Language models for the prediction of SARS-CoV-2 inhibitors

Andrew E Blanchard, John Gounley, Debsindhu Bhowmik, Mayanka Chandra Shekar, Isaac Lyngaas, Shang Gao, Junqi Yin, Aristeidis Tsaris, Feiyi Wang and Jens Glaser

Abstract
The COVID-19 pandemic highlights the need for computational tools to automate and accelerate drug design for novel protein targets. We leverage deep learning language models to generate and score drug candidates based on predicted protein binding affinity. We pre-trained a deep learning language model (BERT) on ~9.6 billion molecules and achieved peak performance of 603 petaflops in mixed precision. Our work reduces pre-training time from days to hours, compared to previous efforts with this architecture, while also increasing the dataset size by nearly an order of magnitude. For scoring, we fine-tuned the language model using an assembled set of thousands of protein targets with binding affinity data and searched for inhibitors of specific protein targets, SARS-CoV-2 Mpro and PLpro. We utilized a genetic algorithm approach for finding optimal candidates using the generation and scoring capabilities of the language model. Our generalizable models accelerate the identification of inhibitors for emerging therapeutic targets.

Keywords
COVID-19, drug design, machine learning, language model, pre-training, fine-tuning, genetic algorithm

1. Justification for Prize
We:

- pre-train a BERT model on a dataset of 9.6 billion molecules, nearly an order of magnitude larger than previous efforts (1.1–1.6 billion) (Jacobs et al., 2021; Xue et al., 2020),
- achieve 603 petaflops in mixed precision on 4032 Summit nodes, reducing pre-training time-to-solution from days to hours, and
- train a general model for protein binding affinity, accelerating the search for drug candidates relevant to SARS-CoV-2.

2. Performance attributes

| Performance attribute          | Our submission                      |
|--------------------------------|-------------------------------------|
| Category of achievement       | Time-to-solution, scalability       |
| Type of method used            | Machine learning                    |
| Results reported for           | Whole application with and without I/O |

3. Overview of the problem
The COVID-19 pandemic has drastically altered living conditions in countries throughout the world over the past 2 years. To date, approximately 230 million people have been infected and 4.7 million have been killed by variants of...
the SARS-CoV-2 virus (Dong et al., 2020). It is not unrealistic to assume that another event like this is possible; several infectious diseases with the potential for global impact have been documented in recent years, including SARS, MERS, Ebola, and Zika (Reperant and Osterhaus, 2017). Within this broader context, the current pandemic highlights the need for the development of therapeutic agents to combat emerging infectious diseases. Unfortunately, the speed at which antivirals have been developed has not maintained pace with the frequency of outbreaks. For example, although vaccines have been developed as an effective means to prevent SARS-CoV-2 infection, no clinically tested therapeutics have been approved for widespread use except for antibody treatment (WHO, 2021; NIH, 2021; HHS, 2021). Furthermore, recent clinical trials have highlighted the need for antivirals (Fischer et al., 2021). Therefore, the timely development of drugs to treat emerging viral threats, in combination with preventive vaccines, poses a key challenge with global implications.

Although many previous efforts in drug discovery have been successful, the process can be prohibitively long (i.e., 10–15 years) for response to an emerging pandemic. The approval of a single compound for widespread use typically involves the screening of small molecules for potential candidates, hit-to-lead testing followed by extensive multi-stage clinical trials (Hughes et al., 2011). The initial step of determining interesting molecules for further investigation is pivotal due to the vast size of chemical space, which prevents an exhaustive search using costly experiments and trials. To accelerate the screening process, tools from machine learning (ML) and high-performance computing (HPC) have been increasingly used to guide the selection of promising drug candidates (Minnich et al., 2020; Sanchez-Lengeling and Aspuru-Guzik, 2018; Vanhaelen et al., 2020). Although computational methods can partially alleviate some of the associated experimental costs, typical approaches require the creation of a large compound library with measured properties for ML model training (Minnich et al., 2020; Vanhaelen et al., 2020). Therefore, a timely response to an emerging pandemic also poses a challenge for computational methods, as custom models and datasets must be quickly generated for the new targets of interest.

To overcome the challenges associated with accelerating the discovery of drug candidates for novel protein targets, a computational approach is needed that satisfies the following criteria: (i) leverages existing large compound libraries without the need for chemical property measurements; (ii) predicts affinities for novel protein targets with limited or no additional experimental data; (iii) explores chemical space to efficiently identify compounds for further investigation. To satisfy the three criteria, we leverage HPC to train generalizable ML models for both candidate generation and affinity prediction.

To take advantage of large existing compound libraries, we utilize a text representation for molecule data known as SMILES, Simplified Molecular Input Line Entry System (Weininger, 1998). Using Enamine REAL database (Enamine, 2020) as a starting point, we generate a novel dataset of approximately 9.6 billion unique molecules. The dataset is used to pre-train a Transformer model (i.e. BERT), using the mask prediction task commonly found in natural language processing applications. During pre-training, subsequences of a given molecule are replaced by a mask, and the model must predict the appropriate sequence based on context. Therefore, the model learns a representation for chemical structure in a completely unsupervised manner that does not require additional property measurements.

To predict affinities for protein targets, we fine-tune the pre-trained molecule model on a dataset with over a million known protein and ligand binding affinities. The fine-tuned model utilizes two pre-trained language models to generate embeddings for a given molecule and protein. For the protein embedding, we utilize a recently published Transformer model for protein sequences (Elmagar et al., 2021). By using models for molecules and proteins trained in an unsupervised manner on large datasets, the fine-tuned model leverages the structural information in the respective embeddings. A final cross-attention layer is added on top of the embeddings to generate an affinity score for any given protein and molecule combination. The fine-tuned model, therefore, can be used to predict affinities for novel proteins outside the training set and/or can be additionally fine-tuned given new experimental data.

The pre-trained and fine-tuned models enable both the generation and scoring of new candidates. For a given input molecule, the pre-trained model can be used to predict viable sub-sequence rearrangements similar to the mask prediction task. The fine-tuned model can then be used to predict the binding affinity for a newly generated molecule with a provided protein sequence. We utilize a genetic algorithm to automate rounds of molecule generation, scoring, and the selection of high scoring candidates.

The large scale of the pre-training and fine-tuning datasets necessitates the HPC resources of a leadership computing facility. Similar to natural language processing applications, the scale of the dataset (i.e., billions of molecules) enables the pre-trained model to learn generalizable features of molecule structure that are useful for affinity prediction. Fortunately, after pre-training and fine-tuning, the models can be used for inference or genetic algorithm optimization with modest resources (i.e., a single GPU). Therefore, our work provides models that can generalize to new protein targets, accelerate screening of potential candidates with limited or no additional fine-tuning, and be utilized throughout the research community for drug discovery efforts.
4. Current state of the art

4.1. Drug discovery pipelines

The complexity and challenges associated with the drug discovery process have motivated the development of pipelines to collect data and organize research efforts (Gentile et al., 2020; Minnich et al., 2020). The computational techniques in such pipelines are primarily organized around the generation and scoring of new molecules. For molecule generation, multiple different representations have been used (e.g., SMILES and graph) along with several ML model architectures (e.g., generative adversarial networks (GAN, VAE, RNN) (Arús-Pous et al., 2019; Blanchard et al., 2021; De Cao and Kipf, 2018; Jacobs et al., 2021; Segler et al., 2018). In addition, manually defined rules (e.g., add an atom, change atom type) have been used to generate new candidates starting from an initial population of molecules (Jensen, 2019; Virshup et al., 2013). For scoring candidates, both ML models and docking simulations have been used along with hybrid approaches (Acharya et al., 2020; Gentile et al., 2020; Minnich et al., 2020). The features used in ML models for scoring range from learned embeddings to chemical descriptors and molecular fingerprints (Minnich et al., 2020).

Determining the correct metric for scoring and optimizing molecules is a key difficulty for the practical application of computational drug discovery pipelines. Cheminformatics packages, such as rdkit (RDKit, 2021), provide standard heuristic metrics for chemical properties, including solubility (Wildman and Crippen, 1999), synthesizability (Ertl and Schuffenhauer, 2009), and quantitative estimation of drug-likeness (Bickerton et al., 2012). However, these metrics are not specific for a given therapeutic target. Alternatively, a supervised ML model for scoring can provide customized optimization metrics but requires a suitable experimentally measured dataset (Martins et al., 2012; Subramanian et al., 2016). To overcome this difficulty, we here utilize a strategy from natural language processing, where a model is initially trained in an unsupervised manner before being fine-tuned to make specific predictions.

4.2. Transformers

Over the past few years, the field of natural language processing (NLP) has undergone a paradigm shift powered by the use of Transformer-based models (Devlin et al., 2019). Previously, the application of ML models was largely task specific, with a single model being trained in a supervised manner for each task (e.g., classification, similarity, entity recognition). However, with the introduction of Transformer models (e.g., BERT), training was split into two distinct stages. In the first stage (i.e., pre-training) the model is typically trained on a large corpus of text in an unsupervised manner. Unsupervised training was accomplished by using a mask prediction task, in which the model was trained to predict a given word based on context. In the second stage (i.e., fine-tuning), the pre-trained model is trained in a supervised manner on a relatively small labeled dataset. In this way, a single pre-trained model can be fine-tuned for any number of specific tasks. Models developed according to this two stage approach have achieved state-of-the-art results for a number of NLP tasks (Devlin et al., 2019; Gu et al., 2020; Liu et al., 2019).

Advances in NLP can be directly applied to drug discovery efforts, as proteins and molecules can be represented as sequences of text. Recent efforts have indeed trained Transformer models using molecules in SMILES format for chemical property prediction tasks (Chithrananda et al., 2020; Honda et al., 2019; Wang et al., 2019; Xue et al., 2020). Most previous work, however, has focused on a language model vocabulary of individual characters or atoms within a sequence, limiting the ability of the model to concisely represent commonly occurring chemical structures. Furthermore, the largest dataset used for pre-training contained approximately 1 billion molecules (Xue et al., 2020), with most investigations using fewer than 100 million (Chithrananda et al., 2020; Honda et al., 2019; Wang et al., 2019). With the success of previous Transformer models using SMILES and text data, we were motivated to increase the number of pre-training samples by an order of magnitude and utilize different model vocabularies.

Transformer models have also been trained using protein sequence data. A recent study investigated the performance of multiple model architectures on protein prediction tasks (Elmaggar et al., 2021). Furthermore, the outputs of pre-trained models for both molecule data and protein data can be used as embeddings for additional downstream tasks (Gurbych et al., 2020). In the context of drug discovery, this enables the pre-trained models to be fine-tuned on a dataset consisting of many different protein and ligand combinations with experimentally determined binding affinity. Notably, Transformer-based approaches have shown significant performance improvements for affinity prediction over alternative architectures, such as convolutional neural networks (Gurbych et al., 2020; Öztürk et al., 2018). In the current work, we leverage the Transformer architecture to develop a fine-tuned model capable of predicting binding affinity for novel protein targets.

4.3. Deep learning at scale

Using increasingly large training datasets poses a substantial challenge in terms of time-to-solution. Data parallelism enables many deep learning models to be trained efficiently at the scale of current supercomputers (Kurth et al., 2018; Laanait et al., 2019). Transformers are one such
model; for example, large scale data parallelism has been used to dramatically reduce BERT pre-training times (Rasley et al., 2020; You et al., 2019; Zheng et al., 2020). Larger Transformer models, such as Megatron-LM, have achieved performance of over 500 petaflops in mixed precision on Nvidia’s Selene supercomputer (Narayanan et al., 2021). Inasmuch as a previous effort to train a BERT model using 1 billion molecules required approximately 4 days for pre-training (Xue et al., 2020), the potential advantage of using large scale data parallelism to enable pre-training on larger datasets with faster turnaround times is clear.

However, extreme scale data parallelism necessarily leads to extremely large batch sizes and large batch sizes can lead to instability during training which degrades model evaluation performance. While this problem is general (Goyal et al., 2017), it has also specifically been observed as a scaling bottleneck in the context of developing deep learning models for drug discovery: a recent study found that an overall batch size above 4096 (across eight or 16 GPUs) was detrimental to model training on molecule data for a variational autoencoder (Jacobs et al., 2021). The recently developed LAMB optimizer has been shown to address this problem for batch sizes of up to 96 thousand, maintaining similar evaluation performance as batch size increased (You et al., 2019; Zheng et al., 2020).

### 4.4. Genetic algorithms

Inspired by the mutation and selection observed in natural systems, genetic algorithms provide a useful framework for solving optimization problems across scientific and engineering disciplines (Brown et al., 2004; Eiben and Smith, 2015; Hornby et al., 2011; Morse and Stanley, 2016). Specifically, for drug discovery, genetic algorithms have been used in several studies to search chemical space. For example, Virshup et al. proposed a set of hand-crafted rules for mutation and recombination (e.g., add an atom, modify an atom type) to generate new compounds. The generated compounds were then selected based on diversity criteria to expand to unexplored regions of chemical space (Virshup et al., 2013). Additional studies have used genetic algorithms to optimize for drug-specific metrics (e.g., solubility and quantitative estimation of drug-likeness) (Brown et al., 2004, 2019; Jensen, 2019; Yoshikawa et al., 2018). Typically, mutation operators are manually defined based on the application and not learned from the data. However, comparisons with alternative ML optimization techniques have shown that genetic algorithms perform well across a range of drug discovery tasks (Brown et al., 2019).

As an alternative to the manually defined mutation and recombination operators, molecule rearrangements can be determined by a ML model. Generative models, such as GANs can be used to produce molecules with desired properties (Blanchard et al., 2021; De Cao and Kipf 2018). Furthermore, masked language models provide a useful modeling framework in which to learn viable rearrangements of molecular sequences. During pre-training the language model learns to predict missing sequences based on context. The predictions provide a ranked list of all possible substitutions for a given sub-sequence. Therefore, by sampling from the predictions, a set of mutations can be generated for a molecule without the need for manually defined rules. A similar procedure has been used to find adversarial examples for NLP applications (Li et al., 2020a, 2020b). In this work, we utilize a masked language model to generate candidate molecules and then apply selection based on scoring from the fine-tuned model for binding affinity.

### 5. Innovations realized

Our strategy for accelerating computational drug discovery is summarized in Figure 1. We begin by constructing the largest molecule dataset to date for pre-training a masked language model. Pre-training is performed at scale using a batch size of over a million molecules for two different tokenization schemes. We then fine-tune the language model on a dataset with binding affinities for thousands of protein targets. After developing the general pre-trained and fine-tuned models, we search for drug candidates that optimize the predicted binding affinities for a given protein.

#### 5.1. Pre-training molecule language models

**5.1.1. Dataset generation.** Motivated by the success of Transformer models (i.e., BERT) for a range of natural language processing tasks, recent efforts have investigated using the SMILES text representation of molecules to train a masked language model (Chithrananda et al., 2020; Honda et al., 2019; Wang et al., 2019; Xue et al., 2020). Large compound libraries such as Enamine REAL database (Enamine, 2020) can be used for the unsupervised pre-training stage, before the model is fine-tuned for a desired prediction task. Although compound libraries provide a valuable source of training data, the overwhelming size of chemical space ensures that many potentially useful compounds will be excluded from current collections. Current state-of-the-art generative and masked language models have reached a training data size of approximately 1.1–1.6 billion compounds, pushing the boundaries of currently available compound libraries and compute resources (Jacobs et al., 2021; Xue et al., 2020). As a step toward enabling larger explorations of chemical space, here, we utilize the Enamine REAL database as a starting point to generate a training dataset with ~9.6 billion unique molecules.
Our strategy for dataset augmentation is motivated by the pre-training stage for masked language models. During pre-training, random sequences of an input molecule SMILES are masked, and the model is trained to predict the identity of the masked sequences based on the surrounding context. We, therefore, use a pre-trained model, developed using Enamine as the training data, to predict possible structural rearrangements for a given molecule as shown in Figure 1. We also use the pre-trained model to combine two molecules; initial sequences are chosen from each respective molecule, and a mask is placed in between. To be included in the training set, all produced molecules must be valid (RDKit, 2021) and have a normalized synthesizability (De Cao and Kipf, 2018; Ertl and Schuffenhauer, 2009) score above 0.3. To arrive at the final dataset of 9.57 billion molecules, we applied random structural rearrangements to molecules in the Enamine dataset with a maximum of five masks per molecule. The top five predicted rearrangements from the pre-trained model were considered, resulting in the dataset growing from approximately 1.34 billion unique compounds to 4.14 billion unique compounds. Another round of rearrangements for the expanded dataset was accompanied by combinations of molecules to generate the final training set. All molecules were converted to canonical form using rdkit (RDKit, 2021) and only unique molecules were retained in the final dataset. As shown in Figure 2, the histograms for drug-likeness, normalized synthesizability, and solubility did not substantially change between the original data and the augmented data, although the total number of compounds substantially increased.

5.1.2. Tokenization. The process of tokenization is used to convert any given sequence of text into a format that can be recognized by the model. This is accomplished by constructing a vocabulary for the model, which consists of a mapping between sub-sequences of text and unique integer ids. One common method, WordPiece tokenization (Schuster and Nakajima, 2012; Wu et al., 2016), builds the model vocabulary by assembling all unique single characters and then including commonly occurring sequences of increasing length. Although a collection of different tokenization methods have been used for NLP tasks, for molecule language models a simple vocabulary based on single atoms and characters has predominantly been used (Arús-Pous et al., 2019; Grisoni et al., 2020; Honda et al., 2019; Schwaller et al., 2018; Segler et al., 2018; Wang et al., 2019; Xue et al., 2020). Here, we utilized two different tokenization methods, the standard single atom and character Regex (Schwaller et al., 2018) and WordPiece tokenization. As shown in Figure 3, the vocabulary generated by WordPiece tokenization enables large sub-sequences of a SMILES string (e.g., a benzene ring) to be represented as a single token. Notice that for the Regex tokenizer, the vocabulary will contain only individual characters and atoms, so commonly occurring chemical structures cannot be assigned to a single token. Also, the size of the vocabularies for the two tokenizers is drastically different, with the WordPiece tokenizer having $3 \cdot 10^4$ different tokens, while the Regex tokenizer has around 200. Given the substantially different representations of molecules produced by the two tokenizers, we utilized both methods for pre-training and

![Figure 1. Our strategy for developing general models for drug discovery contains three components.](image1)

![Figure 2.](image2)

![Figure 3.](image3)
5.1.3. Pre-training with large batch sizes. We trained BERT models using the LAMB optimizer with both the Regex and WordPiece tokenizers utilizing different numbers of nodes on Summit. Each node contained six GPUs, each having a single partition of $3.95 \cdot 10^5$ unique molecules. The batch size per GPU was set to 240 (80 with three gradient accumulation steps); therefore, the total batch size is given by $1440$ (i.e., $240 \cdot 6$) multiplied by the number of nodes (Lin et al., 2017). At 1000 nodes, this results in over 1.4 million molecules per batch. As shown in Tables 1 and 2, even with large batch sizes, the model can be trained successfully, as evidenced by the validation accuracy for mask prediction. Validation accuracy was determined by evaluating the model on a hold-out set of $1.05 \cdot 10^6$ molecules; for each molecule a random number of masks (up to five for Regex and up to three for WordPiece) were sampled and used to replace tokens. Each pre-training run consisted of seven epochs, with model checkpoints saved and validation accuracy determined after each epoch. The maximum accuracy across checkpoints is shown. Notice that a comparison of accuracy between Tables 1 and 2 should not be made, as the mask prediction task is substantially easier for the Regex tokenizer (i.e., only single atoms or characters are predicted).

For the full dataset, we used two different training configurations. First, we used 1008 nodes, with four partitions of $3.95 \cdot 10^5$ unique molecules per GPU. For comparison, we also performed pre-training on over 4000 nodes for the WordPiece tokenizer with a single partition per GPU (last row of Table 1). Both pre-training runs used a warmup of 1 epoch. As expected from previous studies (You et al., 2019), the increased batch size for the 4032 node run resulted in decreased validation accuracy; however, it is notable that a batch size of nearly six million incurred only a slight decrease in performance, suggesting that distributed training for even larger molecule datasets is possible. The 1008 node runs were completed in 8 h each for 7 epochs; the 4032 node run was stopped after failing to increase validation accuracy (maximum was at 5 epochs), taking less than 2 h. For downstream tasks, such as fine-tuning, we used the models trained on 1008 nodes for the WordPiece and Regex tokenizers.

5.2. Predicting binding affinities

To determine whether a given drug molecule binds to a target molecule, that is, a protein, both the candidate molecule and the amino acid sequence of the protein need to be embedded. Then, to predict the binding affinity, hidden layers are added that accept the concatenation of the two embeddings as inputs. The predictive power of such an ensemble model is chiefly determined by the expressive power of the individual embeddings. Therefore, we expect that using powerful pre-trained models to embed the molecules results in superior performance on the downstream task. Here, we focus on regression to predict the numerical value of the binding affinity, however, the model architecture lends itself equally well to classification. Pre-trained embeddings and extra layers are fine-tuned simultaneously, that is, with all weights adjustable, on a labeled dataset of $1.67 \cdot 10^6$ receptor amino acid sequences and ligand SMILES, with binding affinities. Note that this dataset used for fine-tuning is much smaller than the datasets used for pre-training (Tables 1 and 2).
5.2.1. Binding affinity dataset. We curated a dataset (Glaser 2021) of binding affinities by concatenating the entries of the BindingDB (Liu et al., 2007), PDBBind-cn (Liu et al., 2015), BindingMOAD (Hu et al., 2005), and BioLIP (Yang et al., 2012) databases, following the example of Ref. (Oleksandr et al., 2020). Records containing $K_i$, $1/K_{i^h}$, $K_{i^h}$, and IC$_{50}$ values were retained and converted to \( \text{pK}_d = -\log_{10} K_{d[M]} \) units, and MACCS fingerprints were calculated on the molecules to remove duplicates, resulting in 1,670,637 protein sequences, SMILES strings and binding affinities.

5.2.2. Model architecture. Figure 4 shows the neural network architecture used to predict affinities. For embedding molecules, we use the tokenizers and pre-trained models discussed above. For embedding proteins, we make use of the readily available pre-trained ProtBERT model (Wolf et al., 2020), where every token is a letter in the amino acid alphabet. The embeddings are fed to a cross-attention module (Koyama et al., 2020). The purpose of the cross-attention layer is that molecule sub-units attend to amino-acids in the protein sequence, and vice versa. This architecture represents the physical situation in which the molecule makes well-defined atom-atom contacts with the protein. However, the model is not constrained to learn real physical contacts, and importantly, it is not given any information about which residues belong to the active site of the protein, which it has to learn by itself from the given correlation between structures and binding affinities. Despite the physical motivation behind its architecture, the model is still to be considered as a “black-box.” It cannot be expected that the cross-attention weights directly correspond to observable physical contacts, as sometimes suggested (Koyama et al., 2020; Oleksandr et al., 2020).

The hidden layer outputs of the cross-attention module are concatenated, their mean is taken over the sequence length and they are connected to a linear layer to predict the binding affinity. The model is fine-tuned by minimizing the mean-squared error (MSE) between the predicted and the experimental affinity. We validated the model on a hold-out set from the training data as well as three additional datasets as shown in Tables 3 and 4. We characterize the ability of the model to correctly reproduce the order of the experimental affinity values by the Spearman-$\rho$ rank correlation coefficient (higher is better) (Spearman, 1961), as well as the mean-squared error for the predicted affinity (lower is better). We calculate the uncertainty in the reported values using the bootstrap method with \( n = 500 \) samples. Notably, the Regex tokenizer (and

| Table 1. Validation accuracy for pre-training runs with WordPiece tokenizer. | Table 2. Validation accuracy for pre-training runs with regex tokenizer. |
|---|---|---|---|
| Nodes | Molecules | Batch size | Accuracy | Nodes | Molecules | Batch size | Accuracy |
| 1 | $2.4 \cdot 10^6$ | $1.4 \cdot 10^3$ | 0.760 | 1 | $2.4 \cdot 10^6$ | $1.4 \cdot 10^3$ | 0.845 |
| 10 | $2.4 \cdot 10^7$ | $1.4 \cdot 10^4$ | 0.783 | 10 | $2.4 \cdot 10^7$ | $1.4 \cdot 10^4$ | 0.866 |
| 100 | $2.4 \cdot 10^8$ | $1.4 \cdot 10^5$ | 0.797 | 100 | $2.4 \cdot 10^8$ | $1.4 \cdot 10^5$ | 0.878 |
| 1000 | $2.4 \cdot 10^9$ | $1.4 \cdot 10^6$ | 0.798 | 1000 | $2.4 \cdot 10^9$ | $1.4 \cdot 10^6$ | 0.882 |
| 10008 | $9.6 \cdot 10^7$ | $1.5 \cdot 10^6$ | 0.808 | 10008 | $9.6 \cdot 10^7$ | $1.5 \cdot 10^6$ | 0.801 |
| 4032 | $9.6 \cdot 10^7$ | $5.8 \cdot 10^6$ | 0.801 |

Figure 4. Architecture of the affinity prediction model used for fine-tuning. It uses two independent models for protein sequences (Protein Language Model) and molecule SMILES (Molecule Language Model) connected to a cross-attention module that predicts the logarithm of the binding affinity.
associated ensemble) outperforms the WordPiece tokenizer for certain datasets (i.e., Hold-out and Kinases), but under-performs for others (i.e., PLpro), suggesting that different molecule representations may be suitable for different affinity prediction tasks.

Figure 5 demonstrates the performance for the validated and transferable model on a binary classification task, using the metrics of precision and recall. Here, we impose a threshold of 5\(\mu\) M (Mpro) and 1\(\mu\) M (PLpro) on the experimental IC50 value (lower is better) to label active molecules. These thresholds are typical of more potent non-covalent inhibitors for the Mpro and PLpro targets. We use sampling from the normal distribution of affinities implied by the mean and the variance of the ensemble model to estimate the confidence intervals, as well as the standard error from \(n = 500\) bootstrap samples. Remarkably, the model achieves a maximum precision of 0.60 both for Mpro and PLpro, meaning that 60% of the highest scoring molecules are true actives, which suggests excellent virtual-screening performance.

5.3. Searching for drug candidates

Similar to the strategy we used for data augmentation, the pre-trained model can easily be adapted to generate rearrangements for a given population of molecules. Here, we utilize three different types of rearrangements for a given tokenized molecule: insertion, deletion, and replacement. For insertion, a mask is randomly inserted in between existing tokens or at the beginning or end. For deletion, a mask randomly replaces two adjacent existing tokens. With replacement, a single existing token is randomly masked. To search for new drug candidates, we randomly sampled up to five masks and a rearrangement type for molecules in the population. In addition, we consider recombination by randomly sampling two molecules, selecting a sub-sequence from each and inserting a mask in between. A canonical and randomized SMILES were used to represent each molecule before masking, and the top 10 molecules predicted by the pre-trained model were used as candidates. As a starting population, we used molecules from the validation set for pre-training. Only unique molecules were retained in the population, as determined by canonical SMILES computed using rdkit (RDKit, 2021).

To score the molecules generated through rearrangements, we utilized three metrics: normalized synthesizability, quantitative estimation of drug-likeness, and the affinity predictions of the fine-tuned model. The predicted score for the affinity was divided by 10 and clipped between 0 and one to generate a normalized affinity metric. The harmonic mean of the three metrics was then used to define the fitness of a given candidate. To find optimized candidates, an initial population of \(10^4\) molecules was used from the validation set for pre-training. Then, masked rearrangements were applied to \(5 \times 10^3\) samples and recombination was applied to \(5 \times 10^5\) sampled pairs. The resulting molecules were added to the population and ranked according to fitness; the top \(10^4\) overall were retained as the starting population for the next generation. Based on the Hold-out fine-tuning results, we selected Model 3 with a Regex Tokenizer to predict scores. For molecule

| Test set              | Ensemble \(\rho\) | Model 1 \(\rho\) | Model 2 \(\rho\) | Model 3 \(\rho\) |
|----------------------|------------------|------------------|------------------|------------------|
| Hold-out             | 0.881(2)         | 0.69(1)          | 0.866(3)         | 0.77(1)          |
| Kinases (Davis et al., 2011) | 0.38(1)         | 1.67(3)          | 0.35(1)          | 1.80(3)          |
| Mpro (Achdout et al., 2020) | 0.39(2)         | 1.82(5)          | 0.28(2)          | 1.79(5)          |
| PLpro (Shen et al., 2021) | 0.54(8)         | 0.8(1)           | 0.55(8)          | 0.9(1)           |

| Test set              | Ensemble \(\rho\) | Model 1 \(\rho\) | Model 2 \(\rho\) | Model 3 \(\rho\) |
|----------------------|------------------|------------------|------------------|------------------|
| Hold-out             | 0.864(3)         | 0.72(1)          | 0.830(3)         | 0.94(2)          |
| Kinases (Davis et al., 2011) | 0.30(1)         | 2.03(3)          | 0.23(1)          | 2.90(4)          |
| Mpro (Achdout et al., 2020) | 0.37(2)         | 1.23(5)          | 0.33(3)          | 1.23(5)          |
| PLpro (Shen et al., 2021) | 0.57(8)         | 0.71(10)         | 0.51(8)          | 0.45(7)          |
rarrangements, we used the pre-trained model with WordPiece Tokenizer as it generated higher fitness scores than the corresponding Regex Tokenizer.

As shown in Figure 6 (top two rows), 50 generations of optimization to search for Mpro inhibitors resulted in a substantial shift in the distributions of the three optimized metrics. Notably, the mean for the affinity score increases, with the maximum generated molecule having a normalized affinity score greater than 0.9. By optimizing for synthesizability and drug-likeness in addition to affinity, the generated molecules are constrained by useful heuristic scoring functions for drug discovery (Bickerton et al., 2012; Ertl and Schuffenhauer, 2009). We also show the top scoring molecule in the final population for each respective metric. The three examples show that optimization successfully found molecules with higher predicted affinities while maintaining high synthesizability and drug-likeness scores. The bottom two rows of Figure 6 show the corresponding results for PLpro. The only change to the genetic algorithm is the input protein sequence, highlighting the generality of our approach.

6. How performance was measured

6.1. Applications used to measure performance

In this study, we performed pre-training, fine-tuning, and a genetic algorithm on Transformer models. All models are written in PyTorch using the Hugging Face Transformers API (Wolf et al., 2019). Models are pre-trained and fine-tuned with DeepSpeed (Rasley et al., 2020), a high-performance wrapper for distributed Transformer training. Model training is performed with data parallelism using DeepSpeed’s fused-kernel LAMB optimizers. Sharded I/O is performed using the WebDataset library (Aizman et al., 2019). As pre-training of the molecule language model is by far the most computationally expensive stage of the study, it is the focus of the performance analysis.

The architecture used for the molecule language model is BERT-base, which has approximately 109 million learnable parameters. Pre-training of the model is performed with data parallelism, in which each GPU trains the model on separate data. Communication takes the form of a global asynchronous AllReduce which is performed during back-propagation on each batch.

6.2. Measuring performance

Performance of molecule language model pre-training was measured in two respects. First, sustained performance was measured using built-in timers which report the total wallclock time elapsed during training and the time for I/O operations (specifically, saving checkpoints and trained models). Additionally, to measure peak application performance relative to theoretical machine peak, mixed precision floating point operations per second (FLOPs) are computed using the DeepSpeed FLOPS Profiler.

6.3. System

6.3.1. Hardware. Performance was measured on the Summit supercomputer at the Oak Ridge Leadership Computing Facility at ORNL (Vazhkudai et al., 2018). Summit is comprised of 4674 IBM Power System AC922 nodes which are arranged in a non-blocking Fat Tree topology with dual-rail EDR InfiniBand interconnect. Each node has two IBM Power9 CPUs, six Nvidia 16 GB V100 GPUs, and 512GB of main memory. The V100 device has an estimated peak performance of 14 teraflops for single precision (FP32) and 112 teraflops for mixed precision using the Tensor Cores, which are capable of performing matrix multiply in FP16 with FP32 accumulation for some kernels. Consequently, Summit’s peak performance for mixed precision is approximately 3.1 exaflops.

6.3.2. Software. Summit runs the Red Hat Enterprise Linux eight operating system and uses the IBM LSF job scheduler. Our Python-based software stack uses Open Cognitive Environment v1.2.0, PyTorch v1.7.1, Transformers v4.5.1, DeepSpeed v0.4.5, and WebDataset v0.1.62. The GPU libraries include CUDA 11.0.3, NCCL 2.7.8, and cuDNN 8.0.4.
7. Performance results

7.1. Node level optimization

The results of node level optimization for molecule language model pre-training are shown in Figure 7, in terms of runtime per epoch plotted versus a series of successive optimizations. In this setting, data parallelism is applied to train the molecule language model across the six GPUs of a Summit node. The baseline case uses the Adam optimizer, single precision arithmetic, and the largest batch size per device (96) for which the model fits in GPU memory. Enabling mixed precision with the V100 Tensor Cores decreases runtime by approximately 43%. In addition, mixed precision enables a larger per device batch size (128) to be used while keeping the model within GPU memory, further decreasing runtime per epoch by about 15%. While the decision to switch to the LAMB optimizer was motivated by large batch sizes, as discussed in section 5.1.3, DeepSpeed’s fused LAMB optimizer implementation also improves performance by roughly 15%. Finally, leveraging LAMB’s stability for very large batches, we changed the batch size to 80 and added three gradient accumulation steps, for an effective batch size per device of 240. While the addition of gradient accumulation does not improve node level performance, this configuration enables better scaling at very high node counts due to reduced communication frequency and, after hyperparameter optimization, maintains comparable accuracy.

7.2. Scaling

As this study incorporates the largest molecule dataset ever used for pre-training, the primary focus for scalability was weak scaling. In Figure 8, we assess the weak scaling of pre-training the molecule language model on Summit. For weak scaling, the problem size per device is kept constant at $3.95 \times 10^5$ molecules. The training configuration is that identified in section 7.1, extended to the multi-node setting, with data parallelism used to train a single model across the given number of nodes. These runs are the production runs in section 5.1.3 with the WordPiece tokenizer, and therefore include I/O operations to save checkpoints and the final trained model. Parallel efficiency for weak scaling from 1 to 4032 Summit nodes is measured at 68.0%. However, a significant amount of performance degradation at large node counts is due to I/O; when I/O time is subtracted out, parallel efficiency over the same interval improves to 83.3%. In combination with the validation accuracy results from Tables 1 and 2, this clearly indicates that pre-training can be
extended to incorporate unprecedented molecule dataset and batch sizes without significantly compromising computational efficiency or accuracy.

Strong scaling of the molecule language model pre-training is shown in Figure 9 from 25 to 1600 Summit nodes. The total problem size is kept constant at approximately 1.9 billion molecules, with the same training configuration as for weak scaling. However, as full runs could not be completed within facility wallclock limits for many node counts, the reported runtime per epoch is measured for the first 0.25 epochs for the 25 node job and for the first epoch for all other node counts. Strong scaling is near linear from 25 to 400 Summit nodes and maintains approximately 67.4% parallel efficiency at 1600 nodes versus the 25 node baseline.

7.3. Peak performance

Table 5 shows the peak performance achieved for molecule language model pre-training on Summit during the largest weak scaling run from section 7.2. On 4032 nodes, peak performance of approximately 603.4 petaflops in mixed precision is achieved. As Summit’s theoretical peak at this node count is projected at ~2.71 exaflops, our result represents about 22.3% of this mixed precision peak.

8. Implications

8.1. ML models for drug discovery

Supervised training of ML models for drug discovery poses a key difficulty in terms of both time and resources, as a labeled dataset must be created for each new therapeutic target. As demonstrated by language models for both text (Devlin et al., 2019) and chemical sequence data (Xue et al., 2020), an alternative approach is to leverage large unlabeled datasets to train a general model. The general model is then fine-tuned on a relatively small dataset for a specific task of interest. Although fine-tuning still requires supervised training, unsupervised pre-training has enabled state-of-the-art results across a range of tasks with limited labeled data (Devlin et al., 2019; Gu et al., 2020). Here, we have taken...
the shift towards general models a step further by using pre-training and fine-tuning tasks that can generalize to any protein and molecule sequence.

The ability to pre-train molecule language models with large batch sizes enables an unprecedented exploration of chemical space. Current chemical databases provide hundreds of billions of molecules, but contain only a small fraction of potentially synthesizable molecules (Reymond, 2015). Through the process of tokenization and mask prediction, language models can leverage large datasets to automatically learn commonly occurring sub-sequences (i.e., structural components) and possible rearrangements for effective searches of chemical space.

By combining a pre-trained model for molecule and protein sequences, the fine-tuning task can leverage data from many previous experimental investigations. Furthermore, additional modeling techniques, such as docking simulations, could be used to augment the training data in novel regions of interest. Recent development in protein structure prediction (Jumper et al., 2021) leverages both sequence and spatial information to increase predictive performance. Although there is still much work to be done to make a truly general model for drug discovery, the increase in unsupervised and semi-supervised approaches to training along with the increase in available experimental and simulation data for protein and ligand interactions makes possible the development of off-the-shelf models that generalize across therapeutic targets. Developing a generalizable model is key to reducing the time for discovering and screening new targets in an emerging pandemic, such as COVID-19.

Using a genetic algorithm coupled with a pre-trained language model for optimization enables incremental exploration and refinement from known drugs as well as population searches. For example, a certain sub-sequence of a known compound can be masked, and the language model can predict the most likely rearrangements. Heuristic metrics and ML models can then be used to analyze the expected impact of structural changes. The intuitive process of making incremental changes during exploration can be used to complement and guide researcher intuition during the drug discovery process. Therefore, our suggested optimization strategy provides a natural framework for both fully-automated and user-guided exploration of chemical space.

8.2. HPC resources for model development

The pre-training phase of developing a language model requires substantial computational resources; here, we utilized thousands of GPUs and corresponding node hours to complete training. Similarly, as the labeled dataset and the compound library for screening grows, fine-tuning and inference can also necessitate the resources of a leadership computing facility. Although these resource requirements provide an excellent use case and motivation for the continued development of HPC systems, they generate challenges for the utilization and training of models throughout the research community. Fortunately, the pre-trained models can be leveraged for inference and to some extent fine-tuning applications with only a single GPU.

For fine-tuning tasks with a specific protein target or chemical property, training can typically be done without the need for HPC resources. For inference, our results show that a genetic algorithm using the pre-trained model for mutations can be used to generate optimized candidates without the need for large scale model evaluations. Furthermore, the genetic algorithm approach provides an interpretable scheme for modifying a single molecule. The mask predictions and scores can be inspected to determine single mutations that lead to higher scores for a given metric. Also, the reported genetic algorithm runs from this work used only a single V100 GPU for less than 10 h for optimization. Therefore, although the pre-trained models require substantial computational resources for training, the models can be used for exploration throughout the research community.

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**ORCID iDs**

Andrew E Blanchard  
https://orcid.org/0000-0001-7330-9656  
Jens Glaser  
https://orcid.org/0000-0003-1852-3849

**Note**

1. Pre-trained models are available on Hugging Face with BERT and regex tokenization, along with three fine-tuned models (1, 2, and 3). Repositories with pre-training and fine-tuning code are also available.

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**Author biographies**

**Andrew Blanchard** is a research scientist in the Biostatistics and Multiscale System Modeling Group at Oak Ridge National Laboratory. He received his Ph.D. in Physics from University of Illinois at Urbana-Champaign. His research focuses on using physics-based simulations and machine learning models to develop a quantitative understanding of biological and chemical systems for engineering applications.

**John Gounley** is a computational scientist in the Computational Sciences and Engineering Division at Oak Ridge National Laboratory. He received a PhD in computational and applied mathematics from Old Dominion University in 2014. His research focuses on algorithms and scalability for biomedical simulations and data.

**Debsindhu Bhownik** is a computational Scientist in Computational Sciences and Engineering Division (CSED) at the Oak Ridge National laboratory (ORNL). He holds B.Sc. & M.Sc. in Physics from Jadavpur University, India and a Ph.D. from Université Pierre et Marie Curie (UPMC), France. Upon finishing his Ph.D. he joined as Postdoctoral Research Fellow at the Donostia International Physics Center, Spain and later at the Wayne State University, USA. His current work lies on the interface of implementing Deep Learning (DL) techniques, running Molecular Dynamics (MD) simulations and performing scattering experiments for problems related to Biomedical and biological sciences.
to understand the underneath physics of bio(macro)molecular function, activity, folding, microscopic structure and dynamic behavior at different length and time scale.

Shang Gao received his PhD in Data Science and Engineering at the University of Tennessee. He worked as a research scientist at Oak Ridge National Laboratory, where he was the primary technical lead of a major research collaboration between the National Cancer Institute and U.S. Department of Energy on utilizing AI and HPC to automate cancer surveillance. Shang is currently a senior machine learning researcher at Casetext, where he develops and deploys NLP models for legal search and retrieval.

Mayanka Chandra Shekar is a Research Scientist in Bio-medical Natural Language Processing and AI at Oak Ridge National Laboratory. In ORNL, she worked on natural language processing for free-text electronic health records under the Veteran Affairs’ MVP prostate cancer exemplar and VISION, NCI-DOE MOSSAIC, and Cincinnati Children’s Hospital Medical Center’s Mental Health Trajectory projects. She received her Ph.D. from the University of Missouri-Kansas City in transfer learning and domain adaptation in image and text datasets. During her Ph.D., she did a research internship in Pacific Northwest National Lab on Natural Language Processing, specifically worked on the DARPA Modeling Adversarial Activity (MAA) project on graph generation.

Isaac Lyngaas is a computational scientist in the Life Science and Engineering group in the Oak Ridge Leadership Computing Facility (OLCF) at Oak Ridge National Laboratory (ORNL). He joined ORNL in 2019 after earning his PhD in Computational Science at Florida State University. His research focuses on using machine learning models for natural language processing tasks with emphasis on their computational performance.

Junqi Yin is a computational scientist in Analytics & AI Methods at Scale (AAIMS) group of National Center for Computational Sciences at ORNL. He obtained his Ph.D. in Computational Physics and M.S. in Statistics from the University of Georgia, and B.S. in Physics and Computer Science from Zhejiang University. His research interests range from scalable machine learning/deep learning to materials science, statistical sampling algorithms, HPC, and workflows. He is working with domain scientists to scale up their machine learning applications on DOE leadership-class computing platforms.

Aristeidis Tsaris is a research scientist in large-scale data science and learning at National Center for Computational Sciences at ORNL. His research focus is on scalable machine learning applications on HPC systems, benchmarking, and imaging. He has been working with scientific applications to scale their analytics algorithms on HPC systems. Before he joined ORNL, Aristeidis received his PhD on Medium Energy Particle Physics from Florida State University, working at Jefferson Lab. He also worked as a postdoc at Scientific Computing Division (SCD) at Fermi National Lab where he contributed to a wide variety of deep learning applications for neutrino and collider physics.

Feiyi Wang received his Ph.D. in Computer Engineering from North Carolina State University (NCSU). He is a Senior Computer Scientist and Group Leader of Analytics and AI methods at Scale Group (AAIMS) at National Center for Computational Sciences of ORNL. His current research interests include large-scale data analytics, distributed machine learning and benchmarking, high performance storage system, parallel I/O and file systems. Dr. Wang is a Joint Faculty Professor at University of Tennessee and a senior member of IEEE.

Jens Glaser is a computational scientist at Oak Ridge National Laboratory’s National Center for Computational Sciences, where his research involves developing novel algorithms for biological and soft matter simulation, with application to nanoparticles and synthetic biomaterials made from protein building blocks, and drug discovery. Jens obtained his doctorate in physics from Leipzig University in Germany. He worked as a postdoc at the University of Minnesota and as assistant research scientist at the University of Michigan. A common thread in his research connects the simulation of collective phenomena in many-particle systems and the search for parallel approaches to molecular simulation, using GPUs. He’s an active developer for the open-source particle simulation toolkit HOOMD-blue.