “n” stands for the number of assays. (B) Bars indicate de average (± S.E.) of the density of clusters of monolayers under control (white bar), as well as from: monolayers treated with 10 nM ouabain (red bar); monolayers pre-treated with PP2, then ouabain (blue bar) and monolayers pretreated with PD98059 before adding ouabain for 1 h (green bar). Statistical significance was estimated via one-way analysis of variance (ANOVA) followed by Bonferroni’s multiple comparison or Student’s t-test. (**) denotes P<0.005 and (***) P<0.001. “n” stands for the number of images analyzed.

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nM ouabain (Fig. 3B). Figure 3C shows the corresponding statistical analysis, indicating that in fact 10 nM ouabain promotes a significant displacement of Cnx43 towards the membrane. To further support this conclusion, we made Western blots assays from plasma membrane extracts obtained from MDCK monolayers (Fig. 4). A statistically significant difference in the protein density of Cnx43 between samples under control and treatment with 1 hour incubation of 10 nM ouabain was obtained. This difference was not observed, in contrast, on the density of gp135, a membrane protein that was therefore taken as standard. These results may be taken as further indication that Cnx43 is relocated to the plasma membrane.

Ouabain-promoted Cnx43 recruitment is regulated by a signaling route involving c-Src and ERK1/2

Extracellular Regulated Kinases 1 and 2 (or ERK1/2) are two related extracellular signal–regulated kinases that participate in the Ras-Raf-MEK-ERK signal transduction cascade. It is known that this cascade participates in the regulation of a large variety of processes that includes cell adhesion [48]. On the other hand, c-Src is a tyrosine kinase that is mutated in several cancers. We have previously found that ouabain mediates its effect on GJIC by a signal transduction cascade that includes c-Src and ERK1/2 [44]. To determine whether such signal transduction cascade is also involved in the relocation of Cnx43 induced by ouabain, we probed whether commercial inhibitors of c-SRC or ERK1/2 impairs any of the effects caused by ouabain, either the increase of Cnx43 protein density from Western blot assays of membrane extracts, or the increase of the density of clusters of cells observed in immunohistochemical assays. Figure 5A shows representative stains (above) and the statistical analysis that compares the densitometry analyses of Western blot assays made with membrane extracts of MDCK cells under control (white bar), treatment with 10 nM ouabain (red bar) and ouabain plus 10 µM PP2 (an inhibitor of c-Src kinase, blue bar). As shown previously, ouabain increased significantly (p<0.005) the relative density of Cnx43, but treatment with PP2 significantly inhibited this increase (p<0.05). Figure 5B shows the effect of those inhibitors on the density of clusters of cells with enhanced Cnx43 label. Monolayers treated with 10 µM PP2 had significantly reduced cluster density (p<0.05) as compared with those treated with ouabain only. Monolayers treated with 25 µM PD98059, a substance that impairs the activation of ERK1/2 [49], also show significantly reduced cluster density (p<0.01). Therefore, these results indicate that ouabain modulates the pattern of Cnx43 distribution via c-Src and ERK1/2.

Discussion

Ouabain, a substance originally identified in plant extracts and used as poison and medication, has received renewed interest since proved to be endogenously present in mammal species including humans. As part of our interest in the effect that ouabain, in a nanomolar range, has on epithelial (MDCK) cells, we evaluated in a previous work, how it influences gap junctional intercellular communication (GJIC) by dye transfer assays and by electrical capacitance measurements [44]. We found that it, triggers an enhanced GJIC response, that is significantly noticeable from 10 minutes after addition of ouabain to the bathing media and reaches a maximum at about one hour, followed by a slow decaying, then a sustained response that last, at least 24 hrs. The finding that such enhanced GJIC response is inhibited by PP2 and PD98059, indicates that ouabain activates a signaling cascade that includes activation of c-Src and ERK1/2. Further, in order to determine whether Cnx43 is involved in the GJIC enhancement that ouabain triggers in MDCK cells, we performed silencing assays and found that this procedure effectively impairs it, indicating that Cnx43 is a determinant player in such response.

Such effect of ouabain on GJIC can be accounted for, either by modulation on the gating properties of connexon units already assembled, or by an increase in the number of connexons available or both. In turn, an increase in the number of connexons may be
due to promotion of synthesis of new units, or to membrane recruitment of Cnx43 subunits previously synthesized. However, the fact that its significant response occurs as quickly as 10 min, along with the finding that this response is not impaired by Actinomycin D that blocks the synthesis of RNA, nor cycloheximide that inhibits the synthesis of proteins, suggest that it is due to the recruitment of subunits already in stock.

In the present work we performed immunofluorescence and immunoblots assays, focusing on Cnx43 because, as mentioned above, we found that Cnx43 is a participating subunit type in the ouabain enhanced GJIC. As we show by densitometric analysis of immunoblots, treatment with 10 nM ouabain for 1 hour does no produce a significant increase of Cnx43 label. This result supports the hypothesis that no synthesis of new Cnx43 is required at least for the rapid phase of GJIC triggered by ouabain, but rather to the relocation of previously synthesized Cnx43. However, this conclusion does not exclude, the possibility that synthesis of new subunits occurs in order to maintain GJIC at later times, or that ouabain could additionally modulate the gating of connexons already assembled.

On the other hand, it is interesting that Cnx43 is expressed in clusters of cells scattered over the monolayer and that ouabain increases the density of such clusters, as well as the average number of cells per cluster. The fact that a significant increase in the density of clusters occurs concomitantly with enhancement of GJIC, along with the finding that such response is impaired by PP2 and PD98059 as in the case of the enhancement of GJIC, suggest that both phenomena are related. It does not mean however that GJIC is a phenomenon exclusively occurring in cells belonging to clusters, because an enhanced GJIC may depend not only on more connexin subunits available at the membrane, but on a gating mechanism regulating the opening of gap junctions. In this regard, it is possible, as already mentioned above, that in addition to promoting relocation to the membrane of Cnx43 that enhances GJIC, ouabain would modify directly the gating of gap junctions already assembled. This would explain why ouabain enhances GJIC as early as 10 minutes after its addition to the bathing media, although we have not evaluated such possibility yet.

In summary, the present results further support the emerging role of ouabain on the regulation of cellular processes that depend on cell contacts. As we have shown, challenging MDCK cells with 10 nM, a concentration within its physiological range triggers a series of changes in the epithelial phenotype, which includes tight junctions, ciliogenesis and gap junctional communication. The effects of hormone ouabain start with the binding of ouabain to the α-subunit of Na+/K+-ATPase, that acts as a receptor [14, 15], which in turn, activates a signaling pathway involving c-Src and ERK1/2 [50]. This indicates that the same signaling route mediates all the hormonal effects on cell contacts so far tested. This includes changes of molecular composition of the tight junction [28], to the displacement of Cnx43 towards the plasma membrane implied in GJIC studied in our previous work [44]. Given that ouabain modulates both tight junctions and apical/basolateral polarity, which are taken as the most specific differentiated features of the epithelial transporting phenotype, we may say that ouabain is a hormone that modulates the expression of the transporting epithelia phenotype. We can even foresee a role of this hormone in clinical medicine, because 90% of all human deaths involve the collapse of an epithelium. Therefore, further research on hormone ouabain may lead to achievements of biomedical relevance.

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Disclosure Statement

We declare that we have no conflict of interest

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