Foundation Models of Scientific Knowledge for Chemistry: Opportunities, Challenges and Lessons Learned

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Abstract

Foundation models pre-trained on large corpora demonstrate significant gains across many natural language processing tasks and domains e.g., law, healthcare, education, etc. However, only limited efforts have investigated the opportunities and limitations of applying these powerful models to science and security applications. In this work, we develop foundation models of scientific knowledge for chemistry to augment scientists with the advanced ability to perceive and reason at scale previously unimagined. Specifically, we build large-scale (1.47B parameter) general-purpose models for chemistry that can be effectively used to perform a wide range of in-domain and out-of-domain tasks. Evaluating these models in a zero-shot setting, we analyze the effect of model and data scaling, knowledge depth, and temporality on model performance in context of model training efficiency.

Our novel findings demonstrate that (1) model size significantly contributes to the task performance when evaluated in a zero-shot setting; (2) data quality (aka diversity) affects model performance more than data quantity; (3) similarly, unlike previous work (Luu et al., 2021) temporal order of the documents in the corpus boosts model performance only for specific tasks, e.g., SciQ; and (4) models pre-trained from scratch perform better on in-domain tasks than those tuned from general-purpose models like Open AI's GPT-2.

1 Introduction

The emergence of foundation models (Bommasani et al., 2021) such as large-scale autoencoding models (*e.g.*, BERT (Devlin et al., 2018), RoBERTa (Liu et al., 2019)) and autoregressive language models (*e.g.*, GPT-2 (Radford et al., 2019), GPT-3 (Brown et al., 2020), Megatron-Turing (Smith et al., 2022) and Gopher (Rae et al., 2021)) as well as multimodal vision and language models, such as FLAVA (Singh et al., 2021) and Perceiver (Jaegle et al., 2021), established a paradigm shift in Artificial Intelligence (AI). These foundation models, also called *neural platforms*, are built using self-supervised pretraining at scale. They are then able to be easily adapted to a wide range of downstream tasks via transfer learning (Bommasani et al., 2021) and finetuning (Lee et al., 2019).

The wide community adoption of foundation models can be explained by their key properties, two of which are emergent behavior and homogenization - which also make foundation models appealing for adaption across science and security domains. Emergence, or emergent behavior, reflect new behaviors that a model introduces or is capable of that it was not explicitly trained to perform. Homogenization is the consolidation of methods for building machine learning systems across a wide range of tasks. Another key advantage of scaling language models is that they perform competitively on language tasks using in-context learning without fine-tuning or gradient updates. Thus, in-context learning allows foundation models to be effectively used across new downstream tasks with only simple instructions and a few optional examples.

In this work we focus on a science domain (chemistry) and demonstrate the value and limitations of large-scale language models evaluated across a wide range of in-domain (science-focused) and out-of-domain tasks. Unlike the majority of work on foundation models that focuses on pretraining these models on book corpora, web pages, Wikipedia and mixed sources, e.g., the Pile (Gao et al., 2020), we pretrain our models on scientific literature. Using scientific literature presents unique opportunities and challenges. Opportunities include the scale and diversity of scientific literature, the explicit structure, and explicit alignment across different modalities in the papers, e.g., table and figure references. Challenges include limited benchmarks that can be used to perform model

evaluation, model prompting and interactions.

There are three major contributions of this work: (1) we collect and release a 0.67TB dataset covering research publication data across 10+ sources for chemistry; (2) we release 28 auto-regressive foundation models for chemistry that have been pretrained from scratch; and (3) we present a rigorous evaluation of model performance on 15+ indomain and out-of-domain tasks that investigates the effects of model and data scaling, knowledge depth (aka diversity), and temporal order on performance as described in research questions below.

(**RQ1**) Science-Focused Benchmarks What are the strengths and weaknesses of foundation models pretrained on scientific literature when evaluated on out-of-domain vs. in-domain tasks?

(**RQ2**) Scaling Effect How does model scale affect the downstream performance? Do neural scaling laws presented in (Kaplan et al., 2020) hold for the foundation models for science?

(**RQ3**) **Diversity Effect** How does the depth of scientific knowledge, *e.g.*, from paper abstracts vs. full text, affect downstream performance?

(**RQ4**) **Temporal Effect** How does the recency of scientific knowledge, *e.g.*, when manipulating the temporal order of the documents processed by the model, affect downstream performance?

2 Related Work

In this section we summarize previous efforts in two categories: *mixed-domain continual pretraining* that continues pretraining of a base model on domain data and *in-domain pretraining from scratch* that pretrains a from scratch on domain data. We present a model summary in Table 1.

Mixed-Domain Continual Pretraining Many efforts have focused on continual pretraining of a BERT (Devlin et al., 2018) base model. Several models have been developed for the biomedical domain and the most frequently used corpora for domain-specific continual preraining are PubMed abstracts and PubMed Central full-text articles (PMC) (Lee et al., 2020; Peng et al., 2019; Phan et al., 2021). In the Chemistry domain, Guo et al. (2021) performed continual pretraining of a base BERT model on 200K chemistry journal articles for product extraction (ChemBERT) and reaction role labeling (ChemRxnBERT).

In-Domain Pretraining from Scratch Previous work has shown that pretraining models from scratch on domain-specific data has a significant benefit over continual pretraining of generaldomain language models (Gu et al., 2021). This is mainly due to the availability of in-domain data for both generating the vocabulary and pretraining. SciBERT (Beltagy et al., 2019) is pretrained according to this procedure using the vocabulary generated from computer science and biomedical domains. PubMedBERT (Gu et al., 2021) is another example of pretraining the base BERT model from scratch using PubMed. Unlike any previous work, we use both continual and from scratch pretraining to build the largest foundation model for Chemistry (1.47B) on the largest (0.67TB) and the most diverse corpus (10+ sources) collected to date.

3 Model Pretraining

Unlike the majority of related models that rely on a base BERT (or variant) model, we adapt the Open-AI's GPT-2 transformer decoder architecture (Radford et al., 2019) to train autoregressive language models for Chemistry. To understand the impact of model size (RQ2), we experiment with four different Transformer sizes: small (S), medium (M), large (L), and extra-large (XL). These models differ in the number of decoder layers, hidden size of the model, and the number of attention heads in transformer blocks as shown in Table 2.

Our experiments leverage the GPT-NeoX Python library (Andonian et al., 2021) developed with Megatron (Shoeybi et al., 2019) and Deep-Speed (Rasley et al., 2020). We optimize the autoregressive log-likelihood (*i.e.*, cross-entropy loss) averaged over a 2048-token context. We set the micro batch size per GPU as 4, and the learning rate to 2×10^{-4} , and rely on the cosine decay. We use an Adam optimizer with $\beta_1 = 0.9$, $\beta_2 = 0.99$, and $\sigma = 10^{-8}$ and clip the gradient norm at 1.0. In addition, ZeRO optimizer (Rajbhandari et al., 2019) was used to reduce memory footprint by distributing optimizer states across several processes.

To reduce memory and increase training throughput, we use mixed-precision training (Rasley et al., 2020) and the parallel attention and feed-forward implementations available in GPT-NeoX (Black et al., 2022). We also use the Rotary positional embeddings (Su et al., 2021) instead of the learned positional embeddings used in the GPT-2 model (Radford et al., 2019) because they offer performance

Mode	el	Data Source	Pretraining	Corpus	#Params (B)	
Lee et al. 2020	BioBERT	Wiki + Books	continual pretraining	PubMed	0.11	
Alsentzer et al. 2019	ClinicalBERT	Wiki + Books	continual pretraining	$MIMIC^1$	0.11	
Peng et al. 2019	BlueBERT	Wiki + Books	continual pretraining	PubMed + MIMIC	0.11	
Liu et al. 2021	MATH-BERT	Arxiv	continual pretraining	Arxiv	0.11	
Guo et al. 2021	Chem(Rxn)BERT	Wiki + Books	continual pretraining	Chemistry Journals	0.11	
Phan et al. 2021	SciFive	C4	continual pretraining	PubMed	0.22	
Filall et al. 2021	SCIFIVE	C4	continuar pretraining	rubivieu	0.77	
Naseem et al. 2021	BioALBERT	Wiki + Books	continual pretraining	PMC + MIMIC-II	0.02	
Lewis et al. 2020	BioRoBERTa	Wiki + Books	continual pretraining	PMC + MIMIC-III	0.30	
Yuan et al. 2021	KeBioLM	PubMed	continual pretraining	PubMed + UMLS ²	0.34	
Shin et al. 2020	DiaManatara	PubMed	from scratch	PubMed	0.80	
Shih et al. 2020	BioMegatron		continual pretraining	Publyled	1.20	
Kanakarajan et al. 2021	BioELECTRA	PubMed	from scratch	PubMed	0.11	
Miolo et al. 2021	ELECTRAMed	PubMed	from scratch	PubMed	0.11	
Beltagy et al. 2019	SciBERT	PMC + CS	from scratch	PMC + CS	0.11	
Liu et al. 2021	OAG-BERT	OAG	from scratch	OAG	0.11	
Gu et al. 2021	PubMedBERT	PubMed	from scratch	PubMed	0.34	
Our U	Varte (autorearaasius) +	10+ sources	from scratch	10+ sources	1.47	
Our w	ork (autoregressive) †	(Chemistry)	continual pretraining	(Chemistry)	1.47	

Table 1: Foundation models for science focus on the biomedical, math, computer science and chemistry domains. We use † to indicate models trained for chemistry.

Table 2: Our model configurations: d_L is the number of decoder layers, d_{dim} is the hidden size of the model, d_{heads} is the number of attention heads. We compare model configurations between GPT-NeoX and OpenAI's GPT-2. GPT-NeoX architecture is originally from GPT-3 (Brown et al., 2020)

Size	Model	d_L	d_{dim}	d_{heads}	#Params (B)	
S	GPT-NeoX	12	768	12	0.18	
3	GPT-2	GPT-2 12 768 12		12	0.18	
М	GPT-NeoX	24	1024	16	0.40	
IVI	GPT-2	24	1024	16	0.40	
L	GPT-NeoX	24	1536	16	0.80	
L	GPT-2	36	1280	20	0.80	
XL	GPT-NeoX	24	2048	16	1.47	
ΛL	GPT-2	48	1600	25	1.47	

advantages in tasks with longer texts by capturing relative position dependency in self-attention.

Our models are pretrained across multiple workers with data parallelism. As the largest model in our experiments fit on a single GPU, we didn't use the model (tensor) or pipeline parallelism. Models are pretrained from scratch for a total of 320K steps. The original GPT-2 models are fine-tuned for 150K steps. We perform experiments in a single DGX-A100 machine with 8 80Gb GPUs.

4 Data Collection and Processing

We collected a large corpus of 53.45 million chemistry-focused scientific articles and abstracts, resulting in 670GB of text data. As shown in Table 3, our corpus was collected from 10 different data sources: Arxiv, Aminer (AMiner), CORD-19 (Wang et al., 2020b), CORE (Pontika et al., 2016), Microsoft Academic Graph (MAG) (Wang et al., 2020a), OSTI, PubMed (Gao et al., 2020) (abstracts and fulltexts), and the Web of Science (WoS). See Appendix A for full data descriptions.

Table 3: Dataset statistics: combined datasets are after the de-duplication process. We split datasets to those that include abstracts $\langle A \rangle$ vs. full texts $\langle FT \rangle$.

Source	#Articles (M)	#Tokens (B)	Size (Gb)
MAG $\langle A \rangle$	34.26	7.43	46
Aminer $\langle A \rangle$	18.50	5.80	35
S2ORC $\langle A \rangle$	10.44	2.05	32
WoS $\langle A \rangle$	7.90	3.31	18
CORD-19 $\langle A \rangle$	< 0.01	< 0.01	0.2
OSTI $\langle A \rangle$	0.05	< 0.01	0.1
Arxiv $\langle A \rangle$	0.38	0.04	0.4
PubMed $\langle A \rangle$	0.28	0.08	0.5
PubMed $\langle FT \rangle$	0.70	7.34	32
$CORE \langle FT \rangle$	7.27	215.50	743
Combined $\langle A \rangle$	46.94	16.18	67
Combined $\langle FT \rangle$	6.52	184.42	603
Combined (A+FT)	53.45	200.61	670

Because the data sources we relied on comprise research publications from many science domains, we sampled articles using a list of domain-specific keywords for chemistry to create the dataset summarized in Table 3. These keywords were extracted by using a Correlation Explanation (Gallagher et al., 2017) topic model followed by manual filtering by subject matter experts. This resulted in a list of more than 1K chemistry-related entities, ranging from compound names like *ethyl acetate*, *methyl methacrylate*, *sulfoxide*, etc. to experiment and procedures like *tunneling microscopy*, *neutralization*, *enzymatic hydrolysis*, etc.



Distribution within Sources with <1% (upper right, black box)

Figure 1: Summary of data source representation within the *Combined* A+F data sample. Coloring illustrates whether a data source contains *peer reviewed* (Blue), *mixed* (Purple), or *not peer reviewed* (Red) articles.

Data Cleaning Recent research has shown that duplicates in training data can significantly impact the downstream task performance of LLMs (Lee et al., 2021; Carlini et al., 2022). To this end, we performed deduplication of our corpus based on overlap of titles within and across data sources. We processed titles to strip punctuation and casefold and considered two articles A_1 and A_2 to be duplicates if they had the same processed title. With this technique, we were able to remove significant amounts of duplicate scientific articles both within and across sources. The deduplication process reduced our corpus from 875GB to 670GB (67.8M to 53.5M publications), removing 14.3M duplicates.

Tokenization As used in GPT-2 model, we use a Byte Pair Encoding (BPE) tokenizer. We train BPE tokenizers for each data sample with a vocabulary size of 64K as preliminary experiments varying vocabulary sizes from 64K to 256K for smaller scale model pretraining did not show significant differences in performance. We compare the GPT-2 vocabulary generated from the WebText and the in-domain vocabularies generated from our corpora and find that the in-domain vocabulary breaks chemical entities into fewer tokens. For example, *dimethylnitroxide* was tokenized into #dimethyl, #nitr, #oxide using the in-domain vocabulary and #dim, #ethyl, #nit, #rox, #ide using the GPT-2 vocabulary.

5 Analysis and Results

This section presents the analysis of 28 pretrained models evaluated on 15+ in-domain and out-ofdomain downstream tasks (RQ1, Section 5.1). We investigate the effects of model and data scaling (RQ2, Section 5.2), knowledge diversity (RQ3, Section 5.3), and temporal order (RQ4, Section 5.4) on the downstream performance. We also compare the results from continual vs. from scratch pretraining (Section 5.5) and present the analysis of large-scale training efficiency (Section 5.6).

Baseline Models As we use a similar model architecture, we identify Open AI's GPT-2 (Radford et al., 2019) as a baseline comparison model. We compare our performance with four variants of the original GPT-2 models, corresponding to small (S), medium (M), large (L), and extra-large (XL) sized transformer architectures shown in Table 2. We note that GPT-2 models were pretrained on Web-Text – 8 million web documents (40Gb). Thus, we also include a base GPT-2 model (medium) that has been updated with continual pretraining using our *Combined* $\langle A+FT \rangle$ dataset.

Our Models We pretrained models with individual datasets (AMiner, CORE, MAG, PubMed, S2ORC, WOS) and combined abstracts and fulltexts. Our goal is to systematically study data biases in the model performance when pretraining models with individual datasets. For example, PubMed publications cover mostly bio-medicinal terms (Gu et al., 2021), while the majority of S2ORC publications are from medicine, biology, physics, and mathematics (Lo et al., 2020). We only use 4 GPUs for the models pretrained with individual datasets and 8 GPUs for the rest. This is to control the number of tokens seen during model pretraining (320,000 steps * 4 GPUs * 4 micro batch size * 2,048 context size = 10B tokens) relative to the maximum number of tokens available in the respective datasets (as reported in Table 3). We also trained one XL (4x) model with 4x larger batch size than what used in XL model to evaluate the impact of the number of training tokens.

5.1 Zero-shot Performance

We evaluate our models using several benchmarks to assess the effectiveness in both in-domain and out-of-domain tasks. The benchmarks we include are described in Appendix B. We use the Imevaluation-harness Python repository (Gao et al.,

Table 4: Downstream Zero-shot In-Domain Task Performance. We use \ddagger to indicate the baseline model tuned from the base GPT-2 model. Pile performance is reported using perplexity, with all other tasks reported using accuracy. We highlight the top-4 performance per task in bold, with top performance indicated with an underline. XL (4x) model is trained with 4x larger batch size that used in other models.

Model	Size	HT-HC	HT-CC	ARC-E	ARC-C	SciQ	OpenBookQA	Pile
Baseline	S	0.22	0.25	0.44	0.19	0.75	0.16	96.50
	М	0.18	0.27	0.49	0.22	0.77	0.19	61.26
	L	0.18	0.28	0.53	0.22	0.80	0.19	48.86
	XL	0.18	0.26	0.58	0.25	0.83	0.22	42.29
	M‡	0.19	0.31	0.35	0.19	0.61	0.13	87.57
	S	0.18	0.27	0.43	0.21	0.70	0.17	38.40
4 M	М	0.18	0.34	0.45	0.20	0.74	0.16	30.55
AMiner	L	0.23	0.34	0.49	0.23	0.78	0.18	24.18
	XL	0.23	0.34	0.50	0.23	0.77	0.17	25.52
	S	0.19	0.28	0.36	0.19	0.69	0.15	78.24
CODE	М	0.22	0.34	0.40	0.20	0.71	0.15	59.19
CORE	L	0.17	0.30	0.41	0.19	0.75	0.14	52.95
	XL	0.20	0.28	0.47	0.21	0.78	0.15	39.46
	S	0.24	0.28	0.41	0.20	0.66	0.17	38.03
MAG	М	0.18	0.27	0.45	0.21	0.68	0.17	30.88
MAG	L	0.19	0.36	0.51	0.24	0.80	0.18	24.78
	XL	0.20	0.36	0.50	0.22	0.80	0.20	26.09
	S	0.26	0.30	0.41	0.20	0.60	0.16	56.03
PubMed-F	М	0.19	0.27	0.43	0.21	0.68	0.18	45.69
Publyled-F	L	0.18	0.28	0.43	0.22	0.74	0.17	37.22
	XL	0.18	0.27	0.48	0.21	0.77	0.16	35.14
	S	0.26	0.33	0.31	0.21	0.31	0.17	59.20
CODC	М	0.27	0.22	0.33	0.18	0.31	0.16	45.60
S2ORC	L	0.28	0.23	0.32	0.21	0.31	0.17	42.14
	XL	0.24	0.31	0.33	0.19	0.30	0.18	42.35
WoS	S	0.22	0.31	0.33	0.22	0.37	0.17	54.41
	М	0.25	0.32	0.32	0.20	0.34	0.16	48.31
	L	0.27	0.30	0.32	0.21	0.37	0.17	46.44
	XL	0.23	0.34	0.34	0.21	0.39	0.16	45.86
Combined-A	XL	0.17	0.28	0.54	0.23	0.83	0.18	22.77
Combined-F	XL	0.20	0.30	0.48	0.21	0.79	0.15	40.18
Combined-A+F	XL	0.18	0.30	0.48	0.22	0.79	0.17	31.03
Combined-A+F	XL (4x)	0.18	0.25	0.55	0.24	<u>0.84</u>	0.17	23.01

2021) for the benchmark implementation.

In-domain Evaluation We consider five existing chemistry benchmarks, specifically HendrycksTest (Hendrycks et al., 2020) for high school (HT-HC) and college (HT-CC) levels, and sciencefocused – ARC (Clark et al., 2018), SciQ (Welbl et al., 2017), OpenBookQA (Mihaylov et al., 2018), Pile-PubMed-Abstracts (Gao et al., 2020)). As shown in Table 4, one or more of our models outperform baseline GPT-2 models for the two chemistry tasks, general science QA (SciQ) and the sciencefocused language modelling. Of the remaining tasks, our models perform within 1-4% of GPT-2 baselines.

Out-of-domain Evaluation We evaluate outof-domain performance using 9 commonly used LLM benchmarks: BoolQ (Clark et al., 2019), CB (De Marneffe et al., 2019), WIC (Pilehvar and Camacho-Collados, 2018), WSC (Levesque et al., 2012), MathQA (Amini et al., 2019), PIQA (Bisk et al., 2020), PubMedQA (Jin et al., 2019), Lambada (Paperno et al., 2016) and WikiText (Merity et al., 2016). As shown in Table 5, our models outperform baseline GPT-2 models for CB, WIC and WSC and match the best accuracy for BoolQ but the GPT-2 baselines outperform on the remaining tasks, particularly Lambada and Wikitext – the two general language modeling tasks.

5.2 Scaling Effect

Previous work (Kaplan et al., 2020) has shown that upstream cross entropy loss scales as a power-law with model size, dataset size, and the amount of compute. In this section, we revisit these claims on scaling Transformer architectures.

Analyzing upstream cross entropy loss During pretraining, we group each dataset into training/validation/test (949/50/1) splits. We report the

Table 5: Downstream Out-of-domain Task Performance. We use \ddagger to indicate the baseline model tuned from the base GPT-2 model. Performance on Lambada and Wikitext is reported using perplexity, all other tasks report accuracy. Top-4 performance highlighted in bold, with best performance indicated with underlines. XL (4x) model is trained with 4x larger batch size that used in other models.

Model	Size	BoolQ	CB	WIC	WSC	MathQA	PIQA	PubMedQA	Lambada	Wikitext
	S	0.49	0.41	0.49	0.43	0.21	0.63	0.44	40.06	37.37
	М	0.59	0.43	0.50	0.40	0.23	0.68	0.53	18.25	26.75
Baseline	L	0.60	0.45	0.50	0.46	0.23	0.70	0.54	12.97	22.61
	XL	0.61	0.39	0.50	0.50	0.24	<u>0.71</u>	0.59	<u>10.63</u>	<u>20.38</u>
	M‡	0.62	0.34	0.50	0.36	0.20	0.55	0.55	2834.51	126.55
	S	0.41	0.39	0.50	0.44	0.22	0.56	0.46	2825.84	158.85
AMiner	Μ	0.40	0.39	<u>0.51</u>	0.41	0.21	0.57	0.43	1802.35	116.93
Alviniei	L	0.61	<u>0.48</u>	0.50	0.47	0.22	0.58	0.36	661.81	87.23
	XL	0.50	0.39	0.50	0.37	0.21	0.58	0.43	786.22	91.28
	S	0.62	0.41	0.50	0.37	0.20	0.55	0.55	671.43	100.53
CORE	М	0.62	0.41	0.50	0.37	0.21	0.56	0.55	273.06	77.96
CORE	L	0.61	0.41	0.50	0.37	0.21	0.57	0.51	173.15	69.62
	XL	0.61	0.38	0.50	0.37	0.22	0.58	0.45	79.95	50.47
	S	0.41	0.23	0.50	0.40	0.21	0.56	0.43	1142.83	118.40
MAG	Μ	0.38	0.07	0.50	0.37	0.21	0.57	0.41	628.72	91.36
MAO	L	0.51	0.14	0.50	0.35	0.22	0.59	0.39	282.39	67.74
	XL	0.40	0.11	<u>0.51</u>	0.62	0.22	0.59	0.34	364.54	70.71
	S	0.58	0.41	0.50	0.45	0.21	0.57	0.54	2670.39	148.88
PubMed-F	Μ	0.61	0.39	0.50	0.38	0.20	0.58	0.49	1742.00	119.74
r ubivieu-r	L	0.57	0.41	0.50	0.38	0.21	0.59	0.42	843.83	95.75
	XL	0.60	0.41	0.50	0.39	0.22	0.59	0.49	679.80	90.38
	S	0.38	0.41	0.50	0.63	0.20	0.57	0.34	122739.30	403.48
S2ORC	Μ	0.38	0.43	0.50	<u>0.63</u>	0.22	0.56	0.34	80151.10	330.56
320KC	L	0.38	0.46	0.50	<u>0.63</u>	0.21	0.56	0.34	89136.68	327.53
	XL	0.38	0.50	0.50	0.63	0.20	0.56	0.33	107065.48	351.81
	S	0.38	0.39	0.50	0.63	0.21	0.55	0.34	140552.69	556.00
WoS	М	0.38	0.45	0.50	<u>0.63</u>	0.19	0.54	0.34	182967.37	498.36
	L	0.41	0.36	0.47	0.54	0.21	0.56	0.42	148609.73	480.91
	XL	0.57	0.34	0.50	0.37	0.20	0.55	0.56	192970.64	509.06
Combined-A	XL	0.56	0.16	0.50	0.37	0.21	0.60	0.50	250.88	61.07
Combined-F	XL	<u>0.62</u>	0.38	0.50	0.37	0.22	0.57	0.55	72.50	48.96
Combined-A+F	XL	0.61	0.41	0.50	0.39	0.23	0.59	0.48	71.43	48.65
Combined-A+F	XL (4x)	0.61	0.41	0.50	0.37	<u>0.24</u>	0.60	0.56	30.40	33.05

model performance on validation data using cross entropy loss in nats. This measure will be averaged over the 2048-token context. We find that the cross entropy loss decreases as we increase the model size (as shown in Figure 2). Larger models reach a given loss value in a higher rate than the smaller models. This observation illustrates the relationship between model performance (as measured by the upstream cross entropy loss) and model size, confirming (Kaplan et al., 2020).

Analyzing downstream task performance Can we speculate downstream task performance of a model from the pretraining performance? First, we find that the models perform considerably well on *Pile* in comparison to the *Lambada or WikiText*. There is a 48% performance advantage in this task over the best performing baseline GPT-2 model. This may be due to the models capturing scientific language better than general language. It is im-



Figure 2: Distribution of validation loss by model size: performance improves as the model size increases.

portant to note that we exclude PubMed Abstracts in the individual data collection to avoid potential contamination between the training and *Pile* testing data. As shown in Table 4, larger models perform well on these language modeling tasks.

Second, we noticed that the XL (4x) model trained for more tokens performs significantly better than the similar sized XL model. Specifically, XL (4x) model was trained with 128 total batch

size compared to the 32 total batch size used in XL model. XL (4x) model achieves the lowest *Lambada and WikiText* perplexity values across all our models trained from scratch (as shown in Table 5). The same model also achieves the best SciQ performance with 0.84 accuracy and comparable in other tasks performance with the XL model. This experiment highlights the importance of training models with larger batch size. We note that the baseline models (Radford et al., 2019) were trained with 4x larger batch size (total batch size 512) than what used in XL (4x) model. We believe that the XL (4x) model can reach the similar perplexity values when trained for this data scale.

Third, we find that zero-shot task performance in SciQ, HT-CC and ARC-E increases as we increase the model size (see Table 5). However, there is no clear relationship between the task performance and the model sizes in the rest of benchmark datasets. We suggest that pretraining performance may not be the ideal indicator to speculate the overall downstream task performance, especially in the zero-shot setting. However, model size significantly contributes to the task performance.

5.3 Diversity Effect

While abstracts often provide a summary of scientific publications, the full text contains more details. In this section, we analyze the performance of models trained on paper abstracts versus full texts.

First, the XL models trained with the combined abstract dataset achieve the lowest perplexity score (22.77) on the Pile – a 45% performance advantage over the full text version. There are might be several factors that contribute to this, but one may be the focused language in abstracts.

Second, the model trained with the combined abstracts achieves the second best accuracy (0.83 in comparison to 0.79 for the full text model) in SciQ. Some of the models pretrained on individual abstract data achieve comparable performance in SciQ, *e.g.*, MAG and AMiner models achieve 0.8 and 0.78 accuracy, respectively. We believe the diversity of scientific knowledge provided from the abstract data is useful since SciQ questions span biology, chemistry, earth science, and physics.

Third, we compare model performance trained with abstracts vs. full texts in the HT task and see that the best accuracy is achieved using the MAG and S2ORC datasets rather than the combined abstracts. This suggests the importance of contextual knowledge provided by different data sources.

Finally, combined full text model performs better than the model trained with the abstracts in all outof-domain tasks except PIQA. This performance difference may be due to the more expressive and diverse language presented in the full texts than in the abstracts. Thus, expanding full text coverage may improve out-of-domain task generalization.

5.4 Temporal Effect

Scientific knowledge evolves over time reflecting new research ideas, innovations, and findings. In this section, we test how continual pretraining on temporal-aligned scientific publications impacts downstream performance. For this experiment, we maintain two variants of the MAG dataset with random-ordered and temporal-ordered articles, splitting each into ten equal subsets. We continue pretraining a base medium (M) sized model iteratively with the subsets in the order they appeared in the respective data variant. For example, in the temporally-aligned experiments, we first pretrain a model with 3.4M (10%) articles from before 1978, and then use it as the base model to continue pretraining with another 3.4M (10%) articles from between 1978 and 1989. We train the initial model for 150K steps and each subsequent model for 10K steps with additional data. Figure 3 shows the performance of model checkpoints across in-domain and out-of-domain tasks.

There are two key findings. First, SciQ and ARC-E zero-shot task performances improve over time with the models trained with temporally-ordered scientific texts (as shown in Figure 3b). For example, SciQ accuracy improves from 0.64 to 0.73 from the base model checkpoint to the final model checkpoint. Similarly, ARC-E accuracy improves from 0.43 to 0.45. This is due to the temporal order of the knowledge acquired by the model. When the model was pretrained with random-ordered data subsets, we observe only a slight (< 1%) performance increase (as shown in Figure 3a).

There are mixed patterns in performance across out-of-domain tasks. For example, a slight performance increase in the PIQA, CB, PubMedQA, and WIC over time with the models trained with temporally-ordered scientific texts. On the other hand, there is a performance drop in the BoolQ and WSC over time. This may be due to the *catastrophic forgetting* prevalent in continual learning (Ramasesh et al., 2021). Future work will in-



Figure 3: The effect of temporal order of publications during pretraining. We align publications in the MAG corpus by year and split them into ten equal subsets. We repeat the process in a randomly-ordered corpus for comparison, recording model checkpoints after performing *continual pretraining* on each data subset.

vestigate other confounding factors that may contribute to this performance patterns.

5.5 Continual vs. From Scratch Pretraining

In this section, we test whether the continual pretraining of a base GPT model with additional domain-specific data is helpful in the downstream task performance. We report the zero-shot performance of the tuned model across in-domain (Table 4) and out-of-domain (Table 5) tasks. We have two main observations from this experiment.

First, fine-tuned models fall behind other baselines in a majority of in-domain tasks. HT-CC is the only in-domain task that the tuned model outperforms the rest of models, yet fails to outperform the best performing model trained from scratch.

Second, fine-tuned models have a significant performance drop in the general language modeling tasks (Lambada and Wikitext). For example, the tuned model records 6x performance drop in the Wikitext compared to the best performing model. There are several factors in the continual pretraining that may contribute to this. As the tuned model uses the original GPT-2 vocabulary, it must use the fragmented general subwords to tokenize the chemistry terms available in our corpora. On the other hand, the tuned model starts with the suboptimal initialization from the general-domain language model (Gu et al., 2021). This initialization may diverge the model in the optimization process that may not be recovered.

5.6 Training Efficiency

We use several dimensions to describe the training efficiency, *i.e.*, #FLOPs, throughput (speed), and memory. We compare these compute dimensions





Figure 4: GPU system performance during pretraining.

across the four model sizes described in the Table 2. The smallest (S) model has 59% FLOPs of the largest (XL) model, twice the speed (steps/s), 32% per device GPU memory savings, and 76% total parameter savings (see Figure 4). With such compute budget, small (S) models only outperforms the XL model in 21% in-domain and 34% out-of-domain evaluation tasks. This suggests the importance of compute budget required in scaling foundation models.

6 Conclusions

In this paper, we collected and released 0.67TB of research publication data collected across 10+ sources for chemistry. We pretrained and released 25+ foundation models for chemistry. We rigorously analyzed model performance on 15+ indomain and out-of-domain tasks.

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A Data Descriptions

AMiner ArnetMiner (AMiner) is a service that crawls research publications, performs profile extraction of scientists, models academic networks by integrating publication data from the existing libraries. For the experiments described in this work, we use a sub-sampled version of the data presented in the Open Academic Graph (OAG) version of the AMiner dataset, which originally consisted of more that 172M articles, with 18.5M chemistry-related abstracts.

CORE COnnecting REpositories (CORE) (Pontika et al., 2016) is a large-scale aggregation system which provides an open access to the global network of scientific journals and publications. CORE currently contains more than 207M openaccess articles collected from over 10 thousand data providers, out of which more than 92M are open access full-text research papers. We sub-sampled the original collect into our chemistry-specific corpus consisting of more than 7M full-text articles.

CORD-19 CORD-19 corpus contains COVID-19 (Cord19) and other coronavirus-related publications (e.g. SARS, MERS, etc.) from PubMed's PMC open access corpus, bioRxiv, and medRxiv pre-prints, in addition to COVID-19 articles maintained by the World Health Organization (WHO).

MAG Microsoft Academic Graph (MAG) is a heterogeneous graph created by extracting knowledge from scholarly publications on the web (Wang et al., 2020a). The data used in this work is a subsample from the OAG version of the MAG dataset, which originally consisted of > 208M articles, with 34M chemistry-related articles with abstracts.

PubMed PubMed is a domain-specific data source that allows for search and retrieval of the biomedical and life sciences literature. It is maintained by the National Centre for Biotechnology Information (NCBI) at the U.S. National Library of Medicine (NLM). For this work we utilized the PubMed Central data provided in the Pile corpus (Gao et al., 2020). As presented in Table 3 the sub-sampled data consists of documents with more than 280K abstracts and 700K full text articles.

S2ORC The Semantic Scholar Open Research Corpus (S2ORC) (Lo et al., 2020) is a large academic corpus consisting of 81.1M documents. The data includes the metadata, abstracts, bibliographical references and full-text publications for over 8M open access research articles. In this work, we utilize the sub-sampled version of the original data specific to chemistry, which includes more than 10M abstracts.

WoS The Web of Science (WoS) is a multidiscipline citation database produced by the Institute of Scientific Information. The platform hosts over 171M records across various disciplines, which, when sub-sampled for our chemistry domain, rounded to more than 7M records with abstracts available.

B Task Descriptions

HendrycksTest-Chemistry The Hendrycks Test (Hendrycks et al., 2020) is a large scale collection of multiple choice questions covering 57 subjects. In our experiments, we subsampled college chemistry (HT-CC) and high school chemistry (HT-HC). HT-CC contains 100 questions related to analytical, organic, inorganic, physical, etc. and HT-HC contains 203 questions related chemical reactions, ions, acids and bases, etc.

ARC The ARC dataset (Clark et al., 2018) contains 7,787 genuine grade-school level, science MCQs and is partitioned into a Challenge Set (ARC-C) and an Easy Set (ARC-E). Additionally, 14M science-related sentences are provided with relevant knowledge to answer the ARC questions.

SciQ The SciQ dataset (Welbl et al., 2017) contains 13,679 crowdsourced multiple-choice science exam questions about Physics, Chemistry and Biology, among others.

OpenBookQA The OpenBookQA (Mihaylov et al., 2018) dataset consists of 5,957 multiple choice questions and 1,326 elementary-level science facts. The facts alone do not contain enough information to correctly answer the multiple choice questions, therefore the task is designed to evaluate systems beyond paraphrase matching.

Pile PubMed Abstracts The Pile dataset (Gao et al., 2020) contains 800GB of diverse text sources for benchmarking language models. We limit this task to only include abstracts from the Pile's PubMed collection. As this is framed as a language modeling task, we report word level perplexity.

BoolQ BoolQ (Clark et al., 2019) is a reading comprehension dataset comprised of 16k real, naturally formed queries to the Google search engine with a yes or no answer. Each question-answer pair is accompanied by a Wikipedia article providing evidence to support the correct answer.

CB Commitment Bank (CB) (De Marneffe et al., 2019) is a 3-way classification of textual entailment (true, false, unknown) from 1,200 short text segments where at least one sentence contains an embedded clause. The dataset contains passages from three sources: the Wall Street Journal, the British National Corpus, and Switchboard.

WIC The Word-in-Context dataset (WIC) (Pilehvar and Camacho-Collados, 2018) is a benchmark for evaluating context-sensitive word embeddings. The task is to classify if a target word has the same meaning in two context sentence.

WSC The Winograd Schema Challenge (WSC) (Levesque et al., 2012) dataset is a collection of 804 sentences in which the task is to resolve coreferences.

MathQA MathQA (Amini et al., 2019) is a dataset containing 37k multiple choice math word problems built from the existing dataset, AQuA (Ling et al., 2017).

PIQA The Physical Interactions: Question Answering (PIQA) (Bisk et al., 2020) benchmark dataset provides 21k questions about the physical world and plausible interactions encountered by humans. Annotators provided correct and incorrect answers to questions extracted from instructables.com, a website of instructions for completing many everyday tasks.

PubMedQA The PubMedQA dataset (Jin et al., 2019) is a collection of 273.5k biomedical research questions and related PubMed articles with yes/no/maybe answers.

Lambada Lambada (Paperno et al., 2016) contains passages and target sentences from 5,325 novels collected from Book Corpus (Zhu et al., 2015), and the goal is to predict the last word of the target sentence given the context passage. This task was designed to test genuine language understanding since accurate prediction of the final word would be improbable without the context passage.

WikiText The Wikitext benchmark (Merity et al., 2016) is a language modeling dataset of 29k articles from Wikipedia. Only articles classified as *Good* or *Featured* by Wikipedia editors are included since

they are considered to be well written and neutral in language. All results are reported on Wikitext-2.