Array analysis of gene expression in connexin-43 null astrocytes

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CONNEXIN-43 (Cx43, also termed gap junction alpha-1, GJA1) was the second gap junction protein whose cDNA was cloned and sequenced (3), which was accomplished by low-stringency screening of heart cDNA libraries with Cx32 cDNA. It is the most widespread gap junction protein in mammals, where it occurs in almost every tissue (3, 5, 6). Cx43 is the primary component of intercellular gap junction channels in ventricular myocytes of the heart (50) and in astrocytes (9, 43), the most abundant type of glial cells in the brain. Cx43 also occurs between neural progenitors in early development (14, 45, 46) and in leptomeningeal, ependymal, and vascular cells (10, 11).

Cx43-null mice die at birth due to a developmental cardiac abnormality, where hyperplasia blocks blood flow exiting from the right ventricular outflow tract to the lungs (44); other congenital abnormalities include coronary artery patterning defects (28). Brains of Cx43-null mice are grossly normal (8, 39), although it has been reported that migration of cardiac neural crest cells is altered compared with wild-type (WT) littermates (57) and that neocortical neuronal migration is delayed in Cx43-null mice (15).

Because of its requirement for proper cardiac function, alterations in Cx43 expression and/or function are associated with a large variety of cardiac arrhythmic disorders (27, 40), and mice deficient in Cx43 gene expression exhibit abnormal cardiac conduction and are prone to reentry (35). Alterations in Cx43 expression have also been detected in transcriptomic studies of Huntington’s and Alzheimer’s diseases (33, 54), in autoimmune inflammation (6, 32), and in astrocytomas (17). Mutations in other gap junction genes result in syndromic deafness (31), and oculodentodigital dysplasia (38); Cx43 mutations have also been reported in metastatic colon tumors (12).

Although the association of Cx43 missense mutations with the complex disorder heterovisceral atriocoxia has been controversial (4, 48), more recent studies have suggested that Cx43 mutations are responsible for hypoplastic left heart syndrome (7), rare cases of non-syndromic deafness (31), and oculodentodigital dysplasia (38); Cx43 mutations have also been reported in metastatic colon tumors (12).

One phenotypic difference between astrocytes cultured from Cx43-null mice compared with wild types is a profound reduction in their growth rate (8, 36). Previous studies on Cx43-transfected glioma cells had demonstrated that some highly expressing stable clones exhibited decreased proliferation, which has subsequently been attributed to the increased expression in Cx43 transfectants of genes encoding secreted growth suppressive molecules (see Ref. 37 for overview of these studies). Moreover, recent studies have suggested that the growth suppression in Cx43-transfected cells may be independent of the formation of intercellular gap junction channels (34, 42), although in some studies intercellular communication is re-
quired or enhances the expression of growth suppression molecules (16, 59).

Cx43 expression has also recently been linked to programmed cell death. Cx43 overexpression in human glioblastoma cells was shown to increase sensitivity of the cells to a variety of chemotherapeutic agents and under conditions of low serum, which was correlated with decreased expression of the anti-apoptotic bcl-2 protein (18, 20). However, transfection with Cx43 has also been reported to protect the cells from cell injury, either through a mechanism involving nonjunctional hemichannels or not requiring functional channels at all (29, 41).

A growing number of studies suggest that connexin proteins may play functional roles supplemental to their unique mediation of direct intercellular communication. In addition to effects on growth rate and apoptosis considered above, purinergic receptor expression is altered in Cx43-null astrocytes from brain and spinal cord (47, 52), and cell adhesion appears to depend on connexin expression but not on coupling strength (30). Although some of these effects may be mediated by connexin-protein interactions, which normally assemble a macromolecular complex (13, 49), it is also possible that connexin expression may affect cell phenotype through alterations in the expression of other genes. Indeed, differential mRNA display analysis has identified a number of gene alterations in Cx43-transfected glioma cells (35), and use of cytokine arrays has identified changes in certain proteins following Cx43 transfection (19). To test this hypothesis for a large number of genes, we have compared gene expression in sibling WT and Cx43-null mouse astrocytes using cDNA microarrays. Our findings indicate a surprisingly high degree of alteration in gene expression in astrocytes from Cx43-null mice, with many of the known genes on the arrays being statistically up- or downregulated. Moreover, a number of these genes encode proteins involved in cell cycle control and apoptosis, consistent with phenotypic differences in growth rate of Cx43-deficient astrocytes and with previous linkage of Cx43 expression to cellular sensitivity to apoptotic stimuli.

**MATERIALS AND METHODS**

The study was conducted according to standards developed by the Microarray Gene Expression Data Society (MGED), and data complying with the “minimum information about microarray experiments” (MIAME) have been deposited in the National Center for Biotechnology Information (NCBI) Gene Expression Omnibus (GEO) database, and can be accessed at [http://www.ncbi.nlm.nih.gov/geo/](http://www.ncbi.nlm.nih.gov/geo/) (platform GPL369, samples GSM8889 to GSM8896).

**RNA sources.** Heterozygous Cx43+/- mice (44), obtained from Jackson Laboratories (Bar Harbor, ME), in the C57BL/6j background, were interbred in our AALAC-accredited animal facilities to obtain WT and Cx43-null progeny. At birth, brains of individual mice were dissected free of meninges in cold PBS, minced, and dissociated in trypsin (0.1%). Cells were then resuspended in DMEM with 10% fetal bovine serum and 1% penicillin-streptomycin and seeded in tissue culture flasks. One week later, flasks were shaken overnight to remove less adherent cells. After trypsinization (0.05% trypsin), cells were plated in 100-mm culture dishes for 1 wk (WT) or 2 wk (Cx43 null), at which time they were confluent (about 225 cells/mm²). Medium was replaced every 2–3 days during culture. Genotype of each animal was determined by PCR of tail DNA as described (8). At 2 days after the last feeding, RNA was isolated using the TRIzol method (see [http://www.aecom.yu.edu/home/molgen/fungeneomic.html](http://www.aecom.yu.edu/home/molgen/fungeneomic.html)), parallel dishes were used for cell density determination and staining with glial fibrillary protein antibodies (Sigma polyclonal anti-GFAP, 1:500). Purity of cultures was 91–92% GFAP-positive cells; neurons and oligodendrocytes were absent as judged by morphology of GFAP-negative cells. No difference in percentage of GFAP-positive cells was detected between WT and Cx43-null astrocyte cultures (see Ref. 8).

Mouse reference RNA (hereafter denoted by R) was prepared at one time in sufficient quantity for the entire projected experiment from selected amounts of total RNA extracted from 10 adult male and female mouse tissues (aorta, brain, heart, kidney, liver, lung, ovary/testicles, spleen, and stomach), in a combination that provided a very high diversity of moderately well-expressed genes.

**Construction and hybridization of labeled cDNAs.** A total of 40 µg RNA from R and extracted from each of four confluent 100-mm-diameter dishes of cultured astrocytes, each isolated from different neonatal WT or Cx43-null mice, was reverse transcribed into cDNA using fluorescent dUTPs (Cy3-dUTP, excited at 532 nm (green), and Cy5-dUTP, excited at 635 nm (red)). The labeled cDNAs were hybridized overnight at 50°C on 10 aminosilane-coated Corning glass slides, spotted with 8,975 selected mouse DNA sequences in 16 pen-domains (provided by the Albert Einstein College of Medicine Microarray Facility; array code 15M) in the combinations indicated in Table 1. After hybridization, the slides

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**Table 1. Experimental design**

<table>
<thead>
<tr>
<th>Step</th>
<th>Hybridization</th>
<th>Purpose</th>
</tr>
</thead>
<tbody>
<tr>
<td>α</td>
<td>RrKg</td>
<td>“Yellow test” to check the printing quality, verify the hybridization efficiency, test the relevancy of gene expression within the reference, and determine the PMT settings</td>
</tr>
<tr>
<td>β</td>
<td>RrWT/1g</td>
<td>Estimate the expression variability and stability in WT cortical astrocytes</td>
</tr>
<tr>
<td>γ</td>
<td>RrKO/1g</td>
<td>Estimate expression variability and stability in Cx43-null cortical astrocytes and determine expression ratios and scores</td>
</tr>
</tbody>
</table>

CNS cDNA array spot intensities were analyzed in three steps (α, β, γ), using the number of arrays and type of hybridization indicated. In the α-step, two arrays were hybridized with reference (R) cDNA, half of which was labeled with the red fluorescent Cy5 (r) and half with green Cy3 (g), to correct for differences in photomultiplier voltages (PMT settings). In the β-step, RNA extracted from cultured wild-type (WT) astrocytes from four animals (indicated as 1...4) was labeled with green fluorescent and hybridized against red labeled reference (R) to establish the variability of hybridization to each spotted sequence among the WT astrocytes. In the γ-step, green-labeled cDNA from four connexin-43 (Cx43)-null mice (KO, indicated 1...4) was hybridized against the red-labeled reference to measure variance. Finally, expression ratios (r) were determined for each spot, and expression scores (σ) were calculated based on the hybridization variances, to test the statistical significance of the observed regulations.
were washed at room temperature, using solutions containing 0.1% sodium dodecyl sulfate (SDS) and 1% SSC (3 M NaCl + 0.3 M sodium citrate) to remove the nonhybridized cDNAs. The experimental design and the usage of an invariant reference eliminate the bias introduced by the direct incorporation of fluorescent dyes during the reverse transcription reaction (24), since the extracts which are compared have the same label (here green) and the reference unit (here red labeled) is constant.

Scanning, data acquisition, and normalization. The microarrays were scanned with an Axon GenePix 4000A scanner using the same PMT settings (750 V at 635 nm, 670 V at 532 nm) and maintained throughout the entire experiment, and data were acquired through GenePix Pro 4.0 software (http://www.axon.com). We considered only spots $s$ (of each array $γ$) for which the means of the foreground signals ($P$) were significantly ($P < 0.001$) higher than those of the background signals ($B$) in both channels. Thus the cut-off conditions (denoted as “635cut-off” and “532cut-off”) of valid spots were

$$
\tau(s; W \rightarrow K; P) = \sqrt{\frac{[n_B(s) - 1] \text{Var}(\psi_{532}(\gamma; s))}{\text{Var}(\psi_{635}(\gamma; s))}} + \frac{[n_B(s) - 1] \text{Var}(\psi_{635}(\gamma; s))}{\text{Var}(\psi_{532}(\gamma; s))} > 3,
$$

$$
\sqrt{\frac{[F_532\text{mean}(\gamma; s) - B_532\text{mean}(\gamma; s)]^2}{F_532\text{pixels}(\gamma; s) + B_532\text{pixels}(\gamma; s)}} > 3.
$$

where symbols are those from the gpr file provided by GenePix Pro 4.0 software, and “3” is the critical value of the standard score establishing that the means of the foreground and background pixel intensities are statistically different ($P < 0.001$).

The raw data were first corrected according a previously published procedure (25) to minimize various noise sources (26), then normalized to make them comparable (4). In addition to the composite within-array normalization (58) intended to balance the net reference channels of the two channels within each array, a similar composite between-array normalization was used to balance the net reference channels from all arrays.

Gene expression indicators. The expression ratio of the gene identified by the spot $s$ in the Cx43-null cortical astrocytes ($K$) compared with the WT ($W$) was computed as

$$
x(s; W \rightarrow K) = \frac{\psi_{532}(\gamma; s)}{\psi_{635}(\gamma; s)} \text{all } RrKg \text{ chips} - \frac{\psi_{635}(\gamma; s)}{\psi_{532}(\gamma; s)} \text{all } RrWg \text{ chips}.
$$

where $γ$ indicates the array hybridized with cDNAs from $R$ and $W$ or $R$ and $K$, and $ψ_{532}(γ; s)/ψ_{635}(γ; s)$ designates the arithmetic mean of the corrected fluorescence net ratio of the spot $s$ in the array $γ$ hybridized with the red-labeled reference and green-labeled extract of Cx43-null astrocytes, $RrKg$, or with the red-labeled reference and the green-labeled extract of WT astrocytes, $RrWg$.

The $P$ value of the up- or downregulation can be determined through the one-tailed $t$-test for different (unknown) population variances

$$
P(s) = \frac{\psi_{532}(\gamma; s) \text{all } RrKg \text{ chips}}{\psi_{635}(\gamma; s) \text{all } RrWg \text{ chips}} = t_{\text{test}}(\psi_{532}(\gamma; s) \text{all } RrKg \text{ chips}, \psi_{635}(\gamma; s) \text{all } RrWg \text{ chips}, 1, 3)
$$

where $[ψ_{532}(γ; s)]$ is the set of corrected green readings of the spot $s$ in all arrays hybridized with $RrKg$ or $RrWg$.

The observed regulation is significant if the expression score ($τ$) falls in the critical region

$$
\left( -\infty, t\left(n_B(s) + n_W(s) - 2, \frac{P}{2}\right) \right) \cup \left( t\left(n_B(s) + n_W(s) - 2, \frac{P}{2}\right), \infty \right)
$$

where $t(n_B(s) + n_W(s) - 2, P/2)$ is the negative $P$-significant Student critical value, and $n_B(s)$ is the number of arrays hybridized as $RrKg$ (or $RrWg$) in which the spot $s$ was adequately quantified.

Transcription patholog. Using the expression scores, we built the pre-Hilbert space of the transcriptomic profiles having as many dimensions as distinct genes that were adequately quantified; every point of this space is a possible transcriptome (here of cortical astrocytes), and every curve is a possible transcriptomic evolution (22). For each significantly regulated gene, we determined the ratio between the absolute score, $|x(s; W \rightarrow K)|$, and the positive limit of the critical region, $t(n_B(s) + n_W(s) - 2, P/2)$, and then we summed the differences between each ratio and the unit. The percent ratio between the obtained sum and the number of adequately quantified genes was termed the transcription patholog, which quantifies the overall significant alteration of the expression of adequately quantified genes (23).

Variability (REV) and stability (GES) of gene expression. We defined the $P$-significant relative estimated variability (REV) of the gene identified by the spot $s$ in the specimen $A$ ($= W, K$) as the midrange chi-squared ($χ^2$) interval estimate of the standard deviation compared with the average corrected signal.

Because of the asymmetry of the REV distribution (expressed by the skewedness value, see RESULTS), this indicator cannot be used to categorize the genes according to their expression variability. Therefore, we introduced the gene expression stability (GES) as the percentile that locates the
gene’s REV in the inversely ordered sequence of REV values of all adequately quantified genes.

These measurements of variability and stability of transcription are not redundant; REV provides an estimation of transcriptional control, whereas GES integrates the studied genes in the general picture of GES within a particular specimen, eliminating the variability introduced by preparations of different types of samples.

Coordinated transcriptomics. The analysis of expression coordination makes it possible to group the genes that regulate each other in various experimental conditions and may ultimately allow identification of “command gene” within every group of interest.

The Pearson correlation coefficient \( r \) was used to determine the degree of expression coordination of the genes probed by the spotted sequences \( s_1 \) and \( s_2 \) within the specimen \( A \) (= W, K). The correlation coefficient yields values from -1 to +1. Positive correlation indicates that the expressions of both genes increase and decrease simultaneously in a series of extracts, and negative correlation indicates inverse expression tendencies, whereas a value close to zero indicates that the variations of expression levels of the two genes are independent. The statistical significance of the correlation coefficient is strongly dependent on the number of replications. In our case of four replications, the 5% cutoffs are \( |r| > 0.9 \) for coordination and \( |r| < 0.05 \) for independence, whereas the 10% cutoffs are \( |r| > 0.8 \) for coordination and \( |r| < 0.1 \) for independence. Moderate correlations \((0.1 \leq |r| \leq 0.8)\) were eliminated from the analysis as not statistically significant.

A major problem in the coordinated transcriptomic analysis is the manipulation of hundreds of millions of correlation coefficients at a time. To limit the calculations to manageable data sets, it is currently necessary to preselect the classes of genes to be considered and then to construct the coordination topology by aggregating the small groups. We hypothesized that transcription coordination among the genes related in a functional pathway might be conserved among genotypes. Therefore, we studied the transcription coordination of the genes related to the cell cycle-shape-differentiation and cell death pathways (see RESULTS) within the WT astrocytes that were found to be significantly up- or downregulated in the Cx43-null astrocytes.

We defined the coordination power (COP) of a gene to show the percentage contribution of that gene to the total correlations within a group of selected genes for each genotype. One goal of our studies has been to introduce an objective criterion by which to choose the most prominent (or “command”) gene for every group and report the coordination of all other genes to it. Therefore, we identified the most prominent gene as the one combining the highest coordination (COP) power against the others within its group with the maximum expression stability (GES); the term given for the product of COP and GES is gene prominence (GP).

\[
\rho(s_1, s_2; A) = \frac{\text{CoVar}(\log_2 [\psi_{532}(\gamma; s_1)], \log_2 [\psi_{532}(\gamma; s_2)])_{\text{all exp. chips}}}{\sqrt{\text{Var}(\log_2 [\psi_{532}(\gamma; s_1)])_{\text{all exp. chips}}} \times \text{Var}(\log_2 [\psi_{532}(\gamma; s_2)])_{\text{all exp. chips}}} 
\]

\[ \text{(6)} \]

RESULTS

The \( \alpha \)-step (Table 1) revealed that hybridization of 6,909 out of the 8,975 spots were adequately quantifiable and therefore useful for further analysis. For affected genes, accession numbers were reblasted using http://www.informatics.jax.org and http://www.ncbi.nlm.nih.gov to update the gene description from that given on the platform file for the 15M mouse microarray (http://biotech.aecom.yu.edu:7778/web/aecomarray.html).

Expression variability for individual spots and genotype-specific differences. To determine the significance of changes in observed gene expression ratio, the normal variability for each quantifiable spot among individual WT and Cx43-null astrocytes was first determined (\( \beta \)- and \( \gamma \)-steps in Table 1). The histograms of REV are shown in Fig. 1A for 3,032 spots that were quantifiable in all four arrays from WT astrocytes and in Fig. 1B for the 3,593 spots that were quantifiable in each of four arrays from Cx43-null astrocytes. A comparison of these histograms indicates that the mean and standard deviation REV values were lower for WT than for Cx43-null astrocytes. Moreover, the histogram of REV values for the WT astrocytes displayed a...
slightly sharper peak than for Cx43-null astrocytes (as quantified by skewness and kurtosis values presented in Fig. 1, A and B). Both distributions were slightly positively oriented (as quantified by skewness values).

The slight apparent differences in overall distribution of REV values raised the question of whether the REV values of individual genes were similar in WT and Cx43-null astrocytes. To address this issue, the REV value for each spot from WT astrocytes was plotted against the REV value for the same spot from Cx43-null astrocytes (Fig. 1C). This analysis revealed a surprising degree of independence of spot variability in WT compared with Cx43-null astrocytes. This independence was most apparent when comparing REV values of spots that were highly variable (REV > 100) in one or the other genotype. For example, in both WT and Cx43 knockout (KO) astrocytes, virtually all genes with REV > 100 in one genotype showed REV < 100 in the other genotype. Regression analysis revealed that the two sets of REV values are independent at P < 0.05, consistent with the apparent independence for spots with the highest REV values.

The most and least variable spots for which the genes were identifiable in WT astrocytes are listed in Tables 2 and 3, where REV values are given for each gene in Cx43-null astrocytes as well. In addition, values are presented for GES, a parameter that indicates the normalized rank order of stability in each genotype (see MATERIALS AND METHODS).

As expected from the independence of REV values illustrated in Fig. 1C, the degree of expression stability of certain genes was dramatically different in the WT and Cx43-null astrocytes. For example, some genes converted from being highly controlled to highly fluctuant (e.g., immunoglobulin κ-chain joining region and procollagen-lysine 2-oxoglutarate 5-dioxygenase 1; notes A and B in Table 2), and some genes changed from highly fluctuant to highly stable (e.g., UDP-glucose dehydrogenase and ribosomal protein L44; notes A and B in Table 3). However, a small number of genes preserved their character of being highly stably or highly unstably transcribed in the astrocytes following deletion of the Cx43 gene. For instance, ganglioside-induced differentiation-associated protein 2 (note 1 in Table 2), zinc finger protein 275 (note 2, Table 2), proteasome subunit-α type 4 (note 3, Table 3), ubiquitin conjugating enzyme 7 interacting protein (note 4, Table 2), torsin family 3 member 1A (note 5, Table 2), and adaptor-related protein complex AP1 μ-subunit 1 (note 6, Table 2) were very stable in astrocytes of both genotypes, whereas aquaporin 1 (note 1 in Table 3), biglycan (notes 2a, 2b, and 2c, Table 3), secreted modular calcium binding protein 2 (note 3, Table 3), and low-density lipoprotein receptor (note 4, Table 3) were highly unstable in both groups. It should be noted that one of these genes (biglycan, bolded and notes 2a, 2b, and 2c in Table 3) was represented by three DNA spots; in all three cases and in both genotypes, this gene exhibited a high degree of variability. The similar REV values for each of these spots in WT and in Cx43-null astrocytes provided further validation of the reproducibility of the measurements.

As indicated below, we have analyzed the coordination among expression of genes related to programmed cell death. Interestingly, disruption of Cx43 increased the average transcription variability among these genes from 37.6% to 58.9%, suggesting a relaxation of the transcription control of these genes in the Cx43-null astrocytes.

Differences in expression levels of individual genes in WT and Cx43-null astrocytes. Hybridization of the cDNA microarrays with labeled RNA extracts led to quantifiable signals for at least 3 of 4 samples from both genotypes for 4,998 of the spots. Of these, 206 (4.1%) of the spotted sequences showed significantly (P < 0.05) less hybridization in the Cx43-null astrocytes, and 468 (9.4%) showed significantly higher hybridization. When corrected for redundancy and expressed sequence tag (ESTs) removed, the number of significantly altered genes in Cx43 KO astrocytes was reduced to 252. Of the 252 identified genes that were altered in Cx43-null astrocytes, 35% (88 genes) were downregulated and 65% (164) were upregulated (Fig. 24).

A complete list of the 252 identified genes whose expression was significantly different in Cx43-null and WT astrocytes is given in the Supplemental Table A (available at the Physiological Genomics web site), along with the expression ratio (x), its binary logarithm (log2 x) and P values. To begin to assign the functional pathways to which these gene products belong, we have classified them according to biological functions within the cell (2). The 10 categories in which we have grouped the altered genes are as follows: junction-adhesion-extracellular matrix (JAE), which includes antigens, globulins, integrins, claudins, cadherins, connexins, desmosomal components, laminin, proteoglycans, etc.; cytoskeletal (CY, which includes intermediator filaments, microtubules, centrosomes, actin, and their associated proteins); transport into the cell (T1, which includes channels, transporters, and ionotropic receptors); transport within the cell (T2, which includes proteins of vesicles, cellular motors, endosomes, lysosomes, nuclear transport, protein folding, etc.); cell signaling (CS, which includes G-protein-coupled receptors, protein kinases, SH2 and SH3 domain proteins, calcium binding proteins, etc.); cell cycle-shape-differentiation (CSD, which includes growth factors, apoptosis-related genes, cytokines, etc.); transcription (TR, which includes DNA binding proteins, DNA repair, RNA, transcription factors, oncogenes, etc.); energy and metabolism (EnMet, which includes oxidants, peroxisomes, respiratory chain, glycolysis and glycogenesis, enzymes, etc.); organelle genetics (OG, which con-
Table 2. The least variably (i.e., most stably) expressed genes in WT neonatal mouse cortical astrocytes and corresponding values in Cx43-null astrocytes

<table>
<thead>
<tr>
<th>Notes</th>
<th>GenBank ID</th>
<th>Description</th>
<th>REV-WT</th>
<th>GES-WT</th>
<th>REV-KO</th>
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<td>AA032414</td>
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<td>procollagen-lysine, 2-oxoglutarate 5-dioxygenase 1</td>
<td>7.75</td>
<td>99.21</td>
<td>134.64</td>
<td>3.67</td>
</tr>
</tbody>
</table>

REV, relative estimated variability; GES, gene expression stability; KO, knockout. Notes are addressed in text.
equally distributed among the other three categories (Fig. 2B).

Pie charts depicting the distributions of up- and downregulated genes in Cx43-null compared with WT astrocytes are presented in Fig. 2, C and D. Comparison of the proportions of genes that were upregulated and downregulated in each functional category reveals overall similarity as well as some interesting differ-

### Table 3. The most variably expressed genes in WT astrocytes and corresponding values in Cx43-null astrocytes

<table>
<thead>
<tr>
<th>Notes</th>
<th>GenBank ID</th>
<th>Description</th>
<th>REV-WT</th>
<th>GES-WT</th>
<th>REV-KO</th>
<th>GES-KO</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>AA38426</td>
<td>translocated promoter region protein</td>
<td>317.10</td>
<td>0.03</td>
<td>121.74</td>
<td>4.98</td>
</tr>
<tr>
<td></td>
<td>AA387076</td>
<td>A kinase (PRKA) anchor protein (gravin) 12</td>
<td>193.29</td>
<td>0.20</td>
<td>109.38</td>
<td>6.76</td>
</tr>
<tr>
<td></td>
<td>AA109041</td>
<td>calmodulin 3</td>
<td>185.07</td>
<td>0.26</td>
<td>60.40</td>
<td>32.09</td>
</tr>
<tr>
<td></td>
<td>W97837</td>
<td>protein kinase, DNA activated, catalytic polypeptide</td>
<td>172.19</td>
<td>0.33</td>
<td>67.00</td>
<td>26.41</td>
</tr>
<tr>
<td>2a</td>
<td>AA000678</td>
<td>biglycan</td>
<td>150.07</td>
<td>0.69</td>
<td>228.78</td>
<td>1.03</td>
</tr>
<tr>
<td></td>
<td>AA172527</td>
<td>ATP-binding cassette, sub-family G (WHITE), member 1</td>
<td>147.68</td>
<td>0.73</td>
<td>189.56</td>
<td>1.81</td>
</tr>
<tr>
<td></td>
<td>AA145458</td>
<td>fibronectin 1</td>
<td>143.88</td>
<td>0.79</td>
<td>199.98</td>
<td>1.42</td>
</tr>
<tr>
<td></td>
<td>W98883</td>
<td>procollagen, type III, alpha 1</td>
<td>141.08</td>
<td>0.86</td>
<td>376.91</td>
<td>0.06</td>
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<td></td>
<td>W83904</td>
<td>peptidylprolyl isomerase C</td>
<td>140.99</td>
<td>0.89</td>
<td>222.57</td>
<td>1.11</td>
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<td></td>
<td>W98037</td>
<td>proteasome (prosome, macropain) 26S subunit, non-ATPase, 5</td>
<td>139.95</td>
<td>0.92</td>
<td>90.25</td>
<td>11.75</td>
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<tr>
<td></td>
<td>W85177</td>
<td>type IV collagenase</td>
<td>139.61</td>
<td>0.96</td>
<td>227.98</td>
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<td></td>
<td>AA989099</td>
<td>cadherin 11</td>
<td>136.20</td>
<td>1.06</td>
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<td>18.93</td>
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<td></td>
<td>AA388435</td>
<td>peroxisomal sarcosine oxidase</td>
<td>134.85</td>
<td>1.09</td>
<td>131.51</td>
<td>4.04</td>
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<td>2b</td>
<td>AA200264</td>
<td>biglycan</td>
<td>130.61</td>
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<tr>
<td></td>
<td>AA059909</td>
<td>secreted modular calcium binding protein 2</td>
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<td>1.42</td>
<td>368.60</td>
<td>0.11</td>
</tr>
<tr>
<td></td>
<td>AA220047</td>
<td>zinc bromatosis 2; schwannomin; moesin-ezrin-radixin-like protein</td>
<td>121.48</td>
<td>1.48</td>
<td>99.28</td>
<td>9.32</td>
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<td></td>
<td>AA105153</td>
<td>neuro</td>
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<td>2.77</td>
<td>63.57</td>
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<td></td>
<td>AA051390</td>
<td>protein (peptidyl-prolyl cis/trans isomerase) NIMA-interacting 1</td>
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<td>1.42</td>
<td>368.60</td>
<td>0.11</td>
</tr>
<tr>
<td></td>
<td>AA060905</td>
<td>guanosine diphosphate (GDP) dissociation inhibitor 1</td>
<td>124.61</td>
<td>1.42</td>
<td>368.60</td>
<td>0.11</td>
</tr>
<tr>
<td></td>
<td>AA116505</td>
<td>CD53 antigen</td>
<td>124.61</td>
<td>1.42</td>
<td>368.60</td>
<td>0.11</td>
</tr>
<tr>
<td></td>
<td>AA176035</td>
<td>polycystic kidney disease 2</td>
<td>124.61</td>
<td>1.42</td>
<td>368.60</td>
<td>0.11</td>
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<tr>
<td></td>
<td>AA119255</td>
<td>ribosomal protein L13a</td>
<td>124.61</td>
<td>1.42</td>
<td>368.60</td>
<td>0.11</td>
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<tr>
<td></td>
<td>AA177363</td>
<td>ectonucleotide pyrophosphatase/phosphodiesterase 2</td>
<td>124.61</td>
<td>1.42</td>
<td>368.60</td>
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<tr>
<td></td>
<td>AA110412</td>
<td>myeloid differentiation primary response gene 88</td>
<td>124.61</td>
<td>1.42</td>
<td>368.60</td>
<td>0.11</td>
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<tr>
<td></td>
<td>AA239404</td>
<td>nidogen 1</td>
<td>124.61</td>
<td>1.42</td>
<td>368.60</td>
<td>0.11</td>
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<tr>
<td></td>
<td>AA172598</td>
<td>ATPase, Na+/K+ transporting, alpha 4 polypeptide</td>
<td>124.61</td>
<td>1.42</td>
<td>368.60</td>
<td>0.11</td>
</tr>
</tbody>
</table>

**REV and GES are defined in Table 2. Note the similarity of the values obtained for the three distinct spotted sequences homologous to biglycan: AA000678 [HS = 407, p(N) = 1.1 × 10^{-3}], AA200264 [HS = 2014, p(N) = 1.1 × 10^{-3}], W54287 [HS = 2970, p(N) = 7.2 × 10^{-20}], which reinforces confidence in the accuracy of our transcription stability analysis.**

Notes are discussed in the text.
ences. For example, a higher percentage of TR genes (32 vs. 25%) and almost three times the percentage of T2 genes (12 vs. 4.5%) were upregulated than downregulated in the Cx43-null astrocytes, whereas a higher percentage of JAE genes (11 vs. 7.3%) and a greater percentage of EnMet genes (24 vs. 17%) were downregulated in the Cx43-null astrocytes. With regard to transcripts encoding proteins involved in cell cycle and cell growth control, we detected about the same percentage upregulated (7.3%) as downregulated (9.1%). These regulated genes included upregulation of fibroblast growth factor 1 (FGF-1), fractalkine, BMP1, growth arrest and DNA-damage-inducible 45p, and Bcl2-like; the most highly downregulated CSD genes included platelet factor 4, septin 3, and LIM only 2. Moreover, transcription factors acting on the cell cycle were affected; notably, axin (a component of the Wnt signaling pathway, which presumably is attributable to the more homogeneous population of Cx43 expressing cells in cultured astrocytes than in the whole brain.

Overall change in transcriptome of Cx43-null astrocytes. The global alteration of the transcriptome was evaluated by calculating the value of the transcription pathol (see MATERIALS AND METHODS). When all genes were considered, the calculated pathol was 2.4%. This value is twice that calculated for the pathol in Cx43-null brains (D. A. Iacobas, unpublished results), which presumably is attributable to the more homogeneous population of Cx43 expressing cells in cultured astrocytes than in the whole brain.

Coordinated transcriptomics of cell cycle/shape/differentiation genes. We and others (8, 34) have previously reported that the growth rate of astrocytes cultured from Cx43-null mice is dramatically slower than that of WT astrocytes. To determine whether genes involved in growth and differentiation were coordinately regulated in WT astrocytes and to examine genotype-related changes in this regulation, we have calculated correlation coefficients for each pair of the CSD genes found to be significantly altered for all arrays.

Of the 20 significantly regulated genes classified as encoding proteins with CSD functions, 16 were adequately quantifiable on all four arrays from both genotypes and therefore suitable for the coordination analysis. Correlation coefficients for each pair of selected genes are presented in the form of a coordination diagram in Fig. 3, where lines between the boxed genes indicate significant correlation in WT astrocytes and numbers represent the correlation coefficients between the WT expression ratios (highlighted values indicate antagonistic coordination). Colors indicate whether the gene was found to be significantly upregulated (red) or downregulated (green) in Cx43-null astrocytes. This analysis revealed 17 significant correlations, including 3 triangular associations, where the 3 correlation coefficients were consistent with the coordination of each of the others (Fig. 3A). In 4 cases out of 17 (all within the bottom cluster in Fig. 3B), regulations in Cx43-null astrocytes were opposite to that predicted by the correlation coefficients. Nevertheless, 77% of the significantly coordinated pairs of genes in WT astrocytes accurately predicted the significant regulation of these genes in the Cx43-null astrocytes.

Based on the correlation coefficients and the GES, we have calculated the COP and the GP of the CSD genes, which are listed in Table 4. Interestingly, the coordination powers (COP values) are confined in a relatively narrow interval (4.91% to 6.81%), with cyclin-dependent kinase inhibitor 2C (Cdkn2c) showing the least coordination with the other CSD genes, and growth arrest and DNA-damage-inducible 45p (Gadd45b) having the highest COP score. By contrast, the expression stability (GES) of these genes spans a large range of percentiles (12.63 to 88.26), with N-myc...
downstream regulated 3 (Ndr3) the least stable and Gadd45b having the highest stability.

As a consequence of its highest coordination power (COP = 6.81%) and highest stability (GES = 88.26%), Gadd45b was identified as the most prominent gene of the CSD class (GP = 11.81%), followed closely by Cdk2 (GP = 11.62%), Cdk9 (GP = 10.04%), and similar to mitotic control protein dis3 (Mcpd3) (GP = 10.66%). As predicted by the antagonistic coordination with Gadd45b and Cdk9, Cdk2 was downregulated in Cx43-null astrocytes while the other two were upregulated. Mcpd3 is remarkable in that its downregulation in Cx43-null astrocytes while the other two were upregulated. Mcpd3 is remarkable in that its downregulation in Cx43-null astrocytes while the other two were upregulated. Mcpd3 is remarkable in that its downregulation in Cx43-null astrocytes while the other two were upregulated. Mcpd3 is remarkable in that its downregulation in Cx43-null astrocytes while the other two were upregulated. Mcpd3 is remarkable in that its downregulation in Cx43-null astrocytes while the other two were upregulated. Mcpd3 is remarkable in that its downregulation in Cx43-null astrocytes while the other two were upregulated.

Coordinated transcriptomics of apoptotic genes. As an additional strategy to search for coordinated gene expression within functional pathways and to investigate the relationship between Cx43 expression and apoptosis, we extended our coordinated transcriptomic analysis to genes related to programmed cell death. For this analysis, we selected all adequately expressed genes related to programmed cell death within WT astrocytes, using as a guide the pathway map compiled by Apotech (http://www.apotech.ch), with gene synonyms obtained from Online Mendelian Inheritance in Man (OMIM; NCBI).

Our analysis in WT astrocytes of 24 programmed cell death-related genes whose expression was quantifiable
on all arrays revealed that for most of these genes there was highly significant synergistic or antagonistic expression with at least one other gene in this group. In addition, gene expression was significantly independent for some pairs. The significant positive and negative relationships are summarized in the coordination diagram shown in Fig. 4, where the solid lines indicate correlations that are highly significant \((P < 0.05); \) numbers represent correlation coefficients calculated in WT astrocytes. As can be seen from the solid lines in Fig. 4, the majority of highly significant correlation coefficients were for positive regulation \(20\) were significantly synergistic (red lines), compared with 8 antagonistic relations (blue lines).

The prominence sequence of the analyzed apoptotic genes (where numbers indicate GP values) is: Apaf1 \((7.49)\) > Tnfrsf10b \((6.61)\) > Map2k1ip1 \((6.57)\) > Ube2i \((6.47)\) > Tnfrsf10a \((6.45)\) > Casp2 \((6.23)\) < Prkacb \((6.17)\) > Hspb2 \((5.67)\) > Bin2 \((4.40)\) > Prnp \((4.24)\) > Prkaca \((4.18)\) > Akt1 \((4.17)\) > Akt2 \((0.9)\). Therefore, for these apoptotic pathway genes, our analysis indicates that

![Fig. 4. Significant coordinations among the expression of genes related to programmed cell death in WT cortical astrocytes. Gene designations are within boxes; lines between boxes indicate significant coordinations of the expressions of paired genes in the WT cortical astrocytes (values on lines provide correlation coefficients). Colors of boxes indicate whether a gene was significantly upregulated (red) or downregulated (green) in Cx43-null astrocytes, while colors of lines indicate whether the coordination was synergistic (blue) or antagonistic (red).](image-url)
the most prominent genes are apoptotic protease activating factor 1 (Apafl) and tumor necrosis factor receptor super-family member 10b (Tnfrsf10b or Trail2), whereas the least prominent is thyoma viral protooncogene 2 (Akt2).

Expression of 8 of the 24 quantifiable apoptotic genes was significantly altered in Cx43-null astrocytes (Fig. 4, red boxes indicate upregulation, green boxes downregulation). In all cases but one, significantly regulated genes were highly correlated with expression of at least one other gene in this pathway. Whereas some of these up- and downregulations in Cx43-null astrocytes were as expected from the correlation linkages determined in WT astrocytes (e.g., the Nfkb2 cluster), others were not, suggesting that correlations among the genes might be altered in Cx43-null astrocytes. Calculations of correlation coefficients between each pair of the 24 quantifiable apoptotic genes revealed global changes, with 10.6% of the correlations being significantly increased and 16.2% being significantly reduced.

We further found that the proportion of the coexpressed genes was higher in Cx43 KO astrocytes (13.4% compared with 7.1%), indicating an enhanced coupling among apoptotic genes in the Cx43-deficient cells. It is also interesting to note that the proportion of antagonistically coexpressed pairs increased from 2.7% in WT to 8.9% in KO, suggesting an enhanced reciprocal negative control of these genes; the proportion of significant synergistic coordination was slightly reduced (from 6.1% in WT to 4.7% in Cx43-null astrocytes). The most stably and the most unstably expressed genes in the WT astrocytes were apoptotic protease activating factor 1 (GES_Apafl = 98.9%) and thyoma viral protooncogene 2 (GES_Akt2 = 12.5%).

Correlation patterns in the Cx43-null astrocytes were similar to those seen in WT astrocytes. Functional stability (FS), defined as the weighted average of GES values by the COP values, was calculated for WT and Cx43-null astrocytes. Functional stability of the moderately stable apoptotic pathway (FS = 56.8%) was found to be slightly diminished by the disruption of Cx43 (FS = 46.5%).

DISCUSSION

We have quantitatively analyzed 9k DNA arrays in which labeled cDNA from four WT and four Cx43-null sibling mice were hybridized against a pooled reference obtained from a variety of mouse tissues. To statistically compare expression levels in each gene between genotypes, we have determined the variability in expression ratio (REV) for each spot in each genotype and then incorporated the variability in calculations of expression ratio (x) to obtain gene expression scores (r). Because these scores incorporate variability, they permit statistical comparison of gene expression without using arbitrary cutoff thresholds in ratio values.

One finding from these calculations was that the variability of individual gene expression ratios spanned a wide range in both WT and Cx43-null astrocytes. Although low REV values indicate loose transcriptional control and high REV values indicate loose control, we have not yet attempted to address the implied issue of whether high or low variability might reflect genes whose functions segregate with cellular housekeeping or with cellular specialization. However, the REV values for individual spots in one genotype were statistically independent of REV values for the same spots in the other genotype. This is an interesting finding, suggesting that “compensation” or “adaptation” for the loss of one gene may change the degree of transcriptional control for a wide range of individual genes with unrelated functions.

When expression scores were analyzed, a large number of spots were found to differ significantly between genotypes, and the significantly altered spots surprisingly corresponded to a large number of identified genes with functions apparently unrelated to intercellular communication. As noted in the Introduction, this finding is consistent with a number of recent reports describing effects of connexin expression on cell growth, apoptosis, and purinergic receptor expression (16, 34, 42, 59, 31, 47, 52, 30), not all of which require formation of functional gap junction channels. Numerous mechanisms may be responsible for such profound changes in gene expression, including those related to developmental events that may depend on intercellular communication or functionally compensate for its loss. Because Cx43 is a gap junction protein that is expressed at earliest times during embryogenesis (53), loss of Cx43 might be predicted to exert particularly global effects on subsequent development, where changes in expression might reflect functional adaptation. Studies to evaluate this possibility are underway using astrocytes treated acutely with antisense oligonucleotides to reduce Cx43 expression (52). Alternatively, or in addition, there may be direct feedback, so that Cx43 or its binding partners may act directly or indirectly to transactivate or inactivate the other genes. Such a mechanism has been identified involving β-catenin interaction with Cx43, whereby β-catenin exerts a transcriptional control that is modulated by the interaction between β-catenin and Cx43 at the junctional membrane (1). Moreover, “connexin response elements” have been proposed to modulate gene expression in osteocytes (51).

To evaluate whether Cx43 deletion selectively affected genes with specific functions, we have categorized the identified genes according to 10 functions of encoded proteins. This simple classification allowed us to select 16 regulated genes involved in cell cycle regulation for detailed analysis of coordinated expression. Our analysis of the coordination among the genes and alterations in the Cx43-null astrocytes may be interpreted to provide a mechanistic basis for the slowed growth rates of Cx43-null than WT astrocytes in culture (8, 36). For example, upregulation of Gadd45b (which activates p38 and JNK pathways), Ndr3 (which may decrease cyclin D1), FGF1 (which promotes exit from cell cycle), and requiem (which is required for apoptosis) may all favor decreased cell proliferation. Moreover, downregulation of Cdk2 (which is necessary
for G2-M phase transition), Lmo2 (which is an important factor in early development), Myadm (a myeloid differentiation factor), Septin 3 (which is a key factor in late steps of cytokinesis), and the relative of Dis3 (Mcpd3, which is another required mitotic signal) may act directly to slow cell growth. Indirect effects through transcription factors may also participate, as indicated by the upregulation of axin and p53 (which directly and indirectly slow cell growth).

The central hypothesis of our coordinated expression analysis is that genes encoding proteins with synergistic or antagonistic functions might be coordinately expressed, in contrast to genes encoding proteins with independent functions. Consistent with this hypothesis, the coordinated transcriptomic analysis predicted most of the up- and downregulated CSD genes in the Cx43-null astrocytes, with the exception of the interactions of Set3 and Mcpd3 with Cx3cl1 and Bcl2-like. This finding that the correlations of variations in the expression of a gene of interest and other genes among different animals largely predicts changes in the knockout further indicates that the correlations are robust. This type of analysis of other pathways may provide a methodological key to promote general understanding of synergistic, antagonistic, and independent intracellular signaling.

Both increased and decreased resistance to cell injury have previously been associated with Cx43 expression (18, 20, 29, 41). Our analysis of the coordinated expression of genes associated with programmed cell death involved calculations of correlation coefficients among all possible pairs of the 24 apoptosis-associated genes adequately expressed on all 8 of the DNA arrays used in these studies. This analysis revealed that both pro-apoptotic and anti-apoptotic genes were up- and downregulated in Cx43-null astrocytes. Upregulated pro-apoptotic genes include Nfkb2 and Map4k5, both of which selectively activate the JNK pathway, whereas the anti-apoptotic protein Bcl2-like (BclXL, BAX) was also upregulated. The coordination analysis revealed that expressions of other cell death-related genes were strongly correlated, many of which were further correlated with expression of those genes that were altered in Cx43-null astrocytes. For example, Map4k5 was negatively correlated with the Trail-2 receptor (Tnfrsf10b), which was positively correlated with two proteins implicated in neuronal death (Hspb2, Bircla), as well as a protein induced by inflammatory cytokines (Tnfaip2) and an activator of the ERK signaling pathway (Map3k3). These associations suggest that the genes involved in the TNFα and Trail death receptor pathways may be synergistic within the pathways and antagonistic between them. Moreover, these opposite regulations of genes encoding pro- and anti-apoptotic proteins and their complex linkages to the expression of other death pathway proteins may explain in part the reported differences in sensitivity of Cx43 transfectants and Cx43-null cell lines to stimuli that may selectively activate one or another of these pathways. It is important to point out that although these studies have been limited to cultured astrocytes, the findings imply that gene expression changes secondary to Cx43 expression may either enhance or diminish the neuroprotective effects of astrocytes on neurons.

Through the efforts of several laboratories, a number of gap junction genes have now been deleted through homologous recombination, and prominent phenotypic changes have been observed in many but not all of these transgenic mice (see Ref. 56 for recent review). Until now, all phenotypic changes observed in connexin-deficient mice have been attributed to the absence of intercellular channels in tissues where they are normally expressed. A caveat of such studies, and of the present analysis as well, is the lack of control for the neomycin phosphotransferase gene inserted to disrupt the coding region in the Cx43-null mouse. Nevertheless, the findings reported in this paper indicate that astrocytes from Cx43-null mice show up- and downregulation of many other genes with diverse functions and thus suggest that gap junction genes regulate multiple cellular processes. Functional changes observed in either deletion or overexpression studies may thus not only reveal the function of gap junction channels in the organ of interest, but may also reflect changes in expression of other genes induced by the manipulation.

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Gene expression in Cx43 knockout astrocytes


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