Purinergic receptor (P2X7) activation reduces cell–cell adhesion between tubular epithelial cells of the proximal kidney

Eleftherios Siamantouras, PhD, Gareth W. Price, PhD, Joe A. Potter, PhD, Claire E. Hills, PhD, Paul E. Squires, PhD∗

Joseph Banks Laboratories, School of Life Sciences, Green Lane, University of Lincoln, UK

Rev 6 October 2019

Abstract

Loss of epithelial (E)-cadherin mediated cell–cell adhesion impairs gap junction formation and facilitates hemichannel-mediated ATP release in the diabetic kidney. Linked to inflammation and fibrosis, we hypothesized that local increases in inter-cellular ATP activate P2X7 receptors on neighboring epithelial cells of the proximal tubule, to further impair cell–cell adhesion and ultimately exacerbate tubular injury. Immunoblotting confirmed changes in E-cadherin expression in human kidney cells treated with non-hydrolysable ATPγS ± the P2X7 antagonist, A438079. Atomic force microscopy based single-cell force spectroscopy quantified maximum unbinding force, tether rupture events, and work of detachment. Confocal microscopy assessed cytoskeletal reorganization. Our studies confirmed that ATPγS downregulated E-cadherin expression in proximal kidney cells, loss of which was paralleled by a reduction in intercellular ligation forces, decreased tether rupture events and cytoskeletal remodeling. Co-incubation with A438079 restored loss of adhesion, suggesting that elevated extracellular ATP mediates tubular injury through P2X7 induced loss of E-cadherin mediated adhesion.

© 2019 . Published by Elsevier Inc. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).

Key words: Kidney; Adherens junction; Cell adhesion; ATP; P2X7; AFM-SCFS

Diabetic nephropathy is the leading cause of End Stage Renal Disease (ESRD) in people with diabetes.1 Characterized by multiple structural and functional disturbances, tubulointerstitial fibrosis of the proximal region of the kidney represents a major underlying pathology of diabetic nephropathy and develops in response to a number of morphological and phenotypic changes.2 We have previously reported that glucose-evoked increases in the beta1 isoform of the profibrotic cytokine Transforming Growth Factor (TGFβ1), drive a loss of epithelial associated markers, with a concomitant increase in proteins more commonly associated with fibroblasts.3 Often referred to as partial epithelial-to-mesenchymal transition (pEMT), this series of events is initiated in response to impaired expression of epithelial (E)-cadherin and the functional loss of cell–cell adhesion.4–6 Decreased cell–cell adhesion prevents docking of connexons on neighboring membranes and decreases Gap Junction-mediated Intercellular Communication (GJIC).7 In the absence of neighboring binding partners, uncoupled connexons form hemichannels that release nucleotides, e.g. adenosine triphosphate (ATP), in to the intercellular space.8 Locally released ATP binds purinoceptors, notably P2X7, and is linked to the progression and development of inflammation and fibrosis in multiple tissue types,9–11 including the diabetic kidney.12–15 With our recent studies linking increased hemichannel mediated ATP release to both inflammation and fibrosis, it seems likely that increased intercellular ATP may accelerate progression of tubular injury by exacerbating the further loss of cell–cell tethering.

Part of the multi-protein adherens junction (AJ), cadherins are calcium-dependent trans-membrane cell adhesion proteins that connect the cell interface to actin cytoskeleton (CSK). The extracellular domains of individual cadherins form weak (40pN) binding pairs16 with cadherins on adjacent cells, while the cytoplasmic domain bind β-catenin, linking cadherins to the cytoskeleton via α-catenin.17,18 By regulating ligation clustering, interaction of cadherin with F-actin, via catenins, is crucial...
in maintaining adhesive strength and acts as a signaling ‘node’ for proteins that influence adhesiveness & initiate intracellular signaling.\textsuperscript{19,20} In recruiting catenins, cadherins trigger signals that modulate the actin cytoskeleton at the point of cell–cell contact.\textsuperscript{21}

The intricate molecular structure of the adherens junction and viscoelastic properties of the cytoskeleton suggest that separation of adherent cells is a complex process. Recent efforts to systematically investigate the effects of mechano-chemical changes in cell–cell adhesion in diabetic nephropathy\textsuperscript{6} and other pathological states have been reported.\textsuperscript{22–24} It is clear that the assembly of AJJs that modulate the actin cytoskeleton at the point of cell–cell contact, -

Although initially used to study molecular function of binding receptors,\textsuperscript{26} AFM-based force spectroscopy has been applied to investigate the separation behavior of whole cells.\textsuperscript{27,28} In the current study we applied AFM-based single-cell force spectroscopy (SCFS) to investigate the effects of ATP in mediating loss of adhesion between tubular epithelial cells of the proximal kidney via P2X7 purinoceptors. The AFM-based instrument incorporates an extended displacement range (100 μm) to allow complete separation of adherent cells for nanoscale resolution. Our novel analytical data clarify the mechanism by which the loss of cell–cell adhesion is regulated by the physicochemical interplay between the extracellular and intracellular domains of E-cadherin, and provide compelling evidence that ATP-mediated paracrine signaling is important in progression of early tubular injury in kidney disease.

Methods

Reagents

Fibronectin and ATPγS were obtained from Sigma (Poole, UK). P2X7 receptor antagonist A438079, tetramethyl rhodamine isothiocyanate (TRITC)-conjugated phalloidin, DAPI (4′,6-
diamidino-2-phenylindole) and goat serum were purchased from Bio-Technne (Abingdon, UK). Immobilon-Fi PVDF membrane was from Millipore (Watford, UK), while Odyssey blocking buffer and secondary fluorescent antibodies were purchased from LI-COR (Cambridge, UK). Anti-E-cadherin was purchased from Cell Signalling Technologies (Hertfordshire, UK), while alpha-tubulin was purchased from Sigma (Poole, UK).

Culture of HK2 cells

Human derived proximal tubule kidney (HK2) epithelial cells were purchased from the American Type Culture Collection (ATCC; Gaithersburg, MD 20878). Tissue culture media and plasticware were from Invitrogen Life Technologies (Paisley, UK). HK2 cells (passage 18–30) were maintained in DMEM/Ham’s F12 (DMEM/F12) medium, supplemented with 10% fetal calf serum (FCS), glutamine (2 mmol/l), 2% penicillin–streptomycin, and epidermal growth factor (EGF; 5 ng/ml). Cells were seeded onto 40 mm petri-dishes, T25 flask (for suspended cells) and cultured at 37 °C in a humidified atmosphere of 5% CO2 in air. Prior to treatment, cells were cultured in DMEM/F12 low glucose (5 mM) for 48 h. Basal (5 mM) glucose culture media were generated as described previously.\textsuperscript{3} Cells were then treated with ATPγS (100 μM) + A438079 (50 μM) for 48 h. In all experiments, cells were serum starved overnight before stimulation.

Western blotting

Preparation of protein, its separation by SDS-gel electrophoresis and transfer onto Immobilon-Fi PVDF membranes have been described previously.\textsuperscript{3} Membranes were blocked with Odyssey blocking buffer overnight (LI-COR), then probed simultaneously with a polyclonal antibody against E-cadherin (1:1000) and alpha-tublin (1:20,000) for 1 h at room temperature. Bands were visualized using an Odyssey FC and semi-quantified using Image Studio (v5.2, LI-COR).

AFM single-cell force spectroscopy

AFM is a powerful tool for high-resolution single cell force measurements. Briefly, a piezo actuator moves the base of a cantilever towards the surface of the sample in the vertical direction and then retracts it again, while the deflection of the cantilever is measured continuously. An SCFS force curve is the result of mechanical interaction between the tip of the cantilever and the surface of the sample. When the cantilever is retracted from the sample adhesion occurs, in which the cantilever is still in contact with the sample. During this process the cantilever is deflected downwards, and adhesion can be detected in a force curve by a negative force peak (Figure 1). As the cantilever is further retracted from the surface, adhesion forces will be disrupted, and the tip will be completely separated from the sample. Using spring stiffness, the deflection of the cantilever provides information about the elastic properties of the sample and a direct measure of the adhesion forces. Force spectroscopy between single, soft biological cells includes more complex interactions between the tip and the sample. As the cantilever is retracted, hysteresis is observed, which is common for
viscoelastic materials. In addition, adhesion of the tip with long 
surface molecules requires an extended displacement range to 
avoid extendable contacts. The cantilever-attached cell was 
brought in contact with substrate-attached cell, until a pre-set 
contact force of 1nN was reached. The cells were allowed to 
form ligation during a contact time of 10 s, after which the 
cantilever was retracted using a velocity 5 μm/s until the two 
cells were completely separated. The procedure was repeated 
three times for each cell tested, with 45 s intervals between each 
measurement. Velocity was kept constant during the extend and 
retract processes. Data were collected from three separate 
experiments (n = 3).

**Instrumentation**

Latest advancements in AFM-force spectroscopy allow for an 
effective displacement range (100 μm), sufficient to disrupt 
adsorption forces needed for complete cell separation and facilitate 
the investigation of cell-to-cell contact. Experiments were 
performed using the CellHesion®200 module (JPK Instruments, 
Berlin, Germany) that was installed on an Eclipse TE 300 
inverted microscope (Nikon, USA). During each experiment, 
cells were maintained at 37 °C by incorporating the BioCell™ 
temperature controller (JPK, Berlin, Germany) into the AFM 
stage. Phase microscopy images were acquired using a CCD 
camera (Orca, Hamamatsu) connected on the side port of the 
microscope. To avoid hydrodynamic drag when performing F-d 
curves in liquid a speed of 5 μm/s was used. The AFM-SCFS set-
up with the CCD camera was driven by JPK's CellHesion200 
software. Images were captured at x20 magnification. The 
system was supported on an anti-vibration table (TMC 63-530, 
USA). Changes in the temperature of the room were less than 
0.5-1 °C during experimental measurements. Deflection of the 
cantilever was measured by the difference in the reflection of a 
laser beam between the upper or lower parts a quadric-sected 
photodiode.

**Functionalization**

To perform cell–cell adhesion experiments, suspended cells 
were introduced into the media of testing cells and a single cell was 
attached on a tipless cantilever. An arrow geometry is designed to 
ease manipulation of individual cells among others in close 
proximity. Arrow sensors (TL1, Nanoworld AG, Switzerland) 
with force constant of 0.03 N/m were used to attach cells to the free 
end of the sensor. Tipless cantilevers are more suitable for adhesion 
experiments since any contact of the tip with the surface of the cell 
or the substrate will disrupt the measurements. The attached cell 
was probing an adherent cell on the substrate in order to investigate 
the disruption forces between two cells. Tip-less cantilevers were 
chemically functionalized so that a single suspended cell could be 
attached. Initially the cantilevers were sterilized by UV treatment 
(10mins). Next, they were incubated in poly-L-lysine (25 μg/ml in 
PBS) for 30mins at room temperature (RT). Subsequently, the 
cantilevers were transferred in fibronectin solution (20 μg/ml in 
PBS) and they were incubated for 2 h at 37 °C. After 
functionalization cantilevers were stored in PBS solution at 4 °C 
and used within 3 days.

![Figure 1. (A) Phase microscopy images of probe cells attached on the cantilever from each treatment. Clusters of treated cells are firmly attached in the substrate. The probe cell was brought in contact above the central point of the nucleus for 10 s with a contact force of 1 nN. The probe cell was then retracted at a 5 μm/s while force vs displacement was continuously recorded. (B) A retraction F-d curve of a control cell illustrating the experimental process. In phase 1 the probe cell is moving vertically downwards until the preset force is reached. During phase 2, interaction between the two cells occurs in which ligation is formed. Lastly, in phase 3 the probe cell is retracted until the force returns to the baseline where the two cells are separated.](image-url)
Calibration

For small deflections, the cantilever approximates a Hookean spring, hence the deflection is linearly related to the acting force. When the cantilever spring constant is calculated, the deflection can be converted into the corresponding force \((F = k \cdot x)\). Each functionalized cantilever was calibrated prior to experimentation to determine the actual spring constant value, using the manufacturer’s software (JPK Instruments, Germany) based on the thermal noise amplitude.\(^{29,30}\) This method measures the thermal fluctuations of the cantilever deflection and uses the equi-partition theorem to calculate the cantilever spring constant. Initially the deflection of the cantilever is displayed as the output of the photodetector in Volts (\(>1\) V). To record a force curve for calibration, the cantilever was configured to approach the base of a cell-free petri-dish once, to minimize the loss of coating (set-point <1 V). This linear curve was used to determine sensitivity, or the distance of cantilever deflection for a given voltage difference measured by the photodiode, which was below 100 nm/V for all cantilevers used. The amplitudes of the thermal fluctuation were measured at \(37^\circ\)C. Since the calibration process was conducted in culturing media a correction factor of \(0.251\) was used.\(^{30}\) The mean spring constant was \(0.02024\) N/m, range: \(0.0098, \text{STD:} 0.00348\). The position of the cantilever on the glass holder and the alignment parameters were maintained throughout experiments as any disturbance would result in recalibration.

Immunocytochemistry

Cells were seeded onto glass-coverslips (22 mm diameter) in low glucose (5 mM) for 48 h, serum-starved overnight, and then incubated with ATP\(\gamma\)S (100 \(\mu\)M) \(\pm\) A438079 (50 \(\mu\)M) for 48 h. Cells were fixed with 4% paraformaldehyde, blocked with goat serum, and stained with DAPI (1 mmol/l) for 3 min, before being incubated with tetramethyl rhodamine isothiocyanate (TRITC)-conjugated phalloidin (1:400 in PBS-Triton) for 1 h. Cells were visualized using an inverted Leica confocal microscope, using a hybrid Leica HyD detector. Leica Application Suite X (LAS X) software was used to operate the confocal hardware and overlay resulting images.

Data processing and analysis

Three force-displacement curves were acquired per cell during experiments and were processed using the JPK data processing software. The separation parameters analyzed in this study are the maximum unbinding force \(F_{\text{max}}\), the work of detachment \(W_d\) and the number of tether rupture events (TREs). To signify statistical significances data were evaluated using univariate ANOVA followed by Tukey’s multiple comparisons post-test using SPSS (version 24). Data from SCFS are expressed as mean \(\pm\) SEM in the text and as median and quartile range at the graphs. Sample numbers refer to separate cell passages made up of recordings from multiple individual cells and are represented with ‘\(n\)’ for passage number. \(P < 0.05\) was taken to indicate statistical significance.

Results

ATP\(\gamma\)S downregulates E-cadherin expression via activation of P2X7 receptors

We recently demonstrated that glucose-evoked changes in TGF-\(\beta\)1-evoked cytoskeletal remodeling and reduced cellular adhesion,\(^{6}\) events that culminated in a loss of gap junction intercellular communication and increased hemichannel mediated ATP release.\(^{8}\) To determine whether ATP propagates this process, HK2 cells were cultured in 5 mM glucose for 48 h, serum-starved overnight, and subsequently incubated with the non-hydrolysable ATP\(\gamma\)S (100 \(\mu\)M) for 48 h. In control cells, TRITC-conjugated phalloidin confirmed that the actin cytoskeleton was made up of a diffuse transcellular network of F-actin filaments that spanned the cytosol. Treatment with ATP\(\gamma\)S produced denser peripheral stress fibers (Figure 2, A), an effect reversed when co-incubated with the P2X7 antagonist A438079 (50 \(\mu\)M). Immunoblotting confirmed that ATP\(\gamma\)S decreased whole-cell expression of E-cadherin to 54.4 \(\pm\) 4% (Figure 2, B; \(P < 0.01, n = 3\)), an effect reversed when HK2 cells were co-incubated with ATP\(\gamma\)S (100 \(\mu\)M) and the selective P2X7 inhibitor, A438079 (50 \(\mu\)M, 113.6 \(\pm\) 9.7%; Figure 2, B; \(n = 3\)).

ATP\(\gamma\)S decreases maximum unbinding force \((F_{\text{max}})\) via activation of P2X7 receptors

AFM-based single cell force spectroscopy (SCFS) was used to quantify the unbinding forces required to separate two cells. The minimum negative point during the pulling phase of the \(F-d\) curve represents the maximum downward deflection of the cantilever as the probe cell is retracted from the substrate cell in the vertical axis. Therefore, the minimum force value of a retraction curve corresponds to the maximum unbinding force \(F_{\text{max}}\) (Figure 3, B). Nanoscale forces are correlated with any alteration in the number of E-cadherin ligations and the associated formation of clusters that promote adhesion. Using SCFS the maximum unbinding forces were quantified after P2X7 activation. Force and contact time remained constant throughout the experiments, while the approach-retract cycle was repeated three times for each cell. To determine \(F_{\text{max}}\), the baseline in which the cells were completely separated was identified in order to calculate the difference \(\Delta F\) (nN) to the maximum negative point of force.

Results showed that when cells were treated with ATP\(\gamma\)S (100 \(\mu\)M) (\(\mu = 1.6nN \pm 0.093, 95\% \text{ CI} [1.42, 1.786]\)), a 26.8% reduction in \(F_{\text{max}}\) occurred, in comparison to control (\(\mu = 2.17nN \pm 0.077, 95\% \text{ CI} [2.023, 2.325]\)) (Figure 3, A, \(P < 0.001, 95\% \text{ CI} [0.287, 0.856]\); Control: 27 cells, ATP\(\gamma\)S: 19 cells, \(n = 3\)). This correlates to weakening of intercellular ligation forces as a consequence of diminished E-cadherin expression (Figure 2, B). To delineate a role for P2X7 in mediating these effects, cells were co-incubated with ATP\(\gamma\)S \(+\) A438079 (50 \(\mu\)M). Mean value of \(F_{\text{max}}\) was increased to 2.45nN \(\pm\) 0.085, 95% CI [2.284, 2.621] showing a restoration of cell–cell ligation forces when co-incubated with A438079 (\(P < 0.001, 95\% \text{ CI} [0.551, 1.148]\); +A438079: 22 cells, \(n = 3\)). Box plots and histograms of the results are shown in Figure 3, A & C respectively.
ATPγS decreases tether rupture events (TREs) via activation of P2X7 receptors

Retraction $F-d$ curves of soft cells have a unique pattern that resemble a step-like profile, corresponding to the unbinding of finite size adhesion clusters that extend as displacement increases. In the early part of the separation process (10 μm) complex unbinding events occur ($j'$ events) that are preceded by a force ramp. As the pulling distance increases a plateau in the displacement indicates that membrane tethers extrude rupture of ligation clusters ($t'$ events). The number of tether rupture events (TREs) was detected by identifying sharp steps of force that correspond to cluster ruptures. The retraction $F-d$ curve of Figure 4, B illustrates the number of TREs as detected by the step fitting function. During the continuous pulling speed only upwards events are anticipated, hence only positive steps were detected to avoid drifts of the cantilevers.

Analysis of retraction $F-d$ curves showed that ATPγS (100 μM) treated cells ($μ = 44.69 ± 2.4$, 95% CI [39.989, 49.382]), exhibit a 22.3% reduction in the number of TREs compared to control ($μ = 57.49 ± 2.0$, 95% CI [53.508, 61.478]), (Figure 4, A, $P < 0.001$, 95% CI [5.432, 20.184]); Control: 25 cells, ATPγS: 18 cells, $n = 3$). Coincubation with ATPγS ± A438079 ($μ = 68.75 ± 2.2$, 95% CI [64.436, 73.064]) restored the number of the rupture events, increasing TREs by 53.84% as compared to ATPγS alone ($P < 0.001$, 95% CI [16.428, 31.702]; +A438079: 22 cells, $n = 3$). These data are in agreement with $F_{max}$ and E-cadherin expression results. Box plots and histograms of the results are shown in Figure 4, A & C respectively.

ATPγS decreases the work of adhesion ($W_d$) via activation of P2X7 receptors

The total energy consumed during the pulling process prior to complete separation (defined as the baseline above), was calculated by the integration of the retraction $F-d$ curve and is referred as the work of adhesion $W_d$. Since the pulling force is
integrated over the separated distance (gray area in Figure 5, B), $W_d$ is driven by molecular changes in E-cadherin ligation and F-actin cytoskeleton. Results showed that ATP$\gamma$S (100 μM) treated cells ($\mu = 13.32fJ \pm 1.13$, 95% CI [10.702, 15.93]) exhibited a 39.04% reduction in $W_d$ compared to control ($\mu = 21.85fJ \pm 1.1$, 95% CI [19.677, 24.013]), (Figure 5, A, $P < 0.001$, 95% CI [4.462, 12.596]; Control: 27 cells, ATP$\gamma$S: 19 cells, $n = 3$), an effect negated when co-incubated with the P2X7 antagonist ($\mu = 23.73fJ \pm 1.22$, 95% CI [21.323, 26.133]). Coincubation with ATP$\gamma$S ± A438079 ($\mu = 23.73 \pm 1.219$, 95% CI [21.323, 26.133]) restored the work of adhesion as compared to ATP$\gamma$S alone ($P < 0.001$, 95% CI [6.158, 14.665]; +A438079: 22 cells, $n = 3$). No statistical significance between control and + A438079 was found. Box plots and histograms of the results are shown in Figure 5, A & C respectively.

Discussion

Hemichannel-mediated release of ATP into the intercellular space around tubular epithelial cells of the diabetic kidney has been linked to disease progression. Understanding the effects of ATP in exacerbating the loss of cell–cell adhesion is fundamental to advancing therapeutic strategies that may help alleviate the effects of this debilitating complication. Nanoscale force-displacement measurements provide a multi-variable signature of the cell–cell separation process that captures the complexity of the adherens junction and its integrated response to purinergic receptor activation. We have used SCFS to investigate the effects of ATP$\gamma$S on cell–cell adhesion in renal proximal tubule (HK2) epithelial cells. The HK2 cell line maintains functional characteristics of human proximal tubular epithelium and has been previously used to characterize alterations in E-cadherin mediated cell–cell adhesion. The organization of surface ligation bindings into finite sized clusters is evidenced by the step-like pattern of the retraction curve. After initial cell–cell contact, a variety of processes, such as junction formation and intercellular signaling are initiated. The initial recognition event involves the formation of trans-dimers between cadherin monomers located on the interface of junctional contact followed by the clustering of cadherin into larger structures to form adhesion. The organization of E-cadherin in to clusters depends on links with cytoskeleton. SCFS has the advantage of functional characterization of adhesion at the cellular level by examining E-cadherin ligation at the EC domain in correlation with the machinery that regulates its function. Adhesion parameters derived from separation experiments showed a variable response to integrated changes of the AJ-CSK architecture. As evidenced by changes in $F_{max}$ and TREs, the AJ-CSK complex regulates cell adhesion by physico-chemical modulation of the number of competent ligations and associated clustering organization. Changes in mechanical properties of cells, due to cytoskeletal re-organization, affect their separation response by modulating deformation as shown by changes in $W_d$. Previous data suggest that single cell mechanics strongly correlate with changes in the adhesion energy rather than binding forces at the whole cell level.
The concentration of ATP in the cytoplasm of cells is high (1-10 mM), while levels in the extracellular space are extremely low, ranging between 1 and 10 nM. In diabetic nephropathy, glucose-evoked TGFβ increases hemichannel-mediated ATP release. Designed to maintain cellular communication following a loss in gap junction intercellular communication in disease, the local rise in ATP activates P2X7 receptors to exacerbate renal injury. In the current study, we present novel evidence that treatment of HK2 cells with ATPγS dramatically reduces all adhesion parameters measured. The maximum unbinding force is the force required to break ligation between E-cadherin clusters on coupled cells. Reduction of $F_{\text{max}}$ clearly suggests that the number of cadherin dimers was reduced after P2X7 activation in agreement with whole cell expression of E-cadherin. In addition, ATPγS reduced the number of tether rupture events, suggesting that the number of competent binding molecules available to form clusters was reduced and the organizational function of ligation was compromised, a finding in agreement with reduced unbinding forces, E-cadherin expression and cytoskeletal re-organization. Mean values showed a negative percentage change of 26.8% and 22.6% for $F_{\text{max}}$ and TREs respectively. Reduced rupturing between cadherin clusters demonstrates the inability of ATPγS-treated cells to organize binding molecules into clusters to ensure appropriate ligation. Evidence in the literature suggests that this loss of this function is associated with cytoskeletal re-organization. The work of detachment ($W_d$) was reduced dramatically (39%) after ATPγS treatment. Considering that actin polymerization mostly affects $W_d$ during separation due to mechanics, the considerable loss in work of detachment in comparison to unbinding force (39% vs 27%) can be explained by changes in the cytoskeleton after treatment rather than any effects on the extracellular interface associated with ligation. Therefore, apart from its role to reinforce ligation clustering, re-arrangement of the cytoskeleton into peripheral stress fibers biophysically affected the work of detachment. This is clearly shown in $F-d$ curves, where $W_d$ is a function of maximum unbinding forces and the last unbinding event, which is a measure of distance. As a result, the underlying molecular machinery associated with the determination of cell deformation during separation, involved changes in both surface avidity and actin cytoskeleton, with the latter dominating the process as shown by SCFS.

Cytoskeletal reorganization following ATPγS treatment affected biophysically the work of detachment and molecularly the organization of ligation clustering. Inhibition of P2X7 restored cell–cell adhesion. Force-displacement ($F-d$) retraction curves suggest that membrane molecular binding and architecture of the AJ-CSK complex were re-established; $F_{\text{max}}$ and TREs were significantly increased (53%) in agreement with an increase in whole cell expression, suggesting that the number of competent adhesion ligations and clustering organization was restored, in correlation with a markedly increase in $W_d$ (78%) after P2X7 inhibition. Biophysically, restoration of the deformation ability of cells after cytoskeletal polymerization explains the increase in $W_d$. Determining strength of adhesion in disease
is challenging due to the underlying molecular assembly that regulates AJ formation. Force-displacement measurements during cell–cell separation can provide valuable information about the response to ligands and their antagonists. Nanoscale SCFS retraction curves capture the molecular activity underlying ATP evoked changes in cell adhesion, loss of which appears to be mediated by downstream P2X7 receptor activation and supports a role for P2X7 as a potential therapeutic target in managing progression of diabetic nephropathy.

References


