Metal–organic frameworks (MOFs) as highly efficient agents for boron removal and boron isotope separation†

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The boron adsorption and isotopic separation on water-stable metal–organic frameworks (MOFs) were evaluated. These materials, to date, exhibit fairly high to highest adsorption capacities for boron. Furthermore, unprecedentedly high isotopic separation factors (S) were observed in some materials.

Global climate change and energy framework uncertainties have propelled a transition in the energy structure. Although there is a continuous and growing effort to adopt more sustainable energy systems with the ambition to reduce greenhouse gas emissions, the development of renewables still needs intensive investigation. However, the position of nuclear power keeps increasing while expanding its share in the global energy mix in spite of all its pros and cons.

Boron has two stable isotopic compositions, $^{10}\text{B}$ and $^{11}\text{B}$, with an abundance of 19.1–20.3% and 79.7–80.9%, respectively. The $^{10}\text{B}$ isotope, with a larger attenuation cross-section for thermal neutrons, is irreproachably used in nuclear power plants for neutron shielding and in radioactive waste disposal equipment, such as the components used for compact fuel storage racks and transportation baskets. It also plays a critical role in treating cancers that cannot be controlled effectively by conventional means, which is known as boron neutron capture therapy (BNCT). Although methods for boron isotope separation are in high demand for the nuclear power industry and medical treatment, the separation process is practically very difficult because of the tiny differences between its isotopes. Some methods have been observed including exchange distillation, adsorption-based separation, thermal ionization mass spectrometry and the laser assisted retardation of condensation (SILARC) method, among which only isotopic exchange distillation has been successfully applied in practical $^{10}\text{B}$ production. Except for a small separation factor of around 1.03, disadvantages including the instability of the anisole–boron trifluoride complex, severe causticity to the equipment and insecurity as a result of the toxicity of boron trifluoride greatly impede the process. Furthermore, technologies such as mass spectrometry and the laser are too expensive to be adopted on an industrial scale.

A promising alternative for boron isotope separation is an adsorption-based process using a boron adsorbent, which is more efficient, easier to operate and inexpensive. In addition, the increasing demand for freshwater also promotes the development of novel boron adsorbents because the ingestion of excess boron is detrimental to plants and human beings. For the health of human beings, the boron concentration limit in drinking water was set at 2.4 ppm by the WHO in 2011. However, boron concentration is usually around 5 ppm in seawater and can be as high as 119 ppm in some groundwater of active volcanic and geothermal activities. In addition, wide applications of boron compounds in industry lead to an enrichment of boron in water. It is also important to develop adsorbents with high boron adsorption capacities for boron removal from water.

Highly efficient adsorbents are desirable for both boron isotope separation (Section S1, ESI†) and boron removal (Section S2, ESI†). Various boron adsorbents, such as anion resins, cation resins, boron-specific resins and other boron adsorbents have been explored for their isotopic separation capacity in static adsorption experiments and fixed in the chromatography column as the solid phase to study dynamic separation properties.

Among the various boron adsorbents, only N-methyl-$\alpha$-glucamine type resins including Amberlite IRA-743 have been widely applied in boron removal, which also exhibit the highest boron isotopic separation factor ($S = 1.027$) to date. This small separation factor makes it impossible to employ chromatography to separate boron isotopes.

Metal–organic frameworks (MOFs), constructed from different metal ions and organic ligands with coordination bonds, have attracted great interest as a result of their good porosity, high specific surface and chemical tenability. A large

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number of MOFs have already been widely studied in areas such as gas storage, catalysis, optics, drug delivery, separation, and chemical sensors.29 MOFs exhibit strong potential in contaminant adsorption. Unfortunately, most metal–organic frameworks are unstable in aqueous solutions due to the water-unstable coordination bond, which limits their practical applications. In recent years, research on water-stable MOFs has made great advances, and MOFs with high water stability have been reported as effective agents for the removal of contaminants,30–34 such as ZIF-8, UiOs, and MILs. Nevertheless, investigations on MOFs as agents for boron isotope separation and adsorption have yet been reported. Herein, our goal was to identify the potential of water-stable MOFs as agents for both boron isotope separation and boron adsorption. These MOFs with good hydrothermal and chemical stability35,36 were prepared based on the reported methods.37–43 The structures and pore properties were confirmed by X-ray diffraction, scanning electron microscopy (SEM), nitrogen adsorption/desorption and thermogravimetric analysis (Section S3–S9, ESI†). Static adsorption experiments of boron acid were carried out to evaluate the boron isotope separation capacity. Seven activated MOFs were added into a 0.5282 mol L−1 aqueous solution (20 mL) at a dosage of 5 g L−1 and the mixtures were stirred in a water bath at 25 °C for 24 h. Then, the MOFs were filtered and the residual solutions were analysed for the boron concentration using ICP-OES and the boron isotopic abundance (10B/11B) using ICP-MS. A standard boronic acid aqueous solution with a known boron concentration was employed to calibrate the instrument. Each sample was analysed three times and an average value was determined.

The 10B/11B abundance of the initial boronic acid aqueous solution (α0) was 0.24779 and the 10B/11B abundance and boron concentration of the residual boronic acid solution after the adsorption experiments are summarized in Fig. S8 (ESI†). The boron isotopic separation factors and boron adsorption capacities of all seven MOFs were calculated (Section S10, ESI†) and shown in Fig. 1 and 2. When compared with other boron adsorbents (Table S2, ESI†), all the seven MOFs perform well for boron adsorption. In Fig. 1, the adsorption capacities were compared with a commercial Amberlite IRA-743 resin and waste sepiolite, which has the highest capacity among the previously reported boron adsorbents. All of them exhibit much higher adsorption capacity than Amberlite IRA-743, particularly ZIF-8, which exhibits a boron adsorption capacity of 191 mg g−1, which is, to the best of our knowledge, the highest capacity ever reported. The other six MOFs have similar boron adsorption capacities as UIO-66 > MIL-96(Al) > MIL-100(Fe) > MIL-100(Cr) > MIL-53(Cr) > MIL-101(Cr), which are also much higher than the capacity of Amberlite IRA-743 resin. The Dubinin–Radushkevich (D–R) isotherm model,44 which is applicable for both homogeneous and heterogeneous surfaces, was used to describe boron sorption on ZIF-8. Linear fitting based on the D–R model estimated the maximum amount of boron that can be adsorbed by ZIF-8 to be 234.6 mg g−1. The adsorbed ZIF-8 and adsorbed MIL-101(Cr) were characterized with XRD to observe their stability. It can be observed from the spectra (Fig. 3) that the structure of ZIF-8 was completely changed after boron adsorption, while MIL-101(Cr) retained its structure under the adsorption conditions.

Based on the boron isotopic abundance of the residual solutions shown in Fig. S8 (ESI†), it can be observed that the boron isotopic abundance (10B/11B) of all the residual solutions was higher than that of the initial boron solution (0.24779), which suggests that 10B was enriched in the solution phase and 11B was enriched in the adsorbents during the adsorption process. It was also proven that the MOFs have good selectivity between the boron isotopes (10B and 11B).

The boron isotopic separation factors were compared with commercial Amberlite IRA-743 and are shown in Fig. 2. All

![Fig. 1](image1.png) The boron adsorption capacities of seven MOFs compared with Amberlite IRA-743 and waste sepiolite.

![Fig. 2](image2.png) The boron isotopic separation factors of seven MOFs and Amberlite IRA-743.

![Fig. 3](image3.png) A comparison of the XRD spectra between the prepared materials and adsorbed materials: (A) ZIF-8, (B) MIL-101(Cr).
seven metal–organic frameworks exhibit higher separation factors than the conventional boron-specific adsorbent Amberlite IRA-743 (1.022)\textsuperscript{a} and isotopic exchange distillation factors (1.039)\textsuperscript{b}. Among them, MIL-101(Cr) and MIL-100(Fe) exhibit boron isotope separation factors of 2.00 and 1.75, respectively. It is worth mentioning that MIL-101(Cr) has by far the highest boron isotope separation factor, which is an astonishing 35 fold increase of the present best result (1.027)\textsuperscript{b} held by N-methyl-D-glucamine type resin. The adsorption separation can be operated under mild conditions with aqueous solutions of boric acid. Compared with a traditional chemical exchange distillation process, it will strongly improve the separation efficiency, reduce the cost of equipment and strengthen the safety.

Furthermore, the effects of temperature and pH on boron adsorption and isotopic separation were investigated. Four temperatures (25 °C, 35 °C, 45 °C and 55 °C) and five different pHs were investigated in this study. The results obtained for the adsorption capacity and isotopic separation factors are summarized in Fig. 4. It was found that the boron adsorption capacity increased upon increasing the temperature (Fig. 4A), which demonstrated that boron adsorption on MIL-101(Cr) was an endothermic process. On the contrary, the boron isotopic separation factor decreased significantly upon increasing the temperature. This can be attributed to the stronger interaction between boric acid and MIL-101(Cr) at higher temperature, which decreases the selectivity for the two isotopes. In Fig. 4B, it can be observed that the boron adsorption capacity displayed an increasing trend with increasing the pH from 2.05 to 6.91, up to 146.1 mg g\textsuperscript{-1}, then decreased with a further increase in pH. Similar to the study of temperature, the isotopic separation exhibited an opposite trend.

To facilitate understanding of mechanism for the boron adsorption process, \textsuperscript{11}B MAS NMR spectroscopy (Fig. 5) was performed to evaluate the chemical environment of boron in the adsorbed MIL-101(Cr). The structure of B(OH)\textsubscript{3} dominates the interaction between boric acid and the metal sites. The XPS patterns of the as-synthesized and adsorbed MIL-101(Cr) were also characterized using X-ray photoelectron spectroscopy (XPS) measurements to observe the interaction between boric acid and the adsorbent. The XPS patterns (Fig. 6) show the change in binding energy at the Cr 2p peaks after boron adsorption on MIL-101(Cr), which indicates the interactions between boric acid and the metal sites.

To further explore the separation mechanism, an adsorption simulation with molecular dynamics was employed. We simulated the boric acid adsorption density distribution\textsuperscript{47} (Fig. S11†) in both pentagonal and hexagonal windows of MIL-101(Cr). It can be proven that boric acid was adsorbed in the tetrahedron cages irrespective of whether the window was pentagonal or hexagonal. This result can be attributed to boric acid being closer to the metal sites in the tetrahedron cages and further confirms the interaction between boric acid and the metal sites. A similar observation was made for ZIF-8 in which boron adsorption was a chemical process, which mainly occurred between the boron and zinc sites (Section 12, ESI†).

The regeneration and recyclability of materials are crucial for practical applications. Both hydrochloric acid (5%) and hydrofluoric acid (0.4%) were employed as regeneration agents. However, it turned out that only part of the boric acid can be eluted from the adsorbed MIL-101(Cr). The adsorbed material cannot be regenerated completely.

![Fig. 4](image-url) The effect of temperature (A) and pH (B) on boron adsorption and isotope separation on MIL-101(Cr).

![Fig. 5](image-url) The \textsuperscript{11}B NMR spectra of aqueous solutions of boric acid and the adsorbed MIL-101(Cr).

![Fig. 6](image-url) The XPS patterns of the as-synthesized and adsorbed MIL-101(Cr).
In conclusion, we examined the boron isotope separation and boron adsorption capacity of seven water-stable MOFs. All of them exhibit higher separation factors than the previously reported techniques, particularly MIL-100(Fe) and MIL-101(Cr). The separation factor of MIL-101(Cr) is 2.00, which is significant in the boron isotope separation process. In addition, all of the MOFs demonstrated excellent performance compared with other boron adsorbents, particularly ZIF-8. The high isotope separation and boron removal capacities make it possible that both problems, boron removal and isotopic separation, can be solved in a dual step process. Further research on both boron adsorption and the isotope separation capacity of other water-stable MOFs and a deeper understanding of the separation mechanism are highly desirable.

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