META-LEARNING INITIALIZATIONS FOR LOW-RESOURCE DRUG DISCOVERY

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ABSTRACT

Building in silico models to predict chemical properties and activities is a crucial step in drug discovery. However, drug discovery projects are often characterized by limited labeled data, hindering the applications of deep learning in this setting. Meanwhile advances in meta-learning have enabled state-of-the-art performances in few-shot learning benchmarks, naturally prompting the question: Can meta-learning improve deep learning performance in low-resource drug discovery projects? In this work, we assess the efficiency of the Model-Agnostic Meta-Learning (MAML) algorithm – along with its variants FO-MAML and ANIL – at learning to predict chemical properties and activities. Using the ChEMBL20 dataset to emulate low-resource settings, our benchmark shows that meta-initializations perform comparably to or outperform multi-task pre-training baselines on 16 out of 20 in-distribution tasks and on all out-of-distribution tasks, providing an average improvement in AUPRC of 7.2% and 14.9% respectively. Finally, we observe that meta-initializations consistently result in the best performing models across fine-tuning sets with $k \in \{16, 32, 64, 128, 256\}$ instances.

1 INTRODUCTION

Drug discovery is a multi-parameter optimization process requiring efficient exploration of chemical space for compounds with desired properties. In a typical project, medicinal chemists propose structural changes to compounds in an effort to improve their therapeutic effects without compromising other properties. Validating these changes are costly – e.g. compounds need to be purchased or synthesized, assays need to be developed and validated – and thus in silico models are often used to prioritize experiments. Following the Merck Molecular Activity Challenge, there has been significant interest in applying deep learning to property prediction. More recently, by directly learning molecular features from chemical graphs, novel architectures in the graph neural networks family have demonstrated improved predictions in quantum chemistry and various property prediction benchmarks (Lusci et al., 2013; Duvenaud et al., 2015; Kearnes et al., 2016; Gilmer et al., 2017; Feinberg et al., 2018; Yang et al., 2019).

The successes of deep learning, however, hinge on an abundance of data: For instance, ImageNet (Deng et al., 2009) contains over 14M images and the English Wikipedia database commonly used to pre-train language models has over 2,500M words. On the contrary labeled scientific data in drug discovery projects often consists of many small, sparse, and heavily biased datasets, consequently limiting the applications of deep learning in this setting. Recent works approach this problem by using pre-training and multitask learning to leverage data from multiple sources (Ramsundar et al., 2015; Wenzel et al., 2019; Hu et al., 2019).

In parallel, the problem of learning in low-data domain has been tackled vehemently by the few-shot learning community. A prominent solution is the meta-learning paradigm, which aims to learn a learner that is efficient at adapting to new task (Thrun & Pratt, 1998; Vilalta & Drissi, 2002; Vanschoren, 2018). Matching Networks (Vinyals et al., 2016), a member of this family, have been previously applied to property prediction in one-shot learning settings by (Altae-Tran et al., 2017).
A related approach is the Model-Agnostic Meta-Learning (MAML) algorithm (Finn et al. [2017]), which has particularly been successful at producing state-of-the-arts results on few-shots classification, regression, and reinforcement learning benchmarks, resulting in numerous follow-ups that expand on this elegant framework. In this work we evaluate the MAML algorithm, its first-order approximation (FO-MAML), and the Almost-No-Inner-Loop (ANIL) variant (Raghu et al., [2020]) for learning to predict molecular properties and activities from chemical graphs in low-data domains. Specifically we aim to answer the following questions:

1. Does meta-initializations offer improvements over multitask pre-training in this setting?
2. How little data can meta-initializations learn efficiently from?

Using ChEMBL20 (Bento et al., [2014]), performances of meta-initializations on in- and out-of-distribution tasks are benchmarked with multitask pre-training baselines, showing that they perform favorably across fine-tuning set sizes of $k \in \{16, 32, 64, 128, 256\}$ instances.

2 BACKGROUND

MAML, FO-MAML, and ANIL MAML’s approach is to directly optimize for a set of initial parameters that is efficient at learning from new data. The algorithm consists of an outer loop that learns an initialization $\theta_0$, and an inner loop that adapts to new tasks starting from $\theta_0$. In this setting we have a set of tasks $\{T_1, T_2, ..., T_K\}$ – denoted as $T^{tr}$ – that is available to obtain $\theta_0$, from which we would like to learn the set of tasks $T^{test}$. Following the nomenclature in Finn et al. [2017], we call the process of obtaining the initialization $\theta_0$ meta-training, and the process of adapting to $T^{test}$ meta-testing. More formally, we define a task $T_j$ with $K$ instances as $T = \{(x_i, y_i) | i \in \{1, ..., K\}\}$ which is divided in to a training set $D^{tr}_{T_j}$ and a test set $D^{test}_{T_j}$, also referred to as the support and query set, respectively. The inner loop adaptation to $T_j$ for a function $f$ parameterized by $\theta$ using $N$ gradient descent updates is expressed as

$$\theta^j_n = \theta^j_{n-1} - \alpha \nabla_{\theta} L_{D^{tr}_{T_j}}(f_{\theta_{n-1}})$$

where $\theta^j_n$ are the network’s weights after $n$ steps toward task $T_j$, $\alpha$ is the inner loop learning rate, and $L_{D^{tr}_{T_j}}$ is the loss on the training set of task $T_j$. The loss is calculated using the function $f$ after $n - 1$ updates. The inner loop repeated for a batch of $B$ tasks sampled from $T^{tr}$.

For the outer loop, the meta-loss is defined as the sum of task-specific losses after inner loop updates:

$$L_{meta}(\theta_0) = \sum_{j=1}^{B} L_{D^{test}_{T_j}}(f_{\theta_0})$$

The task-specific loss $L_{D^{test}_{T_j}}$ is calculated on the test set of task $T_j$. We then minimize the meta-loss using stochastic gradient descent to optimize the initialization $\theta_0$, with updates expressed by

$$\theta_0 \leftarrow \theta_0 - \eta \nabla_{\theta} L_{meta}(\theta_0)$$

where $\eta$ is the outer loop learning rate. Intuitively, the meta-loss $L_{meta}(\theta_0)$ measures how well $\theta_0$ adapts to new tasks, and minimizing this loss enables the algorithm to learn good initial parameters.

Updating $\theta_0$ is computationally expensive since it requires the use of second-order derivatives to compute $\nabla_{\theta} L_{meta}(\theta_0)$. FO-MAML sidesteps this problem by omitting the second-order terms, effectively ignoring the inner loop gradients. On the other hand, [Raghu et al., 2020] proposes the ANIL algorithm, which reduces the number of second-order gradients required by limiting inner loop adaptation to only the output layer of the network. ANIL and FO-MAML have both demonstrated significant speedup over MAML.

Graph Neural Networks The graph neural networks framework enables representation learning on graph structured data by learning node-level representations which are aggregated to form graph-level representations. Throughout our experiments, we use a variant of the Gated Graph Neural Networks framework. The Gated Graph Neural Networks (GGNN) model is a type of graph neural networks that introduces a gate mechanism to control the flow of information between nodes. This allows the model to learn more expressive representations for graph-structured data. The model uses message passing between nodes, where each node aggregates information from its neighbors and uses this information to update its own representation. Additionally, the model uses a gate mechanism to control the flow of information, allowing the model to learn more expressive representations for graph-structured data. Throughout our experiments, we use a variant of the Gated Graph Neural Networks framework that includes a gate mechanism to control the flow of information between nodes. This allows the model to learn more expressive representations for graph-structured data.
Table 1: Summary of dataset split

|   | A  | T  | P  | B  | F  |
|---|----|----|----|----|----|
| T_tr | 0  | 0  | 0  | 126 | 737 |
| T_val | 0  | 0  | 0  | 10  | 10  |
| T_test | 1  | 1  | 1  | 10  | 10  |

Network (GGNN) architecture (Li et al., 2017), a member of the message passing neural network (MPNN) family (Gilmer et al., 2017). Similar to other MPNNs, the GGNN architecture operates in two phases: a message passing phase and a readout phase. For an undirected graph $G$ with $V$ nodes where each node has $F$ features, the message passing phase updates the hidden representation of node $v$ at layer $t$ according to

$$m_{v}^{t+1} = A_{e_{vu}} h_{v}^{t} + \sum_{w \in N(v)} A_{e_{vw}} h_{w}^{t}$$

$$h_{v}^{t+1} = \text{GRU}(h_{v}^{t}, m_{v}^{t+1})$$

where $A_{e_{vw}} \in \mathbb{R}^{F \times F}$ is an edge-specific learnable weight matrix, $N(v)$ denotes neighbors of $v$, GRU is the Gated Recurrent Unit (Cho et al., 2014), and $m_{v} \in \mathbb{R}^{F}$ is a message used to update the hidden representation of node $v$ denoted by $h_{v} \in \mathbb{R}^{F}$. The message $m_{v}$ is often interpreted as aggregating information across central and neighboring nodes. A deviation from Li et al. (2017) comes in our choice to remove weight sharing between GRUs in different layers. Following $T$ updates, the readout phase pools node representations according to

$$\hat{y} = \text{MLP}\left(\sum_{v \in G} h_{v}^{T}\right)$$

3 E XPERIMENTAL SETTINGS

ChEMBL20 Dataset We evaluate the effectiveness of meta-initializations for low-resource tasks using a subset of ChEMBL20. More specifically, the dataset processed by Mayr et al. (2018) is filtered so that only tasks with at least 128 instances remain. The resulting dataset contains 902 binary classification tasks from 5 distinct task types denoted in the assay type field: ADME (A), Toxicity (T), Physicochemical (P), Binding (B), and Functional (F).

The dataset is further divided into $T_{tr}$, $T_{val}$, and $T_{test}$. To construct $T_{val}$, we randomly select 10 B and F tasks. We randomly select another set of 10 B and F tasks and all A, T, and P tasks to construct $T_{test}$. The rest of B and F tasks are included in $T_{tr}$ for meta-training. A summary of task types in each set is shown in Table 3. From $T_{test}$, we use B and T tasks to assess in-distribution performance, and A, T, and P tasks for out-of-distribution performance. For the baselines, we combine $T_{tr}$ and $T_{val}$, which is randomly split into $D_{tr}$ baseline for training and $D_{val}$ baseline for early stopping. This setup gives the baselines access to more tasks in during training.

Each molecule is represented as an undirected graph where nodes and edges are atoms and bonds. We use 75 atomic features for each node, as provided by DeepChem [Ramsundar et al., 2019].

Baselines We include the Finetune-All, Finetune-Top, and k-NN baselines as proposed by Triantafillou et al. (2019). To ensure the baselines are competitive, we perform hyperparameter tuning using the Tree-of-Parzen Estimator implementation of Hyperopt (Bergstra et al., 2015) to optimize performance on $D_{val}$ baseline (see Appendix A for details). The resulting architecture has 7 GGNN layers, 1 fully-connected layer with 1024 units, and Dropout applied with a probability of 0.2 at every layer except for the output layer. We use the Adam optimizer (Kingma & Ba, 2017) with a learning rate of $10^{-3.75}$, batch size of 512, and patience of 20 epochs for early stopping during pre-training.
Table 2: Performance on in-distribution tasks measured in AUPRC. The top and bottom halves of the table are tasks with type $B$ and $F$, respectively. Standard deviation are obtained from repeating experiments with five random seeds. The best and second best values are in bold and regular text, respectively. Statistically significant difference from the next best is denoted by (*).

| CheMBL ID | K-NN | Finetune-ALL | Finetune-Top | FO-MAML | ANIL | MAML |
|-----------|------|--------------|-------------|---------|------|------|
| 2363236  | 0.317 ± 0.097 | 0.324 ± 0.026 | 0.324 ± 0.022 | 0.335 ± 0.021 | 0.323 ± 0.008 | 0.333 ± 0.010 |
| 1614469  | 0.458 ± 0.009 | 0.491 ± 0.017 | 0.508 ± 0.021 | 0.491 ± 0.018 | 0.446 ± 0.044 | 0.512 ± 0.029 |
| 2363146  | 0.564 ± 0.014 | 0.649 ± 0.032 | 0.665 ± 0.022 | 0.547 ± 0.012 | 0.596 ± 0.034 | 0.582 ± 0.031 |
| 2363366  | 0.483 ± 0.015 | 0.557 ± 0.024 | 0.535 ± 0.021 | 0.529 ± 0.027 | 0.571 ± 0.030 | 0.587 ± 0.042 |
| 2363553  | 0.741 ± 0.006 | 0.720 ± 0.008 | 0.745 ± 0.021 | 0.697 ± 0.014 | 0.665 ± 0.022 | 0.690 ± 0.011 |
| 1963818  | 0.628 ± 0.037 | 0.744 ± 0.024 | 0.615 ± 0.156 | 0.679 ± 0.032 | 0.692 ± 0.081 | 0.755 ± 0.059 |
| 1963945  | 0.815 ± 0.017 | 0.835 ± 0.023 | 0.817 ± 0.022 | 0.773 ± 0.032 | 0.756 ± 0.035 | 0.827 ± 0.021 |
| 1614423  | 0.481 ± 0.024 | 0.615 ± 0.041 | 0.650 ± 0.049 | 0.767 ± 0.029 | 0.728 ± 0.079 | 0.833 ± 0.040 |
| 2114825  | 0.682 ± 0.012 | 0.751 ± 0.035 | 0.754 ± 0.033 | 0.844 ± 0.032 | 0.761 ± 0.076 | 0.894 ± 0.007* |
| 1964116  | 0.685 ± 0.012 | 0.758 ± 0.043 | 0.762 ± 0.038 | 0.893 ± 0.019 | 0.903 ± 0.019 | 0.912 ± 0.011 |

| Average Rank | 5.4 | 3.3 | 3.4 | 3.2 | 4.1 | 1.7 |

The k-NN baseline uses the activations from the penultimate of the pre-trained neural network to perform classification from 3 nearest neighbors. The Finetune-Top baseline reinitializes and trains a new output layer while the Finetune-All baseline updates weights at every layer in the neural network. In our experiments each baseline is repeated with five random seeds.

| CheMBL ID | K-NN | Finetune-ALL | Finetune-Top | FO-MAML | ANIL | MAML |
|-----------|------|--------------|-------------|---------|------|------|
| 1804798  | 0.319 ± 0.015 | 0.335 ± 0.023 | 0.329 ± 0.010 | 0.367 ± 0.015 | 0.301 ± 0.029 | 0.369 ± 0.021 |
| 2095143  | 0.291 ± 0.015 | 0.167 ± 0.042 | 0.302 ± 0.076 | 0.579 ± 0.024 | 0.501 ± 0.035 | 0.518 ± 0.044 |
| 918053   | 0.308 ± 0.048 | 0.538 ± 0.065 | 0.556 ± 0.050 | 0.884 ± 0.087 | 0.417 ± 0.184 | 0.747 ± 0.076* |

| Average Rank | 5.7 | 4.3 | 3.7 | 2.3 | 3.3 | 1.7 |

The k-NN baseline uses the activations from the penultimate of the pre-trained neural network to perform classification from 3 nearest neighbors. The Finetune-Top baseline reinitializes and trains a new output layer while the Finetune-All baseline updates weights at every layer in the neural network. In our experiments each baseline is repeated with five random seeds.

**Meta-Learning** The same GGNN architecture as the baselines is used for all three meta-learning algorithms. Other hyperparameters are hand-tuned for performance on $T_{val}$. For MAML and ANIL we use an inner loop learning rate of 0.05, 2 inner gradient steps, and inner batch size of 32, while the outer loop has a learning rate of 0.003 and a batch size of 32. FO-MAML uses an outer loop learning rate of 0.0015. We use the Learn2Learn (Arnold et al., 2019) and PyTorch (Paszke et al., 2019) libraries for our implementation and collect performances from five random seeds.

**Evaluation** For task $T_j$ in $T_{test}$, we fine-tune each method on $k$ randomly selected instances from $D_{val}^T$ using the Adam optimizer with learning rate of $10^{-4}$ and batch size of $b = \min(64, k)$. We use $D_{val}^T$ for early stopping with patience of 10 epochs and report the final performances on $D_{T_j}^{test}$.

### 4 Results & Discussions

**Performances on $T_{test}$** The performance of each method on 23 test tasks is reported in Table 2 and 8. Since random splits have been shown to be overly optimistic in scientific applications (Kearnes et al., 2017; Wu et al., 2018), we emphasize relative ranking over absolute performance throughout our benchmark. We observe that meta-initializations generally exhibit similar or better perfor-
Figure 1: Average ranks of each method performance after fine-tuning with \( k \in \{16, 32, 64, 128, 256\} \) instances for in- (left) and out-of-distribution tasks (right). Rankings are based on mean AUPRC measured from five random seeds. MAML and ANIL are consistently ranked as the best method across all \( k \) for in-distribution and out-of-distribution tasks, respectively.

Performances over baselines despite having been trained on fewer tasks. For in-distribution tasks, MAML performs comparably to or outperforms other methods on 16 out of 20 tasks, 3 of which shows significant improvement over the next best method, making it the top performer with an average rank of 1.7. ANIL and FO-MAML, while benefitting from a shorter training time (Appendix B), rank 3.2 and 4.1 on average, respectively. Similar to observations by Triantafillou et al. (2019), Finetune-All and Finetune-Top baselines prove to be strong competitors, both ranking above ANIL in our benchmark. Given their significantly shorter training time, we suspect both baselines to remain crucial in compute-limited settings. In out-of-distribution settings, meta-initializations outperform baselines on all 3 tasks. Again, MAML is ranked as the best method, followed by FO-MAML and ANIL. Overall, compared to the best baselines, meta-initializations learned by MAML provide an average increase in AUPRC of 7.2% for in-distribution tasks and 14.9% out-of-distributions tasks.

Effect of Fine-tuning Set Size From \( T_{test} \), we select all tasks with at least 256 instances in \( D_{tr} \), resulting in 18 tasks available for evaluation (as opposed to 9 when a threshold of 512 instances is used). The average ranking of each method after fine-tuning on \( k \in \{16, 32, 64, 128, 256\} \) instances is reported in Figure 1 (see Appendix C for performance on each task). As the best performing baselines from the previous experiment, Finetune-Top and Finetune-All are selected for comparison. We observe that the baselines benefit greatly from having more data, with Finetune-All rising from fifth to second in in-distribution tasks and Finetune-Top rising from fourth to second in out-of-distribution tasks. Nonetheless, MAML remains the best method, consistently ranked first across fine-tuning set sizes for both sets of tasks.

## 5 Conclusion & Future Directions

In this work, we explore meta-learning as a tool for learning to predict chemical properties and activities in low-resource settings. Emulating this setting using the ChEMBL20 dataset, we demonstrate that GGNN’s initializations learned by MAML perform comparably to or outperform multitask pre-training baselines on 16 out of 20 in-distribution tasks and on all 3 out-of-distribution tasks. Improved performances of meta-initializations are further shown to remain consistent across fine-tuning sets of size \( k \in \{16, 32, 64, 128, 256\} \).

While the ChEMBL20 dataset enables differentiating between in- and out-of-distribution tasks, we recognize that its chemical space is biased towards compounds that have been reviewed and selected for publications. Moreover, our analysis does not include initializations obtained using self- and unsupervised approaches such as those described in Velikovi et al. (2018), Hu et al. (2019), and Sun et al. (2020). We leave experiments with additional datasets and methods to future work. Overall, we believe our contributions open opportunities in applying deep learning to ongoing drug discovery projects where limited data is available.
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Table 4: Wall clock time to train each method

| Method    | Time (Hours) | Speedup |
|-----------|-------------|---------|
| MAML      | 57.9 ± 0.8  | 1×      |
| ANIL      | 48.0 ± 0.6  | 1.2×    |
| FO-MAML   | 27.0 ± 0.9  | 2.1×    |
| PRE-TRAINING | 1.4 ± 0.1  | 41.4×   |

A Baselines Hyperparameter Tuning

Using Hyperopt, we allow a maximum of 50 evaluations and provide the following search space:

- Number of GGNN layers: \{3, 7, 9\}
- Fully connected layer dimension: \{1024, 2048\}
- Batch size: \{128, 256, 512\}
- Learning rate: \(10^{(-4.0, -3.75, -3.5, -3.25)}\)

B Training Time

Training time was measured as the total time required to reach best performance on the \(D_{\text{val}}^{\text{baseline}}\) for baselines and \(T_{\text{val}}^{\text{val}}\) for MAML, FO-MAML, and ANIL on 1 NVIDIA Tesla V100 GPU. We report the recorded times in Table 4. The mean and standard deviation are calculated by repeating the training process with five random seeds.

C Effect of Fine-tuning Set Sizes on Performances

We report the performances used to create Figure [4] below. Figure [2] and [3] show in-distribution tasks, while Figure [4] shows out-of-distribution tasks.
Figure 2: Performances on in-distribution tasks
Figure 3: Performances on in-distribution tasks (continued)
Figure 4: Performances on out-of-distribution tasks