Protein exchange is reduced in calcium-independent epithelial junctions

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Desmosomes are cell–cell junctions that provide mechanical integrity to epithelial and cardiac tissues. Desmosomes have two distinct adhesive states, calcium-dependent and hyperadhesive, which balance tissue plasticity and strength. A highly ordered array of cadherins in the adhesive interface is hypothesized to drive hyperadhesion, but how desmosome structure confers adhesive state is still elusive. We employed fluorescence polarization microscopy to show that cadherin order is not required for hyperadhesion induced by pharmacologic and genetic approaches. FRAP experiments in cells treated with the PKCa inhibitor Go6976 revealed that cadherins, plakoglobin, and desmoplakin have significantly reduced exchange in and out of hyperadhesive desmosomes. To test whether this was a result of enhanced keratin association, we used the desmoplakin mutant S2849G, which conferred reduced protein exchange. We propose that inside-out regulation of protein exchange modulates adhesive function, whereby proteins are “locked in” to hyperadhesive desmosomes while protein exchange confers plasticity on calcium-dependent desmosomes, thereby providing rapid control of adhesion.

Introduction

Many vital cellular processes including gene expression, cell division, and motility, are dependent on macromolecular complexes. Higher-level features of these complexes including protein architecture, order, organization, and dynamics, are all critical regulators of function. Importantly, complexes that appear static can adopt multiple conformational states (Vrabioiu and Mitchison, 2006), act as depots of regulatory proteins (Ray et al., 2007), and support exchange of protein components (Daigle et al., 2001; Griffiths et al., 2003). Understanding this multifaceted regulation is key to deciphering the functions of macromolecular complexes in health and disease.

Cell junctions represent a class of plasma membrane-associated macromolecular complexes with roles in adhesion, force transmission, and electrical connections (Garcia et al., 2018; Goodenough and Paul, 2009; Parsons et al., 2010). To perform these myriad functions, cell junctions have complex architectures that are key in signal integration and dynamic regulation (Bertocchi et al., 2017; Kanchanawong et al., 2010; Kaufmann et al., 2012; Mehta et al., 2016; Nahidiazar et al., 2015; Stahley et al., 2016). Epithelial cells have two similar yet distinct adhesive junctions that span neighboring cells: desmosomes and adherens junctions. These junctions share the role of mediating cell–cell adhesion and are architecturally analogous, with adhesive cadherin cores linked to the cytoskeleton through a network of proteins. Despite these similarities, adherens junctions and desmosomes are molecularly and functionally distinct (Rübsam et al., 2018).

One key functional difference is the ability of desmosomes to adopt a calcium-independent, or hyperadhesive, state (Wallis et al., 2000). Whereas adherens junctions and calcium-dependent desmosomes disassemble and lose function upon chelation of extracellular Ca2+, hyperadhesive desmosomes maintain adhesion when Ca2+ has been removed (Garrod, 2010; Garrod et al., 2005; Wallis et al., 2000). These two functional states allow rapid and precise tuning of adhesion to balance tissue strength and plasticity in a multitude of processes. For example, during development and tissue remodeling, desmosomes are calcium-dependent and plastic, but ultimately become static and hyperadhesive in mature tissue (Kimura et al., 2012). In the epidermis, desmosomes have different adhesive strengths in basal versus suprabasal cells (Garrod and Kimura, 2008; Harmon and Green, 2013). During wound healing, desmosomes in suprabasal keratinocytes revert to a calcium-dependent state to promote cell migration and wound closure (Garrod et al., 2005; Owen et al., 2008). Conversion between these adhesive states is controlled by PKCa. Inhibition of PKCa induces hyperadhesion, likely owing to the loss of phosphorylation of desmoplakin (DP; Garrod et al., 2005; Wallis et al.,
Hyperadhesion can also be conferred by overexpression of the DP mutant S2849G, which cannot be phosphorylated at that site (Albrecht et al., 2015; Hobbs and Green, 2012). Conversely, hyperadhesive desmosomes can be converted to calcium-dependent by activation of PKCa (Wallis et al., 2000). In this way, regulation of PKCa allows for rapid and precise control of the desmosome adhesive state.

It is not known how desmosome architecture impacts the adhesive state. Because cadherins mediate adhesion by mechanically coupling neighboring cells, they are an obvious candidate for defining function. Classic and desmosomal cadherins are type I transmembrane proteins with five extracellular cadherin (EC) domains with interdomain Ca\(^{2+}\) binding sites. The cadherin tertiary structure is rigid when Ca\(^{2+}\) is bound and disorganized without Ca\(^{2+}\) (Harrison et al., 2016; Pokutta et al., 1994; Sotomayor and Schulten, 2008). Structurally, desmosomal cadherins have a more “bent” conformation (Harrison et al., 2016) and exhibit greater flexibility (Tariq et al., 2015) than classical cadherins, both features that have been proposed to play roles in accommodating hyperadhesion. In tissues, desmosomes have a characteristic dense midline bisecting the extracellular space, as characterized by EM. This dense midline corresponds with overlapping EC1 domains at the site of trans binding and is found solely in hyperadhesive desmosomes (Garrod et al., 2005; He et al., 2003; Shimizu et al., 2005). An ordered and periodic organization of cadherins in the extracellular space of hyperadhesive desmosomes has been proposed (Rayns et al., 1969; Al-Amoudi et al., 2005, 2007). These characteristics support the hypothesis that the arrangement of the cadherin EC domains into quasi-crystalline ordered arrays is a structural feature defining hyperadhesive desmosomes, which may drive function (Garrod, 2013).

Herein we determine the unique architectural features driving hyperadhesion by quantifying cadherin order, plaque organization, and protein dynamics in calcium-dependent and hyperadhesive states. We found that cadherin order and desmosome architecture did not correlate with adhesive state. Surprisingly, cadherins lost order upon Ca\(^{2+}\) chelation even in hyperadhesive desmosomes. Desmosomes with disordered cadherins maintained adhesion, eliminating cadherin order as the driving factor behind hyperadhesion. Using fluorescence recovery after photobleaching (FRAP), we identify protein exchange in and out of desmosomes as a key difference between calcium-dependent and hyperadhesive states. We show that protein exchange can be controlled via DP phosphorylation, providing a rapid switch in protein dynamics between static hyperadhesive and dynamic calcium-dependent desmosomes. These findings reveal a potential novel mechanism for conferring desmosome hyperadhesion and regulating cell adhesion.

**Results**

**Cadherin order is not required for adhesion**

Desmosomes can be switched from calcium-dependent to hyperadhesive by inhibition of PKCa (Garrod et al., 2005; Kimura et al., 2007; Wallis et al., 2000). To study changes associated with hyperadhesion, we used the PKCa inhibitor GÖ6976 (Hobbs and Green, 2012). Immortalized human keratinocyte (HaCaT) cells were treated with 50 nM GÖ6976 or vehicle (DMSO; mock). To confirm hyperadhesion, cells were incubated in low-Ca\(^{2+}\) medium (Ca\(^{2+}\)-free DMEM with 10% chelated FBS and 3 mM EGTA) for 90 min, following the protocol set forward by Garrod and colleagues (Garrod, 2013; Garrod et al., 2005). Hyperadhesive desmosomes persist following this treatment, whereas calcium-dependent desmosomes do not. To observe desmosomes and overall cellular morphology, cells were fixed and immunostained for desmoglein 3 (Dsg3), a desmosomal cadherin, and DP (Fig. 1 A). Dsg3 and DP colocalized in puncta at cell borders when cells were maintained in normal-Ca\(^{2+}\) medium (~1.8 mM), indicating the presence of desmosomes at cell–cell junctions. After a 90-min incubation in low-Ca\(^{2+}\) medium, mock-treated cells had a rounded morphology, lack of cell–cell contacts, and loss of desmosomes. In contrast, GÖ6976-treated cells retained their morphology, protein localization, and cell–cell contacts, suggesting they had acquired hyperadhesion. To test the adhesive function of these cells, intact epithelial sheets were lifted off the culture dish and subjected to mechanical stress to induce fragmentation in a dispase cell adhesion assay (Fig. 1 B). We observed that cells treated with GÖ6976 resisted fragmentation compared with mock-treated cells in both high- and low-Ca\(^{2+}\) medium. Together, these data demonstrate that GÖ6976 treatment leads to hyperadhesion in our experimental system.

Our initial goal was to determine whether there were architectural features of desmosomes that promote hyperadhesion. The leading model is that the arrangement of cadherin extracellular domains in an ordered crystalline array is necessary for hyperadhesion (Garrod, 2013; Garrod et al., 2005). To test this, we used excitation-resolved fluorescence polarization microscopy to measure cadherin order (Bartle et al., 2017; DeMay et al., 2011; Kress et al., 2011). In this approach, the protein of interest is rigidly tagged with a fluorophore, such that the fluorophore orientation reflects the protein orientation. When illuminated with linearly polarized light, fluorophores with dipoles oriented parallel to the excitation polarization are excited, whereas those perpendicular are not (Fig. 1 C). If proteins within a complex are ordered, fluorescence intensity is modulated by changing the orientation of the excitation polarization (Fig. 1 D, top). If the proteins are not ordered, there is no modulation of intensity (Fig. 1 D, bottom). The order of many copies of the tagged protein within a diffraction limited (~250-nm) spot can therefore be quantified from the fluorescence intensity recorded at four unique excitation polarizations.

To measure the order of desmosomal cadherins, we replaced the Dsg3 extracellular anchor domain (EA) with GFP (Dsg3-ΔEA-GFP), as previously described (Bartle et al., 2017). This construct reports the order of the most membrane-proximal extracellular domain of Dsg3. To test whether expression of Dsg3-ΔEA-GFP disrupted desmosome function, cells were subjected to a dispase cell adhesion assay. Cells expressing Dsg3-ΔEA-GFP had increased adhesion following treatment with GÖ6976 (Fig. S1 A). The degree of fragmentation was not significantly different from that in control cells expressing either Dsg3 with GFP attached to the C terminus with an unstructured linker or cytosolic GFP,
Figure 1. Hyperadhesion does not protect against loss of Dsg3 order in low-Ca²⁺ media. (A) Representative images of HaCaT cells treated with either vehicle (DMSO; mock) or Go6976 and maintained in normal-Ca²⁺ medium (left) or switched into low-Ca²⁺ medium (right) for 90 min and labeled with antibodies for Dsg3 and DP. Scale bar = 20 µm. (B) Quantification of adhesive strength by dispase fragmentation assay in HaCaT cells with the same treatment conditions as in A (n = 3; representative of four independent experiments; mean ± SD). (C) Schematic of the fluorescence polarization microscope. Samples are sequentially illuminated with four different angles of polarized excitation, and the total emission is collected. (D) Ordered cadherins (top) are differentially excited by distinct excitation polarizations, resulting in modulated intensity across the four images. Disordered cadherins (bottom) are equally excited regardless of excitation polarization, resulting in equal intensity. Intensity was plotted as a function of excitation polarization, and the amplitude was used to calculate pixel-by-pixel order factor (OF). (E and F) HaCaT cells were transfected with Dsg3-ΔEA-GFP, mock (E) or Go6976 (F) treated and switched from normal to low-Ca²⁺ medium. Dsg3 intensity and OF images of representative cells before and 30 min after the switch to low calcium with region of interest (ROI) indicated and intensity and Dsg3-ΔEA-GFP OR in the ROI over a 30-min time course after the switch to low-Ca²⁺ medium. Scale bar = 5 µm; ROI scale bar = 1 µm. (G–I) Population analysis of cells mock treated in normal-Ca²⁺ (n = 6) or low-Ca²⁺ (n = 11) and Go6976 treated in normal-Ca²⁺ (n = 4) or low Ca²⁺ (n = 15). (G) Normalized integrated intensity at t = 30 min (mean ± SD). (H) OF plotted as a function of time (mean ± SD). (I) OF at t = 30 min (mean ± SD) compared by one-way ANOVA with multiple comparisons. Data were acquired from three independent experiments (ns, not significant; P > 0.05; *, P ≤ 0.05; **, P ≤ 0.01; ***, P ≤ 0.001; ****, P ≤ 0.0001).
indicating that Dsg3-ΔEA-GFP does not act as a dominant negative in desmosome adhesion.

Dsg3-ΔEA-GFP was transfected into HaCaT cells and imaged with fluorescence polarization microscopy. Cells were imaged in normal-Ca2+-medium before and after Gö6976 treatment, which was found to have no effect on Dsg3-ΔEA-GFP order factor (Fig. S1, B and C). In mock-treated cells with calcium-dependent desmosomes, Dsg3-ΔEA-GFP was ordered in the presence of Ca2+, and order was rapidly lost when cells were switched into low-Ca2+-medium (Fig. 1 E; Bartle et al., 2017). The switch to low-Ca2+-medium also led to desmosome disassembly, as shown by a reduction of fluorescence intensity, rearrangement of the cell junctions, and changes in cell morphology (Fig. 1 E).

We next wanted to know whether cadherins remain ordered in hyperadhesive desmosomes without Ca2+. Surprisingly, when Gö6976-treated cells were switched to low-Ca2+-medium, order factor rapidly decreased (Fig. 1 F). Desmosomes in Gö6976-treated cells had a consistent morphology before and after the switch to low Ca2+, and we were able to track individual desmosomes throughout the time course. This was in contrast to the changing morphology and loss of desmosomes observed in the mock-treated cells. To examine whether proteins were lost from desmosomes over the experiment, we quantified the integrated fluorescence intensity of desmosomal puncta. The normalized fluorescence intensity at t = 30 min did not change significantly for Gö6976-treated cells between high and low Ca2+, whereas in mock-treated cells, the intensity was significantly reduced in low Ca2+ (Figs. 1 G and SI, D and E). This indicates that the amount of Dsg3-ΔEA-GFP in desmosomes was decreased in mock- but not Gö6976-treated cells 30 min after the switch to low-Ca2+-medium. The retention of protein and maintenance of morphology for Gö6976-treated cells in low-Ca2+-medium suggests that these represent hyperadhesive desmosomes.

The dynamics of the decreasing order factors were similar between mock- and Gö6976-treated cells in low-Ca2+-medium (Fig. 1 H). In contrast, order factor remained stable for cells maintained in normal-Ca2+-medium in both treatments. The mean order factor 30 min after the change to low-Ca2+-medium was not significantly different between mock- and Gö6976-treated cells (Fig. 1 I). To confirm that this reduced order factor was persistent, Gö6976-treated cells were fixed either 30 or 90 min after a switch to low-Ca2+-medium. When imaged with fluorescence polarization, the Dsg3-ΔEA-GFP order factor was not significantly different at 30 or 90 min in low-Ca2+-medium (Fig. SI F). These experiments reveal that cadherin order, specifically the order of the most membrane-proximal extracellular domain of Dsg3, is not determined by adhesive state, but by Ca2+. Intriguingly, our data suggest that cadherins in hyperadhesive desmosomes can be simultaneously disordered and adhesive. Therefore, a quasi-crystalline arrangement of cadherins cannot be the driving mechanism of hyperadhesion.

Given the unexpected result that cadherin order was dependent on Ca2+, not adhesive state, we wanted to further investigate this relationship. Because of the importance of Ca2+ in cadherin tertiary structure, we hypothesized that order is indicative of Ca2+ binding, and that if Ca2+ is reintroduced, order will be restored. To test this, we used fluorescence polarization microscopy to measure the dynamics of Dsg3-ΔEA-GFP order in Ca2+-dependent desmosomes after a “low-Ca2+” pulse. We chose to use calcium-dependent desmosomes for this assay because the cadherins are disordered after 10 min in low Ca2+, but desmosomes are still present. This allowed us to ask whether reintroducing Ca2+ will restore cadherin order or halt desmosome disassembly. For the low-Ca2+ pulse, cells in normal-Ca2+-medium were switched into low-Ca2+-medium for 10 min and then switched back to normal-Ca2+-medium. Fluorescence polarization imaging revealed that Dsg3-ΔEA-GFP order was lost following the switch to low-Ca2+-medium as anticipated, yet switching the cells back to normal-Ca2+-medium did not rescue order (Fig. 2 A). We tracked individual junctions and found that the majority were present for the duration of the time course and did not represent newly formed junctions. The decreased order factor persisted for 90 min after the low-Ca2+ pulse (Fig. 2 B). Loss of order was further quantified by comparing the order factor from the cells in normal-Ca2+-medium before the low-Ca2+ pulse (0.37 ± 0.07; mean ± SD), after the 10 min low-Ca2+ pulse (0.23 ± 0.04), and 90 min after the pulse in normal-Ca2+-medium (0.18 ± 0.03; Fig. 2 C). We found that there was an initial loss of Dsg3-ΔEA-GFP fluorescence intensity from puncta after the switch to low-Ca2+-medium. However, no further loss of intensity occurred over the 90-min experiment, indicating that the amount of Dsg3-ΔEA-GFP in the puncta was consistent once the cells were returned to normal-Ca2+-medium (Fig. 2 D). This revealed that loss of cadherin order is not reversible by reintroducing Ca2+. Once the membrane-proximal domain of Dsg3 was disordered following removal of Ca2+, it did not return to its original ordered arrangement. These results show that desmosomes can be manipulated to have disordered cadherins in both calcium-dependent and hyperadhesive states.

Desmosome architecture is maintained in Gö6976-induced hyperadhesion

Given that cadherin order did not correlate with adhesive state, we hypothesized that changes in the nanoscale architecture of the desmosomal plaque may provide additional insight toward the mechanism of hyperadhesion. We used superresolution direct stochastic optical reconstruction microscopy (dSTORM; Heilemann et al., 2008) to quantify desmosome protein architecture in different adhesive states (Stahley et al., 2016). We conducted dSTORM on HaCaT cells with calcium-dependent (mock-treated) or hyperadhesive (Gö6976-treated) desmosomes. To test whether extracellular Ca2+ impacted plaque architecture, cells were either maintained in normal-Ca2+-medium or switched to low-Ca2+ medium. Mock-treated samples were switched to low-Ca2+ medium for only 10 min, which preserved junctions for imaging, whereas Gö6976 treated cells were incubated in low-Ca2+-medium for 90 min. Cells were fixed and labeled with antibodies for either the C-terminal keratin binding domain of DP or plakoglobin (PG), an armadillo protein that binds to the cadherin tails, allowing us to assess architecture throughout the different regions of the plaque. After dSTORM imaging, protein localization and architecture were quantified by measuring plaque-to-plaque distance (Fig. 3, A and B). The quantification of plaque-to-plaque distance for

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both DP C-terminus and PG showed no significant difference across treatment conditions (Fig. 3 C). These results show that the localization of PG in the membrane proximal outer dense plaque and the localization of the C-terminus of DP in the inner dense plaque are not altered within the resolution of dSTORM microscopy.

To further investigate desmosome ultrastructure, transmission EM was conducted on cells following the same treatment paradigms (Fig. 3 D). There were no obvious alterations of the plaques by treatment with Gö6976 or incubation in low Ca²⁺. Quantification of the width of the intermembrane space showed no significant difference between groups (Fig. 3 E). We conclude that plaque architecture and desmosome ultrastructure are maintained under the conditions tested and thus cannot explain the mechanism of hyperadhesion.

**Desmosomal cadherins are stabilized by hyperadhesion**

To further investigate architectural changes that could drive the switch between adhesive states, we next explored cadherin exchange in and out of desmosomes using FRAP. First, we measured cadherin exchange in cells with calcium-dependent desmosomes transfected with Dsg3-ΔEA-GFP in normal-Ca²⁺ medium. A region along the cell border containing multiple desmosomes was photobleached, and fluorescence recovery was measured over 20 min (Fig. 4 A). We found that fluorescence recovery localized to desmosomes, as highlighted by tracking individual puncta within the bleach region over the time course (Fig. 4, B and C). The mobile fraction, calculated from the fluorescence recovery curve, represents the amount of Dsg3-ΔEA-GFP exchange in and out of desmosomes (Fig. 4 D). Next, FRAP was conducted on cells treated with Gö6976 and incubated in low-Ca²⁺ medium for 90 min to select for hyperadhesive desmosomes (Fig. 4 E). There was minimal recovery of Dsg3-ΔEA-GFP fluorescence localized to desmosomal puncta within the bleach region (Fig. 4, F–H). To test whether this reduced Dsg3 exchange in hyperadhesion was conserved for different cadherin isotypes, we performed FRAP on cells transfected with Dsg2-GFP under the same treatment conditions (Fig. S2). The mean normalized intensity was fitted to a one-phase association, revealing reduced exchange of Dsg2 and Dsg3 in hyperadhesive desmosomes (Fig. 4 I).

To quantify this difference the recovery curve from each cell was fitted, and the resulting mobile fractions were averaged to determine the mean mobile fraction. The mean mobile fraction of Dsg3-ΔEA-GFP in cells with calcium-dependent desmosomes (22 ± 11%) was significantly higher than in hyperadhesion (10 ± 2%; Fig. 4 J). Similarly the mean mobile fraction of Dsg2-mCherry in cells with calcium-dependent desmosomes (27 ± 14%) was significantly higher than in cells with hyperadhesive desmosomes (9 ± 3%; Fig. 4 J). There was no significant difference in the mobile fraction between cadherin isotypes within the same treatment group.

To confirm that removal of Ca²⁺ from the medium was not responsible for the reduced protein exchange, we performed FRAP of Dsg3-ΔEA-GFP in cells treated with Gö6976 and maintained in normal-Ca²⁺ medium. The mean mobile fraction (9 ± 8%) was not significantly different from hyperadhesive desmosomes in low-Ca²⁺ medium (Fig. 4 J). This significant decrease in desmosomal cadherin mobility in hyperadhesive cells represents a reduced exchange of proteins in and out of desmosomes.
To test whether the reduced mobile fraction of desmogleins in response to Go6976 treatment is specific to desmosomal cadherins, we measured the mobility of the adherens junction protein E-cadherin (E-cad). FRAP experiments were conducted on cells transfected with E-cad-GFP. The mobile fraction did not depend on treatment (57 ± 15% mock and 51 ± 10% Go6976) and was significantly higher than that of the desmosomal cadherins (Fig. 4, I and J).

Plaque proteins are stabilized in hyperadhesion

We next asked whether the mobility of desmosomal plaque proteins was also reduced in hyperadhesive desmosomes. DP is an intermediate filament binding protein that anchors the desmosomal plaque to the cytoskeleton. First, FRAP experiments were conducted on cells transfected with E-cad-GFP. The mobile fraction did not depend on treatment (57 ± 15% mock and 51 ± 10% Go6976) and was significantly higher than that of the desmosomal cadherins (Fig. 4, I and J).

Figure 3. Desmosome architecture is maintained in Go6976-induced hyperadhesion. (A and B) HaCaT cells were mock or Go6976 treated and incubated in normal or low-Ca²⁺ medium. Cells were stained with antibodies for DP (A) or PG (B) and imaged by dSTORM. Representative cell borders are shown for each condition with single desmosome regions of interest (ROIs) indicated by the red squares. Scale bar = 500 nm; ROI scale bar = 100 nm. Linescans of fluorescence intensity perpendicular to the desmosome axis were used to quantify plaque-to-plaque distance; examples are shown for the ROIs. (C) Quantification of plaque-to-plaque distance shows no significant difference across treatment groups for DP or PG (DP left to right, n = 58, 40, 44, 41; PG left to right n = 41, 30, 48, 31, each from three independent experiments). (D) Electron micrographs of desmosomes under the same treatment conditions. Scale bar = 100 nm. (E) Quantification of intermembrane distance shows no difference across treatment groups (n = 15 desmosomes per group). Bars represent mean ± SD (n.s., not significant, P > 0.05; one-way ANOVA followed by Tukey’s multiple comparison test).

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These experiments revealed that the mobility of both DP and PG was reduced in hyperadhesive desmosomes (Fig. 5 Q). The mobile fractions of DP and PG in cells with hyperadhesive desmosomes (DP, 5 ± 4%; PG, 8 ± 6%) was significantly lower than in calcium-dependent desmosomes (DP, 28 ± 17%; PG, 34 ± 17%).
There was no significant difference in mobile fraction between PG and DP within each adhesive state (Fig. 5 R). These results demonstrate that there is a loss of plaque protein mobility in hyperadhesive desmosomes.

**Trans binding contributes to cadherin order and reduced exchange in hyperadhesion**

Because cadherin trans binding mediates desmosome adhesion, we next tested whether trans interactions influence cadherin order or exchange. Cadherin trans binding occurs through a strand-swap mechanism involving insertion of a tryptophan residue on the EC1 domain (W2) into a hydrophobic binding pocket of a cadherin on the opposing cell (Häußinger et al., 2004; Overduin et al., 1995; Shapiro et al., 1995). Site-directed mutagenesis was used to mutate this key tryptophan residue to alanine (W2A) in Dsg3-ΔEA-GFP (Dsg3-W2A), a mutation that has been demonstrated to abrogate trans binding (Harrison et al., 2016; Lowndes et al., 2014). Dsg3-W2A was transfected into HaCaT cells and observed to localize to puncta at the plasma membrane, indicating incorporation into desmosomes. The order factor of Dsg3-W2A (0.22 ± 0.04) was significantly lower than wild-type Dsg3 (0.37 ± 0.08) in calcium-dependent desmosomes maintained in normal-Ca²⁺ medium (Fig. 6, A and B). How Dsg3-W2A is arranged in the desmosomes was determined by FRAP experiments (Fig. 4). Desmosomal cadherin exchange is reduced in hyperadhesive desmosomes. (A–D) FRAP experiments were conducted on HaCaT cells transfected with Dsg3-ΔEA-GFP and mock treated in normal-Ca²⁺ medium. (A) Representative cell (inverted intensity) with bleach (red) and reference (orange) regions of interest (ROIs). Bleach ROI shown with individual desmosomal puncta indicated. Scale bar = 5 µm; ROI scale bar = 1 µm. (B) Bleach ROI over time with dashed lines underscoring individual puncta, highlighting recovery. Scale bar = 1 µm. (C) Linescans through the bleach region indicate relative intensities before bleaching (black), immediately after bleaching (gray), and at 20 min after bleaching (red). (D) FRAP recovery curves of the bleach (red) and reference (orange) ROIs. The bleach ROI intensity was fitted to a one-phase association to determine mobile fraction (MF). (E–H) FRAP experiments were conducted on HaCaT cells transfected with Dsg3-ΔEA-GFP, G66976 treated, and switched to low-Ca²⁺ medium for 90 min. (E) Representative cell (inverted intensity) with bleach (red) and reference (orange) ROIs. Bleach ROI shown with individual desmosomal puncta indicated. Scale bar = 5 µm; ROI scale bar = 1 µm. (F) Bleach region over time with dashed lines underscoring individual puncta. Scale bar = 1 µm. (G) Linescans through the bleach region indicate relative intensities before bleaching (black), immediately after bleaching (gray), and at 20 min after bleaching (red). (H) FRAP recovery curves of the bleach (red) and reference (orange) ROIs. The bleach ROI intensity was fitted to a one-phase exponential association to determine the MF. (I) FRAP recovery curves (mean ± SD) for cells transfected with Dsg3-ΔEA-GFP, Dsg2-GFP, or E-Cad-GFP and mock treated in normal Ca²⁺ (n = 15, 11, 5) or G66976 treated in low Ca²⁺ (n = 11, 10, 4). (J) Mean mobile fraction ± SD for treatments in I and Dsg3-ΔEA-GFP G66976 treated in normal Ca²⁺ (n = 11). All data were acquired from three independent experiments (ns, not significant, P > 0.05; **, P ≤ 0.01; ***, P ≤ 0.001; ANOVA).
intermembrane space is not known, but the decreased order factor could be attributed to a range of Dsg3-W2A orientations, perhaps restricted by the ordering of wild-type cadherins or a mixed population of disordered and ordered Dsg3-W2A. We conclude that Dsg3-W2A is recruited to desmosomes but does not adopt the same ordered conformation as wild type, indicating that trans binding contributes to establishing cadherin order.

Figure 5. Plaque protein exchange is reduced in hyperadhesive desmosomes. (A–D) FRAP experiments were conducted on HaCaT cells transfected with DP-mCherry and mock treated in normal-Ca\(^{2+}\) medium. (A) Representative cell (inverted intensity) with bleach region of interest (ROI; red). (B) Individual puncta over the time course underscored by dashed lines. (C) Linescans through the bleach region indicate relative intensities before bleaching (black), immediately after bleaching (gray), and at 20 min after bleaching (red). (D) Fluorescence intensity over time and mobile fraction of the bleach ROI (MF). (E–H) Representative cell from FRAP experiments conducted on HaCaT cells transfected with DP-mCherry, treated with Go6976, and switched into low-Ca\(^{2+}\) medium for 90 min. (I–L) Representative cell from FRAP experiments conducted on HaCaT cells transfected with PG-mEmerald and mock treated in normal-Ca\(^{2+}\) medium. (M–P) Representative cell from FRAP experiments conducted on HaCaT cells transfected with PG-mEmerald, treated with Go6976, and switched into low-Ca\(^{2+}\) medium for 90 min. (A, E, I, M) Scale bar = 5 µm. (B, F, J, N) Scale bar = 1 µm. (Q and R) FRAP recovery and fit to one-phase association curves (mean ± SD; Q) and mobile fraction (mean ± SD; R) of PG-mEmerald in mock (n = 9) and Go6976 (n = 8) treated cells and DP-mCherry in mock (n = 14) and Go6976 (n = 7) treated cells. All data from three independent experiments. (ns, not significant, P > 0.05; *, P ≤ 0.05; **, P ≤ 0.01; ANOVA).
To investigate whether cadherin mobility depends on trans binding, we conducted FRAP experiments on cells cotransfected with Dsg3-W2A and DP-mCherry. In cells with calcium-dependent desmosomes, Dsg3-W2A and DP-mCherry fluorescence recovered and was localized to desmosomal puncta (Fig. 6, C and D). In Go6976-treated cells incubated in low-Ca\textsuperscript{2+} medium for 90 min, there was no recovery of DP-mCherry, as seen in previous experiments. However, Dsg3-W2A fluorescence recovered as much as in the calcium-dependent desmosomes (Fig. 6, E and F). Quantification showed that hyperadhesion did not significantly change the Dsg3-W2A mobile fraction (calcium-dependent, 24 ± 11%; hyperadhesive, 19 ± 15%), while in the same cells, the mean mobile fraction of DP-mCherry was significantly reduced in hyperadhesive desmosomes (10 ± 7%) compared with calcium-dependent desmosomes (26 ± 14%; Fig. 6, G and H). These data show that expression of Dsg3-W2A does not have a dominant negative effect on DP exchange. These results indicate that trans binding is an essential component of reduced cadherin mobility in hyperadhesion (Fig. 6 I).
DP phosphorylation is a molecular switch controlling protein exchange

Gö6976 treatment is hypothesized to block DP phosphorylation at S2849, a PKCα consensus site. We confirmed this by immunoblot, which revealed that the percentage of total DP phosphorylated at S2849 decreased by 22.4% in HaCaT cells after treatment with Gö6976 (Fig. 7 A). Lack of phosphorylation at this serine has been shown to increase the affinity of DP for keratin (Godsel et al., 2005). Blocking this phosphorylation with a point mutant (S2849G) has been shown to increase the affinity of DP for keratin and induce hyperadhesiveness (Albrecht et al., 2015; Hobbs and Green, 2012; Meng et al., 1997). We set out to use the DP mutant S2849G to test whether the order and mobility results obtained when Gö6976 was used to induce hyperadhesion could be recapitulated by blocking phosphorylation of DP at S2849.

We first wanted to see whether mutant DP-S2849G-mCherry impacted the order of the Dsg3-ΔEA-GFP extracellular domain. Cotransfection with DP-mCherry caused no significant difference in Dsg3-ΔEA-GFP order factor while maintained in normal-Ca2+ medium (wild-type DP, 0.37 ± 0.12; mutant DP, 0.39 ± 0.11; Fig. S3, A–C). Cells transfected with DP-S2849G-mCherry had increased adhesion compared with wild-type DP-mCherry (Fig. S3 D). After a switch into low-Ca2+ medium, cell morphology was unchanged, and puncta were maintained at cell borders over the 30-min imaging window, indicating the presence of hyperadhesive desmosomes. Dsg3-ΔEA-GFP order decreased rapidly following the switch to low-Ca2+ medium (Fig. 7, B and C). The final order factor after 30 min in low-Ca2+ medium (0.19 ± 0.07) was not significantly different from that measured in cells treated with Gö6976 (0.18 ± 0.05; see Fig. 1; Table S1).

Next, we tested whether expression of DP-S2849G was sufficient to block protein exchange. FRAP experiments conducted on cells cotransfected with Dsg3-ΔEA-GFP and DP-S2849G-mCherry and maintained in normal-Ca2+ medium showed minimal exchange of either protein (Fig. 7, D–F). The mobility of DP-S2849G was significantly lower than that established earlier for wild-type DP (Student’s t test; Table S2). We next wanted to investigate whether cadherin trans binding was required for this reduced exchange. The trans-binding mutant Dsg3-W2A retained mobility when cotransfected with DP-S2849G-mCherry (Fig. 7, G–I). The mean mobile fractions of DP-S2849G-mCherry (7 ± 3%) and Dsg3-ΔEA-GFP (6 ± 3%) were not significantly different from each other in the same cells (Fig. 7, J and K). In contrast, the mean mobile fraction of Dsg3-W2A (30 ± 18%) was significantly higher than that of the DP-S2849G-mCherry (10 ± 7%) it was cotransfected with. The mobility of DP-S2849G-mCherry was similar to that of wild-type DP in Gö6976-treated cells with hyperadhesive desmosomes (Table S2).

These findings demonstrate that blocking DP phosphorylation at S2849 is sufficient to prevent protein exchange in desmosomes. However, this reduced exchange requires cadherin trans binding. Together these results show that DP phosphorylation at S2849 acts as a key molecular switch in regulating desmosome adhesive state by controlling protein dynamics.

Discussion

Modulating the adhesive state allows desmosomes to adjust their strength to facilitate modulation of tissue integrity and plasticity in processes such as embryonic development and wound healing. A key hypothesis is that the arrangement of cadherins into an ordered array at the adhesive interface is a defining property of hyperadhesion (Al-Amoudi et al., 2007; Garrod, 2013; Rayns et al., 1969). Our data show that order of the most membrane-proximal extracellular domain of Dsg3 is lost upon removal of Ca2+ in hyperadhesive desmosomes. Order was also shown to be independent of Ca2+ binding, as cadherin order was not restored by reintroduction of Ca2+ to calcium-dependent desmosomes following a low-Ca2+ pulse. Our data uncouple cadherin order and desmosome adhesion, suggesting that cadherin order is not a defining structural feature of hyperadhesion. It is interesting to consider how hyperadhesive desmosomes maintain function in the absence of Ca2+. It is possible that interdomain flexibility could allow disorder cadherins to engage in trans binding in a hyperadhesive state (Tarq et al., 2015). An important consideration is that this approach measures the order of the tagged protein domain, in this case the most membrane-proximal extracellular domain of Dsg3. This leaves open the possibility that cadherins could retain order in other domains, closer to the adhesive interface. However, if a crystalline ectodomain were driving hyperadhesion, we would expect no loss of order from any cadherin domain when Ca2+ is removed from hyperadhesive desmosomes. Given this uncoupling of order and adhesion, why and how cadherin order is conferred to desmosomes is an intriguing question for future investigation.

Fluorescence polarization microscopy is a powerful approach with the unique ability to probe the order of proteins within subcellular structures (DeMay et al., 2011; Rocheleau et al., 2003; Swaminathan et al., 2017; Valades Cruz et al., 2016). This unique insight has allowed us to investigate critical questions about cadherin organization in this work. An important limitation of our approach is that we are unable to determine the type of order or identify specific structural changes. In addition, dynamics, disorder, and some unfavorable GFP orientations can all contribute to lowering the order factor (Bartle et al., 2017). Ultimately, order factor offers a readout for structural organization within the desmosome, but it does not currently indicate the ultrastructure conveying order.

In light of our discovery that cadherin order is lost with the removal of Ca2+, it is interesting to consider the role of Ca2+ in hyperadhesion. Hyperadhesion has been defined by resistance to incubation in low-Ca2+ medium for 90 min (Garrod and Kimura, 2008; Kimura et al., 2007). However, we present here the destructive effect removal of Ca2+ has on cadherin order, regardless of adhesive state. After treatment with Gö6976, Dsg3 mobility was significantly reduced in cells that were subjected to the 90-min incubation in low-Ca2+ medium or maintained in normal-Ca2+ medium. Interestingly, when Ca2+ was kept in the medium, we saw a larger distribution of mobile fractions, possibly because there was a mix of calcium-dependent and hyperadhesive desmosomes resulting from an incomplete effect of Gö6976 on conferring that state. The dynamic, single-cell sensitivity of FRAP experiments could prove helpful in
Figure 7. DP S2849G conferred reduced protein exchange. (A) Representative immunoblot and densitometry analysis of total DP and phospho-DP in HaCaT cells mock or Go6976 treated. For densitometry, the phospho-DP/total DP ratio was normalized to mock treated (****, P < 0.0001 by unpaired t test; data are mean ± SD of three independent experiments). (B and C) HaCaT cells were cotransfected with Dsg3-ΔEA-GFP and DP-S2849G-mCherry, switched from normal to low-Ca²⁺ medium, and imaged with fluorescence polarization microscopy. (B) Representative cell fluorescence and order factor images over time (scale bar = 5 µm; region of interest [ROI] scale bar = 1 µm). (C) Mean Dsg3-ΔEA-GFP order factor over time (mean ± SD; n = 8 cells). (D) Representative cell expressing DP-S2849G-mCherry (magenta) and Dsg3-ΔEA-GFP (cyan) with bleach ROI indicated. Scale bar = 5 µm. (E) ROI (inverted intensity) with dotted lines underscoring individual puncta of DP-S2849G-mCherry and Dsg3-ΔEA-GFP over the time course (scale bar = 1 µm). (F) Bleach ROI of DP-S2849G-mCherry and Dsg3-ΔEA-GFP intensity plotted as a function of time and fitted to a one-phase association. (G) Representative cell expressing DP-S2849G-mCherry (magenta) and Dsg3-W2A-ΔEA-GFP (cyan) with bleach ROI indicated. Scale bar = 5 µm. (H) ROI (inverted intensity) with dotted lines underscoring individual puncta of DP-S2849G-mCherry and Dsg3-W2A-ΔEA-GFP over the time course (scale bar = 1 µm). (I) Bleach ROI of DP-S2849G-mCherry and Dsg3-W2A-ΔEA-GFP intensity plotted as a function of time and fitted to a one-phase association. (J and K) Normalized intensity over time fitted to a one-phase association (mean ± SD; J) and mobile fraction (mean ± SD; K) for cotransfections with quantified protein in bold and coexpressed protein in parentheses. DP-S2849G-mCherry (Dsg3-ΔEA-GFP; n = 8) and Dsg3-ΔEA-GFP (DP-S2849G-mCherry, n = 8) and DP-S2849G-mCherry (Dsg3-W2A-ΔEA-GFP, n = 6) and Dsg3-W2A-ΔEA-GFP (DP-S2849G-mCherry, n = 8). All data from three independent experiments (ns, not significant, P > 0.05; *, P ≤ 0.05; **, P ≤ 0.01; ANOVA).
future work investigating hyperadhesion under native Ca\(^{2+}\) conditions.

In this work, we show that a defining physical characteristic correlating with adhesive state is protein exchange, not cadherin order (Fig. 8 A). Cadherins and plaque proteins in desmosomes engaged in calcium-dependent adhesion exhibited significant mobility. This indicates that there is exchange between two pools of proteins: desmosomal and non-desmosomal. We propose that sequestration, degradation, or endocytosis of cadherins not associated with desmosomes could “tip the balance” of the ratio between these pools, with the potential to promote disassembly and impact desmosome retention in cells. Assuming that cadherin mobility out of the complex remains unchanged, a reduction in the ability to recruit or retain new cadherins would ultimately result in desmosome disassembly. This state is represented by our low-Ca\(^{2+}\) treatment, in which the ability of cadherins to engage in trans binding is reduced by removal of Ca\(^{2+}\). In this case, the diffusion of cadherins out of desmosomes is unchanged, but the ability to incorporate new protein is reduced. Over time, cadherins are extracted from desmosomes and not replaced, leading to dissolution of the structures. A similar state could be achieved by depleting the free pool of proteins, reducing the availability of proteins to incorporate into desmosomes, while not altering the rate of loss. In this way, protein exchange provides a mechanism for desmosomes to rapidly adapt to changing conditions by either maintaining or disassembling cell adhesion structures.

Figure 8. Protein exchange controls adhesive state. Models illustrating how reduced protein exchange confers hyperadhesion. (A) Model illustrating dynamics of DP phosphorylation, protein exchange, and cadherin order in calcium-dependent adhesion (left) and hyperadhesion (right) with (top) and without (bottom) Ca\(^{2+}\). In calcium-dependent desmosomes with Ca\(^{2+}\), cadherins are ordered, and desmosomal proteins can diffuse in and out of the complex. In the absence of Ca\(^{2+}\), cadherin order is lost, proteins diffuse out but do not reenter, and desmosomes disassemble. In hyperadhesive desmosomes with Ca\(^{2+}\), cadherins are ordered, but desmosomal proteins are unable to diffuse in or out of the complex. In the absence of Ca\(^{2+}\), cadherin order is lost, but adhesion is maintained because proteins are unable to diffuse out of the complex. (B) We suggest that protein mobility is blocked by G\(\theta\)6976 or DPS2849G through modulation of DP–keratin interactions via an inside-out mechanism. PM, plasma membrane.
In contrast, we found that there was limited exchange of proteins in hyperadhesive desmosomes. Because there is limited exchange, there is less interaction between the two pools of desmosomal proteins. Therefore, in hyperadhesion, alterations to the non-desmosomal pool of proteins do not have the opportunity to impact the stability of cell adhesion. When treated with low-Ca\(^{2+}\) medium, cadherins became disordered, but desmosomes remained, and adhesion was not lost. We postulate that this is because the proteins are “locked” within the complex by the combination of trans interactions and plaque associations. This is supported by our data showing that when trans binding is abrogated, cadherins are no longer locked in to hyperadhesive desmosomes. This suggests that in the absence of Ca\(^{2+}\), the balance between the probability of cadherins engaging in trans binding and diffusing out of the complex favors retention within desmosomes and maintenance of functional integrity.

Evidence suggests that PKCa endogenously regulates desmosome adhesive state (Thomason et al., 2012), facilitated by its localization to keratin and trafficking with DP (Bass-Zubek et al., 2008; Kröger et al., 2013). It has been previously established that the phospho-null mutant DP S2849G has an increased affinity for keratin, resulting in delayed DP trafficking and increased keratin integration at desmosomes (Godsel et al., 2005). This behavior, and the evidence presented here and by others showing that PKCa is responsible for phosphorylating DP at S2849 (Wallis et al., 2000), suggests that both PKCa inhibition and DP S2849G act through similar means. We posit that increased integration of DP with keratin due to reduced DP phosphorylation is responsible for the loss of protein mobility, but do not eliminate the possibility of other downstream post-translational modifications of DP or other desmosomal proteins acting in the same pathway. We also leave open a role for cellular mechanics, which could be similarly altered by the change in DP and keratin association. Increasing the affinity of DP for keratin resulted in reduced protein exchange transmitted throughout the complex. This inside-out pathway results in a desmosome structure in which proteins are essentially locked in. In this way, DP phosphorylation can act as a fast and powerful molecular switch for regulating desmosome adhesion by controlling protein exchange (Fig. 8 B).

Our findings shed new light on the cadherin diffusion rates found in the literature. The mobility of Dsg2 and Dsg3 reported from cell culture models is generally consistent with the values we report here in calcium-dependent desmosomes (Gloshankova et al., 2003; Lowndes et al., 2014; Vielmuth et al., 2018; Windoffer et al., 2002), while the lower mobile fraction reported in mouse epidermis is consistent with what we measured in hyperadhesion (Foote et al., 2013). The reduced mobility of desmosomal cadherins is in direct contrast to adherens junction proteins, which have been found to be mobile in mature epidermis (Cao and Schnittler, 2019; Foote et al., 2013). These data agree with our finding that the mobile fraction of E-cad was higher than that of the desmosomal cadherins and not altered by hyperadhesion. Interestingly, the mobile fraction of E-cad in vivo has been shown to correlate with junction integrity and cell invasiveness (Canel et al., 2010; Erami et al., 2016). Vascular endothelial cadherin has also been shown to have high mobility, which could be restricted by protein–protein interactions (Nanes et al., 2012). Together, these data point to a global mechanism in which protein mobility in and out of cell junctions plays a central role in defining stability versus plasticity.

This work provides a novel framework for understanding the regulation of cell–cell adhesion and raises new and exciting questions. Our results demonstrate that hyperadhesion is established by a loss of protein dynamics and plasticity. This mechanism allows cells to rapidly and precisely control cell cohesion and tissue integrity by modulating protein exchange and desmosome stability through posttranslational modifications. Epithelial tissues can thus rapidly respond to both intrinsic and extrinsic cues to control processes such as cell migration and proliferation that require tuning of adhesive function. This regulation is critical for maintaining tissue homeostasis, with aberrant adhesive function contributing to cancer invasiveness, skin fragility, and cardiac arrhythmias. Assessing adhesive function by measuring cell junction protein dynamics is a potential avenue for investigation of adhesion in situ, which would provide a platform for integrating adhesion with the broader context of signaling events, morphological changes, and other regulatory mechanisms. Future work focused on measuring and modulating junctional protein mobility could shed light on the function and dysfunction of junctions, possibly resulting in new therapeutic avenues.

**Materials and methods**

**Cell culture**

The human immortalized keratinocyte cell line HaCaT (Addex Bio) was cultured in optimized DMEM (Addex Bio) supplemented with 10% FBS (Corning) and 100 U/ml penicillin/streptomycin (Corning) and maintained at 37°C and 5% CO\(_2\). For imaging, cells were seeded into either eight-well #1.5 coverslip bottom dishes (Ibidi) or 25-mm #1.5 coverslips (Electron Microscopy Sciences).

**Cloning and constructs**

Dsg3-ΔEA-GFP, Dsg3-link-GFP, and DP-mCherry were previously published (Bartle et al., 2017). PG-mEmerald was a gift from Michael Davidson (Florida State University, Tallahassee, FL; Addgene plasmid 54133), and E-cad-GFP was a gift from Jennifer Stow (University of Queensland, Brisbane, Australia; Addgene plasmid 28009; Miranda et al., 2001). Site-directed mutagenesis was conducted on the Dsg3-ΔEA-GFP backbone using QuikChange II (Agilent) to generate Dsg3-ΔEA-W2A using sense primer 5′-GGTACAAACGTGAAGCGGTGAAATTTGC-3′ and antisense primer 5′-GCCAAATTTCACCGCTTCAGTTTG TACC-3′. Site-directed mutagenesis was conducted on the DP-mCherry backbone using QuikChange II to generate DP-S2849G-mCherry mutants using sense primer 5′-GGGTCCCGGAGA TCTCCGGGACCC-3′ and antisense primer 5′-GGGTCCCGGAGA TCTCCGGGACCC-3′. All constructs were verified by sequencing and demonstrated to be expressed at the expected molecular weight (Fig. S4).

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Concentration. Mock treatment used 1 µl/ml of DMSO in medium to achieve a 50-nM final concentration.

Gö6976 inhibitor (Abcam or Biovision) was diluted in DMSO to 50 µM and used at 1 µl/ml in medium to achieve a 50-nM final concentration. Mock treatment used 1 µl/ml of DMSO in medium to achieve a 50-nM final concentration. Mock treatment used 1 µl/ml of DMSO in medium to achieve a 50-nM final concentration.

Inhibitor treatment
Gö6976 inhibitor (Abcam or Biovision) was diluted in DMSO to 50 µM and used at 1 µl/ml in medium to achieve a 50-nM final concentration. Mock treatment used 1 µl/ml of DMSO in medium to achieve a 50-nM final concentration.

Calcium switch
Low-Ca²⁺ medium was generated by incubating FluoroBrite DMEM (Thermo Fisher Scientific) with Chelex resin (Bio-Rad) for 1 h, rotating at 4°C, followed by sterile filtration and supplementation with 3 mM EGTA (Millipore). For live-cell fluorescence polarization experiments, cells were switched from normal-Ca²⁺ FluoroBrite DMEM (~1.8 mM) to low-Ca²⁺ medium while on the microscope stage. For all other experiments, the cells were incubated with low-Ca²⁺ medium at 37°C and 5% CO₂ for 90 min while the treatment (mock or Gö6976) was maintained if applicable.

Dispase fragmentation assay
Cells were seeded, transfected (if applicable), and cultured to confluence in 12-well tissue culture plates. After drug and Ca²⁺ treatment (as described above), cells were incubated with 1 U/ml dispase (Sigma-Aldrich) for 30–45 min. The released monolayer was gently rinsed with PBS and subjected to mechanical stress by pipetting. Fragments were fixed in PFA and counted using a dissecting microscope.

EM
HaCaT cells were plated on 10-mm Transwell inserts with 0.4-µm pores (Corning). Once grown to 80% confluence, cells were either treated with 50 nM Gö6976 or mock treated for 1 h at 37°C. Cells were switched into either normal-Ca²⁺ medium (for 90 min) or low-Ca²⁺ medium (for 10 min [mock] or 90 min [Gö6976]) at 37°C. After treatment, cells were fixed with EM-grade fixative (2.0% glutaraldehyde in 0.2 M NaCacodylate, pH 7.2) postfixed in 0.2 M Na Cacodylate buffer containing 1% OsO₄, washed in 0.1 M NaCacodylate buffer (3×), treated with 1% low molecular weight tannic acid in 0.05 M NaCacodylate for 30 min at RT, washed in 0.05 M NaCacodylate containing 1% NaSO₄ for 5 min at RT, and dehydrated using a graded EtOH series. After dehydration, samples were embedded (Embed 812, Electron Microscopy Sciences), thin sectioned at 70–100 nm, and placed on formvar-coated copper mesh grids for transmission EM imaging. After drying, the sections were poststained with a 1:1 mixture of 1% uranyl acetate/EtOH for 5 min in the dark, followed by 5 min in Reynolds’s lead citrate solution.

Images were acquired with a Tecnai Spirit T12 120-kV transmission electron microscope (FEI) operating at 80 kV at 42,000× nominal magnification on a BioSprint 29 charge-coupled device camera (AMT). Desmosomes in which both leaflets of the plasma membrane were clearly visible were used for quantification of intermembrane distance. Intermembrane distance values were determined by taking the average of five measurements made at different points along the length of each desmosome in ImageJ (National Institutes of Health).

Fluorescence polarization microscopy
Fluorescence polarization microscopy was conducted as described previously (Bartle et al., 2017). Experiments were conducted using a Nikon Ti-2 microscope, equipped with a motorized stage, stage-top incubator to maintain 37°C and 5% CO₂ (Tokai Hit, INUBG2SF-TIZB), and 60× 1.49-NA objective. A cleanup polarizer, half-wave plate, and lens were used to polarize and focus a 488-nm laser (Coherent) at the microscope back focal plane (ThorLabs). A motorized mount (PRM1Z8; ThorLabs) controlled half-wave plate rotation and the orientation of the excitation polarization. Images were captured with an ORCA-Flash 4.0 v3 complementary metal–oxide–semiconductor camera (Hamamatsu). The system was controlled with Nikon Elements software. A series of four images were collected at 0°, 45°, 90°, and 135° excitation polarization for each order factor image.

Fluorescence polarization microscopy analysis
Fluorescence polarization images were analyzed using custom-built Polarized Order Detection Software (PODS) v1.0 for Matlab (MathWorks), as previously published (Bartle et al., 2017). Briefly, data images were corrected with the mean of three flatfield images acquired by imaging a green fluorescent slide (Chroma) to control for uneven laser illumination and polarization-dependent intensity changes. A binary mask was then generated on the average intensity image to identify desmosomes. Pixel-by-pixel order factor was calculated as previously described and displayed within the mask.

FRAP microscopy
FRAP experiments were conducted on a Nikon Ti-2 A1R confocal microscope, equipped with a motorized stage, stage-top incubator to maintain 37°C and 5% CO₂ (Tokai Hit), and 60× 1.49-NA objective. Images were acquired in Nikon Elements at Nyquist resolution with 488- and 568-nm excitation lasers, gallium arsenide phosphide detectors, and 525/50 and 595/30 emission filters. Five prebleach images were acquired, and postbleach images were acquired every 2 min for 20 min.

FRAP analysis
Using Fiji, intensity was measured for the bleach, reference, photobleaching correction, and background regions for each cell at each time point. The bleach and reference intensities were corrected for background and photobleaching. GraphPad Prism (GraphPad Software) was used to fit the intensities to an exponential and calculate the mobile fraction and t½ for each bleach region. For population analysis, mean normalized intensity and
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**Immunofluorescence staining**
Fixation and labeling protocols were done as described previously (Stahley et al., 2016). Briefly, cells were preextracted for 60 s with 0.2% Triton X-100 and 300 mM sucrose in PBS with calcium and magnesium at 37°C. Cells were then fixed in 4% PFA prepared fresh from 16% EM-grade material (Electron Microscopy Sciences) for 12 min, followed by 30-min blocking and permeabilization with 5% normal horse serum, 5% normal goat serum, 1% BSA, and 0.25% Triton X-100, with multiple washes between steps. Cells were incubated in primary antibody for 3 h and secondary antibody for 1 h.

**Cell lysate preparation, immunoblots, and antibodies**
Cells were homogenized in radioimmunoprecipitation assay buffer containing SDS, NP-40 (EMD Millipore), and protease/phosphatase inhibitor (Complete, Roche). All lysates were clarified by centrifugation. Protein concentrations were measured by the bicinchoninic acid method (Pierce, Thermo Fisher Scientific). A quantity of 65 µg per sample was loaded in 8% SDS-polyacrylamide gels under denaturing conditions. The size-fractionated proteins were electrobotted to polyvinylidene difluoride (Bio-Rad) membranes. The membranes were treated with Intercept Blocking Buffer (LI-COR) for 1 h and incubated overnight with appropriate primary antibodies sequentially in Intercept Antibody Diluent (LI-COR). Membranes were washed and incubated with secondary antibodies (goat anti-rabbit RDye 800 CW and goat anti-mouse 680 LT; LI-COR) for 1 h and washed with TBS with Tween. Images were captured using the Odyssey CLx Imaging system (LI-COR). Primary antibodies were Dsg3 (clone 5G11, Thermo Fisher Scientific), DP I-II (ab127941, Abcam), DP I & II (CBL173, Millipore), DP pS2849 (600-401-J65, Rockland), Vinculin (V9131, Millipore Sigma), β-tubulin (2146, Cell Signaling Technology), GFP (ab290, Abcam), and mCherry (ab167453, Abcam).

**Antibodies**
Primary antibodies for microscopy were anti-DP (DPI/II, Bethyl Labs), anti-γ-catenin (H-80, Santa Cruz Biotechnology), and anti-Dsg3 (G194, Progen Biotechnik). Secondary antibodies were Alexa Fluor 488- and 647-conjugated goat anti-mouse or goat anti-rabbit (IgG H&L) from Invitrogen.

**dSTORM analysis**
Reconstructed dSTORM images were rendered in Nikon Elements. Photoswitching imaging buffer included glucose oxidase (Sigma-Aldrich), glucose (Sigma-Aldrich), catalase (Roche), and β-mercaptoethanol (Sigma-Aldrich; Rust et al., 2006).

**Statistics**
All statistical analysis was performed with GraphPad Prism. P values were determined by Student’s t test when measuring significance between two conditions or by one-way ANOVA with multiple comparisons when measuring significance between more than two conditions. Curve fitting for FRAP analysis was done using one-phase association-fitting algorithms, or linear regression in the absence of recovery, where the plateau determined the mobile fraction.

**Online supplemental material**
Fig. S1 contains controls for fluorescence polarization microscopy. Fig. S2 has representative images of Dsg2-mCherry–expressing cells showing that exchange is reduced in hyperadhesions. Fig. S3 shows that Dsg3 order factor is not impacted by cotransfection with DP-S2849G-mCherry. Fig. S4 contains blots showing expression of tagged proteins. Table S1 is a summary of all Dsg3-ΔEA-GFP order factors, and Table S2 is a summary of all mobile fractions.

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Author contributions: E.I. Bartle carried out the majority of experiments and data analysis. T.C. Rao carried out the biochemical experiments and analysis, generated reagents, and provided cell culture support. R.R. Beggs and W.F. Dean carried out additional key experiments and analysis. T.M. Urner collected data, developed dSTORM and fluorescence polarization analysis methods, and provided critical discussions of the results. A.P. Kowalczyk provided critical discussions of the results and experimental design. E.I. Bartle and A.L. Mattheyses conceived and designed the study, interpreted results, and wrote the manuscript. All authors provided intellectual input, revised, edited, and approved the manuscript.

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Supplemental material

Figure S1. **Fluorescence polarization microscopy controls.** (A) Dispase fragmentation assay for cells expressing Dsg3-ΔEA-GFP, Dsg3-link-GFP, or GFP. Dsg3-ΔEA-GFP did not impact adhesive strength or the ability to acquire hyperadhesion (n = 3; mean ± SD). (B and C) HaCaT cells transfected with Dsg3-ΔEA-GFP were imaged by fluorescence polarization microscopy before and 1 h after treatment with Go6976 while maintained in normal-Ca$^{2+}$ medium. (B) Representative intensity and order factor images. Scale bar = 5 µm. (C) Order factor quantification (mean ± SD; n = 15 cells; n.s., not significant P > 0.05; Student’s t test). (D) HaCaT cells were transfected with Dsg3-ΔEA-GFP, mock treated, maintained in normal-Ca$^{2+}$ medium (top) or switched to low-Ca$^{2+}$ medium (bottom), and imaged for 30 min. Representative intensity images at each time point are shown. Scale bar = 5 µm. (E) Intensity over time was quantified by masking each image and calculating the integrated intensity. Integrated intensity was normalized and averaged across the population of cells (D and E are the same cells with analysis presented in Fig 1; normal-Ca$^{2+}$ medium mock [n = 6] or Go6976 [n = 6] treated; low-Ca$^{2+}$ medium mock [n = 11] or Go6976 [n = 15] treated; mean ± SEM). (F) Order factor from Go6976-treated cells fixed at 0, 30, or 90 min after the switch to low-Ca$^{2+}$ medium (mean ± SD). Order factor was not significantly different at 30 or 90 min in low Ca$^{2+}$ (n = 7 cells each condition; n.s., not significant P > 0.05; ANOVA).
Figure S2. **Representative images of cells expressing Dsg2-mCherry showing that exchange is reduced in hyperadhesion.** Representative cells from data presented in Fig 4. (A) HaCaT cell transfected with Dsg2-mCherry, mock treated, and maintained in normal Ca\(^{2+}\) with bleach region of interest (ROI) indicated. Scale bar = 5 µm. (B) Zoom-in of cell border in bleach ROI over time. Recovery of individual puncta is underscored by the dashed lines. (C) Fluorescence intensity plotted as a function of time. Mobile fraction = 20%. (D) Representative HaCaT cell transfected with Dsg2-mCherry, Gö6976 treated, and maintained in low-Ca\(^{2+}\) medium for 90 min. Scale bar = 5 µm. (E) Zoom-in of cell border in bleach ROI over time with individual puncta underscored by the dashed lines. (F) Fluorescence intensity plotted as a function of time. Mobile fraction = 7%.

Figure S3. **Dsg3 order factor is not impacted by DP-S8249G-mCherry transfection.** (A and B) Representative images of HaCaT cells cotransfected with (A) Dsg3-ΔEA-GFP and DP-wt-mCherry or (B) Dsg3-ΔEA-GFP and DP-S2849G-mCherry and maintained in normal-Ca\(^{2+}\) medium. Scale bar = 5 µm. (C) Order factor of Dsg3-ΔEA-GFP in cells expressing DP-wt-mCherry (n = 27) or DP S2849G-mCherry (n = 26; ns, not significant, P > 0.05; Student’s t test). (D) Quantification of adhesive strength by dispase fragmentation assay in HaCaT cells expressing either wt or mutant DP with Gö6976 and Ca\(^{2+}\) conditions indicated (n = 3; mean ± SD; **, P ≤ 0.01; Student’s t test).
Figure S4. **Tagged proteins are expressed at the expected molecular weights.** Cells were transiently transfected with plasmids as indicated. 48 h after transfection, cells were harvested for SDS-PAGE and Western blot analysis. Each blot includes an untransfected control. (A) Cos7 cells transfected with GFP, Dsg3-ΔEA-GFP, or Dsg3-ΔEA-W2A-GFP blot were probed for Dsg3 or GFP. Tubulin was used as a loading control. (B) Cos7 cells transfected with mCherry, DP-mCherry, or DP-S2849-mCherry. Blot was probed for DP or mCherry with tubulin as a loading control. (C) A431 cells transfected with E-cad-GFP. Blot was probed with GFP and tubulin. (D) A431 cells transfected with PG-mEmerald. Blot was probed with GFP, and tubulin was used as a loading control. (E) A431 cells transfected with Dsg2-mCherry. Blot was probed with mCherry, and GAPDH was used as a loading control.

Provided online are two tables. Table S1 is a summary of Dsg3-ΔEA-GFP order factors. Table S2 is a summary of mobile fractions.