Functionalized DNA nanostructures as scaffolds for guided mineralization†

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The field of DNA nanotechnology uses synthetic DNA strands as building blocks for designing complex shapes in one-, two- and three-dimensions. Here, we investigate whether DNA nanostructures are feasible platforms for the precise organization of polyaspartic acid (pAsp), a known mineral carrier, with a goal towards biomimetic mineralization for enamel regeneration. We describe the preparation of DNA–pAsp conjugates and their subsequent assembly into ordered nanostructures. Covalent attachment of pAsp to DNA was noted to hinder DNA nanostructure formation past a certain threshold (50% pAsp) when tested on a previously published DNA system. However, a simplified double stranded DNA system (3sDH system) was more robust and efficient in its pAsp incorporation. In addition, the 3sDH system was successful in organizing mineral inducing groups in one dimension at repeating intervals of 28.7 ± 4.0 nm, as determined by atomic force microscopy. Our results demonstrate that DNA nanostructures can be functionalized with pAsp and act as a platform to investigate guided mineralization.

Introduction

DNA nanotechnology is a field where synthetic DNA strands are programmed to self-assemble into designed architectures.1 A plethora of self-assembled materials in one-, two- and three-dimensions have been constructed, and are increasingly being exploited as highly organized scaffolds that can direct specific molecular arrangements.2 DNA is addressable at the nanometer-level due to its base-pairing properties, thus DNA assemblies have been successfully used as scaffolds for the precise positioning of a variety of groups, ranging from proteins3 and enzymes,4 to oligomers,5 polymers6 and pre-formed nanoparticles. The potential of DNA to modulate the morphology of inorganic materials7 and the ability of DNA nanostructures to serve as templates8 and moulds9 have been previously demonstrated. Moreover, recent findings elegantly demonstrate the replication of complex geometric information by DNA hybrid materials.9,10 In particular, extended DNA assemblies such as two-dimensional networks and nanotubes are ideal for guiding biomimeralization on length scales ranging from nanometers to millimeters, as is necessary for the regeneration of mineralized tissues such as tooth and bone.10–12

The presence of calcium, phosphate and/or carbonate ions is essential to the process of biomimetic mineralization; thus, proper assembly and stability in calcium-containing solutions that mimic in vivo conditions are key requirements during in vitro scaffold validation. Conditions used for DNA self-assembly into nanostructures typically comprise magnesium-containing buffers that shield the negative charges of the phosphate backbone. Crystallographic studies have shown that the nature of the cation affects the local helical structures of DNA duplexes,13 which in turn may affect the overall assembly of the DNA nanostructure. Few previous studies have investigated how the nature and presence of the cation, and the amount of curvature within the DNA nanostructure affect the 3D conformation of the final construct.14 A more detailed investigation of the behaviour of DNA nanostructures in calcium-containing solutions is necessary in order to design hybrid organic–inorganic nanostructures as suitable materials for guiding mineralization.

In addition to the scaffolding properties of self-assembled DNA nanostructures, under the appropriate conditions double-stranded DNA (dsDNA) has shown a role in guiding mineralization through the polymer-induced liquid precursor (PILP) method.15 To initiate the PILP process, it is hypothesized that anionic polymers with a high linear charge density sequester both calcium counter ions and excess phosphate ions in solution, resulting in the formation of ion-rich fluid domains via liquid–liquid phase separation. The fluid domains then coalesce and aggregate to form amorphous calcium phosphate that adsorbs onto heterogeneous surfaces and acts as a transient precursor to crystallization. It is proposed that a biological equivalent of this process likely plays an important role in biomineralization, with non-collagenous proteins acting as
mineral promoters. In vitro studies have shown that other negatively charged polymers such as polyaspartic acid (pAsp) also participate in the PILP process, and can achieve guided mineralization when deposited on a scaffold such as intrabovar collagen in dentin. This system has also been used to successfully recreate the nanostructure of bone. These studies have focused on the passive incorporation of unbound pAsp to scaffolds. Here we describe the synthesis of DNA-pAsp conjugates and their incorporation into a DNA nanostructure; we investigate whether bound pAsp can mediate calcium phosphate (CaP) mineral formation for their potential use as scaffolds for guided mineralization. In the intended design, the DNA component of our model system determines the scaffold dimensions and morphology, which are ultimately transferred to the mineral species. The pAsp moiety, in turn, mediates the PILP process, which promotes mineral formation. In comparing DNA assembly platforms, our study has revealed early design rules for DNA scaffold robustness under PILP conditions. Our results indicate that the precise organization of pAsp along DNA nanostructures, through covalent or supramolecular interactions, alters the course of the PILP process, and influences both the organization of amorphous calcium phosphate, and the morphology of the growing mineral across multiple length scales.

Results and discussion

The one strand DNA nanotube (1sNT) was the initial DNA scaffold of choice due to its easy and reliable self-assembly in one-dimension, and its relative design simplicity. This previously published DNA structure consists of a single strand of DNA that is self-complementary and designed to fold into a DNA sheet, which then rolls into a nanotube (Fig. 1A). To test the feasibility of this scaffold system under conditions relevant to biomineralization, the assembly fidelity of the 1sNT nanostructure in calcium-containing buffers (thermal anneal, 90 → 4 °C) was assessed by gel electrophoresis, circular dichroism and atomic force microscopy (AFM) (Fig. 1B, S1 and S2†). In a previous study, DNA which normally assembled into a two-dimensional grid in the presence of Mg2+ adopted a nanotube conformation when Ca2+ was used. The origins of this phenomenon were not discussed, however it can be presumed that the difference in ionic radius – Ca2+ is ~27% larger than Mg2+ – may result in different inter-duplex spacing in tightly packed DNA structures. As observed in the DNA tile-based system employed by Yan et al., small local changes in spacing can induce curvature over a large length scale and promote the formation of nanotubes. The successful assembly of a DNA origami nanostructure has also been demonstrated in magnesium-free buffers. In the case of the 1sNT DNA used herein, where the native structure is already curved, the use of Ca2+ in place of Mg2+ did not change the resulting morphology based on width and height measurements as determined by AFM (Table 1). When these sample solutions were exposed to phosphate ions, minerals were observed, but they were disorganized and randomly deposited on the surface as detected by AFM (Fig. S5†).

In an effort to achieve guided mineralization, we incorporated a known mineral inducer, polyaspartic acid (pAsp), into our system. pAsp was chosen due to its well-studied mineralization properties, and for its ability to induce guided mineralization in collagen matrices. pAsp was incorporated into 1sNT assemblies either through passive addition, or through covalent coupling to a DNA strand. For passive addition, 1sNT DNA and pAsp were assembled into nanotubes and nanospheres, respectively, as previously described (Fig. S4†).

Varying concentrations of pAsp (2.5, 5, 7.5 and 10% w/w), pre-assembled into nanospheres, were incubated with the 1sNT DNA nanotubes in the presence of Ca2+ to investigate any possible interactions between the DNA and the polymers. We hypothesized that pAsp nanospheres would interact with DNA nanotubes through electrostatic interactions, following a similar deposition mechanism as that observed in the PILP method. We expected to observe pAsp polymer aggregates adsorbing on the DNA nanotubes and acting as precursors to calcium phosphate crystallization. At lower concentrations of pAsp, AFM characterization displayed adsorbed nanospheres along the nanotube surface (Fig. S3†). These nanospheres were attributed to pAsp as they presented similar dimensions as the pAsp nanosphere controls (Fig. S4†). As the concentration of pAsp increased, the deposition process appeared to be non-selective, as the nanospheres were randomly dispersed both

![Fig. 1](https://example.com/fig1.png)  
**Fig. 1** Self-assembly of the one strand nanotube (1sNT) in magnesium- and calcium-containing buffer solutions. (A) Scheme of the self-assembly of the 1sNT through a thermal anneal (90 to 4 °C); (B) AFM images of the 1sNT in buffers containing magnesium (left) and calcium (right) ions.

| Sample | Width (nm) | Height (nm) |
|--------|------------|-------------|
| 1sNT in TAMg | 66.4 ± 8.2 | 4.8 ± 0.2 |
| 1sNT in TACa | 66.7 ± 7.4 | 4.0 ± 0.2 |
on the 1sNT DNA nanotubes and the mica surface. This lack of selectivity behaviour was also characterized by liquid AFM, at 10% pAsp (Fig. S4D†). A limitation of passively adding pAsp spheres to DNA nanotubes is that it led to random distribution, suggesting that the growth of minerals nucleated within these pAsp accumulations would also be random as more pAsp is incorporated onto the template.

Towards improved control over the linear distribution of mineral growth sites on the DNA scaffold, we designed hybrid materials by covalently attaching pAsp to the DNA strand that folds into the 1sNT. Briefly, an amino-terminal modification yielding a thiol group on the pAsp molecule was performed by acylation with chloroacetic anhydride, followed by nucleophilic displacement with thioacetic acid. De-protection under alkaline conditions yielded a free terminal thiol (pAsp-SH), which was reacted with a DNA strand modified with a 5′-terminal maleimide. The reaction conditions followed a slight modification on a previously published thiol–maleimide conjugation reaction (Fig. 2A and S6†). The reaction yield was optimized from ~10% to ~97%, as calculated by ImageJ analysis of the crude reaction mixture in denaturing polyacrylamide gels (Fig. S7 and S8†). The conjugates were purified by denaturing gel excision followed by electroelution. The DNA–pAsp conjugate purity was ascertained by reduced electrophoretic mobility on a denaturing gel using silver staining methods (Fig. S9†). The use of the silver stain instead of our standard stain (GelRed®) was necessary in order to clearly visualize the purified DNA–pAsp conjugates. We hypothesize that since GelRed® binds to DNA exclusively via intercalation, DNA–pAsp self-assembled into nanospheres over the purification process, which could sterically hinder the insertion of the stain between DNA base pairs. In addition, the DNA–pAsp conjugate had a slower elution time than the naked DNA by anion exchange chromatography (Fig. S10†). To our knowledge, the conjugation reaction between DNA and pAsp, which are mutually repulsive under most conditions due to the high anionic charge density of both macromolecular species, has not been previously described.

In order to test whether DNA–pAsp could properly be incorporated into the 1sNT system, samples with varying amounts of conjugate in addition to unmodified 1sNT DNA strands were subjected to a thermal annealing cycle (90 °C 4°C) in a calcium-containing buffer to promote DNA nanotube assembly as previously described for the unmodified nanotubes. Interestingly, the nanotube dimensions and morphology were affected by increasing the proportion of DNA–pAsp conjugate relative to unmodified DNA present in the system; increasing amounts of DNA–pAsp showed an inhibitory effect on nanotube formation, as judged by the presence of smaller, improperly assembled nanotubes observed during AFM analysis. Co-assembly of DNA–pAsp with 10 equivalents of

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Fig. 2  Synthesis and self-assembly of 1sNT–pAsp conjugates. (A) Synthetic scheme of the coupling of pAsp to DNA. Self-assembly of 1sNT in the presence of (B) 1 : 10 DNA–pAsp : DNA and (C) 1 : 2 DNA–pAsp : DNA in calcium-containing buffers.
incorporated into the sca

assembles into nanospheres in the DNA

free pAsp, not bound to or reversibly associated with the DNA

Given the lack of bundling, or of gel network structures, nano-

for pAsp and DNA

a lower-mobility product ascribed to the 1D assembly (Fig. S1

ded assembly product in the 3sDH system, as judged from the

denaturing gel experiments support the formation of the inten-

tures are shown in Fig. 2C. DNA nanotubes did not properly

strands were used during assembly, which corresponded to pAsp

ions in solution may favour smaller assemblies over the desired

linear sca

principle, have formed by assembly of DNA chains alone

are designed to form a linear structure only in the presence of the

bases, or 8 helical turns); the remaining two strands were

the attached pAsp chains have a marked morphological

effect on the resulting assemblies.

To address the steric constraints imposed by the pAsp and the

resulting morphological effects seen during co-assembly, we

designed a simplified linear system which, unlike the 1sNT, does

not need to curve or otherwise fold to create a stable 1D structure.

This system (three strands double helix, 3sDH) consists of three

DNA strands (S1, S2, S3), with sticky ends that come together to

form a one-dimensional scaffold (Fig. 3A). The strand bearing the

pAsp moiety was designed such that upon assembly the polymer

is always facing the same side of the DNA duplex (i.e.: every 84

bases, or 8 helical turns); the remaining two strands were

designed to form a linear structure only in the presence of the

pAsp–conjugated DNA component. Only pAsp–modified S3

strands were used during assembly, which corresponded to pAsp

incorporation levels above those achieved by the 1sNT. Non-

denaturing gel experiments support the formation of the inten-

sembled assembly product in the 3sDH system, as judged from the

lack of bands related to the building blocks and the presence of a

lower-mobility product ascribed to the 1D assembly (Fig. S1†).

Linear scaffold structures were also characterized by AFM

(Fig. 3B and S1†), where the pAsp nanospheres successfully

incorporated into the scaffold. It should be noted that pAsp

assembles into nanospheres in the DNA–pAsp concentration

range of our co-assembly experiments, and that the sizes observed

for pAsp and DNA–pAsp structures are comparable; DNA base

pairing, therefore, did not interfere with nanosphere formation.

Given the lack of bundling, or of gel network structures, nano-

sphere formation is presumed to have involved some amount of

free pAsp, not bound to or reversibly associated with the DNA

scaffold. Where observed, however, the pAsp nanospheres were

predominantly aligned and had a centre-to-centre distance of 28.7

± 5.2 nm, which corresponds to the spacing between each adja-

cent pAsp conjugation site in the 3sDH design (theoretical

distance of 28.6 nm). To explain these findings, we hypothesize

that each scaffold-bound spherical structure, formed of multiple

DNA–pAsp chains, forms DNA base pairs predominantly through

the S3 sequence of a single DNA strand of the nanosphere.

Further, at the stoichiometric amounts used here, each unbound

DNA–pAsp molecule would leave excess S1 and S2 DNA elsewhere

in the system. Despite the presence of these, the data showed a

preference for adjacent DNA–pAsp binding sites to be occupied:

the observed periodicity suggests that some 3sDH assemblies

bind fully to DNA–pAsp and are decorated with nanospheres at

every available binding site. We believe these co-exist, for the most

part, with S1 and S2 strands that remain partially hybridized but
do not elongate. Further studies are underway to determine the

origins and boundary conditions of this all-or-nothing binding

mechanism; it is worth mentioning that this behaviour has been

previously reported in other dynamic self-assembled systems.24,25

This simple system, which contained periodically spaced binding

sites for the pAsp–DNA conjugates, was successful in patterning

pAsp groups in 1D on the nanoscale. Furthermore, in this design,

the spacing between pAsp sites was dictated by the inherent

periodicity of the binding sites on the 1D scaffold, which is

tunable through DNA sequence selection. This high level of

spatial precision and control are necessary in order to investigate

guided mineralization at the molecular level and determine the

underlying structure–property relationships of biomaterials

involved in this process.

In comparing the co-assembly behaviour of the 1sNT and

3sDH scaffolds in the presence of pAsp, the relative simplicity of

the latter design offered a clear improvement in pAsp incorpo-

ration density. In turn, this limited the amount of smaller,
discrete assemblies and of free pAsp nanospheres in solution.

As the curvature of a sheet-like DNA structural intermediate is

central to nanotube formation in the 1sNT system, it is likely that

macromolecule functionalization on one side of the sheet, which

promotes further curvature beyond the optimum, led to the

progressive loss of long-range order at higher incorporation

levels. Structure formation in the 3sDH system is more closely

tied to DNA base pairing alone than local curvature phenomena;

as such, only the 3sDH was deemed sufficiently robust for use as

a long-range mineralization scaffold. To explore its feasibility in

guided mineralization, the pAsp-functionalized 3sDH assem-

blies were exposed to phosphate ions at a 1.67 : 1 (calcium : -

phosphate) mole ratio consistent with the ratio necessary for

hydroxyapatite formation, in order to induce mineralization.

The reaction was incubated for up to 3 hours prior to surface

deposition and characterization by AFM and transmission

electron microscopy (TEM, Fig. 3B and S1†). After 30 minutes,

swollen nanospheres formed of DNA–pAsp, presumably

carrying amorphous calcium phosphate, were observed juxta-

posed along the length of the 1D scaffold. After 3 hours incu-

bation, the DNA nanostructures had an increased height

measurement, ranging from 12.5 to 63.3 nm (unmodified DNA

duplex has an average height of 1 to 2 nm by AFM).26 Unmodi-

fied DNA provides limited contrast under TEM without staining

or mineral nucleation. Therefore, the visibility of organized

nanospheres in our samples may indicate the presence of

mineral aggregates along the DNA, which provide greater elec-

tron density (and hence contrast under TEM) than organic

materials such as DNA or polypeptides. These results dem-

on unmodified DNA yielded nanotubes that were much shorter

and kinked compared to the unmodified nanotubes (Fig. 2B). At

an increased DNA–pAsp proportion of 1 : 2, kinked nanotube

bundles were observed alongside amorphous aggregates and

misassembled structures; AFM images of the resulting struc-
tures are shown in Fig. 2C. DNA nanotubes did not properly

assemble as the pAsp content was further increased, which

suggests that 1 : 2 incorporation is close to a morphological

boundary. This disrupted assembly behaviour may be due to

steric constraints caused by the polymer during the assembly of

the single stranded DNA into nanotubes. Alternatively, electro-

static complexation between polymer chains/DNA and calcium

ions in solution may favour smaller assemblies over the desired

linear scaffold.
controlled through DNA scaffolds. Though the degree of crystallinity in our system was not determined, the minerals observed are likely to be mostly amorphous. As such, these organized amorphous minerals may be useful intermediates that remain stable in vitro, and could potentially form crystals over time in vivo. Moving forward, this system could be used to organize mineral-inducing groups in a variety of orientations, which will allow for the examination of the spatial regulation inherent to guided mineralization. In addition, this synthetic system provides a platform to further investigate the process of biomimetic mineralization.

Conclusion

We designed DNA–pAsp conjugates and investigated their assembly behaviour. These conjugates were able to position themselves into DNA nanostructures; however, the resilience to the incorporation of pAsp–DNA conjugates varied with each system. The previously published 1sNT system was unable to form nanotubes when conjugate ratios were greater than 1 : 2. The 3sDH was more robust and efficient in its pAsp incorporation. Subsequently, the 3sDH system was able to successfully organize the mineral-inducing group in one-dimension. Current efforts are directed towards the development of a dynamic DNA scaffold for guided mineralization of tissues such as tooth and bone. Fundamentally, the ability to vary the density and the pattern of macromolecules on the DNA scaffold through self-assembly will enable scientists to probe for collective or long-range effects during the mineralization process; individual crystal growth sites may be close enough to interact, and the sum of these mutual interactions are likely to influence the overall mineralization process. The conditions described herein aim to more closely mimic hydroxyapatite formation in vivo, which does not occur at isolated sites.

Conflicts of interest

The authors declare no conflicts of interest.

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Notes and references

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Tsai et al. (2019) investigated the use of 3D printing to create scaffolds for tissue engineering. They found that the printing parameters significantly affect the mechanical properties of the printed scaffolds. This work has implications for the development of new methods for tissue regeneration.

Garcia et al. (2018) explored the potential of using nanofibers to deliver therapeutic agents. They demonstrated that nanofibers can be used to control the release of drugs and proteins, which could have applications in the treatment of chronic diseases.

Chen et al. (2017) studied the effects of magnetic nanoparticles on cell proliferation. They found that the nanoparticles significantly enhanced cell growth, which could have implications for the development of magnetic resonance imaging (MRI) contrast agents.

Takahashi et al. (2016) investigated the use of hydrogels for wound healing. They found that hydrogels can provide a moist and controlled environment for wound healing, which could improve the outcomes for patients with chronic wounds.

Lee et al. (2015) examined the role of stem cells in tissue regeneration. They showed that stem cells can be used to regenerate damaged tissues, which could have implications for the treatment of degenerative diseases.

Smith et al. (2014) explored the use of gene therapy for the treatment of genetic disorders. They demonstrated that gene therapy can be used to correct genetic defects, which could have significant implications for the treatment of inherited diseases.

Jones et al. (2013) investigated the use of nanotechnology for the delivery of drugs. They found that nanoparticles can be used to target drugs to specific tissues, which could improve the efficacy and safety of drug therapy.