We report a general cell surface molecular engineering strategy via liposome fusion delivery to create a dual photo-active and bio-orthogonal cell surface for remote controlled spatial and temporal manipulation of microtissue assembly and disassembly. Cell surface tailoring of chemoselective functional groups was achieved by a liposome fusion delivery method and quantified by flow cytometry and characterized by a new cell surface lipid pull down mass spectrometry strategy. Dynamic co-culture spheroid tissue assembly in solution and co-culture tissue multilayer assembly on materials was demonstrated by an intercellular photo-oxime ligation that could be remotely cleaved and disassembled on demand. Spatial and temporal control of microtissue structures containing multiple cell types was demonstrated by the generation of patterned multilayers for controlling stem cell differentiation. Remote control of cell interactions via cell surface engineering that allows for real-time manipulation of tissue dynamics may provide tools with the scope to answer fundamental questions of cell communication and initiate new biotechnologies ranging from imaging probes to drug delivery vehicles to regenerative medicine, inexpensive bioreactor technology and tissue engineering therapies.

Results and discussion

To generate tissue assemblies containing multiple cell types for a range of fundamental cell behavior, cell imaging and tissue engineering applications, we used a bottom-up synthetic approach to rewrite cell surfaces. Cell surface tailoring was achieved by a straightforward new liposome fusion method to incorporate complementary bio-
orthogonal molecules capable of an intercellular click chemical reaction upon physical cell–cell contact\(^{13,14}\). The external cell surface click conjugation between cells proceeds at physiological conditions in the presence of serum and allows for stable cell interconnectivity. The limited suite of bio-orthogonal click reactions is increasingly becoming important tools in chemical biology and cell biological research\(^{15,16}\). To access spatial and temporal control of cell–cell interactions, the synthetic ligation tether between cells was engineered to contain a photochemical cleavage site\(^{17}\). Remote controlled tissue disassembly proceeds by a programmed photo-initiated cleavage of the intercellular ligation tether (Fig. 1 top). The key features are the delivery of synthetic chemical groups to cell surfaces (via liposome fusion)\(^{13}\), the intercellular oxime click ligation bond\(^{16}\) (bio-orthogonal) and a photo-cleavage site contained within the oxime lipid tether (Fig. 1 bottom)\(^{17}\).

To demonstrate temporal control of tissue assembly and disassembly, we first delivered the key functional groups to different cell populations (Fig. 2). We generated three liposome populations containing the photo-oxime lipid (1), ketone lipid (2) and oxime lipid (11) respectively (Supporting scheme S1, Supporting Fig. S1–S4). By mixing these lipid-like molecules with background lipids (palmitoyl-oleoyl phosphatidylcholine (POPC) and positively charged, 1,2-dioleoyl-3-trimethylammonium-propane (DOTAP)), we generated liposomes that can easily fuse into a variety of cells to deliver the photo-oxime, ketone and oxime groups (Fig. 2A–C). As a demonstration, we applied this liposome fusion method to non-adherent Jurkat cells to generate different two cell populations; one presenting photo-oxime, ketone groups and the other presenting oxime groups. As controls, when regular Jurkat cells were mixed with ketone tailored Jurkat cells (Jurkat-ketone), no cell assembly was observed (Fig. 2D). When oxime tailored Jurkat cells (Jurkat-ONH\(_2\)) or photo-oxime tailored Jurkat cells (Jurkat-photo-ONH\(_2\)) were mixed with Jurkat-ketone, rapid multi-cell spheroid assemblies were generated via the intercellular oxime click ligation (Fig. 2E–F). When mild UV illumination was applied (365 nm, 10 mW/cm\(^2\), 5 min), the photolabile oxime linkage connecting cells was cleaved resulting in the complete disassembly of the microtissue into individual cells.

To generate and control the size of co-culture aggregate cell assemblies, the concentration of the mixed tailored cells in solution and duration of interaction were varied. As expected, higher concentrations of cells and longer durations resulted in larger co-culture spheroids (Supporting Fig. S5). As an important control, cell viability studies showed no difference between cell populations that were tailored with and without functional groups via liposome fusion (Supporting Fig. S6). Due to the photo-cleavable nature of the intercellular oxime ligation, spheroid disassembly proceeded upon illumination with UV light (365 nm, 10 mW/cm\(^2\), 5 min). The behaviors of newly disassembled cells were indistinguishable from control cells (viability, migration, proliferation). This strategy may

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**Figure 1** | Schematic describing the molecular level control of tissue assembly and disassembly via a chemoselective, bio-orthogonal and photoswitchable cell surface engineering approach. (top) Two or more different cell types are engineered via liposome fusion to present complementary and bio-orthogonal molecules on their cell surfaces. Upon interaction, a stable and covalent click (oxime) type reaction occurs and tissue like assemblies are formed. After a defined period of time, tissue disassembly can be remotely controlled by photo-cleavage of the intercellular ligation tether. (bottom) A schematic showing the interfacial molecular lock and cleave sequence for conjugating and releasing cell associations. The lipid lock molecule contains both a photocleavable group and an amineoxo group for bio-orthogonal oxime ligation to ketone presenting cells. Disassociation of cell assemblies via photo-cleavage occurs upon UV illumination.
Figure 2 | The design of liposomes to deliver chemoselective, bio-orthogonal and photo-active groups to cell surfaces and the subsequent application to assemble and disassemble microtissues on demand. (A) A photolabile oxime molecule is mixed with POPC and positively charged DOTAP to generate photo-active liposomes that are delivered to cell surfaces via liposome fusion. (B) Dodecanone is mixed with POPC and DOTAP molecules to generate ketone presenting liposomes that are delivered to cell surfaces via liposome fusion. (C) O-dodecyl oxime is mixed with POPC and DOTAP molecules to generate oxime presenting liposomes that are delivered to cell surfaces via liposome fusion. (D) No cell assembly occurred when control Jurkat cells were mixed with ketone presenting Jurkat cells. (E) Programmed microtissue assembly proceeded upon mixing of jurkat-photo-ONH$_2$ with jurkat-ketone cells. Microtissue disassembly to single cells proceeded upon cleavage of the intercellular linkage through uv light illumination. (F) Cell Assembly proceeded upon mixing of jurkat-photo-ONH$_2$ with jurkat-ketone cells. Microtissue disassembly to single cells proceeded upon cleavage of the intercellular linkage through uv light illumination.

allow for temporal control for a range of autocrine and paracrine signaling studies and provide new ways to study co-culture and multi-cell type associations.

As a further application, cells that were rewired with the photo-oxime were seeded onto patterned materials presenting aldehyde groups (Fig. 3). Cells specifically attached to the material through the interfacial oxime ligation bypassing non-specific associations used in regular tissue culture and native ligand-receptor based adhesion (integrin-extracellular matrix)18–20. Upon exposure to UV light (365 nm, 10 mW/cm$^2$ 5 min) the photo-oxime was cleaved and the cells detached from the substrate (Supporting Fig. S7). This strategy allows for non-adherent cells to become adhesive to tailored materials and in combination with microfluidic technology may allow for new cell sorting, cell patterning and tissue capture/release biotechnologies.

Fig. 4 shows the construction of a multi-layered tissue co-culture system based on rewiring cell surfaces via liposome fusion. Two different types of cells (hMSCs and fibroblasts), tailored with ketone and photo-oxime respectively, were peeled from tissue culture plates and assembled via the oxime ligation to form a stable multilayered tissue co-culture system. Images of the co-culture tissue showed regions of multilayer and monolayer. The cell layers only adhered if the complementary chemistry was present on their cell surfaces (Supporting Fig. S8). These results show that large tissues can be easily assembled via specific cell surface ligation and may provide new efficient and inexpensive bio-reactor routes to generate complex tissues and organs.

Furthermore, red fluorescent protein (RFP) fibroblasts and green fluorescent protein (GFP) fibroblasts were used in combination with the bio-orthogonal liposome fusion method to demonstrate that multilayer co-cultures can be assembled with control of thickness and orientation.

Since the linkage between cell layers contained a photo-cleavage site, the co-culture tissue could be separated upon UV exposure (Supporting Fig. S9). These results show that the oxime ligation between cells can be scaled from initial liposome fusion (nanometer scale) to small clusters of cells (micrometer scale), to large tissue patches (centimeter scale). Furthermore, upon addition of induction media, the click ligated tissue patches containing hMSCs could differentiate to adipocytes, fibroblasts and osteoblasts (Supporting Fig. S10). Because a range of cell lines may be integrated with stem cells to generate co-culture multi-layers, new stem cell differentiation studies and higher order 3-dimensional tissues may be possible. This strategy is general and may be used to produce complex multi-cell type structures for a range of regenerative medical applications (stem cell co-cultures, tissue grafts, etc…) and as a high-throughput tissue chip screening technology.

We used flow cytometry to quantify and characterize the amount of photo-oxime lipid delivered to cells via liposome fusion, for subsequent photo-oxime ligation induced microtissue assembly (Fig. 5). The photo-oxime was designed to have 3 components that are essential for spatial and temporal control of cell interconnectivity (Fig. 5A). A lipid hydrophobic component to insert into membranes, a photo-cleavable center and a bio-orthogonal oxime group for cell surface ligation of a range of ligands or other cells presenting ketone groups.

Liposomes containing the photo-oxime lipid were synthesized and then delivered to fibroblasts in culture. To measure the amount of photo-oxime incorporated, the cells chemoselectively reacted via oxime formation with a fluorescent calcein dye containing a ketone group. FACS analysis determined the amount of photo-oxime molecule present at the cell surface after various time
points after liposome fusion (Fig. 5B–E). As expected, the cells initially had the highest amounts of photo-oxime molecule, which gradually decreased over time as the cells grew and divided. This decrease is due to multiple cell divisions and therefore dilution of the photo-oxime lipid molecule over subsequent generations of cells. Depending on the cell line, this dilution took weeks and many rounds of cell divisions. After dilution, the cell lines were indistinguishable from untreated cells and could again be used in normal cell culture or for future liposome fusion enhancement or new cell surface tailoring. These results and observations are analogous to classical CDNA transient transfections but in this case with photo-active bio-orthogonal lipids being the transfected biomolecule. Furthermore, FACS analysis is able to quantitate the amount of lipid at different time points and shows the amount of lipid can be controlled by adjusting the liposome fusion conditions (duration, concentration) and allows for multiple fusions of different lipid like molecules (for eg. different bio-orthogonal lipids for hydrazone, huisgen, diels-alder, thiol-ene type conjugations or lipids with fluorescent, spin label, radiolabel etc. properties) or serial fusions at different time points.

As further characterization of cell surface presenting photo-oxime groups, we developed a novel mass spectrometry method (Fig. 5F). Lipid transfected cells were added to SAM surfaces presenting aldehyde groups, which allow cells to attach to the surface via the interfacial oxime ligation. Cells were then removed from the surface (lysed) with vigorous agitation through a stream of PBS solution and H2O. Due to the covalent nature of the interfacial oxime bond between the cells and SAM surface, the lipid was essentially pulled out of the membrane and remained conjugated to the substrate. Because the substrate is conductive, MALDI mass spectrometry could be performed directly on the substrate to characterize the interfacial reaction. Fig. 5G shows MALDI analysis and clearly shows the presence of the oxime molecule on the substrate. This strategy may be used to characterize a range of lipid presenting molecules on cell surfaces and as a new pull-down method of proteins if the lipid molecule spans the plasma membrane and covalently associates with cytoplasmic proteins.

To demonstrate the spatial and temporal control of tissue assembly using the intercellular photo-oxime strategy, we generated multi-layers of adipocytes and hMSC cells (Fig. 6). We first added a dodecanone liposome to hMSCs, which readily presented the ketone from the cell surface. Addition of adipocytes presenting photo-oxime to the hMSCs rapidly generated a two layer structure. In comparison, no layers were formed without the complementary chemistry presented from either cell type. Fig. 6A–C shows images and confocal representations of the layers. Upon UV illumination through a mask, the adipogenic cells were released from the hMSCs due to the cleavage of the intercellular photo-oxime ligation tether.

In order to extend this strategy for potential stem cell differentiation and tissue engineering applications, we generated cell layers through the photo-oxime ligation between hMSCs and fibroblasts (Fig. 6D–I). After incubating in stem cell induction media, the hMSCs differentiated to adipocytes and subsequent UV illumination and media exchange resulted in the release of the fibroblasts generating relatively pure adipogenic cells. It should be noted that the liposome fusion method is general and can be introduced to many cell lines.

**Conclusion**

We have developed a novel liposome fusion system to deliver bio-orthogonal photo-active lipid and ketone-lipid like molecules to cell membranes. Upon mixing these cells in different formats, co-culture spheroids and multilayers could be generated due to an intercellular oxime (click) ligation. Furthermore, we also showed the feasibility of producing macroscopic-scale tissue and achieving multiple tissue-tissue assembly, which could lead to a series of tissue engineering, especially organ printing technologies. Since the ligation tether contains a photo-cleavable group, remote control of disassembly could be achieved upon UV light illumination. We demonstrated this system in several cell lines to generate switchable co-culture spheroids and multi-layers. Flow cytometry and mass spectrometry analysis quantified and characterized the interfacial cell surface reaction. The ability to engineer cell surfaces with a straightforward and inexpensive liposome fusion strategy will find wide use in fundamental studies of membrane biophysics, paracrine signaling and adapt to generate new biomaterials and as a biotechnology platform for screening complex cell behaviors in tissue microarrays. Several other bio-orthogonal chemical ligation strategies including diels-alder, huisgen, oxime, hydrazone, thiol-ene, etc. may be used to tailor cell surfaces with nanoparticles, redox groups and a range of other molecules for targeted delivery and as cell tracking and imaging beacons. The spatial and temporal control of cell interactions between mul-
multiple different cell types will lead to new studies of dynamic cellular communication. Furthermore, the combination of bioreactor technologies with intercellular ligation methods may provide new ways to generate large-scale complex multi-cell type tissues. When combined with traditional polymer scaffolds, molds or printing technologies, a range of complex 3D tissues and organs may be possible for an array of biomedical diagnostic and transplantation applications.

Methods

Structure and list of molecules, liposomes and cells used in this study are shown in Supplementary Information (scheme S1). Synthesis and other experimental procedures and characterizations are detailed in Methods and Supplementary Information. O-Dodecylamine was synthesized as previously described. Egg palmitoyl-oleoyl phosphatidylcholine (egg-POPC) were purchased from Avanti Polar Lipids (Alabaster, AL), and all other chemicals were obtained from Sigma-Aldrich or Fisher. Swiss 3T3 albino mouse fibroblasts were obtained from ATCC. Human mesenchymal stem cells (hMSCs), basic medium, growth medium and differentiation medium were obtained from Lonza.

Liposome preparation. Liposomes were prepared as previously reported. To generate photo-oxyamine (1) or ketone (2) liposome, photo-oxyamine or dodecanone (60 µL, 10 mM solution in CHCl₃) were dissolved with egg-POPC (450 µL, 10 mg/mL in CHCl₃) and 1,2-dioleoyl-3-trimethylammonium-propane (DOTAP, 10 µL, 10 mg/mL in CHCl₃) in chloroform followed by concentration under high vacuum for 4 h. The dried lipid samples were then reconstituted and brought to a final volume of 3 mL in PBS buffer, pH 7.4. The contents of the vial were warmed to 50°C and sonicated for 20 min, in a tip sonicator, until the solution became clear, and liposomes containing photo-oxyamine or ketone groups were formed.

Cell culture. Human mesenchymal stem cells (hMSCs) were cultured as instructed by the vendor. After cells were washed with PBS and trypsinized for 3–5 minutes, they were centrifuged in serum containing medium and followed with gentle resuspending in serum free medium. The cells were then seeded onto transparent glass substrates and then incubated at 37°C in a humidified atmosphere of 5% CO₂ overnight. Adipogenic differentiation was induced by adipogenic induction medium and kept by induction/maintenance cycles as described in the Lonza protocol. Osteogenic differentiation was induced by osteogenic induction medium provided by Lonza. 3T3 Swiss Albino Fibroblasts, RFP Expressing Human Neonatal Dermal Fibroblasts, and C3H/10T1/2 cells were cultured in Dulbecco’s modified Eagle medium (DMEM) with...
10% fetal bovine serum (FBS) and 1% penicillin/streptomycin. NIH3T3/GFP cells were cultured in DMEM containing 10% FBS, 0.1 mM MEM Non-Essential Amino Acids, 2 mM L-glutamine, 10 μg/mL Blasticidin, and 1% penicillin/streptomycin. These cells were incubated at 37 °C in a humidified atmosphere of 5% CO2, and released from tissue culture plates using 0.05% trypsin in 0.53 mM EDTA.

**Immunohistochemistry.** The substrates for confocal imaging were fixed with formaldehyde (3.2% in PBS) and permeated (PBS containing 0.1% Triton X–100). A fluorescent dye mixture, containing phalloidin-TRITC (actin) and DAPI (nucleus) was then made in PBS containing 5% normal goat serum and 0.1% Triton X –100.

Cells were incubated with the dye solution for 2 h. The substrates were then secured in fluorescence mounting medium (Dako, Carpinteria, CA, USA), which enhances the visualization of cells when viewed under a fluorescent microscope with a glass cover slip. The substrates for adipogenic differentiation were washed by PBS and fixed in 3.2% formaldehyde for 30 minutes, followed with sterile water and 60% isopropanol for 5 minutes. Samples were then stained by Oil Red O for 5 minutes followed by Harris Hematoxylin for 1 minute. The substrates for collagen differentiation were fixed with formaldehyde and permeated with 0.1% Triton X–100. Monoclonal antibody of collagen I was applied for 1 h, then incubated with secondary antibody anti-mouse IgG (FITC conjugate) for 30 min, and followed with DAPI for 10 min.

**Figure 5 | Characterization of the photo-oxime reaction at cell surfaces.** (A) NMR characterization of the photo-oxime molecule. Each part of the molecule is designed to optimize membrane insertion, photo-cleavability and availability of the oxime group for intercellular ligation. (B) Flow cytometry analysis to determine the amount of photo-oxime molecules per cell surface. Fibroblast cells were cultured with or without (control) photo-oxime containing liposomes for different time periods. The cells were then reacted with a ketone containing fluorescent calcein dye. The cells were then tested against a standard bead (~10^7 beads/mL) with known fluorescein molecule density. Approximately 25 × 10^7 cells were counted for all samples. Samples were run in triplicate, and the mean fluorescence intensity values are displayed. (C, D) Histograms relating the number of cells counted as a function of fluorescence intensity are shown and labeled as control (without photo-oxime) and 12 h, 36 h, 72 h, and 96 h (with photo-oxime). (E) Micrograph of cells presenting photo-oxime reacted with calcein. When no photo-oxime is present in the cell surface, no fluorescence is observed. (F) A cartoon describing the interfacial reaction between the surface-engineered cells and a self-assembled monolayer (SAM) on a gold substrate. Cells tailored with the photo-oxime group are seeded onto a SAM presenting aldehyde groups to form an interfacial oxime ligation. After washing and removing the cells, the photo-oxime lipid is pulled out of the cell membrane and remains ligated to the SAM via the covalent oxime bond. (G) MALDI-MS characterization of the resulting substrate shows an interfacial reaction occurs between the rewired cell surfaces and the aldehyde substrate via oxime ligation.
30 min for nucleus staining (reagents from Fisher Scientific). The substrates for osteogenic differentiation were stained with sigma ALkaline Phosphatase (ALP) kit (sigma kit 85).

Confocal microscopy. Cell clusters (spheroids) and tissue formation were visualized with a Nikon Eclipse TE2000-E inverted microscope (Nikon USA, Inc., Melville, NY). Data was analyzed by Metamorph software and a spectral confocal microscope (Leica Microsystems, Bannockburn, IL). Three-dimensional reconstructions of fluorescent images were generated using Volocity software.

Flow cytometry. Fluorescence-activated cell sorting (FACS) analysis was performed to quantify the approximate number of photo-oxyamine lipids at the cell surface after membrane fusion. Liposomes were cultured with Fbs (3 mM in tris buffer, 400 μL added to 4 mL) to present photo-active group on cell surface (9). Ketone-conjugated fluorescein was then reacted with the engineered cells (0.15 mM in tris buffer, 2 h). This time course assay was conducted to determine whether the chemistry was being carried on after cell growth and division. A control cell population (not displaying photolabile lipids) was incubated with the ketone-fluorescein (0.15 mM in tris buffer, 2 h). After culturing for the appropriate time, the different cell populations were washed with PBS (3 × 5 mL), trypsinized (1 mL, 5 min, 37°C, 5% CO2), centrifuged (5 min, 1000 rpm), and resuspended in RPMI (~10^5 cells/mL). Flow cytometry measurements were calibrated using RCP-5-30 beads (~10^7 beads/mL). Data was analyzed with Summit 4.3 software. The error bars are represented as the mean fluorescence intensity SD of 3 trials.

Matrix-assisted laser-desorption/ionization mass spectrometry (MALDI-MS). Gold-coated MALDI sample plates (123 × 81 mm) (Applied Biosystems, Foster City, CA) were prepared by electron-beam deposition (Thermionics Laboratory Inc, Hayward, CA) of titanium (5 nm) and then gold (12 nm). In order to form self-assembled monolayers (SAM) of alkanethiolates on the plates, the slides were
immersed in a 1 mM solution of 1:1 ratio mixture of 11-mercaptoundecanol and tetra(ethylene glycol)-terminated undecanethiol in EtOH for 12 h, rinsed with EtOH and dried, and then partially oxidized to aldehyde by mild oxidant pyridinium chlorochromate (PCC), as previously reported\(^1\). Once removed from solution, the surfaces were rinsed with EtOH and dried before use. Cells tailored with photolabile oxynine group are seeded onto the SAM presenting aldehyde group to form oxime ligation between the cell membrane and gold substrate. After washing and removing the cells, the bonded residue on the gold substrate was traced by MALDI-MS. MALDI analysis was carried out using an AB SCIEX TOF/TOF™ 5800 System (Applied Biosystems, Foster City, CA).

**Cell patterning.** Self-assembled monolayers (SAMs) presenting aldehyde and tetra(ethylene glycol) (EG4) groups were patterned on gold substrate at a ratio of 1:9 using microfluidic oxidation to ensure that fbs were only adhering to the patterned surface regions that presented 10% aldehyde groups\(^2\). Fbs were cultured with photo-liposomes (4 h) and then seeded (~10^4 cells/mL, 2 h) to the patterned aldehyde surfaces. Media containing 10% calf bovine serum (CBS) and 1% penicillin/ streptomycin was then added, and the substrates were incubated at 37°C in 5% CO_2 for 4 d. Cells cultured with liposomes, not containing the key functional groups, did not attach to the patterned surfaces. Substrates were then imaged by fluorescence microscopy.

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**Author contributions**

M.N.Y. designed the study. W.L., A.P. and D.D. performed the experiments. M.N.Y., W.L., M.N. and B.M.L. analyzed the data. M.N.Y. wrote the manuscript.

**Additional information**

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