Three Functions of Cadherins in Cell Adhesion

Jean-Léon Maître¹ and Carl-Philipp Heisenberg²

Cadherins are transmembrane proteins that mediate cell-cell adhesion in animals. By regulating contact formation and stability, cadherins play a crucial role in tissue morphogenesis and homeostasis. Here, we review the three major functions of cadherins in cell-cell contact formation and stability. Two of those functions lead to a decrease in interfacial tension at the forming cell-cell contact, thereby promoting contact expansion — first, by providing adhesion tension that lowers interfacial tension at the cell-cell contact, and second, by signaling to the actomyosin cytoskeleton in order to reduce cortex tension and thus interfacial tension at the contact. The third function of cadherins in cell-cell contact formation is to stabilize the contact by resisting mechanical forces that pull on the contact.

Introduction

The ability of cells to adhere to one another is a fundamental property in the evolution of multicellularity. Cadherins are transmembrane cell-cell adhesion molecules conserved among metazoan organisms, which play essential roles in tissue morphogenesis and homeostasis [1–5]. During morphogenesis, tissues can change in size and shape, and form distinct cell layers. Cadherins function in tissue morphogenesis by controlling both cell-cell adhesion and cell signaling. For example, cadherins are involved in determining cell shape and position within the ommatidium of the Drosophila melanogaster retina [1]. Cadherins have also been implicated in germ cell positioning and migration in zebrafish and Caenorhabditis elegans [4,6], oocyte positioning within the egg-chamber of Drosophila [7], epithelial folding [3,8], and mesoderm/endoderm cell internalization in Drosophila and Caenorhabditis elegans [2,9]. Cells also use cadherins to mediate signals that can control cell fate specification [10–12], cell polarity [13,14] and cell proliferation [15–17]. Considering this apparent complexity of cadherin function in morphogenesis, it is difficult to always clearly distinguish between the different morphogenetic functions of cadherins. In this review, we aim at dissecting the different major functions of cadherins in cell-cell contact formation and stabilization.

When cells contact each other, cadherins from the opposing cells located at the site of contact form trans-bonds across the contact. Once engaged in trans-bonds, cadherins can regulate the formation of the cell-cell contact in three distinct ways: by reducing the local interfacial tension directly through adhesion tension and indirectly through signaling to the actomyosin cytoskeleton, and by establishing the mechanical coupling of contacting cells.

Adhesion Tension Function of Cadherins

At the macroscopic level, cell and tissue shapes are regulated by surface tension-like properties (Box 1) [18–23]. Similar to the rounding up of liquids, surface tension-like properties can explain the rounding up of cells during cytokinesis [24,25] or of tissues when explanted [23,26,27]. Furthermore, in analogy to the arrangement of soap bubbles, surface tension-like properties have been used to explain cell packing in epithelia [13,28–30] and the configuration of a subset of cells within tissues [1,31]. Reducing the surface tension of a specific interface, such as the cell-cell contact, tends to increase its size. One function of cadherin in cell-cell adhesion is to promote contact formation by directly reducing the surface tension at the cell-cell interface via adhesion tension. The adhesion tension arises from cadherin binding over the contact thereby generating a negative tension that expands the contact area [32,33].

At the microscopic scale, adhesion tension is thought to originate from the chemical binding of cadherins, which is energetically favorable. As a consequence of binding, cadherins accumulate at the cell-cell contact where they are stabilized, because for cadherins to exit the contact zone, the cadherin trans-bond would need to be broken. Cadherin accumulation at the contact might influence contact size in several ways: cadherins could promote contact expansion via zippering of the contact edge (Figure 1). Consistent with this, accumulations of cadherins are often found at the edges of the cell-cell contact [33,34]. Zippering of the cell-cell contact by cadherins could be achieved by shortening the cadherin trans-bond length after a catch phase and/or stabilizing random encounters of opposing plasma membranes [35,36]. Alternatively or in addition, cadherins could control contact size via protein crowding (Figure 1). The crowding of proteins at the cell-cell contact can lead to the building of a lateral pressure in the plasma membrane [37], which forces the contacting plasma membrane to spread. Such lateral pressure from protein crowding is unlikely to arise solely from the action of cadherins, as cadherins commonly cluster and do not occupy the whole membrane surface of the cell-cell contact [33,34,38]. However, other proteins associated with cadherins, or proteins that require cadherins to accumulate at the cell-cell contact could build an osmotic pressure that leads to the depletion of unbound cadherins and other proteins that are not yet sequestered at the cell-cell contact (Figure 1). Such osmotic depletion of unbound adhesion molecules would be expected to decrease the lateral pressure within the cell-cell contact and thus reduce adhesion tension and limit contact expansion.

While the exact microscopic mechanisms underlying the generation of macroscopic adhesion tension are yet unknown, several attempts were made to measure or calculate adhesion tension. For instance, contact separation experiments have been used to obtain information about the adhesion energy at cell-cell contacts. Adhesion energy is the work required to build or cleave an area of contact at the surface of passive materials and scales with adhesion tension, granted that the process of adhesion is reversible, i.e., that the energy of formation and separation of an adhesive contact are the same. However, while the assumption that the energy of formation and separation of an adhesive contact are the same was validated for passive viscoelastic materials
interfacial tension at cell–cell junctions [41,42]. In zebrafish shapes using sophisticated image analysis tools, together sion contributes. In 1. Cortext tension: the contribution of the actomyosin cortex to the tension of a given surface or interface. In isolated cells or at contact-free surfaces in a tissue, it generally constitutes most of the tension [33,45,46,113].

Adhesion tension: the contribution of adhesion molecules to the tension of a given interface. Adhesion molecules are typically expected to promote the expansion of cell–cell contacts. Therefore, adhesion tension is expected to be a negative tension or, as formulated above for surface tension, it is the energy required to increase a given interface by a unit area.

Adhesion coupling: the force opposed by adhesion molecules to intra- or extra-cellular forces. It can be studied by measuring the force required to separate adhesion molecules (in the pN range [115,116]) or cells (in the nN range [33,46,95]).

Adhesion signaling: the chemical modifications resulting from cadherin trans-binding. Signalings from cadherins can have consequences on a broad range of cellular processes, from rapid cytoskeletal rearrangements, which can modify interfacial tension [33,34] and adhesion coupling [81,82], to the long-term differentiation of tissues [11].

Box 1

Glossary.

**Surface or interfacial tension**: the energy required to decrease by a unit area a given surface (or interface). Similar to liquids, surface tension gives cells and tissues an apparent stiffness, which resists mechanical stresses. Tensions are measured in N.m⁻¹. Typical tensions associated with cells range from 10⁻⁵–10⁻⁴ N.m⁻¹ [45,46,113] while tissue surface tensions are usually found in the 10⁻³ N.m⁻¹ range [23,114].

**Cortex tension**: the contribution of the actomyosin cortex to the tension of a given surface or interface. In isolated cells or at contact-free surfaces in a tissue, it generally constitutes most of the tension [33,45,46,113].

**Adhesion tension**: the contribution of adhesion molecules to the tension of a given interface. Adhesion molecules are typically expected to promote the expansion of cell–cell contacts. Therefore, adhesion tension is expected to be a negative tension or, as formulated above for surface tension, it is the energy required to increase a given interface by a unit area.

**Adhesion coupling**: the force opposed by adhesion molecules to intra- or extra-cellular forces. It can be studied by measuring the force required to separate adhesion molecules (in the pN range [115,116]) or cells (in the nN range [33,46,95]).

**Adhesion signaling**: the chemical modifications resulting from cadherin trans-binding. Signalings from cadherins can have consequences on a broad range of cellular processes, from rapid cytoskeletal rearrangements, which can modify interfacial tension [33,34] and adhesion coupling [81,82], to the long-term differentiation of tissues [11].

[39,40], this might be different for living cells [33]. In fact, cell–cell adhesion is an active process that involves extensive molecular reorganization of the contact zone (see below), suggesting that the energies associated with contact forma- tion and separation are different. It is thus questionable whether measuring the energy of cell–cell contact cleavages represents a suitable approach for obtaining information about the adhesion energy involved in contact formation.

More recently, the shape of adhering cells was used to directly calculate adhesion tension by determining the interfacial tension at the cell–cell contact, to which adhesion tension contributes. In *Drosophila* embryos, the analysis of cell shapes using sophisticated image analysis tools, together with modeling of cell mechanics, allowed determining the interfacial tension at cell–cell junctions [41,42]. In zebrafish embryos, the specific contribution of adhesion tension to the interfacial tension at the cell–cell contact was determined by measuring the interfacial tension at the contact in the presence and absence of adhesion tension [33]. Notably, adhesion tension was found to contribute only little to the total interfacial tension at the contact between these cells. This suggests that adhesion tension plays only a minor role in setting the cell–cell contact size. Future experiments addressing the contribution of adhesion tension in other cell types will be needed to determine whether adhesion tension can also play a more prominent function in cell–cell contact formation and stabilization.

**Signaling Function of Cadherins**

Besides adhesion tension, cells can also use different strategies to reduce interfacial tension at the cell–cell contact and thereby increase the size of the contact. Actomyosin contractility is a decisive factor influencing surface tension of cells in a broad variety of organisms [43–46]. Once cells contact each other, they often reorganize their actomyosin cytoskeleton at the cell–cell interface [34], resulting in a reduction of interfacial tension at the contact, thereby expanding the contact size (Box 1) [33]. Contact-mediated reorganization of the actomyosin cytoskeleton is commonly attributed to signaling from the cadherin adhesion complex [47], although other cadherin-independent processes also might control the organization of the actomyosin cytoskel- eton at the contact [48,49].

Upon trans-binding of cadherins, the local chemistry changes at the cell–cell contact (Figure 2) [50]. For example, plasma membrane phosphatidylinositol (3,4,5)-trisphosphate (PIP₃) accumulates, which in turn leads to local activation of Rac1 at the contact [34,47]. Likewise, p120-catenin binds to cadherins engaged in trans-binding, leading to local down-regulation of RhoA [51]. Also, β-catenin becomes recr uited to the cadherin adhesion complex at the cell–cell contact and competes with Arp2/3 [52] and/or recruits RhoGEFs by interacting with centralspindlin [53]. Since both Arp2/3 and Rho GTPases, such as RhoA and Rac1, are known regulators of the actin cytoskeleton, changing their function leads to alterations in cytoskeleton organiza- tion at the cell–cell contact and, consequently, defects in contact formation [34,51,52].

Taken together, cadherins, in particular by recruiting cate nin, trigger signals that modulate the actomyosin cytoskeleton at the cell–cell contact. Given that interfacial tension at the cell–cell contact is strongly influenced by actomyosin contractility, cadherin-mediated signaling is thus expected to modulate cell–cell contact interfacial tension and thus contact formation (Figure 2).

Cell–cell contact formation is typically followed by contact maturation. In epithelial cells, cell–cell contact maturation in- volves the establishment and stabilization of apico-basal cell polarity and the formation of tight junctions [54]. Interestingly, while the cortical actomyosin cytoskeleton is usually reduced at the contact during initial cell–cell contact formation and expansion [34], junctional myosin appears to be essential for epithelial contact maturation [55,56]. Myosins can have distinct functions in mature epithelial junctions with non-muscle myosin 2 heavy chains A and B controlling cadherin clustering and actin dynamics, respectively. These distinct functions of myosin isoforms in epithelial junctions are thought to be regulated by the GTPases RhoA and RhoG. Recently, Rap1 was also identified as a regulator of epithelial folding by regulating cytosskeletal anchoring of cadherins through β-catenin [8]. Further studies will be needed to understand the temporal and spatial regulation of the molecular composition of the adhesion complex and associated cytoskeleton during cell–cell contact maturation.

Cells not only make but also break contacts with their neighboring cells during various morphogenetic events,
N-cadherin is specifically required for mediating contact inhibition of locomotion in neural crest cells [63]. Other adhesion molecules, such as the atypical cadherins Fat and Dachs, or cadherin-associated proteins, such as nectins and DDR1, also seem to control cell–cell contact formation by regulating actomyosin contractility at the contact [13,67–69]. To understand how cadherin-mediated signaling differently controls contact growth and shrinkage, it will be key to identify and characterize contact- and cell type-specific effector proteins modulating the actomyosin cortex.

**Mechanical Scaffolding Function of Cadherins**

By influencing cell–cell interfacial tension, the contractile actomyosin cytoskeleton plays a key role in contact formation and stabilization. In addition, the actomyosin cytoskeleton exerts forces pulling on the cell–cell junctions [9,70–72]. Recently, these pulling forces were directly visualized using a stretchable cadherin–FRET sensor [73]. Pulling forces generated by the actomyosin cytoskeleton are resisted by the mechanical coupling function of the cadherin adhesion complex at the contact, thereby preventing contact separation (Box 1) [2].

To mediate the mechanical coupling between cells (the force or load that adhesive bonds can withstand), cadherins anchor to the actomyosin cytoskeleton. When under mechanical load, cadherins can transmit through their cytoskeletal anchor forces to the whole interconnected cytoskeleton. The ‘basic’ cytoskeletal anchoring of cadherins is thought to be mediated by β- and α-catenin [10,74,75].

Interestingly, within the cadherin mechanical scaffold, the weakest link appears to be in the cytoplasmic rather than in the extracellular part of the scaffold (Figure 3). Upon separation, cells often remain connected by plasma membrane tethers at the base of which cadherins accumulate, suggesting that detachment occurs within the cell and that cadherins trans-bonds remain intact [33,60,76,77]. Similarly, retraction fibers at the back of migrating cells or attached to dividing cells and adhesion plaques suggest that cell-matrix attachment is limited by the anchoring of integrins to the actin cytoskeleton rather than by the integrin-matrix binding strength [78]. These observations suggest that morphogenetic processes that involve cell–cell separation, such as collective cell migration and cell ingestion, may be primarily influenced by the coupling of cadherins to the actomyosin cytoskeleton rather than by the strength of cadherin trans-bonds. Therefore, the regulation of the cytoskeletal anchoring of cadherins is a critical process in morphogenesis.

Precisely which bond limits the cytoskeletal anchoring of cadherins remains unclear. Interestingly, the fusion of E-cadherin and α-catenin can efficiently substitute for a loss of α-catenin and rescue cell–cell adhesion and embryonic development in *Drosophila* α-catenin or E-cadherin mutants [10,75]. This suggests that E-cadherin, β- and α-catenin unbinding is not required for efficient separation of cell–cell contacts and that the cadherin scaffold can break downstream of α-catenin (Figure 3). However, exogenously forced cell–cell contact separation leads to the detachment of the adhesion complex from the actomyosin cytoskeleton between β- and α-catenin, suggesting that the weakest link is upstream of α-catenin [33].

The mechanical scaffolding function of cadherins can be regulated via chemical signaling. Different kinases (Abl, Fer, Fyn) were shown to regulate the interaction of β- and α-catenin via tyrosine phosphorylation of β-catenin [79,80].

---

**Figure 1. Adhesion tension.**

Upon cadherin trans-binding, opposing plasma membranes are brought closer together, thereby expanding the contact through zipping. Accumulation of cadherins and other proteins at the contact zone can also result in membrane crowding and expansion of the contact zone via lateral pressure. Finally, cadherin accumulation at the contact zone might result in an osmotic-driven movement of proteins outside of the contact zone, resulting in lateral pressure shrinking the contact. The combination of these effects results in the adhesion tension function of cadherins in cell–cell adhesion, which has a direct effect on the surface tension at the cell–cell contact and thereby regulates its size.
p120-catenin also plays an important role in regulating the phosphorylation of β-catenin upstream of these kinases \cite{81,82}. However, the p120-catenin binding site of E-cadherin appears to be dispensable for normal Drosophila oogenesis and embryonic development \cite{83}.

Cytoskeletal anchoring of cadherins can also be achieved by recruiting additional actin-binding proteins such as vinculin and/or EPLIN \cite{84,85}. Interestingly, recruitment of these actin-binding proteins is mechanosensitive \cite{71,84,86,87}. This mechanosensitive recruitment of additional actin binders enables cells to adapt the strength of their mechanical scaffolding function of cadherins to the magnitude of external and internal mechanical stresses (Figure 3). At the molecular level, mechanosensing of the adhesion complexes is thought to be achieved by the unwinding of protein domains upon stretching, which would reveal cryptic binding sites \cite{88}. When tension is applied to adherens junctions, vinculin-binding sites of α-catenin become exposed upon protein stretching, promoting vinculin recruitment to the adhesion complex \cite{86}. Vinculin then further connects the adhesion complex to the actin cytoskeleton, thereby strengthening the cadherin mechanical scaffold \cite{84,86,89}. Interestingly, vinculin, whose structure closely resembles the one of α-catenin, may also stretch under mechanical load and thereby recruit further proteins to the cell junction in a mechanosensitive manner \cite{90}. In addition to α-catenin and vinculin, β-catenin and the extracellular domain of cadherin can unfold upon tension, as suggested by both in vitro and in silico studies \cite{91–93}. However, to understand the molecular mechanisms of mechanosensing and subsequent reinforcement of the mechanical coupling of the adhesion complex, further information on the response of other molecules of the adhesion complex to mechanical load will be needed.

To probe the mechanical coupling of the adhesion complex within cells, several methods can be employed.

**Dissecting the Different Functions of Cadherins in Morphogenesis**

Cell sorting is a morphogenetic event driven by successive breaking and making of cell–cell contacts. Therefore, all three functions of cadherins described above are potentially involved in cell sorting. The differential adhesion hypothesis gives a thermodynamic description of cell sorting. In view of the differential adhesion hypothesis, cells reorganize their cell–cell contacts according to the amount of adhesion molecules available, until an energetic equilibrium is reached \cite{100}. However, in various cases, the number of adhesion molecules expressed in cells does not scale with their sorting behavior, contrary to the predictions by the differential adhesion hypothesis \cite{33,46,101–103}.

In zebrafish, distinct germ layer progenitor cells sort primarily according to their ability to regulate the actomyosin cytoskeleton at the cell–cell contact and the cytoskeletal anchoring strength of cadherins \cite{33}. Notably, the
contribution of adhesion tension to progenitor cell sorting appears to be negligible when compared to cytokoskeletal reorganization regulating cell–cell contact formation. Therefore, cadherins appear to primarily function in zebrafish germ-layer progenitor–cell sorting through their signaling and mechanical coupling functions rather than via their adhesion tension function.

In *Xenopus*, cadherins do not seem to be the only molecules at the cell–cell contact regulating sorting of germ layer progenitors [60,101]. During gastrulation, ephrins were shown to regulate contact repulsion between ectoderm and mesoderm [60]. There, the regulation of contact repulsion is mediated by the small GTPases Rac and RhoA, which in turn might modulate the actomyosin cytoskeleton. Alternatively, ephrins were shown to regulate cell–cell adhesion via ADAM10-mediated cleavage of the extracellular domain of E-cadherin [104]. Similarly to the situation in *Drosophila* and zebrafish [2,33], contact separation of germ layer progenitors in *Xenopus* leads to the formation of membrane tethers, suggesting that cadherin-mediated cell–cell adhesion breaks in the cytoplasmic part of the cadherin adhesion complex [60].

Another widespread morphogenetic event in which all three functions of cadherins are at play is epithelial morphogenesis. During epithelial morphogenesis, tissue integrity needs to be preserved. For this, mechanical coupling of the junctional adhesion complex is critical. In fact, when cytoskeletal coupling of the junctional adhesion complex is affected, such as in the ventral furrow cells of *Drosophila* β-catenin mutants, adhesion complexes are extruded from the cell body and morphogenesis halts [2]. Cytoskeletal anchoring of cadherins is also critical for maintaining tissue integrity during cytokinesis within *Drosophila* epithelial tissues. At early stages of cytokinesis, cadherins coupled to the cytokinetic actomyosin ring towards the apical junction of the dividing cells. Only shortly before a new cell–cell junction between the daughter cells is formed, cadherins disconnect from the actomyosin ring [105].

Besides mechanical coupling, cadherins also regulate interface tension at cell–cell boundaries to regulate cell shape and positioning. In the *Drosophila* retina, for example, cell shape is regulated by the presence of E-cadherin and/or whether cadherins directly lower interface tension via adhesion tension or indirectly by reorganizing the actomyosin cortex remains to be investigated. Similarly, in the extending germ band of the *Drosophila* embryo, cadherins accumulate at expanding cell–cell junctions, whereas they are depleted from shrinking boundaries [106,107]. At shrinking junctions, actomyosin appears to be the main factor determining interfacial tension controlling cell neighbor exchange [43,59,108]. However, the specific contribution of cadherins to the interfacial tension in this process is still unknown. Recently, the differential localization of cadherins along junctions with different orientations was shown to direct the flow of actin and myosin from the cell apex towards those junctions. Interestingly, actomyosin flows towards the junctions containing less E-cadherin, suggesting that the flow results from unbalanced coupling of the actomyosin network to the junction rather than its total coupling strength [59]. It will be interesting to further investigate how cadherins, through their distinct functions, can direct flows of actomyosin in different morphogenetic movements [9,109,110].

**Conclusion**

Since their discovery [111,112], the functions of cadherins in morphogenesis have been intensively studied. Recent technological advances gave us a more comprehensive biophysical description of how cadherins function in morphogenesis. It appears that cadherins can have rather distinct functions in cell adhesion and that therefore using the term ‘adhesion’ in conjunction with cadherin function might be confusing. We thus would like to propose to use the term ‘adhesion tension’ when referring to the direct effect of cadherins on interfacial tension (Box 1). For the indirect effect of cadherins on interfacial tension through signaling to the cortical cytoskeleton, we would propose to use the term ‘adhesion signaling’. Finally, to refer to the function of cadherins in coupling the contractile cell cortices at the contact, the term ‘adhesion coupling’ might be most suitable. Obviously, any other terms might be equally good as long as they clearly distinguish between the specific adhesion functions of cadherins.

As a final note, we would like to emphasize that while cadherins clearly display different adhesion functions, it is experimentally very difficult to specifically regulate any of
those functions alone. Changing adhesion signaling, for example, will predictably also alter adhesion coupling by affecting the cytoskeleton to which the adhesion complex is coupled. Future studies will have to address the interplay between the different functions of cadherins and their co-regulation in tissue morphogenesis.

Acknowledgments

We are grateful to Ewa Paluch and Guillaume Saiboux and Nicolas Borghi for advice and reading earlier versions of this review. J.-L.M. is supported by an EMBO-LTF 1195-2012. Borghi for advice and reading earlier versions of this review. J.-L.M. is coupled. Future studies will have to address the interplay between the different functions of cadherins and their co-regulation in tissue morphogenesis.

References

1. Hayashi, T., and Cartew, R. (2004). Surface mechanics mediate pattern formation in the developing retina. Nature 431, 647–652.
2. Martin, A.C., Gelbart, M., Fernandez-Gonzalez, R., Kасhubev, M., and Wiescha, E.F. (2010). Integration of contractile forces during tissue invagination. J. Cell Biol. 188, 735–749.
3. Wang, Y.C., Klar, Z., Kasprzak, M., Wieschaus, E.F. (2012). Differential adhesion-cadherin junctions are associated with initiation of epithelial folding. Nature 484, 390–393.
4. Chihara, D., and Nance, J. (2012). An E-cadherin-mediated hitchhiking mechanism for C. elegans germ cell internalization during gastrulation. Development 139, 2547–2556.
5. Arboleda-Estudillo, Y., Krieg, M., Sohme, J., Lotica, N.A., Muller, D.J., and Heisenberg, C.-P. (2010). Movement directionality in collective migration of germ layer progenitors. Curr. Biol. 20, 161–169.
6. Kardash, E., Reichman-Fried, M., Maitre, J.-L., Boldajipour, B., Papusheva, A., Wassmer, J.-J., Hanenberg, M., Krieg, M., Stühmer, J., Licata, N.A., Muller, D.J., and Heisenberg, C.-P. (2010). Movement directionality in collective migration of germ layer progenitors. Curr. Biol. 20, 161–169.
7. Godt, D., and Tepass, U. (1998). Drosophila oocyte localization is mediated by differential cadherin-based adhesion. Nature 395, 376–391.
8. Wang, Y.-C., Khan, Z., and Wieschaus, E.F. (2013). Distinct Rap1 activity states control the extent of epithelial invagination via a-catenin. Dev. Cell 25, 299–309.
9. Roh-Johnson, M., Shemer, G., Higgins, C.D., McClellan, J.H., Werts, A.D., Krieg, M., Stühmer, J., Licata, N.A., Muller, D.J., and Heisenberg, C.-P. (2010). The role of adhesion energy in controlling cell–cell contacts. Curr. Opin. Cell Biol. 23, 508–514.
10. Bath, A.J., and Wadman, M. (2005). Cell–cell interaction controls cell patterning in the Drosophila retina. Proc. Natl. Acad. Sci. USA 104, 2519–2526.
11. Stephenson, R.O., Yamanaka, Y., and Rossant, J. (2010). Disorganized tissues control the extent of epithelial invagination via a-catenin. Development 137, 3383–3391.
12. Wang, Y.-C., Khan, Z., and Wieschaus, E.F. (2013). Distinct Rap1 activity states control the extent of epithelial invagination via a-catenin. Dev. Cell 25, 299–309.
13. Wang, Y.-C., Khan, Z., and Wieschaus, E.F. (2013). Distinct Rap1 activity states control the extent of epithelial invagination via a-catenin. Dev. Cell 25, 299–309.
14. Wang, Y.-C., Khan, Z., and Wieschaus, E.F. (2013). Distinct Rap1 activity states control the extent of epithelial invagination via a-catenin. Dev. Cell 25, 299–309.
15. Wang, Y.-C., Khan, Z., and Wieschaus, E.F. (2013). Distinct Rap1 activity states control the extent of epithelial invagination via a-catenin. Dev. Cell 25, 299–309.
16. Wang, Y.-C., Khan, Z., and Wieschaus, E.F. (2013). Distinct Rap1 activity states control the extent of epithelial invagination via a-catenin. Dev. Cell 25, 299–309.
17. Wang, Y.-C., Khan, Z., and Wieschaus, E.F. (2013). Distinct Rap1 activity states control the extent of epithelial invagination via a-catenin. Dev. Cell 25, 299–309.
18. Manning, M.L., Foty, R.A., Steinberg, M.S., and Schoetz, E.-M. (2010). A role for Rho GTPases and cell–cell adhesion in single-cell motility in vivo. Nat. Cell Biol. 12, 47–53.
19. Godt, D., and Tepass, U. (2002). The Differential Interfacial Tension Hypothesis (DITH): a comprehensive theory for the self-rearrangement of embryonic tissues and cells. J. Biomech. Eng. 124, 188–197.
20. Brodland, G. (2002). The Differential Interfacial Tension Hypothesis (DITH): a comprehensive theory for the self-rearrangement of embryonic tissues and cells. J. Biomech. Eng. 124, 188–197.
21. Brodland, G. (2002). The Differential Interfacial Tension Hypothesis (DITH): a comprehensive theory for the self-rearrangement of embryonic tissues and cells. J. Biomech. Eng. 124, 188–197.
54. Kishikawa, M., Suzuki, A., and Ohno, S. (2008). aPKC enables development
70. de
71. Liu, Z., Tan, J.L., Cohen, D.M., Yang, M.T., Sniadecki, N.J., Ruiz, S.A.,
50. Zaidel-Bar, R. (2013). Cadherin adhesome at a glance. J. Cell Sci.
126
55. Miyake, Y., Inoue, N., Nishimura, K., Kinoshita, N., Hosoya, H., and Yone-
56. Shewan, A., Maddugoda, M., Kraemer, A., Stehbens, S., Verma, S., Kovacs,
57. Smutny, M., Cox, H., Leerberg, J., Kovacs, E., Conti, M., Ferguson, C., Ham-
69. Marcinkevicius, E., and Zallen, J.A. (2013). Regulation of cytoskeletal orga-
65. Marlow, F., Topczewski, J., Sepich, D., and Solnica-Krezel, L. (2002).
63. Theveneau, E., Marchant, L., Kuriyama, S., Gull, M., Moepps, B., Parsons,
876–884.
64. le Duc, Q., Shi, Q., Blonk, I., Sonnenberg, A., Wang, N., Leckband, D., and
61. Speirs, C.K., Jernigan, K.K., Kim, S.-H., Cha, Y.I., Lin, F., Sepich, D.S.,
620
84. le Duc, Q., Shi, Q., Blonk, I., Sonnenberg, A., Wang, N., Leckband, D., and
87. Taguchi, K., Ishiuchi, T., and Takeichi, M. (2011). Mechanosensitive EPLIN-
88. del Rio, A., Perez-Jimenez, R., Liu, R., Roca-Cusachs, P., Fernandez, J.M.,
90. Carisey, A., Tsang, R., Greiner, A.M., Nijenhuis, N., Heath, N., Nazgiewicz,
91. Valbuena, A., Vera, A.M., Oroz, J., Mene ´ndez, M., and Carrio´n-Va´zquez, M.
92. Liu, R., Wu, F., and Thiery, J.-P. (2012). Remarkable disparity in mechanical
95. Chu, Y., Thomas, W., Eder, O., Pincet, F., Perez, E., Thiery, J., and Dufour, S.
96. Yonemura, S., Wada, Y., Watanabe, T., Nagafuchi, A., and Shibata, M. (2010).
99. Tominaga, J., Fukunaga, Y., Abelardo, E., and Nagafuchi, A. (2008).
72. Maruthamuthu, V., Sabass, B., Schwarz, U.S., and Gardel, M.L. (2011). Cell-
84
167
85. Borghi, N., Sorokina, M., Shcherbakova, O.G., Weis, W.I., Pruitt, B.L., Nelson,
86. Yonemura, S., Wada, Y., Watanabe, T., Nagafuchi, A., and Shibata, M.
87. Taguchi, K., Ishiuchi, T., and Takeichi, M. (2011). Mechanosensitive EPLIN-
138
167
92
115
88
86
85
84
83
82
81
80
79
78
77
76
75
74
73
72
71
70
69
68
67
66
65
64
63
62
61
60
59
58
57
56
55
54
53
52
51
50
49
48
47
46
45
44
43
42
41
40
39
38
37
36
35
34
33
32
31
30
29
28
27
26
25
24
23
22
21
20
19
18
17
16
15
14
13
12
11
10
9
8
7
6
5
4
3
2
1

Current Biology Vol 23 No 14 R632
96. Grashoff, C., Hoffman, B.D., Brenner, M.D., Zhou, R., Parsons, M., Yang, M.T., McLean, M.A., Sligar, S.G., Chen, C.S., Ha, T., et al. (2010). Measuring mechanical tension across vinculin reveals regulation of focal adhesion dynamics. Nature 466, 263–266.

99. Conway, D.E., Breckenridge, M.T., Hinde, E., Gratton, E., Chen, C.S., and Schwartz, M.A. (2013). Fluid shear stress on endothelial cells modulates mechanical tension across VE-cadherin and PECAM-1. Curr. Biol. 23, 1024–1030.

100. Foty, R., and Steinberg, M. (2005). The differential adhesion hypothesis: a direct evaluation. Dev. Biol. 278, 255–263.

101. Ninomiya, H., David, R., Damm, E.W., Fagotto, F., Niessen, C.M., and Winklbauer, R. (2012). Cadherin-dependent differential cell adhesion in Xenopus causes cell sorting in vitro but not in the embryo. J. Cell Sci. 125, 1877–1883.

102. Shi, Q., Chien, Y., and Leckband, D. (2008). Biophysical properties of cadherin bonds do not predict cell sorting. J. Biol. Chem. 283, 28454–28463.

103. Moore, R., Cai, K.Q., Escudero, D.O., and Xu, X.-X. (2009). Cell adhesive affinity does not dictate primitive endoderm segregation and positioning during murine embryoid body formation. Genesis 47, 579–589.

104. Foty, R.A. (2008). Quantitative differences in tissue surface tension influence zebrafish germ layer positioning. HFSP J. 2, 42–56.

105. Perret, E., Leung, A., Feracci, H., and Evans, E. (2004). Trans-bonded pairs of E-cadherin exhibit a remarkable hierarchy of mechanical strengths. Proc. Natl. Acad. Sci. USA 101, 16472–16477.

106. Baumgartner, W., Hinterdorfer, P., Ness, W., Raab, A., Vestweber, D., Schindler, H., and Drenckhahn, D. (2000). Cadherin interaction probed by atomic force microscopy. Proc. Natl. Acad. Sci. USA 97, 4005–4010.