

Supplementary Information

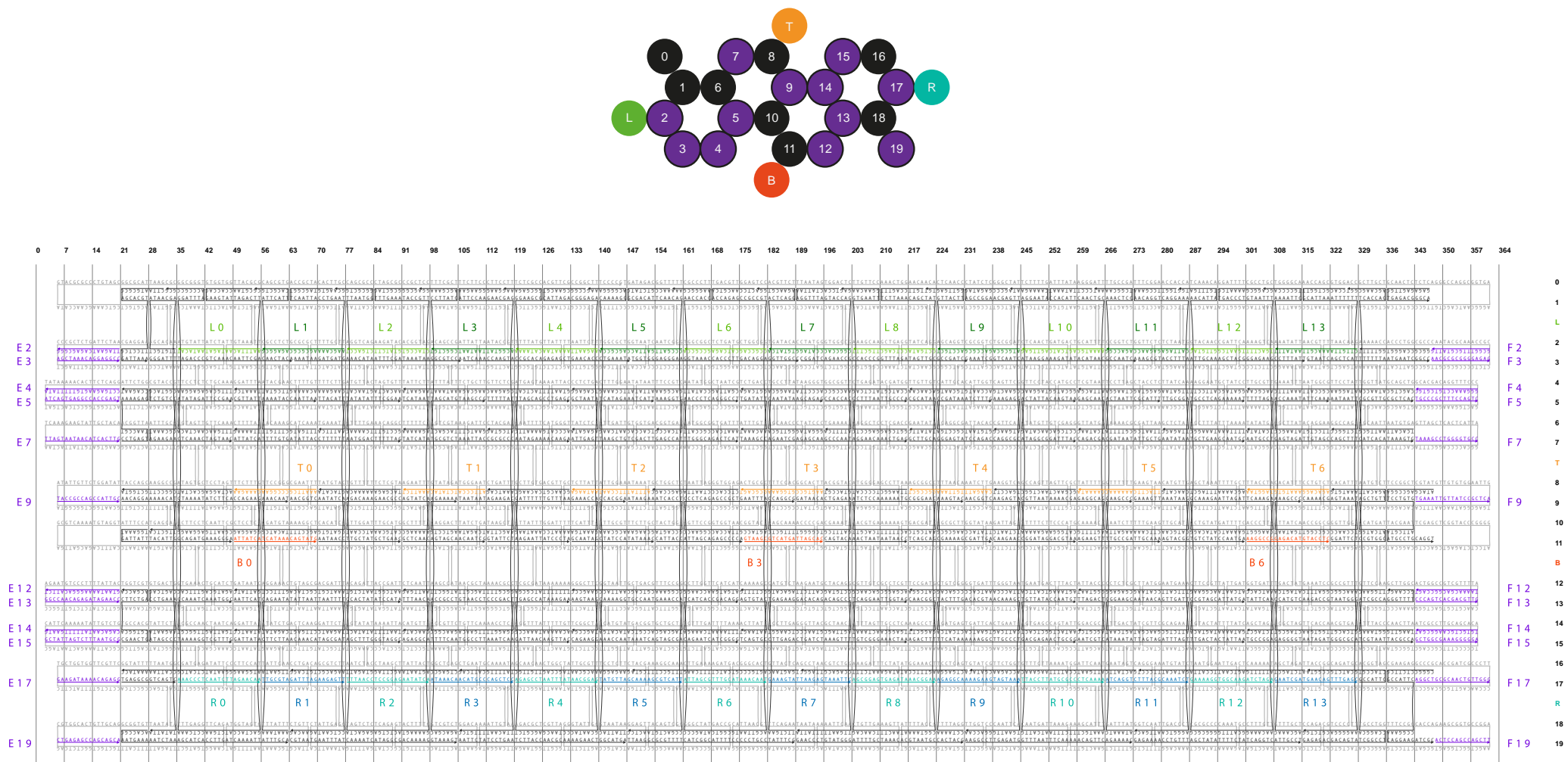
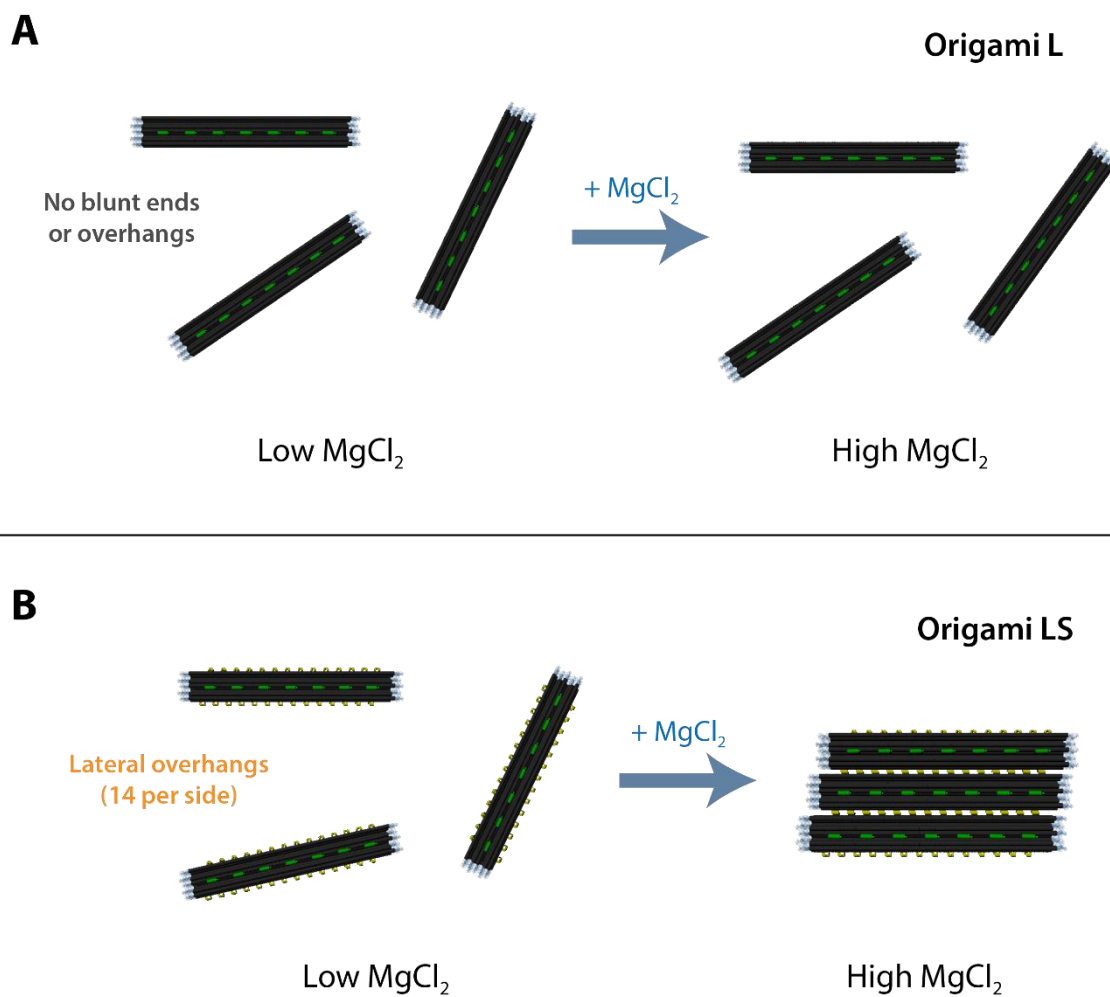


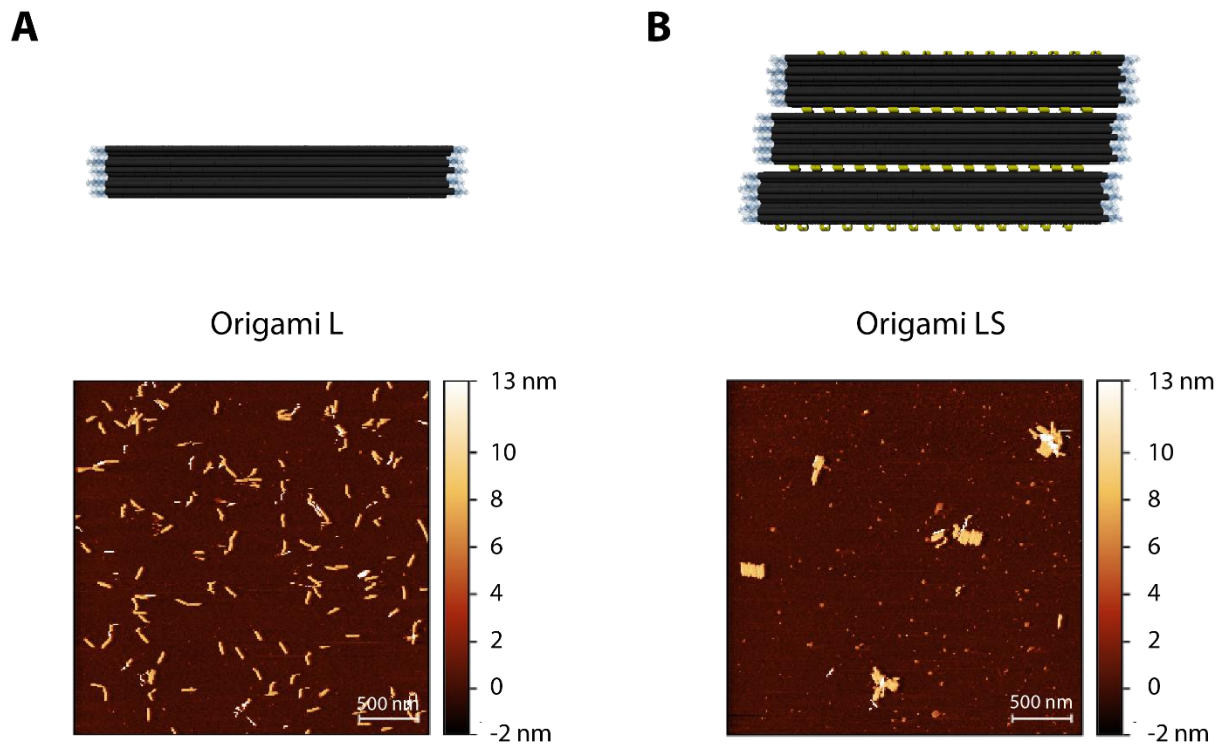
Figure S1 | Cadnano design and oligonucleotide sequences of the various origami L structures.

Top positions T0-T6 are coloured in gold, while bottom positions B0, B3, B6 are coloured in dark orange. Lateral positions L0-L13 and R0-R13 are coloured in green and blue, respectively. Edge positions E2-5, E7, E9, E12-15, E17, E19 and F2-5, F7, F9, F12-15, F17, F19 are coloured in purple. Core staples are coloured in black; M13 p7249 scaffold is coloured in grey. List of functionalized staples can be found in Table S1.



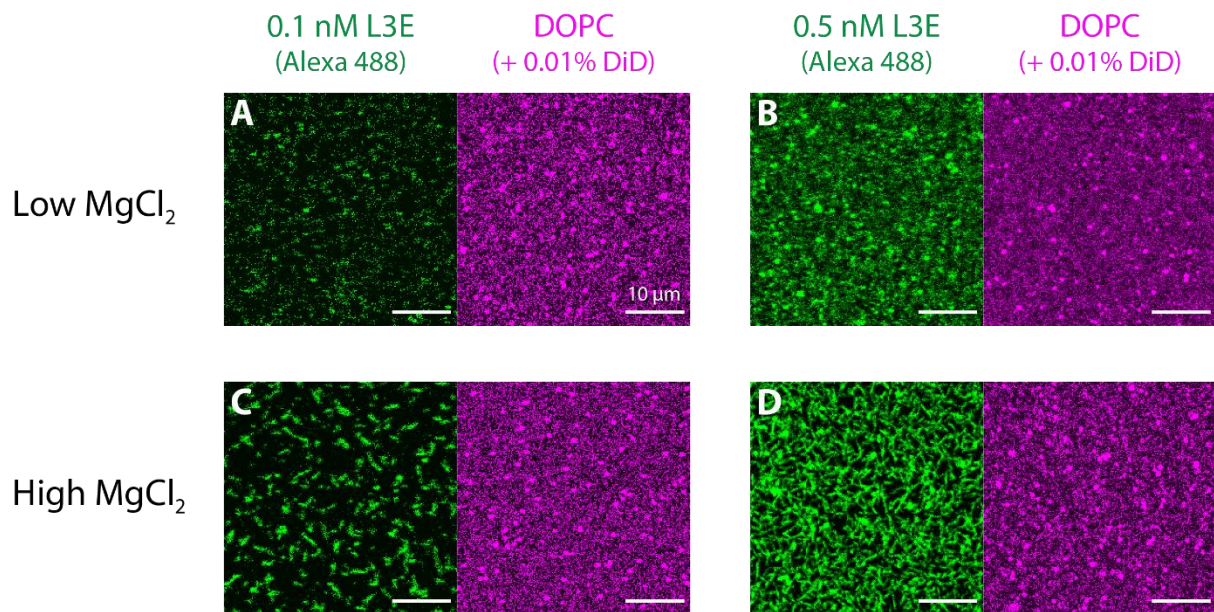
**Figure S2 | Depiction of the self-assembly patterns of origami L and LS upon increasing  $\text{MgCl}_2$ .**

(A) Origami L does not possess the ability to polymerize, as it lacks blunt ends or lateral overhangs for establishing intermolecular interactions. (B) Origami LS displays 14 lateral self-complementary single-stranded overhangs on both sides, and is therefore able to polymerize into sheet-like oligomers at high  $\text{MgCl}_2$  concentrations.



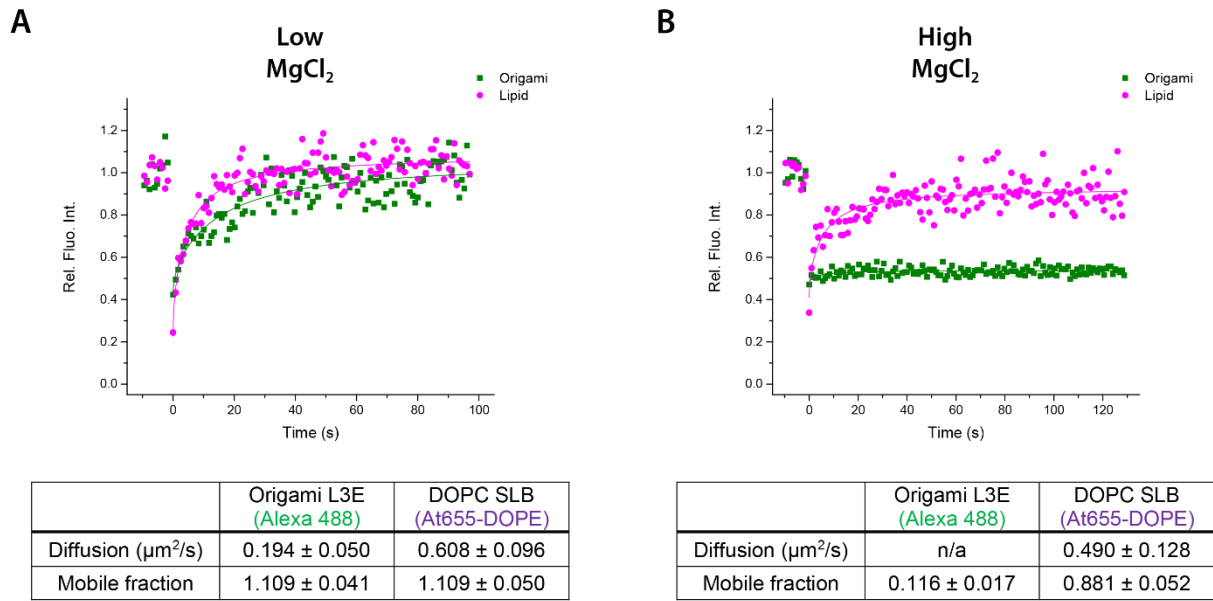
**Figure S3 | Self-assembly properties of origami L and origami LS at high MgCl<sub>2</sub>.**

Atomic force microscopy (AFM) images on PLL-mica of origami L lacking blunt ends (**A**) and origami LS displaying 14 lateral self-complementary single-stranded overhangs on both sides (**B**) after incubation with a high MgCl<sub>2</sub> buffer (70 mM MgCl<sub>2</sub> + 187.5 mM NaCl). As here depicted, whereas origami L (**A**) stays in a monomeric form, origami LS (**B**) polymerizes into sheet-like oligomers.



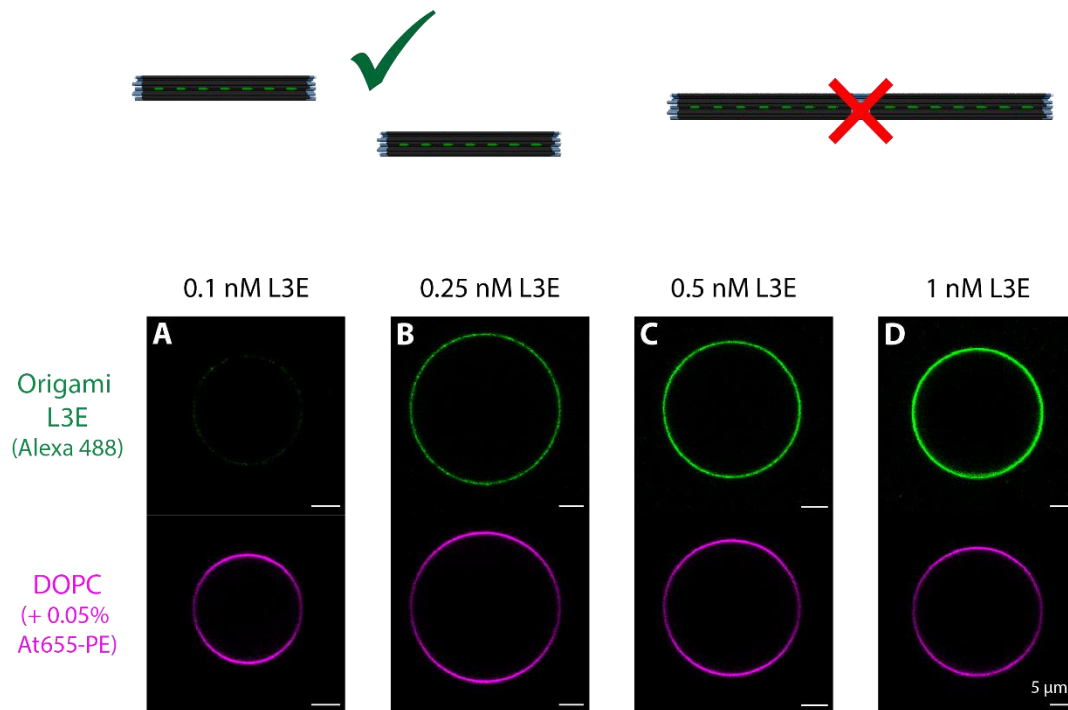
**Figure S4 | Binding and polymerization of DNA origami L3E on top of supported lipid bilayers.**

Zoomed-out images depicting the interaction of 0.1 and 0.5 nM Alexa488-labelled DNA origami L3E displaying 3 TEG-chol anchors (for membrane binding) and blunt ends (for end-to-end self-assembly) with DOPC SLBs (doped with 0.01% DiD). **(A-B)** At low  $\text{MgCl}_2$  (5 mM  $\text{MgCl}_2$  + 300 mM NaCl), a mostly homogenous distribution of origami L3E was observed on top of the lipid bilayers. **(C-D)** At high  $\text{MgCl}_2$  (70 mM  $\text{MgCl}_2$  + 187.5 mM NaCl) origami filaments were observed on top of the lipid bilayer. While at 0.1 nM, origami L3E formed individual short filaments **(C)**, at 0.5 nM origami L3E self-assembled into a mesh of longer and bundled filaments **(D)**.



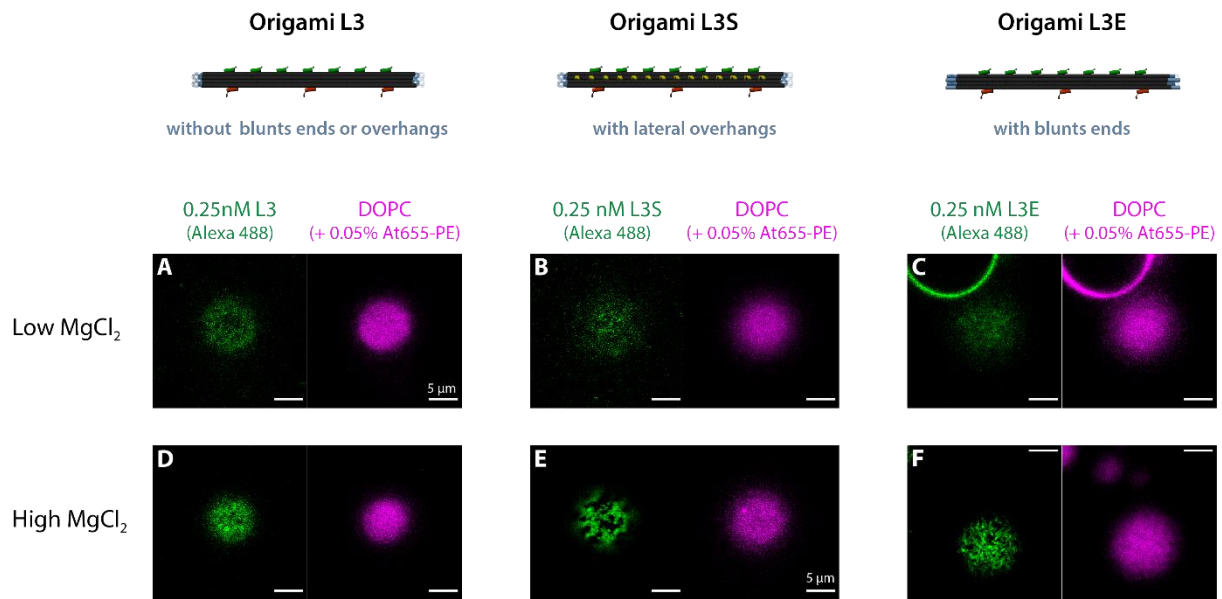
**Figure S5 | Fluorescence recovery after photobleaching (FRAP) of membrane-bound origami L3E on DOPC supported lipid bilayers (SLBs).**

FRAP data (from Movies S1 and S3), fitted results and calculated diffusion coefficients/mobile fractions of 0.5 nM origami L3E on a DOPC SLB, in the presence of **(A)** low MgCl<sub>2</sub> (5 mM MgCl<sub>2</sub> + 300 mM NaCl) and **(B)** high MgCl<sub>2</sub> (70 mM MgCl<sub>2</sub> + 187.5 mM NaCl).



**Figure S6 | Interaction of origami L3E with giant unilamellar vesicles at low  $MgCl_2$ .**

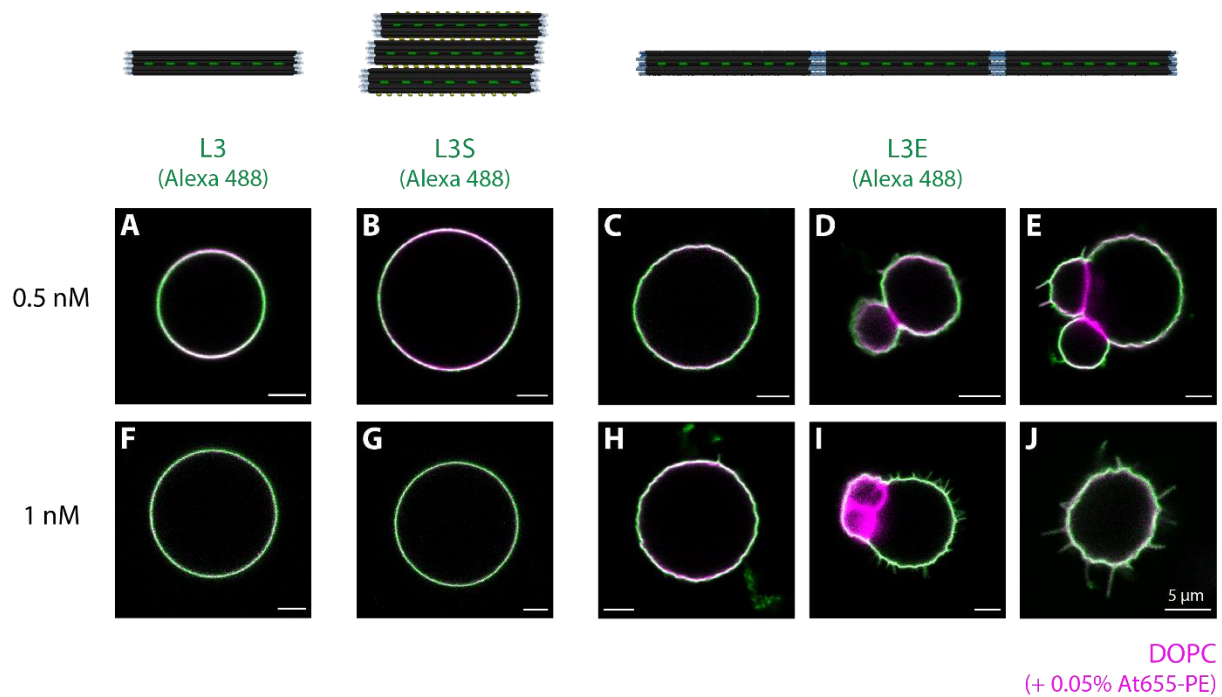
Membrane attachment of different bulk concentrations (0.1-1 nM) of Alexa488/TEG-chol-modified origami L3E displaying blunt ends to DOPC GUVs (doped with 0.05% Atto655-DOPE) in the presence of low  $MgCl_2$  buffer (5 mM  $MgCl_2$  + 300 mM NaCl). Images correspond to equatorial plane slices of GUVs.



**Figure S7 | Triggering self-assembly of membrane-bound origami L3, L3S and L3E.**

As depicted for the pole of selected GUVs, at low  $\text{MgCl}_2$  (**A-C**), origami L3, L3S and L3E are homogeneously distributed on top of DOPC GUVs, corroborating their predominant monomeric state under these conditions. Upon increasing the amount of  $\text{MgCl}_2$ , membrane-bound origami L3 (lacking blunt ends and lateral overhangs) remains homogeneously distributed (**D**); origami L3S (displayed lateral overhangs) can engage into lateral self-assembly, giving rise to large platforms (**E**); and finally origami L3E (displaying blunt ends) can polymerize end-to-end, giving rise to a mesh of filaments (**F**).





**Figure S8 | Membrane deformations by origami L3, L3S and L3E at high  $MgCl_2$ .**

Equatorial plane images of GUVs incubated with 0.5 nM (**A-E**) and 1 nM (**F-J**) origami L3/L3S/L3E at least 90 min prior addition of additional  $MgCl_2$ . For origami L3 lacking the ability to polymerize (**A**, **F**) no significant membrane deformations were reported. Similar results were observed for vesicles incubated with origami L3S, able to form lateral origami platform (**B**, **G**). On the contrary, for vesicles with membrane-bound origami L3E, extensive remodelling as rough (**C**, **H**) and spike-like **tubular** (**D**, **E**, **I**, **J**) deformations were observed, after  $MgCl_2$ -triggered end-to-end self-assembly of L3E into linear origami aggregates/filaments.



**Table S1 | List of functional staples used for various origami L structures.**

Oligo	Sequence	Description	Partner staple
TD_00	AAATTCGCCCGGAACAAAGAAAAAACACCAAACCC	staples with extension for Alexa488 dye	<i>5'-Alexa488-</i>
TD_01	ATTCATCTATACAAATTCTAAAAACACCAAACCC	<i>used in all origami</i>	<i>GGGTTTGGTGTTTTTT</i>
TD_02	ATTTATTTCCAATAATAAGAAAAAACACCAAACCC		
TD_03	AAGTGCCGTGGAAAGCGCAGTAAAAACACCAAACCC		
TD_04	CAAGATTTGTTAAAGGCCGCTAAAAACACCAAACCC		
TD_05	TTACTTCAAAAAACAAAATAAAAAACACCAAACCC		
TD_06	AGACAGGAAATGTGTAGGTAAAAAACACCAAACCC		
B18_00	ATTATCATCATAAACAGTATGGCTATGGGTGGTCTGGTT	staples with extension for TEG-Chol(18)	<i>5'-Chol-TEG-</i>
B18_03	GTAAGCGTCATGATTAGCACGCTATGGGTGGTCTGGTT	<i>used in origami L3, L3E and L3S</i>	<i>AACCAGACCCCATAGC</i>
B18_06	AAGGCCGGAGACATGTACCTCGCTATGGGTGGTCTGGTT		
E_02	TTAGAATCAGAGCGGG	staples for tip-to-tip blunt end interactions	
F_02	GCGGTTTGCGTATTG	<i>used in origami LE and L3E</i>	
E_03	AGCTAAACAGGAGGCC		
F_03	AACGCGGGGAGAG		
E_04	CCTGAGAAGTGTTTTATA		
F_04	GGGAAACCTGTCGTGC		
E_05	ATCAGTGAGGCCACCGAGT		
F_05	TGCCCGCTTTCAGTC		
E_07	TTAGTAATAACATCACTTG		
F_07	TAAAGCCTGGGGTGCC		
E_09	TACCGCCAGCCATTGC		
F_09	TGAAATTGTTATCCGCTCA		
E_12	GTAATAAAAGGGACATTCT		
F_12	TAAAACGACGGCCAGT		
E_13	GGCCAACAGAGATAGAACC		
F_13	CCCAGTCACGACGTTG		
E_14	CAGACAATATTTTGAATG		
F_14	TGTGCTGCAAGGCGAT		
E_15	GCTATTAGTCTTTAATGCG		
F_15	GCTGGCGAAAGGGGA		
E_17	GAAGATAAACAGAGG		
F_17	AGGCTGCGCAACTGTTGGG		
E_19	CTGAGAGCCAGCAGCA		
F_19	ACTCCAGCCAGCTT		
LS_00	TATATATTTAATTTACAATAGATAATACAT	staples for lateral oligomerization	
LS_01	TATATATTTAAGCAAAGCGCGCAGAGGCG	<i>used in origami LS and L3S</i>	
LS_02	TATATATTTCTACCGTGATCTTCTGACCT		
LS_03	TATATATTTACGGTATTAATAATCGGCTGT		
LS_04	TATATATTTAAGAATTAATAAACATAAAA		
LS_05	TATATATTTCCCGATTGATTACCAGCGCCA		
LS_06	TATATATTTCCGCCAGCATCAGAGCCGCCA		
LS_07	TATATATTTTCGGCCACCCATAGGTGTATCA		
LS_08	TATATATTTCTGATACCTCAGCTTGCTTT		
LS_09	TATATATTTATGCGCAGACCGGACCTGCT		
LS_10	TATATATTTAAAAATGCAGTCATCAGTTGAG		
LS_11	TATATATTTTCATTAGAGAGAACCAGACCGG		
LS_12	TATATATTTTGACTTTTGAATCGGTTGTAC		
LS_13	TATATATTTCTTGTTAAAACGTTAATATTT		
RS_00	TATATATTTAAACCCTCAATCTTAGAACAA		
RS_01	TATATATTTTTCGGTAGATTTAGAAGAGTT		
RS_02	TATATATTTTAAACCTCCGGAGAATATCA		
RS_03	TATATATTTATAACAACATGCCAGCTCC		
RS_04	TATATATTTAGAGCCTAATTTATAACGGAG		
RS_05	TATATATTTTATGTTAGCAAAAGCGTCATT		
RS_06	TATATATTTATTAGCGTTGCATAAACAAAT		
RS_07	TATATATTTGAAAGTATTAAGAGTAAATTC		
RS_08	TATATATTTAGCGGAGTGAGATAAACGGAA		
RS_09	TATATATTTAGAGGCAAAAGAAGTAGTAAA		
RS_10	TATATATTTTACCTTATGCGCCCTCAAAA		
RS_11	TATATATTTATCAGGTCTTTACGCAAAATCT		
RS_12	TATATATTTGAAAAGGTGGCAAGATCTAGA		
RS_13	TATATATTTGAATCGATGAACAGTTTGAGC		

**Table S2 | Fraction of deformed vesicles, upon increasing MgCl<sub>2</sub>, as a function of total L3E concentration.**

<b>[L3E] (nM)</b>	<b>% Deformed vesicles (± st. dev.)</b>	<b>Independent repeats</b>	<b>Total number vesicles (<math>N_{total}</math>)</b>
0.1	10.9 ± 10.1%	4	174
0.25	37.1 ± 20.9%	5	316
0.5	65.6 ± 12.1%	5	401
1	70.2 ± 9.6%	4	350

## Movie Captions

**Movie S1** | FRAP of 0.5 nM origami L3E (Alexa488-labelled, green) on top of DOPC SLB (doped with Atto655-DOPE, magenta), in the presence of a low  $\text{MgCl}_2$  buffer (5 mM  $\text{MgCl}_2$  + 300 mM NaCl). Corresponding data represented in Figure 3SA. Scalebar is 5  $\mu\text{m}$ .

**Movie S2** | Time-series of  $\text{MgCl}_2$ -triggered polymerization of 0.5 nM origami L3E (Alexa488-labelled, green) on top of DOPC SLB. Addition of  $\text{MgCl}_2$  happened at timepoint 5:00. Scalebar is 10  $\mu\text{m}$ .

**Movie S3** | FRAP of 0.5 nM origami L3E (Alexa488-labelled, green) on top of DOPC SLB (doped with DiD, magenta), in the presence of a high  $\text{MgCl}_2$  buffer (70 mM  $\text{MgCl}_2$  + 187.5 mM NaCl). Corresponding data represented in Figure 3SB. Scalebar is 5  $\mu\text{m}$ .

**Movies S4 & S5** | Diffusion of 0.1nM origami L3E (Alexa488-labelled, green) on the pole of GUVs (doped with Atto655-DOPE, magenta), after  $\text{MgCl}_2$ -mediated polymerization into membrane-bound end-to-end self-assembled filaments.

**Movie S6** | Diffusion of 0.1nM origami L3S (Alexa488-labelled, green) on the pole of GUV (doped with Atto655-DOPE, magenta), after  $\text{MgCl}_2$ -mediated polymerization into membrane-bound laterally self-assembled platforms.

**Movie S7** | Characteristic wrinkled membrane deformations on DOPC GUV (doped with Atto655-DOPE, magenta) induced by membrane-bound origami L3E (Alexa488-labelled, green) at 1 nM bulk concentration, after  $\text{MgCl}_2$ -mediated polymerization into filaments. Scalebar is 5  $\mu\text{m}$ .

**Movie S8** | Characteristic spike-like tubular deformations on DOPC GUV (doped with Atto655-DOPE, magenta) induced by membrane-bound origami L3E (Alexa488-labelled, green) at 1 nM bulk concentration, after  $\text{MgCl}_2$ -mediated polymerization into filaments. Scalebar is 5  $\mu\text{m}$ .