We describe the use of sulfonate-terminated self-assembled mono-
layers as very efficient surface-confined nucleation agents for rapid
growth of dense, thick and well-percolated ZIF-8 films.

Introduction

Metal-organic frameworks (MOFs) belong to a relatively new
class of hybrid organic–inorganic materials featuring high
surface areas, a wide range of pore sizes, pore shapes, and a
great variety of chemical functionalities exposed in the pore
walls.1–5 These interesting features are possible due to an
overwhelming number of organic linkers and metal cluster
combinations, thus yielding a very broad variety of materials
suitable for diverse applications such as; e.g., separations,6,7
gas storage,8,9 heterogeneous catalysis,10 or biological related
uses when biocompatibility is ensured.11–13 Zeolitic imidazo-
late frameworks (ZIFs) constitute a subclass of MOFs, in which
the zeolite-type topology (with its proverbial fixed Si–O–Si 145°
angle) is achieved by the formation of a geometrically similar
M–N–M bond; with N symbolizing a nitrogen atom from an
imidazolate or substituted imidazolate linker, and M a metal
ion such as Co2+ or Zn2+ tetrahedrally coordinated. In partic-
ular, ZIF-8 is an appealing material for water based applica-
tions due to its demonstrated thermal and water stability, its
high surface area (approx. 1400–1500 m<sup>2</sup> g<sup>−1</sup>), and well defined
–CH<sub>2</sub> terminated spherical hydrophobic large pores (inner
pore diameter 1.16 nm)<sup>14</sup> with small pore apertures (pore
window size 0.34 nm).15 The initial step for homogeneous
nucleation of a crystalline solid (such as ZIF-8 MOF) from a
solution of its precursors can be described as a random
process.16–18 This implies that, given that supersaturation
conditions are met, critical size nuclei leading to solid
formation do not appear immediately; and the time needed for
this process to occur is commonly referred as the induction
time. If surfaces where heterogeneous nucleation can occur
are present, then the induction time can be drastically
reduced. Moreover, the use of chemically modified surfaces
featuring terminal groups compatible with MOFs building
blocks, were reported to have both a strong effect on induction
time reduction, and also to cause preferential growth accord-
ing to a certain crystalline direction.19,20 Supported MOFs films
are widely used for sensor (usually thin films are desired),21–25
and separation applications (e.g., pervaporation membranes,
which require relatively thick films).26–28 Most common
synthetic approaches to create MOF thin films involve layer-by-
layer (LbL) growth, secondary growth (SG), and sequential one-
pot (SOP) growth. In the LbL approach precursor solutions are
kept separated, and the substrate is sequentially dipped into
linker and metal ion solutions.29–32 SG method consists in a
pre-synthesis step where MOFs nanocrystals are synthesized
and separated. Precursors are then seeded over the chosen
support (e.g., via spin-coating), and exposed to a (usually
slightly super-saturated) solution containing the precurs-
ors.33,34 This yields thick films, growing from seeded material
acting as heterogeneous nuclei. Usually, cracks can be
observed in areas where growing film segments merge
(sometimes visible by electron microscopy) causing a reduced
control over percolation. On the other hand, sequential one-
pot (SOP) strategy consists in simply dipping the substrate in
a solution of precursors during a preset reaction time and
temperature, which will depend on the MOF type. Homoge-
neous and heterogeneous nucleation processes occur simultane-
ously, and the (maximum) film thickness can be
controlled by the amount of SOP cycles performed. This
ultimately constrains the method flexibility, which is a pay-off for
the much simpler implementation.29,35 It is important to note
that SOP can be greatly improved if a surface-confined chem-
ical primer is used to enhance heterogeneous over
homogeneous nucleation. In this way, additional control over formation kinetics and morphology of the film would be gained.94† Despite the mentioned exciting developments in this field, the vast majority of ZIF-8 films grown today are exclusively based on the use of amine-terminated surfaces. However, this vision can change if we consider that Zn$^{2+}$ ions can also be strongly coordinated by sulfonate groups, thus introducing a new variable to control the heterogeneous nucleation of these materials and modulate the film growth.92 To the best of our knowledge the use of sulfonate groups as primers for enhancing ZIF-8 thin film growth remains fully unexplored. Herein, we describe for the first time the use of sulfonate-terminated self-assembled monolayers (SAMs) as very efficient nucleating agents enhancing the rapid growth of thick ZIF-8 films.

Results and discussion

Synthesis of ZIF-8 thin films was accomplished using the SOP approach from methanolic mother solutions at room temperature (see ESI† file for further details). Surfaces bearing different primers were prepared by chemisorbing SAMs of N$_2$N$_2$N$_2$-trimethyl(3-mercaptopropyl)-ammonium chloride (NTMAC), cysteamine, and 3-mercaptopropanesulfonic acid (MPSA) on gold substrates. To evaluate the influence of these contrasting functional groups on ZIF-8 films growth, in situ monitoring of growth kinetics using quartz crystal microbalance technique was carried out. Fig. 1 displays time evolution of frequency change ($\Delta f$) for films grown on freshly prepared gold surfaces. Remarkable differences can be observed, not only in terms of the mass/thickness of ZIF-8 films grown on the sensor surface, but also regarding the induction times observed, i.e.: the time required to the start of film growth after mixing of precursor solutions. As is well known the time required for triggering heterogeneous nucleation can be reduced if favourable interactions between exposed surface moieties and solvated MOFs precursor exist. The critical role of surface chemistry becomes evident if we compare the induction time for ZIF-8 films grown on bare-Au substrates (~35 min) with values obtained for substrates modified with quaternary ammonium (NTMAC, ~15 min), primary amine (cysteamine, ~10 min) and sulfonate (MPSA, ~10 min) functionalities. The relation between induction times for heterogeneous ($t_{het}$) and homogeneous nucleation ($t_{hom}$) plays a fundamental role in determining the morphology of the films obtained. Consequently, as $t_{het}$ becomes increasingly larger than $t_{hom}$ it also increases homogeneous nucleation contribution to film morphology; i.e., growth occur preferentially from seeded particles on the surface, instead of growth from precursors by direct interaction with surface. This in turn leads to less dense and scarcely percolated films. It is important to highlight that, even when similar $t_{het}$ values were observed for modified gold QCM sensors, surfaces bearing sulfonate groups exhibited a significant increase in mass. This reveals that the presence of MPSA favours a more extensive film growth, even when comparing with traditionally used primary amine-terminated surfaces (cysteamine-modified).72

Fig. 1 inset shows scanning electron microscopy images of films grown on MPSA-modified and bare Au substrates. Films grown on cysteamine-modified surfaces exhibit similar morphological features to films grown on MPSA modified substrates, while NTMAC images are similar those obtained when using bare Au (not shown, see ESI†). It is important to note that –NH$_2$ and –SO$_3$ are able to strongly interact with Zn$^{2+}$ whereas quaternary amines do not interact with any constituent of the MOF architecture. Hence this comparison illustrates the marked difference in growth kinetics when non-chelating and strongly chelating groups are employed to grow the film. It is evident that homogeneous nucleation dictates rhombic dodecahedron morphology observed for bare Au,72 while much denser films with less defined morphology were obtained for MPSA modified substrates. Further characterization of films presented in Fig. 1 was carried via powder X-ray diffraction (PXRD). Fig. 2 displays diffractograms obtained for bare and surface modified Au substrates, together with calculated ZIF-8 bulk material diffractogram. It can be observed that only films grown on MPSA- and cysteamine-modified substrates display the main features corresponding to bulk material after just one SOP synthesis cycle, while films grown over bare gold and NTMAC-modified substrates produce weak nearly amorphous diffractograms, in line with the less denser films observed via SEM (see ESI† for diffractogram evolution with increasing...
number of SOP cycles). To gain further insight into the nucleation process and the interplay between homogeneous and heterogeneous nucleation, we investigated the ZIF-8 bulk particle size time evolution by dynamic light scattering (DLS). Results presented in Fig. 3 indicate that homogeneous nucleation induction time ($t_{\text{Hom}}$) is within the range of previously QCM determined $t_{\text{Het}}$.

Moreover, DLS experiments for hydrodynamic diameter time evolution ($D_H$) indicates that heterogeneous nucleation on cysteamine and MPSA-modified Au surfaces takes place in the presence of homogeneously nucleated ZIF-8 particles of $D_H > 50$ nm ($t_{\text{Het}}$ ~ 10 min), whereas in the case of bare gold substrates exhibiting larger induction times, film growth takes places in the presence of particles with $D_H > 500$ nm. These results illustrate how important surface chemistry is to control $t_{\text{Het}}$ and, consequently, to avoid undesired side effects from homogeneous nucleation. In addition, the experimental evidence shows that it is possible to introduce drastic changes in the chemical identity of the surface, e.g.: from $-NH_2$ to $-SO_3^-$, without significantly altering the induction time for MOF film growth. Film homogeneity depends on the relative importance of heterogeneous and homogeneous growth. If heterogeneous process dominates, then a compact film could be expected; while if the opposite is true, cracks and less dense arrangements will appear. The reduction in $t_{\text{Het}}$ observed for surfaces modified with cysteamine and MPSA suggest that these primers will produce more compact films.

In order to test such hypothesis, permeation experiments were carried using cyclic voltammetry (CV) technique. To assess film percolation and compactness, CV experiments were conducted using the negatively charged Fe(CN)$_6^{3-}$/Fe(CN)$_6^{4-}$ redox probe. The inherent microporosity of ZIF-8 material features pore windows of ~0.34 nm width, which are not large enough to fit the redox probe. Due to this fact, CV experiments can be employed to evaluate the presence of loosely packed films with large intergrain mesoporosity as well as the presence of dense films in which grains are tightly packed. Given the hydrophobic nature of the ZIF-8 material, and the relative sizes of redox probe used and micropores; the permeation behavior can be rationalized in terms of the effective assembly of the ZIF-8 grains within the film. Dense arrangements of grains should lead to scarce or none intergrain mesoporosity, while the contrary presumption applies for less dense films; this would result in low and high electrochemical signals of the redox probe, respectively. As expected, the results obtained show that the films synthesized on quaternary amine-modified or bare gold surfaces are highly permeable to the redox probe. The CV results suggest that full percolation of surface-confined grains was not achieved for films grown on NTMAC-modified and bare Au surfaces (Fig. 4). On the other hand, MPSA- and cysteamine-
modified surfaces displayed a much lower electrochemical signal, which can, in turn, be related to the presence of rather compact films precluding the passage of the redox probes through the MOF layer (Fig. 4).

Conclusions

An important requirement in the preparation of interfacial architectures based on MOFs is devising strategies to control their nucleation and growth. Over the last years, sequential one-pot growth of ZIF-8 thin films has been mostly accomplished using amine-terminated surfaces as primers. But it remains a practical challenge to manipulate the chemical nature of the substrate in order to enhance the growth process without altering the structural properties of the synthesized films. In this work we showed for the first time the use of sulfonate-terminated primers as efficient surface-confined nucleating agents for rapid growth of ZIF-8 MOF.

We observed that favourable interactions between solvated MOF precursors and surface exposed moieties are crucial to manipulate the dynamics of film growth. It was demonstrated that under comparable conditions, MPSA-modified surfaces promote a remarkable ~3.5 fold increase in Δf values (and therefore on the amount of material deposited) as compared with cysteamine-modified substrates without altering the induction time for heterogeneous nucleation. Remarkably, PXRD of films obtained after just one SOP cycle over SO3−-modified and (to a lesser extent) NH2-terminated surfaces, already display ZIF-8 bulk material main features. In line with these findings, CV experiments provide evidence of a clear difference between MPSA- and cysteamine-modified well percolated films with low intergrain mesoporosity, versus bare Au and NTMAC-modified non-percolated films. From a synthetic viewpoint, introducing new primers is a particularly interesting strategy to optimize the growth of MOF thin films. Much more important—and also technologically relevant—as our array of synthetic strategies grows, so does our repertoire to “engineer” thin film architectures using metal-organic frameworks as active and functional building blocks. In this context, we believe that these results will further broaden the range of possibilities to design MOF-based hybrid materials as well as heterosupramolecular architectures.

Experimental

Zn(NO3)2·6H2O, 2-methyl-imidazole, and 3-mercapto-1-propanesulfonic acid, sodium salt (MPSA) were purchased from Sigma Aldrich. 2-Aminoethanethiol hydrochloride (cysteamine) was purchased from Acros Organics. All reagents were used as received. The N,N,N-trimethyl(3-mercaptopropyl) ammonium chloride was synthesized as described in literature.46

Film synthesis and surface modification of Au substrates

ZIF-8 films were synthesized over chemically modified Au substrates using methanolic solutions of precursors according to previously published procedure.48 Briefly, it consists on the following steps: (i) Au substrate was modified using solutions of the selected alkanethiol (concentrations used were 5 mM cysteamine in ethanol, 1 mM NTMAC in ethanol, and 20 mM MPSA in 10 mM H2SO4) at room temperature overnight. Treated substrates were, then, rinsed, and dried with N2, (ii) clean modified Au substrates were immersed in a fresh mixture of Zn(NO3)2·6H2O (10 mL, 25 mM) + 2-methyl-imidazole (10 mL, 50 mM) stock solutions for appropriate time according to the primer used (see Section 3) at room temperature, (iii) substrates whereas then rinsed with fresh methanol and dried with N2 prior to the next cycle.

Film characterization

ZIF-8 film growth over bare and modified Au substrates was followed using a Quartz Crystal Microbalance (QCM 200) from Stanford Research Systems. For post-synthesis characterization morphology of the films was studied by scanning electron microscopy (ESEM FEI-Quanta 200) and powder X-ray diffraction (Phillips X’Pert X-Ray Diffractometer); size analysis were assessed by DLS using a Zetasizer Nano ZS from Malvern.

Cyclic voltammetry experiments

Cyclic voltammetry experiments were carried out using a Reference 600 potentiostat from Gamry, with a three-electrode setup. Ag/AgCl and Pt wire were used as reference electrode and counter electrodes, respectively. In all the electrochemical experiments 1 mM Fe(CN)6 3−/4− + 100 mM KCl solutions were used.

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Notes and references
