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Synthesis of discrete catalytic oligomers and their potential in silica-supported cooperative catalysis†

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Cooperative catalysis on solid surfaces relies primarily on two or more catalytic partners being close enough to each other to sustain the catalytic cycle. We describe here the synthesis and preliminary investigation of discrete homo-oligomers as flexible scaffolds to inflect the intersite distance and blur the compositional heterogeneity in a silica-grafted catalytic triad.

Supported multifunctional catalysts provide significant advantages over their parent homogeneous systems for more sustainable and efficient approaches in cooperative and cascade catalysis.^{1–12} However, although it is now becoming possible to combine most activation modes on a single support and compete with the original molecular catalysts, controlling the surface distribution of the different functional groups remains challenging.^{13–18} As exemplified by biological catalysts, the perfect spatial preorganization of catalytic centers within a controlled and confined environment is central to enable optimal synergistic interactions. Hence, the current limitation of grafting strategies to random, entropy-controlled systems is an important issue that prevents the fully rational design of active multifunctional surfaces.

The utilization of mixed monolayers for the preparation of silica-supported multifunctional catalysts intrinsically leads to a statistical distribution of the active components on surface.¹⁹ This eventually limits the potential of these systems since it creates a statistical distribution of pairings and spacings between the functional groups, with some catalytic partners being found at unmatched stoichiometry and distance within the mixed monolayer. Hence, it becomes important to develop methods to optimize the spatial positioning of functional groups in supported multifunctional catalysts to overcome the limitations associated with random functionalization. So far, only a few strategies have been proposed to reach a more homogeneous spatial distribution of multiple functional groups on surface. Molecular imprinting or imprint coating methods have allowed to better control the relative arrangement of surface-bound functional groups.^{20–25} For instance, Davis demonstrated that paired thiol/sulfonic acid obtained by molecular imprinting on silica surpassed the same randomly

distributed cooperative catalytic system.²³ In another way, the complementary functional groups can be assembled on a single molecular scaffold prior to immobilization.²⁶ In this direction, we recently reported the use of short sequence-defined oligomers for the preorganization of the (pyta)Cu/TEMPO/imidazole (pyta = pyridyltriazole, TEMPO = 2,2,6,6-tetramethylpiperidine-*N*-oxyl) catalytic triad on mesoporous silica and their superior activity in the aerobic oxidation of alcohols compared to a mixed monolayer of the triad.^{27,28}

The homogeneous (bpy)Cu/TEMPO/NMI catalyst system (bpy = bipyridine, NMI = *N*-methyl-imidazole) developed by Stahl and co-workers,^{29–32} provides a practical, efficient and environmentally benign alternative compared to common methods based on stoichiometric and toxic oxidizing reagents for the transformation of alcohols to aldehydes and ketones.³³ The mechanism involves the formation of a (bpy)Cu^I(NMI) complex that is then oxidized by molecular oxygen to provide the active oxidizing Cu^{II} species. The role of TEMPO is still unclear as whether it interacts directly or not with the Cu center;^{34–36} however, the formation of the (bpy)Cu^I(NMI) complex remains central in the catalytic cycle. Hence, in our approach, it is important to promote interactions between the supported Cu and imidazole sites while the TEMPO is located in close vicinity of the Cu complex to enable high turnovers. Here, we present another approach to minimize the effect of local compositional heterogeneity in randomly-mixed monolayers, based on the use of multivalent flexible short chains which dynamically blur the local composition of the grafted layer, and thereby achieve a statistically more efficient pairing between functional units.

In that aim, we first describe the simple iterative synthesis of azide-functionalized monodisperse oligomers as versatile platforms for the anchoring of the three catalytic units (pyta)Cu, TEMPO and imidazole. After grafting a stoichiometric mixture of the oligomers on mesoporous silica, we find that the catalytic activity per unit amount of catalytic center is strongly enhanced for longer oligomeric chains. This is ascribed to the larger volume explored by longer chains, resulting in a more efficient coupling between the three elements of the catalytic triad despite the grafting heterogeneity of the layers. The functional homo-dimers **13**, **15** and **17** and homo-trimers **14**, **16** and **18**

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bearing pendant TEMPO, imidazole and pyridine active centers, respectively, share the identical dimeric and trimeric precursors **9** and **10**, respectively (Scheme 1). The oligomeric platforms **9** and **10** were straightforwardly synthesized from racemic propargyl glycidyl ether **1** following an iterative epoxide opening/CuAAC (copper-catalyzed azide-alkyne cycloaddition) sequence.^{27,37,38} Precisely, the opening of the oxirane ring in **1** ($n = 0$) with the azide anion releases a secondary hydroxyl function that would later serve for the installation of the catalytic units. The azide functionality in **2** subsequently allows the re-introduction of the epoxide motif by CuAAC reaction with propargyl glycidyl ether. Reiterating this elongation sequence allows to access monodisperse oligomers with lateral $-OH$ handles.

The “living” oligomers **4** and **6** were terminated by reaction of the end-chain azide with phenylacetylene under CuAAC conditions to afford dimer **7** and trimer **8**, respectively. In order to install the catalytic components of the triad, the lateral hydroxyl groups were treated with *N,N'*-disuccinimidyl carbonate (DSC) followed by coupling with 2-azidoethylamine, to provide the dimeric (**9**) and trimeric (**10**) platforms with accessible azide groups. The later were reacted either with TEMPO **11**, imidazole **12**, or 2-ethynylpyridine under CuAAC conditions to provide the TBDMS-protected multivalent homooligomers that were ultimately treated with TBAF (tetra-*n*-

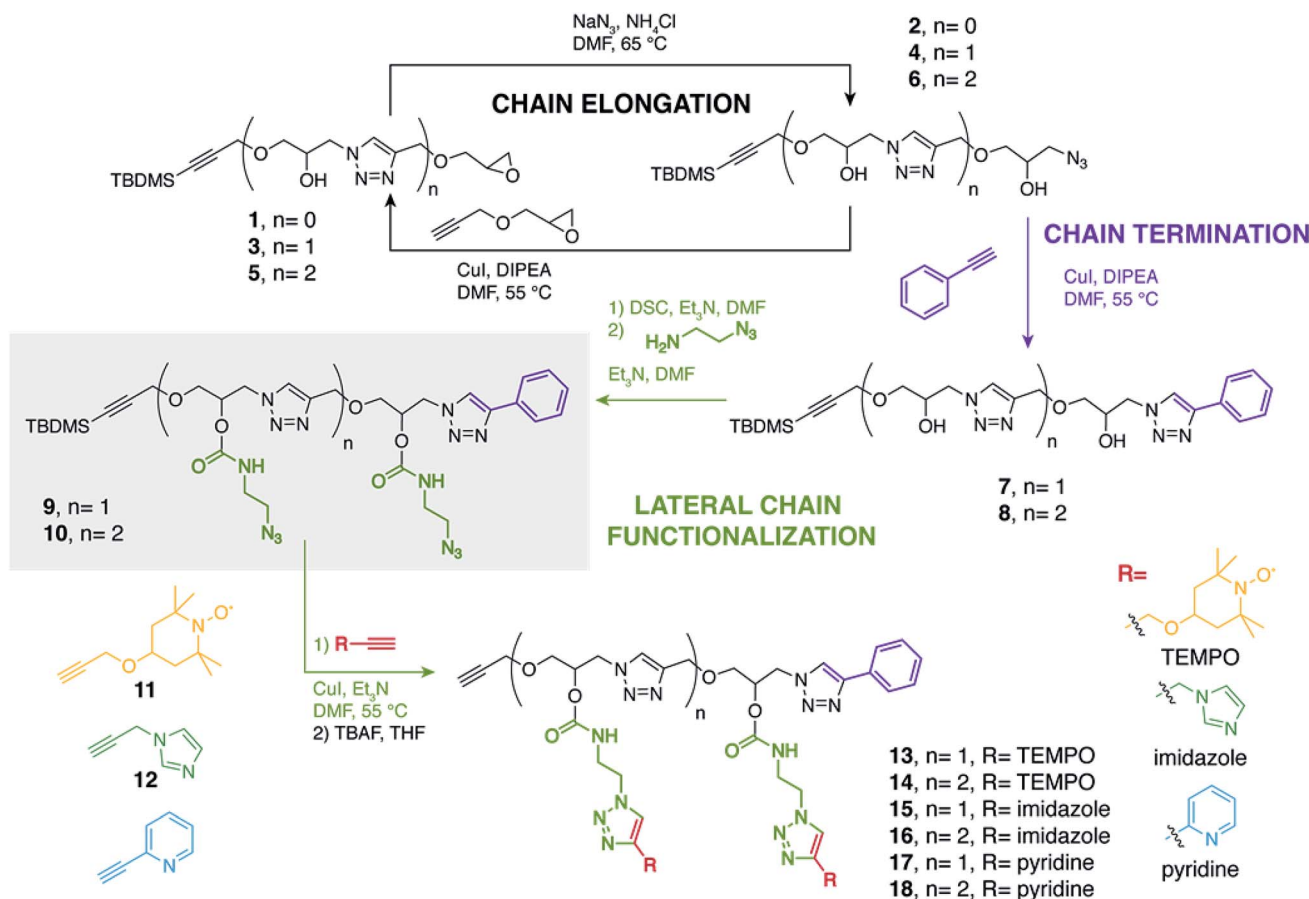
butylammonium fluoride) to afford the alkyne-terminated oligomers **13–18**.

The two-step post-polymerization modification strategy developed here for the installation of lateral functional groups mirrors previous approaches based on labile ester bounds.^{27,37} Precisely, the lateral carbamate bounds ensure here enhanced chemical stability compared to ester analogues that are partly or totally cleaved during the TBAF-mediated alkyne deprotection while the carbamate variants remain unaltered.

Trifunctional (pyta)Cu/TEMPO/imidazole supported catalysts of increasing valency were prepared using our typical CuAAC protocol by reacting stoichiometric mixtures of the corresponding alkyne-terminated catalytic groups with azide-functionalized mesoporous silica in presence of Cu(I) (Scheme 2); the Cu(I) used for the CuAAC grafting also serving for the realization of the (pyta)Cu sites of the supported catalysts.^{39,40}

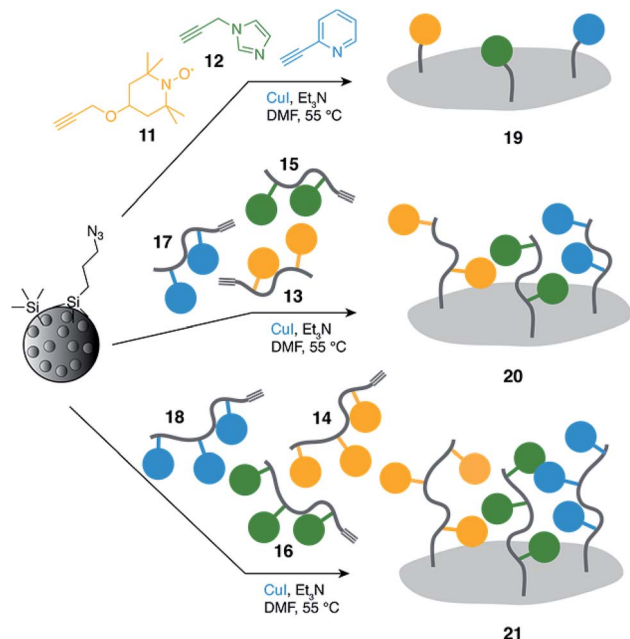
In order to further study the effect of the oligomeric scaffolds, the more traditional “monomeric” supported catalyst **19** was prepared from **11**, **12** and 2-ethynylpyridine (1 : 1 : 1 molar ratio) and azide silica (0.29 ± 0.04 mmol azide per g) in presence of CuI and Et₃N.

Similarly, catalyst **20** was obtained by reacting a mixture of dimers **13**, **15** and **17** in a 1 : 1 : 1 molar ratio with azide silica; catalyst **21** was in turn prepared from a stoichiometric mixture of trimers **14**, **16** and **18**.



Scheme 1 Preparation of the discrete catalytic oligomers **13–18**.





Scheme 2 Preparation of silica-supported multifunctional catalysts 19–21.

The grafting reactions were monitored by thermogravimetric analysis (TGA) and FT-IR. TGA (ESI, Fig. S41a†) provided loadings of 0.27 ± 0.01 , 0.15 ± 0.01 and 0.12 ± 0.01 mmol g⁻¹ for **19**, **20** and **21**, respectively, which corresponds to grafting efficiencies of 89 ± 11 , 53 ± 11 and $43 \pm 9\%$ with respect to the initial azide loading. The DTG curves (ESI, Fig. S41b†) also showed the persistence of thermal events corresponding to the azide layer for catalysts **20** and **21**, also indicative of partial grafting. As can be expected, the grafting efficiency decreased with the length and bulkiness of the oligomers as steric hindrances enter into consideration for the surface CuAAC reaction. FT-IR confirmed this observation with the residual azide signal (2100 cm^{-1}) being more prominent with the length of the immobilized oligomers, synonymous with incomplete CuAAC grafting (ESI, Fig. S42†). The apparition of a C=O vibration band at 1730 cm^{-1} in the FT-IR spectra of **20** and **21** provided additional evidence for the covalent grafting of the multifunctional oligomers bearing carbamate bounds (Fig. S42†). The difference between the loadings of di- and trimeric chains is small, which results from their (unperturbed) radii of gyration being close (the theoretical ratio being $(2/3)^{1/2} = 0.82$). In contrast, the “monomeric” units are essentially rigid rods that can be grafted to much higher densities. The content of Cu for each silica catalysts was quantified by ICP-AES, giving Cu loadings of 0.08, 0.09 and 0.10 mmol g⁻¹ for **19**, **20** and **21**, respectively. The experimental Cu contents fit well with the theoretical values calculated from the measured grafting density and the expected formation of a 1 : 1 pyta/Cu complex (0.09, 0.10 and 0.12 mmol g⁻¹ for **19**, **20** and **21**, respectively). Noticeably, owing to the reduction of grafting efficiencies with chain length, catalysts **19–21** have nearly identical Cu loadings. XPS analysis (ESI, Fig. S43†) of supported catalysts **19–21** confirmed the presence of Cu(I) species with Cu $2p_{1/2}$ and Cu $2p_{3/2}$ peaks at *ca.* 952.4 and

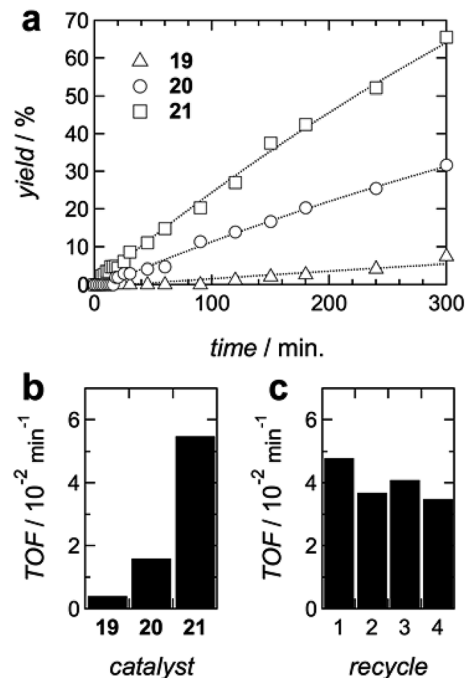


Fig. 1 (a) Activity of catalysts **19–21** (5 mol% based on Cu) in the aerobic oxidation of benzyl alcohol. Conditions: BnOH (0.2 mmol) in acetonitrile (0.2 M), O₂ bubbling (5.5 mL min⁻¹), 60 °C. (b) Initial turnover frequencies (TOFs). (c) Recycle of catalyst **21**.

932.5 eV, respectively (Fig. S44†). In addition, analysis of the high-resolution XPS C 1s region showed an increase of the high-energy components at *ca.* 286.4 eV and the apparition of a peak of higher energy at 289.0 eV in catalysts **20** and **21** compared to **19**, which are attributed to the extra electron deficient C–O and C=O carbons of the oligomers, respectively (ESI, Fig. S44†).

The supported catalysts were tested in the model aerobic oxidation of benzyl alcohol^{29,30} using a 5 mol% Cu loading (Fig. 1a). Catalyst **21**, with the mixed homo-trimer monolayer, showed the highest activity with an initial turnover frequency (TOF) more than one order of magnitude superior to the more traditional mixed monolayer catalyst **19** (Fig. 1b).

Catalyst **20** showed itself a 4-fold increase in initial TOF compared to **19**. Catalyst **21** could be easily recovered and reused for five consecutive runs with moderate erosion of its activity (Fig. 1c).

The remarkable increase of catalytic efficiency of the oligomeric chains can only be explained by the local mixing of the three catalytic units, which is provided by the flexibility of the chains combined with the larger volume of space statistically sampled by the random fluctuations of longer chains. In contrast, the rigid monomeric units do not permit an optimal pairing of the catalytic centers, resulting in a strongly depreciated TOF.

Conclusions

In conclusion, we described here the synthesis of short discrete oligomers using an iterative epoxide opening/CuAAC chemistry. Derivatization of the resulting lateral hydroxyl groups into azide



handles provided unique platforms for the integration of catalytic units. Homo-dimers and -trimers bearing the (pyta)Cu, TEMPO and imidazole active centers were obtained and grafted on mesoporous silica. The supported oligomeric catalysts surpassed the activity of a more conventional mixed monomeric version of the triad, with the trimeric catalysts showing a 14-fold enhanced TOF in the model aerobic oxidation of benzyl alcohol.

The utilization of oligomeric scaffolds opens opportunities in supported cooperative catalysis by affording ways to expand the probability of synergistic interactions within confined spaces. Further refining the tacticity of the chains and the relative arrangement of the pendant functional groups is under investigation.

Conflicts of interest

There are no conflicts to declare.

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