Collective cell migration critically depends on cell–cell interactions coupled to a dynamic actin cytoskeleton. Important cell–cell adhesion receptor systems implicated in controlling collective movements include cadherins, immunoglobulin superfamily members (L1CAM, NCAM, ALCAM), Ephrin/Eph receptors, Slit/Robo, connexins and integrins, and an adaptive array of intracellular adapter and signaling proteins. Depending on molecular composition and signaling context, cell–cell junctions adapt their shape and stability, and this gradual junction plasticity enables different types of collective cell movements such as epithelial sheet and cluster migration, branching morphogenesis and sprouting, collective network migration, as well as coordinated individual-cell migration and streaming. Thereby, plasticity of cell–cell junction composition and turnover defines the type of collective movements in epithelial, mesenchymal, neuronal, and immune cells, and defines migration coordination, anchorage, and cell dissociation. We here review cell–cell adhesion systems and their functions in different types of collective cell migration as key regulators of collective plasticity.

Multicellular organisms form and maintain their bodies through the ability of individual cells to adhere to neighbor cells by cell–cell junctions, which are mechanically both stable and flexible and secure cell position and function over time. Stable junctions anchor cells in their tissue niche and define cell–cell cooperation and mechanical function such as contraction or cell–cell signaling. These junctions are the basis of all polarized epithelia, vessels, muscle, neuronal tissue, as well as cell organization in connective tissue. Dynamic cell–cell junctions enable cells to change position relative to their neighbors or as multicellular groups; they are relevant during morphogenesis and phases of tissue activation, for example, in response to injury or inflammation (Collins and Nelson 2015). By regulating junction “fluidity,” the ag-
Aggregate state and dynamics of cells can change remarkably and, accordingly, alter collective functions (Collins and Nelson 2015; Park et al. 2016).

Depending on the cell type and activation state, a range of adhesion receptor and cytoskeletal adaptor systems are involved in securing short- or long-lived, dynamic or stable cell–cell interactions. These include cadherins and protocadherins, immunoglobulin (Ig) superfamily members, desmosomal and tight junction (TJ) proteins, as well as integrins, selectins, ephrin/ephr receptors, and, likely, connexins, which all directly or indirectly couple to the intracellular cytoskeleton and mediate distinct cell–cell adhesion types (Theveneau and Mayor 2012a; Collins and Nelson 2015). Controlled by upstream signaling, each receptor type can undergo context-dependent alteration in surface expression, ligand interaction, and cytoskeletal coupling, and mediate a range of dedicated types of cell–cell coupling.

Many types of collective cell–cell behaviors depend on stable cell–cell anchorage to form layered cell sheets or complex forms of tissue organization, including barrier function mediated by epithelia and endothelia toward the extra- and intracorporal spaces, intercellular signaling network functions as in neuronal networks, or large-scale contraction and force generation as in muscle or purse–string contraction of epithelia (Tada and Heisenberg 2012; Sunyer et al. 2016). Most dynamic multicellular functions, which depend on long-lived cell–cell junctions lasting hours to days or weeks, can be categorized as collective movements in which clusters, sheets, or strands of cells move as a multicellular unit across or through tissue for developing and maintaining epithelial structures (Friedl and Gilmour 2009; Shamir and Ewald 2015). More dynamic cell–cell junctions lasting in the range of minutes are critical in mediating multicellular crowd behaviors in which groups of cells move individually, but coordinate their directionality and speed by less stable and comparably short-lived adhesions and cell–cell sensing (Theveneau and Mayor 2013). Last, immune cells use even more short-lived cell–cell junctions for coordinating their migration and transient clustering with other leukocytes for signal exchange, which depends on very dynamic physical and chemical cell–cell interactions (Malet-Engra et al. 2015).

By combining different adhesion systems in a modular manner in time and space, cells respond to extracellular triggers and tune their levels of cell–cell cooperation. We here summarize the range of cell–cell junction types expressed by different cell types, their morphologies and kinetics, and implications for collective migration, anchorage, and cell dissociation. We further review how different types of cell–cell interaction-based dynamics and collective cell migration are “tunable” and allow for adaptive strategies of cell movements for different physiological and pathological contexts and discuss their implications for classifying collective and single-cell behaviors.

**CELL–CELL ADHESION SYSTEMS**

Common to all adhesion systems is the requirement for an initial interaction between transmembrane cell-surface receptors on adjacent cells, which usually are followed by the recruitment of intracellular adaptor and cytoskeletal proteins. This complex regulates the shape and mechanical stability of the adhesion junction, its interaction with intracellular effectors, and adhesion-mediated activation of downstream signaling pathways. Typically, cells use several complementary adhesion systems in parallel, resulting in a cell–cell interactome (Porterfield and Prescher 2015).

**Adherens junctions (AJs).** AJs are protein complexes found at cell–cell junctions of epithelial and endothelial tissues that connect the actin cytoskeleton of adjacent cells (Shapiro and Weis 2009). AJs depend on the homophilic binding of calcium–dependent cadherins, which interact via their intracellular domains with several regulatory and cytoskeletal proteins such as p120-, α-, β-, γ-catenin, and vinculin, among others (Harris and Tépass 2010). Although AJs are usually associated with epithelial and endothelial tissues, it has been shown that mesenchy-
mal cells form transient adhesion complexes in which type I N-cadherin, together with the full repertoire of intracellular adhesion proteins (p120, α-, β-, γ-catenin, and vinculin), are engaged (Théveneau and Mayor 2012a). Both E- and N-cadherin-based AJs control apicobasal, as well as front–rear, polarity of interacting cells (Venhuizen and Zegers 2016).

The main functional difference between epithelial and mesenchymal AJs is their stability: epithelial junctions tend to be more stable (in the range of hours to days), whereas mesenchymal junctions are transient (minutes to hours) (Scarpa et al. 2015). The stability of AJs is controlled by several mechanisms, including endocytosis and cytoskeletal regulation. Endocytosis of AJ receptors and adapters occurs both by clathrin-dependent and -independent mechanisms (Delva and Kowalczyk 2009; Schill and Anderson 2009), which cooperate with regulation by Rho family GTPases. For example, Cdc42 works upstream of Par6/αPKC and Cdc42-interacting protein 4 (CIP4), which control actin dynamics at the internalization site (Harris and Tépass 2010). Besides controlling the stability of cell–cell interactions, Rho GTPases, via PAK and βPIX, are reciprocally controlled by AJs in which they play an essential role on actin dynamics (Zegers et al. 2003; Zegers and Friedl 2014). Interaction between cadherin–catenin clusters leads to the recruitment of the Rac guanine nucleotide exchange factor (GEF) TIAM1, which activates the Rho GTPase Rac1, and the activation of Rac1 in leader cells is, in turn, required for the formation of cell protrusions and traction forces observed at the edge of a cell cluster during collective cell migration (Hordijk et al. 1997; Kovacs et al. 2002; Mertens et al. 2005). Another activator of Rho GTPases within the AJ is Nectin, and Nectin-like proteins, a family of Ig-like cell adhesion molecules (CAMs) (Takai et al. 2008). To aid the formation of AJ, nectin recruits afadin and ponsin, which lead to the activation of Cdc42 and Rac and cytoskeletal remodeling at the site of cell–cell contact (Fukuyama et al. 2006). The interaction between AJs and actin is mutual, leading to an increase in the stability of cortical actin toward the maturing AJ complex (Baum and Georgiou 2011). Consequently, AJ are central hubs controlling cell–cell cohesion and collective cell migration underlying tissue dynamics and remodeling.

**Tight junctions (TJs).** TJs are adhesion complexes in which the plasma membranes of adjacent cells become closely associated, forming an impermeable barrier within the tissue. TJs are indispensable for creating a barrier between different regions of the body, and their main role is to function as paracellular gates that restrict diffusion on the basis of size and charge. TJs are composed of transmembrane proteins (claudin, occludin, tricellulin, and marveld3) that seem sufficient to trigger at least some of the aspects required in TJ formation, including mechanical junction stability and apicobasal polarity of connected cells (Zihni et al. 2016). Other TJ transmembrane adhesion proteins comprise the junctional adhesion molecules (JAMs), which enhance TJ stability and turnover (Ebnet et al. 2004; Luissint et al. 2014). The intracellular function of TJs depends on a dense network of proteins, composed of ZO1, ZO2, ZO3, plus a large number of other adaptor proteins (Van Itallie and Anderson 2014). By binding several transmembrane and adaptor proteins, TJs control various signaling pathways involved in actin organization, cell polarity, as well as transcriptional regulation (Gonzalez-Mariscal et al. 2014). The interaction of TJ proteins with the actin cytoskeleton seems to be essential for junction formation and turnover. For example, myosin light chain kinase stimulates TJ remodeling and occludin internalization during inflammation (Herrmann and Turner 2016). Rho GTPase signaling is also controlled by TJ-associated proteins: RhoA, Cdc42, and Rac are regulated by GEFs recruited to cingulin, ZO1, and tricellulin (Otani et al. 2006; Terry et al. 2011; Oda et al. 2014). Thereby, TJs form a central hub between cell–cell interactions and actin dynamics (Balda and Matter 2016).

**Gap junctions (GJs).** GJs are intercellular membrane channels made up of a multigene family, called connexins in vertebrates (Willecke et al. 2002). GJs are specialized junctions char-
characterized by close apposition of the plasma membranes between neighboring cells and contain a hydrophilic channel that mediates the intercellular passage of molecules >1 kDa in size. The extracellular domains of connexins form a tight connection between adjacent cells contributing to cell-cell adhesion. Connexins interact with several proteins to form multiprotein complexes, which are important in cell-cell junction stability and function. For example, Cx43 interacts with N-cadherin and other members of the AJ complex (Xu et al. 2001), as well as cytoskeletal proteins such as microfilaments and microtubules (Wei et al. 2004). Phosphorylation of the cytoplasmic domain of connexin is critical in regulating GJ assembly, trafficking, channel gating, and turnover. GJs contribute to cell migration during development and in homeostatic processes such as wound healing (Kotini and Mayor 2015), and it has been proposed that their channel activity could be important for cell coupling and coordination during migration (Lorraine et al. 2015).

**IgCAMs.** IgCAMs correspond to immunoglobulin-like cell-adhesion molecules containing one or more Ig-like domains in their extracellular regions. IgCAMs are expressed in a wide variety of cell types, such as cells of the nervous system, leukocytes, and epithelial and endothelial cells (Cavallaro and Christofori 2004). By homophilic and heterophilic interactions of their Ig-like domains IgCAMs mediate adaptive cell–cell interactions and play an important role in cell migration (Cavallaro and Christofori 2004). IgCAM adhesion is regulated by lateral oligomerization, which in turn depends on phosphorylation of their Ankyrin-binding domain (Garver et al. 1997). A second mechanism that controls IgCAMs-mediated adhesion is based on their internalization or recycling from the plasma membrane; for example, the internalization of aplysia cell adhesion molecule (apCAM) is controlled by phosphorylation by mitogen-associated protein (MAP) kinase (Bailey et al. 1997). A third mechanism that regulates IgCAM-based cell adhesion is their proteolytic cleavage. For example, the leukocyte adhesion molecule L-selectin is cleaved by sheddases of the metalloprotease and ADAM families, and is protected from this cleavage by intracellular regulators, which engage with its cytoplasmic domain, including calmodulin and moesin (Kahn et al. 1998; Ivetic et al. 2002). IgCAMs have been reported to associate with a range of other proteins at the cell membrane, including growth-factor receptors, integrins, and cadherins, and with intracellular proteins, such as effectors of signal transduction pathways and cytoskeletal proteins (Juliano 2002), and thus contribute to a range of signaling programs involved in cell adhesion and migration.

**Slit/Robo.** Slit/Robo corresponds to the Roundabout receptors (Robo) and their Slit ligand. Robo receptors belong to the superfamily of IgCAMs and engage in both homophilic and heterophilic interactions (Hivert et al. 2002). Slits are the principal ligands for the Robo receptors (Kidd et al. 1999), which, together with heparan sulphates, form a ternary complex required for signaling (Ypsilanti et al. 2010). The cytoplasmic domains of Robo do not possess catalytic activities and, therefore, interact with different signaling molecules to exert their specific effects; these include netrin and several GTPase activating proteins (GAPs) and GEFs that control actin cytoskeletal dynamics by regulating the activity of RhoA, Rac1, and Cdc42 (Ypsilanti et al. 2010). The activity of Slit/Robo, including adhesion, is controlled by transcriptional regulation and endocytosis and degradation (Keleman et al. 2005). Slit–Robo interaction typically mediates cell repulsion, but in some cases also supports cell–cell adhesion. The formation of cranial ganglia requires the adhesion to different cell types, and increased adhesion between neural crest and placodes is promoted by an interaction between Robo2/Slit1, which increases the N-cadherin-dependent adhesion between these cells (Shiau and Bronner-Fraser 2009). Slit/Robo interactions are also involved in collective cell migration of neural crest cells during development and endothelial cells in angiogenesis (Legg et al. 2008).
**Ephrin/Eph receptor.** Ephrin/Eph receptor corresponds to a pair of ligands and receptors involved in short-distance cell–cell signaling. Eph are Tyr kinase receptors and ephrins are membrane-tethered ligands, which can elicit signaling that affects the cytoskeleton, mediating primarily cell repulsion but, depending on context, also cell–cell adhesion (Kania and Klein 2016). Phosphorylation of the intracellular domains of Ephs regulates the recruitment of effector proteins, such as the noncatalytic regions of Tyr kinase adaptor protein 1 (Nck1) and Nck2, Vav2 and Vav3, Src, α2-chimerin, and ephexins, which directly regulate actin or modulate the activity of the small GTPases RhoA and Rac1 (Kania and Klein 2016). Normal morphogenesis of the neural tube and neural progenitors requires ephrin-dependent cell–cell adhesion (Arvanitis et al. 2013), and alternative usage of different splice forms of Eph receptor was implicated in mediating cell–cell coupling during embryonic development. Eph signaling promotes the formation of AJ through interaction with E-cadherin and TJs via the interaction with claudin (Dravis and Henkemeyer 2011). Likely, the consequences of Ephrin/Eph interactions for cell–cell contact stability depend on the overall junction protein repertoire expressed by the cell. Ephrin/Eph has been shown to be important for collective movement; for example, the formation of the thymus anlage requires EphB/ephrin B, which seems to support collective mobility by a collective separation mechanism (Foster et al. 2010).

**Integrins.** Integrins are transmembrane proteins that connect the cytoskeleton with the extracellular matrix (ECM). ECM ligands for integrins include fibronectin, vitronectin, collagen, and laminin, among others, which, beyond their well-established function as structural connective tissue scaffolds, also may be located between cells and contribute to cell–cell interactions (Barczyk et al. 2010). Integrins interact with F-actin and intermediate filaments allowing a mechanical coupling between the cytoskeleton and the ECM, and act as important transducers of mechanical forces (Fagerholm et al. 2005). Integrin engagement results in the formation of focal adhesion complexes of varying sizes and functions, which interact with F-actin and recruit FAK and Src, leading to the activation of signaling pathways involving extracellular signal–regulated kinase (ERK), c-Jun N-terminal kinase (JNK), and small GTPases (Bouvard et al. 2013). Interaction of integrins with cadherins and selectins has been proposed to be required for the participation of integrins in cell–cell adhesion (Bouvard et al. 2013).

**CELL–CELL ADHESION STATES AND DYNAMICS**

The type and durability of cell–cell adhesion and cytoskeletal interaction systems that are engaged by stationary and moving cells provide a range of adhesion strategies between cells, which jointly define the level of collective adhesion and polarity, junction dynamics, and the type of collective migration. The spectrum of tissue fluidity can be found to vary, in a cell- and context-dependent manner, from fully immobilized, highly contractile to loosely connected but highly mobile collective cell–cell organizations and kinetics (Fig. 1).

**Myoblast fusion and myofiber formation.** Myofibers are multicellular syncytia that develop by the fusion of individual myoblasts. Rather than forming a collectively migrating group, myoblasts remain stably anchored to the substrate while establishing stable cell–cell junctions that enable contractility across many cells but show little junction dynamics. Myoblast interactions engage multiple receptor systems in parallel, including focalized high-density accumulation of M- and N-cadherin, neural cell adhesion molecule (NCAM), vascular cell adhesion molecule (VCAM-1), meltrin, and integrins (Fig. 1A) (Charrasse et al. 2006; Abmayr and Pavlath 2012; Ozawa 2015). Once myoblasts connect with each other, individual mobility is largely disabled, whereas collective contractility and force transmission across cell–cell junctions are gained, particularly through the actomyosin cytoskeleton, which develops prominent stress fibers under the
**Figure 1.** Composition of cell–cell adhesions and related types of collective cell behaviors. Morphological organization of cell–cell interactions, related collective cell dynamics (upper panels), and their respective stability (cyan), molecular composition (blue), and multicellular function outcomes (pink; bottom panels). Different examples show variable levels of cell adhesion (blue dots) and cell movement (arrows, direction of migration). (A) Myofiber network. (B) Gut epithelium renewing. (C) Branching morphogenesis. (D) *Drosophila* border cells. (E) Neural crest. (F) Leukocytes. IgCAMs, immunoglobulin-like cell-adhesion molecules; Robo, roundabout receptors.
control of RhoA and bridges multiple cell bodies for coordinated rhythmic contractility of the multicellular ensemble (Charrasse et al. 2006). Thus, in fusing myoblasts high junction stability mediated through overlapping adhesion systems and cytoskeletal linkages support mechanically very stable junctions, which mediate collective contractility and eventually cell fusion, but discourage position change of the group as a whole and individual cells within the group.

**Epithelial sheet migration.** Mature monolayered epithelia display stable cell–cell interactions, established by E-cadherin- and β-catenin-based AJs, combined with apical desmosomal junctions and TJs; these jointly mediate mechanically robust multicellular integrity, apicobasal polarity, and barrier function (Wong et al. 1998; Takeichi 2014). Whereas the epithelium as tissue remains anatomically stable to sustain live-long mechanical and functional integrity, epithelial renewal in several tissues, including the intestine and the epidermis, depends on constitutive collective sheet migration coupled to cell proliferation and sheet expansion (Wong et al. 1998; Nanba et al. 2015). The renewal of the gut epithelium is initiated by releasing precursor cells from the stem cell pool, which resides at the bases of the crypts, and that then change position and move upward along the basement membrane toward the tips of the villi (Ritsma et al. 2014). Because moving cells remain fully coupled to their neighbors by lateral junctions and the intestinal basement membrane via their basal plane, they move as cohesive sheets in the upward direction (Fig. 1B) and, additionally, undergo a controlled number of cell divisions (Wong et al. 1998; Nanba et al. 2015). The mechanics of lateral sheet migration is not fully resolved. Likely, cryptic lamellipodia generate traction force toward the substrate along the sheet, whereas the apical cell–cell junctions transmit collective actomyosin contractility to enable slow movement along the basement membrane (Farooqui and Fenteany 2005; Zegers and Friedl 2014; Bazellieres et al. 2015), but the role of additional rotational and turbulent movements remains to be clarified (Nanba et al. 2015). By coupling apicobasal polarity with high junction stability, sheet migration along a two-dimensional (2D) substrate layer as guidance cue is a conserved and important type of collective migration of a mature epithelium. Accelerated variants of epithelial sheet migration are observed as sheet migration during the wound closure of epithelial defects and epithelial morphogenesis.

**Sprouting strands.** When invading 3D tissues, epithelial and endothelial cells typically move collectively to form linear, branched, or network-like strands (Fig. 1C). Collective sprouting underlies the branching morphogenesis of epithelial tissue and organs with branched or lobular structure, including the trachea, kidney, thymus, and the mammary gland (Gray et al. 2010). Sprouting hemo- or lymphangiogenesis occurs during revascularization of tissue after injury (Senger and Davis 2011). As a biomechanical principle of sprouting in differentiated endothelia and epithelia, one or several leader cells form a leading tip or terminal end bud that protrudes forward, whereas the rear cells undergo apicobasal polarization to form a vascular lumen or duct (Huebner et al. 2016). The cell–cell junctions of invading epithelial strands are dependent on E-cadherin-based AJs, desmosomal, and TJs (Shamir and Ewald 2015), whereas vascular strands depend on VE-cadherin and TJs (Senger and Davis 2011). Commonly, these cell strands preserve apicobasal polarity, form lumen, and deposit an abluminal basement membrane as they move, as has been observed during vascular, mammary gland, kidney, and tracheal development (Riggins et al. 2010; Nguyen-Ngoc et al. 2012). When apicobasal polarity is lacking, as in dedifferentiated tumor cells, collective strand invasion occurs as solid finger-like multicellular protrusions with no lumen formed (Wolf et al. 2007; Nguyen-Ngoc et al. 2012).

**Moving sheets and clusters.** In morphogenesis and during cancer invasion, cell sheets and
detached groups of variable size, which retain cohesive cell–cell junctions between cells, migrate along 2D and three-dimensional (3D) tissue scaffolds (Fig. 1D) (Friedl et al. 1995; Alexander et al. 2008; Montell et al. 2012). Epithelial sheet movement is initiated by a row of leader cells coupled to follower cells by AJs containing E- or P-cadherin (Chapnick and Liu 2014; Plutoni et al. 2016), and sheet displacement depends on coordinated traction force generation between leader and follower cells, which are distributed across cell–cell junctions by the actomyosin cytoskeleton (Brugues et al. 2014; Reffay et al. 2014; Bazellieres et al. 2015). Moving clusters can be epithelial, such as the border cells moving along the boundaries of large nurse cells of the developing Drosophila ovary, or mesenchymal, such as moving neoplastic sheets in rhabdomyosarcoma explant culture (Friedl et al. 1995). The activity of leader cells depends on extracellular stimuli, such as ECM ligand or soluble factors (e.g., transforming growth factor β, TGF-β), inducing MAP kinase signaling and downstream Rac1 for protrusion formation and direction sensing (Khalil and Friedl 2010; Chapnick and Liu 2014). Leader cell polarity is further supported by AJ signaling, which controls leader cell polarization and anterior protrusion dynamics (Khalil and Friedl 2010; Mayor and Etienne-Manneville 2016). In contrast to homeostatic sheet migration, which forms a continuum without leading and trailing edges, the mechanisms defining retraction of the rear cells in moving clusters remain unclear.

Moving cell networks. A more flexible type of collective migration, used by neural crest and other mesenchymal cells, as they move in a coordinated manner as loosely cohesive group in a cell-type and context-dependent way, with a variable tendency to individualize (Fig. 1E) (Scarpa et al. 2015). Examples are the migration of neural crest cells in developing embryos, neuronal/astrocyte networks (Scarpa et al. 2015), glioma cells retaining filamentous cell–cell interactions while moving through complex brain parenchyma (Osswald et al. 2015), and mesenchymal tumor cells moving through confining tissue (Ilina et al. 2011; Haeger et al. 2014). Collectively, moving loose networks are mediated by complex morphological junctions mediated by N-cadherin for cell–cell adhesion and additional receptors mediating repulsive signals, including Ephrin/Eph receptors (Theveneau and Mayor 2011). As a consequence, cells coordinate their polarity and respond to extracellular signals as a group, but also retain the remarkable ability of moving individually. Mesenchymal tumor cells develop ALCAM-positive cell–cell junctions when moving through confined space, and gain many properties of collective invasion, including shared migration path, lateral cell–cell junctions, and alignment of front–rear polarity and mitotic planes (Haeger et al. 2014). But they also retain the ability to rapidly detach in response to extracellular signals such as growth factors, altered tissue geometry, or matrix metalloproteinase (MMP) availability (Wolf et al. 2007; Ilina et al. 2011). Conceptually, collective networks retain properties of both collective and individual cell movements, as well as multicellular streaming, and further show a high level of stochasticity (“noise”) in switching between individual and collective behaviors, which render experimental classification sometimes challenging and requires quantitative analysis as well as mathematical modeling (Huang et al. 2015). As a defining characteristic for collective cooperativity, the cells moving as a network can respond more efficiently to external signals when they are part of groups rather than as individual cells (Theveneau et al. 2010).

Leukocyte swarming and aggregation. When activated, moving leukocytes show a strong tendency to interact with each other, coordinate their migration for “swarming” behaviors, and aggregate. In activated lymphoblasts, αLβ2 integrin/ICAM-1–dependent cell–cell junctions transiently interact and coordinate their migration as cell pairs or small clusters of variable stability (Gunzer et al. 2004; Malet-Engra et al. 2015). As moving myeloid leukocytes converging toward damaged or infected tissue regions, chemoattractant guidance leads to highly coordinated crowd behaviors with head-to-tail ori-
entation and frequent cell–cell interactions, which can eventually transit to firm clustering that depends on β2 integrin availability (Waite et al. 2011; Lammermann et al. 2013). Thus, at the low end of cell–cell adhesion, individually moving leukocytes may coordinate their amoeboid movements with neighboring cells by short-lived cell–cell junctions (seconds to minutes), and rapidly transit toward contact strengthening and aggregation into a multicellular cluster. Depending on experimental context, such clusters may be immobile and transiently close an interstitial wound (Lammermann et al. 2013) or, under chemotactic and free-space conditions, even show collective coordination and migration (Malet-Engra et al. 2015).

In summary, for very different cell types and biological contexts, the organization and stability of cell–cell junctions, together with environmental parameters, determines the morphological and functional types of collective movement. Consequently, alteration of adhesion and environmental parameters may impose an adaptation response and significant plasticity of collective behaviors.

TUNING COLLECTIVE MIGRATION: VARIABILITY AND TRANSITIONS

The molecular variability of cell–cell junctions and their different linkages to the cytoskeleton not only explain distinct types of collective movements; when regulated within the same cell type, by extracellular chemical or physical signals, collective migration may be induced or modulated, with consequences for group behavior and tissue integrity and function.

Tissue Morphogenesis and Regeneration

Morphogenesis involves the complex and coordinated rearrangement of tissues and massive movement of cells. In particular, the initial formation of the body shape, the early separation of the principal tissue types, and the organization of specific organs depend on different types of collective cell migration. Similar morphogenetic processes are implicated in regeneration.

A fundamental morphogenetic process that allows tissues to develop and remodel and that depends on the regulation of cell–cell adhesions is the epithelial-to-mesenchymal transition (EMT). In response to extracellular triggers, including cytokines, growth factors, and metabolic stress, EMT induces the down-regulation of E-cadherin but up-regulates N-cadherin, which lowers cell–cell adhesion strength and apical-basal cell polarity, and favors migratory and invasive properties (Thiery et al. 2009). Traditionally, full EMT has been considered essential for the migration as it was thought that only mesenchymal, but not epithelial cells, were able to migrate. However, recent work indicates that EMT should be considered as a spectrum of intermediary phases, ranging from full EMT to partial, or even quite subtle states of EMT with very variable effects on cell mobility and migration type (Fig. 2) (Nieto et al. 2016). Well-studied examples of collective cell migration during development, which are initiated by or maintain some degree of EMT, include the migration of border cells in *Drosophila*, the lateral line primordium in zebrafish, and the neural crest in amphibian and zebrafish (Mayor and Etienne-Manneville 2016). The precursors of all these tissues are epithelial cells that undergo an EMT to initiate their migration; the degree and pattern of collective cooperation during migration thereafter reflects varying cell–cell junction organization and dynamics maintained by varying coexistence of epithelial and mesenchymal programs (Fig. 2). The zebrafish lateral line primordium represents collective cell migration, which retains both epithelial and mesenchymal characteristics as a default state (Fig. 2A,B) (Ghysen and Dambly-Chaudiere 2004). Cells of the primordium express E-cadherin and show foci of the TJ protein ZO1 and aPKC at the center of the tightly packaged primordium (Fig. 2A,B) (Ghysen and Dambly-Chaudiere 2004). Cells of the primordium express E-cadherin and show foci of the TJ protein ZO1 and aPKC at the center of the tightly packaged primordium (Revenu et al. 2014), similar to mature epithelium. On the other hand, cells particularly located at the edge of this cell group display typical mesenchymal characteristics, such as reduced apicobasal polarity and the presence of highly dynamic actin-rich lamellipodia-like protrusions at the basal interactions to the tissue (Lecaudey et al. 2008; Hava et al.
Thus, within the same moving cell group, different interaction types are found, with strong cell–cell adhesions at the center of the migrating cluster and more mesenchymal cells with weaker adhesions but higher actin-mediated mobility at the periphery and in leading cells.

Branching morphogenesis in the lactating mammary gland and cell movements during gut homeostasis are also seemingly equivalent to the initial branching morphogenesis observed during development, as both processes require E-cadherin-mediated cell–cell cohesion for collective sprouting and tubule elongation; depletion of E-cadherin interferes with the integrity of these tissues, with E-cadherin deficient cells being excluded (Shamir and Ewald 2015). The leading front of the tube, which drives collective mobility, undergoes a loosening (but not ablation) of cell junction stability, for example, by maintaining a partial EMT in which E-cadherin-based junctions gain flexibility and increase their turnover; concurrently cells in the rear position, which form the extending tube, retain stable cell–cell junctions and apico basal polarity, and gradually downscale their migration ability (Shamir and Ewald 2015). Likely, similar reprogramming of leader cells toward loosened junction organization is active during neoplastic invasion of
breast cancer cells (Cheung et al. 2013; Cheung and Ewald 2014). Thus, tuning cell–cell junction stability regulates the degree of collective dynamics.

As an example of a very transient EMT followed by epithelial collective migration, Drosophila border cells initiate their delamination from the epithelium by down-regulation of DE-cadherin for a short time period (few hours) (Fig. 2C) (Montell et al. 2012). A network of transcription factors, including Jing, SIX4, Yan, Similar (also known as HIF1α), Hindsight (HNT), and Jun-related antigen, is activated and controls the levels of DE-cadherin during border cell delamination and migration (Montell et al. 2012). Levels of DE-cadherin need to be precisely regulated and deviations impair border cell migration; consequently, after initially reducing DE-cadherin levels during delamination, moving border cell clusters still maintain substantial levels of both DE-cadherin and its binding partner armadillo (β-catenin) between neighboring border cells to maintain collective migration as a cluster (Peifer 1993; Niewiadomska et al. 1999; Sarpal et al. 2012).

As an example for even stronger mesenchymal properties with few epithelial features retained, in many species the neural crest undergoes EMT to initiate migration (Thiery et al. 2009; Theveneau and Mayor 2012b). In addition, after delamination a collective network-type migration is retained whereby loosely connected individually moving cells and more tightly connected clustered cells both depend on cell–cell interactions for their migration (Fig. 2D) (Teddy and Kulesa 2004; Carmona-Fontaine et al. 2008). A complex genetic network is activated in the neural crest that leads to and maintains EMT. A BMP4-Wnt1 signaling pathway activates a set of transcription factors, including Snail1/2, Sox5/9/10, Foxd3 and Ets1, that modify the expression of cell–cell and cell–matrix adhesion molecules (Sauka-Spengler and Bronner-Fraser 2008; Theveneau and Mayor 2012c). Thus, although neural crest shows all the hallmarks of mesenchymal cells, they obviously form transient AJs (Scarpa et al. 2015). The endocytosis of N-cadherin at the AJs is essential for neural crest migration in vivo, as it confers sufficient fluidity on the cell cluster for it to migrate under physical constrains (Kuriyama et al. 2014). This increase in tissue fluidity allows a dynamic exchange of neighbors while retaining cell–cell interaction.

In conclusion, collective cell migration during morphogenesis involves a wide spectrum of cell adhesion strength, with highly cohesive cell sheets and clusters at one end and relatively loose groups of cells at the other end. By spatially and temporally tuning cell–cell junction stability, a range of collective migration modes and patterns with different levels of cell–cell cohesiveness is achieved to build tissue.

Cancer Invasion and Metastasis

Collective invasion is an important strategy for local tissue infiltration, as well as metastatic evasion in epithelial tumors such as breast cancer, squamous cell carcinoma, colon cancer, and others, as well as in mesenchymal tumors (Ilina and Friedl 2009; Cheung and Ewald 2016). Similar to morphogenesis, the phenotypic and junctional organization of moving cancer cell groups varies greatly (“collective plasticity”). In experimental live-cell models, all types of collective movements can be adopted by tumor cells including (1) cohesive sheets or strands, typically detected in epithelial cancers; (2) isolated clusters detached from the primary/metastatic lesion such as epithelial tumors and melanoma; (3) neuronal-like networks of connected cells, detected in neuroectodermal tumors, such as glioblastoma; or (4) as “jammed” collective cohorts induced by spatially narrow tissue boundaries (confinement) of otherwise transiently/loosely connected (single) cells in experimental melanoma and sarcoma models (Friedl et al. 1995, 2012; Nguyen-Ngoc and Ewald 2013). Similarly, histological examination of both patient lesions and mouse models in vivo shows that the collective invasion patterns develop striking morphological and molecular variability, depending on tumor type and the tissue that is invaded (Weigelin et al. 2012; Bronsert et al. 2014).

Consistent with cellular diversity of collective invasion programs, a range of cell–cell
adhesion mechanisms supports collective invasion of cancer cells. Epithelial tumors invade collectively, with duct-like patterns and E-cadherin and β-catenin positive cell–cell junctions, with or without lumen formation, and with or without up-regulation of EMT markers, including Twist and Zeb1 (Cheung et al. 2013; Bronsert et al. 2014). Furthermore, both E- and N-cadherin can orchestrate AJs and cell–cell interactions in cancer cells (Bronsert et al. 2014; Zucchini et al. 2014). Besides cadherins, Ig superfamily members and ephrins/EpH receptor systems were implicated in mediating more labile or transient cell–cell interactions in cancer cells (Cavallaro and Christofori 2004; Haeger et al. 2014; Krusche et al. 2016). As well, connexins may enable communication through GJs between connected tumor cells and their inhibition reduces collective migration in prostate cancer cells (Zhang et al. 2015). Similar to morphogenetic and homeostatic migration, collectives of migrating cancer cells display leader cells that engage with surrounding tissue structures via Rac-driven filopodal protrusions and integrin-mediated substrate adhesion (Hegerfeldt et al. 2002; Yamaguchi et al. 2015). Collectively moving cancer cells retain a range of actin dynamics, substrate adhesions, and ECM remodeling functions, which typically are shared and coordinated between neighboring cells, generate tissue alignment and remodeling as an integrated process; these combined parameters can further define the degree of cell–cell cohesion and individualization as an integrated function of cell–cell junction stability, MMP activity and tissue organization, and space (Scott et al. 2010; Friedl et al. 2012; Te Boekhorst et al. 2016).

In recapitulation of morphogenesis programs, EMT signaling enhances cancer invasion and metastatic progression of epithelial cancers by reprogramming cell–cell junctions (Kalluri and Weinberg 2009). EMT weakens or fully dissolves cell–cell junctions between cancer cells, including AJs, desmosomes, and TJ’s. EMT also up-regulates the expression of stromal proteases, which cleave cadherins; deregulates integrin adhesion systems, for example, by switching β1 to β3 integrin expression and enhancing αV integrin signaling; and can redirect Rho-mediated actomyosin contractility from cell–cell junctions toward cell–matrix interactions (Kalluri and Weinberg 2009; Parvani et al. 2013; Truong et al. 2014). These molecular reprogramming events result in deregulated cell–cell contacts, loss of apicobasal polarity, including degeneration of the lumen of otherwise ductal and glandular structures, gain of front–rear polarity, and ultimately favor the gradual transition from epithelial to mesenchymal migration modes (Bryant et al. 2014).

In addition to tumor cell individualization caused by full EMT, which allows for single-cell dissemination and metastasis, recent cell-based and modeling work indicates that EMT also contributes to collective invasion with a high likelihood of mixed behaviors after EMT, including intermediate (e.g., metastable or hybrid) phenotypes such as detached collective or loosely connected migrating groups (Jolly et al. 2015). With such EMT-associated reprogramming, or partial EMT, moving tumor cell clusters may still maintain cell–cell contacts but simultaneously undergo a differentiation switch toward embryonic features (Jolly et al. 2015; Nieto et al. 2016). Thus, similar to morphogenesis, the adaptability of collective invasion programs allows a range of coping strategies for cancer invasion and metastasis in different tissue environments (discussed in Te Boekhorst and Friedl 2016).

CONCLUDING REMARKS

Cell–cell junctions emerge as central regulators of the type, efficiency, and fate of collective cell movements. Here, we have proposed an integrative view to frame collective cell functions that is distinguished by modular and often gradual cell–cell adhesion regulation. We hope that this perspective may facilitate the understanding of multicellular dynamics with mixed phenotypes, which are frequently observed in wet-laboratory experiments and also in mathematical modeling (Jolly et al. 2015; Te Boekhorst et al. 2016). By modulating the composition of cell–cell adhesions, collective movements are adaptive in time and space in response to soluble and
structural tissue-derived signals, as well as geometric tissue properties (Haeger et al. 2015; Te Boekhorst et al. 2016). Thus, similar to single-cell migration modes, which can switch between different types of mesenchymal and amoeboid movements, collective migration modes can interconvert and adapt to local and global signals.

Based on their range of stability, cell–cell junctions may be classified as (1) stable, cohesive with disabled cell position change, (2) stable, but dynamic, allowing cells to move relative to neighbors without resolving the junction, (3) partially stable, allowing transient detachment and reintegration, and (4) short lived and partially repulsive. Emergent collective behaviors, that is, the ability of a cell group to perform tasks beyond the abilities of a single cell, are reached as long as cell–cell junctions suffice to coordinate behavior across scale. Examples are collective chemotaxis and durotaxis, which allow cell groups to respond to more shallow chemical or physical gradients for directional migration (Malet-Engra et al. 2015; Sunyer et al. 2016). The gradual range of kinetic states complicates simple classifications as “collective” movement versus multicellular streaming versus predominantly single-cell dynamics. Typical collective cell migration is considered when stable cell–cell junctions support supracellular coordination of cytoskeletal activity across multiple cell bodies and even passive cell transport as part of group behaviors (Friedl et al. 2012). Likewise, emergent collective behaviors can also be observed when cell–cell junctions are transient junctions, particularly in chemotactic gradient sensing and signal integration. Thus, multicellular streaming behavior depends on the active movement of every cell but still retains multiple reciprocal cell–cell interactions, which are limited in duration and stability but enable collective gradient sensing (Theveneau et al. 2010; Ellison et al. 2016). Last, individually moving cells may still engage with other cells in short-lived junctions, which may or may not induce repulsion and directional change, and thereby retain cooperative input from neighboring cells (Ellison et al. 2016). As a special case, an otherwise loosely connected cell may upscale cooperation as part of, for example, a cell-jamming transition when cells are confined in tissue space, and thereby adopt emergent behaviors, including persistent intercellular adhesion and signal integration (Haeger et al. 2014; Sarkar et al. 2016). Thus, different junction states and environmental conditions enable unique sets of emergent mechanical and signal integration beyond single cell behavior. Future classification of different types of collective versus single-cell behaviors will depend on careful dissection of each functional module associated in a junction-, cell-type-, and tissue-specific context.

Conceptual frameworks based on classifying types of collective movements and their links to single-cell migration and other types of tissue dynamics, such as tissue folding and convergent extension, will further allow us to integrate molecular signaling concepts, for example, on EMT or stemness, with varying degrees of cell–cell cooperation. Based on their central function in defining the shape, molecular composition, and duration of cell–cell cooperation, multiscale analysis integrating simultaneously engaged junction mechanisms and their signaling cross talk will be required to delineate which individual and cooperative functions guide or compromise collective decision-making and outcome.

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