
Scientific Knowledge Graph and Ontology Generation using Open Large Language Models

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Abstract

Knowledge Graphs (KGs) are powerful tools for structured information modeling, increasingly recognized for their potential to enhance the factuality and reasoning capabilities of Large Language Models (LLMs). However, in scientific domains, KG representation is often constrained by the absence of ontologies capable of modeling complex hierarchies and relationships inherent in the data. Moreover, the manual curation of KGs and ontologies from scientific literature remains a time-intensive task typically performed by domain experts. This work proposes a novel method leveraging LLMs for zero-shot, end-to-end ontology, and KG generation from scientific literature, implemented exclusively using open-source LLMs. We evaluate our approach by assessing its ability to reconstruct an existing KG and ontology of chemical elements and functional groups. Furthermore, we apply the method to the emerging field of Single Atom Catalysts (SACs), where information is scarce and unstructured. Our results demonstrate the effectiveness of our approach in automatically generating structured knowledge representations from complex scientific literature, in areas where manual curation is challenging or time-consuming. The generated ontologies and KGs provide a foundation for improved information retrieval and reasoning in specialized fields, opening new avenues for LLM-assisted scientific research and knowledge management. ¹

1 Introduction

Knowledge Graphs (KGs) are a powerful tool for representing structured information, enabling inference, retrieval, and analysis over large amounts of data. KGs model the real world in a graph representation, where nodes represent entities and edges represent relationships between them¹³. Recently, KGs have gained traction in the field of Natural Language Processing (NLP), demonstrating their potential to enhance the capabilities of Large Language Models (LLMs)^{1,22,38}. It is well-established that LLMs are prone to ‘hallucinations’, where they generate incorrect or misleading information⁵³. In this context, KGs have been shown to help mitigate this problem through Retrieval Augmented Generation (RAG), where graph structure is embedded as textual information as input to the LLM^{17,27,43,52}. Moreover, KGs have also been shown to improve the reasoning capabilities of LLMs^{33,51,54}.

¹Source code is available under MIT license at <https://github.com/schwallergroup/ontorag/tree/main/src/OntoGen>.

Knowledge Graph Extraction (KGE) from text is usually performed through triplets extraction of the form (subject, predicate, object), where entities are extracted from the text using Named Entity Recognition (NER) and relationships between entities are inferred from the context². Currently, LLMs are state-of-the-art in many NLP tasks, including NER^{25,37}, making them promising candidates for unsupervised KGE^{20,36,47}. KGs based on triplets, however, often fail to model scientific fields such as bioinformatics or chemistry where more complex structures are needed^{6,19,44}. Additionally, most KGE methods are based on the extraction of triplets from short sentences or paragraphs⁵. These methods are typically pre-trained on specific text corpora or domains or require labeled subsets of samples, which hampers their ability to generalize to different contexts^{12,39}. Other approaches, such as co-occurrence-based extraction of relations, may fail to extract true causal relationships⁴⁸. All these factors limit the application of KGE to scientific literature, which remains considerably understudied⁵. In complex domains, the enrichment of KGs with ontologies can significantly improve information representation. Ontologies themselves are composed of classes, organized in a hierarchical structure denoted as taxonomy, containing relationships between classes. Each of the entities in the KGs can be considered as an instance of one or more classes in the ontology. Thus, ontologies provide a high-level schema for the knowledge present in the KG^{13,14}. Ontologies however are commonly built and curated manually by domain experts, a process that is typically time-consuming and labor-intensive. Given the success of LLMs in many NLP tasks, some solutions have been proposed for ontology generation using LLMs. These methods, however, often have limitations: they may require pre-defined training sets for fine-tuning^{8,35}, cannot generate full ontologies¹¹, do not scale beyond smaller problems^{3,21}, or solely rely on the LLMs knowledge, making their application to scientific fields unpractical or unclear and hindering generalisation into other domains.^{7,55}

There is a clear need in the field to generate hallucination-free knowledge extraction from scientific text in a way that is human-interpretable, consistent, and does not require efforts in fine-tuning language models on niche subjects. In this work, a method is proposed for end-to-end zero-shot ontology and KG generation from scientific literature using LLMs. To evaluate the performance, we measure the ability of the proposed method to reconstruct an existing KG and ontology of chemical elements. We then apply the method to the scientific domain of Single Atom Catalysis (SAC)^{10,28,41,45}, an emerging field in catalysis with no previously existing KG. In doing so, we create what is, to our knowledge, the first example of a domain-specific ontology for SAC. We demonstrate the value general-purpose LLMs add to the automation of the typically laborious process of ontology construction. The presented work therefore represents an advancement in the use of LLMs to not only perform domain-specific question answering but also extract domain-specific knowledge in a way that can be interpreted by experts. Finally, to demonstrate the utility of our work, we show that the combination of an ontology and a KG improves the quality of RAG-based solutions that rely only on KGs, such as GraphRAG¹⁷.

2 Methods

2.1 Ontologies as a Form of Knowledge Representation

From a practical engineering perspective, ontologies are a specification of conceptualization using description logic. They model relationships between concepts and the words used to represent them. In their most basic form, an ontology can be thought of as a series of "isA" relationships formed in a hierarchical tree structure called a taxonomy. This taxonomy serves as the backbone of an ontology and can be engineered to include relationships, instances, properties, and axioms. For the purpose of this work, an ontology can be formally defined as follows:

Definition 2.1 An ontology²⁹ is a tuple $\{\mathcal{C}, \mathcal{R}, \mathcal{I}, \mathcal{P}, \mathcal{A}\}$ where:

- \mathcal{C} is the set of all concepts present in the ontology such that $\mathcal{C} = \{c_1, c_2, c_3, \dots, c_n\}$
- \mathcal{R} is the set of all relationships present in the ontology such that $\mathcal{R} = \{(c_i, r_s, c_j) | c_i, c_j \in \mathcal{C} \ \& \ r_s \in \mathcal{R}_f\}$ where \mathcal{R}_f is the set of all possible relations
- \mathcal{I} is the set of all instances of concepts present in the ontology such that $\mathcal{I} = \{i_1, i_2, i_3, \dots, i_n\}$ and $\forall(i)\exists(c)|i \in c$
- \mathcal{P} is the set of all possible properties in an ontology such that $\mathcal{P} = \{p_1, p_2, p_3, \dots, p_n\}$ and $p : \mathcal{I} \rightarrow \mathcal{V}$ or $p : \mathcal{C} \rightarrow \mathcal{V}$ where \mathcal{V} is the set of all possible values for a property

- and \mathcal{A} is the set of axioms in the ontology.

A key practical application of ontologies is the construction and integration of databases. We demonstrate the usefulness of our pipeline by using constructed ontologies to create graph databases, otherwise known as knowledge graphs. To relate knowledge graphs to ontologies, we utilize the following definition:

Definition 2.2 A knowledge graph²⁹ is set of tuples $\{\mathcal{S}, \mathcal{P}, \mathcal{O}\}$ where

- \mathcal{S} is the set of subjects where $s \in \mathcal{S}$ is an instance of a concept from the ontology $\Rightarrow s \in \mathcal{I}$
- \mathcal{P} is the set of predicates where $p \in \mathcal{P}$ is a relation from the ontology $\Rightarrow p \in \mathcal{R}_f$
- \mathcal{O} is the set of objects where $o \in \mathcal{O}$ is also an instance of a concept from the ontology such that $p : s \xrightarrow{p} o$

From this definition, it follows that the ontology serves as the conceptual schema for the construction of a knowledge graph. The following methods detail our approach to leveraging zero-shot learning for creating an ontology of SAC and the subsequent construction of an ontology-informed knowledge graph.

Example 2.3 Figure 1 shows an example ontology with four classes (white nodes) and a KG with three instances (shaded nodes). The ontology has three *isA* and is related to the KG through three *instanceOf* relationships. Between the nodes of the KG, one relationship exists of the type ‘supported on’. The term ‘Thing’ is always the root node of an ontology.

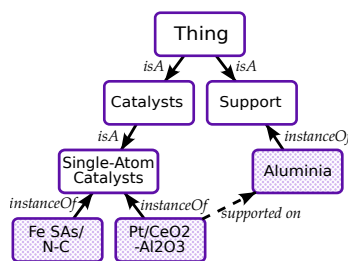


Figure 1: Illustrative example of an ontology and a knowledge graph in the domain of SAC.

2.2 Self consistency

The improvement of LLMs’ capabilities to generate high-quality, hallucination-free answers is currently a highly active area of research. Many generic methods have been proposed that improve LLMs outputs without training data, fine-tuning or reinforcement learning, which includes, among others, self-consistency⁴⁹, debating LLMs¹⁵, and self-refinement³⁴. Research by Huang et al.²⁶ demonstrates that self-consistency offers competitive results while being more computationally efficient compared to other methods. Therefore, in this work, self-consistency is used to improve the quality of answers from a LLM. As utilized in our approach, self-consistency can be defined as:

Definition 2.4 Let $a_1, a_2, \dots, a_n \in \mathbb{A}$ be the answers to a given prompt p generated by a LLM, and r_i the set of tokens generated before the answer a_i .

Self-Consistency (SC) applies a marginalization over r_i by taking the majority vote of the answers a_i , i.e. $a = \arg \max_{a_i} \sum_{j=1}^n \mathbb{1}(a_i = a_j)$, thus giving as a final answer the most “consistent” answer generated by the LLM.

It is important to note that self-consistency was initially proposed to enhance Chain of Thought (CoT) reasoning⁵⁰ in LLMs⁴⁹, to improve performance on generalized problem-solving tasks. In our work, we leverage the generalizability of self-consistency to improve the quality of our knowledge schemas reconstruction.

2.3 In-Context Ontology Generation (ICOG)

Here, we introduce our method for In-Context Ontology Generation (ICOG) and KGE from scientific literature. The method is split into five separate steps: (1) the vocabulary that will be used to generate the ontology is extracted from the text, (2) the higher-level categories of the ontology are generated, (3) the taxonomy of the ontology is extracted, (4) the knowledge graph is instantiated from the extracted ontology, and (5) the relationships between the terms in the vocabulary are extracted.

Vocabulary Extraction The first step in our method is the extraction of all the key vocabulary terms from the text. The extracted terms serve as the base vocabulary that will compose the ontology. It follows that any terms that are not extracted in this step will be absent from the final ontology. Vocabulary extraction is itself split into three steps: (1) terms extraction, (2) acronyms extraction, and (3) lemmatization.

Terms Extraction Traditional models trained for specific NER tasks often fail to generalize to domains not included in their training data. To address this limitation, we utilize open general-purpose LLMs for zero-shot vocabulary identification. The relevant text is first split into sentences to ensure individual words can be obtained in a fine-grained, detailed manner. This sentence splitting is performed using a pre-trained LLM that predicts sentence boundaries. Empirically, we observed that splitting the text into smaller sentences allowed for a more fine-grained extraction and increased the number of terms extracted. For each of the extracted sentences, an LLM is then prompted to list each of the terms present in the sentence. Finally, to avoid terms hallucinated by the LLM or subtle changes in the writing of the terms, a verification step is performed by ensuring that each of the listed terms is present in the original sentence.

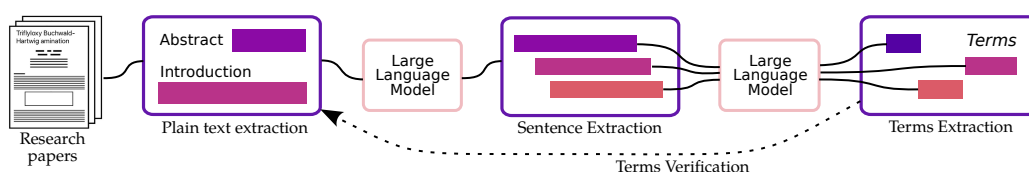


Figure 2: Illustration of the vocabulary extraction process.

Acronym Extraction Acronyms are commonly used in scientific literature to refer to long terms. To be able to associate each of the extracted terms of the previous step with their respective acronyms, we employ a separate procedure similar to terms extraction, but with the LLM prompted specifically to list all acronyms and their corresponding terms. As in the previous step, a verification procedure is used to ensure that the pairs of acronym terms listed are present in the original text.

Lemmatization To handle different writing forms of the same term (e.g. “Single Atom Catalyst” and “Single Atom Catalysts”), we apply lemmatization to each extracted term after removing punctuation. This process reduces words to their root form, avoiding grammatical variations. Terms that are grouped together after lemmatization are considered to refer to the same concept.

Category extraction The first level of the taxonomic hierarchy, which is usually denoted as categories, consists of the main concepts that the ontology will cover. For instance, in the field of chemistry, the first level of the taxonomy might include categories such as elements, compounds, and reactions. However, these high-level categories cannot always be extracted from individual papers, as each scientific paper typically focuses on a particular area within a domain. Therefore, the highest level concepts must be extracted across several of the papers available to the system.

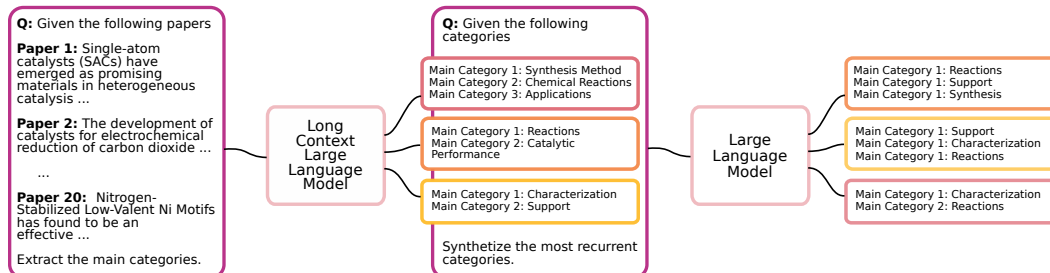


Figure 3: Illustration of the generation and refinement steps from the categories extraction process.

To accomplish this, we propose a two-step process:

1. *Generation*: Generating a list of categories from a set of papers is challenging for conventional LLMs due to their limited context window. To overcome this limitation, we employ

long-context LLMs (LCLLMs)^{9,24,30,40} for this task. First, a random sample of papers is presented to an LCLLM, which is then prompted to extract the main categories across the given papers. As LCLLMs are known to be sensitive to input order³¹, we generate multiple answers, each with a random shuffle in the order of the papers.

2. *Refinement*: Using the answers generated in the previous step, we prompt an LLM to generate a curated list of categories using the most frequently occurring ones. SC is applied in this step to obtain categorical consistency.

It is important to note that the outcome of the previous step is a list of categories that will serve as the seed for the rest of the ontology. However, the correctness of the ontology depends on the downstream application that it is intended for. Therefore, some manual effort may be required at the end of this step to select the most relevant categories from the list of all the generated seeds, according to the requirements of the user.

Taxonomy extraction The relationships in a taxonomy structure consist entirely of *isA* relationships. Once the vocabulary is extracted and the main categories are generated, the vocabulary needs to be organized into a hierarchical structure. Since the first level of the hierarchy is already generated, we introduce a top-down approach where each extracted term is placed in a taxonomic tree structure. While taxonomic information can be extracted from each paper separately, but it must be organised into a global single taxonomy. Therefore, we developed an incremental and sequential approach to build a global taxonomy.

Given a corpus of N papers $\mathcal{P} = \{P_1, P_2, \dots, P_N\}$, a set of terms $\mathcal{V} = \{V_1, V_2, \dots, V_N\}$ extracted from the papers, where V_i is the vocabulary from paper P_i , and a taxonomy $\mathcal{T}^{(k)} = \{(s_1, t_1), (s_2, t_2), \dots, (s_M, t_M)\}$ at iteration k , where each (s_i, t_i) with $s_i, t_i \in \mathcal{V}$ is a pair of terms that represents a *isA* relationship. The procedure to generate the full taxonomy is presented below:

Algorithm 1 Iterative and Incremental Top-Down Taxonomy Generation

Input: Papers \mathcal{P} , Vocabulary \mathcal{V} , Initial Taxonomy $\mathcal{T}^{(0)}$
Output: Reconstructed Taxonomy after M iterations $\mathcal{T}^{(M)}$

- 1: **for** $k = 1, \dots, K$ **do**
- 2: $\mathcal{T}^{(k)} \leftarrow \mathcal{T}^{(k-1)}$
- 3: **for** $P_i \in \mathcal{P}$ **do**
- 4: $R_i \leftarrow \text{query_relationships}(P_i, V_i, \mathcal{T}^{(k)})$
- 5: **for** $(s, t) \in R_i$ **do**
- 6: **if** $\text{is_valid}((s, t), \mathcal{T}^{(k)})$ **then**
- 7: $\mathcal{T}^{(k)} \leftarrow \mathcal{T}^{(k)} \cup \{(s, t)\}$
- 8: **return** $\mathcal{T}^{(M)}$

In this algorithm, `query_relationships` extracts pairs (s, t) of *isA* relationships based on the information presented in paper P_i , where s is a term present in the taxonomy $\mathcal{T}^{(k)}$ and t is a term from the vocabulary V_i . The primary goal of `query_relationships` is to place each term into the existing taxonomy. Note that `query_relationships` may not necessarily return a single relationship for each term, but rather a subset of relationships. The `is_valid` function checks that no loop is created in the taxonomy if a given relationship is inserted. This procedure builds a taxonomy incrementally. Multiple iterations are necessary because a term might not be placed in one of the early taxonomies but in a later one.

In our implementation, `query_relationships` prompts an LLM with the content of the paper, the complete list of terms in the taxonomy, and the vocabulary terms to be queried. An example of such a prompt and its corresponding answer can be found in Appendix B. Given that the quality of the generated taxonomy heavily depends on the performance of this specific function, we employ self-consistency to mitigate the number of hallucinations generated.

Knowledge Graph Instantiation Once the full taxonomy is constructed from the vocabulary terms and papers, we consider the leaf nodes of the taxonomy as instances for the KG. We reason that since leaf nodes have no children, they should be treated as the lowest-level concepts in our knowledge representation schema, and thus can be assumed to be instances of the KG. Conversely, non-leaf

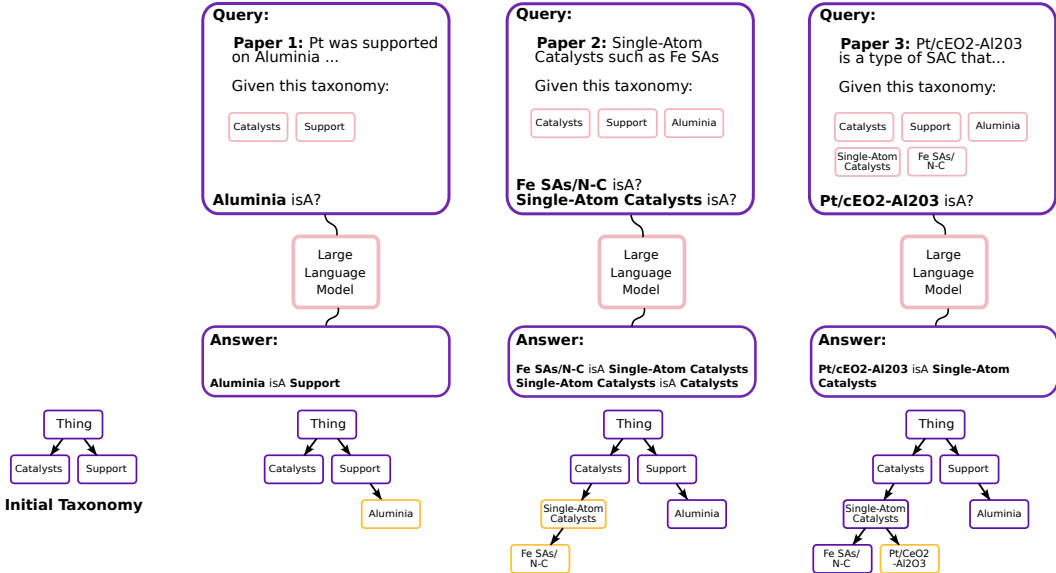


Figure 4: Illustration of the iterative and incremental taxonomy generation process.

nodes, having children related through *isA* relationships, are considered higher-level classes in the ontology rather than instances of the KG.

Relationship extraction Finally, we perform relationship extraction using a conventional triplet extraction approach. Specifically, we prompt an LLM to generate triplets from a given text, considering the extracted vocabulary. The resulting triplets are then filtered to retain only those involving terms from the vocabulary.

Details on the packages and models used in this pipeline are found in the Appendix. The code will be released in open source with the deanonymized version of the manuscript.

2.4 Ontology Reconstruction Evaluation

Evaluating a reconstructed ontology is not a trivial task, as the quality of the ontology is highly dependent on the downstream application for which it is intended. Additionally, many correct ontologies can be generated from the same text. To overcome the limitations of common metrics such as *term accuracy*, the percentage of the terms in the ground truth taxonomy that appear in the reconstructed taxonomy, and *knowledge graph accuracy*, the percentage of the instances in the ground truth KG that are also instances of the reconstructed KG, we propose the hierarchical accuracy as an evaluation metric for reconstructed ontologies.

Hierarchical Accuracy The main idea behind hierarchical accuracy is that if a node in the reconstructed taxonomy shares a common ancestor with a node in a ground truth taxonomy, then that node is considered to be correctly placed in the taxonomy. The hierarchical accuracy is then defined as the average number of nodes from the ground truth taxonomy that are correctly placed in the reconstructed taxonomy. This metric accounts for reconstructions that are more general than the ground truth, but penalizes reconstructions with missing nodes or misplaced nodes.

Generally, it is accepted that a term can appear more than once in a taxonomy. As a consequence, taxonomies are not modeled as trees but rather as Directed Acyclic Graphs (DAGs). Below, the hierarchical accuracy metric is formally defined.

Let $\mathcal{G} = (\mathcal{N}, \mathcal{E})$ be a DAG, where \mathcal{N} is the set of nodes and \mathcal{E} is the set of edges, with a root node $r \in \mathcal{N}$, such that any node $n_i \in \mathcal{N}$ is reachable from r . The set of all paths from r to n_i , which is denoted $\mathcal{P}(r, n_i, \mathcal{G})$, is defined as:

$$\mathcal{P}(r, n_i, \mathcal{G}) = \{(r, n_1, n_2, \dots, n_i) \mid \mathcal{G} = (\mathcal{N}, \mathcal{E}), n_i \in \mathcal{N}, (n_j, n_{j+1}) \in \mathcal{E}\} \quad (1)$$

Let $\hat{\mathcal{G}} = (\hat{\mathcal{N}}, \hat{\mathcal{E}})$ be the reconstructed DAG from the text, with this DAG also containing the same root node $r \in \hat{\mathcal{N}}$. The *hierarchical accuracy* is then defined as:

$$\text{Hierarchical Accuracy} = \frac{1}{|\mathcal{N}|} \sum_{n_i \in \mathcal{N}} \text{CA}(n_i, \mathcal{G}, \hat{\mathcal{G}}) \quad (2)$$

with $\text{CA}(n_i, \mathcal{G}, \hat{\mathcal{G}})$ being:

$$\text{CA}(n_i, \mathcal{G}, \hat{\mathcal{G}}) = \begin{cases} 1 & \text{if } n_i \in \hat{\mathcal{N}} \text{ and } \left(\bigcup_{n_j \in \mathcal{P}(r, n_i, \mathcal{G})} \cap \bigcup_{n_k \in \mathcal{P}(r, n_i, \hat{\mathcal{G}})} \right) \setminus r \neq \emptyset \\ 0 & \text{otherwise} \end{cases} \quad (3)$$

Hence, $\text{CA}(n_i, \mathcal{G}, \hat{\mathcal{G}})$ takes a positive value if node n_i shares a common ancestor with the ground truth taxonomy \mathcal{G} in the reconstructed taxonomy $\hat{\mathcal{G}}$. CA does not count the root node r as a common ancestor as this term is a common ancestor to all terms in both DAGs. The hierarchical accuracy is then the average over all nodes of the ground truth taxonomy.

3 Results & Discussion

Here we present the results obtained with the proposed method for ontology and knowledge graph generation from scientific literature. The performance of the method is first evaluated by measuring the ability to reconstruct an existing KG and ontology of chemical elements and functional groups from a corpus of text. The method is then applied to construct the first KG and ontology of the scientific field of SAC. The effectiveness of the ontology is then evaluated by applying it in a RAG-based solution for SACs domain.

3.1 Chemical Elements and Functional Groups Reconstruction

ElementKG¹⁸ is an ontology and KG of chemical elements and functional groups with 272 nodes, of which 188 are instances. This ontology and KG were curated manually by domain experts from information mostly available on Wikipedia. To reconstruct ElementKG, 76 Wikipedia pages corresponding to functional groups (e.g. Alkyne, Alkane, etc.) and groups of elements (e.g. Noble gases, Metalloids, etc.) were collected for our work. Except for the functional groups’ page, only the introduction section of the pages was used for the reconstruction, we observed that it contained most of the relevant information regarding taxonomical information. Additionally, four plain texts were manually curated for cases where no Wikipedia page exists (e.g. ferrous elements) or the information available is not enough to reconstruct ElementKG (e.g. rare metals).

Initially, the extracted vocabulary contained 1735 terms, with an accuracy of 94.30% compared to ElementKG vocabulary. A total of 555 relationships between terms were extracted from the corpus. The evaluation metrics (term accuracy, hierarchical accuracy, and KG accuracy) for the reconstruction of ElementKG are presented in Table 1. The table also reports the metrics for each of the two main categories in ElementKG, elements and functional groups. To account for terms that might be repeated in a taxonomy, repeated terms in the ground truth taxonomy are treated as separate terms.

Table 1: Evaluation metrics for the reconstruction of ElementKG averaged over 2 runs. Overall accuracy (all) and the more finegrained analysis on the element (E) and functional groups (FG) categories.

Evaluation	Iteration 1	Iteration 2	Iteration 3	Iteration 4
Term acc. (all)	0.835 ± 0.108	0.889 ± 0.073	0.898 ± 0.060	0.907 ± 0.052
Term acc. (E)	0.934 ± 0.030	0.952 ± 0.026	0.952 ± 0.026	0.955 ± 0.032
Term acc. (FG)	0.626 ± 0.320	0.659 ± 0.325	0.681 ± 0.293	0.693 ± 0.278
Hierarchical acc. (all)	0.842 ± 0.110	0.868 ± 0.102	0.876 ± 0.089	0.889 ± 0.075
Hierarchical acc. (E)	0.966 ± 0.007	0.965 ± 0.017	0.965 ± 0.017	0.976 ± 0.007
Hierarchical acc. (FG)	0.647 ± 0.291	0.724 ± 0.234	0.746 ± 0.203	0.761 ± 0.182
KG acc. (all)	0.749 ± 0.068	0.727 ± 0.000	0.709 ± 0.011	0.706 ± 0.015
KG acc. (E)	0.900 ± 0.053	0.902 ± 0.020	0.902 ± 0.020	0.911 ± 0.007
KG acc. (FG)	0.450 ± 0.301	0.394 ± 0.151	0.363 ± 0.159	0.344 ± 0.133

As shown in Table 1, our method demonstrates high accuracy in reconstructing the Element taxonomy. The three metrics considered consistently exceed 90% across all iterations, indicating that the initial

iteration provides a robust foundation for the reconstruction process. However, the accuracy for functional groups is comparatively lower. Notably, both term accuracy and hierarchical accuracy for functional groups show improvement over successive iterations, suggesting that new terms are being correctly integrated into the taxonomy structure. Conversely, the KG accuracy tends to decrease after the second iteration, implying that some terms are incorrectly positioned beneath nodes that should be leaf instances in the KG—i.e., terms that should not have any child nodes. We observed considerable variability in the functional groups’ performance across different runs, attributable to the inherent stochasticity of LLMs. This variability in the initial iteration significantly influenced the performance of subsequent iterations, highlighting the sensitivity of the process to initial conditions.

Table 2 shows general descriptive statistics of the reconstructed taxonomy. Note that the original ElementKG is considerably smaller in comparison, as this taxonomy has been narrowed down to only the required terms, while this work reconstructs a general-purpose taxonomy. The table illustrates how the number of *isA* relationships increases constantly over iterations. We observed that the root cause of this is general terms (e.g. Materials) that are placed in the taxonomy, and that, given their general nature, cause a large number of terms to be placed below them.

Table 2: Taxonomical properties of the reconstructed ElementKG across iterations averaged over 2 runs. More runs will be added in a future version.

Property	ElementKG	Iteration 1	Iteration 2	Iteration 3	Iteration 4
#nodes	272	777.0 ± 53.7	1066.5 ± 3.5	1147.5 ± 10.6	1178.5 ± 0.7
#isA	347	1007.0 ± 217.8	1691.0 ± 144.2	2003.0 ± 196.6	2151.5 ± 119.5
#leaves	188	578.5 ± 54.4	684.0 ± 11.3	691.5 ± 2.1	699.0 ± 1.4
Depth	8	17.0 ± 2.8	27.5 ± 9.2	31.5 ± 10.6	37.0 ± 9.8
Max #isA per node	32	134.0 ± 97.6	139.5 ± 98.3	145.0 ± 93.3	148.0 ± 89.0
Mean #isA per node	1.275	1.280 ± 0.190	1.580 ± 0.139	1.740 ± 0.152	1.825 ± 0.102
Mean steps to leaf	3.628	3.690 ± 0.426	4.130 ± 0.570	4.190 ± 0.499	4.200 ± 0.469

3.2 Ontology and KG generation for Single Atom Catalysis

SAC, a recent scientific domain studying the catalytic properties of isolated single metal atoms^{28,41}, presents an ideal case study for our method due to its mostly unstructured literature. We applied our procedure to generate the first ontology and KG of this field, using a corpus of 20 carefully selected papers on the synthesis and characterization of SACs. Focusing on the abstract and introduction sections, where concepts are typically introduced and explained, we extracted 1944 terms with 296 relationships. The category identification process initially yielded 331 categories, refined to 131, and manually curated to 12 key categories. Table 3 presents the main taxonomical properties of the reconstructed taxonomy.

After five iterations, the growth in nodes, *isA* relationships, and leaves slowed significantly, indicating convergence in taxonomy generation, despite not exhausting the entire vocabulary. This suggests the method’s effectiveness in structuring knowledge from an emerging scientific field.

Table 3: Taxonomical properties of the generated SAC ontology.

Property	Iteration 1	Iteration 2	Iteration 3	Iteration 4	Iteration 5
#nodes	1308	1357	1370	1378	1379
#isA	1574	1755	1810	1881	1894
#leaves	1274	1314	1323	1329	1328
Mean #isA per node	1.1888	1.2711	1.285	1.3091	1.3161
Max #isA per node	349	367	378	385	386
Depth	5	6	6	6	6
Mean steps to leaf	2.0259	1.9955	1.9956	1.9956	1.9956

GraphRAG-based downstream evaluation To demonstrate the practical value of our generated ontology, we conducted a comparative study in the SAC domain. We compared two approaches: a standard KG approach and an enhanced KG+Ontology approach, both integrated into a Retrieval-Augmented Generation (RAG) solution based on the GraphRAG procedure¹⁷.

The GraphRAG pipeline is composed of the following steps: (1) the generation of the KG from plain text, (2) description generation for each term and relationship, (3) clustering for community generation,

(4) community description generation, and (5) community retrieval for answer generation. In our enhanced approach, we extended the KG in step (1) with our generated ontology (KG+ontology).

To evaluate performance, we used the same four metrics as the original GraphRAG study¹⁷: comprehensiveness, diversity, empowerment, and directness. We created a dataset of 50 questions using long-context LLMs, covering various SAC topics such as synthesis, characterization methods, and applications, which were then validated by domain experts.

Following the GraphRAG methodology¹⁷, we used an LLM, in our work Anthropic’s Claude 3.5 Sonnet, to evaluate the quality of generated answers for both approaches. Table 4 presents the results, clearly showing that our KG+Ontology approach significantly outperforms the standard KG-only method across all metrics. This improvement demonstrates the ontology’s ability to enhance term relationships, enabling better clustering and retrieval, ultimately resulting in more informative and higher-quality answers. A comparison of answers generated with both approaches can be found in Appendix C.

Table 4: Win-rate according to different criteria in the GraphRAG setting with a dataset of 50 questions.

Model	Comprehensiveness	Diversity	Empowerment	Directness
GraphRAG (KG)	1/50	4/50	1/50	16/50
GraphRAG (KG + Ontology)	49/50	46/50	49/50	34/50

Expert Evaluation To assess the quality of the generated ontology, a panel of experts conducted an evaluation of the taxonomical relationships. The experts randomly sampled relationships from various iterations of the ontology. Their task was to determine whether each sampled relationship was correct within the context of the domain. The results of this evaluation showed that, on average, 64.5% of the examined relationships were deemed correct by the experts. While this indicates a majority of accurate relationships, it also suggests room for improvement in the ontology generation process. Upon analysis of the incorrect relationships, the experts did not identify any clear patterns of errors. However, they provided three key observations for potential enhancement:

- *Specificity*: The quality of relationships could be improved by making them more specific, thereby reducing ambiguity and increasing precision.
- *Context Dependency*: Some extracted vocabulary terms are heavily context-dependent and may lose their intended meaning when placed in a more general context. This suggests a need for better context preservation or clearer domain boundaries.
- *Semantic Redundancy*: The taxonomy contains instances of semantically similar concepts repeated in different parts of the structure. This redundancy could be addressed to streamline the ontology.

These expert insights provide valuable direction for refining the ontology generation process and improving the overall quality of the taxonomical relationships.

4 Conclusion

Extraction of structured knowledge from scientific literature is a challenging task that is usually performed manually by domain experts. Given the success of LLMs in NLP tasks, this work proposes the use of LLMs as an information extraction engine from the scientific literature. A five-step pipeline is proposed that is able to generate ontologies and KGs with an in-context nature and without the need for any training data. The pipeline is then implemented entirely using openly available LLMs and long-context LLMs. The evaluation of the approach is done by reconstructing an existing KG and ontology of chemical elements and functional groups from the literature. The results show good accuracy in reconstructing the taxonomies and KGs with a term accuracy and hierarchical accuracy of over 80% in all iterations, and knowledge graph accuracy of over 70% in all iterations. It was observed, however, that the quality of the reconstruction is dependent on the first iteration generation, and that given the stochastic nature of the LLMs, there is variability in the results. The method was then applied to the emerging domain of SACs. Given the recent development of the field, the literature lacks structured information, which makes the generation of an ontology and KG a challenging task.

It was shown that the in-context generated ontology of SACs was able to improve the performance of the GraphRAG pipeline in question-answering tasks related to SACs. Finally, the quality was evaluated by a group of experts, who evaluated that 64.5% of the relationships were correct. We can thus conclude that the proposed approach is able to extract meaningful information from scientific literature.

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A Technical Stack and Experimental Setting

The following technical stack was used for the pipeline implementation:

- *Plain text extraction*: Plain text was extracted from the paper’s PDF files using Nougat⁴ with model 0.1.0-base. When the extraction with Nougat failed, extraction resorted to PyMuPDF².
- *Terms Extraction*: The split of the initial text into sentences is done with a RoBERTa base model³² from the Spacy suite³. Multiple sentences are then concatenated as long as they don’t exceed a maximum length of 2048 characters. Vocabulary extraction was performed with a Llama 3.1 70B model¹⁶ quantized to 4 bits, with a maximum context length of 1024 tokens. It was observed that a greedy generation (temperature=0) provided the best results. For the other parameters, we set $k = 5$.
- *Category generation*: The *generation* step is performed with Llama-3 8B Gradient Instruct 1048k quantized to 8 bits, context length limit of 128K tokens and temperature=0.7. The *refinement* step was done with Llama-3.1 70B quantized to 4 bits, context limit of 32K tokens and temperature=0.5. After each of the two steps, an extra prompt is performed with the former model to format the output of the LLMs. A total of 80 different samples were generated both in the generation and refinement steps.
- *Taxonomy generation and relationships extraction*: Both taxonomy generation and relationships extraction are done with a Llama 3.1 70B model¹⁶ quantized to 4 bits, with a maximum context length of 32K tokens and temperature=0.1. For each LLM query, self-consistency was applied by taking the majority voting of 3 sampled answers.

For the GraphRAG setting, the following hyperparameters were used:

- *Text split and vectorization*: Chunk length of 1200 characters with a 100 characters overlap. Text embeddings were computed using the all-MiniLM-L6-v2 pretrained model from Sentence Transformers framework⁴².
- *Node2vec*²³: Dimensionality of 1536, with 10 walks of length 40 and window size 2.
- *Clustering*: Clustering was performed with the Leiden algorithm⁴⁶ as in the original GraphRAG work. Maximum graph cluster size of 10 nodes.
- *LLM evaluation*: The LLM evaluation of the answers was done using claude-3-5-sonnet-20240620.

LLM inference was done using the Ollama suite⁴. All the technical decisions made were adjusted to a budget of a single NVIDIA RTX A6000 graphics unit with 48GB. The source code is publicly available under MIT Licence.

B Prompt sample

This Section includes a sample question and answer obtained from the taxonomy generation phase. Note that in the question below, given the restriction in the context length, the whole taxonomy cannot be included in the prompt. As a consequence, only the term and the main category it belongs (in parenthesis) are included. Notice also that, in the answer provided, not all the terms as classified, as some of them are classified as ‘None’.

²github.com/pymupdf/PyMuPDF

³github.com/explosion/spaCy

⁴github.com/ollama/ollama

Sample Question

Given this context:

====

The actinide () or actinoid () series encompasses at least the 14 metallic chemical elements in the 5f series, with atomic numbers from 89 to 102, actinium through nobelium. (Number 103, lawrencium, is sometimes also included despite being part of the 6d transition series.) The actinide series derives its name from the first element in the series, actinium. The informal chemical symbol An is used in general discussions of actinide chemistry to refer to any actinide.

All the actinides are f-block elements. Lawrencium is sometimes considered one as well, despite being a d-block element and a transition metal. The series mostly corresponds to the filling of the 5f electron shell, although as isolated atoms in the ground state many have anomalous configurations involving the filling of the 6d shell due to interelectronic repulsion. In comparison with the lanthanides, also mostly f-block elements, the actinides show much more variable valence. They all have very large atomic and ionic radii and exhibit an unusually large range of physical properties. While actinium and the late actinides (from curium onwards) behave similarly to the lanthanides, the elements thorium, protactinium, and uranium are much more similar to transition metals in their chemistry, with neptunium, plutonium, and americium occupying an intermediate position.

All actinides are radioactive and release energy upon radioactive decay; naturally occurring uranium and thorium, and synthetically produced plutonium are the most abundant actinides on Earth. These have been used in nuclear reactors, and uranium and plutonium are critical elements of nuclear weapons. Uranium and thorium also have diverse current or historical uses, and americium is used in the ionization chambers of most modern smoke detectors. Of the actinides, primordial thorium and uranium occur naturally in substantial quantities.

====

,and given the following taxonomy:

====

Carbon (Element of the periodic table)

Element of the periodic table

Hemiacetals (Functional group)

R2C=O (Functional group)

Aldehyde hydrate (Functional group)

Carbonyl (Functional group)

Aldehyde (Functional group)

Mixed acetal (Functional group)

Ketone hydrate (Functional group)

Carbonyl compounds (Functional group)

Acetal (Functional group)

Esters (Functional group)

Ketone-derived structures (Functional group)

Symmetric acetal (Functional group)

Orthoester (Functional group)

Hydroxyl group (Functional group)

Ketal (Functional group)

Functional group

====

Complete the following list to classify the terms into the taxonomy according to the context.

If a term does not fit in any of the categories, say "None".

If the text is not clear enough to classify a term, say "None".

If the text does not explicitly mention that a term is a type of another, say "None".

If multiple categories apply, choose the most specific one.

In the output include only the classification. Do not include any explanation or additional information.

Do not classify a term to be its own parent, this is, do not output answers such as "A isA A".

The answer should include 'isA'.

Nuclear weapons isA

Lawrencium isA

F-block elements isA

Thorium isA

Radioactive decay isA

Actinides isA

Actinoid isA

Neptunium isA

Actinium isA

Actinide isA

Curium isA

Ground state isA

Primordial thorium isA

Americium isA

6d transition series isA

Plutonium isA

Protactinium isA

Lanthanides isA

Atomic numbers isA

Smoke detectors isA

Nuclear reactors isA

Ionic radii isA

Uranium isA

Electron shell isA

Transition metals isA

D-block element isA

Ionization chambers isA

5f series isA

Valence isA

Interelectronic repulsion isA

Nobelium isA

Metallic chemical elements isA

Sample Answer

Here is the list of classifications:

- Nuclear weapons isA None
- Lawrencium isA Actinide
- F-block elements isA None
- Thorium isA Actinide
- Radioactive decay isA None
- Actinides isA Metallic chemical elements
- Actinoid isA Actinide
- Neptunium isA Actinide
- Actinium isA Actinide
- Actinide isA F-block elements
- Curium isA Actinide
- Ground state isA None
- Primordial thorium isA Thorium
- Americium isA Actinide
- 6d transition series isA Transition metal
- Plutonium isA Actinide
- Protactinium isA Actinide
- Lanthanides isA F-block elements
- Atomic numbers isA None
- Smoke detectors isA None
- Nuclear reactors isA None
- Ionic radii isA None
- Uranium isA Actinide
- Electron shell isA None
- Transition metals isA Metallic chemical elements
- D-block element isA Transition metal
- Ionization chambers isA None
- 5f series isA F-block elements
- Valence isA None
- Interelectronic repulsion isA None
- Nobelium isA Actinide
- Metallic chemical elements isA Element of the periodic table

C GraphRAG sample

Question

What types of supports do scientists commonly use in preparing single atom catalysts?

GraphRAG (KG) Answer

****Common Supports for Single Atom Catalysts****

Scientists employ a variety of supports when preparing single atom catalysts. According to the available data, the most commonly used supports are:

Primary Supports

Nitrogen-doped graphene, graphene oxide, and carbon materials are the most frequently used supports in the preparation of single atom catalysts. These materials have been widely adopted due to their unique properties, which make them well-suited for this application.

Secondary Supports

In addition to the primary supports, defect-rich zirconia is also used as a support in some cases. This material has been found to be effective in certain catalytic reactions, and its use is becoming increasingly popular.

Other Supports

Other supports, such as metallic surfaces, metal oxides, and metal sulfides, are also being explored in the field of single atom catalysis. While these materials are not as widely used as the primary supports, they have shown promise in certain applications and may become more prominent in the future.

It is worth noting that the choice of support material can significantly impact the performance of the single atom catalyst. As such, researchers continue to investigate new and existing materials to optimize their properties and improve catalytic efficiency.

GraphRAG (KG + Ontology) Answer

****Common Supports Used in Single Atom Catalysts****

Scientists commonly use various types of supports in preparing single atom catalysts to provide a stable platform for the single atoms to anchor on, allowing for improved catalytic performance and stability.

****Oxide Supports****

Oxide supports, such as titanium dioxide (TiO₂) and silicon dioxide (SiO₂), are commonly used to isolate metal catalytic sites and provide a stable platform for single atom catalysts. Anatase TiO₂ is also mentioned as a specific example of an oxide support used in single atom catalysts.

****Conductive Carbon Supports****

Conductive carbon supports, such as graphene substrates and carbon nanotubes, are used to provide good electrical connection and performance. These supports can also serve as active sites for catalytic reactions, further highlighting the importance of support material selection.

****Other Types of Supports****

Other types of supports used in single atom catalysts include:

- * Metal hydroxides
- * Zeolites
- * Mesoporous silica
- * Metal-organic frameworks (MOFs)
- * Metallic surfaces
- * N-Doped Carbon Matrix

****Importance of Support Material Selection****

The choice of support material can significantly impact the performance and stability of single atom catalysts. Researchers may select supports based on their surface properties, such as acidity or basicity, as well as their ability to anchor the single atoms and prevent sintering or agglomeration.

****Hybrid Support Materials****

In some cases, scientists may use multiple supports or combine different supports to create a hybrid support material. This can allow for the creation of single atom catalysts with tailored properties and improved performance.

****Reversible Hydrogen Electrodes (RHE)****

Reversible hydrogen electrodes (RHE) are used as reference electrodes in experiments and provide a stable potential for comparison. However, they are not typically used as supports for single atom catalysts.