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Diversity-driven, efficient exploration of a MOF design space to optimize MOF properties†

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Metal–organic frameworks (MOFs) promise to engender technology-enabling properties for numerous applications. However, one significant challenge in MOF development is their overwhelmingly large design space, which is intractable to fully explore even computationally. To find diverse optimal MOF designs without exploring the full design space, we develop Vendi Bayesian optimization (VBO), a new algorithm that combines traditional Bayesian optimization with the Vendi score, a recently introduced interpretable diversity measure. Both Bayesian optimization and the Vendi score require a kernel similarity function, we therefore also introduce a novel similarity function in the space of MOFs that accounts for both chemical and structural features. This new similarity metric enables VBO to find optimal MOFs with properties that may depend on both chemistry and structure. We statistically assessed VBO by its ability to optimize three NH₃-adsorption dependent performance metrics that depend, to different degrees, on MOF chemistry and structure. With ten simulated campaigns done for each metric, VBO consistently outperformed random search to find high-performing designs within a 1000-MOF subset for (i) NH₃ storage, (ii) NH₃ removal from membrane plasma reactors, and (iii) NH₃ capture from air. Then, with one campaign dedicated to finding optimal MOFs for NH₃ storage in a “hybrid” ~10 000-MOF database, we identify twelve extant and eight hypothesized MOF designs with potentially record-breaking working capacity ΔN_{NH_3} between 300 K and 400 K at 1 bar. Specifically, the best MOF designs are predicted to (i) achieve ΔN_{NH_3} values between 23.6 and 29.3 mmol g⁻¹, potentially surpassing those that MOFs previously experimentally tested for NH₃ adsorption would have at the proposed operation conditions, (ii) be thermally stable at the operation conditions and (iii) require only ca. 10% of the energy content in NH₃ to release the stored molecule from the MOF. Finally, the analysis of the generated simulation data during the search indicates that a pore size of around 10 Å, a heat of adsorption around 33 kJ mol⁻¹, and the presence of Ca could be part of MOF design rules that could help optimize NH₃ working capacity at the proposed operation conditions.

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1. Introduction

Metal–organic frameworks (MOFs) are a class of porous materials that could be bestowed with properties that could enable technological breakthroughs in energy, environment, and other fields.^{1–4} The idea is that judicious selection of MOF constituent nodes and linkers could yield whichever architecture and/or

chemistry is required to engender the necessary material property or behavior to enable the breakthrough.⁵ However, one persisting challenge in MOF development has been that the combinatorics of constituent building blocks creates an overwhelmingly large material “design space.”^{6–8} To expedite the navigation of the MOF design space, for longer than a decade, MOF development has been aided by high throughput computation instead of solely relying on experiments.^{9–11}

High throughput computation in MOFs has usually relied on exhaustively predicting key performance-relevant properties in all MOFs in a database—usually using molecular simulation.^{12,13} Some notable databases have been created out of experimentally reported MOF structures (*i.e.* extant MOFs)¹⁴ curated from the Cambridge Structure Database, or hypothesized MOF structures outputted by crystal creation codes (*i.e.* MOF prototypes).^{6,15–17} Notable databases of extant MOFs have been created by Chung *et al.*¹⁴ (~20k MOFs) and by Moghadam *et al.*¹⁸ (~70k MOFs). On the other hand, notable databases of hypothesized MOFs include

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those created by Wilmer *et al.*¹⁵ (~137k MOFs), Colón *et al.*¹⁹ (~13k MOFs), Boyd *et al.*²⁰ (~280k MOFs), among others. Note, however, that the size of these databases is very small compared to the vastness of the MOF design space, which some estimate to span at least one trillion MOFs.²¹

Indeed, current computational capabilities only allow evaluation of a small number of MOFs relative to the MOF design space size. For instance, the work by Simon *et al.*¹³ only managed to predict methane adsorption in ~650k materials, even though methane adsorption is one of the fastest properties to predict by simulation.²² Calculation of other properties have proven even more limiting. For instance, prediction of charge distribution through density functional theory (DFT) by Nazarian *et al.*²³ was limited to ~3k structures. Prediction of band gaps *via* DFT by Rosen *et al.*²⁴ was limited to ~20k structures. Prediction of thermal conductivity by Islamov *et al.*⁹ *via* molecular dynamics was limited to ~10k structures. Predictions of hexane isomer mixture adsorption by Chung *et al.*²⁵ was limited to ~500 structures. Moreover, in the case of adsorption applications, computational limits may be even more restrictive since screening for such properties for a given application may require considering different conditions in temperature, pressure, and composition (in the case of mixtures).

One can argue that the discovery of technology-enabling MOFs have been hampered by the inability to explore the MOF design space at large. One way that researchers have attempted to expand the number of MOFs considered in a given study is through hierarchical screening. But the latter first requires the calculation of an inexpensive descriptor which (hopefully) points to (smaller) regions of the MOF design space where the property of interest may have desirable values.^{25–27} Therefore, hierarchical screening presents caveats such as: (i) requiring extensive “domain knowledge” to identify an effective, inexpensive “descriptor”²⁸ (ii) still being unlikely that the descriptor can be calculated on the MOF design space at large, (iii) due to a probably imperfect correlation, still being possible that the descriptor calculation may overlook regions of design space where the property of interest could have desirable values.

Hence, there is growing interest in methods that allow exploring the MOF design space efficiently, while still relying solely on direct property calculations. For instance, genetic algorithms (GAs) have been explored to evolve an initial small subset of MOFs into new subsets of MOFs with optimized values of the property of interest (*e.g.*, pre-combustion CO₂ capture properties,²⁹ or CH₄ storage properties²¹). However, it is understood that GAs tend to require a larger number of evaluations and are slower than other sophisticated search/optimization methods. GAs thus may become rapidly intractable as property calculation becomes more computationally expensive. In contrast, Bayesian optimization is known to be a more sample-efficient method,³⁰ and hence is finding success in tasks such as screening molecules with high power conversion efficiency for clean energy,³¹ optimizing reactions for molecular synthesis,³² and finding low-energy molecular conformers,³³ among others.^{34–36}

The potential benefits of Bayesian methods to optimize porous materials have been suggested by work by Simon and

coworkers.³⁷ Working with the data from previously screened ~70k covalent-organic frameworks (COFs), these authors showed that Bayesian optimization could find *ca.* 50% of the top-100 adsorbents for methane storage only exploring *ca.* 1% of the COFs. However, the approach used by these authors may not generalize well to searches aiming to optimize other material properties. For instance, their representation of the adsorbent consisted of a 12-component vector of five common (global) textural properties and simple counts of seven specific chemical elements. Such simple representation likely leverages that methane adsorption is primarily a (relatively) smooth function of textural properties. However, it may not be suitable when the property of interest also depends strongly on material chemistry.

On the other hand, traditional Bayesian optimization is designed to find one single optimal solution, which may turn out to correspond to a MOF design that may not be experimentally synthesizable or stable, or for which the performance prediction may have turned out to be unreliable. The task of optimizing a MOF performance metric while ensuring other properties (*e.g.*, synthesizability and stability) also have desirable values can be framed as a multi-objective optimization problem. Such formulation, however, assumes that all relevant metrics are known *a priori* and can be evaluated in similar manners.^{26,38} Multi-objective optimization cannot be realized, however, if some objectives can only be evaluated after screening is completed, or if we cannot anticipate all possible factors that should be accounted for during the search (*i.e.*, prediction reliability for each particular MOF). We thus take a different approach: finding multiple MOFs, different from one another, with desirable predicted values for the primary property of interest.

Specifically, in this work, we build a general and efficient framework for searching and finding several optimal MOF designs that are distinct from each other. Our framework is designed to be amenable to performance metrics that depend strongly on either MOF chemistry or textural properties, or both. More specifically, we combine the traditional tools of Bayesian optimization with the Vendi score—a statistical measure of diversity developed by Friedman and Dieng³⁹—to find a diverse set of promising MOF designs, each yielding a sufficiently high value for the metric of interest, instead of committing to a single optimal MOF that may not be synthesizable or stable. This comes in the form of promoting more exploration in the behavior of our optimization algorithm, selecting MOFs that are diverse from those already inspected. We name this framework Vendi Bayesian optimization (VBO).

We first statistically test the efficacy of combining a chemistry- and structure-aware MOF representation with VBO, using a randomly drawn subset of ~1000 MOFs as a testbed. We conducted these tests on the optimization of three performance metrics depending on the adsorption of NH₃. We chose metrics involving this molecule because NH₃ is important for our society as a precursor to fertilizers,⁴⁰ and could gain further prominence in the near future as an energy vector.⁴¹ From an application perspective, the three chosen metrics are relevant to rank MOFs for their potential to help make the synthesis of NH₃



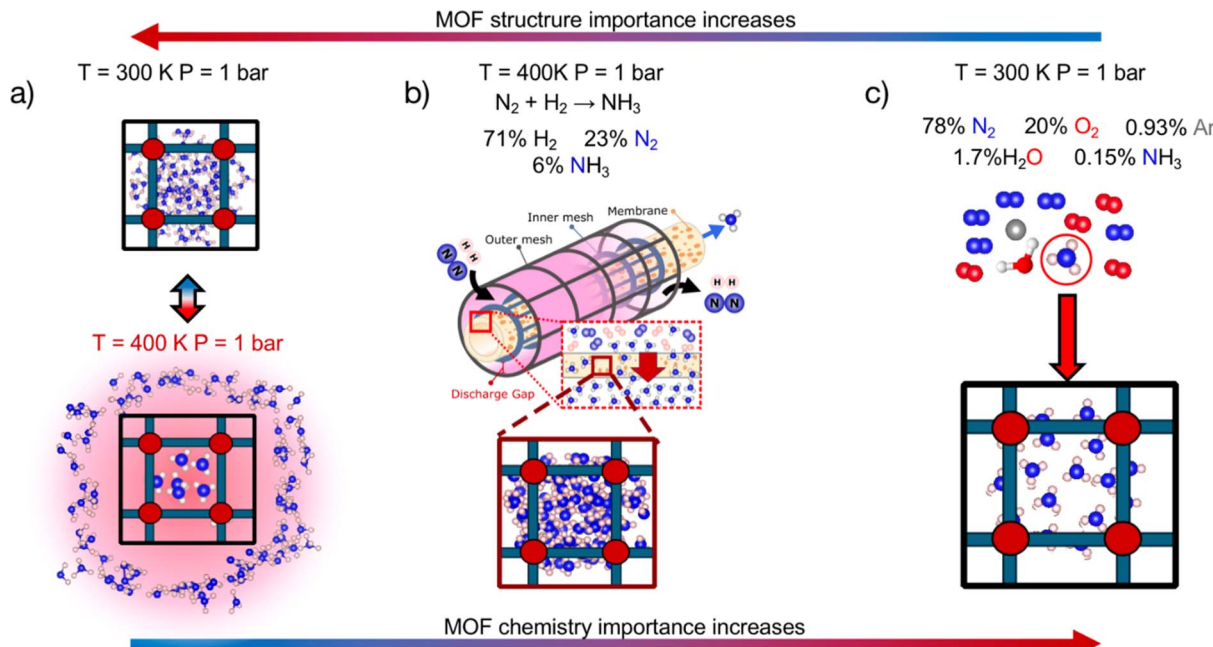


Fig. 1 Applications for which NH₃ adsorption-based MOF performance metrics were optimized to test the efficacy of our Vendi Bayesian optimization (VBO) framework. (a) Adsorptive NH₃ storage at ambient conditions with release at 400 K. (b) Membrane-based NH₃ removal from plasma reactors during NH₃ synthesis at 400 K and 1 bar. (c) Dilute NH₃ capture from air in adsorbent traps at ambient conditions. Gas-phase composition relevant to each application indicated at the top. The three chosen metrics present different levels of dependence on MOF chemistry and structure.

sustainable and carbon-free,⁴² and NH₃ storage and transportation easy, energy-efficient and safe.^{43–45} From a methods perspective, the three chosen metrics pose different challenges to our developed search method. Namely, the polarity of ammonia⁴⁶ and the different adsorption conditions associated with each application (Fig. 1) make the different metrics balance differently their dependence on MOF chemistry and textural properties (*vide infra*). On the other hand, each metric present different (mathematical) complexity on their relation to adsorption loadings.

Upon statistical testing of VBO efficacy, we finish this work with a real search campaign on a ~10 000-MOF hybrid database (*i.e.*, containing extant and hypothesized structures) to find MOFs with outstanding predicted NH₃ storage performance. We chose this application for the real search due to the growing interest of experimentalist chemists in the use of MOFs for NH₃ storage as reflected by the growing number of NH₃ adsorption measurements at 1 bar and 300 K (*i.e.*, ambient conditions) reported in recent years. For instance, Moribe *et al.* reported 10.5 mmol_{NH₃} g_{MOF}⁻¹ in Ga-PMOF,⁴⁷ Guo *et al.* 12.8 mmol_{NH₃} g_{MOF}⁻¹ in MIL-160,⁴⁸ Kim *et al.* 23.9 mmol_{NH₃} g_{MOF}⁻¹ in Mg-MOF-74 (ref. 49) and 23.5 mmol_{NH₃} g_{MOF}⁻¹ in Ni_acryl_TMA,⁵⁰ and Shi *et al.* 33.9 mmol_{NH₃} g_{MOF}⁻¹ in LiCl-MIL-53,⁵¹ among others.^{52,53} But despite growing interest, not much has been done to leverage search algorithms to identify promising MOFs for NH₃ storage. Thus, here we show how our developed VBO, a novel search algorithm for MOFs, can be used to fill such knowledge gaps. Furthermore, our analysis of the MOFs explored by our VBO provides new design rules to guide experimentalists developing MOFs for NH₃ storage.

2. Simulation methods

2.1. MOF database

About 12 000 structures from the 2019 CoRE MOF database¹⁴ and about 3000 structures created earlier using ToBaCCo-3.0 (ref. 6) were used as a starting point to ultimately create a hybrid database of ~10 000 structures. These MOF sources are complementary. CoRE MOFs are extant structures with high, but non-systematic, chemical and structural diversity that tend to feature small pores.⁵⁴ ToBaCCo MOFs are hypothesized structures with systematic, but medium, chemical and structural diversity that feature medium to large pores.⁵⁴ All MOFs underwent characterization of their void fraction, surface area, and pore size distribution using zeo++. A probe radius of 1.3 Å was used by zeo++ to determine the accessibility of pores through the percolation algorithm.⁵⁵ Then a probe of same size was used to determine the characteristic of the accessible pores. Note that the radius of 1.3 Å is adopted to match the kinetic radius of NH₃.⁵⁶ Failures during characterization calculations and assignment of charges to MOF atoms (see Section 2.2) ultimately reduced the total number of structures available for this work to around 10 000.

2.2. Monte Carlo simulations

Monte Carlo simulations were done using RASPA-2.0.^{57,58} Grand canonical Monte Carlo (GCMC) was used to predict adsorption loadings. Temperature and partial pressures of adsorbates in the gas phase were kept constant at the values relevant for the adsorption conditions of interest. Each simulation consisted of



10 000 equilibration cycles, followed by 10 000 production cycles. Each cycle consisted of as many Monte Carlo moves as molecules there are in the simulation box, but never less than 20. Moves corresponded to insertion/deletion, translation, and rotation (and swap for mixture cases). The Widom insertion method,⁵⁹ with at least 10 000 insertion moves, was used to calculate Henry's constants at the temperature of interest. Molecular interactions were modeled using the Lennard-Jones (LJ) and Coulomb potential. A cutoff of 12.8 Å was used for the LJ potential, and 12.0 Å for the Coulomb potential, after which distance Ewald summation was used.^{60,61} LJ parameters and charges for NH₃ and N₂ molecules were assigned according to the TRAPP force field^{62,63} for H₂O according to the TIP4P model,^{64–66} whereas for H₂ were obtained from the work by Darkrim and Levesque, including Feyman–Hibbs corrections.^{67,68} LJ parameters for MOF atoms were assigned according to the Dreiding force field,⁶⁹ or universal force field⁷⁰ if parameters from Dreiding were unavailable. LJ parameters for cross-interactions were obtained using Lorentz–Berthelot mixing rules. Note that the above LJ parameter selection have been used by Snurr and coworkers, and several others, to model NH₃ adsorption in MOFs.^{71–75} Charges for MOF atoms were assigned based on the best method available for each MOF subset. Thus, charges in ToBaCCo MOFs were assigned in earlier work using the MBBB method,⁷⁶ whereas for atoms in CoRE MOFs, charges were assigned using PACMOF.⁷⁷ MBBB is based on DFT calculations on MOF building blocks, which are directly inherited by the MOF, when constructed by ToBaCCo. PACMOF, on the other hand, is a machine learning model that was trained by Snurr and coworkers, from DFT calculations on complete MOF unit cells, to predict charges in MOF atoms, with an accuracy of 0.02e in mean absolute error ($R^2 = 0.99$). Moreover, the higher accuracy of PACMOF over other fast charge assignments was recently shown by Liu and Luan.⁷⁸ Example comparison between simulated adsorption isotherms using the methods herein against experimental ones are shown in Fig. S2.†

2.3. Assessed performance metrics

2.3.1 NH₃ storage. The incumbent method to store NH₃ relies on condensation at temperatures in the 238 to 253 K range, under pressures in the 10 to 15 bar range.^{38,79} Exploration of adsorptive NH₃ storage in the literature coincide on storing NH₃ at ambient conditions (300 K and 1 bar), but do not present consensus on the desired conditions for the release. Importantly, however, the performance of an adsorbent for ammonia storage depends on both the amount of NH₃ trapped at the storage conditions, $N_{\text{NH}_3}^{\text{ads}}$, and that retained in the adsorbent at the release conditions, $N_{\text{NH}_3}^{\text{des}}$. The difference between these two quantities defines the working (effective) storage capacity ΔN_{NH_3} as:

$$\Delta N_{\text{NH}_3} = N_{\text{NH}_3}^{\text{ads}} - N_{\text{NH}_3}^{\text{des}} \quad (1)$$

Due to its technical simplicity, here we consider the release of ammonia to be done simply by heating the adsorbent to 400 K at 1 bar (Fig. 1a). Note that as having enough space in the MOF pore is paramount to this application, ΔN_{NH_3} is expected to

be strongly influenced by MOF textural features such as pore size, void fraction and so forth.

2.3.2 NH₃ removal during plasma-assisted synthesis. The incumbent method to make NH₃ typically uses a pressure of 150 bar and a temperature of 650 K. However, as it turns out, sustainable, carbon-free NH₃ production requires synthesis at mild conditions.^{80–82} A promising method for NH₃ synthesis at 1 bar and 400 K is plasma-assisted synthesis in dielectric barrier discharge (DBD) reactors. In these reactors, low synthesis temperature is enabled by the accelerated breakdown of reactant molecules (N₂ and H₂) due to collisions with high-energy electrons in the plasma. But these electrons can also break down some of the freshly formed ammonia. Thus, a plasma reactor configuration that incorporates a porous membrane that remove ammonia as it forms, could protecting NH₃ from plasma decomposition (Fig. 1b), increasing energy efficiency.^{83,84} One of the desirable characteristics for the porous membrane are high adsorption of ammonia N_{NH_3} at the reaction conditions, but with high adsorption selectivity for ammonia α_{NH_3} over N₂ and H₂, where:

$$\alpha_{\text{NH}_3} = (N_{\text{NH}_3}/y_{\text{NH}_3})/((\sum N_i)/(\sum y_i)) \quad (2)$$

where y_{NH_3} is the molar fraction of NH₃ in the gas phase, and N_i and y_i are the adsorbed loading and molar fractions in the gas phase, respectively, of all other species i. Assuming a 3 : 1 H₂ : N₂ feed ratio and a conversion of 10%, here y_{NH_3} , y_{N_2} and y_{H_2} are assumed to be 0.06, 0.23, 0.71, respectively. Seeking to account for both adsorption and selectivity, here we use M_{ATS} as a performance metric where:

$$M_{\text{ATS}} = \alpha_{\text{NH}_3} \times N_{\text{NH}_3} \quad (3)$$

Note that selectivity, α_{NH_3} , is a reflection of the attraction of the MOF to NH₃ relative to N₂ and H₂, and hence is expected to be strongly influenced by chemistry. On the other hand, the adsorption capacity N_{NH_3} at non-dilute conditions is expected to also be influenced by MOF pore space. Thus, the complete metric M_{ATS} is expected to be influenced by both MOF chemistry and textural features. Also note that diffusion selectivity is an important aspect of choosing a material for a membrane. This selectivity could be incorporated into the performance metric (or could be considered in a subsequent screening step). However, for the purpose of testing the VBO framework, we decided to focus on the adsorption aspects of the membrane.

2.3.3 NH₃ capture from air. NH₃ leakage during storage and transportation is a persistent risk. The maximum NH₃ concentration that individuals can safely breath for 1 h is 1500 ppm.⁸⁵ One way to mitigate the risk is to accompany storage and transportation infrastructure with adsorbent traps that can selectively adsorb substantial amounts of dilute NH₃ over other molecules in air, including water (Fig. 1c). Preferential NH₃ adsorption over H₂O is most challenging because H₂O is polar like NH₃, but it would be present at a higher concentration in air. Thus, we decided to consider a MOF to be potentially useful only if it is hydrophobic, for which we calculate:



$$\delta_{\text{HPHB}} = 1, \text{ if } K_{\text{H}_2\text{O}} \leq 5 \times 10^{-6} \text{ mol kg}^{-1} \text{ Pa}^{-1} \\ \delta_{\text{HPHB}} = 0, \text{ if } K_{\text{H}_2\text{O}} \geq 5 \times 10^{-6} \text{ mol kg}^{-1} \text{ Pa}^{-1} \quad (4)$$

where $K_{\text{H}_2\text{O}}$ is the adsorption Henry's constant of H_2O at 300 K and δ_{HPHB} equal to one (zero) indicates that the MOF is hydrophobic (hydrophilic), in consistency with the $K_{\text{H}_2\text{O}}$ threshold for MOF hydrophobicity determined by Moghadam *et al.*⁸⁶ Then, we use as performance metric:

$$M_{\text{ATSTH}} = M_{\text{ATS}} \times \delta_{\text{HPHB}} \quad (5)$$

where M_{ATS} is calculated from eqn (3) and (2), with i corresponding to N_2 , O_2 and Ar . Adsorption loadings are calculated for a $\text{N}_2/\text{O}_2/\text{Ar}/\text{NH}_3$ mixture with $y_{\text{N}_2} = 0.78$, $y_{\text{O}_2} = 0.21$, $y_{\text{Ar}} = 0.0075$ Ar and $y_{\text{NH}_3} = 0.0015$ (*i.e.*, 1500 ppm NH_3) at 300 K and 1 bar. The above metric circumvents the need to calculate H_2O adsorption in MOFs, which is known to require extremely long simulations.⁸⁷ Note that an analogous strategy to the above was used by Smit and coworkers to discover MOFs for CO_2 capture from wet flue gas.²⁶ Note that as having enough space in the MOF pore to store the dilute quantities of NH_3 originally in air is not a concern, MOF performance, and thus M_{ATSTH} is expected to be primarily influenced by the ability of the MOF to attract NH_3 , and hence by MOF chemistry.

3. Diversity-driven MOF optimization

3.1. Workflow overview

An overview of our diversity-driven MOF optimization/search framework is presented in Fig. 2. To start a MOF (design) optimization campaign, we randomly draw two MOFs and calculate their performance metrics using molecular

simulations. These two datapoints are then used to train a Gaussian Process (GP) regression model⁸⁸ whose kernel is designed to account for both chemistry and physics (see Section 3.2). The GP is trained to predict the performance metric and provide the uncertainty associated with the prediction. This fitted GP is then used to predict the performance of all MOFs in the hybrid database. From these predictions, our Vendi Bayesian Optimization (VBO) algorithm selects the next most promising MOF candidates for which to calculate the performance metric using molecular simulations.

The first candidate that VBO selects is the one corresponding to the most “optimistic” performance prediction made by the trained GP. The remaining candidates are selected only after we prune 10% of the database. The pruning is done by taking out of the database 10% of the MOFs that, if added to the set of MOFs previously chosen by VBO and assessed *via* molecular simulations, would yield the lowest diversity change of that set. In our workflow, diversity of a MOF set is calculated using the Vendi score (see Section 3.3). The lower the Vendi score, the lower the diversity of the set. Thus, the MOFs removed from the database are those that would yield the lowest Vendi score if added to the set of MOFs that have been selected by our VBO algorithm.

Given that for each MOF the GP predicts a distribution of possible performance metric values, our VBO algorithm uses the upper confidence bound (UCB) criterion to assess the “potential” of a MOF. Specifically, the UCB is the mean value (μ) of the distribution of predictions for the MOF plus two times the standard deviation (σ). Ideally, upon evaluation with molecular simulation, some of the MOF selected by our VBO algorithm should have a higher value of the performance metric than the MOFs previously evaluated in this same manner. Regardless,

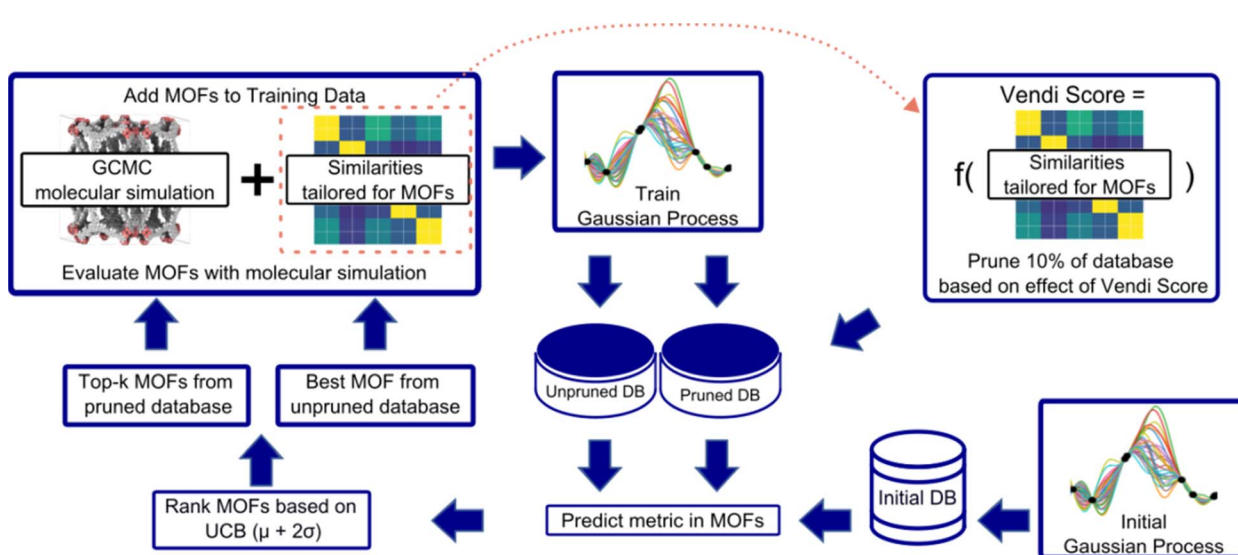


Fig. 2 Workflow for our VBO framework. an initial GP, trained with data for two randomly chosen MOFs, is used to predict the performance metric in the starting database. $k + 1$ MOFs are selected for molecular simulation evaluation based on the upper confidence bound (UCB) acquisition function. One MOF is chosen as the MOF scoring the highest UCB just as in standard Bayes optimization. The remaining k MOFs are selected based on UCB but only after 10% of the database is pruned. The MOFs pruned from the database are the MOFs that would increase the least the Vendi score of the cumulative set of MOFs evaluated by molecular simulation. The top $k + 1$ MOFs selected are then evaluated using molecular simulations. To perform a new iteration, the molecular simulation data for the newly evaluated $k + 1$ MOFs are added to the data for training the GP, and the MOF selection process is repeated.



upon completion of the evaluation with molecular simulation for MOFs that had been selected by the VBO algorithm, a new GP model is trained leveraging the newly generated data, and selection of new candidates is done again using the same procedure as described above. This procedure is repeated until either a preset target number of iterations is achieved or the highest value of the performance metric in the MOFs evaluated with molecular simulation no longer improves.

3.2. MOF representation

Each MOF is chemically characterized by the Morgan fingerprints⁸⁹ of its constituent building blocks (nodes and linkers), which are extracted from each MOF using MOFid.⁷ MOFid provides the SMILES strings⁹⁰ of the building blocks, which are used as input for RDKit to provide the fingerprints. Here, each fingerprint is a vector whose components describe the atom groups of the corresponding node or linker. Each MOF is also structurally characterized by its detailed pore size distribution and global textural properties usually used in the MOF field. Namely, specific pore volume, void fraction, specific surface area, largest and diffusion-limiting pore diameters, and metal-to-nonmetal content ratio. We design a specific similarity kernel for MOFs. This new kernel is the one we use for the GP and the calculation of the Vendi score in our VBO framework. More specifically, if we denote two different MOFs by x_1 and x_2 , then the similarity between these MOFs is given by a specialized kernel function K that is an average of four different kernels, where each kernel K_i specializes in one particular aspect of MOFs and is weighted by a factor w_i . Namely, the kernel similarity between two MOFs x_1 and x_2 is defined as:

$$K(x_1, x_2) = w_1 K_{\text{node}}(x_1, x_2) + w_2 K_{\text{linker}}(x_1, x_2) + w_3 K_{\text{global}}(x_1, x_2) + w_4 K_{\text{PSD}}(x_1, x_2) \quad (6)$$

K_{node} and K_{linker} are each a kernel function computing the Tanimoto similarity⁹¹ between the Morgan fingerprints of either two nodes or two linkers, respectively (Fig. 3a). The Tanimoto similarities between Morgan fingerprints have been found to capture important differences in molecule chemistry, and has been shown effective at guiding machine learning models for search purposes in other areas.⁹² As a MOF could have more than one type of node or linker, we consider all possible pairwise comparisons and use the average value of K_{node} or K_{linker} . On the other hand, K_{global} operates on the global textural properties, and is defined to be the exponential of the Euclidean distance between the two vectors containing the (normalized) values of the above properties for the two MOFs being compared (Fig. 3b). This is analogous to what Simon and coworkers did for COFs.⁹³ Finally, K_{PSD} is a new kernel proposed by us, which computes the difference between the pore size distributions (PSDs) of the two MOFs being compared. We do this by using the Jensen–Shannon divergence (JSD).⁹⁴ Given two PSDs P and Q , this function returns:

$$\text{JSD}(P, Q) = \frac{1}{2}(\text{KLD}(P, M) + \text{KLD}(Q, M)) \quad (7)$$

where $M = \frac{1}{2}(P + Q)$ is a mixture distribution of the original two P and Q and:

$$\text{KLD}(P, Q) = \sum_{s \in S} P(s) \log \left(\frac{P(s)}{Q(s)} \right) \quad (8)$$

refers to the Kullback–Leibler divergence (KLD) between P and Q . Here S is the set of possible pore sizes, and $P(s)$ and $Q(s)$ give the probability of a particular pore size s in each of the two MOFs. JSD computes the distance between the two distributions, giving a symmetric and bounded metric for their difference. Our kernel K_{PSD} subsequently calculates the similarity between the two distributions as $(1 - \text{JSD})$.

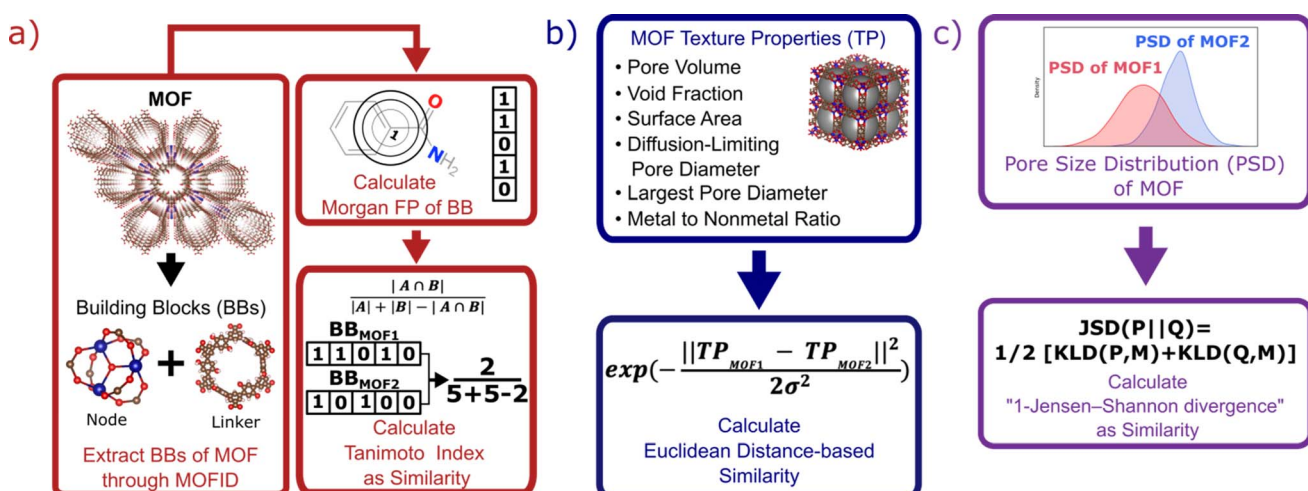


Fig. 3 Schematic representation of methods to calculate kernel similarity between MOFs. (a) Chemical similarity (K_{node} and K_{linker} kernels) obtained by decomposing two MOFs into their building blocks, and calculating the Tanimoto index between the Morgan fingerprints of their building blocks. (b) Global textural properties similarity (K_{global} kernel) obtained by calculating the radial basis function kernel of the Euclidean distance between the property vectors of two MOFs. (c) Detailed pore structure similarity (K_{PSD} kernel) obtained by calculating the difference between the pore size distributions (PSDs) of two MOFs. The different kernels cover different aspects of MOFs, and by tuning the weights of each kernel, the representation is adaptable to prediction of properties with different level of dependence on MOF chemistry and structure.



3.3. Vendi score

The Vendi score (VS) is key to encourage our optimization framework to find many diverse solutions, hence avoiding commitment to a single MOF design “solution” that might be infeasible to produce and test experimentally. The VS is a function whose input is the $n \times n$ similarity matrix K representing data points in a set of size n . The VS is calculated as the exponential of the Shannon entropy of the normalized eigenvalues of K , denoted by λ_i , as follows:

$$\text{VS}(K) = \exp\left(-\sum_{i=1}^n \lambda_i \log \lambda_i\right) \quad (9)$$

Friedman and Dieng³⁹ showed that the VS is a mathematically well-defined diversity metric and quantifies the effective number of unique elements in a set.³⁹ Here, the elements of the similarity matrix are calculated using eqn (6), meaning that the GP model and the VS use the same underlying mathematical object. To keep the output of the kernel function consistent across calculations of the VS, we set the weights w_i in eqn (6) to all be equal to 0.25. However, note that the weights in eqn (6) take different values when training the GP model, where they are optimized for prediction.

3.4. Vendi Bayesian optimization (VBO) framework

3.4.1 Overview. If MOFs are denoted by x and a MOF database by \mathcal{X} , where $x \in \mathcal{X}$, and if f is a “black-box” function that returns the scalar value of the property or performance metric of interest (*i.e.*, $f : \mathcal{X} \rightarrow \mathbb{R}$), then we aim to find the MOF x^* that maximizes the value of the performance metric. More formally, we find x^* such that:

$$x^* = \underset{x \in \mathcal{X}}{\operatorname{argmax}} f(x) \quad (10)$$

The above makes f an objective function that models the mapping between a given MOF and its performance metric. Here f is approximated by a GP that iteratively improves its “understanding” of f based on evaluations of f for specific MOFs x . Here, evaluating $f(x)$ means running molecular simulations to calculate the relevant performance metric for a given MOF x . However, our VBO framework is also amenable to experimental work, where performance metrics are measured *via* experiments instead of molecular simulations. In each case, our VBO framework enables finding the optimal MOF x^* in as few evaluations as possible, to overcome time and/or cost constraints associated with simulations or experiments.

3.4.2 Surrogate model. The first component of our VBO framework is a surrogate model that expresses a belief about f based on previous evaluations of f —*i.e.*, a belief about the relationship between MOF chemistry/structure and performance. Here, the surrogate model is a GP (see comparison with other models in Section S2†), which, as any GP, does not yield a single prediction of f for a given x , but rather a set of predictions that follow a normal (Gaussian) distribution \mathcal{N} such that:

$$f(x) \sim \mathcal{N}(\mu, \sigma^2) \quad (11)$$

where μ and σ are the mean and the standard deviation of the predictions, respectively. When conditioned on a training set, μ and σ are updated to be the posterior predictive mean and standard deviation, reflecting information learned from the training data. Intuitively, μ and σ represent the value that $f(x)$ is most likely to take and the uncertainty about the predicted μ , respectively. We refer to Rasmussen and Williams⁸⁸ for a more thorough treatment of GP learning. To fully specify a GP, one needs a mean value that describes the behavior of f in the absence of data, and a kernel K that calculates similarities between different x ; our choice of K was described in Section 3.2. At each iteration of our VBO framework, the values of the performance metric obtained from molecular simulation are normalized so that they range from -1 to 1 (we use the constant zero mean function). The parameters of the GP model, including the weights w_i in eqn (6) as well as the mean value and a noise factor, are tuned to maximize the fit to the training data, quantified by the marginal log likelihood of the data, as is standard in Gaussian process modeling.⁸³

3.4.3 Acquisition function. The second component of our VBO framework is an acquisition function $\alpha(x)$ that guides us towards promising candidate MOFs that are likely to yield high performance and that have not been evaluated. A good acquisition function should balance exploration (learning about how $f(x)$ behaves across the space) and exploitation (zeroing in on high-performance regions). Here, we opt for the Upper Confidence Bound (UCB) function.⁹⁵ UCB adds μ and σ , the mean and standard deviation of the GP prediction, with the latter multiplied by a trade-off factor β , which we set to 2 here:

$$\alpha(x) = \mu + 2\sigma. \quad (12)$$

This simple expression elegantly captures the balance between exploration of MOFs we are uncertain about (with high σ), and exploitation of MOFs predicted to yield high performance (with high μ). In addition to its interpretability, Taw and Neaton⁹⁶ demonstrated good optimization performance of the above acquisition function to optimize methane uptake capacity of MOFs. At each iteration of Bayesian optimization, we find the MOF that maximizes the UCB score to evaluate $f(x)$ with. We repeat this process until our evaluation budget is depleted, each time updating the GP and the UCB score with the newly observed MOFs.

3.4.4 Solution diversification. Unlike regular Bayesian optimization, VBO iteratively prunes the search space (*i.e.*, the database) by removing remaining candidates that are too similar to those that have been previously selected for evaluation. This removal results in even more exploration than enabled by the acquisition function. Formally, consider a candidate MOF x of unknown performance that we may query. We compute the increase in VS (ΔVS) that we would obtain if we were to evaluate $f(x)$ and add x to the set S containing the MOFs we already selected. That is:

$$\Delta\text{VS} = \text{VS}(S \cup \{x\}) - \text{VS}(S) \quad (13)$$



If x is different from the data points in S , querying x will add more diversity to our data set, as reflected by a large ΔVS . If, on the other hand, x is similar to the points in S , ΔVS will be small. At each iteration, we compute ΔVS for each of the remaining candidate MOFs, and remove the MOFs that yield the lowest ΔVS until the remaining pool of candidates is reduced by ten percent. We thus reduce the effective search space at each iteration, removing candidates that are too similar to those already acquired.

This modification of traditional Bayesian optimization aims at building a diverse set of high-performance MOFs. While this increase level of exploration does not guarantee improved optimization performance, we do not necessarily sacrifice the top MOF either. As the diversity-aware pruning step is reset at each iteration, if we have found a region in our search space that contains very good candidates, our acquisition function allows us to come back to this region (*i.e.*, zeroing in on the top MOF) once other promising regions have been explored. We can also think of this strategy as searching over multiple promising regions at the same time.

4. Results and discussion

4.1. Expressiveness of the MOF-specific kernel

Although the representation of a MOF is inherently multidimensional, the plots in Fig. 4 maps MOFs onto a reduced two-dimensional space, by applying multidimensional scaling (MDS)⁹⁷ to the covariance matrix of the MOFs, which was calculated using the kernel defined earlier by eqn (6). MDS conveys the similarity-dependent original distances between MOFs in multidimensional space, so that in Fig. 4 similar MOFs appear close to each other. From Fig. 4a, the complementary of CoRE MOFs (blue points) and our ToBaCCo MOFs (orange points) is apparent as the groups separate into individual regions. The usual differences between extant CoRE MOFs and hypothesized MOFs such as our ToBaCCo MOF have been pointed out previously in work by others such as Kulik and coworkers.⁵⁴ For instance, CoRE MOFs tend to feature smaller pores and a more diverse selection of metals. ToBaCCo MOFs exhibit a systematic variation in textural properties, focusing on metals Cr, Zr, Mn, Co, Cu, and Zn. Therefore, the observed segregation in Fig. 4a indicates that our kernel captures meaningful similarities/differences between MOFs.

Analogously, we present reduced dimensionality plots but only for a random subset of 1000 MOFs uniformly extracted from the $\sim 10\,000$ hybrid database, and for which the performance metrics pertinent to NH_3 storage, removal during plasma-assisted synthesis, and capture from air (ΔN_{NH_3} , M_{ATS} and M_{ATSTH} , respectively) were calculated using molecular simulation. Upon coloring the points based on the value of each performance metric in the corresponding MOF, it is apparent that segregation also tends to occur on the basis of performance (Fig. 4b-d). For instance, Fig. 4b shows MOFs with $\Delta N_{\text{NH}_3} < 5 \text{ mmol g}^{-1}$ locating in an outer ring, MOFs with $5 \text{ mmol g}^{-1} < \Delta N_{\text{NH}_3} < 15 \text{ mmol g}^{-1}$ locating in the inner region, and MOFs with $\Delta N_{\text{NH}_3} > 15 \text{ mmol g}^{-1}$ locating in a lower-right cluster. Such

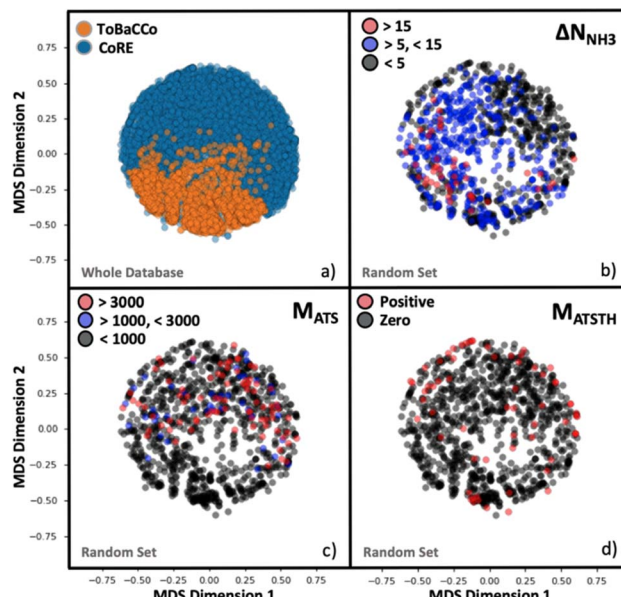


Fig. 4 MOF mapping onto two-dimensional plots by using multidimensional scaling (MDS) representations. (a) All MOFs in the hybrid database colored by their origin (either the ToBaCCo database or the CoRE database). (b-d) 1000 random MOF subset, colored by range of ΔN_{NH_3} (b), M_{ATS} (c), and M_{ATSTH} (d) performance metrics. The extent of segregation observed is a harbinger of the efficacy of our MOF kernel similarity as input to train the GP.

segregation indicates how well our measure of similarity (*i.e.*, our kernel) is conducive to learning.

The extent at which our kernel facilitates learning is illustrated in Fig. 5, which shows parity plots comparing the prediction of the performance metrics N_{NH_3} , M_{ATS} and M_{ATSTH} by corresponding GP models trained on molecular simulation data of the 1000 random MOF subset. The GPs trained to predict ΔN_{NH_3} and M_{ATS} (Fig. 5a and b) present relatively similar correlations between their predictions and the actual values (*i.e.*, ground truth) of the corresponding performance metrics. Namely, R^2 values of 0.59 and 0.37 for ΔN_{NH_3} and M_{ATS} , respectively. On the other hand, the GP trained for the M_{ATSTH} case seems to face higher difficulty in learning to predict the performance metric, which is reflected by an R^2 value of -0.06 (Fig. 5c). Such difficulty is partly due to the exceptional roughness of M_{ATSTH} as a function of MOF chemistry/structure—which partly motivated the selection of this metric for our testing. The roughness of M_{ATSTH} stems from the rather binary character of the metric, which is either zero or positive based on whether the MOF is deemed hydrophobic or not based on the threshold value of $K_{\text{H}_2\text{O}}$, resulting in discrete changes to M_{ATSTH} that are difficult to capture by machine learning models. Yet, as we will demonstrate shortly, our VBO framework remains effective at optimizing these metrics, including, perhaps surprisingly, M_{ATSTH} .

At this point, let us note that the optimized weights (w_i) for the GP models (Table S3†) confirm our hypotheses of what MOF aspects control performance for each application. For instance, the chemical similarity kernel K_{node} weighs 0.97 in the model



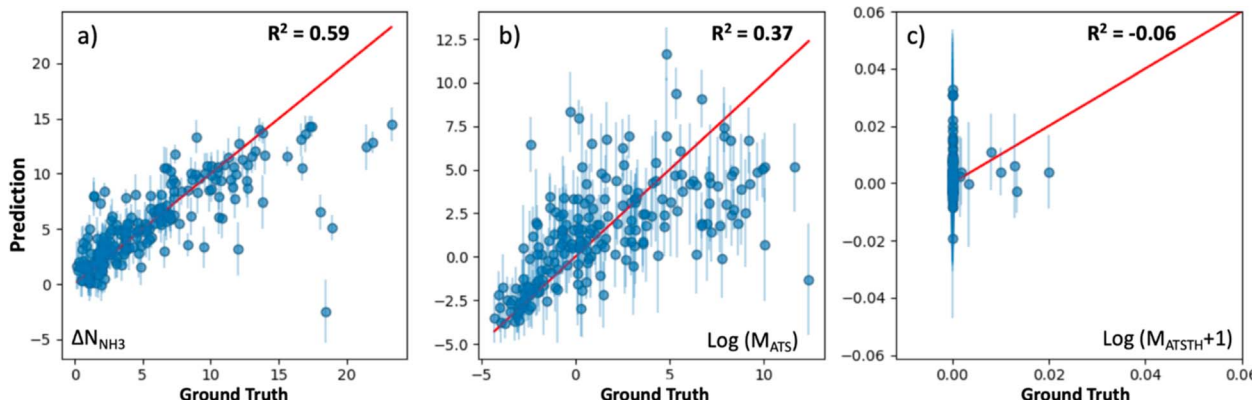


Fig. 5 Prediction performance of GP models (trained on a subset of 1000 random MOFs extracted from the hybrid database) to predict (a) ΔN_{NH_3} , (b) M_{ATS} , and (c) M_{ATSTH} . GP predictions appear on the vertical axis, while the ground truth (from molecular simulation) appears on the horizontal axis. The parity line is presented in red. Each point represents the prediction for a MOF, with the corresponding error bar representing the uncertainty of the predictions based on the prediction standard deviation. The observed prediction performance was found on subsequent statistical testing to be sufficient to make VBO effective.

that predicts M_{ATSTH} but only weighs 0.02 in the model that predicts ΔN_{NH_3} . By contrast, K_{PSD} weighs 0.37 in the model that predicts ΔN_{NH_3} , but only weighs 0.01 in the model that predicts M_{ATSTH} . On the other hand, all kernels weigh rather similarly in the model that predicts M_{ATS} .

4.2. Statistical testing of VBO efficacy

The efficacy of VBO was statistically assessed by simulating our workflow (Fig. 2) ten times on the subset of randomly selected 1000 MOFs, to iteratively optimize MOF design for the ΔN_{NH_3} , M_{ATS} and M_{ATSTH} metrics. During each run, two MOFs were randomly selected to be the initial training set, and 100 MOFs were evaluated in 20 batches of five MOFs each iteration (*i.e.*, when 10% of the MOF subset was evaluated, the run stopped). Each time our VBO workflow was run, an analogous run without the Vendi score-based pruning (*i.e.*, a regular Bayes optimization run) was done in parallel for comparison, as well as random search consisting of the evaluation of 100 randomly selected MOFs within the subset. The lines in Fig. 6 present the average progress of the VBO (blue), Bayesian optimization (green) and random search (orange) runs, whereas the corresponding shaded areas represent the corresponding standard errors.

As evidenced by Fig. 6, although the uncertainty region for VBO and Bayesian optimization tend to overlap, on average VBO did equal or better than Bayesian optimization, when assessed based on the highest value for the metric encountered by the end of 100 evaluations. Notably, VBO outperformed Bayesian optimization for the evaluation of the M_{ATSTH} metric for NH_3 capture from air. On the other hand, both VBO and Bayesian optimization clearly do better on average than random search when compared by the abovementioned criterion. Furthermore, the uncertainty regions for the latter two methods and random search barely overlap, suggesting that in a worst-case scenario VBO and Bayesian optimization would do at least as well as a best-case scenario random search that explores ten percent of the available design space.

But the most significant difference between VBO and Bayesian optimization is the more diverse exploration of the design space by VBO. This fact is evidenced by the consistently higher Vendi score among evaluated MOFs as VBO progresses compared to Bayesian optimization. As expected, random search tends to result in the highest diversity among evaluated MOFs as the search progresses. But it is surprising that for the optimization of M_{ATSTH} our VBO ended up on average with a higher diversity of evaluated MOFs than random search. Ultimately, the average behavior of the Vendi score in VBO *versus* Bayesian optimization is indicative that VBO is bound to create a more diverse pool of promising MOFs for a given application.

4.3. Full database search for MOFs for NH_3 storage

Encouraged by the statistical efficacy of our VBO framework, we decided to perform a full VBO run on the complete hybrid database (*i.e.*, $\sim 10\ 000$ MOFs) to optimize ΔN_{NH_3} . Specifically, to find MOFs with potential for NH_3 storage, considering storage at 1 bar with storage/release through a 300 K to 400 K thermal swing. Fig. 7a presents the progress of the performed VBO run of 20 iterations (each iteration corresponds to a batch of 20 MOFs), comparing it against a random search (technically consisting of the previously randomly selected 1000 MOFs on which VBO was previously tested in Section 4.2). Evidently, VBO greatly outperforms random search, with the former identifying MOFs with ΔN_{NH_3} values approaching as high as $30\ \text{mmol}_{\text{NH}_3}\ \text{g}_{\text{MOF}}^{-1}$, whereas the latter did not identify MOFs with ΔN_{NH_3} values higher than $\sim 23\ \text{mmol}_{\text{NH}_3}\ \text{g}_{\text{MOF}}^{-1}$.

Notably, the outperformance of VBO relative to random search occurred despite VBO terminating early at *ca.* 400 evaluations. This early termination was made because the highest ΔN_{NH_3} value within the evaluated MOFs did not change significantly after around 80 evaluations. However, note that one should not be tempted to consider subsequent MOF evaluations after the 80th evaluation point as wasteful, as these evaluations

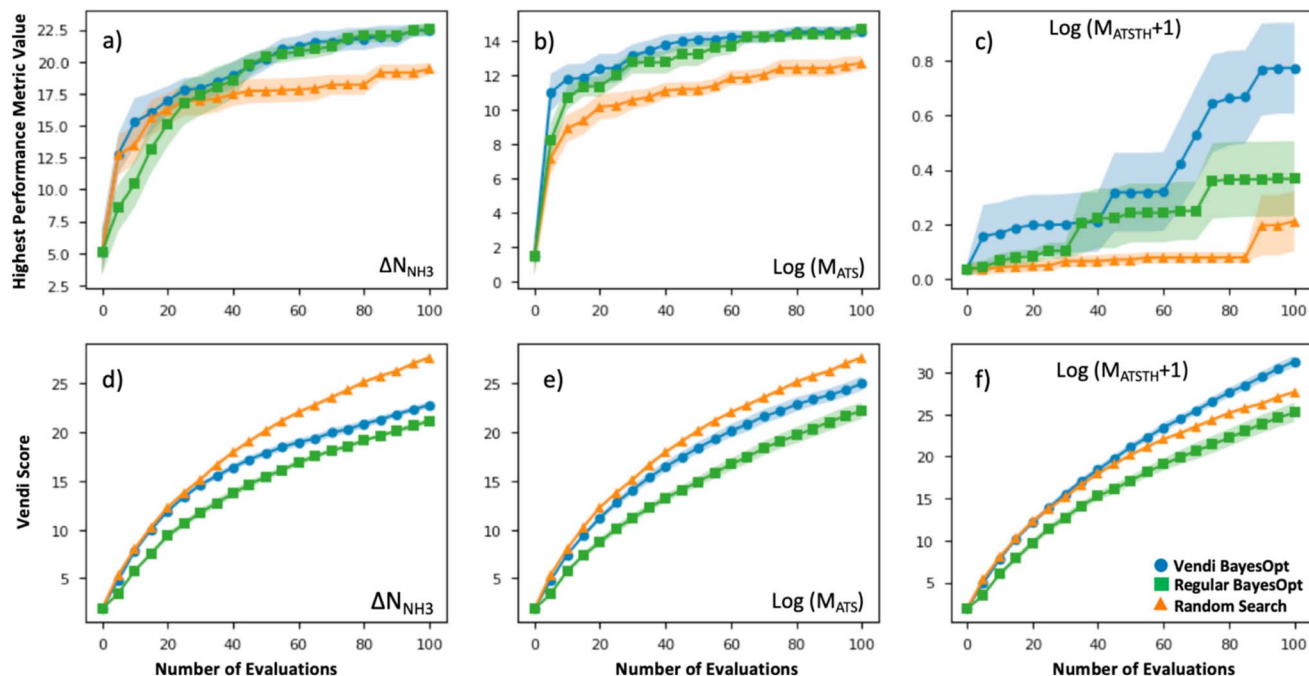


Fig. 6 Efficacy of VBO (blue) applied on a 1000 subset of random MOFs compared to Bayesian optimization (green) and random search (orange). Top row presents the evolution of the highest value of the performance metric as the number of MOF evaluations increases for (a) ΔN_{NH_3} for ammonia storage, (b) M_{ATS} for ammonia removal from plasma reactor, and (c) M_{ATSTH} for ammonia capture from air. Bottom row presents the evolution of the Vendi score for the set of evaluated MOFs as the number of MOF evaluations increases for (d) ΔN_{NH_3} for ammonia storage, (e) M_{ATS} for ammonia removal from plasma reactor, and (f) M_{ATSTH} for ammonia capture from air. Results in (a)–(f) are averaged across 10 repeat runs, the average value is indicated by the solid line, whereas the standard deviation is indicated by the shaded area. Both VBO and Bayesian optimization outperformed random search, but VBO provided higher diversity of MOF "solutions."

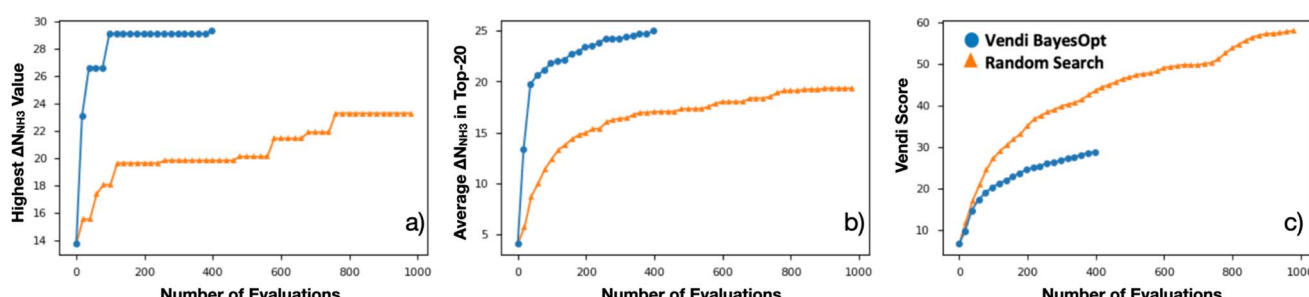


Fig. 7 Evolution of VBO campaign (blue) in the ~10 000 MOF database, when searching for MOFs for NH_3 storage, compared to the evolution of the random search (orange). (a) Evolution of the highest ΔN_{NH_3} found among evaluated MOF at a given point in the campaign. (b) Evolution of the average ΔN_{NH_3} among the top-20 evaluated MOFs at a given point in the campaign. (c) Evolution of the Vendi score of evaluated MOFs at a given point in the campaign. Note that the VBO campaign was ended early due to negligible changes in the highest ΔN_{NH_3} since the 80th evaluation. Once again VBO greatly outperformed random search.

enabled to strengthen the pool of promising MOF designs for NH_3 storage. A fact that is evidenced by the steady improvement in the average ΔN_{NH_3} for the "top-20" evaluated MOFs from the 80th to the 400th evaluation (Fig. 7b). Importantly, this improvement in average ΔN_{NH_3} was accomplished while steadily improving the diversity of the evaluated MOF as indicated by the steady improvement in the Vendi score within the same range of evaluations (Fig. 7c). The latter creates confidence that the pool of promising MOFs to be suggested for future synthesis and experimental testing to be more diverse than provided by other methods.

4.4. Data-driven MOF design rules

As noted earlier, a benefit of computational MOF screening is the emergence of structure–performance relationships, which are useful to establish design rules that experimentalists could leverage to conceive adsorbent designs of their own (not even necessarily for MOFs). Importantly, the emergence of these relationships allows extracting value from computational screenings independently of the success in synthesizing and testing the specific MOF designs recommended by the screening. However, the nature of the emerging relationships is empirical, and thus depend on a sufficiently large number of

observations being made to create clear trends. Conveniently, here, while the number of evaluated MOFs is lower than in other screening studies that relied on exhaustive search, the bias of our selection method towards “good” MOFs allow us to still define well the “interesting” region of the relationship relevant to optimize the performance metric of interest.

For instance, although in Fig. 8a there is a dearth of data for MOFs with average pore diameter (APD) larger than 14 Å, it is apparent that the optimal average pore diameter and for NH₃ storage at the conditions herein proposed is 10 Å. Note that the scarcity of data for MOFs with APD larger than 14 Å is due to reluctance by the VBO algorithm to pick MOFs in that range of APD, probably due to rapidly learning that APDs larger than 14 Å tend not to optimize ΔN_{NH_3} . An APD of 10 Å seems to

compromise confinement effects (*i.e.*, overlap of interaction potentials) to enhance NH₃ attraction to the pore walls and having sufficient space to accommodate NH₃ molecules. To be sure, an APD of 10 Å should be interpreted as necessary, and not as a sufficient condition to optimize ΔN_{NH_3} , as evidenced by the wide range of ΔN_{NH_3} values that can be observed for that APD value. The color coding in Fig. 8a suggests that such variability in ΔN_{NH_3} at APD equal to 10 Å is *partly* explained by variations in MOF void fraction—with MOFs with void fraction around 0.7 tending to appear at the top. In other words, given two MOFs with APD equal to 10 Å, the one with higher void fraction probably corresponds to a higher ΔN_{NH_3} , again partly due to the implication that higher void fraction allows more space to accommodate NH₃ molecules.

On the other hand, note that while attraction of NH₃ to the pore walls (as reflected by the heat of adsorption Q_{st}) is desired, too strong an attraction is detrimental to ΔN_{NH_3} as it prevents the adsorbed NH₃ molecules to be easily released. From our collected data, it seems that a Q_{st} of 33 kJ mol⁻¹ is optimal for ammonia storage at the conditions herein proposed (Fig. 8b). Analogous to our APD analysis, a Q_{st} of 33 kJ mol⁻¹ should be taken only as a necessary but not sufficient condition to optimize ΔN_{NH_3} . Indeed, there is a wide range of ΔN_{NH_3} values at Q_{st} equal 33 kJ mol⁻¹. The color coding in Fig. 8b partly explains this variability on the basis of surface area variations—with MOFs with surface area around 4000 m² g⁻¹ tending to appear at the top, as they provide a larger number of sites with optimal interaction strength. Note that inspecting Fig. S6,† it seems that a Q_{st} value around 33 kJ mol⁻¹ enables recovering up to 95% of the NH₃ molecules adsorbed at the storage conditions.

We acknowledge, however, that a design rule centered around Q_{st} is somewhat abstract as this quantity does not depend only on MOF chemistry, but also on MOF structure. In an attempt to provide some chemistry-based MOF design rules for NH₃ storage, we decided to explore trends in elemental composition among outstanding MOFs. Specifically, for each element in the periodic table, we calculated its average percent content in the top-14 MOFs evaluated with molecular simulation and compared this value with the corresponding average percent content in all ~10 000 MOFs in the database (Fig. S7†). Then we used a *t*-test to assess the statistical significance of observed differences.

Fig. 9 shows the *p*-values for the *t*-test for the elements present in the top-14 MOFs. Using a *p*-value threshold of 0.1, it seems that C, H, and Ca are elements that are, with statistical significance, more abundant within the top-14 MOFs for NH₃ storage than in MOFs at large. Using our understanding of MOF structure, we rationalize that the higher abundance of C, H is probably just a reflection of the optimal APD for ammonia storage being larger than the median APD in the database—*i.e.*, larger pores imply longer linkers, hence more C and H content. On the other hand, we could not find an alternative explanation for the higher abundance of Ca within the outstanding MOFs, suggesting a primarily chemical effect—after all, CaCl₂ is a popular ammonia adsorbent.⁹⁸ To be sure, though, due to the role other MOF features play on ΔN_{NH_3} , the presence of Ca

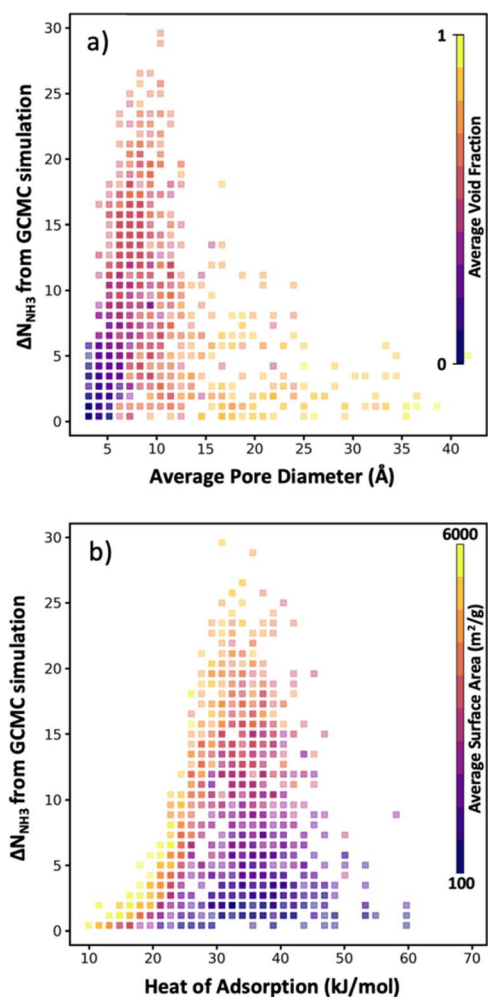


Fig. 8 Plots of structure–performance relationships for NH₃ storage. Each square bin corresponds to a combination of the ΔN_{NH_3} performance metric and MOF property, where the transparency of each square bin is indicative of the number of MOFs in the bin, and the color of each bin reflects the average value of the property in the side color scale across all MOFs in the bin. (a) ΔN_{NH_3} versus MOF average pore diameter (APD), with each bin colored by MOF void fraction. (b) ΔN_{NH_3} versus heat of adsorption Q_{st} , with each bin colored by gravimetric surface area. Optimal APD and Q_{st} appears to be 10 Å and 33 kJ mol⁻¹, respectively.



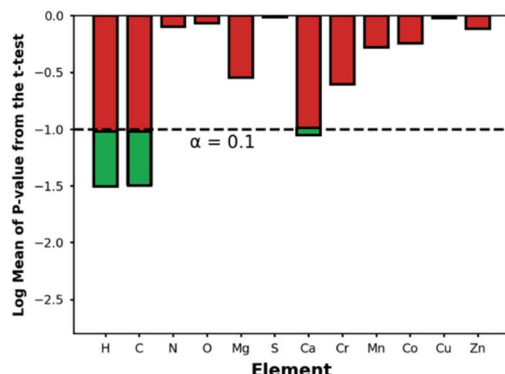


Fig. 9 Statistical significance for comparison of elemental compositions between the top-14 MOFs and the entire database based on the *p*-values derived from the *t*-test. The dashed line represents our chosen critical value for the one-sided *t*-test. Bars that fall below this threshold indicate elements that are statistically significantly more abundant in the top-performing MOFs. Ca is a metal that appears significantly more frequently in the top-14 MOFs than in the full database.

alone, as we will see below, does not guarantee the maximization of ΔN_{NH_3} .

4.5. Promising MOF designs

Contingent on adsorption simulation accuracy, now we proceed to present some promising MOF designs identified by our VBO run. The top-20 MOFs are listed in Table S4,† while the top-6 MOFs are presented in Fig. 10. Three of these MOFs correspond to hypothesized designs (top row), and the remaining

three correspond to extant designs that have been realized synthetically (bottom row). The free energy of the hypothesized designs in Fig. 10 was calculated using the Frenkel–Ladd method as discussed in earlier work,⁹⁹ resulting in free energies below 4.4 kJ mol⁻¹ per atom, which per discussion in ref. 100 suggests high synthesizability likelihood. The MOFs in Fig. 10 present ΔN_{NH_3} values in the 26.6–29.3 mmol_{NH₃} g_{MOF}⁻¹. Consistent with observed structure–performance relationships (Section 4.4), these MOFs exhibit APDs around 10 Å, void fractions around 0.7 and surface areas around 3900 m² g⁻¹. As for metals, note that although Ca was more abundant in the top-14 MOFs than in the whole database, the six best MOF designs featured Cr, Cu, Mn, Zn, and Co instead. Probably, because the textural properties of Ca MOFs were not “ideal.” This situation underscores the importance of optimizing a MOF design *both* structurally and chemically.

To put the predicted ΔN_{NH_3} for MOFs in Fig. 10 in the context of other MOFs experimentally tested in the literature, first let us reiterate that while NH₃ adsorption in MOFs have been consistently evaluated considering 300 K and 1 bar as the storage condition, such consistency has not existed for the release condition. Thus, a direct comparison is not possible. However, note that, with the exception of LiCl-MIL-53, the highest reported NH₃ loading at 300 K and 1 bar is 23.9 mmol_{NH₃} g_{MOF}⁻¹, so that even assuming total recovery at the release conditions, the predicted ΔN_{NH_3} for the MOFs in Fig. 10 is still higher. As for LiCl-MIL-53, its measured 33.9 mmol_{NH₃} g_{MOF}⁻¹ loading at 300 K and 1 bar is accompanied by a reported Q_{st} around 78 kJ mol⁻¹.⁵¹ Based on the relationship between heat of adsorption and percent NH₃ recovered (Fig. S6†)

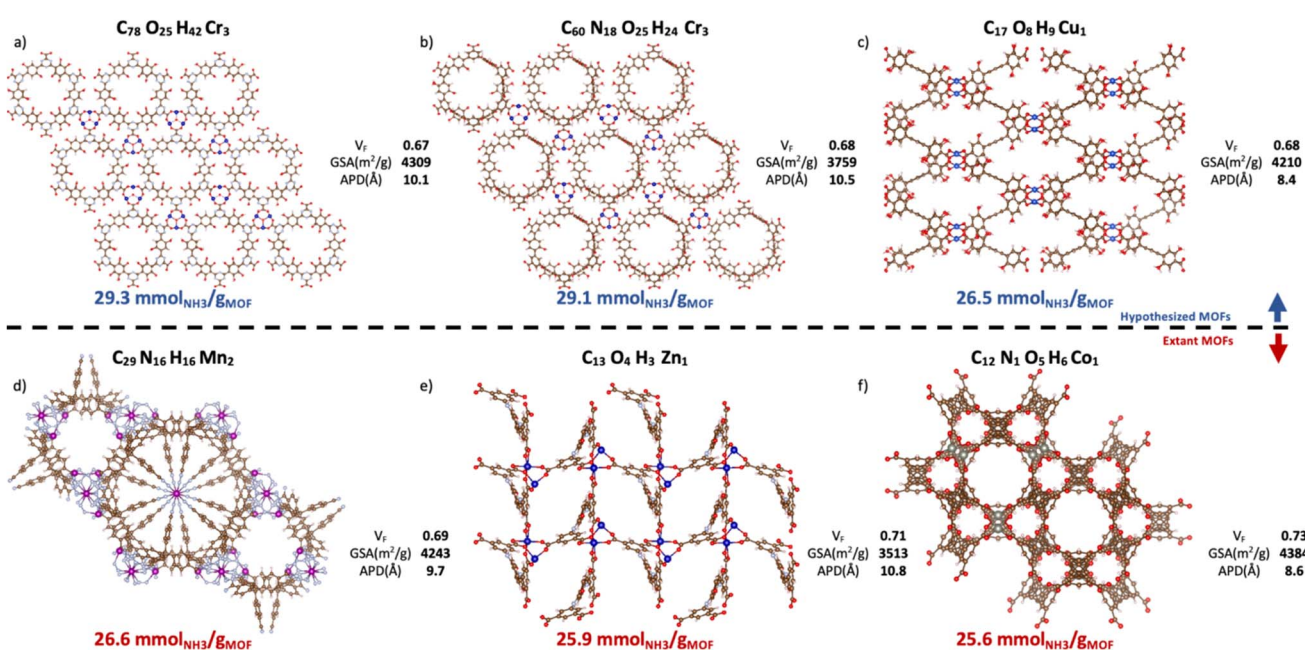


Fig. 10 Top-6 MOFs ranked by ΔN_{NH_3} value. Hypothesized (a–c) and extant MOFs (d–f) are in the top and bottom rows, respectively. V_f, GSA, and APD represent, respectively, the void fraction, gravimetric surface areas, and average pore diameter from pore size distribution. The CSD refcode and corresponding publication can be found in Table S5† with (a) *n* = 1, (b) *n* = 2, (c) *n* = 4, (d) *n* = 3, (e) *n* = 5, (f) *n* = 6, where *n* is the MOF ranking. The three hypothesized MOFs are potentially synthesizable per the free energy criterion by Anderson and Gómez-Gualdrón (Table S6†).¹⁰⁰ Ca MOFs appear in the top-14 but not in top-6 presumably due to suboptimal textural properties.

emerged in the study herein, a best case scenario for this Q_{st} (*i.e.*, 50% recovery) would yield a ΔN_{NH_3} around 16.9 mmol $_{\text{NH}_3}$ g $_{\text{MOF}}^{-1}$ for this MOF, which again is below the predicted ΔN_{NH_3} for the MOFs in Fig. 10.

Although here we focused on optimizing the MOF design to maximize ΔN_{NH_3} , other factors also play a role when using a MOF for a given application. Considering that we propose a thermal swing to release NH_3 , it is important to assess the thermal stability of the MOFs to encourage experimental testing. Accordingly, in Fig. 11a, we show the thermal decomposition temperature T_d of each of the top-20 MOFs (ΔN_{NH_3} ranging from 23 to 30 mmol $_{\text{NH}_3}$ g $_{\text{MOF}}^{-1}$), as predicted by an ANN model developed by Nandy *et al.*¹⁰¹ as available in the MOF-simplify website.¹⁰² This model makes the prediction based on the revised autocorrelation (RAC) descriptors of the MOFs, and was trained using reported thermogravimetric analysis (TGA) data for 3131 MOFs, with a mean absolute error (MAE) of 47 K. Considering this MAE and that the lowest predicted T_d was 466 K (which is 66 K higher than the upper temperature for the thermal swing), it seems that the suggested MOF designs are likely to withstand the proposed operation conditions.

Finally, to inform considerations about energy efficiency and economic viability, we estimated the energy required to release each kilogram of stored NH_3 with the proposed thermal swing, $\Delta Q_{\text{release}}$, using:

$$\Delta Q_{\text{release}} = Q_{\text{st}} + \left(\text{Cp}_{\text{NH}_3} + \frac{\text{Cp}_{\text{MOF}}}{\Delta N_{\text{NH}_3}} \right) \times \Delta T \quad (14)$$

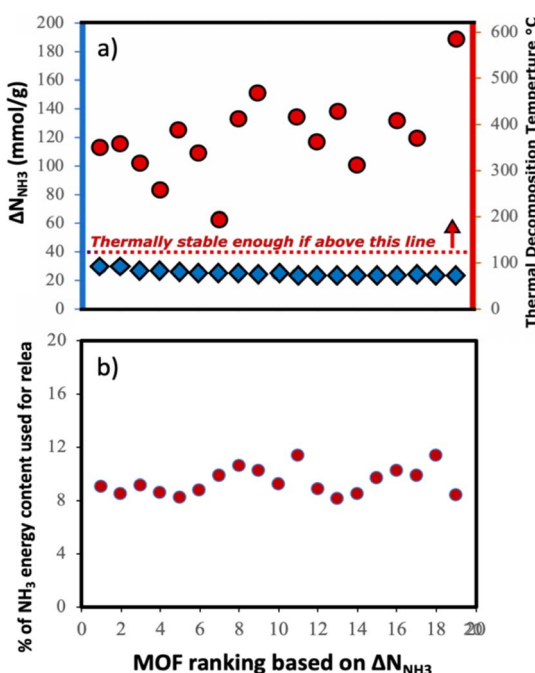


Fig. 11 (a) Thermal stability in top-20 MOFs from VBO campaign. Blue diamonds indicate ΔN_{NH_3} (left-axis) and red circles indicate predicted thermal decomposition temperature (right-axis). Top-20 MOFs appear likely to withstand operation conditions. (b) Estimated energy penalty to release stored NH_3 as percentage of the hydrogen-based energy content of NH_3 (22.5 MJ kg $_{\text{NH}_3}^{-1}$) in the top-20 MOFs. Penalty hovers around 8 to 12 percent on NH_3 energy content.

where, again, Q_{st} is the heat of adsorption of NH_3 , Cp_{NH_3} is the heat capacity of NH_3 in the gas phase (2.2 kJ kg $_{\text{NH}_3}^{-1}$ K $^{-1}$), Cp_{MOF} is the heat capacity of the MOF, and ΔT is the thermal swing magnitude (100 K). Eqn (14) is analogous to that used by Smit and coworkers to estimate the energy to release captured CO_2 from MOFs,²⁶ and essentially accounts for the heating of the MOF along with adsorbed NH_3 molecules from 300 K to 400 K, plus the energy needed to desorb NH_3 from the MOF at 400 K. For all MOFs, we used the average Cp_{MOF} value (0.87 kJ kg $_{\text{MOF}}^{-1}$ K $^{-1}$) previously reported for eleven MOFs,^{103,104} considering that this property seems to have low variability among MOFs.

Assuming that the energy stored in NH_3 corresponds to that of the H_2 that is released from NH_3 *via* cracking, the energy content of NH_3 is 22.5 MJ kg $_{\text{NH}_3}^{-1}$.¹⁰⁵ The latter implies that with the proposed MOF designs a penalty between 8% and 12% of the NH_3 energy content would be used to release the stored NH_3 . For context, an analogous calculation can be done to estimate energy penalty for liquid NH_3 storage, which can be estimated based on the latent heat of condensation for NH_3 (1.4 MJ kg $_{\text{NH}_3}^{-1}$)¹⁰⁶ and the energy to cool down NH_3 from 300 K down to 240 K. The above results in an estimated penalty of 7% of the NH_3 energy content. Considering that adsorptive NH_3 storage at ambient conditions can bypass other technological requirements such as insulation, toxicity, corrosion, or issues such as boil-off,^{43,45} among others, the operation conditions proposed herein for adsorptive NH_3 storage (and materials to achieve so) seem to merit reasonable consideration.

5. Conclusions

In this work, we developed a novel framework for efficiently finding a diverse set of optimal MOFs for applications involving ammonia adsorption. Our framework, called Vendi Bayesian Optimization (VBO), seamlessly combines traditional Bayesian optimization with the Vendi score, a diversity measure rooted in ecology and quantum mechanics. VBO is also made possible by the introduction of a novel similarity function in the space of MOFs that accounts for both chemistry and structure. We used this similarity function both for the GP used by Bayesian optimization and to compute the Vendi score. Our framework enabled the efficient discovery of several optimal MOFs that are distinct from one another, and that perform better than MOFs previously studied experimentally for NH_3 storage. Our analysis of the results of VBO highlights new design rules that MOF experimentalists can leverage to design optimal MOFs for the above application. We believe VBO introduces new useful capabilities for the efficient exploration of the combinatorially large MOF design space for the discovery of MOFs with desired properties. Importantly, our VBO framework is amenable to applications beyond ammonia adsorption. We leave the exploration of these applications as future work.

Data availability

Data for this article, including the code are available at vertaix/VBO at <https://github.com/vertaix/VBO>. Data sources are available at <https://wustl.box.com/s/3jkz8ksu9l3d1hqlkir4olainke9wc5t> and

Jupyter notebooks to reproduce our figures are available at <https://github.com/vertaix/VBO/tree/main/notebooks/Recreate%20figures>.

Author contributions

D. A. G.-G. and A. B. D. designed, supervised and acquired funding for the project. T.-W. L. and Q. N. conducted the work under D. A. G.-G. and A. B. D. supervision. T.-W. L. primarily focused on materials selection and characterization, data acquisition *via* molecular simulation, and analysis of simulation data in the context of target applications. Q. N. primarily focused on building the machine learning model and designing the search algorithm. D. A. G.-G. led the manuscript writing. All authors participated in discussions and intellectually contributed during the progress of the project. All authors edited and contributed to the editing of the manuscript and agreed on its final version.

Conflicts of interest

There are no conflicts to declare.

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