

Cite this: *Digital Discovery*, 2024, 3,
491

Image and data mining in reticular chemistry powered by GPT-4V†

Zhiling Zheng,¹ ^{abc} Zhiguo He,² ^{ab} Omar Khattab,³ ^d Nakul Rampal,⁴ ^{abc} Matei A. Zaharia,⁵ ^e Christian Borgs,⁶ ^{ce} Jennifer T. Chayes⁷ ^{ce} ^{fg} ^h and Omar M. Yaghi ¹ ^{ab} ^{cd} ^{ef} ^{gh} ^{ijkl}

The integration of artificial intelligence into scientific research opens new avenues with the advent of GPT-4V, a large language model equipped with vision capabilities. In this study, we demonstrate that GPT-4V, accessible through the ChatGPT web user interface or an API, offers promising possibilities in navigating and mining complex data for metal–organic frameworks (MOFs) especially from graphical sources (e.g. sorption isotherms, powder X-ray diffraction patterns, thermogravimetric analysis graphs, etc.). Our approach involved an automated process of converting 346 scholarly articles into 6240 images, which represents a benchmark dataset in this task, followed by deploying GPT-4V to categorize and analyze these images using natural language prompts, which can be written by chemists or materials scientists with minimal prior coding knowledge. This methodology enabled GPT-4V to accurately identify and interpret key plots integral to MOF characterization, such as nitrogen isotherms, PXRD patterns, and TGA curves, among others, with accuracy and recall above 93%. The model's proficiency in extracting critical information from these plots not only underscores its capability in data mining but also highlights its potential to aid in the digitalization of experimental data and the creation of datasets for reticular chemistry. In addition, the trends and values of nitrogen isotherm data from the selected literature allowed for a comparison between theoretical and experimental porosity values for over 200 compounds, highlighting certain discrepancies and underscoring the importance of integrating computational and experimental data. This work highlights the potential of AI in accelerating scientific discovery by bridging the gap between computational tools and experimental research.

Received 9th December 2023
Accepted 1st February 2024

DOI: 10.1039/d3dd00239j

rsc.li/digitaldiscovery

Introduction

The integration of artificial intelligence (AI) with the chemical sciences holds immense potential, and is accelerated by the rise of large language models (LLMs).^{1–7} These models have received substantial attention for the fact that they can be intuitively “programmed” or “taught” using daily conversational language, thereby assisting with diverse chemistry research tasks.^{8–20} It is envisioned that the evolution from text-only to more dynamic, multi-modal LLMs will result in even more powerful and convenient AI assistants across various applications.^{5,21–23}

The recent introduction of GPT-4V, with ‘V’ denoting its vision capability, stands as a testament to this progress.^{21,24} Trained on a vast and varied collection of multi-modal data, GPT-4V can process and respond to both textual and visual inputs.^{24–27} Its ability to interpret and analyze scientific literature, especially in identifying valuable data within graphical representations, makes it more attractive than traditional text-only models in natural language processing (NLP).^{10,28–30} These novel capabilities allow researchers from diverse backgrounds, including those with no specialized coding or computer vision expertise, to harness the power of GPT-4V through customized instructions applicable to various areas.^{19,31–33}

¹Department of Chemistry, University of California, Berkeley, California 94720, USA.
E-mail: yaghi@berkeley.edu

²Kavli Energy Nanoscience Institute, University of California, Berkeley, California 94720, USA

³Bakar Institute of Digital Materials for the Planet, University of California, Berkeley, California 94720, USA

⁴Department of Computer Science, Stanford University, Stanford, California 94305, USA

⁵Department of Electrical Engineering and Computer Sciences, University of California, Berkeley, California 94720, USA

⁶Department of Mathematics, University of California, Berkeley, California 94720, USA

⁷Department of Statistics, University of California, Berkeley, California 94720, USA

⁸School of Information, University of California, Berkeley, California 94720, USA

⁹KACST-UC Berkeley Center of Excellence for Nanomaterials for Clean Energy Applications, King Abdulaziz City for Science and Technology, Riyadh 11442, Saudi Arabia

† Electronic supplementary information (ESI) available: Full prompts designed to guide GPT-4V; additional examples showcasing GPT-4V's performance in reading various figure inputs and its corresponding responses; Python code used to automate the data mining and analysis processes; detailed information on the selected papers in this study, including the ground truth and the classification output for each page in a spread-sheet format; extracted nitrogen isotherms in this study. See DOI: <https://doi.org/10.1039/d3dd00239j>



the expanding role of AI in fostering innovation and discovery, which will further bridge the gap between advanced computational tools and cutting-edge chemical research.^{40,41}

Methods

Data preparation

For this study, a comprehensive dataset comprising 6240 pages from 346 scholarly articles published in 22 peer-reviewed journals among 5 different publishers discussing metal–organic frameworks (MOFs) was meticulously curated from the CoRE MOF Database (2019)⁴² to ensure the diversity of writing styles, article formatting and page layout. The selection criteria focused on excluding papers reporting structures with zero accessible surface areas and those without publicly available DOI numbers. In this case, the criteria resulted in the selection of 346 downloadable articles which met the specified requirements on the accessible surface area from the 9147 papers listed in the CoRE MOF database. The selected PDFs were transformed into PNG format utilizing a Python script developed with the assistance of ChatGPT (ESI Fig. S16†). This conversion process included 4423 pages from the main text and 1817 pages from the ESI of those chosen papers (ESI Data Set).†

Prompts and image mining considerations

We wrote instructions in natural language to guide GPT-4V for image classification and analysis. The primary objective was to enable GPT-4V to categorize each image with a label reflecting its content. These labels were designed to represent the most prevalent types of plots encountered in reticular chemistry, encompassing essential data and values. Key plots include the nitrogen adsorption–desorption isotherm (indicating porosity), the PXRD pattern (illustrating crystallinity), the TGA curve (demonstrating thermal stability), crystallographic structure rendering images (depicting crystal structure or topology), and other gas sorption isotherms (relevant for applications and gas uptake). Accordingly, these were established as label choices 1 through 6. It is noteworthy that a single page may contain multiple plots (*e.g.*, PXRD and TGA concurrently), which requires GPT-4V's capability to perform an identification of multiple formats of plots. Therefore, the primary task assigned to GPT-4V involves identifying these specified elements within the image of the whole page, complemented by a structured response template. Furthermore, the crafting of prompts adhered to three guiding principles outlined in our previous publication:¹⁰ (i) minimizing hallucination, (ii) implementing detailed instructions, and (iii) ensuring structured output. The complete prompt is available in ESI Fig. S15.†

Automation in paper reading

With the prompt finalized, GPT-4V's task was to “read” through all the pages from selected papers. This was accomplished by sequentially presenting each image of the whole page, along with the designated prompt, to the model and collecting its responses. This iterative process was automated using a for-loop structure *via* simple python code generated with the

assistance of ChatGPT (ESI Fig. S17†). In particular, two options exist, and both were tested in this study: (i) interfacing with the web-based chatbot (ChatGPT powered by GPT-4V, version dated September 25, 2023) and (ii) utilizing an API to connect with the *gpt-4-1106-vision-preview* model. Both methods applied the same underlying base model and were facilitated through Python scripts capable of operating autonomously (Fig. S17 and S18†). Additionally, the process for parsing and extracting the labels from the responses given by GPT-4V was automated by Python script (Fig. S19 and S20†).

Results and discussion

Initial assessment of GPT-4V's knowledge of reticular chemistry

At the outset of this study, we rigorously evaluated the proficiency of GPT-4V in recognizing and interpreting a range of figures typically found in reticular chemistry literature by asking GPT-4V to describe the respective figure sent along with the chat (Fig. S1–S14†). This knowledge assessment encompassed various physical characterization plots including nitrogen isotherms, PXRD patterns, TGA curves, NMR and IR spectra, as well as illustrative plots such as scatter plots, bar plots, and 2D or 3D molecular structures. In addition, synthesis schemes and real experimental images, like microscope and SEM images, were also analyzed. Each figure was presented to GPT-4V alongside a brief prompt requesting a description of the input figure in a conversational format.

The responses from GPT-4V (ESI Fig. S1–S14†) demonstrated its remarkable capability to not only categorize these images accurately but also to elaborate on specific details, including notations, axis ranges, color coding and shape of symbols and lines, labels, legends, and to draw inferences from the provided information in the figure caption. This advanced level of contextual data interpretation and holistic analysis underscores the suitability of GPT-4V as a potent AI assistant for image and data mining in scientific literature.

Designing prompts for page content labelling

The core objective of this study was to investigate whether GPT-4V could automatically navigate through scientific papers, identifying specific information and aggregating it into a comprehensive dataset for further analysis. Our interest specifically centered on plots pivotal in the physical characterization of MOFs. These plots, namely nitrogen isotherms, PXRD patterns, TGA curves, crystal structure or topology illustrations, and other gas sorption isotherms, are instrumental in deducing key properties of chemical compounds, including permanent porosity, crystallinity, thermal stability, connectivity (topology), and sorbent selectivity towards gases. Successfully extracting and consolidating data from these plots amongst a vast volume of literature holds immense potential for advancing our understanding of structure–property relationships and in accelerating the discovery of novel compounds.^{8,34,37,38,43–48}

To this end, we designed a specific and extensive prompt for GPT-4V, targeting the aforementioned categories (Fig. 1 and ESI



Fig. S15†). Unlike the brief prompts employed in the initial assessment of GPT-4V's knowledge in reticular chemistry (Fig. S1–S14†), the formal prompt used for image mining in this study was developed by employing the prompt engineering strategies that we previously reported,^{10,14} making it more extensive and comprehensive (see Methods section and ESI Fig. S15†). It is noteworthy that the prompt allowed for the possibility of multiple selections on a single page, as it is common for various plots to coexist in scientific literature. Additionally, GPT-4V was instructed to indicate the absence of these five categories when applicable. Therefore, in total six choices were provided for GPT-4V (Fig. 1). The development of these prompts was guided by general principles for prompt engineering in text mining.¹⁰

Performance evaluation of GPT-4V

In our workflow, each page of the selected literature was digitized into an image and then analyzed by GPT-4V (see Methods section). This process involved combining each image with a textual prompt, after which GPT-4V's responses were collected. Particularly, GPT-4V was “programmed” by instructions using human daily conversational language in the prompt to guide it to follow a specific response format (ESI Fig. S15†), allowing for the automatic labeling of each page based on its content. While copyright restrictions preclude the sharing of the actual images used in this study, we provide four representative examples that closely simulate the layout and content of the analyzed pages from actual published literature^{49–53} to illustrate this figure content identification process (Fig. 1). These examples demonstrate GPT-4V's ability to accurately recognize and label the desired plots on each page, regardless of the complexity of the information shown. It is important to note that in the actual evaluation, instead of using the four demo images shown in Fig. 1, we used 6240 single-page images, akin to “screenshots” that contain all the text and associated figures, if any, in that page as a whole image, from 346 articles. Consequently, the respective 6240 output answers from GPT-4V were collected (ESI Data Set†). In other words, during each interaction, each page was submitted separately in a new instance of conversation to yield the classification result of the given page. Essentially, GPT-4V could perform iterative classifications of each page, utilizing the same prompt but varying image content; this process was completed without human intervention (ESI Fig. S17 and S18†).

The evaluation of GPT-4V's classification accuracy involved comparing its responses against a benchmarking ground truth dataset. This dataset was meticulously created by experts in reticular chemistry, who manually reviewed and labeled all 6240 images for the presence of specific content (choices 1 to 6). Performance metrics for each category were calculated individually, considering the multiple-choice nature of the task (ESI Data Set and Table S1†). For example, if the ground truth for an image was labeled as “1, 3, 4” and GPT-4V's response was “1, 4, 5,” the classifications for each category were assessed as follows: nitrogen isotherm (Choice 1) received a True Positive, PXRD pattern (Choice 2) a True Negative, TGA curve (Choice 3) a False

Negative, crystal category (Choice 4) a True Positive, other gas sorption isotherm (Choice 5) a False Positive, and “none of the above” (Choice 6) a True Negative.

The summarized metrics, including accuracy, precision, recall and F1 score, are presented in Table 1. Notably, it displays promising accuracy rates above 94% for all categories. This indicates that GPT-4V can correctly classify a high percentage of figure content across all 6 categories, which encompass 5 different plots types and 1 NOTA category. Precision ranges from 88% to 97% for five of the plot types; however, it is notably lower at 61% for the “other gas sorption isotherm” category. This reduced precision is attributed to the category's broad scope and occasional mislabeling of IR and NMR spectra as such. This highlights an opportunity for further refinement in the content of prompts, aiming to enhance GPT-4V's understanding of the definition of gas sorption isotherm and ensuring it does not mistakenly identify other spectra as isotherm plots. The recall values, between 94% and 99% for all six classes, suggest the model is effective at capturing a high percentage of figure content relevant to each specific class. The F1 scores range from 93% to 95%, demonstrating a well-balanced and robust performance in both precision and recall. However, the “other gas sorption isotherm” is an exception with a 76% F1 score, which is foreseen due to its low recall performance, indicating a certain limitation of the model in this specific class, possibly due to insufficiently detailed instruction in the prompt.

Additionally, similar accuracy rates were observed in GPT-4V's performance *via* both the web user interface (WUI) and API (Table S2†), a testament to the uniformity of the underlying base model (*gpt-4-1106-vision-preview*). Here we acknowledge that the base model is still in preview version, yet the obtained results are very promising. The automated process for reading papers and collecting output classifications through both the WUI and API offers diverse operational choices with consistently high performance. We note that in this study, no fine-tuning strategies were applied, and the performance primarily resulted from carefully crafted text-based instructions after prompt engineering. This approach is becoming increasingly important in the realm of scientific research involving LLMs.^{32,54–57} Impressively, we found that this also holds true for multimodal models handling both text and graphical inputs such as GPT-4V in this study.

Furthermore, the relation between the ground truth and GPT-4V's classification output was visualized in a confusion

Table 1 Summary of GPT-4V performance metrics in classifying different scientific plot types seen in reticular chemistry literature. The definitions of the performance metrics are given in ESI Table S1

Plot type	Accuracy	Precision	Recall	F1 score
Nitrogen isotherm	99.5%	90%	96.7%	93.5%
Power X-ray diffraction	99.2%	94.3%	98.4%	96.3%
Thermogravimetric analysis	99.2%	87.8%	99.3%	93.2%
Crystal structure or topology	98.1%	93.2%	97.1%	95.1%
Other gas sorption isotherm	95.0%	61.4%	99.5%	76.0%
None of the above	94.3%	96.7%	93.7%	95.1%



Actual Choice	Predicted Choice					
	NI	PXRD	TGA	CST	OI	NOTA
NI	232	2	1	3	1	15
PXRD	1	630	0	8	0	20
TGA	0	2	331	4	1	21
CST	1	7	0	1173	2	81
OI	6	3	2	14	495	145
NOTA	1	2	0	19	0	3537

Fig. 2 Confusion matrix displaying the performance of GPT-4V in accurately predicting scientific plot types from reticular chemistry literature. The matrix compares actual choices (ground truth), including Nitrogen Isotherm (NI), Powder X-Ray Diffraction (PXRD), Thermogravimetric Analysis (TGA), Crystal Structure or Topology (CST), Other Gas Sorption Isotherm (OI), and None of the Above (NOTA), against the predicted choices made by the GPT-4V in its outputs. The numbers in the diagonal represent the number of each type of plot that was correctly identified by GPT-4V. The color gradient represents the frequency of predictions, with darker shades indicating higher occurrences.

matrix (Fig. 2). Among the 6240 pages analyzed, GPT-4V identified approximately 232 plots that were with nitrogen isotherms, 630 that were PXRD patterns, and 331 that were TGA traces. It suggests a substantial volume of data within the analyzed literature. Moreover, 3537 pages were classified as lacking the plots of interest, which can be a strategy to be applied in the future to help human researchers streamline the review process for a specific type of literature plot by excluding less important pages and focusing only on the rest of the content-rich pages. We also estimated the time of this automated literature processing approach (ESI Tables S3 and S4†). On average, processing a 20-page paper took approximately 2 minutes at a rate of 5.5 seconds per image *via* API and this process can be parallelized (ESI Table S3†).

GPT-4V's interpretation of nitrogen isotherm data

Building upon the successful labeling of page contents, we next directed our focus towards utilizing GPT-4V for the detailed interpretation and analysis of pages featuring nitrogen isotherm plots. To achieve this, we refined our prompt strategy, incorporating additional and specific verbal instructions (ESI Fig. S15†). These enhanced prompts guided GPT-4V to not only recognize nitrogen isotherms but also to extract and report key descriptors from each plot. These descriptors included the figure number, compound name, surface area or pore volume value reported by the author, the presence of hysteresis in the

adsorption-desorption curve, the saturation plateaus of the isotherm, and an estimation of a bounding box encompassing the figure (Fig. 3). A critical aspect of this approach, aiming to minimize hallucinations, was the instruction for GPT-4V to strictly utilize the information available on the page image, defaulting to answer “N/A” for any unobtainable or ambiguous data. It turned out that GPT-4V demonstrated a remarkable ability to extract these details by analyzing the isotherm, its axes, legends, and accompanying text content.

To validate the accuracy of GPT-4V's nitrogen isotherm analysis, we manually reviewed over 200 responses from pages containing nitrogen isotherms in our selected papers (ESI† data set). This evaluation was conducted for each of the six descriptors independently. The results indicated high accuracy levels for the figure number (96.67%), compound name (90.42%), and porosity analysis (98.33%). We hypothesize that GPT-4V's image processing capabilities, potentially incorporating optical character recognition (OCR) tools, played a pivotal role in these tasks. Given GPT-4V's proficiency with text, it likely excelled in tasks where textual information was directly “readable” from the image. Conversely, the other three descriptors, namely hysteresis presence, saturation plateaus, and bounding box estimation, showed generally satisfactory performance, ranging between 76.25% and 84.58%. These tasks, inherently more challenging and nuanced, required a comprehensive analysis of all image elements.⁵⁸ Nonetheless, the overall performance was impressive, especially considering the simplicity with which researchers could instruct GPT-4V using natural language.

Accelerating the creation of digital datasets in reticular chemistry

Our study further demonstrates the potential of GPT-4V in accelerating the digitalization of experimental data for reticular compounds. This is particularly evident from the sorption measurement outcomes gleaned from community-published literature. Once pages featuring nitrogen isotherm plots are identified, each corresponding curve, typically presented in a non-digital format (*e.g.*, scanned or plotted images), can be meticulously extracted using data extraction tools like Web-PlotDigitizer.⁵⁹ The extracted data points are then systematically compiled and stored (ESI Data Set†). While we currently make all mined sorption data available in a spreadsheet dataset format, it is feasible that digitalized data can be converted into AIF files as a first step toward building the MOF sorption data library.⁶⁰ Fig. 4a presents a collection of nitrogen isotherm data points manually extracted from the qualified pages in our study as a proof-of-concept, showcasing a diverse array of isotherm types and porosity characteristics. Leveraging the CoRE MOF Database,^{36,42} which provides computational results for compounds discussed in these papers, we matched each adsorption-desorption curve's corresponding compound with their CCDC number, accessible surface area, and pore volume. This enabled us to visualize and compare the experimental porosity (indicated by the nitrogen isotherm curves) with the calculated values (derived from the CoRE MOF Database). The



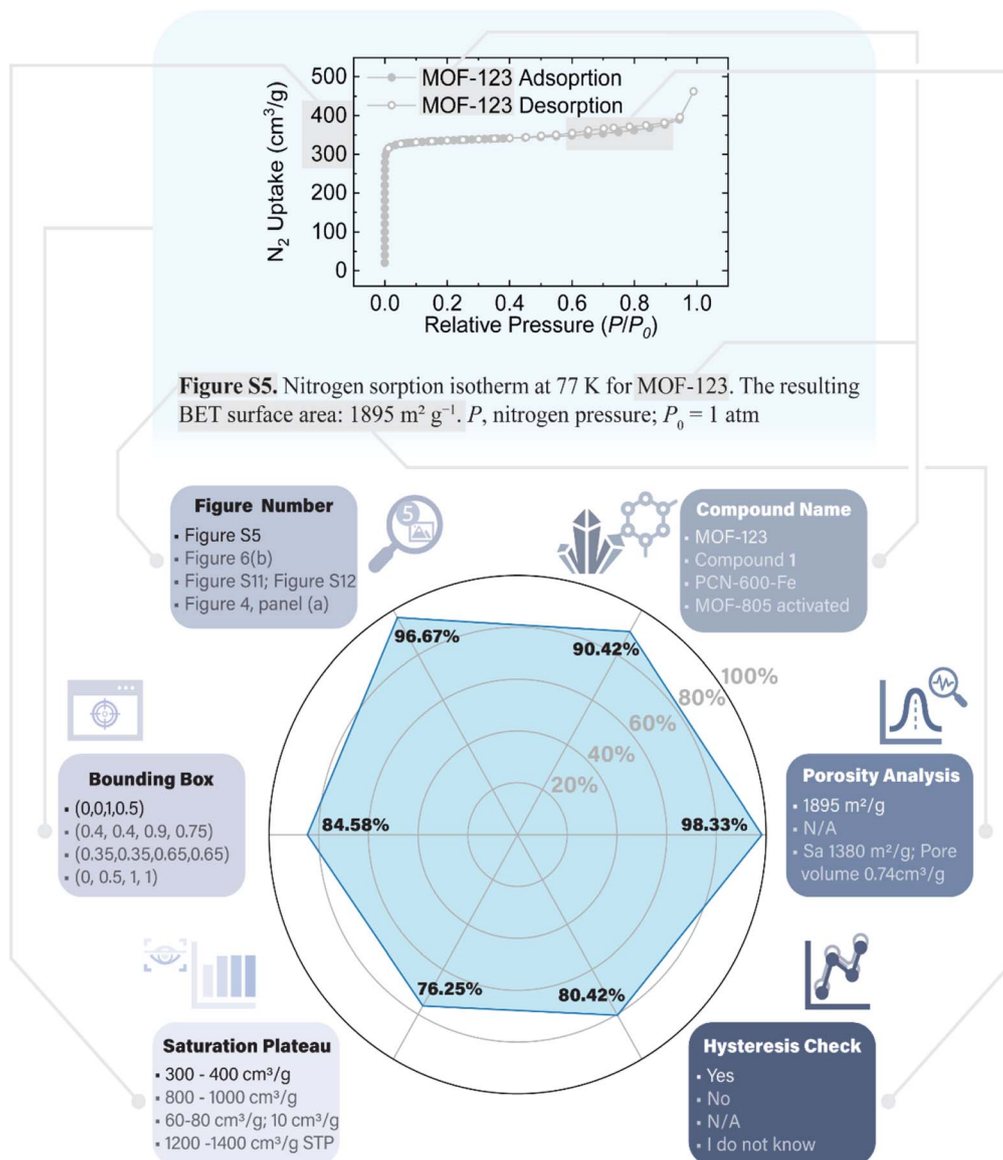


Fig. 3 The upper section presents an exemplary nitrogen isotherm graph, which is styled in a manner consistent with standard scientific publications and has a figure caption. The lower section features a radar chart detailing the accuracy of GPT-4V's performance in extracting and interpreting critical information, such as figure number, compound name, porosity values, hysteresis presence, saturation uptake, and bounding box parameters. Each vertex on the chart corresponds to the accuracy achieved in its associated category, illustrating the advanced image input capability of GPT-4V in processing and analysing graphical data from nitrogen isotherms. The grey lines demonstrate textual and visual cues, such as colour and shape, that guide GPT-4V in identifying specific information, with sample outcomes for each category displayed within each box.

results, illustrated in scatter plots (Fig. 4b and c), represent each compound as a data point, where the x -axis denotes the theoretical values calculated from CCDC CIF structures,^{36,42} and the y -axis represents the experimentally reported surface area or pore volume inferred from the extracted isotherm obtained from the graphs identified by GPT-4V. The distributions of x and y values of the data points are shown in the respective bar charts. It is crucial to note that our focus was more on the general trends across MOF compounds selected for this study rather than on pinpointing the exact surface area and porosity for each compound, as variations may arise due to differences

in calculation models and assumptions, and the re-digitalized data points at very low relative pressures may decrease the accuracy of the calculation.⁶⁰⁻⁶⁵

Interestingly, despite the use of experimentally determined crystal structures in the CoRE MOF database,^{36,42} discrepancies between theoretical predictions and actual experimental outcomes were observed. For instance, some compounds exhibited a high theoretical porosity based on their refined CCDC structures but failed to demonstrate such porosity experimentally, possibly due to factors like structural collapse during activation, inaccessible pore environments, or



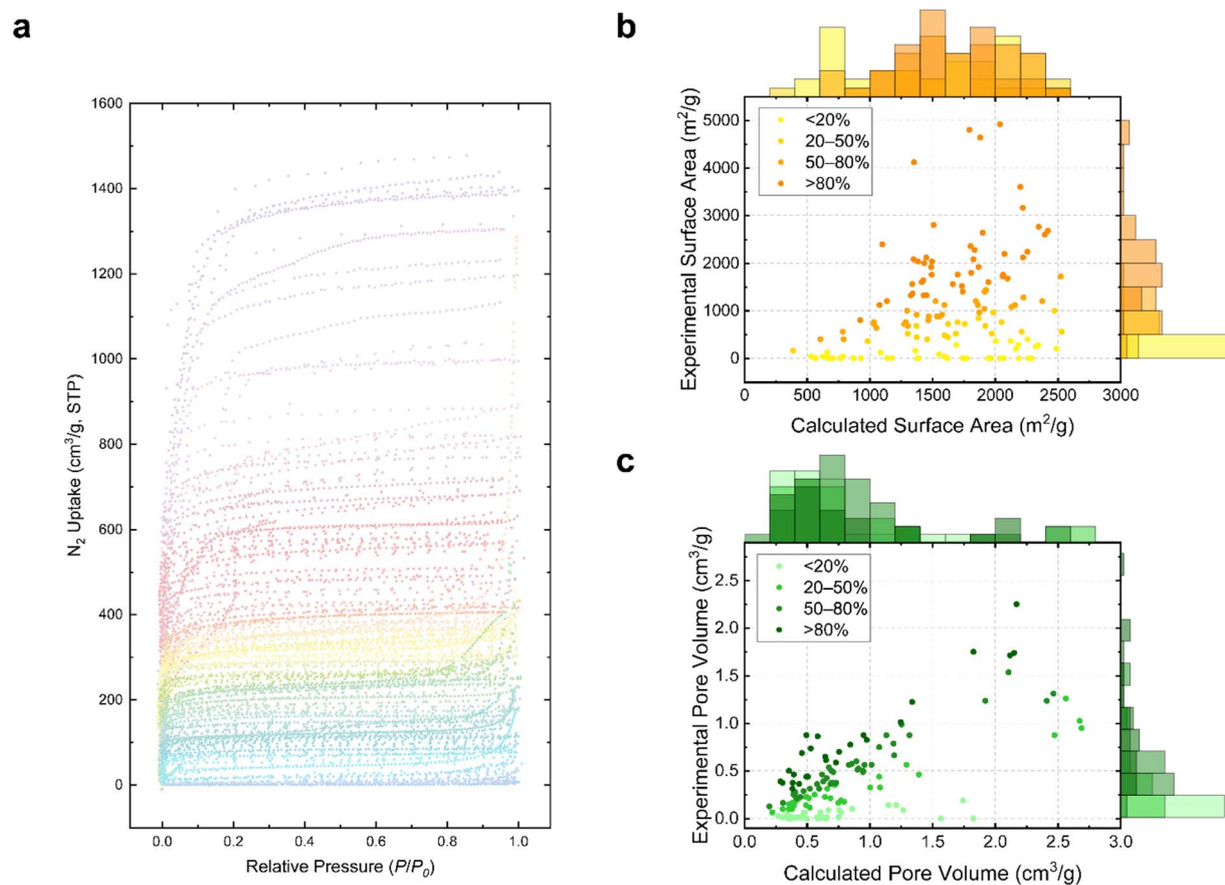


Fig. 4 Comprehensive analysis of nitrogen isotherm data extracted from selected reticular chemistry literature. (a) An overlay scatter plot of distinct adsorption branches for nitrogen isotherms from 346 selected papers, exhibiting a range of uptake behaviors. The data points were picked by WebPlotDigitizer using Automatic Extraction mode while the axes were aligned manually. The desorption branches are omitted for clarity. (b) A plot correlating experimental and calculated surface area values, where each point represents one compound's data, linked via the DOI number from the literature to the corresponding CCDC crystal structure in the CoRE MOF database. (c) A correlational plot for pore volume, again with each point representing a single compound, illustrating the relationship between experimentally measured values and those calculated based on crystallographic data. The color-coding in (b) and (c) indicates the percentage of experimental value reaching the theoretical value. The horizontal bars along the top sides of the plots in (b) and (c) represent the histogram of the distribution of the calculated porosity indicators, namely the surface area and the pore volume, reported in the CoRE MOF database. Similarly, the vertical bars along the right sides of the two plots display the distribution of the experimentally determined porosity values re-digitized from the selected papers.

suboptimal synthesis conditions. While variations of surface area, porosity or measured isotherms may exist among different research groups and preparations,⁶⁶ such a phenomenon does not necessarily indicate a reproducibility issue, but rather a failure to reach the full potential and optimal conditions of the MOF in those studies. As a demonstration, Table 2 presents eight representative examples illustrating these variances, with some compounds showing excellent agreement between theoretical and experimental values (*e.g.*, TAKCAM, OTIHOQ), while others displayed certain deviations (*e.g.*, BOHXED, TAKTAD, and TOCJAY).

Our findings reveal that reliance solely on computational results for material selection can be sometimes misleading, as many compounds exhibit experimental performances that deviate substantially from theoretical predictions, even when based on experimentally determined structures (Fig. 4). It should be noted that these experimentally-determined non-porous compounds are not necessarily useless, they in fact

serve as valuable negative data points in the mined nitrogen isotherm dataset. By acknowledging these limitations and combining computational methods with experimental data, researchers can gain more comprehensive insights into reticular chemistry, discern trends, and make informed predictions.^{75–77} We envision that leveraging GPT-4V's capabilities to search for more experimental data—not limited to nitrogen isotherm but extending to other isotherms like water, CO₂ capture, methane sorption—and other critical plots like TGA curves and PXRD from literature, in tandem with theoretical insights and computational science, can significantly advance the discovery and development of high-performance reticular compounds with desirable properties and functionalities.

To achieve this, it should be acknowledged that one limitation of our workflow is that the efficiency of using GPT-4V hinges significantly on the skillful crafting of prompts.^{10,58} It has always been a challenge to write good instructions to guide



Table 2 Calculated and experimental accessible surface areas for selected compounds in reticular chemistry literature

Representative compound	Metal	CCDC code	Surface area (m ² g ⁻¹)			Literature ^c	Reference
			CoRE MOF database ^a	Re-digitized experimental sorption curves ^b			
ZJU-70a	Cu	DORNAB	2520	1790		1791	67
MOF-802	Zr	BOHXED	2070	0		<20	68
Hf-L7	Hf	TAKCAM	2070	2200		2270	69
MFM-300(Ga ₂)	Ga	TAKTAD	1600	480		400	70
MIL-100 (Al)	Al	BUSPIP	1350	2090		2152	71
[Cd(L4)]·1.5DMF	Cd	OTIHOQ	1330	1320		1376	72
Bio-MOF-101	Zn	TOCJAY	1100	2400		4410	73
{[Ni ₆ (N ₃) ₁₂ L ₆](H ₂ O) ₁₃ } _∞	Ni	NUZCER	790	400		309	74

^a Based on the accessible surface area calculated for each compound's crystal structure as reported in the CoRE MOF 2019 database⁴² using nitrogen as the probe. ^b Based on data extracted from the experimental nitrogen isotherm reported in the respective literature, results are rounded to the nearest ten. ^c Based on the surface area reported by the author in the literature.

LLMs.⁷⁸ Precision in prompt design is essential to achieve specific and accurate results in desirable format, underscoring the need for clear, detailed, and well-articulated instructions. In addition, idiosyncratic and subtle prompt cues can influence the LLMs without changing the underlying meaning, and consequently sometimes equivalent-looking prompts having very different quality.^{79,80} In this context, the emerging tool DSPy (Declarative Self-Improving Python)⁸¹ represents a significant advancement as compared with human written prompts and a future direction for this study. DSPy streamlines the process of prompt creation, combining techniques for both prompting and fine-tuning language models. Applying DSPy for prompt optimization necessitates the existence of a relatively small development set and some data annotation that DSPy will use to conduct the optimization automatically. Conceptually, this innovative system could enable researchers to commence their inquiries with a basic scaffolding of the high-level steps of the design and use DSPy to automate the instruction of LLMs, which can be considered a solution to further strengthen our workflow and improve the performance of LLMs. In the near future, we envision this combined approach can further increase the accuracy of the tasks demonstrated in this study, expand scope of the type of data to be mined from the literature, and hold the promise of making advanced natural language processing tools more accessible and adaptable to diverse research scenarios.

Concluding remarks

This study demonstrated the application of GPT-4V in text, image, and data mining within the realm of reticular chemistry. Our findings reveal that GPT-4V, when paired with carefully constructed prompts, is capable of processing page images and accurately identifying content in a fast and accurate manner. This capability can further extend to isolating pages with specific plots for in-depth analysis and building up datasets for experimental measurements from the literature. Importantly, the key features of LLMs like GPT-4V—being “programmable” through everyday natural language and possessing pre-trained domain knowledge—reduce traditional barriers associated with coding expertise and model training for specific plot or

figure recognition. This adaptability is highlighted by the ease of transitioning from analysing one type of data, such as TGA curves, to another, like water isotherms, simply by modifying the instructional prompt. We envision that this workflow can be transferable and move beyond reticular chemistry to a broader spectrum of scientific disciplines. Nevertheless, there are certain limitations to consider. For instance, the model exhibits reduced accuracy in more complex tasks like the interpretation of graphs, indicating a need for further refinement in prompt engineering strategies. The lack of benchmarking datasets may increase the difficulty in coming up with effective prompts to guide LLMs. On the other hand, we envision that to make the instruction on GPT-4V more effective, the integration of advanced platform like DSPy into our workflow in the future will open new avenues in scientific data mining and points to a future where AI becomes an integral, user-friendly tool in the advancement of scientific knowledge.

Data availability

The code, analysis scripts, and datasets supporting this article have been uploaded as part of the ESI.†

Author contributions

Z. Z. and O. M. Y conceived and designed the study. Z. Z. and Z. H. performed the data curation. Z. Z. conducted the investigation and developed the methodology. Z. Z. and O. M. Y. carried out the formal analysis and visualization. O. M. Y. supervised the project. Z. Z. wrote the initial draft of the paper. All authors contributed significantly and collaboratively to the data analysis, and the review and editing of the manuscript.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This material is based upon work supported by the Defense Advanced Research Projects Agency (DARPA) under contract



HR0011-21-C-0020. Any opinions, findings, and conclusions or recommendations expressed in this material are those of the authors and do not necessarily reflect the views of DARPA. Z. Z. expresses gratitude to Mr Kefan Dong (Stanford University), Mr Jiayi Weng (OpenAI), Ms. Oufan Zhang (UC Berkeley), Mr Zichao Rong (UC Berkeley) and Ms. Zeqi Gu (Cornell University) for their valuable discussions. Z. Z. also acknowledges financial support from a Kavli ENSI Graduate Student Fellowship.

Notes and references

- 1 A. Birhane, A. Kasirzadeh, D. Leslie and S. Wachter, *Nat. Rev. Phys.*, 2023, 5, 277–280.
- 2 A. D. White, *Nat. Rev. Chem.*, 2023, 7, 457–458.
- 3 S. Bubeck, V. Chandrasekaran, R. Eldan, J. Gehrke, E. Horvitz, E. Kamar, P. Lee, Y. T. Lee, Y. Li and S. Lundberg, Sparks of artificial general intelligence: Early experiments with gpt-4, *arXiv*, 2023, preprint, arXiv:2303.12712, DOI: [10.48550/arXiv:2303.12712](https://doi.org/10.48550/arXiv:2303.12712).
- 4 Microsoft Research AI4Science, Microsoft Azure Quantum, The Impact of Large Language Models on Scientific Discovery: a Preliminary Study using GPT-4, *arXiv*, 2023, preprint, arXiv:2311.07361, DOI: [10.48550/arXiv.2311.07361](https://doi.org/10.48550/arXiv.2311.07361).
- 5 OpenAI, GPT-4 technical report, *arXiv*, 2023, preprint, arXiv:2303.08774v3, DOI: [10.48550/arXiv:2303.08774v3](https://doi.org/10.48550/arXiv:2303.08774v3).
- 6 R. Anil, A. M. Dai, O. Firat, M. Johnson, D. Lepikhin, A. Passos, S. Shakeri, E. Taropa, P. Bailey and Z. Chen, Palm 2 technical report, *arXiv*, 2023, preprint, arXiv:2305.10403, DOI: [10.48550/arXiv.2305.10403](https://doi.org/10.48550/arXiv.2305.10403).
- 7 H. Touvron, L. Martin, K. Stone, P. Albert, A. Almahairi, Y. Babaei, N. Bashlykov, S. Batra, P. Bhargava and S. Bhosale, Llama 2: Open foundation and fine-tuned chat models, *arXiv*, 2023, preprint, arXiv:2307.09288, DOI: [10.48550/arXiv.2307.09288](https://doi.org/10.48550/arXiv.2307.09288).
- 8 Z. Zheng, O. Zhang, H. L. Nguyen, N. Rampal, A. H. Alawadhi, Z. Rong, T. Head-Gordon, C. Borgs, J. T. Chayes and O. M. Yaghi, *ACS Cent. Sci.*, 2023, 9, 2161–2170.
- 9 K. M. Jablonka, Q. Ai, A. Al-Feghali, S. Badhwar, J. D. Bocarsly, A. M. Bran, S. Bringuier, L. C. Brinson, K. Choudhary and D. Circi, *Digital Discovery*, 2023, 2, 1233–1250.
- 10 Z. Zheng, O. Zhang, C. Borgs, J. T. Chayes and O. M. Yaghi, *J. Am. Chem. Soc.*, 2023, 145, 18048–18062.
- 11 Y. Kang and J. Kim, ChatMOF: An Autonomous AI System for Predicting and Generating Metal-Organic Frameworks, *arXiv*, 2023, preprint, arXiv:2308.01423, DOI: [10.48550/arXiv:2308.01423](https://doi.org/10.48550/arXiv:2308.01423).
- 12 S. Liu, J. Wang, Y. Yang, C. Wang, L. Liu, H. Guo and C. Xiao, ChatGPT-powered Conversational Drug Editing Using Retrieval and Domain Feedback, *arXiv*, 2023, preprint, arXiv:2305.18090, DOI: [10.48550/arXiv:2305.18090](https://doi.org/10.48550/arXiv:2305.18090).
- 13 A. M. Bran, S. Cox, A. D. White and P. Schwaller, ChemCrow: Augmenting large-language models with chemistry tools, *arXiv*, 2023, preprint, arXiv:2304.05376, DOI: [10.48550/arXiv:2304.05376](https://doi.org/10.48550/arXiv:2304.05376).
- 14 Z. Zheng, Z. Rong, N. Rampal, C. Borgs, J. T. Chayes and O. M. Yaghi, *Angew. Chem., Int. Ed.*, 2023, 62, e202311983.
- 15 M. Thway, A. Low, S. Khetan, H. Dai, J. Recatala-Gomez, A. P. Chen and K. Hippalgaonkar, *Digital Discovery*, 2024, DOI: [10.1039/D3DD00020K](https://doi.org/10.1039/D3DD00020K).
- 16 Z. Zheng, A. H. Alawadhi, S. Chheda, S. E. Neumann, N. Rampal, S. Liu, H. L. Nguyen, Y.-h. Lin, Z. Rong, J. I. Siepmann, L. Gagliardi, A. Anandkumar, C. Borgs, J. T. Chayes and O. M. Yaghi, *J. Am. Chem. Soc.*, 2023, 145, 28284–28295.
- 17 G. M. Hocky and A. D. White, *Digital Discovery*, 2022, 1, 79–83.
- 18 Z. Xie, X. Evangelopoulos, Ö. H. Omar, A. Troisi, A. I. Cooper and L. Chen, *Chem. Sci.*, 2024, 15, 500–510.
- 19 M. Suvarna, A. C. Vaucher, S. Mitchell, T. Laino and J. Pérez-Ramírez, *Nat. Commun.*, 2023, 14, 7964.
- 20 K. Cruse, V. Baibakova, M. Abdelsamie, K. Hong, C. J. Bartel, A. Trewartha, A. Jain, C. M. Sutter-Fella and G. Ceder, *Chem. Mater.*, 2024, 36(2), 772–785.
- 21 Z. Yang, L. Li, K. Lin, J. Wang, C.-C. Lin, Z. Liu and L. Wang, The dawn of Imms: Preliminary explorations with gpt-4v (ision), *arXiv*, 2023, preprint, arXiv:2309.17421, DOI: [10.48550/arXiv.2309.17421](https://doi.org/10.48550/arXiv.2309.17421).
- 22 Z. Yan, K. Zhang, R. Zhou, L. He, X. Li and L. Sun, Multimodal ChatGPT for Medical Applications: an Experimental Study of GPT-4V, *arXiv*, 2023, preprint, arXiv:2310.19061, DOI: [10.48550/arXiv.2310.19061](https://doi.org/10.48550/arXiv.2310.19061).
- 23 C. Wu, S. Yin, W. Qi, X. Wang, Z. Tang and N. Duan, Visual chatgpt: Talking, drawing and editing with visual foundation models, *arXiv*, 2023, preprint, arXiv:2303.04671, DOI: [10.48550/arXiv.2303.04671](https://doi.org/10.48550/arXiv.2303.04671).
- 24 OpenAI, *GPT-4V(ision) System Card*, 2023, accessed 2023-09-25.
- 25 C. Wu, J. Lei, Q. Zheng, W. Zhao, W. Lin, X. Zhang, X. Zhou, Z. Zhao, Y. Zhang and Y. Wang, Can gpt-4v (ision) serve medical applications? case studies on gpt-4v for multimodal medical diagnosis, *arXiv*, 2023, preprint, arXiv:2310.09909, DOI: [10.48550/arXiv:2310.09909](https://doi.org/10.48550/arXiv:2310.09909).
- 26 N. Wake, A. Kanehira, K. Sasabuchi, J. Takamatsu and K. Ikeuchi, GPT-4V (ision) for Robotics: Multimodal Task Planning from Human Demonstration, *arXiv*, 2023, preprint, arXiv:2311.12015, DOI: [10.48550/arXiv:2311.12015](https://doi.org/10.48550/arXiv:2311.12015).
- 27 Y. Shi, D. Peng, W. Liao, Z. Lin, X. Chen, C. Liu, Y. Zhang and L. Jin, Exploring ocr capabilities of gpt-4v (ision): A quantitative and in-depth evaluation, *arXiv*, 2023, preprint, arXiv:2310.16809, DOI: [10.48550/arXiv:2310.16809](https://doi.org/10.48550/arXiv:2310.16809).
- 28 S. Park, B. Kim, S. Choi, P. G. Boyd, B. Smit and J. Kim, *J. Chem. Inf. Model.*, 2018, 58, 244–251.
- 29 H. Park, Y. Kang, W. Choe and J. Kim, *J. Chem. Inf. Model.*, 2022, 62, 1190–1198.
- 30 Y. Luo, S. Bag, O. Zaremba, A. Cierpka, J. Andreo, S. Wuttke, P. Friederich and M. Tsotsalas, *Angew. Chem., Int. Ed.*, 2022, 61, e202200242.
- 31 Z. Wu, J. Chen, Y. Li, Y. Deng, H. Zhao, C.-Y. Hsieh and T. Hou, *J. Chem. Inf. Model.*, 2023, 63(24), 7617–7627.



- 32 D. Vidhani, *The Art of Asking Question: Mastering Human-AI (HAI) Duet in Chemistry Through Prompt Engineering*, 2024, DOI: [10.21203/rs.3.rs-3825267/v1](https://doi.org/10.21203/rs.3.rs-3825267/v1).
- 33 M. Ansari and S. M. Moosavi, Agent-based Learning of Materials Datasets from Scientific Literature, *arXiv*, 2023, preprint, arXiv:2312.11690, DOI: [10.48550/arXiv:2312.11690](https://doi.org/10.48550/arXiv:2312.11690).
- 34 H. Lyu, Z. Ji, S. Wuttke and O. M. Yaghi, *Chem*, 2020, **6**, 2219–2241.
- 35 O. M. Yaghi and Z. Zheng, *Reticular Chemistry and New Materials*, World Scientific, 2024.
- 36 Y. G. Chung, J. Camp, M. Haranczyk, B. J. Sikora, W. Bury, V. Krungleviciute, T. Yildirim, O. K. Farha, D. S. Sholl and R. Q. Snurr, *Chem. Mater.*, 2014, **26**, 6185–6192.
- 37 A. S. Rosen, V. Fung, P. Huck, C. T. O'Donnell, M. K. Horton, D. G. Truhlar, K. A. Persson, J. M. Notestein and R. Q. Snurr, *npj Comput. Mater.*, 2022, **8**, 112.
- 38 S. Kancharlapalli and R. Q. Snurr, *ACS Appl. Mater. Interfaces*, 2023, **15**(23), 28084–28092.
- 39 P. Z. Moghadam, Y. G. Chung and R. Q. Snurr, *Nat. Energy*, 2024, 1–13.
- 40 D. A. Boiko, R. MacKnight, B. Kline and G. Gomes, *Nature*, 2023, **624**, 570–578.
- 41 B. A. Koscher, R. B. Canty, M. A. McDonald, K. P. Greenman, C. J. McGill, C. L. Bilodeau, W. Jin, H. Wu, F. H. Vermeire, B. Jin, T. Hart, T. Kulesza, S.-C. Li, T. S. Jaakkola, R. Barzilay, R. Gómez-Bombarelli, W. H. Green and K. F. Jensen, *Science*, 2023, **382**, eadi1407.
- 42 Y. G. Chung, E. Haldoupis, B. J. Bucior, M. Haranczyk, S. Lee, H. Zhang, K. D. Vogiatzis, M. Milisavljevic, S. Ling, J. S. Camp, B. Slater, J. I. Siepmann, D. S. Sholl and R. Q. Snurr, *J. Chem. Eng. Data*, 2019, **64**, 5985–5998.
- 43 A. Nandy, G. Terrones, N. Arunachalam, C. Duan, D. W. Kastner and H. J. Kulik, *Sci. Data*, 2022, **9**, 74.
- 44 S. M. Moosavi, A. Nandy, K. M. Jablonka, D. Ongari, J. P. Janet, P. G. Boyd, Y. Lee, B. Smit and H. J. Kulik, *Nat. Commun.*, 2020, **11**, 1–10.
- 45 A. Nandy, S. Yue, C. Oh, C. Duan, G. G. Terrones, Y. G. Chung and H. J. Kulik, *Matter*, 2023, **6**, 1585–1603.
- 46 A. Nandy, C. Duan and H. J. Kulik, *J. Am. Chem. Soc.*, 2021, **143**, 17535–17547.
- 47 J. C. Tan, T. D. Bennett and A. K. Cheetham, *Proc. Natl. Acad. Sci. U.S.A.*, 2010, **107**, 9938–9943.
- 48 R. Batra, C. Chen, T. G. Evans, K. S. Walton and R. Ramprasad, *Nat. Mach.*, 2020, **2**, 704–710.
- 49 B. Wang, X.-L. Lv, D. Feng, L.-H. Xie, J. Zhang, M. Li, Y. Xie, J.-R. Li and H.-C. Zhou, *J. Am. Chem. Soc.*, 2016, **138**, 6204–6216.
- 50 W. Song, Z. Zheng, A. H. Alawadhi and O. M. Yaghi, *Nat. Water*, 2023, **1**, 626–634.
- 51 B. F. Abrahams, A. D. Dharma, M. J. Grannas, T. A. Hudson, H. E. Maynard-Casely, G. R. Oliver, R. Robson and K. F. White, *Inorg. Chem.*, 2014, **53**, 4956–4969.
- 52 Z. Zheng, H. L. Nguyen, N. Hanikel, K. K.-Y. Li, Z. Zhou, T. Ma and O. M. Yaghi, *Nat. Protoc.*, 2023, **18**, 136–156.
- 53 Z. Zheng, N. Hanikel, H. Lyu and O. M. Yaghi, *J. Am. Chem. Soc.*, 2022, **144**, 22669–22675.
- 54 K. Hatakeyama-Sato, N. Yamane, Y. Igarashi, Y. Nabae, T. Hayakawa, Prompt engineering of GPT-4 for chemical research: what can/cannot be done?, *ChemRxiv*, 2023, preprint, DOI: [10.26434/chemrxiv-2023-s1x5p](https://doi.org/10.26434/chemrxiv-2023-s1x5p).
- 55 A. G. Parameswaran, S. Shankar, P. Asawa, N. Jain and Y. Wang, Revisiting Prompt Engineering via Declarative Crowdsourcing, *arXiv*, 2023, preprint, arXiv:2308.03854, DOI: [10.48550/arXiv:2308.03854](https://doi.org/10.48550/arXiv:2308.03854).
- 56 L. Reynolds and K. McDonnell, *presented in part at the Extended Abstracts of the 2021 CHI Conference on Human Factors in Computing Systems*, 2021.
- 57 Y. Zhou, A. I. Muresanu, Z. Han, K. Paster, S. Pitis, H. Chan and J. Ba, Large language models are human-level prompt engineers, *arXiv*, 2022, preprint, arXiv:2211.01910, DOI: [10.48550/arXiv:2211.01910](https://doi.org/10.48550/arXiv:2211.01910).
- 58 M. Mitchell, A. B. Palmarini and A. Moskvichev, Comparing Humans, GPT-4, and GPT-4V On Abstraction and Reasoning Tasks, *arXiv*, 2023, preprint, arXiv:2311.09247, DOI: [10.48550/arXiv.2311.09247](https://doi.org/10.48550/arXiv.2311.09247).
- 59 A. Rohatgi, *WebPlotDigitizer*, <https://automeris.io/WebPlotDigitizer>, accessed 2022-09.
- 60 J. D. Evans, V. Bon, I. Senkovska and S. Kaskel, *Langmuir*, 2021, **37**, 4222–4226.
- 61 T. Düren, F. Millange, G. Férey, K. S. Walton and R. Q. Snurr, *J. Phys. Chem. C*, 2007, **111**, 15350–15356.
- 62 G. Hai and H. Wang, *Coord. Chem. Rev.*, 2022, **469**, 214670.
- 63 K. S. Walton and R. Q. Snurr, *J. Am. Chem. Soc.*, 2007, **129**, 8552–8556.
- 64 P. Sinha, A. Datar, C. Jeong, X. Deng, Y. G. Chung and L.-C. Lin, *J. Phys. Chem. C*, 2019, **123**, 20195–20209.
- 65 F. Ambroz, T. J. Macdonald, V. Martis and I. P. Parkin, *Small Methods*, 2018, **2**, 1800173.
- 66 J. W. M. Osterrieth, J. Rampersad, D. Madden, N. Rampal, L. Skoric, B. Connolly, M. D. Allendorf, V. Stavila, J. L. Snider, R. Ameloot, J. Marreiros, C. Ania, D. Azevedo, E. Villarrasa-Garcia, B. F. Santos, X.-H. Bu, Z. Chang, H. Bunzen, N. R. Champness, S. L. Griffin, B. Chen, R.-B. Lin, B. Coasne, S. Cohen, J. C. Moreton, Y. J. Colón, L. Chen, R. Clowes, F.-X. Coudert, Y. Cui, B. Hou, D. M. D'Alessandro, P. W. Doheny, M. Dincă, C. Sun, C. Doonan, M. T. Huxley, J. D. Evans, P. Falcaro, R. Ricco, O. Farha, K. B. Idrees, T. Islamoglu, P. Feng, H. Yang, R. S. Forgan, D. Bara, S. Furukawa, E. Sanchez, J. Gascon, S. Telalović, S. K. Ghosh, S. Mukherjee, M. R. Hill, M. M. Sadiq, P. Horcajada, P. Salcedo-Abraira, K. Kaneko, R. Kukobat, J. Kevin, S. Keskin, S. Kitagawa, K.-i. Otake, R. P. Lively, S. J. A. DeWitt, P. Llewellyn, B. V. Lotsch, S. T. Emmerling, A. M. Pütz, C. Martí-Gastaldo, N. M. Padial, J. García-Martínez, N. Linares, D. MasPOCH, J. A. Suárez del Pino, P. Moghadam, R. Oktavian, R. E. Morris, P. S. Wheatley, J. Navarro, C. Petit, D. Danaci, M. J. Rosseinsky, A. P. Katsoulidis, M. Schröder, X. Han, S. Yang, C. Serre, G. Mouchaham, D. S. Sholl, R. Thyagarajan, D. Siderius, R. Q. Snurr, R. B. Goncalves, S. Telfer, S. J. Lee, V. P. Ting, J. L. Rowlandson, T. Uemura, T. Iiyuka, M. A. van der Veen, D. Rega, V. Van Speybroeck, S. M. J. Rogge, A. Lemaire, K. S. Walton, L. W. Bingel,



- S. Wuttke, J. Andreo, O. Yaghi, B. Zhang, C. T. Yavuz, T. S. Nguyen, F. Zamora, C. Montoro, H. Zhou, A. Kirchon and D. Fairen-Jimenez, *Adv. Mater.*, 2022, **34**, 2201502.
- 67 X. Duan, C. Wu, S. Xiang, W. Zhou, T. Yildirim, Y. Cui, Y. Yang, B. Chen and G. Qian, *Inorg. Chem.*, 2015, **54**, 4377–4381.
- 68 H. Furukawa, F. Gandara, Y.-B. Zhang, J. Jiang, W. L. Queen, M. R. Hudson and O. M. Yaghi, *J. Am. Chem. Soc.*, 2014, **136**, 4369–4381.
- 69 R. J. Marshall, C. L. Hobday, C. F. Murphie, S. L. Griffin, C. A. Morrison, S. A. Moggach and R. S. Forgan, *J. Mater. Chem. A*, 2016, **4**, 6955–6963.
- 70 C. P. Krap, R. Newby, A. Dhakshinamoorthy, H. García, I. Cebula, T. L. Easun, M. Savage, J. E. Eyley, S. Gao, A. J. Blake, W. Lewis, P. H. Beton, M. R. Warren, D. R. Allan, M. D. Frogley, C. C. Tang, G. Cinque, S. Yang and M. Schröder, *Inorg. Chem.*, 2016, **55**, 1076–1088.
- 71 C. Volkringer, D. Popov, T. Loiseau, G. Férey, M. Burghammer, C. Riekel, M. Haouas and F. Taulelle, *Chem. Mater.*, 2009, **21**, 5695–5697.
- 72 B. Liu, H.-F. Zhou, L. Hou, J.-P. Wang, Y.-Y. Wang and Z. Zhu, *Inorg. Chem.*, 2016, **55**, 8871–8880.
- 73 T. Li, M. T. Kozlowski, E. A. Doud, M. N. Blakely and N. L. Rosi, *J. Am. Chem. Soc.*, 2013, **135**, 11688–11691.
- 74 Q. Yu, Y.-F. Zeng, J.-P. Zhao, Q. Yang, B.-W. Hu, Z. Chang and X.-H. Bu, *Inorg. Chem.*, 2010, **49**, 4301–4306.
- 75 L. Gagliardi and O. M. Yaghi, *Chem. Mater.*, 2023, **35**, 5711–5712.
- 76 A. A. AlGhamdi, *Mol. Front. J.*, 2023, 1–4.
- 77 A. S. Rosen, J. M. Notestein and R. Q. Snurr, *Curr. Opin. Chem. Eng.*, 2022, **35**, 100760.
- 78 H. Nori, Y. T. Lee, S. Zhang, D. Carignan, R. Edgar, N. Fusi, N. King, J. Larson, Y. Li and W. Liu, Can Generalist Foundation Models Outcompete Special-Purpose Tuning? Case Study in Medicine, *arXiv*, 2023, preprint, arXiv:2311.16452, DOI: [10.48550/arXiv.2311.16452](https://doi.org/10.48550/arXiv.2311.16452).
- 79 S. Huang, S. Mamidanna, S. Jangam, Y. Zhou and L. H. Gilpin, Can large language models explain themselves? a study of llm-generated self-explanations, *arXiv*, 2023, preprint, arXiv:2310.11207, DOI: [10.48550/arXiv.2310.11207](https://doi.org/10.48550/arXiv.2310.11207).
- 80 G. Yona, R. Aharoni and M. Geva, Narrowing the Knowledge Evaluation Gap: Open-Domain Question Answering with Multi-Granularity Answers, *arXiv*, 2024, preprint, arXiv:2401.04695, DOI: [10.48550/arXiv.2401.04695](https://doi.org/10.48550/arXiv.2401.04695).
- 81 O. Khattab, A. Singhvi, P. Maheshwari, Z. Zhang, K. Santhanam, S. Vardhamanan, S. Haq, A. Sharma, T. T. Joshi and H. Moazam, Dspy: Compiling declarative language model calls into self-improving pipelines, *arXiv*, 2023, preprint, arXiv.2310.03714, DOI: [10.48550/arXiv.2310.03714](https://doi.org/10.48550/arXiv.2310.03714).

