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Graphical abstracts
39x19mm (300 x 300 DPI)
Synthesis, characterization and evaluation of hollow molecularly imprinted polymers for Sudan I

Dong Ren*, Jiang He**, Haixia Zhangab

A novel strategy was developed to prepare hollow molecularly imprinted polymers (HMIPs) with thin and solid shell, in which introduced soft polystyrene core and hard inner shell of SiO, and combined surface molecular imprinting of Sudan I and in situ polymerization. The HMIPs possessed good morphological stability without deformation and broken owing to the existing of SiO. The thin imprinted coating (50 nm) of HMIPs ensured the faster mass transfer and the higher efficiency of active sites utilization during the adsorption. What’s more, it was successfully applied as solid-phase extraction (SPE) sorbent for selective adsorption of Sudan I in chilli powder samples, and the satisfied recoveries were obtained in the range of 95-108% with the spiked samples.

1. Introduction

In recent years, molecular imprinting has been considered as a promising method for synthesizing the materials with memory of the shape and size for template molecule. The materials obtained were named as molecular imprint polymers (MIPs) [1-5]. Owing to the easy preparation, good stability and recognition properties, MIPs have been applied in many fields for the separation and enrichment of target molecules, especially acted as solid phase extraction (SPE) sorbents in sample pretreatment [6-8]. In order to meet the needs in practical applications, various synthesis strategies have been developed, including bulk polymerization, precipitation polymerization, and suspension polymerization etc. Although the MIPs prepared using the above methods exhibited good binding affinity and specificity toward target molecules, they involved in some shortcomings, such as incomplete template removal, low utilization ratio of binding sites, and slow mass transfer because of their highly cross-linked nature [9].

To overcome the above shortages mentioned, methods such as surface imprinting [10], porous imprinting [11], and hollow imprinting have been developed [12-18]. The hollow imprinting polymers (HMIPs) have aroused extensive attention especially, owing to their larger specific surface area, higher utilization ratio of binding sites, and faster mass transfer. There are two main methods for synthesis of HMIPs with sacrificing soft or hard core, respectively. For method with sacrificing soft core, divinylbenzene (DVB) [12-14] was usual as cross-linking agent and polystyrene (PS) particle as core. Because the dissolving and removal of the core was accompanied with its swelling, which led to the collapse of the polymer outside, the thicker imprinting shell had to be made. However, the thick imprinting shell led to low mass transfer and low utilization ratio of binding sites. For method with sacrificing hard core, SiO or TiO were usual as the core [15-18]. With this kind of method, thin imprinting shell (50-80 nm) could be obtained. But, the thin shell was fragile and easy to be broken.

Sudan dyes (There structures are shown in Fig. S1, ESI†) are phenyl-azoic derivatives, with an orange-red appearance [19]. Due to azo dyes considered to be genotoxic carcinogen and mutation for humans, their presence are not permitted in foodstuffs for any purpose at any level [20]. Sudan I was the most common dye used illegally as additives to enhance the gaily-colored appearance of food products containing chilli, curry, curcuma, palm oil and so on [21,22]. Therefore, it is important to inspect the presence of the Sudan I in foodstuffs. A lot of analytical methods have been developed for the determination of Sudan dyes [23-26]. Owing to the complexities of sample matrixes and the low levels of Sudan I, sample pretreatment and enrichment processes have become the crucial steps in the analytical procedures [27-28].

Up to now, some MIPs for Sudan I had been prepared with different methods [28-33]. Among them, no HMIPs have been prepared. In this work, it was the first time to design a new approach to prepare a novel HMIPs with thin and solid MIPs shell. We used PS/SiO particles as core and only Ps part was sacrificed. It is to say, SiO part was kept in the HMIPs as the support, which made it possible to get a thin but solid MIPs shell. The HMIPs prepared was not only avoided the deformation and broken, but also offered high utilization ratio of binding sites and faster mass transfer. The HMIPs were fully characterized and evaluated by scanning electron microscopy (SEM), transmission electron microscopy (TEM), thermogravimetry analysis (TG) and Fourier transform infrared spectrometer (FT-IR). Their molecular recognition capacities were also investigated in detail. The HMIPs was further used as SPE sorbents to selective adsorb and enrich Sudan I from chilli power samples and the satisfied recoveries were obtained.

2. Experimental

2.1 Reagents and chemicals
Methacrylic acid (MAA), acetonitrile (ACN), tetrahydrofuran (THF), styrene, ammonia (25-28%), toluene, and chloroform were from Tianjin Guangfu Fine Chemical Research Institute (Tianjin, China). Ethylene glycol dimethacrylate (EGDMA), tetraethoxysilane (TEOS), 3-methacryloxypropyl trimethoxysilane (MATMS), and 3-aminopropyl triethoxysilane (APTES) were purchased from Alfa Aesar (Beijing, China). Potassium peroxysulfate (K2S2O8) and Azobis-isobutyronitrile (AIBN) were obtained from Chemistry Reagent Factory of Chinese Fuchen (Tianjin, China). Sudan I-IV were purchased from Sinopharm Chemical Reagent Co. Ltd (Shanghai, China). All the chemicals above were of analytical grade. Chilli powder samples were bought from a local market in Lanzhou, China. Ultra pure water was used throughout the whole experiments.

2.2 Instruments

The morphologies of prepared polymers were observed by a JEM-1200EX TEM (Tokyo, Japan), and an S-94800 SEM (Hitachi, Japan). Infrared spectra were collected on a Nicolet 20 NEXUS 670 FT-IR spectrometer (Ramsey, MA, USA). The TG analysis over 50-800 °C was obtained by a STA PT1600 Thermal Analyzer Instruments (Linscis, Germany) with the heating rate of 10 °C min⁻¹ under N2. The chromatographic analytical system consisted of a Model 210 HPLC pump and a UV-Vis detector (Varian Prostar, USA). All separations were carried out on a C18 column (Dikma Technologies, 250×4.6 mm, 5 μm). The UV-Vis detector was operated at 478 nm for Sudan I and 520 nm for Sudan II-IV. The optimal HPLC conditions for Sudan dyes were as follows: mobile phase: ACN-water 95.5 (v/v); flow rate: 1.0 mL min⁻¹; room temperature; injection volume: 20 μL [33].

2.3 Preparation of hollow molecularly imprinted polymer for Sudan I (HMIPs)

2.3.1 Synthesis of PS/SiO2 spheres

First, monodisperse polystyrene (PS) particles (500 nm in diameter) were synthesized by emulsifier-free emulsion polymerization method according to the previous report with some modification [9]. Styrene monomer (4.0 g) was added to 60 mL of water under stirring for 10 min at 70 °C in N2 atmosphere. And then 4.0 mL of K2S2O8 aqueous solution (23.0 mg mL⁻¹) was added. After the reaction was refluxed for 24 h, the monodisperse PS spheres were obtained after centrifugation (8000 rpm, 10 min) and washing with water for several times. Then, the PS spheres were dispersed in 60 mL of concentrated sulfuric acid (H2SO4), the sulfonation reaction was allowed to take place at 40 °C for 4 h. The obtained Sulfonated PS (SPS) spheres were purified by repeat washing with water and ethanol until the pH reached to 6.0.

The obtained spheres were dispersed in 50 mL of water again and 0.5 mL of APTES monomer was added. The mixture was kept stirring for 12 h at 25 °C, and then centrifuged and washed with ethanol to remove the unreacted APTES. The precipitate obtained was dispersed in a solution consisted of 50 mL of ethanol, 10 mL of ammonia and 10 mL of water, followed by adding 1.5 mL of TEOS dropwise under stirring. The reaction was kept for 3 h at 50 °C. The PS/SiO2 spheres were obtained after centrifugation and washing with ethanol [34].

2.3.2 Synthesis of MATMS functionalized hollow SiO2 (MHSiO2)

The surface of SP/SiO2 spheres was modified further with MATMS by following. The SP/SiO2 spheres were dispersed in the solution with 100 mL of ethanol, 1.0 mL of ammonia and 0.2 mL of MATMS, and then the mixture was stirred for 24 h at 25 °C. After the products were centrifuged and washed with ethanol for several times, they were dispersed in THF for 2-3 h to dissolve the PS core. The final products MHSiO2 were dried in vacuum at 50 °C for 24 h.

2.3.3 Synthesis of HMIPs for Sudan I

Typically, Sudan I (24.8 mg, 0.1 mmol) and MAA (34.5 μL, 0.4 mmol) were added in 24 mL of toluene and stirred for 8 h in dark to finish the pre-polymerization. Then under stirring, 375 μL (2.0 mmol) of EGDMA, 20 mg of AIBN, 1 mL of chloroform and 160 mg of MHSiO2 spheres were added in the pre-polymerization solution. After deoxygenized with N2 for 5 min, the polymerization was undertaken at 60 °C for 12 h under N2 protection. The products were collected and washed with ACN. The template molecules were removed by Soxhlet extraction in methanol-acetic acid solution (90:10, v/v) until no Sudan I could be detected by HPLC from the HMIPs. The HMIPs were finally dried in vacuum for 24 h at 50 °C. The preparation route was shown in Scheme 1. For comparison, non-imprinted hollow polymer (HNIPs) was prepared with the same procedure except for the presence of Sudan I.

(Scheme 1)

2.4 Estimation of recognition properties of HMIPs

In order to evaluate the binding capacity of the HMIPs, a static adsorption experiment was performed first by placing 20 mg of HMIPs into 2 mL of ACN containing Sudan I with various concentrations. The mixture were shaken for 24 h at room temperature, then the supernatant solution was collected, the concentrations of Sudan I...
were detected by HPLC method. Every test was done for parallel three times and the experimental data were the mean values of the results.

The data of the static adsorption experiment were further processed with the Langmuir isotherm models (Eq. (1)) and Freundlich isotherm models (Eq. (2)) to estimate the binding parameters of HMIPs prepared [35, 36]. \( Q (\mu\text{mol} g^{-1}) \) was the amount of Sudan I bound to MIPs at equilibrium. \( Q_{\text{max}} (\mu\text{mol} g^{-1}) \) was the maximum binding capacity, \( C_{\text{free}} (\mu\text{mol} \text{L}^{-1}) \) was the equilibrium concentration of Sudan I, \( K_L (\text{mL} \mu\text{mol}^{-1}) \) was the Langmuir binding coefficient, \( K_f (\mu\text{mol} g^{-1}) (\mu\text{mol} g^{-1})^{n/2} \) was the Freundlich binding coefficient and \( n \) was the Freundlich binding constant.

\[
\frac{C_{\text{free}}}{Q} = \frac{C_{\text{free}}}{Q_{\text{max}}} + \frac{1}{K_L Q_{\text{max}}} \tag{1}
\]

\[
\log Q = \log K_L + \frac{1}{n} \log C_{\text{free}} \tag{2}
\]

Then a dynamic adsorption test for Sudan I was carried out. 200 mg of HMIPs were dispersed in 20 mL of ACN containing 1.20 \( \mu\text{mol} \text{L}^{-1} \) or 0.16 \( \mu\text{mol} \text{L}^{-1} \) of Sudan I, respectively. The mixtures were shaken at 25 °C in a thermostat oscillator. At different time interval the concentration of Sudan I in solution was determined. The data were further processed with pseudo-first-order kinetic model (Eq. (3)) and pseudo-second-order kinetic model (Eq. (4)). Where \( Q_e (\mu\text{mol} g^{-1}) \) was the equilibrium uptake, \( Q_t (\mu\text{mol} g^{-1}) \) was the adsorption capacity at different time, \( k_1 (\text{min}^{-1}) \) was the first-order rate constant, \( k_2 (g \mu\text{mol}^{-1} \text{min}^{-1}) \) was the second-order rate constant, \( t (\text{min}) \) is the adsorption time, and \( v (\mu\text{mol} g^{-1} \text{min}^{-1}) \) was initial sorption rate \( (v=k_2Q_e^2) \).

\[
\log(Q_e - Q) = \log Q_e - \frac{kt}{2.303} \tag{3}
\]

\[
\frac{t}{Q_e} = \frac{1}{k_2Q_e^2} + \frac{t}{Q_e} \tag{4}
\]

Third, the selectivity of HMIPs towards Sudan I was evaluated with a series of experiments on HMIPs and HNIPs, using 0.40 \( \mu\text{mol} \text{L}^{-1} \) of Sudan I and its structural analogs (Sudan II-IV). The experiment process was same as the static adsorption experiment.

2.5 Analysis of chilli powder samples

Chilli powder samples were used to demonstrate the applicability of the HMIPs to enrich Sudan I from complicated matrices. Extraction of Sudan I from 2.0 g of spiked or non-spiked chilli powder was carried out by adding 4 mL of n-hexane. The mixture was shaken for 5 min and the supernatants were collected. The residue was washed with n-hexane for another twice. The extracts were pooled together, filtrated with 0.22 \( \mu\text{m} \) filter and diluted to 25.0 mL.

An empty SPE cartridge was packed with 400 mg of the HMIPs, which was conditioned sequentially with 5 mL of methanol and 10 mL of n-hexane at a flow rate of 1.0 mL min\(^{-1}\), and then the cartridge was loaded with the prepared samples at a flow rate of 5 mL min\(^{-1}\). After loading, the column was washed with 4 mL of methanol at 1 mL min\(^{-1}\). Finally, the elution was performed by passing 6 mL of methanol-acetic acid solution (90:10, v/v) [12]. The eluant was collected and evaporated under N\(_2\). The residue was dissolved in 200 \( \mu\text{L} \) of mobile phase for HPLC analysis.

3. Result and discussion

3.1 Preparation of HMIPs

In our preparation, how to coat PS sphere with SiO\(_2\) shell was the key step. Without the synthesis of SPS spheres, it was failed to form the PS/SiO\(_2\) spheres. So the PS sphere was first sulfonated with H\(_2\)SO\(_4\) to form sulfo groups on its surface and then APTES monomer was introduced through the bonding between the sulfo groups and the amino groups, which led further to the successful formation of SiO\(_2\) shell.

The ratio of template molecules and functional monomers, the type of cross-linkers and the kind of solvents were taken into consideration in the preparation of HMIPs. MAA was chosen as the functional monomer because it could form hydrogen bond with Sudan I. The molar ratios of template molecule, functional monomer and cross-linker were set at 1:4:20 according to the references [30, 33]. The HMIPs were prepared in toluene or chloroform to investigate the influence of solvent. As can be seen from table 1, toluene was superior to chloroform as the solvent for the synthesis of HMIPs for Sudan I.
3.2 Characterization of the HMIPs

The SEM and TEM images of PS/SiO\(_2\), MHSiO\(_2\), HMIPs were shown in Fig. 1. As shown in Fig. 1A and Fig. 1a, the PS/SiO\(_2\) spheres with an average diameter of 550 nm were uniform and their surface was smooth. Fig. 1B and Fig. 1b showed the images of the MHSiO\(_2\) spheres, which were obtained by removing the PS core with THF, and the shell thickness of the MHSiO\(_2\) was about 15 nm. A hole was present on the SiO\(_2\) shell. In Fig. 1C and Fig. 1c, the MIP coating was successfully synthesized onto the MHSiO\(_2\) surface and the thickness of the imprinted shell was estimated about 50 nm. It could be seen the HMIPs were rigid and kept perfect spherical shape without breakage and deformation. The hole on the SiO\(_2\) sphere was survived, which ensured that the template molecules could enter the imprinted cavities from both sides of the polymer spheres.

FT-IR spectra were employed to ascertain the successful synthesis of materials (Fig. S2, ESI†). In the spectrum of SPS particles (Fig. S2a), the typical PS absorption bands at 1600, 1492, 1451, 755, and 698 cm\(^{-1}\) were clearly seen [37]. The band at 1068 cm\(^{-1}\) was ascribed to the -SO\(_3\)H group, confirming that the PS spheres were doped with -SO\(_3\)H directly [34]. In the spectrum of PS/SiO\(_2\) particles (Fig. S2b), the asymmetrical stretching vibration peak of siloxane bond (Si-O-Si) was found at 1087 cm\(^{-1}\); and the peaks at 960 cm\(^{-1}\) and 3418 cm\(^{-1}\) were from the stretching vibration of the hydroxyl group in Si-OH, which confirmed that the SiO\(_2\) shell was successfully modified on the surface of PS. The peak of C=O at 1713 cm\(^{-1}\) and the C-O asymmetric stretching vibration of MATMS at 1192 cm\(^{-1}\) in M/PS/SiO\(_2\) spectrum (Fig. S2c) showed the MATMS had been grafted onto the surfaces of the PS/SiO\(_2\). The typical bands of PS were disappeared in the spectrum of MHSiO\(_2\) (Fig. S2d), which indicated that the PS core was removed by THF successfully. HMIPs and HNIPs exhibited strong absorption bands around 1728 cm\(^{-1}\), 1253 cm\(^{-1}\), and 1454 cm\(^{-1}\), which were assigned to the C=O stretching vibration from MAA and EGDMA, the C-O symmetric stretching vibration and the CH\(_2\) stretching vibration from EGDMA, respectively [39,40].

TG analysis results were obtained (Fig. S3, ESI†). PS/SiO\(_2\) could be table below 371 °C. When the temperature was increased to 800 °C, the weight loss was increased to 77.6%, which was resulted from the loss of PS core. For the HMIPs and HNIPs, the weight loss of water below 265 °C was observed. In the range of 265-465 °C, the polymers on the surface of HMIPs and HNIPs were decomposed seriously. At 800 °C, the residue amounts (SiO\(_2\)) were only 5.7% and 7.3% for HMIPs and HNIPs, respectively.

3.3 Binding profiles of the HMIPs

Fig. 2 showed the adsorption isotherm of HMIPs and HNIPs for Sudan I in ACN. It can be noted that the adsorption capacities of HMIPs and HNIPs increased with the increased initial concentration of Sudan I from 0 to 2 μmol mL\(^{-1}\). The imprint factors \(\alpha (Q_{\text{HMIP}}/Q_{\text{HNIP}})\) were near each other at the different initial concentrations, which were in a range of 2 to 3.

The Langmuir and Freundlich isotherm models were employed for studying the adsorption thermodynamics data (Table 2). The linear coefficients (R\(^2\)) were close each other, which indicated that both the monolayer and multilayer adsorption might be coexisted in the complex adsorption process. The maximum adsorption capacities (\(Q_{\text{max}}, \text{μmol g}^{-1}\)) were calculated to be 12.5 for the HMIPs and 6.32 for HNIPs. The Freundlich binding constants (\(n\)) of the HMIPs and HNIPs were 1.51 and 1.39, respectively. \(n\) was larger than 1, indicating the adsorption was easy to occur. The Freundlich binding coefficient (\(K_r, \text{μmol}g^{-1}(\text{mL} \text{μmol}^{-1})^{1/n}\) ) of the HMIPs (10.35) was about 2 times of the HNIPs (4.05), certifying further that the HMIPs possessed higher adsorption capacity than the HNIPs owing to the imprinting effect.

Fig. 3 depicted the kinetic curve of Sudan I on the HMIPs and HNIPs. The adsorption amount of HMIPs and HNIPs increased sharply during the first 10 min for the initial concentration of 0.16 μmol mL\(^{-1}\) or 60 min for 1.21 μmol mL\(^{-1}\) and then reached stable, which meant the equilibration time was depended heavily on the initial concentration. Compared with the similar hollow MIPs materials reported in references, it was found the new HMIPs prepared in the work had the advantage of having faster mass transfer rate compared with [12, 13] and not crushed easily compared with [17, 18]. The detailed comparison with the references was summarized in table 3.

Pseudo-first-order kinetic model and pseudo-second-order kinetic model [41] were used to study the kinetics procedure. For both HMIPs and HNIPs, the experimental data were well fitted to the pseudo-second-order model with a correlation coefficient (R\(^2\)=0.9993). The \(Q_e\) value (μmol g\(^{-1}\)) for the HMIPs was calculated as 11.5, which was near 2 times higher than of the HNIPs (6.16). The v value (μmol g\(^{-1}\) min\(^{-1}\)) for HMIPs was 0.695, which was higher than from the HNIPs (0.432), suggesting that the HMIPs had a higher adsorption rate.

The \(Q_e\) value (μmol g\(^{-1}\)) calculated from Freundlich isotherm model was 12.5 and 11.5 from...
pseudo-second-order kinetic model, which indicated that 92% of binding sites were utilized in the adsorption procedure. The very high utilization ratio of binding sites resulted to the larger adsorption capacity compared with the other HMIPs with PS core [16,17].

3.4 Selectivity of the HMIPs

Solvent could affect the interaction between the template molecules and HMIPs. Pure ACN, 90% and 40% ACN in water solution were chosen as the solvents to carry out the selectivity tests. H₂O could destroy the special binding sites by reducing the hydrogen bond between the binding sites and the template molecules and led to the decreased imprint factor α (1.5 times) for HMIPs but the increased adsorption capacity was obtained (1.6 time) for both HMIPs and HNIPs owing to the decreased solubility of Sudan I in the solution.

Sudan dyes had the very similar structures and Fig. 4 showed the adsorption capacities of the HMIPs and HNIPs to Sudan I-IV in ACN. The HMIPs showed the selectivity factors k (the ratio of α_{Sudan I} and α_{other Sudan dyes}) were between 1.5-2.0, which was larger than our previous report (no more than 1.5) [33]. The higher k values to the substances with similar structures meant the better discernment capacity of the HMIPs, which maybe benefited from the thin shell of HMIPs. Whatever HMIPs or HNIPs, they showed the lowest binding capacity to Sudan II, the possible reason was the poor structural planarity of Sudan II hindered its adsorption.

3.5 Analysis of chilli powder samples

A SPE-HPLC method was developed to determine Sudan I in chilli powder samples, which could attain favorable limits of detection (LOD) of 4.0 μg kg⁻¹ based on the signals as 3-fold the baseline noise under the optimal conditions (shown in experimental). Chilli powder samples spiked with different amount of Sudan I at 0, 4, 10 and 100 μg kg⁻¹ were determined and the results were shown in Table 4. A black experiment was done with the same procedures to evaluate the template leakage. As seen, the template of Sudan I had been removed completely. Sudan I was found in the real chilli powder samples to be 7.0 μg kg⁻¹. For the spiked chilli powder samples recoveries ranged from 95-108% were obtained with high precision (RSD<5%).

3.6 Regeneration studies of HMIPs.

The regeneration of HMIPs was considered to have a great cost benefit on extending their applications. A HMIPs-SPE cartridge was repeated for usage to investigate the reuse capacity. After each use, the cartridge was washed by methanol-acetic acid solution 90:10 (v/v) and methanol. Six cycles were performed and no less recovery was found. The HMIPs were taken out for SEM images and the images before and after usage were compared (Fig. S5, ESI†). It could be seen that after six cycles, the imprinting shell was still stable without broken.

4. Conclusion

A novel strategy was successfully developed to prepare HMIPs, in which introduced both soft PS core and hard SiO₂ inner shell with the help of sulfonation of PS, which ensured to obtain the solid and thinner MIPs shell.

Thinner MIPs shell provided fast mass transfer rate and the residual SiO₂ offered the support for increasing the rigidity of MIPs. The HMIPs could be used repeatedly as SPE sorbent at least 6 times without decreasing the adsorption capacity and being broken. It was expected to be used to enrich Sudan I from the complex samples.

Acknowledgements

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Notes and references

1. State Key Laboratory of Applied Organic Chemistry, Lanzhou University, Lanzhou 730000, P. R. China
2. Testing Demonstration Center for Food quality & safety of the Ministry of Industry & Information, Lanzhou 730000, P. R. China
3. *Corresponding author: Jiang He, Tel.: +86 (931) 891 2591, Fax: +86 (931) 891 2582, E-mail: hejiang@lzu.edu.cn
4. † Electronic Supplementary Information (ESI) available: The structures of Sudan dyes; Characterization and performance evaluation results of HMIPs.
Fig. 1. TEM images (A-C) and SEM images (a-c) of PS/SiO_{2}, MHSiO_{2}, and HMIPs.

Fig. 2. Adsorption isotherm of HMIPs and HNIPs for Sudan I in ACN. Experimental conditions: v=2.0 mL; mass of polymers, 20 mg; adsorption time, 24 h.

Fig. 3. Kinetic uptake of Sudan I onto the HMIPs and HNIPs. Experimental conditions: v=20 mL; mass of polymers, 200 mg.

Fig. 4. Uptake of different Sudan dyes at the same initial concentration of 0.40 μmol mL^{-1}. Experimental conditions: solvent, ACN; v=2 mL; mass of polymers, 20 mg; adsorption time, 1 h; structure references, Sudan I-IV.
Scheme 1. Schematic illustration of the preparation process of HMIPs.

Fig. 1. TEM images (A-C) and SEM images (a-c) of PS/SiO₂, MHSiO₂, and HMIPs.
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conditions: solvent, ACN; v=2 mL; mass of polymers, 20 mg; adsorption time, 1 h; structure references, Sudan I-IV.

Table 1. Comparison of the adsorption capacities of materials synthesized in different solvents

<table>
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<tr>
<th>Solvent</th>
<th>(Q_{\text{HIMPs}})</th>
<th>(Q_{\text{HNIPs}})</th>
<th>(\alpha)</th>
<th>(Q_{\text{HIMPs}})</th>
<th>(Q_{\text{HNIPs}})</th>
<th>(\alpha)</th>
<th>(Q_{\text{HIMPs}})</th>
<th>(Q_{\text{HNIPs}})</th>
<th>(\alpha)</th>
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<tr>
<td>Toluene</td>
<td>9.03</td>
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<td>2.33</td>
<td>10.22</td>
<td>4.49</td>
<td>2.27</td>
<td>9.94</td>
<td>4.57</td>
<td>2.18</td>
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<tr>
<td>Chloroform</td>
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<td>4.68</td>
<td>1.22</td>
<td>7.41</td>
<td>4.66</td>
<td>1.59</td>
<td>7.13</td>
<td>4.37</td>
<td>1.63</td>
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</table>

\(Q:\ \mu\text{mol g}^{-1}\), \(\alpha=\frac{Q_{\text{HIMPs}}}{Q_{\text{HNIPs}}}\). Amount of materials: 20 mg; solvent: 2 mL of ACN; adsorption time: 60 min; room temperature. *initial concentration of Sudan I.

Table 2. Isotherm model constants for HMIPs and HNIPs

<table>
<thead>
<tr>
<th>Materials</th>
<th>Langmuir</th>
<th>Freudlich</th>
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<tr>
<td></td>
<td>(R^2)</td>
<td>(Q_{\text{max}})</td>
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<td>HMIPs</td>
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<td>12.5</td>
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<tr>
<td>HNIPs</td>
<td>0.989</td>
<td>6.94</td>
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Table 3. Comparison of morphological stability and mass transfer of HMIPs with reported

<table>
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<tr>
<th>Template molecule</th>
<th>Sacrificed core</th>
<th>Shell thickness (nm)</th>
<th>Morphological stability</th>
<th>(^a) adsorption capacity ((\mu\text{mol nm}^{-1}))</th>
<th>Concentration of template ((\mu\text{mol mL}^{-1}))</th>
<th>Equilibration time (min)</th>
<th>Ref.</th>
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<td>12</td>
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<tr>
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<td>0.5</td>
<td>50</td>
<td>16</td>
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<tr>
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<td>TiO(_2)</td>
<td>/</td>
<td>crash</td>
<td>-</td>
<td>2.7</td>
<td>240</td>
<td>17</td>
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<td>-</td>
<td>0.54</td>
<td>60</td>
<td>18</td>
</tr>
<tr>
<td>Sudan I</td>
<td>PS</td>
<td>50</td>
<td>Stable</td>
<td>0.25</td>
<td>0.5</td>
<td>25</td>
<td>this work</td>
</tr>
</tbody>
</table>

\(^a\) adsorption capacity: adsorption amount on 1 nm of HMIP thickness.
Table 4. Recoveries of Sudan I in chilli powder samples (n=3).

<table>
<thead>
<tr>
<th>Chilli powder (g)</th>
<th>Sudan I added (μg kg⁻¹)</th>
<th>Found (μg kg⁻¹)</th>
<th>Recoveries (%)</th>
<th>RSD (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>-</td>
<td>Not found</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>2</td>
<td>0</td>
<td>7.0</td>
<td>-</td>
<td>5.8</td>
</tr>
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<td>4</td>
<td>11.3</td>
<td>107.5</td>
<td>3.0</td>
</tr>
<tr>
<td>2</td>
<td>10</td>
<td>16.9</td>
<td>99.0</td>
<td>1.8</td>
</tr>
<tr>
<td>2</td>
<td>100</td>
<td>102.9</td>
<td>95.9</td>
<td>1.1</td>
</tr>
</tbody>
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