The Importance of Cell–Cell Interaction Dynamics in Bottom-Up Tissue Engineering: Concepts of Colloidal Self-Assembly in the Fabrication of Multicellular Architectures

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Supporting Information

ABSTRACT: Building tissue from cells as the basic building block based on principles of self-assembly is a challenging and promising approach. Understanding how far principles of self-assembly and self-sorting known for colloidal particles apply to cells remains unanswered. In this study, we demonstrate that not just controlling the cell–cell interactions but also their dynamics is a crucial factor that determines the formed multicellular structure, using photoswitchable interactions between cells that are activated with blue light and reverse in the dark. Tuning dynamics of the cell–cell interactions by pulsed light activation results in multicellular architectures with different sizes and shapes. When the interactions between cells are dynamic, compact and round multicellular clusters form under thermodynamic control, while otherwise branched and loose aggregates under kinetic control assemble. These structures parallel what is known for colloidal assemblies under reaction- and diffusion-limited cluster aggregation, respectively. Similarly, dynamic interactions between cells are essential for cells to self-sort into distinct groups. Using four different cell types, which expressed two orthogonal cell–cell interaction pairs, the cells sorted into two separate assemblies. Bringing concepts of colloidal self-assembly to bottom-up tissue engineering provides a new theoretical framework and will help in the design of more predictable tissue-like structures.

KEYWORDS: Bottom-up tissue engineering, optogenetics, cell–cell interactions, self-assembly, self-sorting, photoswitchable

The bottom-up assembly of a spatially ordered tissue made from cellular building blocks based on the principles of self-assembly is a highly promising and powerful approach to tissue engineering and an extreme synthetic and biological challenge at the same time. Building multicellular structures requires more than simply putting together a solution with the right composition of cells; it requires specific interactions between the cells and spatial organization of these building blocks into hierarchical structures, which determines how cells work together as a tissue. The bottom-up approach to tissue assembly parallels observations seen during tissue formation in biology, where no template or scaffold is needed and cell–cell interactions are a major driving force that determines their organization. Remarkably, even dissociated cells from different tissues are able to self-aggregate and self-sort again into multicellular structures that resemble their tissues of origin. Moreover, increasing possibilities in organoid and stem cell cultures as well as programmable multicellular structures with synthetic cell to cell signaling speaks for the massive potential of the living cells to self-organize into complex functional architectures and controlling them using synthetic biology.

Going forward, it is indispensable to understand how cells as the basic building blocks of tissue self-assembly. This requires controlling the interactions between cellular building blocks and understanding to what extent the principles of self-assembly and self-sorting known for nonliving colloidal particles apply to cells. Such insight would allow us to build up multicellular architectures with predictable and programmable organization, understand the limits of multicellular structures that can be generated solely based on self-assembly and where further biological signals are required. As model building blocks for materials, colloids provide a valuable framework for the self-assembly of micron-sized objects such as the cells. For colloidal systems, the interactions between colloids are the major driving force behind self-assembly, and different architectures can self-assemble depending on the kinetic and thermodynamic parameters for the interactions between the colloids. While compact and spherical structures at the thermodynamic equilibrium form under reaction-limited cluster aggregation (RLCA), loose and ramified assemblies in kinetically trapped states form under diffusion-limited cluster aggregation (DLCA). This puts forward the importance of not just controlling the interactions between the colloidal/cellular building blocks but also their dynamics. Cell–cell
interactions have been controlled by modifying the surfaces of cells with complementary DNA strands,15 biotin–streptavidin,16 clickable functional groups,17,18 supramolecular interaction partners,18,19 and the photoswitchable protein pair CRY2/CIBN.20 Yet, except for the last two methods, these interactions are not reversible and provide no control over the dynamics of the cell–cell interactions. Moreover, the important role of the cell–cell interaction dynamics for the final multicellular structure has not been considered in any of these studies. This is in contrast to native cadherin-based cell–cell adhesions, which have fast exchange rates and form thermodynamically controlled multicellular structures.21 Therefore, the question of what kind of tissue structures can be generated by employing only the principles of self-assembly and controlling cell–cell interactions to achieve diffusion or reaction-limited assembly remains unanswered.

Another concept where the principles of colloids and cells connect to one another is their self-sorting/sorting-out behavior in multicomponent mixtures.2,23 Observations in in vivo and in vitro multicellular systems led to the differential adhesion hypothesis, which postulates that, if two populations of cells are mixed, the cells sort-out to reach a final organization that approaches a state with a minimal internal free energy and maximum total cell–cell interactions.24 Such self-sorting under thermodynamic control is only possible provided that the cell–cell interactions are dynamic, and this criterion is indeed satisfied for native cadherin-based cell–cell interactions.27 Consequently, in mixtures of dissociated cells that express different types or levels of cadherins, the cells sort-out to form self-isolated, enveloped, and intermixed multicellular structures depending on their preference to bind to cells of the same or opposite type.21 Yet, also other mechanisms of self-sorting that rely on local cell signaling or contractile properties of cells have also been proposed and add to the complexity of multicellular systems.26 Similarly, multicolloidal mixtures self-sort into families of colloids based on multiple molecularly orthogonal homophilic and heterophilic interactions between different types of colloids.27–29 For example, mixtures of four distinct colloids self-sort into two families of colloidal aggregates using two orthogonal heterodimerization pairs by virtue of a behavior named social self-sorting.27,29

Here, we employ concepts known from colloidal self-assembly and explore how these can be used in the context of multicellular structures (Figure 1a). For this purpose, we establish different photoswitchable cell–cell interactions, which can be triggered under blue light illumination and turned off in the dark with different protein–protein interaction dynamics and dark reversion rates. Controlling the cell–cell interaction with light comes with the unique advantage of high spatiotemporal resolution and turning on the cell–cell adhesions remotely using low-intensity biocompatible light without interfering with other cellular processes. Most importantly, regulation with light allows tuning cell–cell interactions dynamically by using pulses of light. These unique features enabled us to investigate how the thermodynamics and kinetics of the interactions between the cellular building blocks impact the multicellular assemblies and achieve self-assembly under kinetic and thermodynamic control, as has been described for colloidal systems. Moreover, combining different orthogonal cell–cell interactions allowed us to not only self-assemble but also self-sort mixtures of four different cell types into separate preferential assemblies.

We first focused on establishing different photoswitchable cell–cell interactions with different binding strengths, protein–protein interaction dynamics, and reversion kinetics in the dark. For this purpose, we expressed different light-dependent protein–protein interaction partners as synthetic adhesion receptors on the surfaces of the breast cancer cell line MDA-MB-231, which do not form strong native cell–cell adhesions.30 In particular, we used three protein pairs that specifically heterodimerize with each other under blue light (450 nm) and dissociate from each other in the dark, named iLID and Nano (dark reversion rate 3.5 × 10^{-2} \text{s}^{-1}, t_{1/2} = 20 \text{s}),31 nMag and pMag (dark reversion rate 1.1 × 10^{-4} \text{s}^{-1}, t_{1/2} = 1.8 \text{ h}), and nMagHigh and pMagHigh (dark reversion rate 4.1 × 10^{-5} \text{s}^{-1}, t_{1/2} = 4.7 \text{ h}).32 These proteins were chosen due to the large range of dark reversion times they cover, their different protein–protein interaction dynamics, the tunability of their interactions with few point mutations (e.g., nMag/ pMag vs nMagHigh/pMagHigh), and their similar size, which presumably will lead to a similar expression level on the cell surface. To express these proteins on the cell surface each of the genes coding for them were cloned into a pDisplay vector.
which, once the protein is expressed, guides it to the cell membrane with an N-terminal murine Igκ-chain leader sequence and anchors it at the cell membrane with a C-terminal platelet-derived growth factor receptor (PDGFR) transmembrane domain (Supporting Information, Figure S1). Plasmids coding for different proteins were individually transfected into MDA-MB-231 cells, and stable monoclonal cell lines expressing these proteins at their surfaces were generated. The cell lines were named after the protein expressed at their surface; e.g., iLID expressing cells were named iLID-MDA. For each photoswitchable protein, a single clone with high protein expression was selected, and the expression of each protein on the cell surface was confirmed by flow cytometry and immunostaining (Supporting Information, Figures S2 and S3). On the surfaces of the cells $6 \times 10^3$ to $5 \times 10^4$ photoswitchable proteins per cell were expressed as measured by quantitative flow cytometry, showing comparable expression levels of the different photoswitchable proteins on the cell surface (Supporting Information, Table S2). In this study, we used MDA-MB-231 cells to demonstrate the concept, yet these genetically encoded photoswitchable proteins could be transfected and used to mediate cell–cell interactions between other cell types too.

To see if the photoswitchable proteins can mediate light-triggered cell–cell interactions, cells expressing complementary interaction partners (iLID-MDA and Nano-MDA, nMag-MDA and pMag-MDA, nMagHigh-MDA, and pMagHigh-MDA) were incubated in suspension in the dark and under blue light illumination for 30 min. The mixed cultures of two complementary cell types aggregated significantly under blue light but remained scattered in the dark, as observed in bright-field images (Figure 1b). To quantify the cell aggregation, large areas of the samples were scanned (1–2.56 cm$^2$ per sample containing about 25 000 cells/cm$^2$), and cell aggregates with a two-dimensional projected area of larger than 5000 μm$^2$, i.e., containing at least 20 cells, were detected as clusters using automated image analysis. In each of the three cocultures, blue light resulted in the assembly of multicellular structures with a significantly higher mean cluster area than in the dark (Figure 1c). Moreover, the blue-light-dependent cell aggregation was due to the specific heterophilic interactions between the different photoswitchable proteins, and homophilic interactions did not contribute to the aggregation, as in none of the monocultures containing just one cell type blue light illumination increased the cell aggregation (Figure 1d, Supporting Information, Figure S4).

One reason the protein pairs, iLID/Nano, nMag/pMag, and nMagHigh/pMagHigh, were selected is due to their different reversion kinetics in the dark. The reversibility of cell–cell interactions is a key feature of cell–cell adhesions in biology and indispensable for the self-sorting following the differential adhesion hypothesis. When cocultures of cells expressing complementary interaction partners were preaggregated for 30 min under blue light illumination and then placed in the dark, all three aggregate types dissociated, yet with different time dependences (Figure 1e, Supporting Information, Figure S5). The aggregates in iLID-/Nano-MDA cocultures disassembled the fastest within 60 min, aggregates in nMag-/pMag-MDA cocultures disassembled within 90 min, and aggregates in nMagHigh-/pMagHigh-MDA cocultures disassembled the slowest over 180 min. This trend corresponds to reversion time at the molecular level, which is iLID/Nano < nMag/pMag < nMagHigh/pMagHigh. The disparity in the absolute values for the reversion for the cell–cell interactions to the protein level could potentially be due to multivalent protein–protein interactions between cells, processes that are coupled to the cell–cell interactions beyond the photoswitching at the molecular level and the display of the proteins on the extracellular cell surface.

The second striking difference between different photoswitchable protein pairs was the morphology of the multicellular aggregates formed (Figures 1b and 2a). iLID-/Nano-MDA cocultures and nMag-/pMag-MDA cocultures formed compact aggregates with smooth edges after 2 h under blue light. On the other hand, in nMagHigh-/pMagHigh-MDA cocultures under the same conditions loose and ramified aggregates with irregular shapes formed. Furthermore, iLID-/Nano-MDA and nMag-/pMag-MDA aggregates were also larger than nMagHigh-MDA/pMagHigh-MDA aggregates. These observations suggest that the dynamics of the protein–protein interactions and their interaction strengths play an important role in the self-assembly of multicellular structures. While iLID-/Nano-MDA and nMag-/pMag-MDA aggregates exemplify the RLCA dominated by thermodynamic control, aggregates of nMagHigh-/pMagHigh-MDA cells are examples of the DLCA and are mostly under kinetic control. This data also mirrors the reaction- and diffusion-limited cluster aggregation observed in colloidal polystyrene particles coated with iLID and Nano or nMagHigh and pMagHigh, respectively. These observations directly correlate with the stronger protein–protein interaction between nMagHigh/pMagHigh and slower on/off rates compared to the weaker

Figure 2. Multicellular assemblies under kinetic and thermodynamic control. (a) Bright-field images of iLID-/Nano-MDA, nMag-/pMag-MDA and nMagHigh-/pMagHigh-MDA cells with different blue light pulsing sequences (120 min continues blue light, 0.5 min on/0.5 min off (only for iLID-/Nano-MDA), 1 min on/1 min off (only for iLID-/Nano-MDA), 5 min on/5 min off, 20 min on/20 min off, 1 min on/19 min off) after 2 h. Scale bars are 500 μm. Quantification of (b) the mean cluster area and (c) fractal dimension with different illumination protocols. Here, 64 images with a total area of 2.56 cm$^2$ were analyzed for each sample, each done in biological duplicates with 3 technical replicates. Error bars are the standard error of the mean cluster area, $p$-value < 0.001 represented as ***.

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and more dynamics protein–protein interaction between nMag/pMag and iLID/Nano.33

Next, we wanted to explore whether we could shift the self-assembled multicellular architectures from kinetically to thermodynamically controlled structures by altering the strength and dynamics of the cell–cell interactions. The photoswitchable cell–cell interactions provide a unique opportunity to address this question as protein–protein interaction strength and dynamics can be tuned using pulses of light.34,35 For this purpose, we incubated different cocultures under blue light illumination with varying on and off times for a total of 2 h (continuous 120 min on, 0.5 min on/0.5 min off (only for iLID-/Nano-MDA), 1 min on/1 min off (only for iLID-/Nano-MDA), 5 min on/5 min off, 20 min on/20 min off, 1 min on/19 min off) (Figure 2a). We observed that different multicellular aggregates formed depending on the illumination frequency. Outstandingly, less total illumination but in pulses lead to an increase in cell aggregation for iLID-/Nano-MDA cells (0.5 min on/0.5 min off), nMag-/pMag-MDA cells (5 min on/5 min off), as well as nMagHigh-/pMagHigh-MDA cells (5 min on/5 min off and 20 min on/20 min off) as also evident by the increase in the mean cluster area (Figure 2a, Supporting Information, Figure S6). This shows that pulsed illumination can lead to increased aggregation if the cell–cell interactions partially revert, and cells can reposition themselves when the light is off such that, upon reactivation with blue light, cells can optimize their position and increase interactions with their neighbors. For this reason, slower pulsing (20 min on/20 min off) enhances aggregation for nMagHigh-/pMagHigh-MDA cells with slower dark reversion and faster pulsing for iLID-/Nano-MDA and nMag/pMag cells (0.5 min on/0.5 min off, 5 min on/5 min off, respectively) with faster dark reversion. On the other hand, a longer off time (20 min on/20 min off) or less photoactivation (1 min on/19 min off) leads to a decrease in aggregation in all three photoswitchable cell–cell interaction pairs. This trend was best observed with nMag-/pMag-MDA cell aggregation, which increased with 5 min on/5 min off pulsing compared to continuous illumination, but decreased with lower pulsing frequency (20 min on/20 min off), although the total light dose was the same and even further if the photoactivation was decreased (1 min on/19 min off). Thus, if the reversion of the cell–cell interactions in the dark is extensive or the reactivation with blue light is not sufficient, aggregates disassemble, and when the dark reversion time of the photoswitchable protein is shorter, this disassembly is more pronounced. Taken together, this data shows that not only the cell–cell interaction strength but also their dynamics here modulated with pulsed illumination are critical for the self-assembly of multicellular structures.

The second aspect that is closely related to the cell–cell interaction dynamics is the morphology of the multicellular assemblies, which vary from loose and ramified to compact and spherical going from DLCA to RLCA.14 As observed above (Figure 2), aggregation increases when the cell–cell interaction is only partially reversed with pulsing, so that the cells could transiently reposition and strengthen their contact with neighboring cells, which represents a shift from kinetically to thermodynamically controlled structures. To rationalize and quantify the relationship between the morphology of the cluster and interaction dynamics, we determined the fractal dimension of the two-dimensional contours of these multicellular aggregates as a measure of cluster shape complexity and size36 (Figure 2c, Supporting Information, Figure S7). For comparison in colloidal systems, the fractal dimension increases from 1.46 for DLCA to 1.55 for RLCA for two-dimensional aggregates.37 For the cellular assemblies, we observed a significant range of fractal dimensions from 1.595 for nMagHigh-/pMagHigh-MDA cells under constant blue light dominated by DLCA to 1.651 for nMag-/pMag-MDA cells with 5 min pluses of blue light dominated by RLCA (Figure 2c). Under constant activation, the fractal dimension was higher for assemblies based on more dynamic protein–protein interactions, (iLID/Nano and nMag/pMag), achieving thermodynamically driven structures. On the contrary, nMagHigh-/pMagHigh-MDA cells formed stronger and less dynamic protein–protein interactions, leading to kinetically trapped structures with a lower fractal dimension. For comparison, we analyzed the fractal dimension of previously reported colloidal aggregates between iLID and Nano as well as nMagHigh and pMagHigh coated 2 μm polystyrene beads,29 and we observed similar trends as with the cellular aggregates. The iLID/Nano-mediated colloidal aggregates had a higher fractal dimension than the nMagHigh/pMagHigh-mediated colloidal aggregates (1.578 vs 1.562) (Supporting Information, Table S3). This analysis further confirms the parallels in the colloidal and cellular aggregates.

Pulsed photoactivation increases the dynamics of the cell–cell interactions and gives the cells an opportunity to rearrange and form a thermodynamically more stable structure, shifting the assembly from DLCA to RLCA as observed for all cell types. As shown in Figure 2c, nMagHigh-/pMagHigh-MDA cells under constant blue light formed branched clusters with a low fractal dimension (1.595), which increased up to 1.61 as the time in the dark increased and the photoactivation time decreased. Likewise, for nMag-/pMag-MDA and iLID-/Nano-MDA assemblies, the fractal dimension increased when 5 min on/5 min off and 0.5 min on/0.5 min off pulsing was used, respectively, compared to continuous blue light illumination. Beyond 5 min on/5 min off pulsing for nMag-/pMag-MDA and 0.5 min on/0.5 min off pulsing for iLID-/Nano-MDA cells, both the cluster size and fractal dimension reduced, suggesting excessive disassembly with an increase in reversion time in the dark. Moreover, the pulsing frequency required to achieve more thermodynamically controlled assemblies, i.e., RLCA, is closely connected to the dark reversion kinetics of the photoswitchable proteins. While 0.5 min on/0.5 min off was best for the iLID/Nano pair with the faster dark reversion kinetics, 5 min on/5 min off pulsing was the best for the nMag/pMag pair and the nMagHigh/pMagHigh interactions with the slowest dark reversion kinetics required longer dark periods (ca. 20 min) and less photoactivation to release the kinetically trapped structures and transform them into more compact assemblies.

Sorting-out/self-sorting is an important mechanism in nature to form multicellular structures out of multiple cell types and organize them in subdomains, as observed during embryogenesis and in vitro reconstitution studies of different tissue types.25 Achieving self-sorting in the context of bottom-up tissue engineering requires multiple orthogonal cell–cell interaction pairs with different interaction strengths, and each of these must be dynamic enough for cells to maximize the interactions with neighboring cells. If the cell–cell interactions are not dynamic enough, kinetically trapped architectures away from the thermodynamic optimum with no self-sorting form could form.
To achieve sorting-out and multicellular structures with subdomains, we mixed four different cell types expressing two orthogonal protein pairs at their surface. In particular, we mixed iLID-/Nano-MDA expressing cells (each stained in red) with either nMag-/pMag-MDA or nMagHigh-/pMagHigh-MDA expressing cells (each stained in green) to check if their orthogonal specificity could result into self-sorting in a mixed culture (Figure 3a). The social self-sorting of the cells was further quantified by analyzing the colocalization of the red and green fluorescent cells using the threshold overlap score (TOS) (Supporting Information, Figure S9). This analysis showed lower colocalization in cocultures of iLID-, Nano- with nMag- and pMag-MDA cells compared to cocultures where nMagHigh- and pMagHigh-MDA cells were used instead. The fact that self-sorting, specifically social self-sorting, was observed combining the more dynamic cell—cell interaction pairs, viz., iLID-/Nano-MDA and nMag-/pMag-MDA, which favor thermodynamically controlled assemblies and not the nMagHigh/pMagHigh pair, which forms kinetically trapped structures, also demonstrates the importance of dynamics in self-sorting. In an attempt to increase the dynamics between the nMagHigh-/pMagHigh-MDA cells, 20 min on/20 min off pulsing was used to achieve self-sorting within the four-component mixture. The pulsing increased the area of the clusters, and yet did not result in the complete self-sorting, and only domains of green- and red-labeled cells formed within the same aggregate, which was also reflected in an intermediate TOS compared to the self-sorting and nonsorting cocultures (Figure 3c, right, Supporting Information, Figure S9).

In this study, we demonstrate the importance of cell—cell interaction dynamics in the assembly and self-sorting of multicellular structures from cells as building blocks under kinetic or thermodynamic control. Blue-light-triggered cell—cell interactions based on different photoswitchable protein interactions (iLID/Nano, nMag/pMag, and nMagHigh/pMagHigh) with various binding strengths, protein-protein interaction dynamics and dark reversion kinetics provide unique tools for modulating cell—cell interaction dynamics. Using different interaction pairs and the temporal control that light as a stimulus provides, we were able to assemble and tune multicellular structures from branched and ramified to compact and spherical. Moreover, in mixtures with four different cell types, we were able to achieve self-sorting provided that the cell—cell interactions were dynamic enough, as also postulated by the differential adhesion hypothesis. These findings showed that concepts of DLCA and RLCA aggregation as well as of self-sorting that are well-established for colloidal systems can also be applied to the self-assembly of cells into tissue-like architectures. While to date, cell—cell interactions have been controlled using chemical and genetic approaches, and the importance of cell—cell interaction dynamics has not been considered. Most chemical approaches using DNA, clickable groups, and biotin—streptavidin form strong interactions with low exchange rates and are hence expected to result in DLCA, which represents kinetically controlled branched structures. On the other hand, introducing different cadherins to the cell surface, which form highly dynamic protein—protein interactions, results in RLCA with round assemblies under thermodynamic control. In terms of dynamics, the photoswitchable cell—cell interactions based on different photoswitchable proteins offer a wide range of interaction dynamics and strengths, which can be modulated to achieve both kinetically and thermodynamically driven multicellular assemblies. In this respect, bringing basic concepts of colloidal self-assembly to bottom-up tissue engineering will help in the design of more predictable and complex microtissue structures.
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