

L²M³OF: A LARGE LANGUAGE MULTIMODAL MODEL FOR METAL-ORGANIC FRAMEWORKS

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ABSTRACT

Large language models (LLMs) have demonstrated remarkable reasoning capabilities across diverse natural language tasks. However, comparable breakthroughs in scientific discovery are more limited, because understanding complex physical phenomena demands multifaceted representations far beyond language alone. A compelling example is the design of functional materials such as metal-organic frameworks (MOFs) — critical for a range of impactful applications like carbon capture and hydrogen storage. Navigating their vast and intricate design space in language-based representations interpretable by LLMs is challenging due to the numerous possible three-dimensional atomic arrangements and strict reticular rules of coordination geometry and topology. Despite promising early results in LLM-assisted discovery for simpler materials systems, MOF design remains heavily reliant on tacit human expertise rarely codified in textual information alone. To overcome this barrier, we introduce L²M³OF, the first multimodal LLM for MOFs. L²M³OF integrates crystal representation learning with language understanding to process structural, textual, and knowledge modalities jointly. L²M³OF employs a pre-trained crystal encoder with a lightweight projection layer to compress structural information into a token space, enabling efficient alignment with language instructions. To facilitate training and evaluation, we curate a structure–property–knowledge database of crystalline materials and benchmark L²M³OF against state-of-the-art (SOTA) closed-source LLMs such as GPT-5, Gemini-2.5-Pro, and DeepSeek-R1. Experiments show that L²M³OF outperforms leading text-based closed-source LLMs in property prediction and knowledge generation tasks, despite using far fewer parameters. These results highlight the importance of multimodal approaches for porous crystalline material understanding and establish L²M³OF as a foundation for next-generation AI systems in materials discovery.

1 INTRODUCTION

Metal-organic frameworks represent a versatile class of porous crystalline materials with high tunability and broad physical properties that promise transformative applications in direct carbon capture (Rohde et al., 2024), clean hydrogen storage (Chen et al., 2020), water harvesting (Alawadhi et al., 2024), and controlled drug delivery (Wu & Yang, 2017). MOF functional design involves intricate reticular synthesis procedures by linking metal atoms and organic molecules into repeating patterns, akin to ‘LEGO building’ at the nanoscale. Scaling-up their design is nevertheless non-trivial, even with machine learning, both due to the large number of possible building-block combinations that give rise to an enormous design space, and because of the expertise-driven nature of the design which heavily relies on domain knowledge (Yaghi et al., 2003).

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Large language models have recently emerged as powerful AI assistants for chemists, demonstrating strong reasoning capabilities in language-related chemistry tasks, such as chemical knowledge integration and tool orchestration, offering promising potential in accelerating the exploration of large design spaces (Mirza et al., 2025). The discovery of new functional materials such as MOFs however, is fundamentally more challenging because unimodal textual representations typically fail to capture complex, high-dimensional reticular phenomena that give rise to different functionalities. Unlike molecules (Zhu et al., 2024) or proteins (Wu et al., 2023), which can be expressed as textual sequences of a relatively small range of elements, MOFs inhabit three-dimensional, periodic structures that resist straightforward representation. In addition to being compositionally much broader, the structure-function problem for MOFs is inherently more complex because the 3-dimensional atomic ‘sequence’ does not encode function alone; rather, it emerges from a combination of factors, including local bonding environments, long-range crystallographic symmetry, pore connectivity, and other topological features (Luo et al., 2024).

Despite the emerging line of research on LLMs for accelerated materials design (Kang et al., 2025; Duan et al., 2025) spanning a broad range of downstream tasks, including property prediction (Niyongabo Rubungo et al., 2025) and de-novo structure generation (Wang et al., 2025a), existing approaches remain restricted to text-centric or file-based representations, such as crystallographic information files (CIFs) and text-based property descriptions (Tang et al., 2025). While effective for sequential reasoning, such encodings fail to capture three-dimensional symmetries, periodicity, and long-range structural correlations that underpin crystalline behavior, often underperforming when compared with geometry- or symmetry-aware models (Alampara et al., 2025b). A collection of existing LLMs and their modeling capacity for crystalline materials is presented in Table 1.

The challenge here extends beyond structural representation; it lies in the ‘machine understanding’ of materials’ functionality. Multimodal integration in learning strategies, that is, leveraging atomic information as well as literature knowledge to interlink structure with function, is therefore key to enable a holistic understanding of materials’ applicability. Whereas molecular modeling has seen initial success in coupling LLMs with graph neural networks or generative models (Jablonka et al., 2024), analogous strategies for crystalline systems remain rare, owing to system and design complexity, as well as the lack of standardized datasets and benchmarks tailored to crystalline materials, rendering rigorous evaluation and reproducibility quite challenging.

This work proposes L^2M^3OF , the first *multimodal LLM for MOF design* that combines multimodal MOFs representations (Park et al., 2023) with curated domain-knowledge from MOFs literature. L^2M^3OF is versatile and inherently designed to be lightweight to allow for an efficient alignment with language instructions, demonstrating SOTA performance on diverse design-critical tasks including property prediction and material application recommendation, rendering it an indispensable AI-assistant for chemists and materials scientists. To train and test L^2M^3OF , we curate the first-ever *structure-property-knowledge MOFs database*, namely MOF-SPK, featuring structural, property and domain-knowledge information for more than 100,000 MOFs materials. L^2M^3OF outperforms leading commercially-available LLMs such as DeepSeek, GPT-4o, and Gemini-2.5-Pro, demonstrating SOTA capabilities not only in capturing essential representational aspects of complex MOFs systems, but also a holistic understanding of their broader functional role and potential applicability. Fig. 1 illustrates the core architectural features of L^2M^3OF .

2 BACKGROUND AND RELATED WORK

Crystal representation. A crystal structure is defined by the geometric arrangement of its atoms within a unit cell. The unit cell represents the smallest repeating block that captures and maintains the complete symmetry and structure of the crystal. A crystal can therefore be uniquely represented as $\mathcal{M} = (\mathcal{A}, \mathbf{X}, \mathbf{L})$ using the following three parameters that characterize its unit cell: i) atom identities $\mathcal{A} = \{a_0, \dots, a_N\} \in \mathbb{A}^N$, where \mathbb{A} denotes the set of all chemical elements, ii) Cartesian coordinates of atoms $\mathbf{X} = [\mathbf{x}_0, \dots, \mathbf{x}_N]^T \in \mathbb{R}^{N \times 3}$ and iii) the lattice matrix that describes the periodicity of the crystal $\mathbf{L} = [\mathbf{l}_1, \mathbf{l}_2, \mathbf{l}_3]^T \in \mathbb{R}^{3 \times 3}$. Crystal information is traditionally encoded in standardized text format in Crystallographic Information Files (CIFs), which have been the key-stone of systematically curated databases of both predicted and experimentally found crystals (Boyd et al., 2019; Zhao et al., 2025) and actively used for materials discovery over the last decades. Fig. 10 exemplifies the structure of a CIF file. CIF databases come with a two-fold challenge: they

Model	CrystalType	LLM	Model Input		Downstream Tasks			
			Text	Multimodal	Structure	Property	Knowledge	Q&A
LLM-Prop (Niyongabo Rubungo et al., 2025)	Inorganic	T5	✓			✓		
CrystLLM (Antunes et al., 2024)	Inorganic	Llama-2	✓			✓		
CrysText (Mohanty et al., 2024)	Inorganic	Llama-3.1	✓		✓			
Mat2Seq (Yan et al., 2025)	Inorganic	GPT	✓		✓			
MatText (Alampara et al., 2025a)	Inorganic	Llama-2	✓			✓		
CrystallCL (Wang et al., 2025b)	Inorganic	Llama-2	✓		✓			
CSLLM (Song et al., 2025)	Inorganic	Llama-3	✓				✓	
deCIFer (Johansen et al., 2025)	Inorganic	Transformer	✓		✓			
MatterGPT (Wang et al., 2025a)	Inorganic	GPT	✓		✓			
Text2Struc (Baibakova, 2025)	Inorganic	CodeGen	✓		✓			
Matterchat (Tang et al., 2025)	Inorganic	Mistral	✓	✓		✓		✓
Chemeleon (Park et al., 2025)	Inorganic	BERT	✓	✓	✓			
MOFGPT (Badrinarayanan et al., 2025)	MOFs	GPT	✓		✓	✓		
ChatMOF (Kang & Kim, 2024)	MOFs	GPT	✓		✓	✓		
L ² M ² OF	MOFs	Qwen2.5	✓		✓	✓	✓	✓
L ² M ³ OF	MOFs	Qwen2.5	✓	✓	✓	✓	✓	✓

Table 1: Model features of LLMs for crystalline materials. ‘Structure’ corresponds to structure prediction or structure extraction; ‘Property’ means property prediction; ‘Knowledge’ means knowledge generation, and ‘Q&A’ stands for question and answering.

describe materials’ structures and isolated properties but lack holistic information on their functionality, which is present in published papers. Importantly, the textual format of CIFs is less amenable to typical ML pipelines posing barriers to streamlining data-driven materials discovery Tian et al. (2022). While this still remains a grand challenge in materials science, recent literature has increasingly focused on the development of either hand-crafted or machine-learned crystal representations that are well suited for machine learning algorithms.

Crystal representation learning. Recent progress in crystal representation learning spans a wide range of representations and modalities, ranging from graph-based to structural and foundation model approaches. CGCNN (Xie & Grossman, 2018) pioneered interpretable crystal graph convolutional networks for property prediction directly from atomic connections, while iCGCNN (Cheng et al., 2021) further enhances this by incorporating Voronoi tessellation and three-body interactions. Physics-guided generative models like PGCGM (Zhao et al., 2023) leverage symmetry-affine transformations to generate diverse, structurally valid crystals, significantly outperforming previous generators. MOFTransformer (Kang et al., 2023) was the first inherently multimodal architecture, combining atom-based and energy-grid embeddings to capture local and global features, achieving SOTA property prediction for MOFs. Similarly, DeepSorption (Cui et al., 2023) integrates global structural awareness via a transformer for highly accurate adsorption predictions in porous materials. More recently, the emergence of foundation models allowed the extension of these practises to broader crystalline systems. MCRT (Feng et al., 2025) multimodally integrates local atomic information with global persistence-image based views of organic molecular crystals, while CLOUD (Xu et al., 2025) employs symmetry-aware, physic-informed string representations for the development of a scalable foundation model, pre-trained on millions of inorganic crystals. Together, these approaches illustrate the shift from local graph-based modeling to multimodal, geometry-aware methodologies, which enable few-shot learning and hold significant promise for leveraging their representations in LLMs.

LLMs for crystalline materials. LLMs have recently drawn much interest from the chemistry and materials community due to their unique capabilities in text generation, chemical knowledge integration, and characterization tool utilization (Zheng et al., 2025). Recent efforts have demonstrated the LLMs’ capacity to process raw CIF files of inorganic crystals to generate textual descriptions for further language-based training exploitation (Alampara et al., 2025a). Beyond text generation, LLMs have demonstrated promising performance in plausible structure generation, such as CrystaLLM (Antunes et al., 2024), which was trained on millions of inorganic crystal CIF files and validated via ab initio simulations on de-novo generated structures. Chemeleon (Park et al., 2025) proposed the integration of text descriptions with 3D structural data using cross-modal contrastive learning and diffusion models, enabling natural language-guided generation of chemical compositions and structures of inorganic crystals. CSLLM (Song et al., 2025), a framework of three fine-tuned LLMs, use a textual representation for crystal material to predict the synthesizability, synthesis method, and precursors of 3D inorganic crystals. Finally, Matterchat (Tang et al., 2025) is

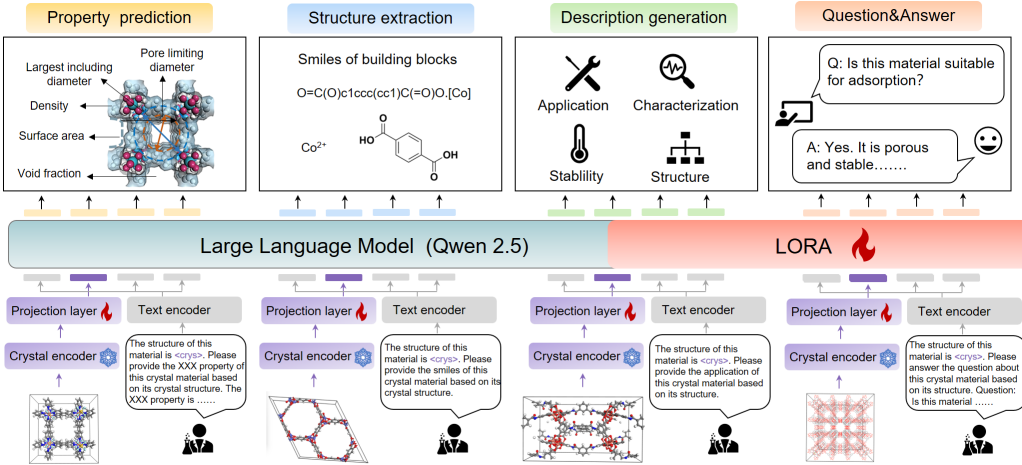


Figure 1: An overview of the L^2M^3OF framework and its applicability in MOFs design.

a structure-aware LLM for inorganic materials, trained on more than 140,000 structures, capable of ingesting textual information from atomic structures to reason answers on material description and property-prediction. While these approaches have demonstrated success to small-scale systems, they struggle to generalise to larger, complex systems, such as MOFs, with hundreds or even thousands of atoms per unit cell and latent functionality cues concealed in their CIF representations (Xiao et al., 2023). MOFGPT Badrinarayanan et al. (2025) is the first LLM for de-novo generation of MOFs, utilizing a GPT generator trained on MOFid sequences and a reinforcement learning framework that steers generation toward target properties—as predicted via MOFormer Kang et al. (2023)—using reward functions. Nevertheless, MOFGPT’s generator relies on 2D MOFid strings, missing thus intrinsic functional information encoded in their 3D structure of MOFs. Our work, further complements this expressivity by not only enabling the use of 3D representation of large MOFs structures, but also enhancing the functionality prediction and comprehension of the LLM via jointly training on domain-specific information from the literature.

3 DATASET AND MODEL ARCHITECTURE

3.1 MOF-SPK DATASET CONSTRUCTION

To train and validate the performance of our model we construct a structure–property–knowledge database for MOFs, namely MOF-SPK. We specifically couple experimentally found MOF structures in CIF format together with curated literature information directly related to these structures. We selected 133,737 CIFs from the Cambridge Crystallographic Data Centre (CCDC) database (Groom et al., 2016) together with the corresponding publications that reported those structures. To ensure data consistency, we removed guest molecules from the MOFs structures using the CSD Python API and applied the StructureMatcher module from the pymatgen Python package (Ong et al., 2013) to identify and eliminate duplicate entries with equivalent crystal structures. We further augmented this data with property calculations and knowledge, leveraging high-throughput computational tools, Python packages and LLMs into a structure–property–knowledge database which formed the test-bench for this study. Based on MOF-SPK we designed four assessment tasks which we then use to evaluate the utility of our LLM as an assistive tool for MOF design. These tasks represent critical and time-consuming characterization procedures in the design of functional materials that can offer valuable acceleration and assistance. The tasks include: i) *property prediction*, ii) *structure extraction*, iii) *description generation*, and iv) *general question & answering*, and form the basis to facilitate model training and evaluation.

Property prediction assesses the LLM’s complex physical perception ability on the absolute and relative positions of atoms in the 3D space. Here we computed key physical characteristics of crystal materials — density, pore limiting diameter (PLD), largest cavity diameter (LCD), accessible

surface area (ASA), and void fraction (VF) — using high-throughput computational tools such as ZEO++ (Willems et al., 2012). The structure extraction task assesses the LLM’s capacity to learn and understand the material’s reticular composition of building units, i.e., by extracting the SMILES representation of its molecular building blocks. The MOFid Python toolbox (Bucior et al., 2019) was used to generate the data for this task. For the description generation task, we curated domain knowledge from more than 35,000 scientific publications on experimentally discovered MOFs, extracting information on applications, characterization methods, stability, and structural features with the assistance of DeepSeek-R1. Application refers to the uses of the material, such as adsorption or catalysis. Characterization method specifies the experimental techniques that should be employed to analyze the material, for example, Powder X-ray Diffraction or X-ray Photoelectron Spectroscopy. Stability describes how stable the material is, for instance, whether it remains stable in water. Structural features summarize the material’s structural characteristics, such as forming a 1D chain or a 3D open framework. The description generation task assesses whether the crystal LLM can learn and establish the relationship between crystal structures and crystal knowledge. Finally, the question & answering task assesses the LLM’s ability to answer materials-related questions on MOFs. We generated five questions and answers for each scientific publication based on their abstract. First, we prompted DeepSeek-R1 to identify five keywords from each abstract and then extract relevant questions and answers pairs around those. Examples of the above tasks are included in Section A.3 of the Appendix. To ensure that the MOF-SPK dataset is chemically well-balanced in terms of material representation we perform comprehensive statistical analyses included in Section A.2 of the Appendix.

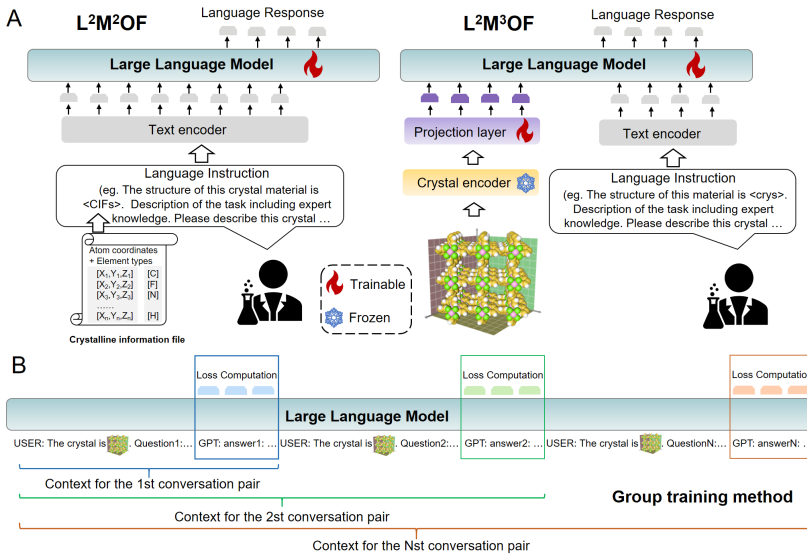


Figure 2: An overview of model architecture and model training methods. (A) The architecture differences between L^2M^2OF and L^2M^3OF . (B) The schematic diagram of the group training method.

3.2 MODEL ARCHITECTURE

We design two complementary models; apart from our main contribution, L^2M^3OF — a multimodal LLM that incorporates structural information through a MOF 3D structure encoder, we further develop a language-only variant, namely L^2M^2OF , which instead represents MOFs in their textual CIF format. The juxtaposition of the two variants against our expertly designed MOF tasks, helps better understand the role of multimodal material representations in LLM-guided discovery. Fig. 2 shows the architectures of the two proposed models.

L^2M^2OF processes a crystal material \mathcal{M} by converting its CIF into a textual sequence $S_{\mathcal{M}}$. This sequence contains the unit cell parameters, space group symmetry, and atomic coordinates with their respective element types. This material representation is concatenated with a task-specific natural

language instruction $I_{\mathcal{T}}$ to form the complete input prompt as $X_{\text{LLM}} = [S_{\mathcal{M}}; I_{\mathcal{T}}]$. The instruction $I_{\mathcal{T}}$ embeds expert domain knowledge to guide the model. For instance, for VF prediction, $I_{\mathcal{T}}$ defines the property and its physical significance. The model then generates the target prediction Y autoregressively. The probability of generating the output sequence of L tokens is given by:

$$P(Y | X_{\text{LLM}}) = \prod_{i=1}^L P(y_i | y_{<i}, X_{\text{LLM}}; \Theta_{\text{LLM}}), \quad (1)$$

where Θ_{LLM} represents the parameters of a pre-trained LLM. A key advantage of this text-only paradigm is its ability to perform inference with SOTA commercial LLMs (e.g., GPT-5, Gemini-Pro, Deepseek-R1) without additional training. In this study, we also fine-tuned an open-source LLM on the MOF-SPK database to serve as a strong text-only baseline for comparison against our multimodal approach.

L^2M^3OF conversely fuses textual instructions with non-textual, geometric structural data. The model consists of three core components:

Crystal Structure Encoder: We employ PMTransformer (Park et al., 2023) as the crystal encoder, which is a GNN pre-trained on 1.9 million hypothetical porous materials. It takes the crystal structure \mathcal{M} and outputs a fixed-dimensional latent representation, or embedding, $\mathbf{z}_{\text{struct}} \in \mathbb{R}^d$:

$$\mathbf{z}_{\text{struct}} = \text{PMTransformer}(\mathcal{M}; \Theta_{\text{PMT}}), \quad (2)$$

where Θ_{PMT} represents the frozen, pre-trained parameters of the encoder.

Multimodal Projection Bridge: This component transforms and compresses the structural embedding for seamless integration with the language model through a compression and projection network. The compression network, $\text{MLP}_{\text{token}}$, then compresses this sequence along the token dimension from length N to a shorter, fixed length M ($M < N$). We empirically found that this compression accelerates training significantly without loss of performance. The projection network, MLP_{feat} , projects the encoder’s output from its native dimension d_{enc} to a sequence of N tokens in LLMs’ embedding space $\mathbb{R}^{d_{\text{LLM}}}$. The entire process is defined as:

$$\mathbf{H}_{\text{struct}} = \text{MLP}_{\text{token}}(\mathbf{z}_{\text{struct}}; \Theta_{\text{token}}), \quad \mathbf{H}_{\text{proj}} = \text{MLP}_{\text{feat}}(\mathbf{H}_{\text{struct}}; \Theta_{\text{feat}}), \quad (3)$$

where $\Theta_{\text{bridge}} = (\Theta_{\text{feat}}, \Theta_{\text{token}})$ denotes the combined parameters of the projection and compression MLPs, and $\mathbf{H}_{\text{struct}} \in \mathbb{R}^{M \times d_{\text{enc}}}$, $\mathbf{H}_{\text{proj}} \in \mathbb{R}^{M \times d_{\text{LLM}}}$.

Large Language Model: The compressed structural token sequence $\mathbf{H}_{\text{struct}}$ is prepended to the tokenized instruction sequence $\text{Tokenize}(I_{\mathcal{T}})$ to form the combined input for the LLM. The LLM then generates the output conditioned on this multimodal input:

$$P(Y | \mathcal{M}, I_{\mathcal{T}}) = \prod_{i=1}^L P(y_i | y_{<i}, \mathbf{H}_{\text{struct}}, I_{\mathcal{T}}; \Theta_{\text{LLM}}, \Theta_{\text{bridge}}). \quad (4)$$

During training, the encoder parameters Θ_{PMT} are kept frozen to preserve its pre-trained knowledge and stabilize training. Only Θ_{Bridge} and Θ_{LLM} are updated.

3.3 TRAINING OBJECTIVE

We trained our models using an instruction-tuning paradigm, tailoring them for property prediction tasks in materials science. The objective is to minimize the negative log-likelihood of the target sequence (e.g., the numerical or classification values) given the input instruction and material data.

For a dataset \mathcal{D} of N examples, each containing an instruction, a material, and a target response $(I_{\mathcal{T}}^{(i)}, \mathcal{M}^{(i)}, Y^{(i)}) i = 1^N$, the loss function \mathcal{L} for L^2M^3OF is defined as:

$$\mathcal{L}(\Theta_{\text{LLM}}, \Theta_{\text{bridge}}) = -\frac{1}{N} \sum_{i=1}^N \log P(Y^{(i)} | \mathcal{M}^{(i)}, I_{\mathcal{T}}^{(i)}; \Theta_{\text{LLM}}, \Theta_{\text{bridge}}). \quad (5)$$

This supervised fine-tuning (SFT) process teaches the model to follow instructions and reason about material properties based on the provided textual and structural information, enabling it to generalize to new, unseen materials and tasks. We further implement a *group training strategy* to enhance

context diversity during SFT. For each mini-batch, we randomly concatenate multiple instruction-answer pairs to form extended conversational contexts. The loss is computed only on the answer tokens of each question within the group, while the preceding Q&A pairs serve as contextual background. This approach effectively increases the diversity of training contexts without substantially increasing computational costs, as the total number of tokens per batch remains unchanged. The method acts as an efficient form of data augmentation, exposing the model to richer contextual patterns during training.

4 EXPERIMENTS

4.1 EXPERIMENTAL RESULTS

Training models on past data and evaluating them on future discoveries is crucial because it mirrors real-world deployment scenarios. To this end, we partition the dataset by material deposition year, i.e., crystal structures deposited on or before 2020 were used for training, while those from 2021 onwards formed the validation set. We further sample 500 crystal structures deposited after 2022 and use as the test set. We evaluate the performance of our models against leading commercial LLMs

Table 2: Performance comparison of commercial LLMs, L^2M^2OF , and L^2M^3OF on property prediction and structure extraction. The best performances are in **bold**, the second best underlined.

Metric	DeepSeek-V3	DeepSeek-R1	GPT-4o	GPT-5 mini	GPT-5	Gemini-2.5-pro	L^2M^2OF	L^2M^3OF
Property Prediction (MAE)								
PLD (↓)	1.99	1.97	2.94	2.93	3.24	2.09	<u>1.21</u>	0.55
LCD (↓)	2.27	3.10	4.14	4.59	4.37	2.28	<u>1.00</u>	0.53
Density (↓)	0.41	0.35	9.86	<u>0.31</u>	<u>0.31</u>	0.31	0.55	0.17
ASA (↓)	762.7	1481.6	745.3	1317.6	726.2	805.9	<u>497.8</u>	253.7
VF (↓)	0.21	0.39	<u>0.13</u>	9.63	0.08	0.88	<u>0.04</u>	0.01
Structure Extraction								
BLEU (↑)	0.27	0.28	0.20	<u>0.38</u>	<u>0.38</u>	0.38	0.41	0.34
EXACT (↑)	0.00	0.01	0.00	0.02	0.02	0.00	0.29	<u>0.18</u>
MACCS (↑)	0.50	0.52	0.49	0.53	0.56	<u>0.57</u>	0.63	0.50
RDK (↑)	0.32	0.40	0.27	0.37	0.43	0.46	<u>0.44</u>	0.25
MORGAN (↑)	0.22	0.25	0.18	0.24	<u>0.28</u>	0.29	0.36	0.23
VALIDITY (↑)	0.34	0.44	0.35	0.44	0.60	0.80	<u>0.78</u>	0.75

by Google (Comanici et al., 2025), DeepSeek Guo et al. (2025) and OpenAI (OpenAI et al., 2024) on the four tasks introduced in Section 3.1 to assess the learning and comprehensive capabilities of LLMs for MOFs. Here we do not compare against other crystal LLMs from the literature as these are not suitable for MOF materials or do not support knowledge generation capabilities which is one of the main scopes of this study.

Property prediction. This task assesses the LLMs’ performance (in terms of mean absolute error) in accurately predicting a wide range of MOF properties. As shown in the top section of Table 2, L^2M^3OF attains the lowest MAE on all five targets, while commercial LLMs consistently underperform, especially on geometry-sensitive metrics such as PLD, LCD, and ASA, which require robust grounding in 3D pore topology. Even on the easier task of density prediction, the best commercial systems (Gemini-2.5-Pro, GPT-5, GPT-5-mini) still perform worse against L^2M^3OF . Importantly, we further observe clear failure modes suggestive of hallucination or unit/normalization errors: GPT-4o reaches MAE = 9.86 on density and GPT-5 mini reaches MAE = 9.63 on void fraction. The L^2M^2OF variant also outperforms the commercial LLMs on property prediction, albeit ‘losing’ against the multimodal L^2M^3OF . Under same number of training steps however, L^2M^2OF is substantially slower because textual crystal descriptions require far more tokens. These findings demonstrate the utility of literature injected domain knowledge in the training of scientific LLMs and further indicate that multimodal training enhances an LLM’s ability to perceive and reason about the 3D spatial information of porous crystalline materials significantly, yielding superior accuracy as well as efficiency.

Structure extraction. Extracting molecular building blocks from MOFs requires a fine-grained perception of local chemical information and structural features. Here we compare the SMILES of

LLM-extracted units against ground-truth (as computed in Section 3.1 to assess accuracy and validity according to the BLEU, EXACT and VALIDITY normalized scores as in (Zhuang et al., 2025)¹. We further assess structural similarity between the extracted molecular units and the ground-truth. We test three different molecular fingerprinting methods, namely MACCS, RDKit and Morgan and use the Tanimoto similarity metric (Szafarczyk et al., 2024). Interestingly, LLMs that use CIFs as textual input achieve even stronger results on this task. In particular, L²M³OF performs best on BLEU, EXACT, MACCS, and MORGAN, while Gemini-2.5-pro leads on RDK and VALIDITY, with a high SMILES VALIDITY score of 0.8, which highlights the strong capabilities of advanced commercially available LLMs in chemical tasks. The success of CIF-based models is not entirely surprising however; the explicit textual representation of CIF directly encodes the elemental composition of materials, which facilitates the inference of constituent molecules and metallic units putting more emphasis on local environment. In contrast, vectorized crystal representations make such local compositional information less transparent.

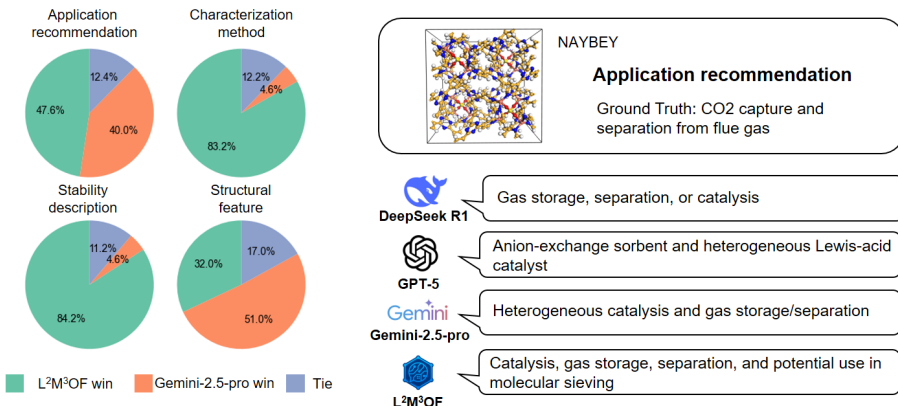


Figure 3: Performance comparison of Gemini-2.5-pro and L²M³OF on the tasks of description generation and a case study of application recommendation task.

Description generation. The description generation task is evaluated across four subtasks: application recommendation, characterization method, stability description, and structural feature. Among these, application recommendation is both the most important and the most challenging, as it requires not only an accurate perception of crystal structures and properties but also sufficient domain knowledge to map materials to plausible use cases. This is an indispensable tool to support scientists in making informed, application-oriented decisions rather than treating structure analysis in isolation. To ensure reliable evaluation and reduce reliance on manual judgment, we employ GPT o4-mini as an impartial chemistry knowledge judge. Following (Wang et al., 2023), we adopt a calibration strategy where o4-mini compares the outputs of two LLMs for each test question. Since LLMs are sensitive to response order, we mitigate positional bias by swapping the order of the outputs and re-evaluating. The final score is computed by aggregating results from both prompt orders. Each evaluation includes the question, a ground-truth answer and two candidate responses. During assessment, the o4-mini is instructed to select best LLM responses based on strict scientific accuracy and factual correctness.

Given its strong preliminary performance on property prediction and structure extraction, we adopt Gemini-2.5-Pro as a baseline for comparisons. As shown in the left-most side of Fig. 3, L²M³OF outperforms Gemini-2.5-Pro across the description generation task. Averaged over the four subtasks, it achieves a 61.8% win rate with 13.2% ties. Notably, for application recommendation L²M³OF wins 47.6% of the head-to-head comparisons versus 40.0% for Gemini-2.5-Pro, indicating stronger grounding from structure to function. The right-most side of Fig. 3 shows an application recommendation example case for a material called NAYBEY (that efficiently separates carbon dioxide

¹The SMILES BLEU score measures the overlap between the LLM-generated and ground-truth SMILES strings. EXACT assesses exact SMILES matches. VALIDITY evaluates the percentage of the LLM-generated molecules that conform to chemical syntax rules (Zhuang et al., 2025).

from nitrogen via molecular sieving (You et al., 2022)), where both DeepSeek-R1 and Gemini-2.5-Pro correctly suggested gas adsorption and separation, whereas GPT-5 failed to do so. L^2M^3OF went further by explicitly identifying the material’s molecular sieving potential. Generally, L^2M^3OF demonstrates decisive gains in characterization method and stability description, while Gemini’s responses often remain overly generic. On the structural feature subtask, Gemini-2.5-Pro maintains edge, consistent with earlier structure-extraction results. This again reflects the advantage of text-based CIF representation in accurately capturing local structural information that are pertinent to structural feature description task.

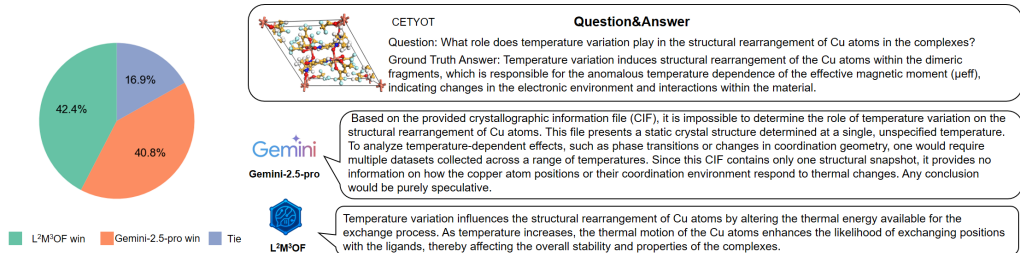


Figure 4: Performance comparison of Gemini-2.5-pro and L^2M^3OF in the tasks of question&answer.

Question & Answering. The question & answering task not only relies on understanding the crystal structure but also emphasizes the model’s mastery of MOF-specific domain knowledge. Figure 4 illustrates an example test of Q&A as well average performance comparisons, highlighting L^2M^3OF ’s clear win rate of 42.4% against Gemini-2.5-pro (40.8%). We empirically observe that overall, Gemini-2.5-pro often produces overly verbose responses and fails to reason toward the correct answer due to its lack of domain knowledge, whereas L^2M^3OF is able to reason and respond concisely and accurately.

Table 3: Ablation study performance comparison.

	Property Prediction					Structure Extraction					
	PLD (↓)	LCD (↓)	Density (↓)	ASA(↓)	VF (↓)	BLEU (↑)	EXACT (↑)	MACCS (↑)	RDK (↑)	MORGAN(↑)	VALIDITY(↑)
L^2M^3OF	0.55	0.53	0.17	253.7	0.01	0.34	0.18	0.50	0.25	0.23	0.75
w/o joint training	3.41	6.13	1.34	1616.3	0.21	0.19	0.01	0.32	0.14	0.11	0.63
w/o group training	0.73	0.72	0.18	275.1	0.02	0.27	0.17	0.48	0.23	0.21	0.77

4.2 ABLATION STUDY

To probe cross-task interactions, we compare joint training and separate training across different tasks using the same data budget and model size (Table 3 and Fig 11). The results show clear, consistent gains from joint training. On property prediction, the jointly trained model achieves substantially lower MAE on geometry-dependent targets. There is also a significant improvement in the structure extraction task. In the description generation task, the head-to-head win rate against Gemini-2.5-Pro rises from 53.3% to 61.8%. The three tasks capture complementary facets of the same underlying material representation. Property prediction forces the model to be numerically faithful to pore geometry; structure extraction sharpens the model’s awareness of local chemistry; description generation ties these cues to functional outcomes. Optimizing them together encourages a holistic, structure-aware embedding that captures both global topology and local chemical context. We also investigate the effect of group training which, without introducing additional training overhead, primarily improves the predictive accuracy of the model on the property prediction task. The above experimental results further demonstrate the importance of enabling the model to jointly learn and capture crystal structures, properties, and knowledge.

5 CONCLUSIONS

We proposed L²M³OF, the first multimodal large language model designed specifically for MOFs. By integrating geometric structure encoding with language-based domain knowledge, L²M³OF outperforms state-of-the-art commercial LLMs across property prediction, description generation, and question answering tasks—despite using fewer parameters. These results highlight the importance of multimodal architectures in capturing the intricate interplay between structure and function in crystalline materials. L²M³OF’s success demonstrates how grounding LLMs in 3D representations and curated literature can bridge gaps in automated materials discovery. As a lightweight and versatile tool, it offers chemists a scalable AI assistant for navigating complex design spaces.

6 REPRODUCIBILITY STATEMENT

To ensure our findings are reproducible, we’ll make all code and processed data publicly available upon paper acceptance. The dataset construction is detailed in Section 3.1, and we will share the processed data to facilitate its use by others. The model architecture is fully described in Section 3.2, and training specifics are provided in Section 3.3. The evaluation protocols are laid out in detail in the Section 4.1. We commit to making both our training and inference code accessible, allowing for full replication of our experiments. This comprehensive approach ensures that our results can be validated and built upon by the research community.

7 ETHICS STATEMENT

While our training data is confined to scientific literature on MOFs, the underlying base model carries potential societal biases and inherent safety risks. Consequently, our model’s outputs do not ensure 100% accuracy and may not comprehensively cover the full spectrum of safety and honesty. The model should, therefore, only be used under professional guidance to prevent the generation of biased, inaccurate, or harmful content in real-world applications.

To mitigate these risks and ensure responsible future development, we recommend a series of safeguards. The deployment of this model in real-world scientific research should be accompanied by professional guidance. Future research should prioritize a more comprehensive evaluation of the base model’s safety and explore integrating formal safety and honesty constraints directly into the trained L²M³OF architecture. These crucial steps can help ensure that further advancements in this field are built upon a strong foundation of ethical responsibility.

8 CODE AVAILABILITY

The code will be made available online after the article is accepted for publication.

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A APPENDIX

A.1 THE USE OF LARGE LANGUAGE MODELS

In this study, large language models were employed in several aspects of our work. During manuscript preparation, we used LLMs for polishing the language. In the research process, LLMs were applied to literature corpus processing, benchmark testing on the MOF-SPK database, and serving as evaluators of experimental results. The specific usage details and the models adopted are provided in the main text. All intellectual contributions such as research ideas, experimental designs, analyses, and conclusions were developed solely by the authors, who take full responsibility for the content of this paper.

A.2 MOF-SPK STATISTICAL ANALYSIS

In this section we examine the underlying chemical balance of the MOF-SPK dataset in its three main aspects, structure, property and knowledge. In terms of elemental diversity, MOF-SPK is quite diverse: excluding noble gases that do not form coordination bonds, the database contains up to 81 chemical elements (Fig. 5A). We further examine the distribution of MOF sizes in the database, expressed in number of LLM tokens, to assess the representational bias of diverse materials when processed by LLMs. For this we used the Qwen2.5-7B (Yang et al., 2025) tokenizer to quantify the token-length distribution of CIF representations for MOFs. The dataset exhibits a unimodal distribution with a peak between 10³ and 10⁴ tokens, indicating that textual serialization of crystal structures typically requires thousands to tens of thousands of tokens (Fig. 5B). Notably, the most verbose case, the CIF of material LELMEW, reaches 94,000 tokens while the structure contains only 3216 atoms, underscoring the substantial redundancy introduced by purely text-based encodings of complex crystalline materials. These observations motivate more compact representations, such as structured symbols or multimodal embeddings, that capture geometric and compositional information without incurring excessive sequence length. We further analyzed the distributions of five key MOF properties, namely LCD, PLD, density, ASA, and VF, crucial for understanding MOFs' physical characteristics. Their distributions exhibit long-tailed behavior, highlighting the inherent challenges in predicting these properties (Fig. 5C). We use Qwen2.5-7B to analyze and summarize the application landscape of MOFs. The results show substantial breadth: beyond uses such

as gas adsorption and separation, catalysis, and chemical sensing, MOFs also function as luminescent materials and as crystalline sponges for host–guest chemistry (Fig. 5D). This diversity creates an opportunity for LLMs to exceed the capabilities of individual human experts, who are typically specialized in a single application area and may overlook cross-domain potential. For example, a framework designed by an adsorption specialist might underperform on uptake targets yet be an excellent catalyst; domain boundaries can mask such good alternative uses.

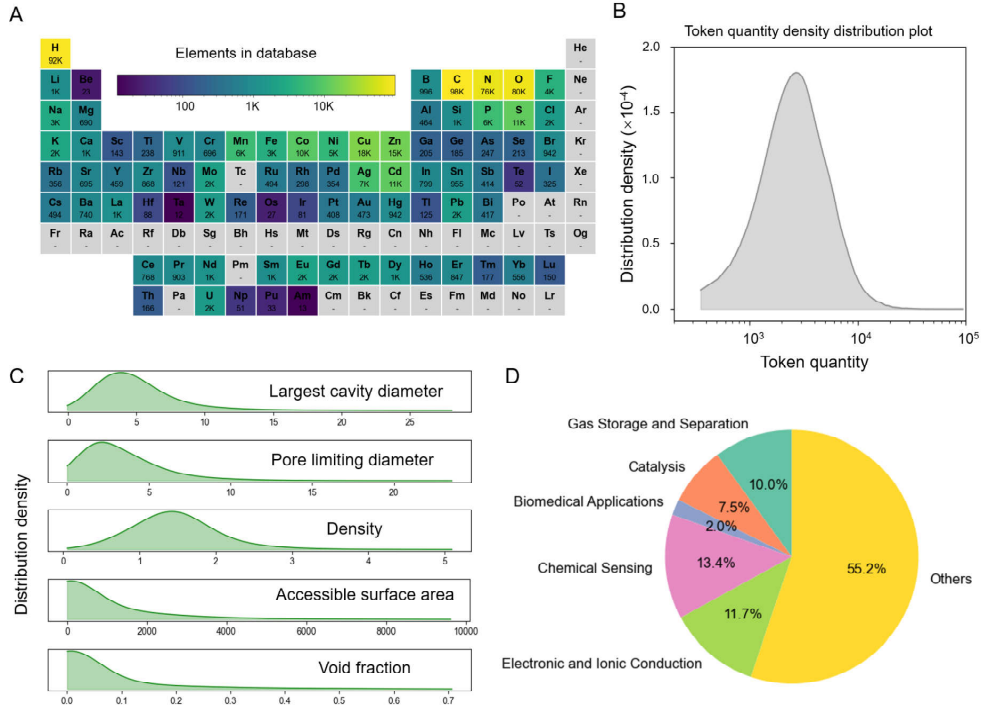


Figure 5: Data analysis of structure–property–knowledge database for crystal materials. (A) Elemental distribution in the dataset. (B) The token quantity density distribution of the CIFs in the dataset. (C) The distribution of properties of crystal material in the dataset. (D) The application distribution of the crystal material in the dataset.

A.3 DATASET EXAMPLE

Prompt

The structure of the crystal material is <crys>. Can you provide the pore limiting diameter of this crystal material according to the structure of this crystal material? The pore limiting diameter (PLD) in crystal materials refers to the smallest diameter of a sphere that can pass through the pore structure of the material. This diameter represents the smallest void space in the pore system, and it's crucial for understanding the material's size-selective capabilities, especially in applications like gas separation. The output format is [[pore limiting diameter]] and the unit of the output is Å. The probe molecule is the Nitrogen molecule, and the radius of the probe molecule is 1.82 Å.

Ground-truth

[[1.5 Å]]

Figure 6: Example of prompt and ground truth for property prediction task.

Prompt

The structure of the crystal material is <crys> .The Simplified Molecular Input Line Entry System (smiles) is a specification in the form of a line notation for describing the structure of chemical species using short ASCII strings. SMILES strings can be imported by most molecule editors for conversion back into two-dimensional drawings or three-dimensional models of the molecules.Can you provide the smiles of this crystal material according to the structure of this crystal material.The format of the output is {{SMILES}}.

Ground-truth

{{O=C1N=NC(=O)c2cnccc21.[Zn]}}

Figure 7: Example of prompt and ground truth for structure extraction task.

Prompt

The structure of the crystal material is <crys>. Please provide the Application of this crystal material and provide the reason in 100 words according to the structure of this crystal material.

Ground-truth

Anti-corrosion coatings for electrogalvanized steel
The reason is: The study demonstrates that binary zinc carboxylate MOFs (e.g., ZnC₁₂C₁₆) form compact, hydrophobic layers on metal surfaces. These coatings act as barriers against corrosion by inhibiting zinc dissolution and reducing solubility. Their performance surpasses single carboxylates (e.g., Zn(C₁₂)₂ or Zn(C₁₆)₂) in electrochemical and climatic tests, aligning with industrial requirements for eco-friendly alternatives to phosphating/chromating processes.

Figure 8: Example of prompt and ground truth for description generation task.

Prompt

The structure of the crystal material is <crys>. Please answer this question in 100 words according to the structure of this crystal material. Question: What advantages do the new zinc-based metal-organic frameworks (MOFs) offer in terms of environmental impact and synthesis?

Ground-truth

The new zinc-based MOFs can be easily synthesized in water, providing an environmentally friendly anti-corrosion treatment for metals, which aligns with sustainable practices in material science.

Figure 9: Example of prompt and ground truth for question&answer task.

Crystallographic Information File(CIFs)

```

_cell_chemical_formula_moiety      '(C72 H48 Cu4 N24 O4 S1 2+)(O4 S1 2-)n'
_cell_chemical_name_systematic    'catena-(tetrakis(m-2,4,6-tris(4-pyridyl)-1,3,5-triazine)-(m-sulfato)-tetra-copper sulfate unknown solvate)'
_symmetry_cell_setting           'cubic'
_symmetry_space_group_name_H-M   'F 2 3'
_symmetry_int_Tables_number     '196'
_space_group_name_Hall          'F 2 2 3'
loop_
_symmetry_equiv_pos_site_id
_symmetry_equiv_pos_as_xyz
 1 x,y,z
 2 z,x,y
 3 y,z,x
.....
 48 1/2-x,1/2+y,-z
_cell_length_a                  21.1951(6)
_cell_length_b                  21.1951(6)
_cell_length_c                  21.1951(6)
_cell_angle_alpha               90
_cell_angle_beta                90
_cell_angle_gamma               90
_cell_volume                    9521.52
_exptl_crystal_colour          'dark black'
_exptl_crystal_description     'octahedral'
_cell_formula_units_Z           4
loop_
_atom_site_label
_atom_site_type_symbol
_atom_site_fract_x
_atom_site_fract_y
_atom_site_fract_z
Cu1 Cu 0.65154(3) 0.34846(3) 0.84846(3)
S1 O 0.7500 0.2500 0.7500
N1 N 0.58469(18) 0.36418(17) 0.78406(19)
N2 N 0.4334(2) 0.3429(2) 0.6036(2)
.....
Cu12 Cu 0.84846(3) -0.15154(3) 1.15154(3)
loop_
_geom_bond_atom_site_label_1
_geom_bond_atom_site_label_2
_geom_bond_site_symmetry_1
_geom_bond_site_symmetry_2
Cu1 N1 1_555 1_555
Cu1 O1 1_555 1_555
Cu1 N19 1_555 1_555
S1 O1 1_555 1_555
.....
C71 H48 1_555 1_555
#END

```

Figure 10: Example of crystallographic information file.

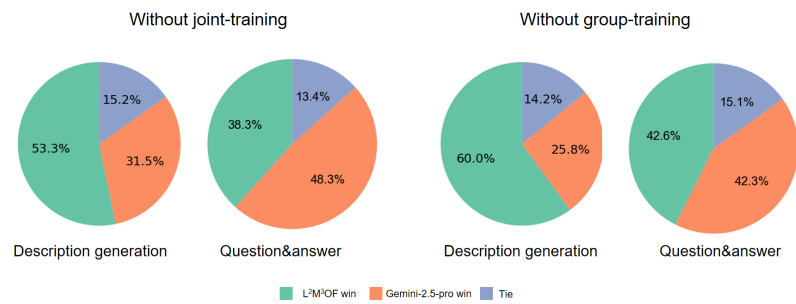


Figure 11: Performance comparison of Gemini-2.5-pro and L²M³OF in the ablation study.