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## **Knowledge Graph Extraction from Total Synthesis Documents**

Anonymous Authors<sup>1</sup>

## Abstract

Knowledge graphs (KGs) have emerged as a pow-011 erful tool for organizing and integrating complex information, making it a suitable format for scientific knowledge. However, translating scientific knowledge into KGs is challenging as a wide va-015 riety of styles and elements to present data and ideas is used. Although efforts for KG extraction (KGE) from scientific documents exist, evalua-018 tion remains challenging and field-dependent; and existing benchmarks do not focuse on scientific 020 information. Furthermore, establishing a general benchmark for this task is challenging as not all scientific knowledge has a ground-truth KG representation, making any benchmark prone to ambiguity. Here we propose Graph of Organic Synthe-025 sis Benchmark (GOSyBench), a benchmark for KG extraction from scientific documents in chemistry, that leverages the native KG-like structure 028 of synthetic routes in organic chemistry. We de-029 velop KG-extraction algorithms based on LLMs 030 (GPT-4, Claude, Mistral) and VLMs (GPT-40), the best of which reaches 73% recovery accuracy and 59% precision, leaving a lot of room for improvement. We expect GOSyBench can serve as 034 a valuable resource for evaluating and advancing 035 KGE methods in the scientific domain, ultimately facilitating better organization, integration, and discovery of scientific knowledge.

## 041 **1. Introduction**

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Knowledge graphs (KGs) have emerged as a powerful tool for representing and organizing complex information, enabling efficient storage, retrieval, and analysis of data across various domains (Hogan et al., 2021). The extraction of knowledge graphs from unstructured data sources, such as text documents, has gained significant attention in recent years due to its potential to unlock valuable insights and facilitate knowledge discovery. KGs have also recently been used in Retrieval-Augmented Generation (RAG) pipelines (Abu-Rasheed et al., 2024), as a strategy to ground text generation from large language models (LLMs) with domainspecific facts, thus improving performance across tasks (Khattab et al., 2023; Khattab & Zaharia, 2020).

#### 1.1. Extraction of Knowledge Graphs

The field of Knowledge Graph Extraction (KGE) has witnessed substantial progress, with numerous approaches being developed to automatically construct KGs from textual data. These methods range from rule-based systems to machine learning-based techniques, and more recently, LLMdriven extraction (Meyer et al., 2023; Shu et al., 2024). Several benchmarks have been proposed to evaluate the performance of KGE systems, from open-domain ones like Open Graph Benchmark (Hu et al., 2020) and Text2KGBbench (Mihindukulasooriya et al., 2023), to more field specific ones like PharmaKG for biomedical data mining (Zheng et al., 2020). These benchmarks focus on evaluating algorithms on the extraction of specific facts from short sentences or paragraphs, while extraction from complete documents, and specially scientific ones, remains largely untested.

Scientific literature contains a wealth of knowledge that can be represented in KGs, the extraction of which would enable more efficient knowledge integration and facilitate discovery. Excellent efforts have been made to extract specific types of scientific information, such as entities and relations in chemical literature (Lowe & Sayle, 2013; Swain & Cole, 2016; Mavračić et al., 2021). While these advances have enabled the extraction of influential reaction datasets (Lowe, 2012), they are tailored to patents, which have a more standardized format and contain less scientific details as journal papers do. Moreover, these methods focus on extracting single reactions or short sequences, mostly ignoring the underlying network of objects and concepts originally expressed in the texts.

The lack of benchmarks specifically designed for evaluating KGE in science poses a challenge, as the diverse nature of scientific knowledge and the absence of ground-truth KGs make it difficult to establish a standardized evaluation frame-

<sup>&</sup>lt;sup>1</sup>Anonymous Institution, Anonymous City, Anonymous Region, Anonymous Country. Correspondence to: Anonymous Author <anon.email@domain.com>.

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*Figure 1.* Example Knowledge Graph and evaluation strategy. **a.** Shows the data representation used for the task, where each node Si in the directed graph represents abstractly a substance, and each edge V ( $i \rightarrow j$ ) expresses that substance Sj is used in a reaction that has substance Si as a product. The goal of the KG is to accurately represent the information presented in the paper. **b.** Evaluation methodology followed in this work. **c.** Summary statistics of the resulting dataset. These highlight aspects critical to graph complexity, like number of substances (nodes), maximum path length, number of head nodes (indegree(Si) = 0), among others. **d.** Algorithm developed for KGE.

work. The heterogeneity of scientific literature, with its wide range of domains, writing styles, and presentation formats, further complicates the development of a comprehensive benchmark.

#### 1.2. KGs in Organic Chemistry

A knowledge graph is defined generally as a graph of data,
intended to convey knowledge. Here, nodes represent entities of interest and edges represent relations between these
entities (Hogan et al., 2021). As such, synthetic sequences
in Organic Chemistry are susceptible of being represented
under such a structure.

Research in synthetic organic chemistry (OC) focuses very generally on the synthesis of organic compounds through a suitable sequence of reactions. Under this conception, substances are *concepts* that are connected through reactions as *relationships*. Each substance may serve as product or reactant for a multitude of different reactions, leading to the natural definition of networks of chemical reactions. This has previously been studied under different models with different levels of depth (Fialkowski et al., 2005). This bare abstraction defines the backbone of a KG, and is this native KG-like structure makes OC an ideal domain for exploring KGE techniques.

But reactions –defined as an experimentally executed transformation that leads from one substance to another– are not the only type of relationships that may exist between substances. In research works in OC, substances are synthesized not only because they will be directly used as building blocks for the synthetic targets, but some are synthesized also to serve as model systems for more complex and valuable structures, some are synthesized but paths need to be abandoned due to unsuccessful reactions, and sometimes even substances are synthesized to facilitate structural elucidation of their precursors. Indeed, many more relationships are built on top of the reaction-graph backbone, that are of interest for organic chemists: these go beyond to inform about strategic aspects of synthesis and multi-level chemistry-driven decision processes.

This work focuses mainly on the extraction of the main backbone from research papers. These are typically given in papers' Supporting Information (SI) files, and contain

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detailed descriptions of synthetic routes and experimental 111 procedures. These documents exhibit a wide variety of 112 representations, designs, and conventions, making it chal-113 lenging to extract consistent and comprehensive KGs, see 114 Appendix A for examples. Despite the heterogeneity in 115 the representation of OC knowledge, the underlying struc-116 ture remains the same: a network of chemical reactions 117 and synthetic plans. This property allows for the definition 118 of a ground-truth KG, making OC a suitable domain for 119 developing and evaluating KGE methods in science.

120 In this paper, we propose **GOSyBench**, a benchmark for 121 KGE from scientific documents in the domain of organic 122 chemistry. By leveraging the native KG-like structure of 123 synthetic routes, we aim to provide a standardized eval-124 uation framework for assessing the performance of KGE 125 algorithms in extracting scientific knowledge. Our KG on-126 tology defines substances as entities, with reference\_key and 127 substance\_name as properties, that are connected by reac-128 tions as relationships. Furthermore, we develop novel KGE 129 algorithms based on LLMs, and conduct extensive exper-130 iments and ablation studies to validate their effectiveness 131 using our proposed benchmark. 132

## 2. Methods

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#### 2.1. Guidance / structured output generation

137 Despite their usefulness in various domains, one of the lim-138 itations of LLMs is their incapacity to generate consistent 139 and controllable outputs that fit use-case specific guidelines. 140 Recent research has focused in steering LLM generation 141 through the enforcement of grammars in the resulting gen-142 erations (Rebedea et al., 2023; Khattab et al., 2023). This 143 not only helps steer models towards non-harmful outcomes, 144 but also enables tool usage in agent-like scenarios (Boiko 145 et al., 2023; M. Bran et al., 2024) and facilitates parsing of 146 the results and integration in existing software (Liu, 2024). 147

#### 2.2. Benchmark dataset curation 149

The dataset curation pipeline used involved a combination of 150 automated knowledge extraction and expert human labeling. 151 Initially, 24 Supplementary Information files (SIs) on total 152 synthesis were manually selected from the Journal of the 153 American Chemical Society (JACS), with the format and 154 content of their SI used as a criterion. The SIs were selected 155 such that the obtained sample represents a wide variety of 156 text formatting, varying use of visual elements, order and 157 location of relevant sections, among others, see Appendix 158 A for examples. 159

160 The SIs were then processed using the KGE method pre-161 sented in Section 2.3, resulting in a collection of 24 knowl-162 edge graphs, where each contains an approximation to the 163 complete network of chemical reactions expressed in the SI. 164

The processed then continued with manual curation, which generally involved node relabeling, node creation/removal, and edge creation/removal. The resulting objects are directed graphs, with individual substances as nodes, and reactions as edges. Some statistics of the dataset are described in Figure 1, which highlights the size and overall complexity of the KGs being extracted.

#### 2.3. KGE method

The Knowledge Graph Extraction method developed for this work has several steps, as shown in Figure 1d. Initially, the SI PDF is pre-processed to select the relevant sections describing the reaction procedures, as explained in more detail in Appendix B. This aims to lower the amount of text that needs to be processed in the steps following, and prevents errors by erroneous addition of spurious nodes to the graph. The PDF is then processed into text and split into single text segments describing chemical reactions. Two methods were tested for this: one based in rule-based text parsing from PDF, and one based in Vision-Language Models (VLMs), namely the recent GPT-40 by OpenAI. The latter method was implemented in view of the variability of representations and interleaved use of visual elements observed in SIs, as shown in Appendix A.

Resulting reaction blocks are then each processed individually by an LLM-powered generation pipeline, that detects and extracts all the substances declared in the input reaction. Each of these substances is represented as a structured object containing three main properties: reference\_key, substance\_name, and role\_in\_reaction. Each collection of substances is converted into a *reaction\_unit*, a structured object resembling a node in a tree, where the head node is the product of the reaction and the children are all the substances with a role different than product.

Finally, a graph is constructed by connecting all the different reaction\_unit objects, using each substance's reference\_key as the node label.

The reported benchmark was used to perform ablations on 3 of the design choices for the algorithm, namely to test the effect of SI preprocessing to select relevant sections, the use of rule-based or vision-based PDF parsing, and the choice of LLM used for structured object generation. The results are shown in Figure 2.

#### 2.4. PDF Parsing methods

Two parsing methods have been tested in this work. One is a simple, rule-based algorithm that is based on general observations from the structure of SIs in organic chemistry papers, while the other is fully driven by a Vision-Language Model (VLM), which aims to recover information by directly processing documents as humans would read it, without loss of 165 visual elements.

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168 This approach consists of parsing the input PDF file using 169 the PyMuPDF package (noa), which yields the complete 170 text from the PDF, including titles and paragraphs, but also 171 formatting details such as bold letters. Unfortunately it also 172 includes spurious formatting details like page numbers and 173 side notes from journals. Using this information, the text 174 is split using "long sequences of bold letters" as a splitting 175 criteria, which leads to a list of text segments. The idea 176 behind this parsing is that most authors state products in 177 bold font with the name of the product (IUPAC, or simply 178 a reference name), followed by a reference key, and then 179 proceed with the description of the reaction procedure in 180 normal font (see Appendix A). This pattern is somewhat 181 consistent and in some cases leads to very nicely parsed 182 documents. 183

1841852.4.2. IMAGE-BASED — VISION

The effectiveness of the rule-based method above is endangered by the variety of formats and representation styles that authors decide to use in their papers, as shown in the Appendix A. Understanding of these documents is heavily dependent on the reader's ability to interpret the visuals and contrast them and connect them with the text, thus the purely rule-based method falls short in some cases.

Leveraging the recent advances in VLM research, we propose directly using one such model for this task. In particular, we use the recently released GPT-40, one of the most powerful end-to-end Large Multimodal Models (LMMs) from OpenAI.

The pipeline starts with the conversion of the input PDF into a suitable format, and for this we simply convert each page from the PDF into a png image using the pdf2image package (Belval, 2024). The images are then processed into overlapping batches of images, each batch in a single VLM call. This process ensures that the VLM sees a more global structure of the paper and thus has better context to give an appropriate response.

The VLM is then queried with all the images from a batch and a prompt with instructions (see Appendix C). The expected output of this is a summary of the relevant information for *each* reaction the VLM can identify in the image context; each reaction separated by a given separator token.

#### 2.5. Evaluation metrics

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A wealth of methods exist to compare graphs, each suitable
for certain sets of use cases (Thompson et al., 2022; Shimada et al., 2016; Hartle et al., 2020). These include direct
comparison of the node or edge sets, subgraph matching,

spectral analysis, and the use of graph kernels, among others. In this work, we take an approach based on subgraph matching, that aims to capture the similarities relevant to synthetic routes in organic chemistry.

Appealing to the specific structure of the types of graphs used in this work, namely directed graphs with mostly a tree-like structure, we use 3 metrics based on the ratio of paths shared between the compared graphs, as shown in Equation 1.

$$S(G, G') = \frac{1}{|G|} \sum_{p \in P_S(G)} \sum_{p' \in P_S(G')} 1_{p=p'}$$
(1)

Where  $P_S(G)$  defines the set of all the linear paths p in G, and the  $1_p = p'$  operator is defined as 1 if the condition p = p' is met, 0 otherwise. The key difference between the methods used here is the definition of the equivalence operator =, which can take multiple forms depending on the property of interest. In particular, two options are defined: exact match and preservation of partial order. Exact match directly compares the two paths based on the exact sequence of nodes defined by each. This method thus directly measures to what extent the exact KG is reconstructed from documents.

The second method aims to capture a more nuanced structure in the retrieved KGs, through a slightly less strict comparison metric based on ordered sets. In this method, two paths are considered equivalent if the order relationships defined by each path are preserved in the other. Take for example the following two paths

$$p_0 = 6 \to S2 \to 7$$
$$p_1 = 6 \to 7$$

Where  $p_0$  defines the order  $6 \succ S2 \succ 7$ . In this example,  $p_0 \neq p_1$  under exact match, however they are under the *PO* equivalence as the order relationship  $6 \succ 7$  exists in both paths. Such a less strict definition is particularly relevant in our case as it is typical in SIs to describe the formation of an intermediate and continue using it "without further purification". In these cases, the complete sequence  $p_0$  with intermediate *S*2 may be reduced by the extraction models to  $p_1$ , which is not necessarily incorrect however missing some information.

A last method is used, which uses exact match as equivalence operator, but both G and G' are preprocessed to remove the leaves (nodes with outdegree(n) = 0), thus only comparing the backbone of the synthetic tree without considering reagents. Figure 1b shows such removed nodes in yellow, and the nodes belonging to the backbone in red.

## 3. Results

The proposed benchmark was used to perform ablations on 3 of the components of the KGE algorithm described in Section 2.3. Namely, we assess the effect of SI preprocessing (Appendix B), the parsing of PDFs using a rulebased approach, or directly through Vision-Language Models (VLMs), and the choice of LLM for parsing of reaction descriptions into formatted reaction units. In addition, we evaluate the performance of multiple LLMs from different providers on the latter task across multiple metrics using a more specific benchmark, aimed at selecting suitable LLMs for this task, without the need to execute the whole extraction pipeline.

#### 3.1. KGE Benchmark

The aim of these experiments is to determine the effectivity of a given system at extracting a KG in the required format, not only at assessing the capabilities of LLMs, hence 2 binary variables are ablated that deal with document preprocessing, parsing and chunking. The latter is the LLM used, however here we have restricted ourselves to only testing models provided by OpenAI, mainly due to rate limit constraints from the other providers.

In Figure 2 we display the per-paper performance for each variation of the system in Figure 1d, across six metrics, all different forms of accuracy (left column=) and precision (right column) of synthetic path recovery. The upper row shows the results on exact path reconstruction, middle row a more relaxed version of this based on comparing the orders defined by each path, and bottom row compares the pruned graphs, assessing the similarity between the tree backbones; see Section 2.5 for details.

For each comparison method,  $S(G_{EX}, G_{GT})$  measures the system's ability to reconstruct Ground Truth paths — highly important for organic chemistry as it defines the specific sequence of reactions, while  $S(G_{GT}, G_{Extracted})$  measures the precision or "purity" of the resulting graphs, thus also accounting for erroneous introduction of nodes or edges in the extraction process.

The results show that the overall performance varies widely as a function of the paper, which is to be expected given 263 the high variability in styles and formats used in these docu-264 ments (see Appendix A). A systematic difference is found 265 between the 2 models tested, with a clear advantage for 266 GPT-4-turbo, the most advanced model, especially on reconstruction accuracy. The gap is nevertheless reduced in reconstruction precision which, as will be shown in the next 269 section, can be attributed to the smaller model being better 270 at detecting wrong inputs, thus introducing less noise into 271 the extracted KG. 272

Interestingly, comparing the pruned graphs demonstrates

GPT-3.5's poor performance on precision, with most values below 0.1, however the corresponding accuracy is relatively high, even surpassing GPT-4 based methods on the same metric. Such results imply that smaller models perform poorly in general conditions, however the information recovered by these is typically valid. More advanced models seem not to have a strong filter and generate valid structured outputs despite noisy filters, which in turns generate accurate but noisy KGs. These observations will be further elaborated in the following section.



Figure 2. Results from Knowledge Graph Extraction benchmark. System performance on GOSyBench for multiple system ablations. The two main columns show accuracy (left) and precision (right). Each sub column shows the result for PDF parsing methods text-based (left) and vision-based (right). Rows present different metrics used for graph comparison, and the color distinguishes between SI pre-processing methods.

From the results presented here it seems that using vision models like GPT-40 (columns in Figure 2), or preprocessing the document before to select the most relevant parts of the SI (colors in Figure) do not improve the system's performance. Vision only helps slightly improve the accuracy of
the system when a smaller model is used, however such
system still underperforms relative to the larger GPT-4.

A more in-depth exploration of the results is needed to
determine how to best leverage vision models for this task.

## 3.2. LLM Performance across tasks

283 To assess the effect of the choice of LLM in the KGE 284 method developed in this work, another benchmark with 285 a narrower scope was produced. The benchmark aims to 286 assess LLM's abilities to recover specific information from 287 reaction description text samples. This involved the creation 288 of 3 smaller datasets, each designed to test the models at 289 specific tasks, namely ability to recognize and retrieve the 290 correct product and reactant sets, ability to produce empty 291 responses whenever a non-reaction text is given, and the ability to correctly retrieve the *reference\_key* of substances. 293

All of these are elements of utmost importance for the algorithm's success at reconstructing a paper's KG, as failure to correctly perform these contaminates the resulting KG with spurious nodes and edges, and leads to the loss of real nodes and edges.

299 For the sake of completeness and ease of implementation, 300 we have tested LLMs from 3 API providers, namely OpenAI, 301 Anthropic and Mistral. Moreover, the models tested span a 302 wide range of sizes and scores on standard benchmarks (). 303 As shown in Figure 3, the top-performing model in terms of 304 product and reactants retrieval accuracy is gpt-4-turbo, on 305 of the most advanced models as shown by benchmarks, in 306 terms of reasoning capabilities. Nevertheless, other models, 307 some smaller and far cheaper, perform almost on-par with 308 gpt-4 on this metric (mistral small and medium, mixtral 309 8x7b, all claude models). 310

Surprisingly, the "smarter" models do not perform as good 311 on other tasks, particularly "Wrong inp" and "Key exact". 312 Smaller, less poweful models, like mistral-small, mixtral-313 8x7b and gpt-3.5-turbo do better in rejecting wrong inputs 314 than their more advanced counterparts despite their less 315 developed reasoning capabilities. An important observation 316 is that, when given a non-reaction text, smaller models give an error as they fail to find the requested information and 318 fail to produce an answer in the requested format, thus being 319 caught as exception during model validation. In counterpart, 320 larger models tend to give a response, despite the input text not containing the desired information, typically through 322 hallucinations. 323

In spite of these observations, the ablations in Section 3.1
have been performed only with OpenAI models as we had
higher rate limits, allowing us to perform multiple experiments concurrently.



Figure 3. Capability-specific benchmark for LLMs. The performance of multiple LLM across multiple scales and providers is shown. Models are evaluated on 4 metrics: **Prod ret** evaluates the accuracy of retrieving the correct product name from an input paragraph (which involves separating product name from its reference key), **React ret** evaluates the same, for retrieval of reactants used in the described reaction, **Wrong inp** assesses how good the models are at rejecting inputs that do not describe a chemical reaction, and **Key exact** evaluates the ability of models to output the exact reference key for products.

#### 4. Conclusions

We have proposed a novel benchmark for knowledge graph extraction in science from full papers. We exploit the native KG-like structure of synthetic organic chemistry and propose a benchmark with 24 manually curated papers. This benchmark is continuously growing to incorporate more high quality samples of challenging papers. We developed an LLM-based algorithm for KGE and evaluate each individual part using a small, handcrafted benchmark to test the capabilities of LLMs for each specific task, and find that advanced models have better recall of input context, however smaller models are advantageous to detect text that can not be identified as a reaction, thus not contaminating the generated KG with spurious nodes. Finally, we perform ablations on our algorithm and show that the usage of Language-Vision Models (LVMs) does not directly improve the system's performance, despite having empirical reasons to believe so. Overall, there is still a lot of room for improvement as our algorithms reach a maximum of 73% average in accuracy, and 59.7% in precision. More work

LLM Performance on individual tasks

needs to go into desiging and optimizing algorithms for this
task, however we believe the release of GOSyBench sets
the field into the right direction by providing a challenging,
diverse and high-quality dataset for benchmarking.

# 5. Future work and outlook

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337 The efforts presented here deal with the extraction and evalu-338 ation of the reaction networks from chemistry papers, which 339 is only the backbone structure of a much richer KG for 340 organic chemistry. However as discussed in Section 1.2, ad-341 ditional relationship types between substances are implicitly 342 reported in papers, such as failed reactions and abandoned 343 synthetic plans, use of substances as model systems, among 344 others. All these are important details that describe not only 345 a successful route to a target substance, but encode also 346 the difficulties, lessons, and other valuable insights that are 347 reported in chemistry papers. From early experiments, we have found that extracting such new connections is possible 349 with LLMs thanks to their summarizing and reasoning ca-350 pabilities. Achieving such a milestone has the potential to 351 unlock promising advances in reaction search and chemical 352 knowledge retrieval in general.

353 In addition to this, the currently presented ontology can 354 further be enhanced with additional substance properties 355 reported in papers. Starting with extraction of the SMILES 356 strings for each molecule (Mavračić et al., 2021; Rajan et al., 357 2021; 2023), along with yields, scalability, and analytical 358 results, the resulting KGs can continuously be populated 359 with more substance-specific details to better represent the 360 knowledge in papers. 361

Additionally, papers report multiple visualizations that display different views, or highlight different aspects of the
molecules and reactions in question. The interplay between
text and image modalities is strong in papers, and leveraging VLMs will be an essential step towards better KGE in
chemistry, as has been shown in this work.

## References

369

- pymupdf/PyMuPDF: PyMuPDF is a high performance
  Python library for data extraction, analysis, conversion
  & manipulation of PDF (and other) documents. URL
  https://github.com/pymupdf/PyMuPDF.
- Abu-Rasheed, H., Weber, C., and Fathi, M. Knowledge graphs as context sources for llm-based explanations of learning recommendations. *ArXiv*, abs/2403.03008, 2024. URL https://api.semanticscholar. org/CorpusID:268249177.
- Belval, E. Belval/pdf2image, May 2024. URL https://github.com/Belval/pdf2image.original-date:
  2017-05-28T19:00:59Z.

- Boiko, D. A., MacKnight, R., Kline, B., and Gomes, G. Autonomous chemical research with large language models. *Nature*, 624(7992):570–578, 2023.
- Fialkowski, M., Bishop, K. J. M., Chubukov, V. A., Campbell, C. J., and Grzybowski, B. A. Architecture and Evolution of Organic Chemistry. *Angewandte Chemie International Edition*, 44(44):7263–7269, 2005. ISSN 1521-3773. doi: 10.1002/anie.200502272. URL https://onlinelibrary.wiley.com/ doi/abs/10.1002/anie.200502272. \_eprint: https://onlinelibrary.wiley.com/doi/pdf/10.1002/anie.200502272.
- Hartle, H., Klein, B., McCabe, S., Daniels, A., St-Onge, G., Murphy, C., and Hébert-Dufresne, L. Network comparison and the within-ensemble graph distance. *Proceedings of the Royal Society A: Mathematical, Physical and Engineering Sciences*, 476 (2243):20190744, November 2020. ISSN 1364-5021, 1471-2946. doi: 10.1098/rspa.2019.0744. URL https://royalsocietypublishing.org/ doi/10.1098/rspa.2019.0744.
- Hogan, A., Blomqvist, E., Cochez, M., D'amato, C., Melo, G. D., Gutierrez, C., Kirrane, S., Gayo, J. E. L., Navigli, R., Neumaier, S., Ngomo, A.-C. N., Polleres, A., Rashid, S. M., Rula, A., Schmelzeisen, L., Sequeda, J., Staab, S., and Zimmermann, A. Knowledge Graphs. ACM Computing Surveys, 54(4):71:1–71:37, July 2021. ISSN 0360-0300. doi: 10.1145/3447772. URL https://dl.acm.org/doi/10.1145/3447772.
- Hu, W., Fey, M., Zitnik, M., Dong, Y., Ren, H., Liu, B., Catasta, M., and Leskovec, J. Open Graph Benchmark: Datasets for Machine Learning on Graphs. ArXiv, May 2020. URL https://www.semanticscholar. org/paper/Open-Graph-Benchmark% 3A-Datasets-for-Machine-Learning-Hu-Fey/ 597bd2e45427563cdf025e53a3239006aa364cfc.
- Khattab, O. and Zaharia, M. A. Colbert: Efficient and effective passage search via contextualized late interaction over bert. *Proceedings of the* 43rd International ACM SIGIR Conference on Research and Development in Information Retrieval, 2020. URL https://api.semanticscholar. org/CorpusID:216553223.
- Khattab, O., Singhvi, A., Maheshwari, P., Zhang, Z., Santhanam, K., Vardhamanan, S., Haq, S., Sharma, A., Joshi, T. T., Moazam, H., Miller, H., Zaharia, M., and Potts, C. Dspy: Compiling declarative language model calls into self-improving pipelines. *ArXiv*, abs/2310.03714, 2023. URL https://api.semanticscholar.org/CorpusID:263671701.

- 385 Liu, J. jxnl/instructor, May 2024. URL https://
  386 github.com/jxnl/instructor. original-date:
  387 2023-06-14T10:42:23Z.
- Lowe, D. M. *Extraction of chemical structures and reactions from the literature*. PhD thesis, University of Cambridge, 2012.
- Lowe, D. M. and Sayle, R. A. Leadmine : A grammar and dictionary driven approach to chemical entity recognition.
   2013. URL https://api.semanticscholar. org/CorpusID:1701882.
- M. Bran, A., Cox, S., Schilter, O., Baldassari, C., White,
  A. D., and Schwaller, P. Augmenting large language models with chemistry tools. *Nature Machine Intelligence*, 6
  (5):525–535, May 2024. ISSN 2522-5839. doi: 10.1038/
  s42256-024-00832-8. URL https://www.nature.
  com/articles/s42256-024-00832-8. Publisher: Nature Publishing Group.
- 404 Mavračić, J., Court, C. J., Isazawa, T., Elliott, S. R., and 405 Cole, J. M. ChemDataExtractor 2.0: Autopopulated On-406 tologies for Materials Science. Journal of Chemical 407 Information and Modeling, 61(9):4280-4289, Septem-408 ber 2021. ISSN 1549-9596. doi: 10.1021/acs.jcim. 409 1c00446. URL https://doi.org/10.1021/acs. 410 jcim. 1c00446. Publisher: American Chemical Soci-411 ety. 412
- Meyer, L., Stadler, C., Frey, J., Radtke, N., Junghanns,
  K., Meissner, R., Dziwis, G., Bulert, K., and Martin, M. Llm-assisted knowledge graph engineering: Experiments with chatgpt. *ArXiv*, abs/2307.06917,
  2023. URL https://api.semanticscholar. org/CorpusID:259847255.
- 420 Mihindukulasooriya, N., Tiwari, S. M., Enguix, C. F.,
  421 and Lata, K. Text2kgbench: A benchmark for
  422 ontology-driven knowledge graph generation from
  423 text. ArXiv, abs/2308.02357, 2023. URL https:
  424 //api.semanticscholar.org/CorpusID:
  425 260611736.

426

- Rajan, K., Zielesny, A., and Steinbeck, C. DECIMER
  1.0: deep learning for chemical image recognition using transformers. *Journal of Cheminformatics*, 13(1):
  61, August 2021. ISSN 1758-2946. doi: 10.1186/
  s13321-021-00538-8. URL https://doi.org/10.
  1186/s13321-021-00538-8.
- 433 Rajan, K., Brinkhaus, H. O., Agea, M. I., Zielesny, 434 A., and Steinbeck, C. DECIMER.ai: an open plat-435 form for automated optical chemical structure iden-436 tification, segmentation and recognition in scientific 437 publications. Nature Communications, 14(1):5045, 438 August 2023. ISSN 2041-1723. doi: 10.1038/ 439

s41467-023-40782-0. URL https://www.nature. com/articles/s41467-023-40782-0. Publisher: Nature Publishing Group.

- Rebedea, T., Dinu, R. L., Sreedhar, M. N., Parisien, C., and Cohen, J. Nemo guardrails: A toolkit for controllable and safe llm applications with programmable rails. In *Conference on Empirical Methods in Natural Language Processing*, 2023. URL https: //api.semanticscholar.org/CorpusID: 264146531.
- Shimada, Y., Hirata, Y., Ikeguchi, T., and Aihara, K. Graph distance for complex networks. *Scientific Reports*, 6(1):34944, October 2016. ISSN 2045-2322. doi: 10.1038/srep34944. URL https://www.nature. com/articles/srep34944.
- Shu, D., Chen, T., Jin, M., Zhang, Y., Zhang, C., Du, M., and Zhang, Y. Knowledge graph large language model (kg-llm) for link prediction. *ArXiv*, abs/2403.07311, 2024. URL https://api.semanticscholar. org/CorpusID:268363911.
- Swain, M. C. and Cole, J. M. ChemDataExtractor: A Toolkit for Automated Extraction of Chemical Information from the Scientific Literature. *Journal of Chemical Information and Modeling*, 56(10):1894–1904, October 2016. ISSN 1549-9596. doi: 10.1021/acs.jcim. 6b00207. URL https://doi.org/10.1021/acs. jcim.6b00207. Publisher: American Chemical Society.
- Thompson, R., Knyazev, B., Ghalebi, E., Kim, J., and Taylor, G. W. On Evaluation Metrics for Graph Generative Models, April 2022. URL http://arxiv.org/ abs/2201.09871. arXiv:2201.09871 [cs].
- Zheng, S., Rao, J., Song, Y., Zhang, J., Xiao, X., Fang, E. F., Yang, Y., and Niu, Z. Pharmkg: a dedicated knowledge graph benchmark for bomedical data mining. *Briefings in bioinformatics*, 2020. URL https://api.semanticscholar. org/CorpusID:229342739.

#### **A. Supplementary Information Files** 440

441 A typical practice in organic chemistry publishing is having 442 Supplementary Information files (SIs) where all information 443 regarding experimental procedures, analytical results, and 444 sometimes computational and theoretical predictions, are 445 reported. In these documents, which all share a general 446 underlying structure, reactions are described with references 447 to other substances in the same document, with a notation 448 shared between the SI and the main manuscript. Hence, a 449 numeration scheme exists for the substances in each paper 450 that can be followed to find the experimental procedure 451 for the preparation of any compound synthesized as part 452 of the research work. Despite of this homogeneity, large 453 differences are noticeable, as is evident from figures 4, 5 454 and 6. 455

456 As these examples show, representations and formats are 457 far from standardized. The SI displayed in Figure 4 shows 458 a common format: compound name and reference in bold, 459 accompanied by the molecular structure of the product sub-460 stance, and followed by the reaction procedure. Notice 461 however that a subsequent reaction is described directly in 462 the same paragraph, without *announcing* the next product. 463

Figure 5 shows an SI with a heavier use of visual elements, 464 where colored marbles are used to reference individual steps 465 in a short reaction sequence. The marbles are then used 466 throughout to refer to specific intermediates, with no refer-467 ence in text to the products' reference keys. Lastly, Figure 468 6 shows another example where the product is not directly 469 announced in the text, but rather a new reaction procedure 470 is presented after a graphical depiction of the reaction in 471 question, making it impossible for a text parser to grasp this 472 information. 473

### **B. SI Preprocessing**

SIs in chemistry research papers contain many sections, 477 however the one of interest for this work is the part on 478 Experimental Methods. For our purposes, it may make 479 sense to extract the most relevant parts of the document 480 and process only that, however no naming convention or 481 guidelines exist for this, making it difficult to identify and 482 isolate the specific sections. 483

484 To address this, we develop a simple rule-based method to 485 identify the relevant sections, partially inspired by ?. For 486 this, we rely on the observation that reaction descriptions 487 typically follow the pattern "reaction setup ightarrow workup ightarrow488 analytics", as the example below. As can be seen, the analyt-489 ics section has a higher ratio of certain special and numeric 490 characters relative to other parts of the text.

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To a solution of alkene 5 (266 mg, 0.92 mmol, 1.0 equiv.) in DCM (30 mL) was bubbled ozone



Stille substrate 145. Dess-Martin Periodinane (35.6 mg, 0.084 mmol) was added to a mixture of alcohol 143 (28 mg, 0.028 mmol) and NaHCO<sub>3</sub> (73 mg, 0.869 mmol) in CH<sub>2</sub>Cl<sub>2</sub> (2 mL) at 0 °C. After 1 h, additional Dess-Martin Periodinane was added (5.0 mg, 0.012 mmol) After 30 min, the reaction mixture was quenched with saturated aqueous NaHCO<sub>1</sub> (1 mL) and saturated aqueous sodium metabisulfite (1 mL). The resulting cloudy mixture was stirred vigorously for 5 min at 0 °C, then for 30 min at rt. The layers were separated and the aqueous layer was extracted with EtOAc (5 x 1 mL). The combined organic layers were washed with brine (1 x 1 mL), dried by passage over a plug of silica gel (EtOAc eluent), and evaporated under reduced pressure. The residue was purified by flash chromatography (19:1 hexanes:EtOAc containing 2% Et<sub>2</sub>N; then 9:1 hexanes:EtOAc containing 2% Et<sub>2</sub>N) to afford aldehyde SI-31 (24.0 mg, 86%) as a colorless foam which was used directly in the subsequent transformation.

NaHMDS (288 uL, 0.288 mmol, 1 M in THF) was added dropwise over 1 min to a mixture of phosphonium salt 1449 (251.5 mg, 0.481 mmol) in DME (4.3 mL) at -78 °C. After stirring 1 h 10 min, aldehyde SI-31 (24.0 mg, 0.0.024 mmol, dried under vacuum over  $CaSO_4$ ) in DME (1 mL) was added. The mixture was maintained at -78 °C for 25 min, then placed in a 0  $^{\circ}\text{C}$  bath for 15 min. The reaction mixture was diluted with H2O (2 mL), brine (2 mL), and EtOAc (2 mL). The mixture was warmed to rt and the layers were separated. The aqueous layer was further extracted with EtOAc (4 x 2 mL). The combined organic layers were dried by passage over a plug of silica gel (EtOAc eluent) and evaporated under reduced pressure. The residue first purified by passage over a second plug of silica gel (4:1 hexanes:EtOAc containing 2%

Figure 4. Example of an SI. Taken from https://pubs.acs. org/doi/10.1021/ja074300t. This example shows

(40mixture was purged with air at -78 °C followed by addition of PPh3 (250 mg, 0.95 mmol, 1.0 equiv.). The mixture was warmed up to room temperature slowly, and stirred at the same temperature for 12 h. After removal of the solvent. the residue was purified by a flash column chromatography on silica gel (hexane/EtOAc = 5:1to 3 : 1) to give compound 6 as a colorless oil (173 mg, 65inconsequential 1.05: 1 mixture. Rf = 0.25 (hexane/EtOAc = 8:1, PMA);  $[\alpha]_{21}$ D = - 4.44 (c 1.31, CHCl3); 1H NMR (400 MHz, CDCl3)  $\delta$  9.77 – 9.70 (m, 1.69H, overlap), 2.63 – 2.48 (m, 2.21H, overlap), 2.42 – 2.18 (m, 9.27H, overlap), 2.18 – 2.06 (m, 3.58H, overlap), 2.00 – 1.82 (m, 5.93H, overlap), 1.82 - 1.72 (m, 4.72H, overlap), 1.71 – 1.60 (m, 3.95H, overlap), 1.58 – 1.49

To leverage this, we split the complete document into sentences, and then calculate the ratio of special characters to normal letters for each. Plotting the values of these ratio with the line index in the x-axis, patterns like those in Figure 7 are apparent. An alogrithm is also applied for smoothing and performing selection by selecting the longest region with a prominent signal as the "relevant" SI. We find that this strategy generally leads to an accurate selection of the relevant parts.



O To a solution of carboxylic acid (20 mmol, 1eq.) in CH<sub>2</sub>Cl<sub>2</sub> (60 mL) was added carbonyldiimidazole (23 mmol, 1.15 eq.) portion wise and the solution stirred at room temperature for 1 h. After this time, N₂ was bubbled through the solution for 30 min. Then, N, O-dimethylhydroxylamine hydrochloride (26 mmol, 1.3 eq.) was added and the reaction mixture was stirred at room temperature for 24 h. Subsequently, the reaction mixture was quenched with saturated aqueous NH<sub>4</sub>Cl and extracted with CH<sub>2</sub>Cl<sub>2</sub> (3 × 70 mL). The combined organic layer was dried over MgSO<sub>4</sub> and concentrated in vacuo to afford the product.

The corresponding amide (1 eq.) was placed in a round bottom flask under N<sub>2</sub>. The flask was cooled to -78 °C and dry THF (5 mL/mmol) was added. 2-Methyl-1-propenylmagnesium bromide solution (1.2 eq.) was added dropwise at -78 °C and the reaction mixture was stirred at -78 °C for 15 min. The reaction mixture was then stirred at room temperature for 12 h. Subsequently, the reaction mixture was quenched with saturated aqueous NH<sub>4</sub>Cl and extracted with CH<sub>2</sub>Cl<sub>2</sub> (3 × 30 mL). The organic layers were collected, dried over MgSOa and concentrated in vacuo. The desired compound

Figure 5. Example of an SI. Taken from https://pubs.acs. org/doi/10.1021/jacs.1c01356. This example shows

## C. VLM

The following prompt was used as a template to pass the images to GPT-40 for the vision-based parsing method exposed in Figure 1.

These are some pages from the SI of an organic chemistry paper. Describe all the reactions shown there, if any. Separate each reaction with {SEPARATOR}, describe products and reactants for each reaction. Ignore all characterization data. Consider work-up and purification as part of the same reaction. Use the following format to represent the products and main reactants: {SUBSTANCE\_FORMAT}. Do not rewrite the reaction procedures, just describe the substances involved. <sup>1</sup>**H NMR** (600 MHz, CDCl<sub>3</sub>) δ 6.76 (d, *J* = 10.0 Hz, 2H), 6.22 (d, *J* = 10.0 Hz, 2H), 3.74 (s, 2H), 1.53 (s, 9H), 1.49 (s, 9H).;

<sup>13</sup>C NMR (151 MHz, CDCl<sub>3</sub>) δ 185.3, 152.0, 152.0, 149.3, 149.1, 128.3, 84.6, 82.3, 63.9, 53.1, 28.12, 28.10 ppm.;

HRMS (m/z) calc'd for C<sub>18</sub>H<sub>26</sub>N<sub>3</sub>O<sub>5</sub>+ [M+H]+ 364.1867, found 364.1873



Experimental: The cyclized product 5 (11 g, 30.3 mmol) was placed in a flame-dried 1 L round bottom flask fixed with a stir bar. The flask was placed under vacuum and backfilled with argon. Dry THF (525 ml) and dry HMPA (81 ml) were added into the flask via syringe. Start stirring at room temperature and then cool the reaction down to -40 °C after the mixture became homogeneous. Add methylmagnesium bromide (50.5 ml of 3 M solution in EtzO, 5 eq) dropwise into the mixture. Wait for 30 mins after the completion of the addition of methylmagnesium bromide then warm the reaction mixture up to 0 °C. Wait for another 30 mins then add sat. NH<sub>4</sub>Cl solution to quench the reaction. Extract the aqueous layer with ethyl acetate twice then combine all organic layers which were then washed with 10% LiCl solution for three times.

The organic layer was first dried over  $MgSO_4$  which was then removed by filtration. The solvent was removed under reduced pressure to give the crude reaction mixture as a yellow solid. Add ethyl acetate (35 ml) and hexane (175 ml) into the resulting solid and heat the mixture with a heat oun to dissolve the solid as much as possible. Let the mixture

Figure 6. Example of an SI. Taken from https://pubs.acs. org/doi/10.1021/jacs.3c01991. This example shows



*Figure 7.* SIs were processed like this. Based on the frequency of special characters etc. Based on the observation that, most commonly, text-summaries of analytical data are given after the end of each reaction, giving a distinctive signal to each line in the document, producing more or less a spectrum that can then be analysed and processed.

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