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Functional Gap Junctions Accumulate at the Immunological Synapse and Contribute to T Cell Activation


Gap junction (GJ) mediates intercellular communication through linked hemichannels from each of two adjacent cells. Using human and mouse models, we show that connexin 43 (Cx43), the main GJ protein in the immune system, was recruited to the immunological synapse during T cell priming as both GJs and stand-alone hemichannels. Cx43 accumulation at the synapse was Ag specific and time dependent, and required an intact actin cytoskeleton. Fluorescence recovery after photobleaching and Cx43-specific inhibitors were used to prove that intercellular communication between T cells and dendritic cells is bidirectional and specifically mediated by Cx43. Moreover, this intercellular cross talk contributed to T cell activation as silencing of Cx43 with an antisense or inhibition of GJ docking impaired intracellular Ca²⁺ responses and cytokine release by T cells. These findings identify Cx43 as an important functional component of the immunological synapse and reveal a crucial role for GJs and hemichannels as coordinators of the dendritic cell–T cell signaling machinery that regulates T cell activation. The Journal of Immunology, 2011, 187: 000–000.

Initiation of an Ag-specific immune response requires productive engagement of TCRs by MHC-peptide (pMHC) complexes on the APC (1, 2). This TCR engagement by cognate pMHC results in the formation of a highly organized protein network known as the immunological synapse (IS), which is required for T cell activation and proliferation (3). The mature IS is characterized by the assembly of specific proteins on the T cell and APC membranes into supramolecular activation clusters. These structures consist of a centralized accumulation of TCRs and pMHC (central supramolecular activation complex [cSMAC]), surrounded by a peripheral ring (peripheral supramolecular activation complex [pSMAC]) containing the integrin LFA-1 and its receptor ICAM-1 (3, 4).

The IS is comprised of a multitude of structures, many of which are mediators of intercellular communication (5). However, it is not known whether communication involving gap junction (GJ) channels, one of the most important mechanisms for cellular cross talk, occurs at the IS assembly site. GJs are clusters of intercellular channels in the plasma membrane that mediate direct intercellular communication between adjacent cells, allowing the passage of soluble molecules, including cAMP, Ca²⁺, ATP, inositol 1,4,5-trisphosphate, and morphogens (6, 7). GJs also mediate electrical and metabolic coupling among cells and tissues, such that signals initiated in one cell can readily propagate to neighboring cells. In mammals, functional GJs are composed of connexin (Cx) proteins. Six Cx proteins form a hemichannel (Hchl) inserted into the membrane of one cell, which then docks with a Hchl from an adjacent cell to establish a GJ channel (8, 9).

Cx- and GJ-mediated intercellular communication (GJIC) have been shown to participate in key immunological processes, such as Ig secretion and cytokine production (10), transendothelial migration of leukocytes (11), peptide transfer and cross-presentation in activated monocytes (12), activation of murine dendritic cells (DCs) (13), and regulatory T cell-mediated suppression through the transfer of cAMP (14). Additionally, we have demonstrated that GJ channels can also mediate the transfer of MHC class I–restricted melanoma peptides between human DCs, triggering T cell-specific immune response against melanoma-associated Ags (15). Recently, GJIC have also been shown to participate in DC-mediated induction of IL-2 release and proliferation of murine T cells (16). T cell activation and proliferation result from intercellular communications mediated by multiple surface molecules located at the IS. However, the accumulation of Cx at the IS, the mechanisms involved in their recruitment, and their role in T cell Ca²⁺ signaling and IFN-γ production have not been elucidated.

In the current study, we describe that Cx43, the main GJ protein of the immune system (10, 12, 17–19), accumulates at the IS and

*Institute of Biomedical Sciences, Faculty of Medicine, University of Chile, 8380453 Santiago, Chile; †Millennium Institute on Immunology and Immunotherapy, University of Chile, 8380453 Santiago, Chile; ‡University College London Cancer Institute, London WC1E 6DD, United Kingdom; ‖Molecular Immunology Unit, University College London Institute of Child Health, London WC1E 6BT, United Kingdom; and †Department of Molecular Genetics and Microbiology, Millennium Institute on Immunology and Immunotherapy, Pontificial Catholic University of Chile, 8331510 Santiago, Chile

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mediates bidirectional cross talk between DCs and T cells in murine and human systems. Moreover, we identify a role for GJs in the regulation of T cell activation.

**Materials and Methods**

**Mice**

OT-II transgenic mice expressing the OVA\(\alpha\)1,\(\alpha\)3,\(\alpha\)9 peptide (ISQAVHAA- HAEINEAGR), TCR (H-2b), and wild-type C57BL/6 mice were obtained from Charles River (Kent, U.K.). Mice were used at 6–12 wk of age, and all experiments were approved by and performed according to Home Office Animal Welfare Legislation.

**Generation of DCs and T cells**

This study was approved by the Bioethical Committee of Human Research, Faculty of Medicine, University of Chile. Informed written consents were given and signed by all patients. Leukocytes from stage IV melanoma patients were isolated by density gradient using Ficoll-Hypaque (Axis-Shield, Oslo, Norway). Human DCs were obtained, as described (20). At day 6, DCs were treated overnight with 150 \(\mu\)g/ml melanoma cell lysate (MCL), which was obtained, as previously described (15), and stimulated with 2 ng/ml TNF-\(\alpha\) (U.S. Biological, Swampscott, MA) (MCL-DCs). DCs stimulated with 1 \(\mu\)g/ml LPS plus 2 ng/ml TNF-\(\alpha\) and loaded with 2 \(\mu\)g/ml gp100\(\alpha\)2,\(\alpha\)17 peptide (gp100-DCs) were used as negative control. DCs from C57BL/6 mice were cultured from bone marrow cells for 7 d in the presence of GM-CSF (20 ng/ml; Invitrogen, Paisley, U.K.). For T cell priming experiments, DCs were matured with LPS (100 ng/ml; Sigma-Aldrich, Steinheim, Germany) in the presence or absence of OVA (100 \(\mu\)g/ml; Sigma-Aldrich).

MT56-4 is a CD4\(^+\) T cell line derived from tumor-infiltrating lymphocytes of a melanoma patient, which specifically recognizes autologous MCL-DCs and autologous melanoma cells, and was isolated and grown, as described (15). CD4\(^+\) T cells from OT-II transgenic mice were isolated from spleen using magnetic bead separation, according to manufacturer’s protocol (murine CD4\(^+\) T cell isolation kit; Miltenyi Biotec, Bisley, U.K.).

**T cell stimulation and immune fluorescence staining**

Dynal M450 beads (DYNal, Lake Success, NY) were coated with 3 \(\mu\)g/ml anti-human CD3 (OKT3) mAb and anti-human CD28 mAb (eBio-science, San Diego, CA), as well as with a control (anti-CD28) Ab (BD Pharmingen, San Jose, CA), according to manufacturer’s recommendations. PBLs (2 \(\times\) 10\(^5\)) from melanoma patients were incubated with 6 \(\times\) 10\(^3\) beads for 1 h at 37\(^\circ\)C. For some experiments, PBLs were pretreated or not with 10 \(\mu\)M cytochalasin D (Calbiochem, Gibbstown, NJ), 5 \(\mu\)M latrunculin A, 10 \(\mu\)M taxol, or 5 \(\mu\)g/ml nocodazole (last three from Sigma-Aldrich) 30 min before incubation with CD3/CD28-coated beads. In this set of experiments, cells were also stained with 5 \(\mu\)g/ml Hoechst 33342 (Invitrogen). Polyclonal anti-Cx43 Ab (21) and/or anti-CD3 mAb, clone HIT3a (BD Pharmingen), were incubated overnight at 4\(^\circ\)C. Samples were stained with Alexa Fluor 647 goat anti-rabbit (Molecular Probes, Invitrogen) and with tetramethylrhodamine isothiocyanate-conjugated rabbit anti-mouse (Sigma-Aldrich) and analyzed by confocal microscopy (LSM 510 META software; Carl Zeiss MicroImaging) using a 

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\text{immersion objective (Carl Zeiss) with } 510 \text{ META software; Carl Zeiss MicroImaging using a }
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\text{63 NA objective (Carl Zeiss).}
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**DC–T cell adhesion assay**

OT-II T cells and OVA-DCs, pretreated or not for 4 h with 40 \(\mu\)M Cx43-sense or Cx43-antisense (AS) or with 300 \(\mu\)M 1848 Cx43-mimetic peptide for 4 h, were stained using anti-Cx43 (21); PE-conjugated anti-CD4, PE-Cx43-sense, or PE-Cx43-antisense Abs. Cells were acquired on a flow cytometer (FACSort; BD Pharmingen) and analyzed using the CellQuest software.

**Flow cytometry analysis**

Flow cytometry experiments were performed, as previously described (15). DCs or T cells were pretreated or not with 40 \(\mu\)M Cx43-sense or Cx43-antisense (AS) or with 300 \(\mu\)M 1848 Cx43-mimetic peptide for 4 h. CD11c\(^+\) cells, corresponding to DCs, were gated, and the levels of different markers, including Cx43, were analyzed using a double staining. Cells were stained using anti-Cx43 (21); FITC-conjugated anti-CD3, CD40, MHC class I, and MHC class II (BD Pharmingen), and PE-Cx43-sense, or PE-Cx43-antisense Abs. Cells were acquired on a flow cytometer (FACSort; BD Pharmingen) and analyzed using the CellQuest software.

**MCL-DCs**

MCL-DCs (3 \(\times\) 10\(^5\)) were coincubated with 9 \(\times\) 10\(^3\) MCL-specific CD4\(^+\) T cells (MT56-4) for 1 h. The cell mixture was incubated with an anti-Cx43 Ab (21). Protein expression was visualized by using the corresponding secondary fluorescence-conjugated Ab Ab Alexa Fluor 647 goat anti-rabbit (Molecular Probes), which was also added alone as negative control. Cells were analyzed by confocal microscopy (LSM 510; \(\times\)63 NA 1.4 oil immersion objective (Carl Zeiss)).

HCCh formation was quantified by using a polyclonal anti-Cx43-Hchl Ab raised to a peptide sequence from the first external loop of Cx43 that recognizes Cx43 in an undocked conformation, and is occluded in docked GJs. T cells displaying >70% of Cx43-Hchl staining in the quadrant contacting DCs were scored as positive. We also analyzed the percentage of cells accumulated at the synapse that were not forming Hchl. To this end, a Cx43 mAb raised against the intracellular loop of Cx43 that recognized total Cx43, as both GJ and Hchls, was used, and again, T cells displaying >70% positive staining in the quadrant contacting DCs were scored as positive. We subtracted the number of cells scored as positive for Cx43-Hchl (Fig. 4E, dark gray) from the total number of cells scored as positive for Cx43, then obtaining the fraction of cells positive for Cx43, but negative for Hchls (Fig. 4E, light gray).

**Three-dimensional reconstructions and projections and quantitative image analysis**

The three-dimensional (3D) reconstruction of the confocal image stacks taken from interacting DCs and T cells stained for Cx43, LFA-1, and TCR was accomplished by using Velocity 4.4.0 analysis software (Improvision, Coventry, U.K.). For 3D reconstructions, 20–25 \(z\) sections were collected at 0.3-\(\mu\)m \(z\) intervals. The en face view of the immune synapse was obtained by an \(x-z\) projection of the 3D image at the cellular interface of the T cell and DC. The ratio of Cx43 fluorescence at the immunological synapse versus Cx43 fluorescence at the plasma membrane was calculated using National Institutes of Health ImageJ software. Statistical analyses were carried out by the nonparametrical Mann–Whitney U test. The \(p\) values <0.05 were considered statistically significant.

**Colocalization analysis**

Manders’ colocalization coefficients (24) were calculated at the site of interaction between OT-II T cells and OVA-DCs for TCR/Cx43, CD3/ Cx43, and LFA-1/Cx43 using NIH ImageJ software with the colocalization analysis plugin JACoP. Manders’ colocalization coefficient calculates the spatial overlap of two proteins, with M1 representing the percentage of Cx43 pixels (red channel) that overlaps pixels in the green channel (CD3 or LFA-1), and conversely for M2; M1 and M2 values range from 0 to 1, with a value of 0 corresponding to nonoverlapping images and the latter reflecting 100% colocalization between both images. Manders’ coefficients are not influenced by differences in absolute signal intensities in each channel because pixel intensity in a particular channel is normalized to total pixel intensity across the image for that label. Values were reported as mean \(\pm\) SEM.

**Fluorescence recovery after photobleaching experiments**

Images of six random fields from three individual samples per condition were analyzed. The recruitment of Cx43 to the contact site was quantified by using the corresponding secondary fluorescence-conjugated Ab Ab Alexa Fluor 647 goat anti-Cx43 (21). Protein expression was visualized by using the corresponding secondary fluorescence-conjugated Abs: hamster anti-mouse 7-amin-4-methylcoumarin-3-acetic acid (Jackson Immunoresearch Laboratories, West Grove, PA), streptavidin Alexa Fluor 488 conjugate (Molecular Probes), and goat anti-rabbit DyLight 549 (Pierce, Thermo Scientific, Basingstoke, U.K.) for 1 h. Fluorescence-labeled secondary Abs were added alone as negative controls. Cells were analyzed by using a Leica inverted TCS SPE (Milton Keynes) confocal microscope (\(\times\)40, 1.15 NA, oil immersion objective). Thirty-five conjugates/ experiment were analyzed in at least three experiments.

**Fluorescence recovery after photobleaching experiments**

GHC was quantitatively assessed in living cells by fluorescence recovery after photobleaching (FRAP) assay. Gap junctional dye transfer was measured using the acetoxymethyl (AM) ester derivative of the fluorescent indicator calcein (calcein-AM; Invitrogen). DCs interacting with T cells,
pretreated or not for 4 h with 40 µM murine AS (25) or human Cx43-AS (sequence: 5′-GTAATGCGGCAAGAAGAATTGTTTCTGTG-3′), 40 µM Cx43-sense (25), 300 µM Gap20 control peptide, 300 µM 1848-mimetic peptide, or 50 µM 18β-glycyrrhetinic acid (β-Ga) were collected and loaded with 1 µM calcein-AM in culture medium for 30 min. Once inside the cell, endogenous esterases cleave the lipophilic AM groups, producing fluorescent calcein molecule that is unable to leak out of cells across cell membranes, but is able to pass between cells connected via GJs.

FRAP was performed on a Leica SPUV (Milton Keynes) confocal microscope (×40, 0.8 NA, water immersion objective), using the FRAP function on the Leica confocal software, 1 h after calcein-AM loading. As a control, 1 mM propidium iodide was added to the media to check cell viability during imaging. A region of interest (ROI), either the T cell or the DC, was chosen, and between 12 and 15 cycles of the 488 nM laser at 12% emission strength were used to photobleach the fluorescence within the ROI. These were the conditions determined for optimal bleaching of the DC–T cell conjugates. The progression of FRAP was followed by continuously acquiring images with a time interval of 5 s for 2 min of total imaging time. Fluorescence of the mobile fraction was quantified using the mean ROI function (ImageJ software). Fluorescence intensities of ROIs were recorded before photobleaching, immediately after photobleaching, and at 5-s intervals after photobleaching. The percentage of fluorescence recovery was calculated using the equation for determining the mobile fraction (26).

Measurement of Ca2+ signals

A Leica SPUV confocal microscope (×63, 1.2 NA, water immersion objective) was used for analysis of Ca2+ transients in MCL-specific CD4+ T cells cocultured with MCL-DCs, as well as in OVA-DCs incubated with OT-II T cells. Cells pretreated or not for 4 h with 40 µM Cx43-AS, 40 µM Cx43-sense, 300 µM Gap20 control peptide, 300 µM 1848-mimetic peptide, or 50 µM β-Ga were collected and stained with 1 µM Fluo4–AM (Molecular Probes, Invitrogen). The 1848-mimetic peptide and Gap20 control peptide were added back after washing the cells to remove excess probe. The Fluo4–AM fluorescence and bright field were monitored simultaneously by taking frames at 10-s intervals. Intracellular Ca2+ signals were reported as total mean fluorescence ± SEM and were quantified as Fluo4 fluorescence at any time point-basal fluorescence obtained by averaging the fluorescence of the entire frame. The emission intensity was displayed on a pseudocolor scale using the Leica Lite browser software.

IFN-γ ELISPOT assay

MultiScreen plates (MAPP1450; Millipore, Watford, U.K.) were coated overnight with 2 µg/ml anti-human IFN-γ capture mAb (1-D1K; Mabtech, Stockholm, Sweden). MCL-DCs and autologous MCL-specific CD4+ T cells were preincubated for 4 h in the presence or absence of 300 µM 1848 Cx43-mimetic peptide (sequence CNTQQPGCENVCY extracellular loop 1; 95% purity), 300 µM Gap20 control peptide (EIKKFKYGIEEHC cyttoplasmic loop; 95% purity) (both from JPT Peptide Technology, Berlin, Germany), 50 µM β-Ga, 40 µM Cx43-AS, or vehicle. After this time, 5 × 10^4 DCS were cocultured with T cells at a 1:1 ratio for another 4 h. Additionally, nontreated DCs and T cells were coincubated at a 1:1 ratio for another 4 h with T cells and DCs, respectively, which were pretreated 4 h with the aforementioned drugs or vehicles. IFN-γ spots were counted using an automated counter ELISPOTScan (A.EL.VIS, Hannover, Germany).

Statistics

Statistical analysis was done using the Statgraphics-Plus 2.1 software. Differences between treatments were tested by one-way ANOVA, using Duncan’s multiple comparison procedure or the Mann–Whitney U test for data sets of multiple comparisons. Results are presented as mean ± SD, except where stated. The p values <0.05 were considered statistically significant.

Results

Cx43 accumulation at the T cell stimulatory interface is dependent on the actin cytoskeleton

Beads coated with Abs directed against CD3 and the costimulatory surface receptor CD28 have been widely used to mimic T cell activation by APCs. Cx43 was found to accumulate at the site of contact formed between T cells and beads after stimulation with anti-CD3 and anti-CD28 beads, whereas a random distribution of Cx43 was observed in T cells incubated with beads coated with an irrelevant Ab (Fig. 1A ). Image analysis confirmed that >60% of T cells recruited Cx43 following activation with anti-CD3 plus anti-CD28 beads (Fig. 1B ). In contrast, only 30% of T cells accumulated Cx43 in the presence of anti-CD3– or anti-CD28–coated beads, similar to what was found when using an irrelevant Ab (Fig. 1B ). Furthermore, accumulation of Cx43 and CD3 was found following T cell engagement, suggesting that Cx43 may be recruited to the stimulatory synapse formed between T cells and APCs (Fig. 1C ).

As recent work has provided evidence for direct targeting of Hch1 to cell-cell junctions through a pathway that is dependent on microtubules (27), we investigated whether Cx43 recruitment after TCR engagement was a cytoskeleton-dependent process. The distribution of Cx43 was analyzed by confocal microscopy in human CD4+ T cells, which were incubated with anti-CD3 and anti-CD28 beads in the presence or absence of specific inhibitors of either microtubule or actin polymerization. When microtubule dynamics were inhibited by incubation with taxol or nocodazole, similar accumulation of Cx43 at the site of contact was observed as in the absence of inhibitors (Fig. 2A, 2B, 2D). In contrast, incubation with either latrunculin A or cytochalasin D (inhibitors of actin polymerization) completely abrogated Cx43 accumulation to the contact area (Fig. 2C, 2D). These results indicate that Cx43 recruitment to the synapse is dependent on the actin cytoskeleton.

Cx43 accumulates at the IS pSMAC in an Ag-specific and time-dependent manner

Cx43 distribution was then investigated by confocal microscopy in conjugates of human DCs loaded with a MCL and cocultured with CD4+ T cells that specifically recognize autologous MCL-DCs (Supplemental Fig. 1A). Cx43 was found to accumulate to the interface between T cells and MCL-DCs (Fig. 3A, arrowhead), but was homogenously distributed when T cells were incubated with DCs pulsed with the control Ag gp100 (Fig. 3A). As the MCL contains a number of unknown antigenic peptides, Cx43 accumulation was further investigated using CD4+ T cells from OT-II mice, which carry a transgenic TCR specifically recognizing the OVA323–339 peptide presented in MHC class II. OT-II CD4+ T cells were cocultivated with mature bone marrow-derived DCs (LPS-DCs) or OVA-pulsed mature DCs (OVA-DCs), and synapse formation was visualized by confocal microscopy. Cx43 was found to accumulate at the synapse formed between OT-II T cells and OVA-DCs (Fig. 3B, arrowhead), but was almost completely absent when T cells were incubated with LPS-DCs (Fig. 3B). Quantification showed that >60% of OVA-DC and OT-II T cell conjugates concentrated Cx43 at the T cell–DC interface, and a 4-fold increase of Cx43 accumulated at the contact area, compared with the plasma membrane (Fig. 3C). Similar amounts of Cx43 recruited to the MCL-DCs and MCL-specific T cell interface, and a 3-fold increase of Cx43 accumulated at the contact area was observed when using the human model (Fig. 3C). In contrast, in the absence of Ag-specific presentation (LPS or gp100), fewer conjugates accumulated Cx43, and reduced Cx43 amounts were found at the site of T cell–DC interaction (Fig. 3A–C).

Spatial segregation of accumulated molecules at the IS leads to the formation of cSMAC and pSMAC within the synapse (4). To establish to which compartment Cx43 was recruited, the distribution of Cx43 was compared with the distribution of CD3 (cSMAC) and LFA-1 (pSMAC). The Cx43 pool that redistributed to the IS was only partially colocalized with CD3 (Fig. 3E, Supplemental Fig. 1B), and was found predominantly colocalized with LFA-1 (Fig. 3F, Supplemental Fig. 1C) in both mouse and human models, indicating a preferential recruitment of Cx43 to the
pSMAC ring. This observation of colocalization was calculated by quantification using overlap coefficient according to Manders’s automatic threshold determination. Cx43 displayed 63.1% colocalization with LFA-1, against only 35.7% with CD3. The measured colocalization coefficients for Cx43-LFA-1 and LFA-1-Cx43 were statistically higher ($p < 0.05$) compared with the values for Cx43-CD3 and CD3-Cx43 (Fig. 3D).

Furthermore, serial optical sections along the $z$-axis for Cx43, LFA-1, and TCR labeling on OVA-DCs or LPS-DCs contacting OT-II T cells were taken, allowing 3D reconstruction and projection on the $x$-$z$ plane. Fig. 3G shows representative en face views illustrating the Cx43 accumulation at the DC–T cell interface. Whereas the TCR clustered in the central zone of a T cell in contact with an OVA-DC, Cx43 was excluded from this area and was found colocalizing with LFA-1 at the peripheral zone of contact (Fig. 3G). Cx43 accumulation was rare at the site of interaction of conjugates formed between OT-II T cells and LPS-DCs (Fig. 3G).

The molecular structure of the IS facilitates Ag recognition and T cell activation. To further investigate whether accumulation of Cx43 at the synapse is Ag specific, as well as to evaluate dynamic changes in Cx43 recruitment, the distribution of Cx43 was analyzed over time in both mouse and human systems. Accumulation of Cx43 to the contact area of DCs interacting with T cells was quantified and analyzed, as previously described (22). Significant higher recruitment of Cx43 to the IS was found in conjugates of OT-II T cells and OVA-DCs (Fig. 4A,4B, $p < 0.01$ and $p < 0.005$; Supplemental Fig. 2A, $p < 0.05$). Maximal accumulation of Cx43 was observed 30–45 min after OT-II T cell/OVA-DC incubation (Fig. 4B,4C), whereas in the human model statistically significant differences were observed 45 min after MCL-DC and MCL-specific T cell incubation (Supplemental Fig. 2A). In contrast, when OT-II T cells were incubated with LPS-DCs, or when MCL-specific T cells were cocultured with gp-100-DCs, substantially fewer conjugates accumulated Cx43 at the site of interaction (Fig. 4A–C, Supplemental Fig. 2A). Taken together, these data identify
were incubated for 30 min in the presence or absence (A) of taxol or nocodazole (B), or cytochalasin D or latrunculin A (C), before incubation with CD3- and CD28-coated beads. Cx43 and Hoechst staining were analyzed by confocal microscopy. The inducible capping of Cx43 to the contact area was impaired in the presence of the inhibitors of actin polymerization. Scale bar, 10 μm. D, The number of T cells that accumulate Cx43 at the site contacting the beads was quantified under the different conditions studied. Values are expressed as the percentage of cells that recruit Cx43 to the IS, relative to the total number of cells examined. Differences are indicated by p values (***p < 0.005).

Cx43 as a component of the IS, and suggest that Cx43 recruitment is time dependent and requires cognate Ag recognition by T cells.

Cx43 accumulates at the IS as Cx Hchls

Besides GJs, Cx can also form stand-alone Hchls; therefore, we analyzed whether Hchls may possibly form and accumulate to the IS. To address this, a polyclonal anti–Cx43-Hchl Ab raised to a peptide sequence from the first external loop of Cx43 that recognizes Cx43 in an undocked conformation, and is occluded in docked GJs, was used to examine the distribution of Hchls in conjugates of T cells and DCs in both mouse and human systems. This Cx43-Hchl Ab was used in combination with a Cx43 mAb conjugates of T cells and DCs in both mouse and human systems.

GJs mediate bidirectional communication between DCs and T cells

Bidirectional communication mediated by GJs between cells of the immune system has been previously described (10). The establishment of bidirectional GJIC between DCs and T cells was then monitored by FRAP. OT-II T cells and DCs were loaded with calcine-AM, a fluorescent GJ channel permeant dye, and bleached, and the recovery of fluorescence was monitored for 2 min at intervals of 5 s. Cell viability was verified by propidium iodide exclusion, which was added to the medium and was present throughout the experiment. Cell communication from DCs to T cells, identified as fluorescence recovery, was confirmed 2 min after bleaching OT-II T cells forming conjugates with OVA-DCs (Fig. 5A, 5C, 5D, Supplemental Video 1). Bidirectional transport, in this case from T cells to DCs, was observed when OVA-DCs were photobleached and fluorescence recovery monitored (Fig. 5B, 5D, Supplemental Video 2). In contrast, photobleaching of T cells contacting LPS-DCs, or bleaching of LPS-DCs contacting OT-II T cells showed no fluorescence recovery (Fig. 5A–D), indicating that this is an Ag-dependent process. Inhibiting Cx43 by means of a Cx43-AS oligodeoxynucleotide that targets Cx43 expression in T cells and DCs (Supplemental Fig. 3A–D) completely blocked fluorescence recovery (Fig. 5), confirming that Cx43 is required for functional GJs to form in either direction. In contrast, intercellular communication was not affected when DC–T cell conjugates were incubated with a Cx43-sense oligo control (Supplemental Fig. 3E, 3F). We also investigated fluorescence recovery after incubation with β-Ga or the Cx43-mimetic peptide 1848 that blocks docking adjacent Hchls. Intercellular communication was dramatically reduced after treatment with these inhibitors, although it was not affected in cells treated with the Gap20 control peptide (Supplemental Fig. 3E, 3F).

Moreover, FRAP analysis using MCL-specific T cells and DCs confirmed our findings in the human model (Supplemental Fig. 3G, 3H). These results provide support for the role of Cx43 in mediating bidirectional intercellular communication between T cells and DCs at the IS.

Cx43 is required for APC-mediated T cell activation

Binding of TCR to specific MHC-peptide complexes triggers downstream intracellular events and oscillations of intracellular Ca2+, essential for T cell activation (28). Because previous studies have described Cx43 participation in Ca2+ influx in various cell types (29, 30), we investigated whether Cx43 was involved in regulating Ca2+ signaling in T cells. Calcium signals were monitored over time in T cells coincubated with MCL-DCs and loaded with Fluo4-AM. Ag-specific T cell stimulation resulted in oscillations of intracellular Ca2+, which was impaired when DCs and T cells were preincubated and cocultured with a 2-fold increase of Cx43 accumulated at the contact area, compared with the plasma membrane, was observed 2 h after cognate CD4+ T cell–DC interaction in both human and mouse model (Fig. 4F, Supplemental Fig. 2C, 2D). These data provide evidence for Cx43 accumulation at the IS as stand-alone Hchls.
presence of Cx43 inhibitors (Fig. 6D). Moreover, incubation with the Cx43-AS did not alter the expression of TCR complexes on T cells (Fig. 6E). These data indicate that inhibition of GJIC does not affect signals 1 and 2 of T cell activation; therefore, the impairment in Ca2+ signaling is most likely the result of reduced GJ or Hchl activity.

As interactions between opposing GJ Hchls are a form of intercellular adhesion, the contribution of Cx43 to adhesion of T cells and Ag-pulsed DCs was also investigated. Silencing of Cx43 did not substantially alter cell adhesion, and similar numbers of conjugates were formed between untreated, Cx43-AS–, and Cx43-sense–treated cells (Fig. 6F).

To further investigate whether recruitment of Cx43 to the IS contributes to T cell activation, the secretion of IFN-γ was evaluated by ELISPOT after both T cells and DCs were incubated (or not) with β-Ga, the 1848 Cx43-mimetic peptide, Cx43-AS, or...
FIGURE 4. Cx43 and Cx43-Hchls accumulate at the IS in a time-dependent and Ag-specific way. A, Representative images of Cx43 and LFA-1 distribution after incubation of OVA-DCs or LPS-DCs with OT-II T cells are shown. Scale bar, 5 μm. B, Cx43 accumulation at the IS was evaluated at different time points based on positive costaining of Cx43 and LFA-1 in OVA-DCs or LPS-DCs cocultured with OT-II T cells. Each plotted point represents mean ± SD of three independent experiments (a, p < 0.01 and b, p < 0.005). C, Cx43 distribution to the synapse was measured as ratio of Cx43 accumulated at the contact site versus at the plasma membrane, and was evaluated at different time points. Cx43 accumulation was significantly higher in T cells cocultured with OVA-DCs versus LPS-DCs (30 min, p < 0.01; 45, 60, and 120 min, p < 0.005). Values are expressed as mean ± SEM; n = 3. D, Hchls and Cx43 accumulate at the site of interaction of OVA-DCs and OT-II CD4+ T cells, but distribute randomly in T cells incubated with LPS-DCs. Scale bar, 5 μm. E, The percentage of cells that accumulated Hchls (dark gray) and Cx43 (light gray) at the synapse was assessed. Values are reported as mean ± SEM (a, p < 0.05 and b, p < 0.01). F, The ratio of Cx43 fluorescence accumulated at the contact area versus at the plasma membrane was quantified 2 h after DC-T cell conjugate formation. Values are expressed as mean ± SD of three independent experiments. Differences are indicated by p values (**p < 0.01).
their respective controls. T cell activation was significantly reduced after incubation with each of the aforementioned drugs (Fig. 6G). In contrast, treatments with control vehicle or irrelevant peptide did not inhibit IFN-γ secretion (Fig. 6G). Similarly, when secretion of IFN-γ or IL-2 was analyzed by intracellular FACS following treatment of OT-II T cells stimulated with OVA-pulsed DCs with different GJ or Cx43 inhibitors, T cell activation was found impaired in the murine system as well (Supplemental Fig. 4C–E). Furthermore, we evaluated the individual contribution of Cx43 from T cells or DCs to the T cell activation process. When GJ activity was inhibited in DCs only, significant reduction of IFN-γ secretion was detected after preincubation with β-Ga or the Cx43-AS ($p < 0.01$ and $p < 0.05$, respectively), but only a slight decrease was obtained after pretreatment with the 1848-mimetic peptide (Fig. 6H). In contrast, T cell activation was more dramatically impaired after preincubation of T cells with any of these
FIGURE 6. Cx43 contributes to T cell activation by DC. A, Ca\textsuperscript{2+} signaling was analyzed using Fluo4-AM by sequential confocal images of MCL-specific T cells cocultured with autologous MCL-DCs and treated with different GJs or Cx43 inhibitors and their respective controls. Phase-contrast images corresponding to the same fields are shown. Scale bar, 5 µm. B, Time course showing changes of intracellular Ca\textsuperscript{2+} signaling in T cells contacting DCs, under different conditions. Ca\textsuperscript{2+} signals are shown as total mean fluorescence ± SEM. C, Overall mean of Fluo4-AM fluorescence over the time ± SEM for each condition (*p < 0.05; n = 3. D and E, Treatments with a Cx43-sense or with the 1848 Cx43-mimetic peptide did not affect the expression of MHC class I and class II, CD40, CD83, and TCR. F, Cell adhesion was evaluated in conjugates of OVA-DCs and OT-II T cells treated or not with a Cx43-sense or Cx43-AS. Cx43 gene targeting did not affect conjugate formation 5 min and 30 min after DC–T cell coincubation. Each bar represents percentage ± SD of four independent experiments. G, IFN-γ secretion was assessed by ELISPOT assay in MCL-specific CD4\textsuperscript{+} T cells coincubated with MCL-DCs, plus a nonspecific GJ blocker (β-Ga), Cx43-AS, or 1848 Cx43-mimetic peptide. Inhibition of GJIC significantly reduced the secretion of IFN-γ by CD4\textsuperscript{+} T cells. Data were expressed as the mean of spots/5 × 10\textsuperscript{3} effector cells ± SD (***p < 0.01, *p < 0.05), n = 2, performed in triplicate. H, MCL-DCs or MCL-
specific T cells were independently pretreated with β-Ga, 1848-mimetic peptide, Cx43-AS, or their respective controls, and then incubated with nontreated T cells (dark gray) or DCs (light gray), respectively. Control, nontreated T cells, and DCs (black) were also evaluated. Graphic represents IFN-γ secretion reported as the mean of spots/5 × 10^3 CD4^+ T cells ± SD (*p < 0.05, **p < 0.01, ***p < 0.005), n = 2, performed in triplicate.
of Cx43-GJ function substantially diminished IFN-γ secretion by primed T cells. In addition, cytokines can positively regulate the surface expression of Cx and GJIC in cells of the immune system (17), suggesting the existence of a positive feedback regulation by cytokines, such as IFN-γ, which can stimulate opening of GJ channels and Cx43 upregulation (13). Such a mechanism may contribute to sustained communication between T cells and APCs, allowing optimal T cell activation. Although we have shown that GJIC is important for T cell activation, we cannot exclude a possible contribution for Hchls to this process. The fact that a Cx43-mimetic peptide has also affected IFN-γ secretion by T cells following incubation with anti-CD3 and anti-CD28 beads suggests that these structures are also involved in T cell activation. This role additionally correlates with our findings of Hchl accumulation at the site of contact between T cells and DCs. As targeting of Cx43 protein expression or the use of a specific Cx43-mimetic peptide blocks both GJ and Hchl, further studies using Cx43 mutants may provide useful tools to discern the contribution of Hchl to Ag-specific T cell activation.

In summary, this work identifies Cx43 as a key component of the IS and provides evidence of a role for GJ and Hchl in T cell activation, opening new questions regarding the involvement of these structures in the regulation and synchronization of immunological processes.

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Disclosures

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