Dimerization and oligomerization of DNA-assembled building blocks for controlled multi-motion in high-order architectures

Ling Xin1,2, Xiaoyang Duan2 & Na Liu1,2✉

In living organisms, proteins are organized prevalently through a self-association mechanism to form dimers and oligomers, which often confer new functions at the intermolecular interfaces. Despite the progress on DNA-assembled artificial systems, endeavors have been largely paid to achieve monomeric nanostructures that mimic motor proteins for a single type of motion. Here, we demonstrate a DNA-assembled building block with rotary and walking modules, which can introduce new motion through dimerization and oligomerization. The building block is a chiral system, comprising two interacting gold nanorods to perform rotation and walking, respectively. Through dimerization, two building blocks can form a dimer to yield coordinated sliding. Further oligomerization leads to higher-order structures, containing alternating rotation and sliding dimer interfaces to impose structural twisting. Our hierarchical assembly scheme offers a design blueprint to construct DNA-assembled advanced architectures with high degrees of freedom to tailor the optical responses and regulate multi-motion on the nanoscale.
Cellular life functions through a collection of highly-controlled dynamic processes involving the self-assembly and organization of diverse molecular building blocks. These biological constructs optimized through billions of years of evolution provide us with the inspiration to create artificial systems, which emulate the structural and functional features of their natural counterparts. In particular, oligomerization is essential in many protein-involved cellular processes. For instance, a GTPase called dynamin is at the heart of endocytic vesicle fission. The dynamin unit is an antiparallel dimer, which can oligomerize into a helical polymer. The ratchet model proposes that GTP hydrolysis powers the relative sliding of the helical turns, giving rise to twisting of the helix and eventually membrane fission. Such a self-association mechanism through dimerization and oligomerization not only enables new functions at the intermolecular interfaces but also elicits a wealth of structural and functional advantages. This offers a blueprint to create mimics that can collectively operate to define functions in artificial machinery.

Among different state-of-the-art techniques, DNA nanotechnology1–7 represents a unique tool to build bio-inspired artificial systems, taking advantage of the precision, addressability, and programmability of DNA on the nanoscale. A variety of DNA-based dynamic devices8–21 that mimic motor proteins22 in living cells has been accomplished, including walkers23–30, rotors31–36, sliders37–40, and assembly lines41. Recently, efforts to constructing bio-inspired DNA-assembled systems are taking a step forward from monomeric to high-order structures42–53 and from simple to complex motion.58,59,54–57. Here, we demonstrate a DNA-assembled building block with rotary and walking modules, which can form dimeric structures to execute sliding, as well as subsequent higher-order architectures linked through dimer interfaces of the rotary and walking modules for controlled multi-motion. The monomeric building block is a chiral system that consists of two interacting gold nanorods (AuNRs) templated by DNA origami.50–63. The two AuNRs can carry out the rotation and walking, respectively. Through dimerization, the walking modules are merged into a sliding module, imposing relative movements between two building blocks. Further oligomerization leads to higher-order structures with alternating dimer interfaces for collective rotation and sliding, imposing structural twisting. Our work delineates one of the rudimentary steps towards self-assembled artificial structures that imitate the collective motion of protein complexes, with the ultimate goal to build and engineer cellular mimics with fully operational structural and functional features de novo. Although it is an incredible adventure, fortunately, we will be able to look at the solutions provided by nature for every problem we encounter.

Results

Structural design of the UNIT. Figure 1a shows the schematic of the DNA-assembled building block, which is composed of a rotary module and a walking module. It is named UNIT. The DNA origami template consists of a double-layer plate (42 nm × 41.2 nm), a 6-helix 90° arc-shaped track (~32 nm in radius), and a 10-helix rotary bundle (45 nm in length). The arc-shaped track is immobilized on the top surface of the plate, whereas one end of the rotary bundle is flexibly linked to the plate through adjacent scaffold crossovers (Supplementary Figs. 1 and 2). The rotary module contains a AuNR (yellow, 10 nm × 38 nm) anchored on the rotary bundle and the arched-shaped track dressed with five rotation footholds (r fh1–r fh5). A foot strand (Fr) is extended from the other end of the rotary bundle, 32 nm away from the flexible linker. The five rotation footholds spaced by 12.6 nm on the arc-shaped track define five rotation positions 1–5 (see Fig. 1a and b). The corresponding rotation angles for positions 1–5 are 15°, 37.5°, 60°, 82.5°, and 105°. The walking module contains an AuNR (blue, 10 nm × 38 nm) fully decorated with foot strands for walking (F w) and six rows of walking footholds (w fh1–w fh6) extended from the bottom surface of the plate. These six foothold rows define five walking positions I–V that are evenly separated by 7 nm (Fig. 1a, b, and Supplementary Fig. 2d). The walker AuNR is assembled on the plate through DNA hybridization between the F w and two neighboring rows of w fh. Figure 1c presents the transmission electron microscopy (TEM) image of the DNA origami template structures. Each structure contains three components: bundle, arc, and plate. Figure 1d shows the TEM images of the UNITs, in which the AuNRs are assembled on the individual DNA origami templates. Additional structural information is presented in Supplementary Figs. 1–3.

The reversible rotation and walking mechanisms are displayed in Fig. 1e and f, respectively. Two independent sets of DNA fuel strands are designed to power the stepwise rotation and walking through foothold-mediated strand displacement reactions, respectively (see Supplementary Figs. 4 and 5). The rotation process proceeds in a “release and capture” manner. Blocking (w fh1–w fh5) and removal (r fh1–r fh5) strands are utilized for activation and deactivation of r fh1–r fh5, respectively. As shown in Fig. 1e, the rotary bundle is first fixed at position 1 through DNA hybridization between Fr (black) and r fh1 (brown), while all other four r fhs are blocked. For rotation from position 1 to 2, r fh2 is activated and r fh1 is blocked upon addition of removal strands r fh2 and blocking strands r fh1, respectively. Then, Fr is hybridized with the activated r fh2. The rotary bundle clockwise rotates about 22.5° and reaches position 2. The walking process carries on through “adhesive rolling” and removal (w fh1–w fh6) and w fh1–w fh6 strands are utilized to control the accessibility of w fh1–w fh6. As shown in Fig. 1f, the walker AuNR first resides at the position I with F w bound to w fh1 (red) and w fh6 (blue). All other four rows of w fhs are blocked. For walking from position I to II, blocking strands w fhB and removal strands w fhC is added to block w fhA and activate w fhC, respectively. The walker AuNR moves forward to interact with the activated w fhC and subsequently reaches position II.

Optical characterizations of the UNIT motion. Altogether, five rotation positions and five walking positions establish 25 different states for the UNIT as depicted in Fig. 2a. The configuration changes among different states on the nanoscale give rise to the near-field coupling changes between the two AuNRs within the UNIT, which is a plasmonic chiral object in nature.58,59. These different states can be optically discriminated against by circular dichroism (CD) spectroscopy.59,64. The CD spectra (see Supplementary Fig. 6) at the 25 states have been measured using a Jasco 1500 CD spectrometer. Figure 2b shows the experimental CD intensities extracted at 721 nm. Figure 2c presents the theoretical results, which agree qualitatively well with the experimental results. Next, the dynamic transitions among different states are optically monitored. As shown in Fig. 2d, path 1 first follows a walking route and then a rotation route. Starting at state I, the walker AuNR strides from the left to the right on the origami plate and halts at state V1. Afterward, the rotary AuNR stepwisely rotates from state V1 to state V5. As presented by the dashed black curve in Fig. 2d, the time-course CD measurements reveal nine distinct states in the expected order. Subsequently, the walking and rotary modules that are operated alternatingly (path 2) and simultaneously (path 3) are examined. As shown in Fig. 2d, the time-course CD measurements along these two paths presented by solid black and gray curves exhibit a close
resemblance, indicating that the UNIT can carry out the rotation and walking motion in concert with high fidelity.

**Dimerization of the UNITs and motion of the DIMER.** Next, the dimer interface of the walking module is activated to create a new structure from two UNITs, called DIMER. As shown in Fig. 3a, wfh in the UNIT at state IV5 are activated by wRa. The interaction between wfh and the walker AuNR at position IV within the same UNIT is hindered, as their relative distance is as large as 19.5 nm. In contrast, the cooperative interaction between wfh in one UNIT and the walker AuNR in the other UNIT induces the crosslinking of the two UNITs in an antiparallel fashion, leading to the formation of a DIMER. Subsequently, wRb are added to stabilize the DIMER at state 5I-IV5. To evaluate the relative orientations between the UNITs within the individual DIMERs after crosslinking, gold nanoparticles (AuNPs, 10 nm) are utilized as markers for structural analysis by TEM. The detailed information is shown in Supplementary Figs. 7–9. Through dimerization, a new type of linear motion, which is conferred at the dimer interface. The working principle of reversible sliding is depicted in Fig. 3b. The two rotary modules in the DIMER can collectively carry out rotations along with two opposite directions with relative angle changes at 45° intervals, irrespective of the motion performed by the sliding module. For instance, as shown in Fig. 3b, starting at state 4I-IV4, the two walker AuNRs are bound in between two origami plates through the same combination of four wfh, in which wfh and wfhb are from one UNIT, as well as wfhc and wfhf are from the other UNIT. Blocking strands wBa and wBd are added to deactivate wfh and wfhc, respectively. The two walker AuNRs are bound to wfh and wfhc in an antiparallel manner to maintain the dimeric structure. Then, removal strands wRc and wRf are added to activate wfhc and wfhb, respectively. The Fw on each AuNR capture the newly activated wfh and the DIMER slides to state 4II-V4. For each sliding step, a 14 nm displacement between the two UNITs is introduced.

**Optical characterizations of the DIMER motion.** Five collective rotation positions and two sliding positions give rise to 10 distinct states of the DIMER. The working principle of one-step rotation from position 1 to 2 and one-step walking from position I to II are shown in Figs. 3e and 3f, respectively. Through dimerization, a new type of linear motion, which is coordinated sliding between the two walking modules, is...
distinct states for the DIMER as depicted in Fig. 4a. Figure 4b (see also Supplementary Fig. 10) shows the TEM images at the representative states 4I-IV4 and 4II-V4, respectively. Four AuNRs are clearly visible in the individual structures. The configurations of the DIMERs are deformed due to the drying process of the structures on the TEM grids. Figure 4c presents the experimental and theoretical CD intensities extracted at 721 nm at these 10 states, which show a good agreement. The experimental CD spectra are presented in Supplementary Fig. 11. The details of CD calculations and the influence of the parallel dimers on CD can be found in Supplementary Fig. 12. The DIMER can be regulated along different routes through independent (path 1) or simultaneous (path 2) control of the rotary and sliding modules as demonstrated by the time-course measurements in Fig. 4d. Notably, the CD intensities and kinetics at the common states along the two paths show close resemblance. This proves that the DIMER can execute as well-controlled motion as the UNIT does.

Oligomerization of the UNITs and regulation of the OLIGOMERS. To generate higher-order architectures that are named OLIGOMERS, the UNITs are oligomerized by activating the dimer interfaces of both the rotary and walking modules. Here, UNIT and DIMER without the anchored rotary AuNRs are named UNITO and DIMERO, respectively. As shown in Fig. 5a, dimerization of the rotary modules is enabled through the functionalization of complementary DNA strands on the rotary bundles in UNITO and cUNITO. UNITO and cUNITO are mixed in a 1:1 ratio and the intermediate dimeric structure (rDIMER) is depicted in Fig. 5a. The TEM images of the rDIMERS and DIMEROs are presented in Supplementary Fig. 13. Together with dimerization of the walking modules, OLIGOMERS are formed, containing helical queues of rDIMERS and helical queues of DIMEROs. The OLIGOMERS can reconfigure from state O-5I-IV5 to state O-3II-V3, and then to O-2I-IV2 through simultaneous rotation and sliding processes (see Supplementary Fig. 14). Figure 5b presents the TEM images of the OLIGOMERS at these
three states in different views. The side views show linear-like structures in various lengths, resulting from different degrees of oligomerization (see Supplementary Fig. 15). Circular structural profiles in different diameters (d) are observed in the top views. In response to the specific DNA fuels, in an OLIGOMER, the collective motion of the sliding modules results in the relative displacements between the adjacent rDIMERS, while the collective motion of the rotary modules leads to the relative twisting between the adjacent DIMERS. As the rotation angle between two adjacent DIMERS within the OLIGOMER at state O-5I-IV5 is 30°, 12 DIMERS complete one full helical turn (see Supplementary Fig. 14b). From position 5 to 3 and then to 2, the relative twisting leads to the decrease of the number of DIMERS per turn from 12 to 7 and to 3.3 accordingly (Fig. 5b and Supplementary Fig. 14b). As a result, the OLIGOMER shrinks its lateral dimension and becomes more compact. The diameter (d) of the OLIGOMER is also influenced by the sliding modules, as the two sliding positions in each DIMER correspond to different relative lateral distances between the adjacent rDIMERS (see Supplementary Fig. 14b). Additional schematics and TEM images of the OLIGOMERs in three-dimensional views are shown in Supplementary Fig. 16. Time-course CD measurements of the dynamic transitions among the three states are presented in Supplementary Fig. 17.

**Discussion**

In this work, we have demonstrated dimerization and oligomerization of DNA-assembled building blocks for controlled multimotion in high-order architectures. The monomeric UNIT contains walking and rotary modules. Activation of the walking...
dimer interface enables the formation of a DIMER from two UNITs and meanwhile introduces the sliding module. Activation of the dimer interfaces of both the walking and rotary modules leads to the formation of OLIGOMERs, which can undergo collective rotation and sliding processes. The collective motion also enforces relative twisting within the structures and leads to varied lateral dimensions. The hierarchical assembly through activation of different dynamic dimer interfaces provides a pathway to achieve complex and controlled multi-motion in high-order structures.

Methods

Materials. Single-stranded scaffold DNA (p8064) was purchased from Thelitib nanosystems. Staple, blocking, and removal strands were purchased from Sigma-Aldrich. Agarose for electrophoresis and SYBR Gold nucleic acid stain were purchased from Life Technologies. Uranyl formate for negative TEM staining was purchased from Polysciences, Inc.

Design and assembly of the DNA origami template structures. The UNIT was designed using caDNAno software. It consisted of a 10-helix bundle, a 6-helix arc, and a double-layer plate arranged in a 'honeycomb' lattice, which was connected through scaffold crossovers (see Supplementary Fig. 1 and Supplementary Dataset 1). The DNA origami was assembled from 10 nM scaffold strands and 100 nM of each of the staple strands in a 0.5× TE buffer with 12 mM MgCl₂ and 5 mM NaCl using a 20 h annealing program (85 °C 5 min, 70 °C–61 °C–1 °C/min, 60 °C–51 °C–1 °C/1h, 50 °C–22 °C–1 °C/20 min, and held at 15 °C). The DNA origami structures were purified by 0.7% agarose gel electrophoresis in a 0.5× TBE buffer with 11 mM MgCl₂ for 3 h at 8 V/cm.

DNA functionalization of the AuNRs and AuNPs, as well as assembly of the UNITs. AuNRs were purchased from Sigma-Aldrich (Cat no. 716812). Functionalization of the AuNRs with thiolated DNA was carried out following a low pH procedure. Basically, thiolated DNA strands were incubated with TCEP (tris(carboxymethyl)phosphine) for 2 h. The ratio of DNA: TCEP was 1:200. Second, the AuNRs (1 nM, 1 mL) were spun down and the supernatant was removed. The AuNRs were then mixed with thiolated DNA strands (100 μM, 30 μL). In total, 845 μL modification buffer (0.59× TBE (tris-(hydroxymethyl)-aminomethane, borate, ethylenediaminetetraacetic acid, 0.023% SDS, pH = 3) was added. 10 mL, 5 M NaCl was added every 10 min for nine times. 10 mL, 5 M NaOH was added subsequently to adjust the pH value to ~8, and the final concentration of NaCl reached 0.5 M. The AuNRs functionalized with DNA were then purified by centrifugation. Five times of centrifugations at a rate of 8000×g for 30 min were carried out. Each time, the supernatant was carefully removed and the AuNRs were resuspended in a 0.5× TBE buffer containing 0.02% of SDS. The supernatant was then removed. AuNPs were purchased from Sigma-Aldrich (Cat no. 741957). Functionalization of the AuNPs with thiolated DNA was carried out following a well-established procedure. Basically, after reduced with TCEP, thiol-modified DNA and BSPP-modified AuNPs were incubated at a molar ratio of DNA: AuNPs of 300:1 in a 0.5× TBE buffer solution overnight at room temperature. The concentration of NaCl was slowly increased to 500 mM in the subsequent 24 h. The AuNPs functionalized with DNA were then washed using a 0.5× TBE buffer solution in 100 kDa centrifuge filters to remove the free DNA strands. DNA-functionalized AuNRs or AuNPs were mixed with the DNA origami structures in a ratio of 10:1. The mixtures were annealed with the procedure of 35 °C 2.5 h, 34 °C–29 °C–1 °C/3 h, 28–26 °C–1 °C/1.5 h, and held at 25 °C. Gel electrophoresis was used to purify the UNITs under the same condition for the purification of the DNA origami template structures.

Dimerization of the UNITs. 250 μM 0.4 μL whfa strands were added in the purified UNITs (1 nM, 100 μL) and the mixture was annealed with the procedure of...
Fig. 5 Oligomerization of the UNITs for high-order structures. 

**a** Schematic of the oligomerization of the UNITs. The rotary bundles in UNIT₀ and cUNIT₀ are functionalized with complementary strands (10 olive strands in UNIT₀ and 10 dark gray strands in cUNIT₀). Both UNIT₀ and cUNIT₀ are at state IV5. The intermediate dimeric structure is called rDIMER. The final OLIGOMER is at state O-5I-IV5, containing both helical queues of rDIMERs and DIMEROs.

**b** Regulation of the OLIGOMERs. Schematics and TEM images of the side views and top views of the OLIGOMERs at different states. The OLIGOMERs reconfigure from state O-5I-IV5 to O-3II-V3, and then to state O-2I-IV2 through simultaneous rotation and sliding processes.
CD characterizations. The CD spectra were measured using a Jasco-1500 CD spectrometer with a quartz suprasil cuvette (path length, 10 mm). All measurements were carried out at 25 °C. The concentrations of the blocking and removal strands were 250 μM.

For the CD spectral measurements in Fig. 2b, one sample at state I13 was divided into 25 copies. Each copy was 60 μL and the respective DNA fuels were added to drive the individual systems to their designated positions (Supplementary Table 1). To keep the concentration constant, an equal volume of H2O was added, if no DNA strands were added. After each addition, the samples were incubated at 25 °C for about 1 h. The final concentrations of the AuNRs in all the systems were about 1 nM.

The time-course measurements for the UNITs were carried out as follows. Two hundred microliter was used for each UNIT sample. The CD signals at 721 nm were monitored using the time-scan acquisition mode with a data pitch of 1 s. The respective blocking and removal strands were added to enable programmed routes (Supplementary Tables 2 and 3). The initial concentrations of the AuNRs were about 1 nM. After the process, the volume of the sample was increased by about 24.4 μL (11.6%).

For the CD spectral measurements in Fig. 4b, one sample at state SI-IVS was divided into 10 copies. The same experimental protocol as that for the UNIT was conducted and the respective DNA fuels were added to drive the individual systems (Supplementary Table 4). After each addition, the samples were incubated at 25 °C for about 1 h. The final concentrations of the AuNRs in all the systems were about 0.2 nM.

The time-course measurements for the DIMERs were carried out as follows. Two hundred microliter was used for each DIMER sample. The CD signals at 721 nm were monitored at the same conditions as those of the UNITs. The respective blocking and removal strands were added to enable programmed routes (Supplementary Table 5). The initial concentrations of the AuNRs were about 0.10 nM. After the process, the volume of the sample was increased by about 26.2 μL (11.6%).

Oligomerization of the UNITs and regulations of the OLIGOMERS. The purified DNA origami template for UNIT1 and cUNIT1 were mixed in a ratio of 1:1. The walker AuNRs and the mixture were annealed with the procedure of 35 °C–2.5 h, 34 °C–29 °C–1 °C/3 h, 28–26 °C–1 °C/1.5 h, and held at 25 °C. Gel electrophoresis was used to purify the rDIMERs under the same condition for the DNA origami template structure. The dimerization yield was 68.7% (see Supplementary Fig. 18).

Table 1). To keep the concentration constant, an equal volume of H2O was added, if no DNA strands were added. After each addition, the samples were incubated at 25 °C for about 1 h. The final concentrations of the AuNRs in all the systems were about 1 nM.

The time-course measurements for the UNITs were carried out as follows. Two hundred microliter was used for each UNIT sample. The CD signals at 721 nm were monitored using the time-scan acquisition mode with a data pitch of 1 s. The respective blocking and removal strands were added to enable programmed routes (Supplementary Tables 2 and 3). The initial concentrations of the AuNRs were about 1 nM. After the process, the volume of the sample was increased by about 24.4 μL (11.6%).

For the CD spectral measurements in Fig. 2b, one sample at state I13 was divided into 25 copies. Each copy was 60 μL and the respective DNA fuels were added to drive the individual systems to their designated positions (Supplementary Table 1). To keep the concentration constant, an equal volume of H2O was added, if no DNA strands were added. After each addition, the samples were incubated at 25 °C for about 1 h. The final concentrations of the AuNRs in all the systems were about 1 nM.

The time-course measurements for the UNITs were carried out as follows. Two hundred microliter was used for each UNIT sample. The CD signals at 721 nm were monitored using the time-scan acquisition mode with a data pitch of 1 s. The respective blocking and removal strands were added to enable programmed routes (Supplementary Tables 2 and 3). The initial concentrations of the AuNRs were about 1 nM. After the process, the volume of the sample was increased by about 24.4 μL (11.6%).

For the CD spectral measurements in Fig. 4b, one sample at state SI-IVS was divided into 10 copies. The same experimental protocol as that for the UNIT was conducted and the respective DNA fuels were added to drive the individual systems (Supplementary Table 4). After each addition, the samples were incubated at 25 °C for about 1 h. The final concentrations of the AuNRs in all the systems were about 0.2 nM.

The time-course measurements for the DIMERs were carried out as follows. Two hundred microliter was used for each DIMER sample. The CD signals at 721 nm were monitored at the same conditions as those of the UNITs. The respective blocking and removal strands were added to enable programmed routes (Supplementary Table 5). The initial concentrations of the AuNRs were about 0.10 nM. After the process, the volume of the sample was increased by about 26.2 μL (11.6%).

Oligomerization of the UNITs and regulations of the OLIGOMERS. The purified DNA origami template for UNIT1 and cUNIT1 were mixed in a ratio of 1:1. The walker AuNRs and the mixture were annealed with the procedure of 35 °C–2.5 h, 34 °C–29 °C–1 °C/3 h, 28–26 °C–1 °C/1.5 h, and held at 25 °C. Gel electrophoresis was used to purify the rDIMERs under the same condition for the DNA origami template structure. The dimerization yield was 68.7% (see Supplementary Fig. 18).

Table 1). To keep the concentration constant, an equal volume of H2O was added, if no DNA strands were added. After each addition, the samples were incubated at 25 °C for about 1 h. The final concentrations of the AuNRs in all the systems were about 1 nM.

The time-course measurements for the UNITs were carried out as follows. Two hundred microliter was used for each UNIT sample. The CD signals at 721 nm were monitored using the time-scan acquisition mode with a data pitch of 1 s. The respective blocking and removal strands were added to enable programmed routes (Supplementary Tables 2 and 3). The initial concentrations of the AuNRs were about 1 nM. After the process, the volume of the sample was increased by about 24.4 μL (11.6%).

For the CD spectral measurements in Fig. 4b, one sample at state SI-IVS was divided into 10 copies. The same experimental protocol as that for the UNIT was conducted and the respective DNA fuels were added to drive the individual systems (Supplementary Table 4). After each addition, the samples were incubated at 25 °C for about 1 h. The final concentrations of the AuNRs in all the systems were about 0.2 nM.
42. Hong, F., Zhang, F., Liu, Y. & Yan, H. DNA origami: scaffolds for creating higher order structures. Chem. Rev. 117, 12584–12640 (2017).

43. Lan, M., et al. Programmable super-assembly of a DNA surface adapter for tunable chiral directional self-assembly of gold nanorods. Angew. Chem. Int. Ed. Engl. 56, 14632–14636 (2017).

44. Ong, L. L. et al. Programmable self-assembly of three-dimensional nanostructures from 10,000 unique components. Nature 552, 72–77 (2017).

45. Song, J. et al. Reconfiguration of DNA molecular arrays driven by information relay. Science 357, eaan3377 (2017).

46. Tikhonov, G., J. & Qian, L. Fractal assembly of micrometre-scale DNA origami arrays with arbitrary patterns. Nature 552, 67–71 (2017).

47. Wagenbauer, K. F., S. & Dietz, H. Gigadalton-scale shape-programmable DNA assemblies. Nature 552, 78–83 (2017).

48. Zhang, Z., Yang, V., Pincet, F., Llaguno, M. C. & Lin, C. Placing and shaping liposomes with reconfigurable DNA nanocages. Nat. Chem. 9, 653–659 (2017).

49. Hen, X. et al. DNA-guided plasmonic helix with switchable chirality. Annu. Rev. Phys. Chem. 70, 11763–11770 (2018).

50. Wang, M. et al. Reconfigurable plasmonic diastereomers assembled by DNA origami. ACS Nano 13, 13702–13708 (2019).

51. Sun, W. et al. Precise pitch-scaling of carbon nanotube arrays within three-dimensional DNA nanotrenches. Science 368, 874–877 (2020).

52. Wickham, S. F. J. et al. Complex multicomponent patterns rendered on a 3D DNA-barrel pegboard. Nat. Commun. 11, 5768 (2020).

53. Yao, G. et al. Meta-DNA structures. Nat. Chem. 12, 1067–1075 (2020).

54. Thuenger, A. J. et al. A cargo-sorting DNA robot. Science 357, eaan6558 (2017).

55. Zhou, L., Marras, A. E., Huang, C. M., Castro, C. E. & Su, H. J. Paper origami-inspired design and actuation of DNA nanomachines with complex motions. Small 14, e1802580 (2018).

56. Benn, F. et al. Chiral DNA origami nanotubes with well-defined and addressable inside and outside surfaces. Angew. Chem. Int. Ed. Engl. 57, 7687–7690 (2018).

57. Urban, M. J., Zhou, C., Duan, X. & Liu, N. Optically resolving the dynamic walking of a plasmonic walker couple. Nano Lett. 15, 8392–8396 (2015).

58. Hentschel, M., Schäferling, M., Duan, X., Giessen, H. & Liu, N. Chiral plasmonics. Sci. Adv. 3, e1602735 (2017).

59. Urban, M. J. et al. Chiral plasmonic nanostructures enabled by bottom-up approaches. Annu. Rev. Phys. Chem. 70, 305–329 (2019).

60. Rothemund, P. W. Folding DNA to create nanoscale shapes and patterns. Nature 440, 297–302 (2006).

61. Dietz, H., Douglas, S. M. & Shih, W. M. Folding DNA into twisted and curved nanoscale shapes. Science 325, 725–730 (2009).

62. Douglas, S. M. et al. Self-assembly of DNA into nanoscale three-dimensional shapes. Nature 459, 414–418 (2009).

63. Pinheiro, A. V., Han, D., Shih, W. M. & Yan, H. Challenges and opportunities for structural DNA nanotechnology. Nat. Nanotechnol. 6, 763–772 (2011).

64. Yin, X., Schäferling, M., Metzger, B. & Giessen, H. Interpreting chiral nanophotonic spectra: the plasmonic Born–Kuhn model. Nano Lett. 13, 6238–6243 (2013).

65. Douglas, S. M. et al. Rapid prototyping of 3D DNA-origami shapes with caDNAno. Nucleic Acids Res. 37, 5001–5006 (2009).

Acknowledgements
This project was supported by the European Research Council (ERC Dynamic Nano) grant and the Max Planck Society (Max Planck Fellow). We thank Marion Kelsch for assistance with TEM. TEM images were collected at the Stuttgart Center for Electron Microscopy.

Author contributions
L.X. and N.L. conceived the concept. L.X. designed the DNA origami template and the working principles. L.X. performed the experiments. X.D. carried out the theoretical calculations and figure illustrations. L.X. and N.L. wrote the paper. All authors discussed the results, analyzed the data, and commented on the paper.

Funding
Open Access funding enabled and organized by Projekt DEAL.

Competing interests
The authors declare no competing interests.

Additional information

Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41467-021-23532-y.

Correspondence and requests for materials should be addressed to N.L.

Peer review information Nature Communications thanks Velkko Linko and the other anonymous reviewers for their contribution to the peer review of this work. Peer reviewer reports are available.

Reprints and permission information is available at http://www.nature.com/reprints

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.