From pico to nano: biofunctionalization of cube-octameric silsesquioxanes by peptides and miniproteins†

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Polyhedral silsesquioxanes are considered valuable conjugation scaffolds. Nevertheless, only a few examples of silsesquioxane-assembled peptide oligomers have been reported to date. We developed a new bioorthogonal cube-octameric silsesquioxane (COSS) scaffold bearing eight aminooxy coupling sites allowing for the conjugation of diverse peptides via oxime ligation. We found that the coupling efficacy depends on the ligand in view of steric hindrance and electrostatic repulsion. For the first time scaffold-based conjugation of cystine-knot miniproteins having a backbone of about thirty amino acids was successfully accomplished without loss of bioactivity. Atomic force microscopy (AFM) provided further knowledge on the size of COSS verifying them as picoscaffolds growing upon bioconjugation to nanodimension.

Introduction

Combining several ligands on a single scaffold often results in improved characteristics of a formed oligomer compared to its individual constituents. Nature extensively uses this phenomenon known as multivalency to effect biomolecular interactions in living organisms by enhancing affinity and specificity of binding. This inspires research efforts towards creation of synthetic molecules in which the benefits of simultaneous multiple contacts are achieved through the oligomerization of bioactive modules – small molecules, carbohydrates, peptides, and proteins. Properties of scaffold-grafted molecular blocks are often governed by the peculiar architecture that implies shape, size, and valency of the framework, as well as spatial orientation of ligands.

In recent years, cube-octameric silsesquioxanes (COSS) have been brought into focus as promising oligomerization scaffolds due to their unique characteristics. These monodisperse particles with a core size of 0.5 nm are considered the smallest known nanoscaffolds with a high degree of symmetry. Their hybrid molecules are composed of a siloxane inorganic core decorated with organic ligands which combine an aliphatic linker with a terminal active group. From a broad repertoire of functionalized COSS molecules reported to date, amine, azide, azide, alkyne, thiol, aldehyde or maleimide bearing particles are potentially applicable for bioconjugations.

To date, several COSS-based bioconjugates have been reported, among them lysine dendrimers for drug delivery, oligomers of peptides and carbohydrates, as well as macrocyclic Gd chelates as potential magnetic resonance imaging contrast agents. The convergent synthesis of peptidic COSS is limited to homopolypeptides where one amino acid unit with a protected or non-reactive side chain is propagated, or to rather short (up to 8 residues) RGD oligomers. In these bioconjugations, copper-catalyzed azide–alkyne cycloaddition (CuAAC) is currently the established method to obtain a linkage between an azide-bearing silsesquioxane and an alkyn-modified peptide ligand usually added in stoichiometric excess. Due to the instability of a COSS core in the presence of aqueous nucleophiles, CuAAC with silsesquioxanes is generally performed in water-free DMF. In this aprotic solvent, the copper catalyst often appeared to be coordinated by an amide backbone or functional side chains of peptides leading to a drastic decrease of active catalytic species and, as a consequence, to prolonged reaction times (up to several weeks) and low yields. Very recently, an elegant alternative approach to silsesquioxane-based peptide conjugations has been reported which utilized the photo-induced free-radical thiol–ene coupling and resulted in eightfold presentation of the tripeptide glutathione on a COSS scaffold. Interestingly, for a tetrapeptide RGDC complete
hydrothiolation of vinyl coupling sites was achieved only after the introduction of elongation linkers.31

Herein, we report bioconjugation on a COSS scaffold in acidic aqueous media based on the oxime ligation33,34 between an aldehyde and an aminooxy functionality (Scheme 1). This approach provides several advantages compared to other methods reported for COSS–peptide conjugations12,27,28,31,32 as it ensures both stability of the silsesquioxane core and good solubility of biological ligands. The introduction of aldehyde moieties into biomolecules is well-established and can easily be achieved via periodate oxidation of an N-terminal serine residue.35 Moreover, building blocks for solid phase peptide synthesis36,37 containing a masked side-chain aldehyde function enable the generation of this moiety at any desired sequence position. Aminooxy functionality can be easily introduced in peptides by N-acylation of side-chain or terminal amines by butoxycarbonyl (Boc)38 or ethoxyethylidene (Eei)39,40 protected aminooxy acetic acid. Protection is essential to avoid side reactions and overacylation of target compounds. In the present research, the applicability of the oxime ligation to the synthesis of hybrid COSS–peptide conjugates was studied highlighting the benefits and the limitations of the method.

Results and discussion

As a conjugation scaffold an octaaminopropyl-COSS 1 was used. The reaction with commercially available mono-Boc aminooxy acetic acid towards 3 appeared strongly dependent on the activation conditions leading to overacylated species (Table 1). Decostaire et al. have recently reported38 that the control over these undesired processes could be achieved by the proper choice of used activator, base, and solvent as well as their excess. Corroborating the reported data,38 2-(7-aza-1H-benzotriazole-1-yl)-1,1,3,3-tetramethyluronium hexa-fluoro phosphate (HATU) activation in the presence of collidine was found to be the optimal reaction conditions, and 20 equivalents of the base per silsesquioxane octamer were used (Fig. 1). Although no overacylated COSS species have been found in both acetonitrile- and DMF-based reaction mixtures (Table 1, entries 5 and 6), dry acetonitrile appeared to be the solvent of choice as no complete conversion was observed in DMF after an overnight reaction. Contrary to this procedure, reaction of 1 with N-hydroxysuccinimide-activated 2-(1-ethoxy-ethylidene-aminooxy) acetic acid40 resulted in octakis Eei-protected aminooxy-COSS 2 without overacylation. Due to the instability of COSS in the presence of aqueous nucleophiles,21,26 the cage integrity of 2 and 3 was verified by 29Si-NMR and IR spectroscopy (see ESI Fig. S3, S4†). The NMR spectra showed exclusively the shifts corresponding to COSS cages 2 and 3 (δ −66.87 and δ −66.75, respectively). The IR spectra showed a Si–O–Si stretch characteristic band at 1115 cm−1. Both Boc and Eei groups were cleaved in 50% aqueous trifluoroacetic acid (TFA) within 3 hours, and one-pot conjugation with aldehyde functionalized peptides35 succeeded. In our study, ligands with diverse primary

![Scheme 1 Synthetic approaches to protected aminooxy COSS 2 and 3. Octakis (Boc-aminooxy) COSS was used as a scaffold for bioconjugation. Ligands: an octapeptide with an RGD motif (p4), a derivative of the antimicrobial peptide Jelleine 1 (p5), mastoparan (p6), MCoTI-based miniproteins (p7, p8) and Kcoil, a basic α-helical coil (p9), were chosen.](image)

![Pep tidic Ligan ds](image)

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<th>Amino Acid Sequence (AAS)</th>
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<td>IPR5DYRGK-OH</td>
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<tr>
<td>LHLKSFPG-OH</td>
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<td>8 3</td>
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<td>CGVSAKELb-NH&lt;sub&gt;2&lt;/sub&gt;</td>
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<th>Table 1 Reaction conditions for the synthesis of 3</th>
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X: detected; nd: not detected. Reaction showed incomplete conversion. Heptameric Boc-aminooxy COSS species were detected via LC-MS analysis.

![Fig. 1 ESI-MS spectra for the synthesis of 3. (a) Reaction in DMF using HBTU and 85 equiv. collidine; (b) reaction in MeCN using HATU and 20 equiv. collidine.](image)
and secondary structures and different bioactivity were used (Scheme 1).

For the initial experiment, an integrin-binding octapeptide bearing an RGD functional motif (p4) was chosen.\textsuperscript{41,42} It was shown previously that this ligand could be coupled onto the COSS scaffold in eightfold copies via CuAAC.\textsuperscript{12} We found that oxime ligation had obvious benefits compared to CuAAC as the reaction time needed for full conversion was drastically reduced – from several weeks to an overnight reaction at room temperature. LC-MS monitoring of the reaction progress showed the presence of COSS species bearing both protected aminooxy coupling sites and oxime-ligated peptides (see ESI Fig. S8†), indicating that Boc cleavage was the rate-determining step. Nevertheless, as soon as full deprotection was achieved, the steric hindrance and electrostatic repulsion caused by the growing number of peptidic ligands seemed to have a major influence on the reaction progress.

Consequently, we explored the ligand-dependent limitations of the proposed COSS-based oximation. It is obvious that direct coupling of bulky peptide ligands onto constrained COSS scaffolds can be hindered due to steric reasons. On the other hand, often the oligomerization of peptides and full-size proteins is desirable without implementation of elongation linkers. Therefore, in a series of coupling experiments the size of bioactive peptidic ligands was progressively increased from eight (p4) to 35 (p9) amino acid residues.

To that end, a derivative of natural antimicrobial peptide Jelleine \textsuperscript{43,44} from the royal jelly of honey bees (p5) was used. The successful eightfold presentation of this octapeptide on the COSS scaffold encouraged us to approach more sophisticated ligands. Thus, an α-helical calmodulin\textsuperscript{45} binding peptide mastoparan (p6)\textsuperscript{46} comprising 14 residues, trypsin inhibitors\textsuperscript{47} p7 and p8 containing 29 and 30 residues, respectively, and the inherent Kcoi part of a heterodimeric coiled coil (p9)\textsuperscript{48–50} with a length of 35 amino acids were used. Moreover, miniproteins\textsuperscript{51} p7 and p8 contained a characteristic tri-disulfide pattern known as a cystine knot\textsuperscript{52} that is absolutely essential for the function; the damage of this motif results in the loss of three-dimensional structure and, as a consequence, of bioactivity.\textsuperscript{52}

All peptidic ligands contained, compared to their parent sequences, an additional N-terminal serine and were assembled by standard microwave-assisted Fmoc-SPPS as previously described.\textsuperscript{53} After chain assembling and cleavage from the support, sodium periodate oxidation resulted in peptide aldehydes.\textsuperscript{35} In the case of miniproteins p7 and p8, oxidative folding\textsuperscript{54} into a cystine knot preceded the formation of an N-terminal glyoxal.

One-step deprotection of 3 followed by conjugation with peptidic ligands p5–p9 was performed in 50% aqueous TFA as described above. The LC-MS monitoring of the conjugation reaction with cystine-free ligands revealed the formation of COSS–peptide intermediates after 10 min. Within 12 hours the reactions were completed. The formation of miniprotein–COSS conjugates 7 and 8 was only observed after an overnight reaction. This might be attributed to a hindered accessibility of the miniprotein aldehyde functionalities.

The amount of ligand copies attached to the COSS core correlated with the primary structure of the peptides and the increasing steric demand. Thus, the conjugation with the Jelleine-derived octapeptide p5 yielded an octameric product. Mastoparan p6 with just 6 additional amino acids formed a hexameric conjugate. In miniproteins p7 and p8 the amount of amino acids is doubled compared to p6 resulting in a decreased coupling efficacy. Accordingly, only dimeric as well as trimeric constructs were observed for these ligands (Fig. 2). Finally, the pronounced steric hindrance as well as the possibility of strong electrostatic

![Fig. 2](image-url)
repulsion allowed only for the formation of mono- and dimeric products with the α-helical Kcoil peptide p9.

To the best of our knowledge, to date the dimerization of miniproteins was only facilitated via chemical cross-linking using a bis-succinimidyl suberate.55,56 Herein, we present the first scaffold-based approach to the oligomerization of miniproteins. To examine whether cystine knot protease inhibitors retained their unique three-dimensional structure after conjugation, we studied the bioactivity of formed hybrid molecules. To that end, an RP-HPLC fraction containing conjugate 7 and lower substituted COSS–p7 conjugates was tested for inhibitory activity against trypsin (Fig. 2).56 Although a precise determination of inhibitor concentrations in the mixture of tri-, bi- and monoconjugated miniproteins was not possible and corresponding inhibition constants were not calculated, the concentration-dependent inhibition of trypsin-mediated proteolysis provided clear evidence that the bioactivity of p7 was preserved under oxidation conditions.

The bioactivity and toxicity of nanoparticles as well as their potential for drug delivery is greatly dependent on the size. Therefore, we conducted atomic force microscopy (AFM) experiments using the particles 1, 4, and 5. We considered mica (negatively charged sheet silica) an optimal substrate, as the eight amino groups of 1 and the peptidic side chains of 4 and 5 promised a good binding to this surface. The dropcasting of an acidic solution of 1 (∼1 ng mL⁻¹) allowed us to visualize singular particles. The enumeration of 60 particles (see ESI Fig. S7† and Fig. 3d) resulted in an average particle size of 743 ± 211 pm. This value is in good accordance with the literature stating ~500 pm for the POSS core.8,9 For further measurements a mica substrate was dipcoated with an RP-HPLC fraction containing 4. An analysis of the particle size distribution revealed an average size of 1810 ± 300 pm. Hence, the peptidic shell induced a measurable size increase of 1050 pm. On a first glance, this value seemed to be rather small compared to a rough simulation of 4 predicting 6.42 ± 0.69 nm as its maximal diameter in solution (see ESI Fig. S12†). However, it has to be taken into account that the particle is absorbed on a mica surface and might adopt a pancake-like structure. Furthermore, the recorded topography of a sample always depends on several parameters. Being operated in tapping mode, the AFM z-piezo detects changes upon the interaction of its tip with an analyzed sample keeping the amplitude of the vibrating cantilever constant. If attractive van der Waals forces change due to differences in the Hamaker constant between two materials, the z-piezo will compensate this difference by varying the mean distance between the tip and the sample. On soft materials, compared to the sample substrate, an additional difference of the indentation depth of the tip into the material is compensated by the z-piezo. As a consequence, even for atomically flat surfaces a non-negligible height profile can be measured for inhomogeneous samples.58 Hence, the measured height differences should only be used to classify the nanoparticle dimension.

The subsequent analysis of 5 revealed, in accordance with the expectations, particles with a similar height profile. Therefore, we chose a random particle and continuously increased its mapping resolution using bimodal atomic force microscopy. Fig. 4a shows the high resolution phase image of the second eigenmode. Phase images in AFM are related to mechanical properties of the sample. Three regions with a distinctive phase shift between cantilever oscillation and excitation were identified. The difference in phase values of mica (Δφ₂ = 120°) and the main body of the particle (Δφ₂ = 119°) can be attributed to the mechanical dissimilarity of the materials. A region with a relatively low phase shift (Δφ₂ = 110°) was found in the right part of the particle (Fig. 4b). An image artefact can be excluded since the same area was successively scanned several times depicting comparable contrast. We presume that this region might visualize the position of the mica-adsorbed silsesquioxane core surrounded by randomly oriented peptide chains (Fig. 4d).

Conclusion

A new bioorthogonal COSS scaffold was developed comprising a highly symmetric silsesquioxane core decorated with aminoxy functional modules. We demonstrated the applicability of this scaffold for the oligomerization of aldehyde-bearing peptidic ligands through oxime ligation. Our scheme has an obvious advantage as all involved reactions proceed in an acidic medium that does not affect the COSS core known to be extremely unstable in the presence of nucleophiles. We found that the coupling efficacy depends on the ligand in view of steric hindrance and electrostatic repulsion. Thus, octapeptide derivatives were eightfold presented on a COSS scaffold, whereas for more bulky ligands lower oligomers were detected. Nevertheless, for the first time the scaffold-based conjugation of miniproteins comprising a backbone of about thirty amino acids with a structure-defining cystine knot was successfully accomplished without the loss of ligand bioactivity.

![AFM topography image of a uniformly distributed COSS–peptide particle 3 on a mica surface; (b) image extension displaying the trend of the section line; (c) profile of four adjacent particles; (d) comparative histogram depicting the size distribution of 1 and 4.](image-url)
AFM experiments conducted in the present study provided further knowledge on the size of COSS allowing us to consider them as picoscaffolds growing upon bioconjugation to nano-dimension.

Detailed studies concerning the bioactivity of COSS-assembled conjugates are currently ongoing, with constructs 6 and 9 being of particular interest. They can provide an interface for the non-covalent oligomerization of proteins fused either to a complementary Ecoil or to calmodulin. Thus, oligomerization can occur via the formation of a Kcoil/Ecoil leucine zipper, or through tight binding to mastoparan, respectively.

**Experimental**

**Peptide synthesis**

Microwave-assisted Fmoc-SPPS of peptide ligands was performed on a CEM liberty® peptide synthesizer equipped with a CEM discover® SPS microwave (CEM GmbH). As solid supports 2-CT-resin-Gly-OH from Iris Biotech GmbH (p4 and p5), AmphiSphere™ 40 RAM resin from Varian Inc. (p6, p7 and p8) and NovaSyn®TGR resin from Merck KGaA (p8) were used. After cleavage, cysteine-free crude products were oxidized using 10 equiv. of sodium periodate in PBS buffer for 10 min. The reaction product was isolated by RP-HPLC using a Varian 940- LC-MS monitoring. Subsequently, the solvent and base were removed in vacuo. The resulting crude product was purified via RP-HPLC using a semi-preparative C4 column (300 × 20 mm, 5 μ, PSS Polymer Standards Service GmbH) and a linear gradient of 90% aq. MeCN (10 → 100% B in 63 min) preceded by 10 min isocratic 10% B at a flow rate of 10 mL min⁻¹ with tR = 48.0 min. Yield after RP-HPLC purification: 5.7 mg (33%).

HR-MS: calc. for C82H166N15O44Si8 (+2): 1133.4224, meas. 1133.4216 [M + 2H]²⁺; A TR-IR: ν 3226 s, 2920 m, 2849 m, 1706 m, 1536 m, 1445 s, 1378 w, 1114 vs, 1075 m, 1018 m, 865 m, 730 w cm⁻¹, ²⁹Si NMR IGATED (99 MHz, CDCl₃): δ −66.85.

**Synthesis of scaffold 3**

50 mg (1 equiv., 0.043 mmol) of 1, 347 mg Boc-aminooxy acetic acid (5.3 equiv. per ammonium group, 1.815 mmol), 674 mg HATU (5.2 equiv. per ammonium group, 1.773 mmol) and 112.5 μL dry collidine (120.8 mg, 2.9 equiv. per ammonium group, 0.977 mmol) were dissolved in 8 mL dry MeCN. After 12 hours of stirring full conversion of 1 into 3 was confirmed via LC-MS monitoring. Subsequently, the solvent and base were removed in vacuo. The resulting crude product was purified via RP-HPLC using a semi-preparative C8 column (250 × 20 mm, 5 μ, Phenomenex Inc.) and a linear gradient of 90% aq. MeCN in 0.1% aq. TFA (40 → 100% B in 20 min) preceded by 5 min isocratic 40% B at a flow rate of 18 mL min⁻¹ with tR = 16.5 min. Yield after RP-HPLC purification: 3.6 mg (4%).

HR-MS: calc. for C₅₀H₈₄N₁₅O₇₄Si₈ (+2): 1140.4375, meas. 1143.4224, [M + 2Na]²⁺; A TR-IR: ν 1718 m, 1540 s, 1445 m, 1378 w, 1286 m, 1115 vs, 1011 m, 850 m, 6293 | 6291

**Conjugation of peptidic ligands on aminooxy COSS 3**

0.1 mg (1 equiv., 0.041 μmol) of 3 were dissolved in 500 μL 50% aq. TFA and 9 equiv. (0.39 μmol) of a corresponding peptidic ligand were added. The reaction process was monitored for 12 hours by LC-MS. The conjugates were isolated via RP-HPLC using an analytical C4 column (50 × 1 mm, 5 μ, Phenomenex Inc.) and a linear gradient of MeCN in 0.1% aq. formic acid (2 → 100% in 10 min) at a flow rate of 0.2 mL min⁻¹.

HR-MS: calc. for 4 C₁₄₂H₂₆₉N₁₄O₁₄Si₈ (9+): 1140.4375, meas. 1140.4393 [M + 9H]⁹⁺ (see Fig. S9 and S10†), calc. for 5

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**Fig. 4** High-resolution image of a single COSS–peptide particle 4 using bimodal atomic force microscopy. (a) Topography image and (b) profile of a single particle. (c) Corresponding second eigenmode phase image. The image reveals a bright region at the right area of the particle. (d) Schematic structure of the particle deduced from the phase image shown in (b).
C448H705N112O124Si8 (+9): 1096.0051, meas. 1096.0042 [M + 9H]+ (see ESI Fig. S14 and S15†), calc. for 6 C472H862N130O124Si8: 5.1065.34, meas. 10.565.37 (see ESI Fig. S17 and S18†). Yields based on the integration of HPLC traces at 220 nm: 90% (conjugate 4) and 99% (conjugate 5).

See the ESI† for full LC-MS and HR-MS data.

Trypsin inhibition assay

The trypsin inhibition assay of 7 was performed according to the reported procedure.56

High resolution imaging of COSS–peptide particles using bimodal atomic force microscopy

Tapping mode atomic force microscopy measurements were performed on COSS–peptide particles randomly distributed on a mica surface. The resonance frequency of the cantilever (PPP-ZEIHR and PPP-NCH from NanoandMore GmbH, Wetzlar, Germany) was f1 = 120–320 kHz and a free amplitude of A01 ≈ 10 nm was chosen. We paid careful attention to operate the AFM in the net attractive regime where van der Waals forces dominate the tip–sample interaction by keeping the free amplitude low and using a relatively high setpoint amplitude A1/A01 ≈ 0.9. This operation regime is comparably gentle to the sample surface avoiding excessive indentation of the tip apex into the surface structure.

High-resolution images on single COSS–peptide particles were accomplished using bimodal atomic force microscopy.60–66 Compared to the conventional tapping mode, the cantilever was excited simultaneously at the first and second oscillations, respectively. The free amplitude of the oscillating cantilever was excited to A01 = 5 nm and A02 = 0.5–1 nm.

AFM image processing

All topography images were 1st order flattened in order to remove image tilt. A Gauss filter was applied to enhance the signal-to-noise ratio.

Acknowledgements

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References