
LLM4Mat-Bench: Benchmarking Large Language Models for Materials Property Prediction

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Abstract

Large language models (LLMs) are increasingly being used in materials science. However, little attention has been given to benchmarking and standardized evaluation for LLM-based materials property prediction, which hinders progress. We present LLM4Mat-Bench, the largest benchmark to date for evaluating the performance of LLMs in predicting the properties of crystalline materials. LLM4Mat-Bench contains about 1.9M crystal structures in total, collected from 10 publicly available materials data sources, and 45 distinct properties. LLM4Mat-Bench features different input modalities: crystal composition, CIF, and crystal text description, with 4.7M, 615.5M, and 3.1B tokens in total for each modality, respectively. We use LLM4Mat-Bench to fine-tune models with different sizes, including LLM-Prop and MatBERT, and provide zero-shot and few-shot prompts to evaluate the property prediction capabilities of LLM-chat-like models, including Llama, Gemma, and Mistral. The results highlight the challenges of general-purpose LLMs in materials science and the need for task-specific predictive models and task-specific instruction-tuned LLMs in materials property prediction ¹.

1 Introduction

With the remarkable success of large language models (LLMs) in solving natural language tasks [12, 31, 32, 1, 39] and different scientific tasks [26, 14, 40, 4, 15, 27], scientists have recently started to leverage LLMs to tackle very important and challenging problems in materials science, including predicting materials properties [35, 24, 44, 11, 30, 6] and discovering new materials [2, 19, 30, 6].

The learning capabilities of LLMs have the potential to revolutionize the field of materials science. For example, recent research by Rubungo et al. [35] has demonstrated the exceptional performance of LLMs in predicting the properties of crystalline materials based on textual descriptions of their structures. In their study, they introduced a novel dataset, TextEdge, which comprises textual descriptions of crystals and their corresponding properties. This dataset was used to fine-tune the encoder component of the T5-small model for the task of materials property prediction. The findings of Rubungo et al. [35] challenge the conventional practice of heavily relying on graph neural networks and using solely either crystal composition or structure as input for property prediction. Their work underscores the significance of further investigating the extent to which LLMs can be harnessed to

¹The Benchmark and code can be found at: <https://github.com/vertaix/LLM4Mat-Bench>

develop innovative techniques for accurately predicting the properties of crystalline materials, thereby enhancing the materials discovery pipeline. Unfortunately, the proposed TextEdge dataset is limited in scope, comprising approximately 145K samples and encompassing only three distinct properties. Furthermore, its lack of diversity, being derived from a single data source (Materials Project [21]), hinders its effectiveness in assessing the robustness of LLMs in materials property prediction.

In this work, we introduce LLM4Mat-Bench, a benchmark dataset collected to evaluate the performance of LLMs in predicting the properties of crystalline materials. To the best of our knowledge, LLM4Mat-Bench is the most extensive benchmark to date for assessing the efficacy of language models in materials property prediction. The dataset comprises approximately two million samples, sourced from ten publicly available materials sources, each containing between 10K and 1M structure samples. LLM4Mat-Bench encompasses several tasks, including the prediction of electronic, elastic, and thermodynamic properties based on a material’s composition, crystal information file (CIF), or textual description of its structure. We use LLM4Mat-Bench to evaluate several LLMs of different sizes, namely LLM-Prop [35] (35M parameters), MatBERT [41] (109.5M parameters), and Llama 2 [39] (7B parameters). And we provide fixed train-valid-test splits, along with carefully designed zero-shot and few-shot prompts to ensure reproducibility. We anticipate that LLM4Mat-Bench will significantly advance the application of LLMs in addressing critical challenges in materials science, including property prediction and materials discovery.

2 LLM4Mat-Bench

2.1 Data Collection Process

We collected the data used to create LLM4Mat-Bench from 10 publicly available materials data sources. In this section, we describe each data source and discuss how we accessed its data.

2.1.1 Data sources

hMOF [42] is a publicly available database² consisting of about 160K Metal-Organic Frameworks (MOFs), generated by Wilmer et al. using computational approaches. Materials Project (MP) [21] is a database with around 150K materials, offering free API access³ for data retrieval, including CIF files and material properties. The Open Quantum Materials Database (OQMD) [23] is a publicly accessible database⁴ of 1.2M materials, containing DFT-calculated thermodynamic and structural properties, created at Northwestern University. OMDb [3] is an organic materials database with about 12K structures and related electronic band structure properties, freely available⁵. JARVIS-DFT [8, 9] is a repository created by NIST researchers, containing around 75.9K material structures with downloadable properties⁶. QMOF [33, 34] is a quantum-chemical property database of over 16K MOFs, accessible via GitHub⁷. JARVIS-QETB [17] is a NIST-created database⁸ of nearly one million materials with tight-binding parameters for 65 elements. GNoME is a database of 381K new stable materials discovered by Merchant et al. [28] using graph networks and DFT, available on GitHub⁹. Cantor HEA [25] is a DFT dataset of formation energies for 84K alloy structures, available on Zenodo¹⁰. SNUMAT is a database with around 10K experimentally synthesized materials and DFT properties, accessible via API¹¹.

2.1.2 Collecting crystal information files (CIFs) and materials property

Crystal structure files (CIFs), material compositions, and material properties were collected from publicly accessible sources described in Section 2.1.1. Data collection was facilitated by APIs and

²<https://mof.tech.northwestern.edu/>

³<https://next-gen.materialsproject.org/api>

⁴<https://www.oqmd.org/>

⁵<https://omdb.mathub.io/>

⁶<https://jarvis.nist.gov/jarvisdft>

⁷<https://github.com/Andrew-S-Rosen/QMOF>

⁸<https://jarvis.nist.gov/jarvisqetb/>

⁹https://github.com/google-deepmind/materials_discovery/blob/main/DATASET.md

¹⁰<https://doi.org/10.5281/zenodo.10854500>

¹¹<https://www.snumat.com/apis>

direct download links provided by the respective databases. For databases such as Materials Project, OMDb, SNUMAT, JARVIS-DFT, and JARVIS-QETB, user registration is required for access, while databases like hMOF, QMOF, OQMD, and GNoME allow direct data access without registration. From each source, we obtained CIFs and associated material properties. Although the Materials Project and JARVIS-DFT databases offer a broader range of properties, we selected a subset—10 and 20 properties respectively—that adequately represents the data within our benchmark, based on the number of data points available for each property. This selection was made to optimize computational efficiency when training models across the 65 properties included in LLM4Mat-Bench.

2.1.3 Generating the textual description of crystal structure

LLMs perform better with textual input, and Rubungo et al. [35], Korolev and Protsenko [24], Qu et al. [30] have demonstrated that LLMs can effectively learn the structural representation of a crystal from its textual description, outperforming graph neural network (GNN)-based models that directly utilize the crystal structure for property prediction. Crystal structures are typically described in file formats such as Crystallographic Information File (CIF) which include predominantly numbers describing lattice vectors and atomic coordinates and are less amenable to LLMs. Instead of directly using these as inputs, we use Robocrystallographer [21] to deterministically generate texts that are more descriptive of crystal structures from CIF files. Robocrystallographer was developed and has been used by the Materials Project team to auto-generate texts for their database. Given a structure, Robocrystallographer leverages predefined rules and existing libraries to extract chemical and structural information, including oxidation states, global structural descriptions (symmetry information, prototype matching, structural fingerprint calculations etc.), and local structural descriptions (e.g. bonding and neighbor analysis, connectivity). This method not only generate deterministic and human-readable texts, but also ensures no data contamination in our fine-tuned LLMs, as the data sources mentioned do not include these crystal text descriptions.

Table 1: LLM4Mat-Bench statistics.

Data source	# Structure files	# Structure-Description pairs				# Tokens (Words)			# Avg. subword tokens/Sample			# Properties
		Total	Train	Validation	Test	Composition	Structure	Description	Composition	Structure	Description	
OQMD [23]	1,008,266	964,403	771,522	96,440	96,441	964K	96M	244M	5.3	635.4	347.3	2
JARVIS-QETB [17]	829,576	623,989	499,191	62,399	62,399	624K	45M	90M	3.5	466.6	202.5	4
GNoME [28]	381,000	376,276	301,020	37,628	37,628	830K	78M	508M	9.7	1185.3	1711.3	6
Materials Project [21]	146,143	125,825	100,000	10,000	10,318	272K	37M	157M	6.8	1611.8	1467.3	10
hMOF [42]	133,524	132,743	106,194	13,274	13,275	449K	96M	581M	14.9	4583.9	5629.3	7
Camtore HEA [25]	84,024	84,019	67,215	8,402	8,402	84K	11M	251M	9.5	868.4	4988.6	4
JARVIS-DFT [8, 9]	75,965	75,965	60,772	7,596	7,597	76K	9M	25M	5.0	786.0	455.9	20
QMOF [33, 34]	16,340	7,656	6,124	766	766	8K	7M	22M	14.0	5876.4	3668.0	4
OMDB [3]	12,500	12,122	9,697	1,212	1,213	66K	8M	14M	14.8	4097.4	1496.6	1
SNUMAT ¹²	10,481	10,372	8,297	1,037	1,038	16K	2M	4M	5.9	1244.5	539.1	7
Total	2,697,779	1,978,985	1,592,315	193,357	193,313	4.7M	615.5M	3.1B	7.9	1559.7	1703.6	65

Table 2: Comparing the LLM4Mat-Bench with other existing benchmarks.

Benchmark	# Data Sources	# Distinct Properties	# Properties/# Samples			# Properties/Task Type		Material Representations		
			<10k	10-100k	100k+	Regression	Classification	Composition	Structure	Description
MatBench [13]	6	10	7	3	3	10	3	✓	✓	✗
TextEdge [35]	1	3	0	0	3	2	1	✗	✗	✓
LLM4Mat-Bench (Ours)	10	45	5	31	29	60	5	✓	✓	✓

2.2 Data Statistics

As Table 1 shows, LLM4Mat-Bench comprises 2,697,779 structure files, which, after pairing with descriptions generated by Robocrystallographer and filtering out descriptions with fewer than five words, result in 1,978,985 composition-structure-description pairs¹³. The reduction in sample count is also due to Robocrystallographer’s inability to describe certain CIF files. The total samples for each dataset in LLM4Mat-Bench are randomly split into 80%, 10%, and 10% for training, validation, and testing, respectively. OQMD has the highest number of samples at 964,403, while QMOF has the fewest with 7,656 samples. On average, each dataset in LLM4Mat-Bench contains approximately 200,000 samples.

In LLM4Mat-Bench, when combined, textual descriptions contain 3.1 billion tokens, crystal structures 615 million, and compositions 4.7 million¹⁴. OQMD leads in composition tokens (964K), while

¹³The total number of pairs were 2,433,688, after removing about 454703 duplicated pairs across datasets, it resulted to 1,978,985 pairs.

¹⁴We used NLTK toolkit as a tokenizer to count the number of words/tokens.

hMOF has the most description tokens (581M). For CIFs, both OQMD and hMOF have around 96M tokens. On average, compositions have 8 subword tokens per sample, CIFs 1600, and descriptions 1700. hMOF averages the longest inputs for compositions (14.9) and descriptions (5629), while QMOF leads in structures (5876.4)¹⁵. JARVIS-DFT has the most tasks with 20 properties, followed by Materials Project with 10, and OMDB with one. Details on sample counts are in Section 3.2.

LLM4Mat-Bench provides the most comprehensive dataset compared to existing benchmarks, with the largest number of samples, properties, and tasks, including 60 regression and 5 classification tasks (see Table 2). It also offers more diverse material representations, incorporating chemical formulas, crystal structures, and crystal text descriptions. In contrast, MatBench [13] and TextEdge [35] have fewer tasks and less representation diversity, with MatBench lacking crystal text descriptions and TextEdge missing material compositions and crystal structures.

2.3 Data Quality

Since Robocrystallographer generates crystal text descriptions in a deterministic manner following predefined and well-validated rules [21], these texts should faithfully describe the crystal structures used to generate them. Regarding the quality of labels, they are calculated from simulations and are usually considered noise-free. Properties data except those from JARVIS-QETB and hMOF are obtained from DFT, which is based on fundamental quantum mechanical equations. While DFT calculations can still be performed with different levels of approximations and fidelity, the DFT-calculated properties are usually considered to be highly reliable and are routinely used as noise-free ground truths for ML models in the materials science community.

3 Results

3.1 Experimental Details

We conducted about 845 experiments, evaluating the performance of five models and three material representations on each property for each data source. Consistent with standard practices in materials science, we evaluated performance separately for each data source rather than combining samples from different sources for the same property. This approach accounts for variations in techniques and settings used by different data sources, which can result in discrepancies, such as differing band gaps for the same material. Below, we will describe each material representation, model, and metric that we used to conduct our experiments.

3.1.1 Material Representations

LLM4Mat-Bench includes three distinct materials representations: Composition, CIF, and Description (see Table 6). The primary goal of using these diverse representations is to identify which best enhances LLM performance in predicting material properties across different data sources.

Composition (Comp.) Material composition refers to the chemical formula of a material. Though it only provides stoichiometric information, studies have shown it can still be a reliable material representation for property prediction [13, 38]. For LLMs, it offers the advantage of being a short sequence that usually fits within the model’s context window, making it efficient to train. To further optimize efficiency, we set the longest sequence of material compositions from each data source as the context window, rather than using the default 512 tokens for fine-tuning while the original length is kept during inference.

CIF We represent the materials structure using CIF files, the conventional way of representing the crystal structure in crystallography [20]. CIFs are commonly used for GNN-based models, but some recent works have demonstrated that it can also work with LLMs [2, 16, 19].

Description (Descr.) As we outlined in Section 2.1.3, we also use textual descriptions of crystal structures as representations for both atomic crystals and MOFs.

¹⁵We used Llama 2 tokenizer to count the number of subword tokens.

3.1.2 Models

We benchmarked different LLM-based models with various sizes, and a GNN-based baseline. Herein, We provide the details of each model.

CGCNN [43] is employed as a GNN baseline which is widely used in the materials science community¹⁶. We trained on LLM4Mat-Bench from scratch with optimal hyperparameters: 128 hidden dimensions, batch size of 256, three message passing layers, 1e-2 learning rate, 8.0 radius cutoff, 12 nearest neighbors, and 500 training epochs, though extending to 1000 epochs improved performance in some cases.

MatBERT [41] is a BERT-base model [12] with 109 million parameters, pretrained on two million materials science articles. We fine-tuned MatBERT on LLM4Mat-Bench, following Rubungo et al. [35], and achieved optimal performance with a 512-token input length, 64-sample batch size, 5e-5 learning rate, 0.5 dropout, and 100 epochs using the Adam optimizer and one-cycle learning rate scheduler [36]. Although training for 200 epochs improves performance, results are reported for 100 epochs due to computational constraints.

LLM-Prop is a model based on the encoder part of T5-small model [32] introduced by Rubungo et al. [35], with 35 million parameters, smaller than MatBERT. It predicts material properties from the textual descriptions of crystal structures. To adapt LLM-Prop on CIF, we employed xVal encoding [18], where we parse an input sequence x to extract numerical values into a list x_{num} , replace them with a [NUM] token to form x_{text} , and then embed x_{text} , followed by multiplying each [NUM] embedding by its corresponding value in x_{num} to get h_{embed} that we feed to the model. xVal encoding ensures that the quantitative value of each number is reflected in the input embedding while reducing the input length caused by the high volume of numerical values in CIF files, which extend the length of the input sequence after tokenization. We fine-tuned LLM-Prop on LLM4Mat-Bench and optimizing with a 1e-3 learning rate, 0.2 dropout, Adam optimizer, and one-cycle learning rate scheduler for 100 epochs, with a 768-token input length, batch size of 64 for training, and 512 for inference. While Rubungo et al. [35] recommended that training for 200 epochs and increasing the number of input tokens improves the performance, we could not replicate this due to computational constraints.

Table 3: Prompt template. <material representation type> denotes “chemical formula”, “cif structure”, or “structure description”. <value> represents the input context (for example *NaCl*, etc.). <property name> denotes the name of the property (for example *band gap*, etc.). <predicted value> represents the property value generated by Llama 2 while <actual value_i> represents the ground truth of the **EXAMPLE_i**. **FINAL PROMPT** and **RESPONSE** denote the input prompt to Llama 2 and its generated output, respectively.

Prompt Type	Template
-	SYSTEM PROMPT: «SYS» You are a material scientist. Look at the <material representation type> of the given crystalline material and predict its property. The output must be in a json format. For example: {property_name:predicted_property_value}. Answer as precise as possible and in as few words as possible. «/SYS»
	INPUT PROMPT: <material representation type>: <value> property name: <property name>.
0-shot	FINAL PROMPT: <s>[INST] + SYSTEM PROMPT + INPUT PROMPT + [/INST] RESPONSE: <property name>:<predicted value>
5-shot	EXAMPLE₁: <material representation type>: <value ₁ > property name: <property name>. <property name>:<actual value ₁ >
	... EXAMPLE₅: <material representation type>: <value ₅ > property name: <property name>. <property name>:<actual value ₅ >
	FINAL PROMPT: <s>[INST] + SYSTEM PROMPT + EXAMPLE₁ + ... + EXAMPLE₅ + INPUT PROMPT + [/INST] RESPONSE: <property name>:<predicted value>

Llama 2-7b-chat To assess the performance of conversational LLMs in materials property prediction, we tested Llama 2-7b-chat (7 billion parameters) using our designed zero-shot and five-shot prompts (see Table 3) without fine-tuning. For the CIF structure prompts, we removed “# generated using

¹⁶Although CGCNN is not state-of-the-art for some properties, it was faster compared to models like ALIGNN [7] and DeeperGatGNN [29], making it suitable for our extensive experiments

pymatgen" comment that is appended to each file. The maximum input length was set to 4000 tokens while the output length was set to 256, with a batch size of 256 samples, temperature of 0.8, and top-K sampling applied with $K = 10$. The details of other models that we compared with Llama 2 can be found in Appendix C. For five-shot examples, we sampled from crystals with shorter structures and descriptions to reduce the context length. We also made sure the property values for those examples are diverse (for instance, they should not all have 0.0 eV as their bandgap values).

We trained all models using NVIDIA RTX A6000 GPUs. Training MatBERT with two GPUs on about 300K data points and 100 epochs took about four days while for LLM-Prop took about 2.5 days. For CGCNN, it took about 7 hours training time on one GPU for 500 epochs. With one GPU, Llama 2 took about a half day to generate the output of 40K samples with 256 tokens maximum length each. We report the test set results averaged over five runs for predictive models and three runs for generative models.

3.1.3 Evaluation Metrics

Following Choudhary and DeCost [7], we evaluated regression tasks using the ratio between the mean absolute deviation (MAD) of the ground truth and the mean absolute error (MAE) of the predicted properties. The MAD:MAE ratio ensures an unbiased model comparison between different properties where the higher ratio the better. According to Choudhary and DeCost [7], a good predictive model should have at least 5.0 ratio. MAD values represent the performance of a random guessing model predicting the average value for each data point. To provide a comprehensive performance comparison across datasets, we also reported the weighted average of MAD:MAE across all properties in each dataset (Wtd. Avg. (MAD:MAE), see Equation 1).

For classification tasks, we reported the area under the ROC curve (AUC) for each task and provided the weighted average across all properties (Wtd. Avg. AUC, see Equation 2).

$$\text{Wtd. Avg. (MAD/MAE)} = \frac{\sum_i^m \text{TestSize}_i \times \frac{\text{MAD}_i}{\text{MAE}_i}}{\sum_i^m \text{TestSize}_i} \quad (1)$$

$$\text{Wtd. Avg. AUC} = \frac{\sum_i^m \text{TestSize}_i \times \text{AUC}_i}{\sum_i^m \text{TestSize}_i}, \quad (2)$$

m denotes the number of regression properties in the dataset.

Table 4: The Wtd. Avg. (MAD:MAE) scores (the higher the better) for the regression tasks in the LLM4Mat-Bench are reported. **Bolded** results indicate the best model for each input format, while **bolded results with blue background** highlight the best model per each dataset. **Inval.** denotes cases where the Llama model failed to generate outputs with a property value or had fewer than 10 valid predictions.

Input	Model	Dataset									
		MP 8 tasks	JARVIS-DFT 20 tasks	GNoME 6 tasks	hMOF 7 tasks	Cantor HEA 4 tasks	JARVIS-QETB 4 tasks	OQMD 2 tasks	QMOF 4 tasks	SNUMAT 4 tasks	OMDB 1 task
CIF	CGCNN (baseline)	5.319	7.048	19.478	2.257	17.780	61.729	14.496	3.076	1.973	2.751
Comp.	Llama 2-7b-chat:0S	0.389	Inval.	0.164	0.174	0.034	0.188	0.105	0.303	0.940	0.885
	Llama 2-7b-chat:5S	0.627	0.704	0.499	0.655	0.867	1.047	1.160	0.932	1.157	1.009
	MatBERT-109M	5.317	4.103	12.834	1.430	6.769	11.952	5.772	2.049	1.828	1.554
	LLM-Prop-35M	4.394	2.912	15.599	1.479	8.400	59.443	6.020	1.958	1.509	1.507
CIF	Llama 2-7b-chat:0S	0.392	0.216	6.746	0.214	0.022	0.278	0.028	0.119	0.682	0.159
	Llama 2-7b-chat:5S	Inval.	Inval.	Inval.	Inval.	Inval.	1.152	1.391	Inval.	Inval.	0.930
	MatBERT-109M	7.452	6.211	14.227	1.514	9.958	47.687	10.521	3.024	2.131	1.777
	LLM-Prop-35M	8.554	6.756	16.032	1.623	15.728	97.919	11.041	3.076	1.829	1.777
Descr.	Llama 2-7b-chat:0S	0.437	0.247	0.336	0.193	0.069	0.264	0.106	0.152	0.883	0.155
	Llama 2-7b-chat:5S	0.635	0.703	0.470	0.653	0.820	0.980	1.230	0.946	1.040	1.001
	MatBERT-109M	7.651	6.083	15.558	1.558	9.976	46.586	11.027	3.055	2.152	1.847
	LLM-Prop-35M	9.116	7.204	16.224	1.706	15.926	93.001	9.995	3.016	1.950	1.656

3.2 Discussion

Table 4 and 5, and Figure 1 and 2 show the main results. The detailed results on each dataset can be found in Appendix E. The main observations are as follows:

Table 5: The Wtd. Avg. AUC scores (the higher the better) for the classification tasks in the LLM4Mat-Bench.

Input	Model	Dataset	
		MP 2 tasks	SNUMAT 3 tasks
CIF	CGCNN (baseline)	0.846	0.722
Comp.	Llama 2-7b-chat:0S	0.491	Invalid.
	Llama 2-7b-chat:5S	0.507	0.466
	MatBERT-109M	0.722	0.712
	LLM-Prop-35M	0.691	0.716
CIF	Llama 2-7b-chat:0S	0.501	0.489
	Llama 2-7b-chat:5S	0.502	0.474
	MatBERT-109M	0.750	0.717
	LLM-Prop-35M	0.738	0.660
Descr.	Llama 2-7b-chat:0S	0.500	Invalid.
	Llama 2-7b-chat:5S	0.502	0.568
	MatBERT-109M	0.735	0.730
	LLM-Prop-35M	0.742	0.735

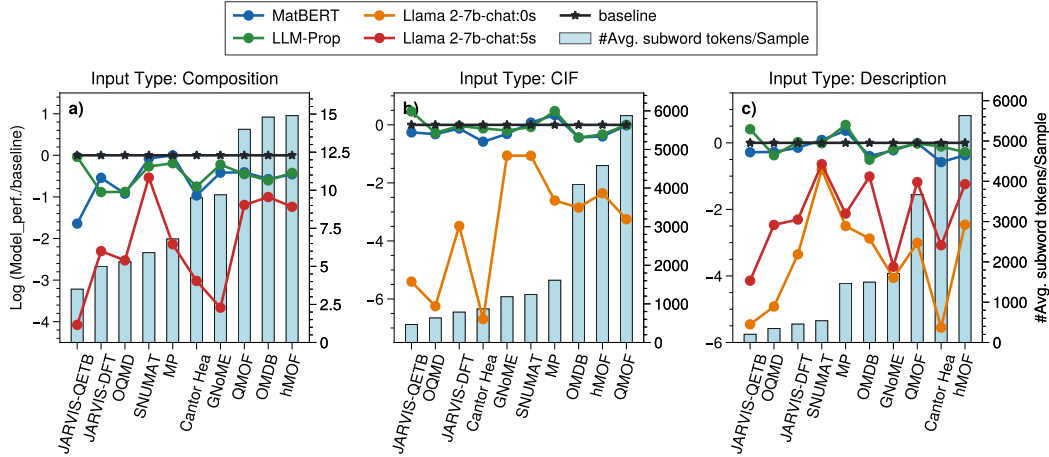


Figure 1: The performance comparison across models for each material representation is presented. The left y-axis shows the log-normalized performance of each LLM-based model relative to the baseline (CGCNN), while the right y-axis (bar plots) displays the average subword tokens per sample for each dataset. Datasets on the x-axis are ordered by increasing average subword tokens. Results for Llama 2-7b-chat:0S and Llama 2-7b-chat:5S are missing in plots (a) and (b), respectively, due to invalid outputs. Higher values in the line plots indicate better performance.

Small, task-specific predictive LLMs exhibit significantly better performance than larger, generative general-purpose LLMs. This performance disparity is evident across both regression (Table 4 and Figure 1) and classification tasks (Table 5) on all 10 datasets. Specifically, LLM-Prop and MatBERT outperform Llama 2-7b-chat:0S and Llama 2-7b-chat:5S by a substantial margin, despite being approximately 200 and 64 times smaller in size, respectively. In regression tasks, LLM-Prop achieves the highest accuracy on 8 out of 10 datasets, with MatBERT leading on the remaining 2 datasets. For classification tasks, both LLM-Prop and MatBERT deliver the best performance on 1 out of 2 datasets. LLM-Prop surpasses MatBERT by 1.8% on the SNUMAT dataset, whereas MatBERT outperforms LLM-Prop by 0.8% on the other dataset. As anticipated, a modest enhancement in average performance is observed across various datasets and input formats when the Llama 2-7b-chat

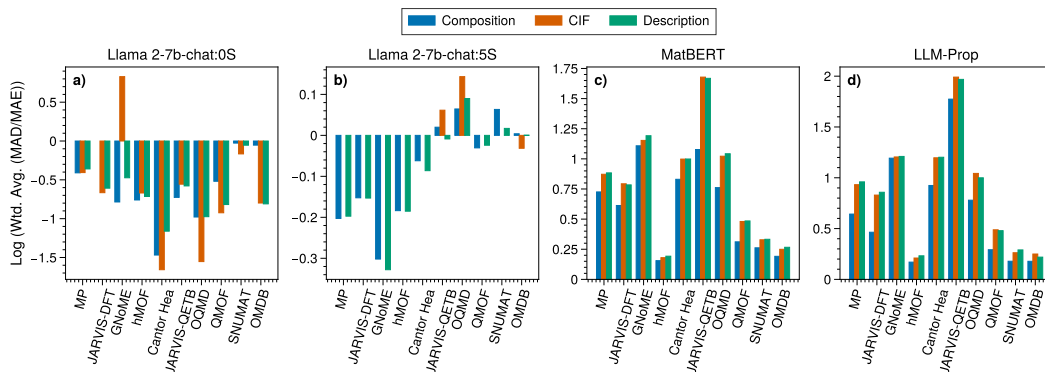


Figure 2: The performance comparison across material representations for each LLM-based model is shown. The y-axis represents the log-normalized Weighted Average (MAD/MAE) score for each representation, while the x-axis displays randomly ordered datasets. In the (a) and (b) plots, some Composition and Structure performance results are missing due to invalid outputs. A higher y-axis value indicates better performance.

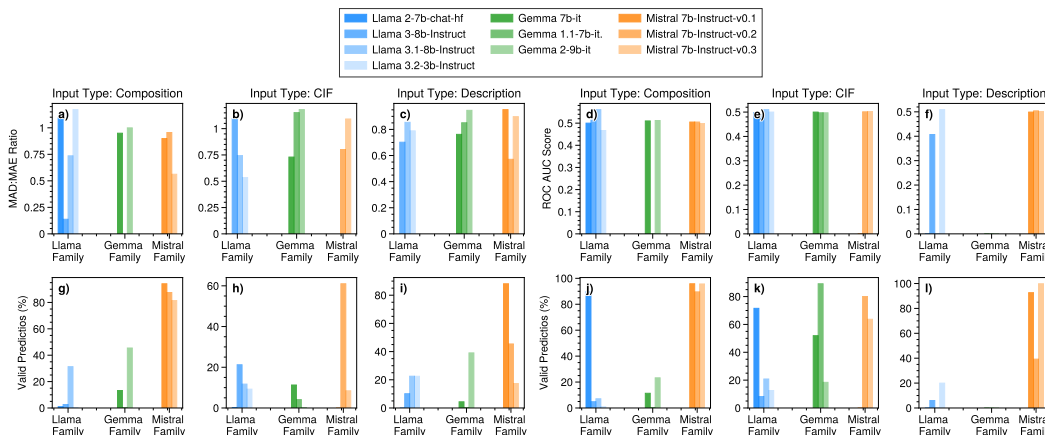


Figure 3: The performance comparison of different chat-based LLM versions is presented with results based on 5-shot prompts, averaged over three inference runs. Panels (a)–(c) and (d)–(f) show each model’s accuracy in predicting band gaps and stability in the MP dataset, respectively, while panels (g)–(i) and (j)–(l) depict the percentage of valid predictions for band gap and stability on the test set.

model is evaluated using 5-shot prompts rather than 0-shot prompts. Determining the optimal number of examples required to achieve peak performance will be the focus of future work.

General-purpose generative LLMs hallucinate and often fail to generate valid property values. As shown in Table 4, Table 5, Figure 2, and Appendix E, Llama 2-7b-chat model produces invalid outputs on multiple tasks, where the expected property value is missing. This issue occurs less frequently when the input is a description or chemical formula, but more commonly when the input is a CIF file. One reason may be that descriptions and chemical formulas resemble natural language, which LLMs can more easily interpret compared to CIF files. This may also indicate that when the input modality during inference differs significantly from the modalities encountered during pretraining, fine-tuning is necessary to achieve reasonable performance. Another key observation is that Llama 2-7b-chat model often generates the same property value for different inputs (i.e. hallucinates), contributing to its poor performance across multiple tasks. These findings highlight the importance of caution when using general-purpose generative LLMs for materials property prediction and emphasize the need for fine-tuned, task-specific LLM-based models.

Representing materials with their textual descriptions improves the performance of LLM-based property predictors compared to other representations. We observe a significant performance

improvement when the input is a description compared to when it is a CIF file or a chemical formula. One of the possible reasons for this might be that LLMs are more adept at learning from natural language data. On the other hand, although material compositions appear more natural to LLMs compared to CIF files, they lack sufficient structural information. This is likely why LLMs with CIF files as input significantly outperform those using chemical formulas.

More advanced, general-purpose generative LLMs do not necessarily yield better results in predicting material properties. In Figure 3, we compare the performance of Llama 2-7b-chat-hf model with advanced versions of Llama of comparable sizes when predicting material’s band gap and its stability. Similar comparisons are also conducted for the Mistral [22] and Gemma [37] models. The results indicate that, despite being trained on substantially larger and higher-quality datasets, more advanced versions of generative LLMs show limited improvements in performance and validity of predictions for material properties. For instance, Llama 3 and 3.1 8b models were trained on over 15 trillion tokens—around eight times more data than the 2 trillion tokens used for the Llama 2 7b models. This finding highlights the ongoing challenges of leveraging LLMs in material property prediction and underscores the need for further research to harness the potential of these robust models in this domain.

The performance on energetic properties is consistently better across all datasets compared to other properties. This is consistent with the trend observed in the community benchmarks such as MatBench and JARVIS-Leaderboard, where energetic properties are among those that can be most accurately predicted [13, 10]. This is not surprising because energy is known to be relatively well predicted from e.g., compositions and atom coordination (bonding), which is inherently represented in GNNs and also presented in text descriptions.

Task-specific predictive LLM-based models excel with shorter textual descriptions, while CGCNN performs better on datasets with longer descriptions. While the focus on this work is on LLMs, a comparison with a simple but widely used GNN-based baseline suggests room for improvement in LLM-based property prediction. For regression tasks, LLM-Prop outperforms CGCNN on only 4 out of 10 datasets (MP, JARVIS-DFT, JARVIS-QETB, and SNUMAT), and MatBERT outperforms CGCNN on just 2 out of 10 datasets (MP and JARVIS-QETB). In contrast, CGCNN achieves the best performance on 5 out of 10 datasets (GNoME, hMOF, Cantor HEA, OQMD, and OMDb). Further analysis reveals that CGCNN tends to perform better than LLM-based models on datasets with relatively longer textual descriptions, while LLM-based models excel on datasets with shorter descriptions (see Table 1). The performance gain on shorter descriptions may stem from LLM-based models’ ability to leverage more context from compact text, while CGCNN consistently benefits from training on the entire crystal structure.

4 Conclusion

LLMs are increasingly being utilized in materials science, particularly for materials property prediction and discovery. However, the absence of standardized evaluation benchmarks has impeded progress in this field. We introduced LLM4Mat-Bench, a comprehensive benchmark dataset designed to evaluate LLMs for predicting properties of atomic and molecular crystals and MOFs. Our results demonstrate the limitations of general-purpose LLMs in this domain and underscore the necessity for task-specific predictive models and instruction-tuned LLMs tailored for materials property prediction. These findings emphasize the importance of using LLM4Mat-Bench to advance the development of more effective LLMs in materials science.

5 Limitations

Due to computational constraints and the number of experiments, we were unable to conduct thorough hyperparameter searches for each property and dataset. The reported settings were optimized on the MP dataset and then fixed for other datasets. For each model, we highlighted hyperparameter settings that may improve performance (see Section 3.1.2). Additionally, we could not include results from SOTA commercial LLMs such as GPT-4o¹⁷ or Claude 3.5 Sonnet¹⁸ due to budget constraints.

¹⁷<https://openai.com/index/hello-gpt-4o/>

¹⁸<https://www.anthropic.com/news/claude-3-5-sonnet>

We also encountered issues with chat-based models, which sometimes failed to follow the output format, producing invalid or incomplete outputs. Extracting property values was therefore challenging. We believe further instruction-tuning chat-based models on the provided prompts could mitigate these issues.

Furthermore, we did not include comparisons with dataset-specific retrieval-augmented generation (RAG) models, such as the recently developed LLaMP [5], a RAG-based model tailored for interaction with the MP dataset. Our work aims to provide a comprehensive benchmark and baseline results to advance the evaluation of LLM-based methods for materials property prediction. Future work should address these limitations.

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Appendices

A Materials Representations

Table 6: LLM4Mat-Bench material representations of Sodium Chloride (NaCl).

Crystal Information File (CIF)						
<pre># generated using pymatgen data_NaCl _symmetry_space_group_name_H-M 'P 1' _cell_length_a 3.50219000 _cell_length_b 3.50219000 _cell_length_c 3.50219000 _cell_angle_alpha 90.00000000 _cell_angle_beta 90.00000000 _cell_angle_gamma 90.00000000 _symmetry_Int_Tables_number 1 _chemical_formula_structural NaCl _chemical_formula_sum 'Na1 Cl1' _cell_volume 42.95553287 _cell_formula_units_Z 1 loop_ _symmetry_equiv_pos_site_id _symmetry_equiv_pos_as_xyz 1 'x, y, z' loop_ _atom_type_symbol _atom_type_oxidation_number Na+ 1.0 Cl- -1.0 loop_ _atom_site_type_symbol _atom_site_label _atom_site_symmetry_multiplicity _atom_site_fract_x _atom_site_fract_y _atom_site_fract_z _atom_site_occupancy Na+ Na0 1 0.00000000 0.00000000 0.00000000 1 Cl- Cl1 1 0.50000000 0.50000000 0.50000000 1</pre>						
Description						
<p>NaCl is Tetraauricupride structured and crystallizes in the cubic $P\bar{m}3m$ space group. Na^{1+} is bonded in a body-centered cubic geometry to eight equivalent Cl^{1-} atoms. All Na-Cl bond lengths are 3.03 Å. Cl^{1-} is bonded in a body-centered cubic geometry to eight equivalent Na^{1+} atoms.</p>						

B Statistics of All Properties in LLM4Mat-Bench

Table 7: Statistics of all datasets in LLM4Mat-Bench. It is important to note that we retain the naming convention of each property from the original data source with the intent to provide the distribution of properties in each dataset.

Property	Task type	# Samples/Data source									
		JARVIS-DFT	Materials Project	SNUMAT	hMOF	GNoME	JARVIS-QETB	Cantor HEA	QMOF	OQMD	OMDB
Bandgap	Regression	-	145,302	-	-	288,209	-	-	16,340	1,007,324	12,500
Bandgap (OPT)	Regression	75,965	-	-	-	-	-	-	-	-	-
Bandgap (MBJ)	Regression	19,800	-	-	-	-	-	-	-	-	-
Bandgap GGA	Regression	-	-	10,481	-	-	-	-	-	-	-
Bandgap HSE	Regression	-	-	10,481	-	-	-	-	-	-	-
Bandgap GGA Optical	Regression	-	-	10,481	-	-	-	-	-	-	-
Bandgap HSE Optical	Regression	-	-	10,481	-	-	-	-	-	-	-
Indirect Bandgap	Regression	-	-	-	-	-	829,576	-	-	-	-
Formation Energy Per Atom (FEPA)	Regression	75,965	145,262	-	-	384,871	829,576	84,024	-	1,008,266	-
Energy Per Atom (EPA)	Regression	-	145,262	-	-	-	829,576	84,024	-	-	-
Decomposition Energy Per Atom (DEPA)	Regression	-	-	-	-	384,871	-	-	-	-	-
Energy Above Hull (Ehull)	Regression	75,965	145,262	-	-	-	-	84,024	-	-	-
Total Energy	Regression	75,965	-	-	-	384,871	829,576	-	16,340	-	-
Elfermi	Regression	-	145,262	-	-	-	-	-	-	-	-
Exfoliation Energy	Regression	812	-	-	-	-	-	-	-	-	-
Bulk Modulus (Kv)	Regression	23,823	-	-	-	-	-	-	-	-	-
Shear Modulus (Gv)	Regression	23,823	-	-	-	-	-	-	-	-	-
SLME	Regression	9,765	-	-	-	-	-	-	-	-	-
Spillage	Regression	11,377	-	-	-	-	-	-	-	-	-
ϵ_r (OPT)	Regression	52,158	-	-	-	-	-	-	-	-	-
ϵ (DFPT)	Regression	4,704	-	-	-	-	-	-	-	-	-
Max. piezoelectri c strain coeff (dij)	Regression	3,347	-	-	-	-	-	-	-	-	-
Max. piezo. stress coeff (cij)	Regression	4,797	-	-	-	-	-	-	-	-	-
Max. EFG	Regression	11,871	-	-	-	-	-	-	-	-	-
Avg. m_e	Regression	17,643	-	-	-	-	-	-	-	-	-
Is Stable	Classification	-	145,262	-	-	-	-	-	-	-	-
Is Gap Direct	Classification	-	145,262	-	-	-	-	-	-	-	-
n-Seedbeck	Regression	23,211	-	-	-	-	-	-	-	-	-
n-PF	Regression	23,211	-	-	-	-	-	-	-	-	-
p-Seedbeck	Regression	23,211	-	-	-	-	-	-	-	-	-
p-PF	Regression	23,211	-	-	-	-	-	-	-	-	-
Density	Regression	-	145,262	-	-	384,871	-	-	-	-	-
Density Atomic	Regression	-	145,262	-	-	-	-	-	-	-	-
Volume	Regression	-	145,262	-	-	384,871	-	-	-	-	-
Volume Per Atom (VPA)	Regression	-	-	-	-	-	-	84,024	-	-	-
Is Direct	Classification	-	-	10,481	-	-	-	-	-	-	-
Is Direct HSE	Classification	-	-	10,481	-	-	-	-	-	-	-
SOC	Classification	-	-	10,481	-	-	-	-	-	-	-
LCD	Regression	-	-	-	133,524	-	-	-	16,340	-	-
PLD	Regression	-	-	-	133,524	-	-	-	16,340	-	-
Max CO2	Regression	-	-	-	133,524	-	-	-	-	-	-
Min CO2	Regression	-	-	-	133,524	-	-	-	-	-	-
Void Fraction	Regression	-	-	-	133,524	-	-	-	-	-	-
Surface Area m2g	Regression	-	-	-	133,524	-	-	-	-	-	-
Surface Area m2cm3	Regression	-	-	-	133,524	-	-	-	-	-	-

C Chat-like Model Inference Details

Table 8: Hyperparameters used during inference. Temp. represents temperature.

Model Type	Model Name	Input Length	Output Length	Batch Size	Temp.	Top_K
Llama Family	Llama 2-7b-chat-hf	4000	256	256	0.8	10
	Llama 3-8b-Instruct	8000	256	256	0.8	10
	Llama 3.1-8b-Instruct	98000	256	128	0.8	10
	Llama 3.2-3b-Instruct	47000	256	128	0.8	10
Gemma Family	Gemma 7b-it	4000	256	256	0.8	10
	Gemma 1.1-7b-it	4000	256	256	0.8	10
	Gemma 2-9b-it	3000	256	256	0.8	10
Mistral Family	Mistral 7b-Instruct-v0.1	20000	256	256	0.8	10
	Mistral 7b-Instruct-v0.2	20000	256	256	0.8	10
	Mistral 7b-Instruct-v0.3	20000	256	256	0.8	10

D Prompt Templates

Zero-shot Prompt: 0S	Five-shot Prompt: 5S
<p>INPUT PROMPT <S>[INST] <<SYS>> You are a material scientist. Look at the chemical formula of the given crystalline material and predict its property. The output must be in a json format. For example: {property_name:predicted_property_value}. Answer as precise as possible and in as few words as possible. <</SYS>></p> <p>chemical formula: NaCl property name: Band gap. [/INST]</p> <p>RESPONSE {Band gap: 3.97 eV}</p>	<p>INPUT PROMPT <S>[INST] <<SYS>> You are a material scientist. Look at the chemical formula of the given crystalline material and predict its property. The output must be in a json format. For example: {property_name:predicted_property_value}. Answer as precise as possible and in as few words as possible. <</SYS>></p> <p>chemical formula: Na3Bi(P2O7)2 property name: Band gap. {Band gap: 0.0 eV}</p> <p>chemical formula: SrCa7Ti2Mn6O23 property name: Band gap. {Band gap: 0.0 eV}</p> <p>chemical formula: LiLa4FeO8 property name: Band gap. {Band gap: 0.46 eV}</p> <p>chemical formula: CaLaTiMnO6 property name: Band gap. {Band gap: 0.24 eV}</p> <p>chemical formula: PmCu2In property name: Band gap. {Band gap: 0.0 eV}</p> <p>chemical formula: NaCl property name: Band gap. [/INST]</p> <p>RESPONSE {Band gap: 3.97 eV}</p>

Figure 4: Prompt templates when the input is a chemical formula.

Zero-shot Prompt: 0S	Five-shot Prompt: 5S
<p>INPUT PROMPT <S>[INST] <<SYS>> You are a material scientist. Look at the cif structure information of the given crystalline material and predict its property. The output must be in a json format. For example: {property_name:predicted_property_value}. Answer as precise as possible and in as few words as possible. <</SYS>></p> <p>cif structure: data_NaCl_symmetry_space_group_name_H-M 'P 1' _cell_length... property name: Band gap. [/INST]</p> <p>RESPONSE {Band gap: 3.97 eV}</p>	<p>INPUT PROMPT <S>[INST] <<SYS>> You are a material scientist. Look at the cif structure information of the given crystalline material and predict its property. The output must be in a json format. For example: {property_name:predicted_property_value}. Answer as precise as possible and in as few words as possible. <</SYS>></p> <p>cif structure: data_Na3Bi(P2O7)2_symmetry_space_group_name_H-M 'P 1' _cell_length... property name: Band gap. {Band gap: 0.0 eV}</p> <p>cif structure: data_SrCa7Ti2Mn6O23_symmetry_space_group_name_H-M 'P 1' _cell_length... property name: Band gap. {Band gap: 0.0 eV}</p> <p>cif structure: data_LiLa4FeO8_symmetry_space_group_name_H-M 'P 1' _cell_length... property name: Band gap. {Band gap: 0.46 eV}</p> <p>cif structure: data_CaLaTiMnO6_symmetry_space_group_name_H-M 'P 1' _cell_length... property name: Band gap. {Band gap: 0.24 eV}</p> <p>cif structure: data_PmInCu2_symmetry_space_group_name_H-M 'P 1' _cell_length... property name: Band gap. {Band gap: 0.0 eV}</p> <p>cif structure: data_NaCl_symmetry_space_group_name_H-M 'P 1' _cell_length... property name: Band gap. [/INST]</p> <p>RESPONSE {Band gap: 3.97 eV}</p>

Figure 5: Prompt templates when the input is a CIF file.

Zero-shot Prompt: 0S	Five-shot Prompt: 5S
<p>INPUT PROMPT</p> <p><s>[INST] <<SYS>></p> <p>You are a material scientist.</p> <p>Look at the structure description of the given crystalline material and predict its property.</p> <p>The output must be in a json format. For example: {property_name:predicted_property_value}.</p> <p>Answer as precise as possible and in as few words as possible.</p> <p><</SYS>></p> <p>structure description: NaCl is Tetraauricupride structured and crystallizes in the cubic P m3m...</p> <p>property name: Band gap. [/INST]</p> <p>RESPONSE</p> <p>{Band gap: 3.97 eV}</p>	<p>INPUT PROMPT</p> <p><s>[INST] <<SYS>></p> <p>You are a material scientist.</p> <p>Look at the structure description of the given crystalline material and predict its property.</p> <p>The output must be in a json format. For example: {property_name:predicted_property_value}.</p> <p>Answer as precise as possible and in as few words as possible.</p> <p><</SYS>></p> <p>structure description: Na3Bi(P2O7)2 crystallizes in the triclinic P-1 space group...</p> <p>property name: Band gap.</p> <p>{Band gap: 0.0 eV}</p> <p>structure description: SrCa7Ti2Mn6O23 crystallizes in the triclinic P1 space group...</p> <p>property name: Band gap.</p> <p>{Band gap: 0.0 eV}</p> <p>structure description: LiLa4FeO8 is (La,Ba)CuO4-derived structured and crystallizes in the...</p> <p>property name: Band gap.</p> <p>{Band gap: 0.46 eV}</p> <p>structure description: CaLaTiMnO6 is Orthorhombic Perovskite-derived structured and crysta...</p> <p>property name: Band gap.</p> <p>{Band gap: 0.24 eV}</p> <p>structure description: PmCu2In is Heusler structured and crystallizes in the trigonal R-3m...</p> <p>property name: Band gap.</p> <p>{Band gap: 0.0 eV}</p> <p>structure description: NaCl is Tetraauricupride structured and crystallizes in the cubic P m3m...</p> <p>property name: Band gap. [/INST]</p> <p>RESPONSE</p> <p>{Band gap: 3.97 eV}</p>

Figure 6: Prompt templates when the input is a crystal structure description.

E Result Details for Each Dataset

Table 9: Results for MP dataset. The performance on regression tasks is evaluated in terms of MAD:MAE ratio (the higher the better) while that of classification tasks (Is Stable and Is Gab Direct) is evaluated in terms of AUC score. FEPA: Formation Energy Per Atom, EPA: Energy Per Atom.

Input	Model	FEPA 145.2K	Bandgap 145.3K	EPA 145.2K	Ehull 145.2K	Efermi 145.2K	Density 145.2K	Density Atomic 145.2K	Volume 145.2K	Is Stable 145.2K	Is Gab Direct 145.2K
CIF	CGCNN (baseline)	8.151	3.255	7.224	3.874	3.689	8.773	5.888	1.703	0.882	0.810
Comp.	Llama 2-7b-chat:0S	0.008	0.623	0.009	0.001	0.003	0.967	0.754	0.747	0.500	0.482
	Llama 2-7b-chat:5S	0.33	1.217	0.239	0.132	0.706	0.899	0.724	0.771	0.502	0.512
	MatBERT-109M	8.151	2.971	9.32	2.583	3.527	7.626	5.26	3.099	0.764	0.681
	LLM-Prop-35M	7.482	2.345	7.437	2.006	3.159	6.682	3.523	2.521	0.746	0.636
CIF	Llama 2-7b-chat:0S	0.032	0.135	0.022	0.001	0.015	0.97	0.549	1.41	0.503	0.499
	Llama 2-7b-chat:5S	Inval.	1.111	0.289	Inval.	0.685	0.98	0.99	0.926	0.498	0.506
	MatBERT-109M	11.017	3.423	13.244	3.808	4.435	10.426	6.686	6.58	0.790	0.710
	LLM-Prop-35M	14.322	3.758	17.354	2.182	4.515	13.834	4.913	7.556	0.776	0.700
Descr.	Llama 2-7b-chat:0S	0.019	0.633	0.023	0.001	0.008	1.31	0.693	0.807	0.500	0.500
	Llama 2-7b-chat:5S	0.394	1.061	0.297	0.247	0.684	0.916	0.782	0.704	0.500	0.504
	MatBERT-109M	11.935	3.524	13.851	4.085	4.323	9.9	6.899	6.693	0.794	0.713
	LLM-Prop-35M	15.913	3.931	18.412	2.74	4.598	14.388	4.063	8.888	0.794	0.690

Table 10: Results for JARVIS-DFT. The performance on regression tasks is evaluated in terms of MAD:MAE ratio (the higher the better). FEPA: Formation Energy Per Atom, Tot. En.: Total Energy, Exf. En.: Exfoliation Energy.

Input	Model	FEPA 75.9K	Bandgap (OPT) 75.9K	Tot. En. 75.9K	Ehull 75.9K	Bandgap (MBJ) 19.8K	Kv 23.8K	Gv 23.8K	SLME 9.7K	Spillage 11.3K	ϵ_{∞} (OPT) 18.2K
CIF	CGCNN (baseline)	13.615	4.797	22.906	1.573	4.497	3.715	2.337	1.862	1.271	2.425
Comp.	Llama 2-7b-chat:0S	0.021	0.011	0.02	0.005	0.92	0.428	0.374	0.148	Inval.	0.18
	Llama 2-7b-chat:5S	0.886	0.011	0.02	1.292	0.979	0.88	0.992	0.456	0.85	1.148
	MatBERT-109M	6.808	4.083	9.21	2.786	3.755	2.906	1.928	1.801	1.243	2.017
	LLM-Prop-35M	4.765	2.621	5.936	2.073	2.922	2.162	1.654	1.575	1.14	1.734
CIF	Llama 2-7b-chat:0S	0.023	0.011	0.02	0.002	0.193	0.278	0.358	0.186	0.702	0.781
	Llama 2-7b-chat:5S	0.859	Inval.	Inval.	1.173	1.054	0.874	0.91	0.486	0.916	1.253
	MatBERT-109M	10.211	5.483	15.673	4.862	5.344	4.283	2.6	2.208	1.444	2.408
	LLM-Prop-35M	12.996	3.331	22.058	2.648	4.93	4.121	2.409	2.175	1.37	2.135
Descr.	Llama 2-7b-chat:0S	0.007	0.011	0.02	0.004	0.94	0.498	0.382	0.07	0.135	0.647
	Llama 2-7b-chat:5S	0.845	0.011	0.02	1.273	1.033	0.87	0.969	0.461	0.857	1.201
	MatBERT-109M	10.211	5.33	15.141	4.691	5.01	4.252	2.623	2.178	1.452	2.384
	LLM-Prop-35M	12.614	3.427	23.509	4.532	4.983	4.128	2.419	2.061	1.307	2.334
		ϵ (DFPT) 4.7K	Max. Piezo. (dij) 3.3K	Max. Piezo. (eij) 4.7K	Max. EFG 11.8K	Exf. En. 0.8K	Avg. m_e 17.6K	n-Seebeck 23.2K	n-PF 23.2K	p-Seebeck 23.2K	p-PF 23.2K
CIF	CGCNN (baseline)	1.12	0.418	1.291	1.787	0.842	1.796	2.23	1.573	3.963	1.59
Comp.	Llama 2-7b-chat:0S	0.012	0.121	0.001	0.141	0.384	0.028	0.874	0.801	0.971	0.874
	Llama 2-7b-chat:5S	1.416	1.289	1.305	0.765	0.512	0.535	1.008	1.04	0.93	0.568
	MatBERT-109M	1.533	1.464	1.426	1.658	1.124	2.093	1.908	1.318	2.752	1.356
	LLM-Prop-35M	1.454	1.447	1.573	1.38	1.042	1.658	1.725	1.145	2.233	1.285
CIF	Llama 2-7b-chat:0S	0.033	0.104	0.001	0.246	0.411	0.041	0.429	0.766	0.83	0.826
	Llama 2-7b-chat:5S	Inval.	Inval.	Inval.	0.796	0.51	Inval.	1.039	1.396	Inval.	Inval.
	MatBERT-109M	1.509	1.758	2.405	2.143	1.374	2.45	2.268	1.446	3.337	1.476
	LLM-Prop-35M	1.578	2.103	2.405	1.936	1.044	1.796	1.955	1.332	2.503	1.399
Descr.	Llama 2-7b-chat:0S	0.08	0.266	0.001	0.138	0.285	0.019	0.769	0.793	0.825	0.829
	Llama 2-7b-chat:5S	1.649	1.174	1.152	0.806	0.661	0.523	1.098	1.024	0.948	0.563
	MatBERT-109M	1.534	1.807	2.556	2.081	1.36	2.597	2.241	1.432	3.26	1.565
	LLM-Prop-35M	1.64	2.116	2.315	1.978	1.168	1.858	2.154	1.364	2.61	1.407

Table 11: Results for SNUMAT. The performance on regression tasks is evaluated in terms of MAD:MAE ratio (the higher the better) while that of classification tasks (Is Direct, Is Direct HSE, and SOC) is evaluated in terms of AUC score.

Input	Model	Bandgap GGA 10.3K	Bandgap HSE 10.3K	Bandgap GGA Optical 10.3K	Bandgap HSE Optical 10.3K	Is Direct 10.3K	Is Direct HSE 10.3K	SOC 10.3K
CIF	CGCNN (baseline)	2.075	2.257	1.727	1.835	0.691	0.675	0.800
Comp.	Llama 2-7b-chat:0S	0.797	0.948	1.156	0.859	0.503	0.484	Inval.
	Llama 2-7b-chat:5S	1.267	1.327	0.862	1.174	0.475	0.468	0.455
	MatBERT-109M	1.899	1.975	1.646	1.793	0.671	0.645	0.820
	LLM-Prop-35M	1.533	1.621	1.392	1.491	0.647	0.624	0.829
CIF	Llama 2-7b-chat:0S	0.346	0.454	1.09	0.838	0.479	0.488	0.500
	Llama 2-7b-chat:5S	Inval.	Inval.	Inval.	Inval.	0.494	0.500	0.427
	MatBERT-109M	2.28	2.472	1.885	1.889	0.677	0.650	0.823
	LLM-Prop-35M	1.23	2.401	1.786	1.9	0.661	0.664	0.656
Descr.	Llama 2-7b-chat:0S	0.802	0.941	1.013	0.779	0.499	0.509	Inval.
	Llama 2-7b-chat:5S	0.774	1.315	0.901	1.172	0.594	0.623	0.486
	MatBERT-109M	2.298	2.433	1.901	1.978	0.683	0.645	0.862
	LLM-Prop-35M	2.251	2.142	1.84	1.569	0.681	0.657	0.866

Table 12: Results for GNoME. The performance on regression tasks is evaluated in terms of MAD:MAE ratio (the higher the better). FEPA: Formation Energy Per Atom, DEPA: Decomposition Energy Per Atom, Tot. En.: Total Energy.

Input	Model	FEPA 376.2K	Bandgap 282.7K	DEPA 376.2K	Tot. En. 282.7K	Volume 282.7K	Density 282.7K
CIF	CGCNN (baseline)	34.57	8.549	2.787	7.443	7.967	56.077
Comp.	Llama 2-7b-chat:0S	0.002	0.177	0.0	0.088	0.455	0.368
	Llama 2-7b-chat:5S	0.194	0.086	0.255	0.765	1.006	0.865
	MatBERT-109M	30.248	4.692	2.787	8.57	13.157	15.145
	LLM-Prop-35M	25.472	3.735	1.858	21.624	16.556	25.615
CIF	Llama 2-7b-chat:0S	0.003	0.045	0.0	0.706	43.331	0.794
	Llama 2-7b-chat:5S	Invalid.	0.087	Invalid.	Invalid.	1.029	0.878
	MatBERT-109M	24.199	9.16	3.716	15.309	16.691	16.467
	LLM-Prop-35M	28.469	3.926	3.344	17.837	17.082	25.615
Descr.	Llama 2-7b-chat:0S	0.002	0.114	0.0	0.661	0.654	0.805
	Llama 2-7b-chat:5S	0.192	0.086	0.106	0.75	1.006	0.891
	MatBERT-109M	30.248	5.829	3.716	18.205	17.824	16.599
	LLM-Prop-35M	28.469	5.27	3.716	17.02	17.02	25.936

Table 13: Results for hMOF. The performance on regression tasks is evaluated in terms of MAD:MAE ratio (the higher the better).

Input	Model	Max CO2 132.7K	Min CO2 132.7K	LCD 132.7K	PLD 132.7K	Void Fraction 132.7K	Surface Area m2g 132.7K	Surface Area m2cm3 132.7K
CIF	CGCNN (baseline)	1.719	1.617	1.989	1.757	2.912	3.765	2.039
Comp.	Llama 2-7b-chat:0S	0.011	0.002	0.009	0.008	0.5	0.454	0.233
	Llama 2-7b-chat:5S	0.679	0.058	0.949	1.026	0.945	0.567	0.366
	MatBERT-109M	1.335	1.41	1.435	1.378	1.57	1.517	1.367
	LLM-Prop-35M	1.41	1.392	1.432	1.468	1.672	1.657	1.321
CIF	Llama 2-7b-chat:0S	0.017	0.003	0.016	0.011	0.549	0.54	0.359
	Llama 2-7b-chat:5S	Invalid.	Invalid.	0.951	1.067	Invalid.	Invalid.	Invalid.
	MatBERT-109M	1.421	1.428	1.544	1.482	1.641	1.622	1.461
	LLM-Prop-35M	1.564	1.41	1.753	1.435	1.9	1.926	1.374
Descr.	Llama 2-7b-chat:0S	0.129	0.014	0.026	0.006	0.382	0.497	0.299
	Llama 2-7b-chat:5S	0.684	0.058	0.955	1.006	0.931	0.571	0.37
	MatBERT-109M	1.438	1.466	1.602	1.511	1.719	1.697	1.475
	LLM-Prop-35M	1.659	1.486	1.623	1.789	1.736	2.144	1.508

Table 14: Results for Cantor HEA. The performance on regression tasks is evaluated in terms of MAD:MAE ratio (the higher the better). FEPA: Formation Energy Per Atom, EPA:Energy Per Atom, VPA:Volume Per Atom.

Input	Model	FEPA 84.0K	EPA 84.0K	Ehull 84.0K	VPA 84.0K
CIF	CGCNN (baseline)	9.036	49.521	9.697	2.869
Comp.	Llama 2-7b-chat:0S	0.005	0.098	0.003	0.031
	Llama 2-7b-chat:5S	0.896	0.658	0.928	0.986
	MatBERT-109M	3.286	16.17	5.134	2.489
	LLM-Prop-35M	3.286	22.638	5.134	2.543
CIF	Llama 2-7b-chat:0S	0.001	0.084	0.0	0.004
	Llama 2-7b-chat:5S	Invalid.	Invalid.	Invalid.	Invalid.
	MatBERT-109M	7.229	17.607	9.187	5.809
	LLM-Prop-35M	8.341	36.015	11.636	6.919
Descr.	Llama 2-7b-chat:0S	0.001	0.101	0.164	0.011
	Llama 2-7b-chat:5S	0.797	0.615	0.938	0.93
	MatBERT-109M	7.229	17.607	9.187	5.881
	LLM-Prop-35M	8.341	36.015	11.636	7.713

Table 15: Results for QMOF. The performance on regression tasks is evaluated in terms of MAD:MAE ratio (the higher the better). Tot. En.: Total Energy.

Input	Model	Bandgap 7.6K	Tot. En. 7.6K	LCD 7.6K	PLD 7.6K
CIF	CGCNN (baseline)	2.431	1.489	4.068	4.317
Comp.	Llama 2-7b-chat:0S	0.901	0.26	0.045	0.009
	Llama 2-7b-chat:5S	0.648	0.754	1.241	1.086
	MatBERT-109M	1.823	1.695	2.329	2.349
	LLM-Prop-35M	1.759	1.621	2.293	2.157
CIF	Llama 2-7b-chat:0S	0.201	0.244	0.02	0.011
	Llama 2-7b-chat:5S	Invalid.	Invalid.	Invalid.	Invalid.
	MatBERT-109M	1.994	4.378	2.908	2.818
	LLM-Prop-35M	2.166	4.323	2.947	2.87
Descr.	Llama 2-7b-chat:0S	0.358	0.217	0.025	0.006
	Llama 2-7b-chat:5S	0.777	0.713	1.125	1.17
	MatBERT-109M	2.166	4.133	2.981	2.941
	LLM-Prop-35M	2.091	4.312	2.831	2.829

Table 16: Results for JARVIS-QETB. The performance on regression tasks is evaluated in terms of MAD:MAE ratio (the higher the better). FEPA: Formation Energy Per Atom, EPA:Energy Per Atom, Tot. En.: Total Energy, Ind. Bandgap: Indirect Bandgap.

Input	Model	FEPA 623.9K	EPA 623.9K	Tot. En. 623.9K	Ind. Bandgap 623.9K
CIF	CGCNN (baseline)	1.964	228.201	11.218	5.534
Comp.	Llama 2-7b-chat:0S	0.003	0.369	0.172	0.21
	Llama 2-7b-chat:5S	0.812	1.037	1.032	1.306
	MatBERT-109M	1.431	37.979	8.19	0.21
	LLM-Prop-35M	2.846	211.757	21.309	1.861
CIF	Llama 2-7b-chat:0S	0.003	0.412	0.656	0.04
	Llama 2-7b-chat:5S	0.8	1.024	1.076	1.71
	MatBERT-109M	24.72	135.156	26.094	4.779
	LLM-Prop-35M	23.346	318.291	48.192	1.845
Descr.	Llama 2-7b-chat:0S	0.003	0.408	0.484	0.16
	Llama 2-7b-chat:5S	0.85	1.015	1.035	1.021
	MatBERT-109M	26.265	122.884	29.409	7.788
	LLM-Prop-35M	22.513	312.218	35.43	1.845

Table 17: Results for OQMD. The performance on regression tasks is evaluated in terms of MAD:MAE ratio (the higher the better). FEPA: Formation Energy Per Atom.

Input	Model	FEPA 963.5K	Bandgap 963.5K
CIF	CGCNN (baseline)	22.291	6.701
Comp.	Llama 2-7b-chat:0S	0.019	0.192
	Llama 2-7b-chat:5S	1.013	1.306
	MatBERT-109M	7.662	3.883
	LLM-Prop-35M	9.195	2.845
CIF	Llama 2-7b-chat:0S	0.009	0.047
	Llama 2-7b-chat:5S	1.051	1.731
	MatBERT-109M	13.879	7.163
	LLM-Prop-35M	18.861	3.22
Descr.	Llama 2-7b-chat:0S	0.025	0.187
	Llama 2-7b-chat:5S	0.991	1.468
	MatBERT-109M	15.012	7.041
	LLM-Prop-35M	16.346	3.644

Table 18: Results for OMDB. The performance on regression tasks is evaluated in terms of MAD:MAE ratio (the higher the better).

Input	Model	Bandgap 12.1K
CIF	CGCNN (baseline)	2.751
Comp.	Llama 2-7b-chat:0S	0.886
	Llama 2-7b-chat:5S	1.009
	MatBERT-109M	1.554
	LLM-Prop-35M	1.507
CIF	Llama 2-7b-chat:0S	0.159
	Llama 2-7b-chat:5S	0.930
	MatBERT-109M	1.777
	LLM-Prop-35M	1.777
Descr.	Llama 2-7b-chat:0S	0.155
	Llama 2-7b-chat:5S	1.002
	MatBERT-109M	1.847
	LLM-Prop-35M	1.656

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