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Transfer of multi-DNA patches by colloidal stamping[†]

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Patchy particles have received great attention due to their ability to develop directional and selective interactions and serve as building units for the self-assembly of innovative colloidal molecules and crystalline structures. Although synthesizing particles with multiple dissimilar patches is still highly challenging and lacks efficient methods, these building blocks would open paths towards a broader range of ordered materials with inherent properties. Herein, we describe a new approach to pattern functional DNA patches at the surface of particles, by the use of colloidal stamps. DNA inks are transferred only at the contact zones between the target particles and the stamps thanks to selective strand-displacement reac-tions. The produced DNA-patchy particles are ideal candidates to act as advanced precision/designer building blocks to self-assemble the next generation of colloidal materials.

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Introduction

Over the past few decades, scientists have aspired to fabricate functional materials by colloidal self-assembly.¹ Although many beautiful examples of self-assembled colloidal molecules² or colloidal crystals³ from particles with well-defined shapes and compositions have been reported so far, colloidal systems cannot be targeted towards most of the sophisticated structures that nature has built. Indeed, the latter requires encoding the building units with information to guide their self-assembly by programming their geometry as well as their directionality, valence, and range of pairing interactions. Several strategies have been developed to address this challenge, including the attachment of molecules that recognize one another on the surface of particles.⁴ Among the wide range of binding groups that have been employed, synthetic DNA strands have been proven to be very versatile and promising as a tremendous number of orthogonal interactions can be programmed based on the design of nucleotide sequences, giving access to highly specific programmable interactions. DNA-coated particles have thus been extensively employed as

building blocks for the self-assembly of clusters with precise symmetries⁵ and crystalline lattices.⁶ To further control both the valence of the particles and the directionality of the bonds they form with their partners, a number of groups have recently proposed strategies to regioselectively pattern particles with DNA patches.⁷ Sleiman *et al.* successfully transferred DNA motifs from a parent 3D DNA template to gold⁸ and polymeric nanoparticles.⁹ Two-¹⁰ and three-dimensional¹¹ DNA origami structures were used as stamping platforms to transfer DNA inks onto gold nanoparticles. In both cases, the printed nanoparticles were released from the frame by a strand-displacement reaction.¹²

In order to create micron-sized particles with several dissimilar patches, we translated this strategy by using colloidal particles coated with DNA inks as stamps. The colloidal stamps can assemble with support colloids *via* DNA hybridization. The injection of eject strands allowed us to transfer the DNA inks at the contact zones between the support and stamp particles (Fig. 1a) leading to patchy particles. We also took advantage of packing constraints to control the number of stamp particles that can park around support particles,^{5a} which finally defines the number of transferred patches.

Experimental

Synthesis of DNA-coated particles

Azidated 3-(trimethoxysilyl)propyl methacrylate (TPM) particles were prepared through the azidation of chlorine groups present at the surface of particles previously synthesized according to the protocol developed by Wang *et al.*^{6e} (see the

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Fig. 1 General scheme and key steps. (a) General schematic representation of the preparation of particles with DNA patches by colloidal stamping following sequentially the subsequent steps. (b) Inking: the ink, formed by the association of $T-X-A^*$ and X^*-B^* , is hybridized at the surface of the bare stamp particle decorated with an A^* DNA brush. (c) Assembly: in the contact zone, formation of duplexes between the support particle decorated by B strands and the stamp thanks to the B* domain exposed by the ink. (d) Passivation: the surface of the support particle, outside of the contact zone is passivated by hybridization between the B strands of the surface and Y^*-B^* strands. (e) Eject: strand displacement reaction to separate the stamp from the support particle, leading to the formation of an X^*-B^* patch at the contact zone and the recovery of patchy particles exposing the stand Y^* on the surface with patches of X^* . (f) Schematic representation of the DNA strands and assemblies used in this study.

ESI[†]). After synthesis the particles were imaged by TEM and SEM. Fig. S1[†] shows that they are spherical and monodisperse in size. Their surface is smooth, which has been shown to be required to allow homogeneous distribution of DNA strands during the former step.13 In order to functionalize monodisperse polystyrene (PS) particles with azide groups, we followed the protocol developed by Oh et al.,^{6d} which relies on the physical entrapment of an azidated PS-b-PEO copolymer (PS-b-PEO-N₃). For this, the PS particles are swollen with tetrahydrofuran (THF) to allow the PS block of the copolymer to penetrate the PS particles (see the ESI[†]). After the evaporation of THF, the PS block of the copolymer is physically trapped in the PS particles while the PEO block and the terminal azide group form a brush at the surface, swollen by water and exposed to the surrounding media. Azidated particles were further functionalized with DNA following the protocol described by Wang et al.^{6e} that ensures a dense surface coverage of the colloids with DNA (see the ESI[†]). The process relies on the strain-promoted azide-alkyne cycloaddition (SPAAC) to graft DNA strands end-functionalized with a dibenzocyclooctyne moiety (DBCO) onto azide functionalized particles.

Characterization

Transmission electron microscopy (TEM) images were taken using a Hitachi H600 microscope operating at an acceleration voltage of 75 kV. The samples for TEM observation were supported on conventional carbon-coated copper grids. Scanning electron microscopy (SEM) images were taken using a Hitachi S4500 microscope at an accelerating voltage of 5 kV. Confocal fluorescence microscopy images were taken using a Leica SP2 confocal laser scanning microscope as well as a ZEISS LSM980 equipped with an Airyscan detector.

Results and discussion

DNA strands A and B (Table S1[†]) have first been grafted onto azidated TPM and PS particles, respectively. The coated particles are referred to as TPMA and PSB. We then functionalized/ inked the stamp TPMA particles with the Ink₅₆₅ (see Table S1⁺ for details) by adding a large excess of ink to a suspension of particles (Fig. 1b). The ink consists of two hybridized DNA strands, T-X-A* and X*565-B*, the latter being modified with the fluorescent dye Atto565. As the domain A* is complementary to sequence A at the surface of the TPM particles, the ink sticks to their surface due to the formation of A/A* duplexes. The particles and strands were maintained in a buffer enriched in magnesium ions at low temperature in order to strengthen the DNA duplexes and prevent strand migration (see ESI[†]). Excess ink was subsequently washed away by centrifugation/dispersion steps. As Ink₅₆₅ is also complementary to strands B, the $TPM_A \sim Ink_{565}$ stamp particles were mixed with a 40:1 excess of PS_B particles to form preferentially small clusters with only one TPMA particle at the core and PSB satellites (Fig. 1c). This ultimately maximizes the number of onepatch PS particles produced in the process. When the assembly is completed, Y*488-B* strands (labelled with Alexa488) are added to hybridize with the remaining B strands available on



Fig. 2 Confocal fluorescence microscopy images of clusters obtained by incubating 1.6 μ m TPM_A ~ Ink₅₆₅ and 1.5 μ m PS_B particles in a 1:40 ratio (a) before and (b) after purification. (c)–(i) Zoom images of the different clusters containing 1 to 7 PS particles. Their relative amounts (in %) in the sample before/after purification are given at the top right corner of each image. (j) Confocal fluorescence microscopy image of the patchy PS particles resulting from the strand-displacement disassembly of TPM_A ~ Ink₅₆₅ ~ PS_B clusters. Scale bar of 10 μ m for (a), (b) and (j), and 5 μ m for (c) to (i).

the PS_B particles and passivate their surface (Fig. 1d). Fig. 2a shows that clusters made of one TPM_A particle surrounded by PS_B particles were obtained. A detailed analysis reveals that the number of PS_B particles in the clusters varies from 1 to 7. Their relative amounts have been determined by a statistical analysis performed over 100 clusters (Fig. 2c-i). Some rare clusters (\sim 5%) formed from one PS_B particle in contact with two TPMA particles could be observed as well as a large amount of free PS_A particles. Thanks to the density difference between TPM (1.2 g cm⁻³ (ref. 14)) and PS (1.06 g cm⁻³), we successfully removed most of these free PS particles (Fig. 2b) by sedimentation in a PBS-based buffer solution of intermediate density (1.07 g cm^{-3}) prepared by mixing H₂O and D₂O (see ESI[†]). The relative proportions of different clusters remained unchanged during this purification stage, as indicated in Fig. 2c-i, proving that the clusters are sufficiently robust and do not break during centrifugation. The final step to form patchy particles consists of disassembling the clusters formed by the stamp and support particles leaving the fluorescent part of the ink on the support particle only at the contact point between them. To do so, we injected the $Eject_X$ strand, which binds to the toehold T of Ink₅₆₅ and replaces the strand X*₅₆₅-B*. This breaks the duplex X/X* that was holding the stamp and support particles and results in the release of the support particle, which still carries the red fluorescent strands X*₅₆₅-B* at the former contact point with the stamp (Fig. 1e and S2[†]). Fig. 2j and Movie S1[†] show that PS particles with one red fluorescent patch are mostly obtained, validating the developed strategy.

Some non-patchy PS particles that were not completely removed by centrifugation, and a few two-patch particles (\sim 5%), which result from the disassembly of the clusters in which one PS particle is in contact with two TPM particles, are also observed.

We firstly extended our strategy to prepare particles with multiple identical patches precisely located at their surface. To do so, we prepared clusters with different controlled morphologies by the random parking^{5*a*} of an excess of large PS spheres functionalized with DNA strands A and Ink₅₆₅ on smaller TPM particles functionalized with DNA strands B. Due to packing constraints inherent in the ratio of radii between large and small spheres, only a fixed number of large spheres can park, leading to a population of clusters with a well-defined coordination. When the assembly is completed, the Y*₄₈₈–B* strands, complementary to DNA strands B, are added to passivate the surface of TPM_B particles. Fig. 3a–c show confocal images of the DNA-colloidal clusters obtained when TPM particles with diameters of 1.1, 1.6, and 2 μ m are employed, respectively. Different clusters made of one TPM core and different numbers of PS satellites are observed (Fig. 3d–g and S3†). The relative proportions of each type of cluster obtained



Fig. 3 Confocal fluorescence microscopy images of the clusters obtained by incubating TPM_B and PS_A ~ lnk_{565} in a 1:40 ratio followed by the addition of the Y*₄₈₈–B* passivation strand. The diameters of TPM particles are: (a) 1.1 µm (α = 4.39), (b) and (d)–(g) 1.6 µm (α = 3.02), and (c) 2 µm (α = 2.42). (h) Confocal fluorescence microscopy image (Alexa488, green channel; Atto565, red channel) with transmission microscopy (grey channel) of PS_A ~ lnk_{565} ~ TPM_B and Y*₄₈₈–B* (α = 3.02) clusters after the strand-displacement reaction using the Eject_X strand. (i)–(l) Zoom images of the patchy TPM particles with an increasing number of patches obtained after the strand-displacement reaction using Eject_X (α = 3.02). Scale bar of 10 µm for (a) to (h) and 2 µm for (i) to (l).

Table 1 Compositions of the batches resulting from the mixing of TPM_B of different sizes with PS_A ~ Ink_{565} in a 1:40 number ratio determined by statistical analysis of confocal fluorescence images over about 100 clusters

α	•	•	000	*	\$
4.39	3	39	50	8	0
3.02	0	20	29	44	7
2.42	0	3	27	39	31

for different values of the size ratio α of PS/TPM, are listed in Table 1. One can note that when α is 4.39, clusters made of one or two PS particles attached to one TPM sphere are mainly formed. Decreasing α to 3.02 and 2.42 led to the formation of higher proportions of clusters containing 3 and 4 PS particles, as expected.

After the injection of the eject strands Eject_X in the cluster suspension, non-fluorescent PS particles and TPM particles with red fluorescent patches are observed, proving the transfer of the fluorescent DNA from PS_A ~ Ink₅₆₅ (Fig. 3h and S3†). More precisely, Fig. 3i–l show that 1.6 µm TPM particles with one to four patches are obtained. Similar results were obtained with 1.1 µm and 2 µm TPM particles (Fig. S5†), validating the strategy based on the combination of colloidal parking and colloidal stamping.

Finally, we further extended our strategy to prepare particles with multiple dissimilar patches. We first divided the PS particles into two batches, and coated one batch with Ink_{565} and the other with Ink_{647} (Table S1†). The two batches were then mixed together and TPM_B particles functionalized with B strands were added in a number ratio $TPM_B: PS_A \sim Ink_{565}: PS_A \sim Ink_{647}$ of 1:20:20. The sample was kept in the fridge at 4 °C for 24 h to maximize the formation of clusters. Then, Y^*_{488} -B* strands were added to hybridize with the B strands outside the contact zones and passivate the surface of the TPM_B particles.

Different clusters made of one TPM core and different numbers of $PS_A \sim Ink_{565}$ and $PS_A \sim Ink_{647}$ are observed (Fig. 4a-j). The relative proportions of each type of cluster are listed in Table 2. After injection of the eject strands $Eject_x$ and E_{ij} Eject_Z in the suspension of clusters, non-fluorescent PS particles and TPM particles with red and/or blue fluorescent patches are observed, proving the transfer of the fluorescent DNA from $PS_A \sim Ink_{565}$ and $PS_A \sim Ink_{647}$ (Fig. 4k and S6 and Movie S2[†]). When we worked with three batches of PS_A particles coated with Ink488, Ink565 and Ink647, respectively, and mixed them with TPM_B particles in a number ratio TPM_B: PS_A $\sim~Ink_{488}\!:\!PS_A~\sim~Ink_{565}\!:\!PS_A~\sim~Ink_{647}$ of 1:13:13:13, we observed the formation of a few clusters made of one TPM particle surrounded by varying numbers of $PS_A \sim Ink_{488}$, $PS_A \sim$ Ink_{565} and $PS_A \sim Ink_{647}$ particles (Fig. 41). After the injection of $Eject_X$, $Eject_Y$ and $Eject_Z$, TPM particles with red and/or blue and/or green fluorescent patches are observed, proving once again the efficiency of our approach (Fig. 4m and Movie S3[†]).



Fig. 4 Confocal fluorescence microscopy images (a–j) of the clusters obtained by incubating TPM_B, PS_A ~ Ink₅₆₅ and PS_A ~ Ink₆₄₇ in a 1:20:20 number ratio followed by the addition of the passivation strand Y*₄₈₈–B*. The diameter of the TPM particles is 1.6 µm (α = 3.02). (k) Confocal fluorescence microscopy image of the patchy particles obtained after injection of Eject_x and Eject_z. The inset shows a TPM particle with one red and one blue fluorescent patch. (l) Confocal fluorescence microscopy image of one TPM particle surrounded by one PS_A ~ Ink₄₈₈, one PS_A ~ Ink₅₆₅ and one PS_A ~ Ink₆₄₇ particle. (m) Confocal fluorescence microscopy image of the patchy 1.6 µm TPM particles obtained after injection of Eject_x, Eject_y and Eject_z. Scale bar of 10 µm for (a), 5 µm for (b) to (m), and 2 µm for (k) in the inset.

Table 2 Compositions of the batches resulting from the mixing of TPM_B with PS_A \sim Ink₅₆₅ and PS_A \sim Ink₆₄₇ in a 1:20:20 number ratio determined by statistical analysis of confocal fluorescence images over about 100 clusters



Conclusions

In conclusion, we have synthesized micron-sized particles with one or several identical and distinct DNA patches by combining colloidal parking and the transfer of DNA strands at the contact zones with colloidal stamps thanks to stranddisplacement reactions. Our strategy is versatile and can be extended to a gallery of hard particles and soft systems whose shape and size could be independently varied, opening the way to the synthesis of new DNA-patchy building blocks and the comprehensive study of their assembly into novel structures, such as alternating polymers or rings, dendrimers or gyroid crystals.

Author contributions

R. Mérindol, E. Ducrot and S. Ravaine conceived this research. R. Mérindol and E. Ducrot designed the experimental process and revised the manuscript. R. Khalaf performed most of the experiments. A. Viamonte helped R. Khalaf to perform some of the experiments. R. Khalaf, E. Ducrot and S. Ravaine performed data analysis. S. Ravaine wrote the manuscript.

Conflicts of interest

There are no conflicts to declare.

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