Functionalized Mesoporous Silicon Nanomaterials in Inorganic Soil Pollution Research: Opportunities for Soil Protection and Advanced Chemical Imaging

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Abstract

“Innovative actions towards a pollution free-planet” is a goal of the United Nations Environment Assembly (UNEA). Aided by both the Food and Agricultural Organisation (FAO) and its Global Soil Partnership under the 3rd UNEA resolution, a consensus from > 170 countries have agreed a need for accelerated action and collaboration to combat soil pollution. This initiative has been tasked to find new and improved solutions to prevent and reduce soil pollution, and it is in this context that this review provides an updated perspective on an emerging technology platform that has already provided demonstrable utility for measurement, mapping, and monitoring of toxic trace elements (TTEs) in soils, in addition to the entrapment, removal, and remediation of pollutant sources. In this article, the development and characteristics of functionalized mesoporous silica nanomaterials (FMSN) will be discussed and compared with other common metal scavenging materials. The chemistries of the common functionalizations will be reviewed, in addition to providing an outlook on some of the future directions/applications of FMSN. The use of FMSN in soil will be considered with some specific case studies focusing on Hg and As. Finally, the advantages and developments of FMSN in the widely used diffusive gradients-in-thin films (DGT) technique will be discussed, in particular, its advantages as a DGT substrate for integration with oxygen planar optodes in multilayer systems that provide 2D mapping of metal pollutant fluxes at submillimeter resolution, which can be used to measure detailed sediment-water fluxes as well as soil-root interactions, to predict plant uptake and bioavailability.

Keywords Functional mesoporous silicon nanomaterials (FMSN) · Diffusive gradient in thin films (DGT) · X-ray fluorescence spectrometry (XRF) · Heavy metals · Soil pollution
Introduction to FMSN

The Development of FMSN and Its Principles of Adsorption

Functionalized mesoporous silica nanomaterials (FMSN) can be ordered by the physical/structural characteristics of the inert hosting material and/or the chemistries of the functionalization. Both factors control the behavior of the material and its suitability for different tasks and conditions. This ability for optimization and fine-tuning has made FMSN an increasingly promising technology for medical and environmental applications. The most popular configurations have a tendency for large pore volumes but variable pore sizes, high surface areas, and a stability that makes them reliable even in challenging matrices (low pH, high organic concentrations) and environmental conditions (low–high temperatures) [1–3]. The crystal state of FMSN is shown in Fig. 1. To develop a better understanding of FMSN, it is instructive to consider how they have developed, why, and the technologies that they have superseded.

Research on FMSN has been ongoing for ca. 30 years. In the early 1990s, the initial aim was to expand beyond the confines of zeolite minerals used for ion exchange and contaminant sorption. These hydrated aluminosilicates, comprising of a tetrahedra patchwork of alumina and silica structures, were the most widely used metal/elemental scavengers. Effective as they were, however, they had their limitations, and consequently, there was a demand for new materials (mesoporous silica) with more structured/controlled pores of sizes between 2 and 10 nm. These designs were needed to accommodate molecules that could not be captured by existing zeolites because they were not small enough to fit in the micropores [4]. It was not until 1992 that the first scanning electron microscopy (SEM) micrograph of mesoporous silica appeared, yielding a greater appreciation of their detailed/intricate structure and how synthesis conditions governed morphogenesis [5]. In 1998, Stucky and colleagues synthesized mesoporous silica with larger pores (30 nm), further expanding the range of molecules accessible to the interiors of the Si crystals [6] and opening up new opportunities for functionalization.

It is also worth noting that FMSN have benefited from the increase of interest in nanotechnologies. In accordance with the International Organization for Standardization definition

Fig. 1 Copper removal by iMoLoX FMSN and its high absorbing capacity. a Partitioning of Cu from 15 ml 98% anhydrous ethanol solution (pH 4–6) containing 30 mg L\(^{-1}\) CuSO\(_4\) into 0.025 g FMSN, removal efficiency reached 92.97%. The color change of the solution is caused by bromophenol blue (0.1–0.3 ml). The color of FMSN changed from colorless to blue after the adsorption was completed. b Si crystal image before and after palladium recovery from a solution (pH 2–5) of 250 mg L\(^{-1}\) PbCl\(_2\), c and d Si dosed with 30 mg L\(^{-1}\) CuSO\(_4\) solution (pH 4–6). e Si crystal image before and after iron recovery from a solution (pH 3–5) of 150 mg L\(^{-1}\) FeCl\(_3\)/FeCl\(_2\). Si crystal particle size is 180–600 microns. The images of FMSN particle were obtained through compound optical microscope (LEICA DM500), provided by GoldenKeys High-tech Materials Co., Ltd.
in ISO/TS 80004, nanomaterials encompass not just materials with any external dimension between 1 and 100 nm but those “having internal structure or surface structure in the nano-scale” [7]. With the development of nanomaterials, researchers have discovered the potential of functionalized silica as an advanced chemical material [8, 9]. Combining an ease of surface modification and synthesis, good biocompatibility and stability FMSN quickly became a viable replacement/alternative to zeolites [10].

The other metal-removing agent traditionally used is activated carbon (AC). It has been successfully used in many industrial applications. AC has been applied successfully before to remove As, Cr, and pharmaceutical compounds from water [11–13]. It is also applied for greenhouse gas capture [14], electrical energy applications (as an ultra-capacitor) [15–17], and as an air pollution mitigator [18, 19].

The adsorption capacity traits associated with AC depend principally on their porosity and surface area [20]. Two traditional methods are used to prepare AC that delineate by either physical or chemical activation. Physical activation involves carbonization of organic matter, and then, the resulting char is modified in the presence of an activator (such as CO₂ or steam). In chemical activation, the activating agents (such as K₂CO₃) is used directly for raw material impregnation. Next, heat treatment is applied in an inert atmosphere (various/high temperatures) [20]. Generally, compared with physically activated AC, chemically activated AC has a larger surface area and a smaller pore size, with a wider range of applications [21].

Activated carbon in general has a high internal surface area due to its special sponge-like structure. Moreover, these adsorbents have reasonable chemical stability [22]. Suitable natural ingredients are continually being sought to make low-cost AC materials. To date, AC has been formed successfully from agricultural wastes as diverse as apricot and bagasse, grape seeds and cherry stones, rice husks, coconut, tomato processing solid waste, etc. [12], which highlights its versatility.

The main benefit associated with AC as a clean-up technology for use in wastewater is its overall chemical flexibility and universality. This stems from its large specific surface area, adsorption capacity, wetting characteristics, and heterogeneous porous structure; all of this can be tailored via regulation of the physical/chemical activation steps [23]. The general adsorption capacity of AC depends principally on physical entrainment and then secondly by chemical properties. However, for metal adsorption specifically, the surface chemistry and configuration of outer ligands on the AC are of more importance, providing some element/species selectivity [24, 25]. The problem, though, is that the distribution of functional groups can also change the structure of the AC [24, 25]. This complicates attempts to further functionalize the char, as chemical changes can similarly cause physical modification, with potentially behavior-altering changes for the metal scavenger. Ideally, an inert/stable host is required for functionalization, as this enables better control of the specific chemistries, which is the key to superior performance. Functionalized Si meets these requirements, and this has been partly the catalyst for the rapid development of FMSN.

Many popular metal clean-up methods rely on materials that utilize ion exchange as the main binding mechanism. Here, the process centers on the exchange of ions between a solid and an associated liquid phase [26]. The solid phase is called the ion exchange resin/ion exchanger, because it carries exchangeable ions, insoluble in the liquid phase and during chemical reactions/interactions maintains its structural integrity [27]. The most useful ion exchangers are reversible enabling the materials to be reused multiple times, and based on the charge characteristics of the ion reactions, the materials can be further defined as either cation or anion exchangers [27]. High-quality ion exchangers should have chemical and physical stability, large surface areas, and large ion exchange capacities, and exhibit a relatively fast ion exchange speed [28].

FMSN structural/chemical attributes make them highly attractive to be configured as ion exchangers. However, specialist ion exchange membranes remain the most popular and widely used technology platform for ion exchange-mediated metal removal. This is despite many studies having shown that membrane-based ion exchangers have numerous practical limitations. The most significant of these being poor chemical stability in alkaline environments [29]. For example, in matrices with a high pH, hydrogen peroxide (including its free radicals) is particularly reactive/damaging for ion exchange membrane modules, making them prone to premature failure [29, 30]. An exception to this is perfluorinated anion exchange membranes, which have outstanding proton conductivity and chemical stability, but suffer from high production costs and a low ion selectivity [30]. In addition, the degradation treatment of waste after the use of ion exchange membranes is still very complicated [31]. As although the elemental target is removed from the matrix, the counter ion still replaces it and this can cause new/secondary problems. In contrast, functional silicas have a more stable chemistry and are less expensive to manufacture. There is also no requirement for a chemical “swap” or exchange to occur, which improves the quality of the treated liquid phase. It is these practical considerations with ion exchange membranes being generally limited by relatively high capital and operating costs, as well as technical or economic constraints [32], that have seen many industries moving away from these systems. Functionalised silica on the other hand have a low operating cost and can be optimized to work in environments with high dissolved organic carbon and low pH [33].

This major increase in the popularity of FMSN research is exemplified by two FMSN in particular: Mobile Crystal Corporation (MCM-41) and Santa Barbara Amorphous
(SBA-15). These two categories of mesoporous silica have received the most attention. Figure 2 shows that the number of FMSN studies published started to increase rapidly from the early 2000s onwards. Today, there are over 70,000 published papers featuring SBA-15 and MCM-41.

The majority of FMSN studies over the last two decades have focused on the synthesis and preparation of new materials [34–39]. During this period, there has been a constant development of novel structures and functionalizations, with major gains in the understanding of the importance of the inner fine structure of mesoporous Si along with the mechanisms of FMSN binding [40–42]. Depending on the pore length/size, the mass transfer and rate of molecular diffusion are modulated [43, 44]. However, the pore sizes do not need to be uniform through the Si either, and this creates additional scope for tailoring the chemistries for a specific environment or need. For example, Peng et al. [45] were able to create specific mesopore channel regions within the Si with larger surface areas and pore diameters (3.0 to 7.3 nm) by adding 1,3,5-trimethyl benzene alongside cetyltrimethyl ammonium bromide (CTAB) during synthesis. This tuneable micelle core-swelling agent provided a template that controlled the shell architecture of the internal cavities, one that could be easily removed from the silica structure prior to use. It was this manipulation that created the 2D hexagonal mesostructure, with smaller pores in the shell and larger spaces in the silica core, providing the material with its unique, 2-step adsorption-desorption characteristics [45].

However, despite the increasing role and complexity of mesoporous structure as a determinant of adsorption behavior, it is still the functionalization process that dominates the materials’ final characteristics. Thus, introducing different functional groups ultimately controls the metal ions’ adsorption both externally and internally within the adsorbent. For example, carboxylate, sulfonate, and phosphate groups promote the adsorption of metal ions through ion exchange, while amine moieties typically operate by complexation. The adsorption of metals by FMSN in water is mainly based on the use of surface-grafted functional groups, such as amino or mercapto, for complex coordination, thereby achieving selective adsorption of metal ions [46]. Studies have also shown that the adsorption capacity of FMSN increases with an increase in the total concentration of functional groups. The surface area of the adsorbent is obviously an important factor determining functional group concentration [47]. However, the branching structures of the functionalizations, and how many terminal groups there are, also play a defining role.

For the adsorption of nonionic organic matter in water, generally, grafting alkanes onto the FMSN increases surface hydrophobicity, which is beneficial to the capture of organic matter. For the adsorption of ionic organics, the charge density of the solid surface is the key factor for adsorption. The degree of interaction between ionic organics and FMSN changes due to variation in solution pH, because the surface charge of the FMSN and the ionic radius of the ion target are controlled by pH. The ionic strength and ionic composition of the solution also affect the adsorption of ionic organics, especially when inorganic ions compete with organic ions for binding sites [46]. With greater appreciation of the unique adsorbing ability of functional silica, this has prompted more studies to consider a broader range of chemical targets, further expanding the utility of FMSN [48–50]. For example, there are specialist FMSN scavengers for both Hg [51] and Cu [52, 53]. Despite the vast number of wastewater treatment and environmental protection technologies available, FMSN have proven themselves to be a sector leader for metal-chemical removal [54].
Synthesis of FMSN

There are three main steps in the production of FMSN, synthesis, surface modification, and stabilization. The synthesis of mesoporous silica is commonly performed by base or acid hydrolysis, using an alkoxysilane precursor. The ordered architecture of pores and channels is obtained from amphiphilic micelles which are trapped under the silica network. This template/mold is then removed by either chemical extraction or heat-assisted treatments. The pore length and diameter can be adjusted by altering the amount and type of templating agent used and the reaction conditions [10]. Protective etching with NaOH on silica coated with PVP molecules can be used to form FMSN, especially for crystalline silica such as quartz, cristobalite, and tridymite [55]. Variation in the hollow silica channels is achieved with different etching speeds and durations, and the targeting of outer or interior regions of the particle. However, fine control of pore structure is difficult, with this method approach being more suited to disordered mesopore synthesis [56].

Post-synthesis surface modification and direct synthesis (one-step synthesis/co-condensation) methods are applied to fix organic functionalities to mesoporous silica [57]. For example, in the post-synthesis surface modification of reactive nano-silica, sodium silicate undergoes a hydrolysis reaction as pH decreases. This then generates Si(OH)₄ monomers which are then developed through a series of dehydration condensation reactions, resulting in a high-density surface functionalization with active hydroxyl groups. At the same time, methoxyl groups of the surface modifier 3-mercaptopropyl-trimethoxysilane (MPTS) then hydrolyze under alkaline conditions to produce the active surhydroxyl group. These active hydroxyl groups generated by MPTS then react with the surface functionalized hydroxyl groups just generated on the silica, forming Si-O-Si bonds and thus providing the mesoporous silica with surface adhered sulfhydryl groups [58].

Through the co-condensation of precursor inorganic framework materials with terminal trialkoxyorganosilanes (such as (R′O)₃SiR), the mesoporous silica phase can be prepared. This results in organic moieties being covalently anchored to the pore/channel walls of the silica in an orientation that enables them to project directly into the pore walls [57]. The advantage to this functionalization approach is that obstructions within the pores themselves caused by the organic units are minimized. An additional merit is that the modifications are more homogenously distributed within the pores, compared with other techniques. However, the degree of mesoscopic order of the FMSN needs to be monitored. As the ratio between the organic and silica increases in the reaction mixture, a more disordered and less reproducible material is produced [57]. These restrictions though can be conquered with the inclusion of structure-directing agents, producing a class of materials termed periodic mesoporous organosilicas (PMOs). Here, the functionalizations are integral components of the silica structure, resulting in a narrow pore radius distribution and well-organized pore network [57].

The final production stage of FMSN is stabilization. Here, functionalization can be used to give different regions within the Si particle different properties. For example, the core can be made hydrophobic and the shell hydrophilic [59]. This is a key part in improving hydrothermal stability. Hydrophobicity enhances the hydrothermal stability of materials in the aqueous environment, and hydrophilicity ensures biocompatibility. The free silanol (Si-OH) groups on the materials can provide a certain degree of hydrophilicity; with the help of condensers, these will be further used to modify some functional groups, biomolecules, or drug molecules [60, 61].

Diversity of FMSN’s Functional Groups

Many studies have reported that FMSN with different functional groups have specific selective adsorption functions. In Georgescu’s study [62], synthesized FMSN (TAC-functionalized silica) used for the removal cadmium and copper from water, adsorption results show that 0.1 g of FMSN can extract 94% of Cd(II) and 98% of Cu(II) from aqueous solutions (pH 6). Li et al. [33], through a one-step method, synthesized LMS-AP-FMSN. In a subsequent wastewater treatment experiment, LMS-AP has good metal removal efficiency, and the removal rate of Al, Pb, and Zn reached more than 80%. But the adsorption of Cr from the same waste water was only ca. 70%, while the removal of As was less still at only 30–40% [33]. The huge variety of mesoporous structures, pore network structures, combined with functionalization configurations makes FMSN a highly versatile yet extremely powerful chemical solution to toxic element containment. However, with the enormity of different FMSN configurations, selecting the right materials for the correct conditions is still challenging, and their effectiveness requires case-by-case validation. In the following sections, the application of FMSN in tackling environmental pollution will be discussed.

Current Applications and Future Directions of FMSN

The Use of FMSN in Water and Food Security

FMSN are currently widely used to absorb numerous metal ions and compounds from aqueous solutions [63–66] (Table 1). For example, Xia et al. [63] synthesized a variety of different FMSN to study comparatively, FMSN adsorption capacity for TTEs in water. In their absorption experiments, Cd(II), Mn(II), Pb(II), and Fe(III) concentrations were all ~ 0.5 mg L⁻¹ in drinking water with a pH of ~ 4; it was found among the types of FMSN tested; S16-1N completely removed these four elemental impurities (< detection limit).
The excellent performance of S16-1N can be ascribed to its high pore volume and specific surface area, along with possessing high accessibility to its functional group ligands [63]. Chatterjee et al. [66], by using a modified Stober process, synthesized cubic mesoporous silica, then it was functionalized to produce a fluorogenic silica probe material (SiO$_2$@AZOL). The adsorption experiment of the fluorogenic silica probe material was performed in a solution with a pH of ca. 7. It was found that this material showed an extraction efficiency of around 99% for HSO$_3^-$, Cd(II), Hg(II), Zn(II), and Cu(II), with adsorption capacities of 873, 633, 630, 412, and 260 mg g$^{-1}$, respectively [66]. For existing industrial applications, in Guizhou, China, the FMSN iMoLboX has been successfully applied to retrieve precious metals in addition to the removal of TTEs from a variety of process waste streams. For example, iMoLboX has achieved a 96.14% removal efficiency of As in medium/strong acid solution (mass fraction of 65–85%) containing 300 µg L$^{-1}$ As. While it possesses a removal efficiency for Cu of 92.97% in aqueous solutions containing 30 mg L$^{-1}$ CuSO$_4$ (Fig. 1). In addition, iMoLboX FMSN can effectively extract Rh from acidic aqueous solutions containing 57 mg L$^{-1}$ Rh(III), and remove Pd from organic alcohol solutions containing 562 mg L$^{-1}$ Pd(II); in both cases, removal efficiency is > 99% [69]. But the adsorption efficiency of ion exchange resins for Rh(III) and Pd(II) from 8.9 mg L$^{-1}$ Rh(III) solution and 43.9 mg L$^{-1}$ Pd(II) solution is only about 90% [70], while the removal ratio of AC for Pd(II) from 100 mg L$^{-1}$ Pd(II) solution is 93.5% [71].

Fluorine is highly reactive and forms a great variety of chemical compounds. It is used extensively in the nuclear and metal processing industries, in electronics, as a surfactant.
features prominently in polymers and in agrochemicals where it features within pesticide formulations. Fluorinated compounds though are major environmental pollutants and can be difficult to remove from waters. Zhou et al. [65] developed a novel silica functionalized nanocomposite (Fe$_3$O$_4$@SiO$_2$-NH$_2$&F$_{13}$); this material showed a good selectivity for perfluorinated compounds (PFCs). In a 1-L water sample fortified with PFC concentrations ranging from 0.5 to 50 ng L$^{-1}$, the composite possessed a much better removal efficiency (86.29%) for PFCs than that (58.61%) of powdered AC [65].

FMSN can also be used to extract drug residues from food [72]. In this example, three different mesostructured silicas (SBA-15-C18, MSU-2-C18, HMS-C18) with added octadecylsilane groups were trialed. Despite the functionalization approach being identical, the different silica structures evolved distinctive binding properties. SBA-15-C18 was the least effective scavenger due to hydrophobic and polar secondary/hydrogen bonding interactions caused from the functionalization leaving a high frequency of non-modified silanol groups in the silica. However, HMS-C18 proved a promising multi-drug sorbent. Out of the panel of 26 veterinary drug residuals tested from bovine meat samples, the extraction efficiency for more than half of these compounds was over 80% [72]. Although the direct use of FMSN in food safety is new, the successes to date provide further evidence of the materials’ wide-ranging applications, across diverse chemical conditions. However, perhaps, it is the use of FMSN to combat soil pollution that might be the materials’ greatest contribution to our future food safety [73].

The Use of FMSN in Soil Research

FMSN has good compatibility for use within soil because it is inert, highly stable, and soils naturally already contain high proportions of silica. The application of FMSN in soil research is a theme that has only emerged relatively recently. The two principle directions being soil remediation (direct amendment) [73] and in situ measurement [74].

Although the use of functionalized silica in soils is relatively new, there have already been a number of successful applications. For example, diacetylmonoxime-functionalized silica gel has been used to remove Cu from fly ash-ameliorated soil samples [75]. In this study through FTIR, element analysis, BET surface area analysis, and $^{13}$C CPMAS NMR spectroscopy, it was shown that the FMSN have high selectively for Cu removal with a preconcentration factor of 250; sorption capacity reached 0.93 mmol g$^{-1}$. Kinetic parameters and isotherms correlated well with the pseudo-second-order and Langmuir models [75]. Another important study reported that FMSN can limit the absorption of toxic metals and As by plants through reducing bioavailability. In this example, a soil with an As concentration of 9.87 ± 1.22 mg kg$^{-1}$ had the fraction of unavailable As increased from 40 to 77% after FMSN was applied [76]. Similarly, Lian et al. [58] used FMSN in contaminated soil to selectively immobilize Pb and Cd. Results showed that the FMSN applications can decrease the pools of bioavailable Cd and Pb from 12 mg kg$^{-1}$ and 1194 mg kg$^{-1}$ to 0.2 mg kg$^{-1}$ and 10 mg kg$^{-1}$, respectively, with the immobilization efficiency of Pb and Cd reaching 99.12% and 98.23%. More importantly, this FMSN exhibited a low effect on other soil parameters. While the Cd and Pb species immobilized by FMSN exhibited greatly increased acid resistance [58], an attribute making FMSN suited for long-term remediation roles.

Recently, a series of studies has shown that FMSN can be used for remediating Cd-contaminated agriculture soil [73, 77]. The stabilization efficiency of Cd by FMSN reached up to 91.2% with just a 1% application. At the same time, the migration rate of Cd in the soil decreased from 45.8 to 19.0%, and the effect of the addition of FMSN on the soil properties was negligible [77]. In addition, FMSN reduced the leaching rate of Cd in the soil (36.0%) and bioavailability (54.3%); the concentration of Cd in crops decreased more than 50% [73].

Directly, addressing pollutant bioavailability with amendments is one solution to improve soil health. However, any implementation of such technologies needs sufficient knowledge of in situ conditions and subsequent pollutant mobility. Over the past couple of decades, the passive sampling technology of DGT has emerged as a leading method for the measurement of bioavailability for inorganic and organic contaminants in soil. However, the connection with FMSN is that they are increasingly being used as DGT substrates, opening up a range of exciting avenues in soil testing, new rapid analysis approaches, and for high-resolution 2D ion-mapping of pore water chemistries.

The Use of FMSN in DGT Research

DGT Introduction

DGT was originally developed for aquatic systems, then evolved into an established technique to understand chemical lability and speciation in soil [78]. Comprising of a base unit, a cap, and thin film stack system (consisting of a membrane filter, diffusive gel, and binding layer), the DGT configuration is robust, easy to work with in the environment, and crucially, units do not require individual calibration [78]. The binding layer provides the device’s selectively, while the diffusive layer controls solute transport and defines the intensity of the metal flux induced by the DGT device. Together, they both determine the measurement characteristics of a specific/individual DGT configuration.

In practice, DGT devices are easy to deploy in water/sediment/soil; for a detailed review of the technique, refer to Davison [78]. After the DGT binding layer is recovered and eluted, quantitative analysis of the eluents can be
achieved through a range of commonly available analytical techniques, including inductively coupled plasma mass spectrometry (ICP-MS), inductively coupled plasma optical emission spectrometry (ICP-OES), atomic absorption spectroscopy (AAS), and X-ray fluorescence (XRF). Environmental samples are prone to a wide range of measurement interferences due to matrix complexity. A further benefit of the DGT approach is that there is a preconcentration of the target analyte. In the following sections, the use of DGT and FMSN for the measurement of TTEs in soil will be discussed from the perspective of Hg and As pollution.

**Hg In Situ Sampling by DGT**

Mercury (Hg) pollution remains a key pollution focus, not only is it a potent human poison, but it can cause widespread environmental damage [16, 17]. Mercury exposure can arise naturally from rock weathering, volcanic activity, and geological deposition [79]. However, the main sources of Hg release are from human activities, particularly waste incinerators, residential coal for heating and cooking, coal-fired power plants, and mining for gold and other metals [16, 17, 79, 80]. The fate and behavior of Hg are difficult to predict, which complicates the assessment of toxicological risk, due to the wide range of species and complexes found in the environment. Elemental mercury (Hg0), referred to as quick silver, is the most volatile species, forming vapor easily at typical ambient temperature/pressure [81]. Hg(OH)2, Hg(OH)Cl, and HgCl2 are common inorganic mercury (iHg) compounds in freshwaters. However, because the Cl concentration in seawaters is higher, Hg speciation trends are also typified by HgCl+, HgCl2, HgCl3−, and HgCl42−. Methylmercury (MeHg) is an organometallic Hg species, characterized by extreme toxicity, making it the critical contaminant/pollutant in the environment [81].

Soil is an important reservoir of anthropogenic Hg release, playing a key role in the Hg cycle [82]. The soil environment is a heterogenous micro-landscape of solid-solution-gas interfaces, and fluxes between these compartments are dominated by the soil phase characteristic/composition, where partitioning of the Hg is concentrated. Mechanistically, the kinetics of Hg mobility can be broadly segregated into two groupings. The first is an immediate/quick reaction scheme forming outer-sphere complexes involving cation exchange, with a multitude of different ligand types, both organic and inorganic. The other group involves the formation of stable colloid complexes that diffuse within the interior of the soil particles. The bonds formed are inner sphere and recalcitrant, characterized by slow release. However, these stores should not be considered as insignificant as they can provide a long-term and sustained source of bioavailable Hg [83, 84].

DGT has been shown to be effective at predicting Hg uptake in plants [85]. It acts to introduce a controlled perturbation into the soil system—replicating the processes evoked by the plant root. The method is also advantageous for in situ measurement of Hg speciation, because it preserves the stability and distribution of the Hg species, while also preconcentrating the target analyte during sampling [86]. DGT and HPLC-ICP-MS are appropriate tools for the evaluation of rhizosphere Hg bioavailability in estimating the risk from contaminated soils [84]. Liu et al. [85] predicted MeHg uptake by rice plants (*Oryza sativa* L.) by using the DGT technique to detect bioavailable MeHg in pore water; the DGT probe used a 3-mercaptopropyl-FMSN. A significant positive correlation between MeHg flux in soil measured by DGT and MeHg flux in rice roots (R = 0.853, p < 0.01) was observed. The correlation between these two flux parameters indicates that DGT can provide a quantitative description of the rate of update of this bioavailable MeHg, and that the DGT can predict the bioavailability of MeHg in rice paddies soil [85]. Ridošková et al. [87] used Ambersep GT74 and Duolite GT73 gels as DGT binding layer for measuring the bioavailability of Hg in contaminated soils. The DGT experiments indicated that Duolite GT73 shows similar Hg accumulation properties as Ambersep GT74; the maximum Hg accumulation for these two resin gels was obtained after 12 days of deployment in the soil, about 27 and 33 ng/disk, respectively. In four different soil deployment experiments, the range of DGT available Hg compared with total Hg ranged between 2 and 10% [87]. Huu Nguyen et al. [88] measured the bioavailability of Hg in soil for the earthworm *Eisenia Fetida* by using DGT which used FMSN (3-mercaptopropyl-functionalized silica) as the binding phase. Soils used in Hg exposure experiments were prepared with different pH (4.6, 5.6, and 6.2) and varying peat moss concentrations of 5, 10, 15, and 20%. DGT was deployed in the soil with approximately 35–40 earthworms and removed across a series of timepoints spanning half-a-day through to 10 days. The calculated uptake efficiency for Hg for this binding gel was 91 ± 3.4%; the DGT was able to predict the Hg concentration of earthworm tissue effectively, unlike comparative measurements of pore water/acid-extractable Hg [88]. It can be seen that DGT is a dependable technique that can be used to measure the bioavailability of Hg in soil. Compared with other DGT, DGT using FMSN as the binding phase has better absorption efficiency for Hg. For more research papers about Hg in situ sampling by DGT, see Table 2.

**In Situ Sampling of As with DGT**

Arsenic is the 20th most abundant element in the earth’s crust and can be present in the environment in four oxidation states As(-III), As(0), As(III), and As(V) and as both organic and
inorganic species [92]. The general order of toxicity of the different arsenic species is as follows: arsenite > arsenate > monomethyl arsenate (MMA) > dimethyl arsenate (DMA) [92, 93]. While, the total content of As in the environment is largely decided by the degree of environmental pollution and associated geology. Localized redox conditions play a main role, along with the microbiome in controlling the concentrations of As(III) and As(V) [92], while methylation/demethylation is subject to biotic controls Su et al. [94]. This creates a diverse range of As chemistries in situ that can be difficult to capture.

In 2012, Bennett et al. performed selective measurement of total inorganic arsenic and As(III) using separate DGT sampler units configured with different binders. Here, it was determined that As(III) was the main substance transferred from the sediment (solid phase) to pore water. They also showed that the DGT technology can be used under a range of deployment conditions to investigate arsenic morphology and its migration potential [95]. Wang et al. [96] used Zn-O DGT to systematically study the concentration variation of unstable As in the sediments of Hongze Lake; the recovery of Zr oxide DGT was 88%. It was found that the concentration of labile As in the sediment profiles varied considerably (ranging from 0.15 to 4.15 μg L\(^{-1}\)) [96]. Gorny et al. [97] used a Zn-ferrite (ZnFe\(_2\)O\(_4\)) substrate as the DGT-binding phase to measure total arsenic in river water and sediment pore water. Results shown ZnFe\(_2\)O\(_4\) have a better sorption capacity for As(III) compared with ferrihydrite and Metsorb binding gels, with the sorption capacity of ZnFe\(_2\)O\(_4\) reaching 54,000 ng for As(III) [97]. In 2011, Bennett et al. first used FMSN as a DGT-binding substrate to selectively measure As(III), and its absorption efficiency for As(III) was > 99%, validating the feasibility of this method. The reason why mercaptopropyl-silica have highly selective adsorption ability for As(III) is because of the strong complexation of H\(_3\)AsO\(_3\) by thiol (S-H) groups [98]. Gorny et al. [99] also reported the use of FMSN as the DGT-binding layer and found that the use of matrix corrected elution factors to rectify the interaction with sulfides can ameliorate the measurement of As(III) in 3-mercaptop-silica (3MP)-binding gels; the concentration of As(III) recovered from the 3MP-DGT samplers reached 103 ± 7.0 and 93.8 ± 8.6% (mean ± standard deviation) of the initial As(III) concentration in seawater and freshwater, respectively [99] (Table 3).

### DGT-Two-Dimensional Chemical Imaging

The acquisition of environmental/soil measurements in the majority of studies to date prioritizes the sampling of

**Table 2** A summary of FMSN and non-FMSN DGT configurations used for in situ sampling of Hg

| Binding phase                        | Species                  | Matrix  | Dominant system/scheme | Key finding                                                                 | Ref.                  |
|-------------------------------------|--------------------------|---------|------------------------|-----------------------------------------------------------------------------|-----------------------|
| Chelex-100 and Spheron-Thiol        | MeHg, CH3CH2Hg+, Hg(II) | Soil    | HPLC-ICP-MS, DGT       | DGT techniques is a suitable tool for the estimation of Hg root availability  | Cattani et al. [84]   |
| Duolite GT73 and Ambersep GT74 gel  | Hg(II), MeHg             | Water   | DGT                    | The agarose gel was found as a diffusion gel for mercury measurement in DGT  | Docekalová and Diviš [89] |
| 3-mercaptopropyl--functionalized silica (Sigma-Aldrich) | Hg(II), MeHg | Soil | DGT                    | A new resin gel Ambersep GT74 was used in DGT technology, which showed similar Hg accumulation characteristics to Duolite GT73. | Ridošková et al. [87] |
| 3-mercaptopropyl--functionalized silica (Sigma-Aldrich) | MeHg                     | Soil    | DGT                    | Research shows that DGT can predict bioavailability of MeHg in paddy soil. | Liu et al. [85]       |
| 3-mercaptopropyl--functionalized silica (Sigma-Aldrich) | Total Hg                  | Soil    | DGT                    | DGT-measured Hg flux is a better tool than conventional methods for predicting Hg bioavailability for earthworms inhabiting diverse types of soil. | Huu Nguyen et al. [88] |
| 3-mercaptopropyl--functionalized silica (Sigma-Aldrich) | Total Hg                  | Soil    | DGT                    | The effectiveness of DGT technology in predicting plant uptake of mercury was confirmed. | Turull et al. [90]   |
| 3-mercaptoppropyl--functionalized silica (Sigma-Aldrich) | Total Hg, MeHg           | Water   | DGT                    | Hg speciation in aquatic systems was investigated, found that Labile MeHg in the sediment of the Gulf represents up to 75% of total labile Hg. | Bratkič et al. [91] |
individual biogeochemical compartments. Far less consideration is given to the speciation, transformation, and exchange of elements across interfaces between different niches/zones, and the processes controlling these events. 2D high-resolution diffusive gradients in thin films (HR-DGT) are a key component of one of the few existing tools that can capture multi-elemental solute dynamics/interactions in appropriate submillimeter scales, to suitably understand/decipher interface fluxes. The usual operation of this chemical imaging approach uses a multi-layered system of DGT and planar optodes to obtain the HR-2D images. This is not intended to be a comprehensive review of this emerging technology; for this, we recommend Santner et al. [101] and Santner and Williams [102]. However, FMSN are increasingly being used in these ion-mapping applications, because of their high capacities, environmental tolerances, and species specificity. For example, Shi et al. [103], through mapping the 2D chemistry of a rice rhizosphere, were able to capture the distribution of Se(IV) and oxygen simultaneously. Here, the DGT substrate used was a bi-functionalized SBA15 FMSN, which provided a selective measurement for Se(IV) (Fig. 3) [103]. What was apparent from these chemical images was that Se(IV) mobility is restricted to the rooting zone and with localized redox conditions controlling release patterns. This is despite the Se amendments having been thoroughly mixed through the soils prior to the addition of the rice plants.

This recent interest in the root-soil interactions within rice rhizospheres is understandable. Rice is so widely cropped and plays a fundamental role in global food security, yet is significantly threatened by trace element imbalances, either deficiencies or over-exposure. The below-ground processes around the roots are very challenging to measure because of the complexity and instability of the environment due to strong zonation in both chemical and biological effects. This means that there is still much to learn about the biogeochemistry of paddy soils and in particular the role of these fine-scale, localized features on plant health and quality. This is demonstrated by the recent
DGT-optode imaging of greatly enhanced fluxes of As, Pb, and Fe(II) mobilization immediately adjacent to lateral root tips (Fig. 4) that forms simultaneously within a zone of P depletion and co-currently with a root-induced oxidation front [104, 105]. 2D HR-DGT applications lend themselves very well to this study, being a low-disturbance sampling system that was initially developed for flooded wetland sediments [105]. Following on from this work, Fang et al. [74] expanded the range of species that can be measured in a rice rhizosphere by developing a mercapto-functionalized SBA-15 FMSN DGT for Sb(III) solute imaging. Compared with commercially available 3-mercaptopropyl-functionalized silica, the newly synthesized FMSN was distributed more evenly within the sampler, making it a better candidate material for LA-ICP-MS analysis. Furthermore, the DGT multi-system was also deployed with an AgI gel layer to study the influences of sulfur application on Sb(III) solubility (Fig. 5) [74]. It was found that sulfur additions were contributing to greater release of Sb(III) compared with the non-amended control, and there was a strong spatial trend between Sb(III) and S(-II). There were also hotspots of mobilization that formed around root tip zones, similar to the patterns observed for As and Pb [104, 105].

Traditionally, to extend the range of element/species measurements captured by the DGT/optode systems, binders have been mixed together, or the combination of binder layers has been increased. What is exciting about FMSN approaches is that these properties can be potentially delivered by one binder phase, given the correct functionalization approach, providing a molecular-based solution to expanding and improving chemical imaging delivery.

**Limitations and Considerations for FMSN Use in Soils**

As documented in this review, FMSN are a diverse and rapidly developing class of materials with a wide array of different physio-chemical properties. However, there is still a lot that remains unknown about the immediate and persistent effects these incredibly versatile and varied materials can have...
on soil/plant health. Or what the interplays are when soil type, management practices and climate all interact. Answering some of these questions just involves more testing. However, as a first principle, the toxicity of inorganics in severely polluted soils would have a dominant influence on soil processes and function. Therefore, reducing the mobility of harmful metal(loids) would render a restorative effect. However, this would of course be dependent on specific FMSN, dosage, and the soil environment.

The greatest uncertainty, though, in terms of long-term safety and behavior lies with the smallest of the FMSN technologies. Those Si materials, at the lowest end of the nanoscale spectrum, even when not chemically functionalized, can give rise to unexpected interactions with cellular biological systems/metabolism. Sometimes, these are beneficial. For example, in addition to the widely known antagonisms associated with Si and As(III) uptake in plants. It has recently been observed that SiO₂ nanoparticles (NPs) provide additional protection to plant cells subject to As stress, mediating a strengthening of plant cell walls by increasing their thickness and altering their composition [106]. Likewise, in maize, field applications of SiO₂-NPs resulted in a large improvement to growth, but interestingly, there were disruptive effects as well with the Si found to induce a pesticide response for the crop during storage [107]. Similarly, some soil invertebrates demonstrate avoidance behaviors to SiO₂NP Santos et al. [108], and it has been shown that they can increase soil bacteria community diversity [109].

SiO₂NP place within the top five commercial nanoproducts globally [108, 110] with applications ranging from paint...
stabilizers through to food agents [111]. Therefore, extensive material flows of SiO$_2$NP into the environment are already ongoing [111]. However, when trying to isolate and understand the already complex mechanisms of FMSN interactions within the plant-soil continuum, it would be prudent to work initially with larger particle size FMSN. Not only would these have more predictable mobility in soils but they would simplify the interpretation of the soil chemical trends, making it easier to match a specific functionalization to its response in situ.

**Outlook/Conclusion**

Global soil health continues to be threatened by trace element pollution. This damage is inextricably linked to our continued intensive agri and industrial activities, demand for natural resources, and poor waste management and material recycling/reuse. This land degradation can be broadly viewed as either a general increase in metal loading into soil systems, or modification of existing elemental reservoirs, or a mixture of both processes which results in enhanced bioavailability and toxicity. New technologies are required to counter these pollution challenges, and the behavior and properties of FMSN are highly suitable for this mission. Firstly, Si is already a major constituent of soils, and is highly stable in the environment. It is those traits that provide a cost-effective and resistant platform from which to optimize the functionalization which in combination with the architecture of the pores and channel networks within the Si controls the binding chemistries and their specificities to desired targets. These dual physical and chemical “levers” enable the production of materials with a...
The other aspect of FMSN applications is in their role in environmental measurement. In this review, we have focused on the use of FMSN in the widely used/popular passive sampler DGT, examining this from the perspective of two soil pollutant priorities Hg and As, alongside the emerging new DGT applications of HR-2D ion-mapping. In conclusion, the benefits FMSN in an environmental setting are gaining increasing attention, as more application successes continue to be shown. The versatility of the FMSN platforms is perhaps best illustrated by their large range of uses, fulfilling a research need across the diverse themes of pollutant characterization, analysis, monitoring, prevention, treatment, and remediation. However, there is still much to be discovered about how FMSN perform in different soil environments, especially in wetland agri-ecosystems like rice paddies where inorganic pollutants such as As are of particular concern [105, 112]. It is in these systems where in order to improve air quality, worldwide bans on rice straw burning have resulted in the repurposing of the straw which was once cycled back into the soil. This export removes millions of tonnes of Si. This is of concern because of links Si depletion has on harvest failures, rampant pest/disease outbreaks, and enhanced plant uptake of As [113]. How FMSN could play a role in efforts to resupply Si to these damaged soils is still unresolved, but certainly for the worst metal-impacted sites, its use carries appeal.

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