MODELLING BIOLOGICAL ASSAYS WITH ADAPTIVE DEEP KERNEL LEARNING

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ABSTRACT

Due to the significant costs of data generation, many prediction tasks within drug discovery are by nature few-shot regression (FSR) problems, including accurate modelling of biological assays. Although a number of few-shot classification and reinforcement learning methods exist for similar applications, we find relatively few FSR methods meeting the performance standards required for such tasks under real-world constraints. Inspired by deep kernel learning, we develop a novel FSR algorithm that is better suited to these settings. Our algorithm consists of learning a deep network in combination with a kernel function and a differentiable FSR algorithm. As the choice of kernel is critical, our algorithm learns to find the appropriate one for each task during inference. It thus performs more effectively with complex task distributions, outperforming current state-of-the-art algorithms on both toy and novel, real-world benchmarks that we introduce herein. By introducing novel benchmarks derived from biological assays, we hope that the community will progress towards the development of FSR algorithms suitable for use in noisy and uncertain environments such as drug discovery.

1 INTRODUCTION

Following breakthroughs in domains including computer vision, autonomous driving, and natural language processing, deep learning methods are now entering the domain of pharmaceutical R&D. Recent successes include the deconvolution of biological targets from -omics data (Min et al., 2017), generation of drug-like compounds via de novo molecular design (Xu et al., 2019), chemical synthesis planning (Segler and Waller, 2017; Segler et al., 2017), and multi-modal image analysis for quantification of cellular response (Min et al., 2017). A common characteristic of these applications, however, is the availability of high quality, high quantity training data. Unfortunately, many critical prediction tasks in the drug discovery pipeline fail to satisfy these requirements, in part due to resource and cost constraints (Cherkasov et al., 2014).

We therefore focus this work on modelling biological assays (bio-assays) relevant in the early stages of drug discovery, primarily binding and cellular readouts. Under the constraints of an active drug discovery program, the data from these assays, consisting of libraries of molecules and their associated real-valued activity scores, is often relatively small and noisy (refer to statistics in Section 5). In many contexts, it can be a struggle to build a training set of even a few dozen samples per individual assay. Modelling an assay is thus best viewed as a few-shot regression (FSR) problem, with many variables (including experimental conditions, readouts, concentrations, and instrument configurations) accounting for the data distribution generated. Practically, these variables make it infeasible to compare data collected across different assays, thereby making it difficult to learn predictive models from molecular structures. Furthermore, as bio-assay modelling is intended to be used for prioritizing molecules for subsequent evaluation (e.g. Bayesian optimization) and efficiently exploring the overall chemical space (e.g. active learning), accurate prediction and uncertainty estimation using few data is critical to successful application in drug discovery.

It is our view that robust FSR algorithms are needed to tackle this challenge. Specifically, we argue that these algorithms should remain accurate in noisy environments, and also provide well-calibrated uncertainty estimates to inform efficient exploration of chemical space during molecular optimization. Fortunately, recent advances in few-shot learning have led to new algorithms that learn efficiently and generalize adequately from small training data (Wang and Yao, 2019; Chen et al., 2019). Most
have adopted the meta-learning paradigm (Thrun and Pratt, 1998; Vilalta and Drissi, 2002), where some prior knowledge is learned across a large collection of tasks and then transferred to new tasks in which there are limited amounts of data. Such algorithms differ in two aspects: the nature of the meta-knowledge captured and the amount of adaptation performed at test-time for new tasks or datasets. Due to the size of the total chemical space accessible when modelling bio-assays (Bohacek et al., 1996), there is a particular need for the meta-knowledge to be sufficiently rich so as to allow for extrapolation and uncertainty estimation in unseen regions of chemical space at test-time (i.e. for new tasks). Given that the same molecule can behave differently across different assays, greater test-time adaptation is also required and must be accounted for during modelling.

In previous work, metric learning methods (Koch et al., 2015; Vinyals et al., 2016; Snell et al., 2017; Garcia and Bruna, 2017; Bertinetto et al., 2018) accumulate meta-knowledge in high capacity covariance/distance functions and use simple base-learners such as k-nearest neighbor (Snell et al., 2017; Vinyals et al., 2016) or low capacity neural networks (Garcia and Bruna, 2017) to produce adequate models for new tasks. However, they do not adapt the covariance functions nor the base-learners at test-time. Initialization- and optimization-based methods (Finn et al., 2017; Kim et al., 2018; Ravi and Larochelle, 2016) that learn the initialization points and update rules for gradient descent-based algorithms, respectively, allow for improved adaptation on new tasks but remain time consuming and memory inefficient. We therefore argue that to ensure optimal performance when modelling bio-assays, it is crucial to combine the strengths of both types of methods while also allowing for the incorporation of domain-specific knowledge when making predictions. We achieve this by framing FSR as a deep kernel learning (DKL) task, deriving novel algorithms that we apply to modelling specific assays and readouts.

Contributions: Our contributions are several-fold. We first frame few-shot regression as a DKL problem and showcase its advantages relative to classical metric learning methods. We then derive the adaptive deep kernel learning (ADKL) framework by learning a conditional kernel function that is task dependant, allowing for more test-time adaptation than the DKL framework. Finally, we introduce two real-world datasets for modelling biological assays using FSR. With this contribution, we hope to encourage the development of subsequent few-shot regression methods suitable for real-world applications (as is the case for few-shot classification and reinforcement learning, each of which have received comparatively greater attention in recent years (Wang and Yao, 2019)).

2 Deep Kernel Learning

In this section, we describe the DKL framework introduced for single tasks by Wilson et al. (2016). We then extend it to few-shot learning and discuss its advantages over the metric learning framework.

Single Task DKL: Let \( D_{trn}^t = \{ (x_i, y_i) \}_{i=1}^m \subset \mathcal{X} \times \mathbb{R} \), a training dataset available for learning the regression task \( t \) where \( \mathcal{X} \) is the input space and \( \mathbb{R} \) is the output space. A DKL algorithm aims to obtain a non-linear embedding of inputs in the embedding space \( \mathcal{H} \), using a deep neural network \( \phi_\theta : \mathcal{X} \rightarrow \mathcal{H} \) of parameters \( \theta \). It then finds the minimal norm regressor \( h^*_t \) in the reproducing kernel Hilbert space (RKHS) \( \mathcal{R} \) on \( \mathcal{H} \), that fits the training data, i.e.:

\[
\begin{align*}
\hat{h}^*_t & := \arg \min_{h \in \mathcal{R}} \| h \|_\mathcal{R} + \ell(h, D_{trn}^t) \quad (1)
\end{align*}
\]

where \( \ell \) is a non-negative loss function that measures the loss of a regressor \( h \) and \( \lambda \) weighs the importance of the norm minimization against the training loss. Following the representer theorem (Scholkopf and Smola, 2001; Steinwart and Christmann, 2008), \( h^*_t \) can be written as a finite linear combination of kernel evaluations on training inputs, i.e.:

\[
\hat{h}^*_t(x) = \sum_{(x_i, y_i) \in D_{trn}^t} \alpha^t_i k_\rho(\phi_\theta(x), \phi_\theta(x_i)), \quad (2)
\]

where \( \alpha^t = (\alpha^t_1, \ldots, \alpha^t_m) \) are the combination weights and \( k_\rho : \mathcal{H} \times \mathcal{H} \rightarrow \mathbb{R}^+ \) is a reproducing kernel of \( \mathcal{R} \) with hyperparameters \( \rho \). Candidates include the radial basis, polynomial, and linear kernels. Depending on the loss function \( \ell \), the weights \( \alpha^t \) can be obtained by using a differentiable kernel method enabling the computation of the gradients of the loss w.r.t. the parameters \( \theta \). Such methods include Gaussian Process (GP), Kernel Ridge Regression (KRR), and Logistic Regression (LR).
As DKL inherits from deep learning and kernel methods, it follows that gradient descent algorithms are required to optimize $\theta$, which can be high dimensional such that seeing a significant amount of training samples is essential to avoid overfitting. However, once the latter condition is met, scalability of the kernel method becomes limiting as the running time of kernel methods scales approximately in $O(m^3)$ for a training set of $m$ samples. Some approximations of the kernel are thus needed for the scalability of the DKL method (see Williams and Seeger (2001); Wilson and Nickisch (2015)).

**Few-Shot DKL:** In the setup of episodic meta learning, also known as few-shot learning, one has access to a meta-training collection $\mathcal{D}_{meta-trn} := \{(D_{trn}^1, D_{val}^1)\}_{j=1}^T$ of $T$ tasks to learn how to learn from few datapoints. Each task $t_j$ has its own training (or support) set $D_{trn}^j$ and validation (or query) set $D_{val}^j$. A meta-testing collection $\mathcal{D}_{meta-tst}$ is also available to assess the generalization performance of the few-shot algorithm across unseen tasks. To obtain a Few-Shot DKL (FSDKL) method for FSR in such settings, one can share the parameters of $\phi_\theta$ across all tasks, similar to metric learning algorithms. Hence, for a given task $t_j$, the inputs are first transformed by the function $\phi_\theta$ and then a kernel method is used to obtain the regressor $h_{trn}^j$, which will be evaluated on $D_{val}^j$. Here, KRR and GP are explored as they are the state-of-the-art algorithms for kernel-based regression. The latter is used to allow our models to provide accurate predictive uncertainty, which is useful when modelling biological assays.

**KRR:** Using the squared loss and the L2-norm to compute $\|\hat{h}\|_2$, KRR gives the optimal regressor for a task $t$ and its validation loss $\mathcal{L}_{\theta,p,\lambda}^t$ as follows:

$$h_{trn}^t(x) = \alpha K_{x,trn}, \quad \text{with} \quad \alpha = (K_{trn,trn} + \lambda I)^{-1} y_{trn}$$

$$\mathcal{L}_{\theta,p,\lambda}^t = \mathbb{E}_{x,y \sim D_{val}^j} (\alpha K_{x,trn} - y)^2,$$

where $y_{trn} = (y_1, \cdots, y_{|D_{trn}^j|})^T$, $K_{trn,trn}$ is the matrix of kernel evaluations and each entry is $k_\rho(\phi_\theta(x_i), \phi_\theta(x_i))$ for $(x_i, \cdot), (x_i, \cdot) \in D_{trn}^j$. An equivalent definition applies to $K_{x,trn}$.

**GP:** Using the negative log likelihood loss function instead, the GP algorithm gives a probabilistic regressor for which the predictive mean, covariance, and loss for a task $t$ are:

$$\mathcal{L}_{\theta,p,\lambda}^t = - \ln \mathcal{N}(y_{val}; \mathbb{E}[h_{trn}^t], \cov(h_{trn}^t)), \quad \mathbb{E}[h_{trn}^t] = K_{val,trn}(K_{trn,trn} + \lambda I)^{-1} y_{trn}, \quad \cov(h_{trn}^t) = K_{val,val} - K_{val,trn}(K_{trn,trn} + \lambda I)^{-1} K_{trn,val}$$

Finally, the parameters $\theta$ of the neural network, along with $\lambda$ and the kernel hyperparameters $\rho$, are optimized using the expected loss on all tasks:

$$\arg\min_{\theta,p,\lambda} \mathbb{E}_{D_{meta-trn}} \mathcal{L}_{\theta,p,\lambda}^t.$$

To summarize, FSDKL finds a representation common to all tasks such that the kernel method (in our case, GP and KRR) will generalize well from a small amount of samples. In doing so, this alleviates two of the main limitations of single task DKL: i) the scalability of the kernel method is no longer an issue since we are in the few-shot learning regime\(^1\), and ii) the parameters $\theta$ (and $\rho, \lambda$) are learned across a potentially large amount of tasks and samples, providing the opportunity to learn a rich representation without overfitting.

Despite shared characteristics with the metric learning framework, the FSDKL framework is more powerful and flexible. It provides better task-specific adaptation due to the inference of the appropriate model using the kernel methods compared to shared model parameters in metric learning. After meta-training, any task-specific model also inherits the generalization guarantees of kernel-based models, and consequently increasing the number of shots for new tasks can only improve generalization performance. The incorporation of prior knowledge through user-specific kernel functions is also a major advantage of DKL over metric learning (e.g. use periodic kernels for periodic function regression tasks).

\(^1\)Even with several hundred samples, the computational cost of embedding each example is usually higher than inverting the Gram matrix.
3 ADAPTIVE DEEP KERNEL LEARNING

FSDKL uses a shared deep kernel function, in which the base kernel $k_\rho$ is user-chosen (although its parameters $\rho$ are learned during meta-training). Given that the choice of the kernel function dictates almost all the generalization properties of any kernel-based method, it may not be optimal to leave it to the user. In addition, for modelling bio-assays, it is not straightforward how to best incorporate task-specific prior knowledge in this shared and user-chosen kernel. We overcome these limitations by developing the Adaptive Deep Kernel Learning (ADKL) framework, illustrated by Fig. 1.

ADKL also aims to obtain a non-linear embedding of inputs using a deep neural network $\phi_\theta$ shared by all tasks before finding the minimal norm task-specific regressor $h^*_{t}$ using either GP or KRR as described in Section 2 (ADKL-GP and ADKL-KRR will refer to our algorithm when using GP and KRR, respectively). The fundamental difference between FSDKL and ADKL lies in the kernel definition, which brings significantly more flexibility in the latter case relative to the former. Specifically, during the meta-training, ADKL learns to learn task-specific kernel functions instead of using one chosen by the user. It does so by learning how to represent tasks with the task encoding network $\psi_\eta$ and then how to leverage task embeddings to build task-specific kernels using a multi-modal neural network $c_\rho$. Given a task $t$, ADKL thus first computes its embedding $z_t = \psi_\eta(D_{trn}^t)$ using its support set $D_{trn}^t$, and deduces the adapted kernel with $c_\rho$. We describe in more detail both the task encoding network $\psi_\eta$ and the network $c_\rho$ responsible for computing the task-specific kernel below.

3.1 TASK ENCODING

The challenge of the $\psi_\eta$ network is to capture complex dependencies in the training set $D_{trn}^t$ to provide a useful task encoding $z$. Furthermore, the task encoder should be invariant to permutations of the training set and be able to encode a variable amount of samples. After exploring a variety of architectures, we found that those that are more complex, such as Transformers (Vaswani et al., 2017), tend to underperform. This is possibly due to overfitting or the sensitivity of training such architectures.

Consequently, inspired by DeepSets (Zaheer et al., 2017), we propose the following order invariant network. It begins its computations by representing each input-target pair $xy_i = r(Concat(\phi(x_i), v(y_i)))$ for all $(x_i, y_i) \in D_{trn}^t$, using neural networks $\phi, v,$ and $r$. The $xy_i$ captures nonlinear interactions between the inputs and the targets if $r$ is a nonlinear transformation. Then, by computing $\mu^t_{xy}$ and $\sigma^t_{xy}$, the empirical mean and standard deviation of the set $\{xy_1, xy_2, \ldots, xy_m\}$, respectively, we obtain the task representation as follows:

$$z_t = \psi_\eta(D_{trn}^t) := [\mu^t_{xy}, \sigma^t_{xy}] .$$

As $\mu^t_{xy}$ and $\sigma^t_{xy}$ are invariant to permutations in $D_{trn}^t$, it follows that $\psi_\eta$ is also permutation invariant. Overall, $\psi_\eta$ is simply the concatenation of the first and second moments of the sample representations,
which were nonlinear transformations of the original inputs and targets. The learnable parameters $\eta$ of the task encoder include all the parameters of the networks $v$ and $r$, and are shared across all tasks.

To help the training of these parameters, we maximize the mutual information between $D^t_{\text{trn}}$ and $D^t_{\text{val}}$, i.e., we expect and encourage the network to produce similar task encodings using any data partitions for a given task. More explicitly, we use the MINE algorithm (Belghazi et al., 2018), which optimizes a lower bound on the mutual information. For two random variables $r, s \sim p(r, s)$ and a similarity measure $f_\phi$ between $r$ and $s$, parameterized by $\phi$, the following inequality holds:

$$ I[r, s] \geq \max_{\phi} \mathbb{E}_{r, s \sim p(r, s)} f_\phi(r, s) - \ln \mathbb{E}_{r \sim p(r)} \mathbb{E}_{s \sim p(s)} e^{f_\phi(r, s)}. $$

(10)

Using a batch of $b$ tasks, and the cosine similarity $c$ as the similarity measure between two task encodings, one obtains and maximizes $I[D_{\text{trn}}, D_{\text{val}}] \geq \tilde{I}_\eta$, where:

$$ \tilde{I}_\eta \overset{\text{def}}{=} \frac{1}{b} \sum_{j=1}^b c(\psi_\eta(D^t_{\text{trn}}), \psi_\eta(D^t_{\text{val}})) - \ln \frac{1}{b(b-1)} \sum_{j=1}^b \sum_{i \neq j} e^{c(\psi_\eta(D^t_{\text{trn}}), \psi_\eta(D^t_{\text{val}}))}. $$

(11)

3.2 Task-Specific Kernel

Let $\mathcal{H}$ and $\mathcal{Z}$ be the output domains of $\phi_\rho$ and $\psi_\eta$, respectively. We define a pairwise function on $\mathcal{H}$, whose outputs are dependent from the task representations in $\mathcal{Z}$ as follows:

$$ c_\rho : \mathcal{H} \times \mathcal{H} \times \mathcal{Z} \to \mathbb{R} $$

$$(\phi_\rho, \phi_\rho', z_t) \mapsto \text{MLP}_\rho\left((\phi_\rho - \phi_\rho')^2, z_t\right),$$

(12)

where $[\cdot, \cdot]$ is the concatenation operator. It bears mentioning that the parameters $\rho$ are shared across all tasks and learned during the meta-training. Also, $c_\rho$ is symmetric and stationary with regard to its inputs ($\phi_\rho$ and $\phi_\rho'$) as their element-wise L2 distances vector is received as input of the fully connected network. Further, by simply concatenating the task representation $z_t$ to this distance vector at the input, $c_\rho$ provides a powerful approach to producing task-specific kernels. However, these kernels are not positive semi-definite (PSD) and cannot be directly used for KRR and GP.

Therefore, without losing any information given by $c_\rho$, we compute task-specific PSD kernels $k_{p, t}$ as the empirical kernel maps with regard to the support set inputs, i.e.:

$$ k_{p, t}(x, x') = C_1(x) \cdot C_1(x'), $$

$$ C_1(x) = (c_\rho(x, x_1, z_1), \cdots, c_\rho(x, x_m, z_1)), \quad \text{and} \quad (x_i, \cdot) \in D^t_{\text{trn}} \forall i = 1, \cdots, m $$

(13)

Using the empirical kernel map of $c_\rho$ to compute $k_{p, t}$ offers the opportunity to improve the kernel evaluations in low data settings using some unlabelled data. More precisely, instead of computing the empirical kernel map with regard to only $D^t_{\text{trn}}$, we could use $(D^t_{\text{trn}} \cup U)$ where $U$ is a set of unlabelled inputs. However, to avoid a significant increase in the computation costs of the PSD kernels, $|U|$ should be kept relatively small (in our experiments $|U| \leq 50$). One must also be careful about the composition of $U$ to avoid overfitting certain tasks and under-fitting others. Therefore, instead of asking the user to provide the set $U$, we propose directly learning them through back-propagation. To do so, we introduce pseudo-input representations (or pseudo-representations) $u_l \in \mathcal{H}$ that are shared by all tasks and learned during meta-training. The function $C_1$, from Eq. (13), becomes:

$$ C_1(x) = (c_\rho(x, x_1, z_1), \cdots, c_\rho(x, x_m, z_1), c_\rho(x, u_1, z_t), \cdots, c_\rho(x, u_l, z_t)), $$

$$ \text{with} \quad u_l \in U \forall l = 1, \cdots, |U| \quad \text{and} \quad (x_i, \cdot) \in D^t_{\text{trn}} \forall i = 1, \cdots, m $$

(14)

These pseudo-representations can be thought of as parameters of the adaptive kernel and the number to be included is a hyperparameter of the algorithm. To prevent their collapse into a single point and ensure that they are well distributed in the feature space $\mathcal{H}$, we add a regularization term to the training loss. Let $p$ and $q$ be the distributions that generate the true input representations and the pseudo-input representations, respectively. We make the assumption that $p$ and $q$ are both multivariate Gaussian distributions with diagonal covariance matrices and have respective parameters $(\mu_\phi, \sigma_\phi^2)$.

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2This yields a small bias on the gradient since the right hand side takes the log of the expectations. Since we are not interested in the precise value of the mutual information, this does not constitute a problem.
and \((\mu_u, \sigma_u^2)\). The parameters of \(p\) are estimated using the running means and variances of all input representations computed over batches of tasks. Those of \(q\) are estimated using \(U\). The training of the pseudo-input representations is then regularized by minimizing the KL distance \(\hat{D}_u\) between \(p\) and \(q\), i.e.:

\[
\hat{D}_u = KL(N(\mu_u, \sigma_u^2) \parallel N(\mu_\phi, \sigma_\phi^2))
\]  

Putting it all together, the ADKL training objective is the following:

\[
\arg \min_{\theta, \eta, p, u, \lambda} E_{t_j \sim B} \mathcal{L}_{\theta, \eta, p, u, \lambda}^{(t_j)} - \gamma_{\text{task}} \hat{L}_{\eta} + \gamma_{\text{pseudo}} \hat{D}_u,
\]

with \(\gamma_{\text{task}} \geq 0\) as a tradeoff hyperparameter for the regularization of the task-encoder, \(\gamma_{\text{pseudo}} \geq 0\) as a tradeoff hyperparameter for the regularization of the pseudo-inputs.

4 RELATED WORK

Across the spectrum of learning approaches, DKL methods lie between neural networks and kernel methods. While neural networks can learn from a very large amount of data without much prior knowledge, kernel methods learn from fewer data when given an appropriate covariance function that accounts for prior knowledge of the relevant task. In the first DKL attempt, Wilson et al. (2016) combined GP with CNN to learn a covariance function adapted to a task from large amounts of data, though the large time and space complexity of kernel methods forced the approximation of the exact kernel using KISS-GP (Wilson and Nickisch, 2015). Dasgupta et al. (2018) have demonstrated that such approximation is not necessary using finite rank kernels. Here, we show that learning from a collection of tasks (FSR mode) does not require any approximation when the covariance function is shared across tasks. This is an important distinction between our study and other existing studies in DKL, which learn their kernel for single task applications instead of multiple task collections.

On the spectrum between NNs and kernel methods we must also mention metric learning. Metric learning algorithms learn an input covariance function shared across tasks but rely only on the expressive power of DNNs. First, stochastic kernels are built out of shared feature extractors and simple pairwise metrics (e.g. cosine similarity (Vinyals et al., 2016), Euclidean distance (Snell et al., 2017)), or parametric functions (e.g. relation modules (Sung et al., 2018), graph neural networks (Garcia and Bruna, 2017; Kim et al., 2019a)). Then, within tasks, the predictions are distance-weighted combinations of the training labels with the stochastic kernel evaluations—no adaptation is done.

In connection with the test-time adaptation capabilities of our method, methods that combine metric learning with initialization-based models are great competitors. In fact, Proto-MAML (Triantaflou et al., 2019), which captures the best of Prototypical Networks (Snell et al., 2017) and MAML (Finn et al., 2017), allows within-task adaptation using MAML on top of a shared feature extractor. Similarly, Kim et al. (2018) have proposed a Bayesian version of MAML where a feature extractor is shared across tasks, while multiple MAML particles are used for the task-level adaptation. Bertinetto et al. (2018) have also tackled the lack of adaptation for new tasks by using Ridge Regression and Logistic Regression to find the appropriate weighting of the training samples for classification tasks. This study can be considered as an instance of the FSDKL framework, though its contribution was limited to showing that simple differentiable learning algorithms can increase adaptation in the metric learning framework. Our work goes beyond by formalizing few-shot DKL and proposing ADKL: a data-driven manner for computing the correct kernel for a task.

Since ADKL-GP learns task-specific stochastic processes, it is related to neural processes (Garnelo et al., 2018a) and the ML-PIP framework (Gordon et al., 2018). Both propose a scalable alternative to learning regression functions by performing inference on stochastic processes. In these families of methods, both Conditional Neural Processes (CNP) (Garnelo et al., 2018b) and Attentive Neural Processes (ANP) (Kim et al., 2019b) learn conditional stochastic processes parameterized by task-specific conditions derived from the support sets, but CNP is the most related to ADKL-GP. CNP is an instance of ML-PIP when the task encoder gives a point estimate of the task parameters instead of a distribution. Finally, the main differences between ANP and CNP are the architecture of the task-encoder and the lack of mathematical guarantees associated with stochastic processes in CNP (as it does not impose any consistency with respect to a prior process). By comparison, ADKL-GP
also learns conditional stochastic processes but has mathematical guarantees thanks to GP and PSD kernels.

5 Datasets

Existing FSR methods have been mostly tested on 1D function regression and pixel-wise image completion tasks with MNIST and CelebA (Kim et al., 2018; Garnelo et al., 2018b,a). On one hand, the 1D regression tasks are all relatively simple, almost noise-less, and homogeneous. On the other hand, methods have been successful for image completion tasks only outside the few-shot regime (i.e. when the number of samples is greater than 500) (Garnelo et al., 2018b,a). For these reasons, we introduce two task collections from a real-world context. Deemed Binding and Antibacterial, these task collections contain data from bio-assays that are representative of real-world FSR tasks in drug discovery. The pre-processed versions of these collections and detailed statistics are available here (anonymized link).

Binding: All tasks in this collection aim to predict the binding affinity of small molecules to a target protein. The characteristics of the proteins thus define different data distributions over the chemical space. The inputs and the targets for each task are molecules that have been tested in a binding assay and the measured binding affinity of the molecule against a given protein. The task collection was extracted from the public database BindingDB and altered by removing bio-assays with correlations above 0.8 or those with less than 10 experimental measurements, leaving us with 5,717 tasks.

Antibacterial: Within this collection, the task is to predict the antimicrobial activity of small molecules against various bacteria. They are characterized by a bacterial strain whose resistance to drug-like molecules was being evaluated. The task collection was extracted from the public database PubChem. After also removing bio-assays with correlations above 0.8 and those with less than 10 samples, we obtain 3,255 tasks.

Their meta-test partitions each contain 500 tasks, with the remaining used in the meta-train and meta-validation. The molecules (represented as SMILES) are converted into vectors using routines available in the RDKit software (more precisely into ECFP6 binary fingerprint vectors of 4,096 dimensions). These inputs were also processed in all methods using the same feature extractor architecture, which is a fully-connected network of $256 \times 256 \times 256$. Due to the high noise-to-signal ratio, the targets are first log2-scaled and then scaled linearly between 0 and 1 to avoid scaling issues during training.

Fig. 2 highlights three aspects of the collections that make them better benchmarks for evaluating the readiness of FSR methods for real-world applications relative to toy collections. First, the distributions of number of samples per task show that they naturally contain few samples, which we believe reflects the costs of acquiring labelled data in a drug discovery setting. In comparison, the number of samples available per task is relatively large in previous benchmarks, with the few-shot regime being achieved artificially through sampling. Second, as illustrated by their noise-to-signal ratio, real-world tasks are inherently noisy, increasing the difficulties associated with few-shot learning. Finally, the input diversity within each task is reduced relative to the total among tasks. Despite this diversity difference, good models should perform relatively well outside the input region they have seen in the support set. This situation challenges the methods to learn strong priors about the input space and to be able to generalize after seeing only a small fraction of it. These collections invite researchers to explore meta-learning with increasingly heterogeneous datasets and in noisy environments, as well as generalisation and extrapolation in large input spaces (such as the drug-like chemical space, which is estimated to be approximately $10^{33}$ molecules (Polishchuk et al., 2013)).

To test our method in a noise-less environment, we also use the Sinusoids collection introduced by Kim et al. (2018). This challenging few-shot regression benchmark consists of 5,000 tasks defined by functions of the form: $y = A \sin(wx + b) + \epsilon$ with $A \in [0.1, 5.0]$, $b \in [0.0, 2\pi]$, and $w \in [0.5, 2.0]$. Sampling inputs $x \in [-5.0, 5.0]$ and observational noise $\epsilon \in N(0, (0.01A)^2)$ and computing $y$ gives the samples for each task. Here, the meta-train, meta-validation, and meta-test contain 56.25%, 18.75% and 25% of all the tasks, respectively, and all methods use the same feature extractor architecture, which is a fully-connected network of $40 \times 40 \times 40$. 
6 EXPERIMENTS

6.1 Benchmarking Analysis

Performance of ADKL is evaluated against a FSDKL instance (R2-D2 of Bertinetto et al. (2018)), CNP (Garnelo et al., 2018b), MAML (Finn et al., 2017), BMAML (Kim et al., 2018), ProtoMAML (Triantafillou et al., 2019) and Learned Basis (Yi Loo, 2019) (all implementations are available here (anonymized link)). These algorithms have all proven to have efficient and effective test-time adaptation routines and therefore constitute strong baselines for benchmarking. Tables 1 to 3 report the average MSE over all tasks and 20 random support and query partitions for each task, for different sized support sets.

For the Sinusoids collection, we observe that DKL-based methods significantly outperform all other methods despite their test-time adaptation capabilities. These results alone demonstrate the effectiveness of DKL-based methods in FSR relative to the current state-of-the-art. Furthermore, of all DKL-based methods, ADKL-KRR shows consistently stronger performance than others. This demonstrates that using ADKL increases test-time performance relative to FS-DKL (as R2-D2 and ADKL-KRR only differ by the kernel definition). It also indicates that attempting to capture the model uncertainty using GP in ADKL (instead of KRR) comes with a significant cost, especially in lower data regimes. This may be due to the inability of GP to differentiate between the observational noise and the model uncertainty as the number of samples get smaller. It is also important to notice that all methods using the task representation significantly outperform those that do not. This shows that adequately capturing the task representation is crucial for this task collection, which ADKL-KRR appears to be well-equipped to handle.

Tables 2 and 3 show that real-world datasets are challenging for most methods, as their MSE only marginally improves when the size of the support set increases. It should be noted that while the scaling of the targets makes the MSE low for all methods, even a decrease of 0.005 can translate into large improvements of the modelling accuracy. However, we still observe that ADKL-KRR outperforms all other methods when the number of samples is greater than 10, again providing evidence of effectiveness of our method for FSR with complex task distributions. The gaps between ADKL and R2-D2 for these collections also confirm that using task specific kernels can be very useful even though inferring the right kernel can become difficult as the size of the support set gets smaller. Finally, it also appears that estimating the model uncertainty using ADKL-GP instead of ADKL-KRR comes with a marginal accuracy cost.

6.2 Active Learning

In this section, we report the results of active learning experiments. Our intent is to measure the effectiveness of the uncertainty captured by the predictive distribution of ADKL-GP for active
learning, as it is critical to our drug discovery use-cases. CNP, in comparison, serves to measure which of CNP and GP better captures the data uncertainty for improving FSR under active sample selection. For this purpose, we meta-train both algorithms using support and query sets of size $m = 5$. During meta-test time, five samples are randomly selected to constitute the support set $D_{trn}$ and build the initial hypothesis for each task. Then, from a pool $U$ of unlabeled data, we choose the input $x^*$ of maximum predictive entropy, i.e. $x^* = \arg\max_{x \in U} \mathbb{E} \left[ \log p(y|x, D_{trn}) \right]$. The latter is removed from $U$ and added to $D_{trn}$ with its predicted label. The within-task adaptation is performed on the new support set to obtain a new hypothesis which is evaluated on the query set $D_{val}$ of the task. This process is repeated until we reach the allotted budget of 20 queries.

Fig. 3 illustrates, for all collections, the MSE after each sample acquisition iteration and under both random and active learning acquisition strategies. Under the active learning strategy, ADKL-GP consistently outperforms CNP. In particular, we observe that very few samples are queried by ADKL-GP to capture the data distribution whereas CNP performance remains far from optimal even when allowed the maximum number of queries. Further, since using the maximum predictive entropy strategy is better than querying samples at random for ADKL-GP (solid vs. dashed line), these results suggest that the predictive uncertainty obtained with GP is informative and more accurate than that of CNP. Moreover, when the number of queries is greater than 10, we observe a performance degradation for CNP while ADKL-GP remains consistent. This observation highlights the generalization capacity of DKL methods, even outside the few-shot regime where they have been trained — this same property does not hold true for CNP. We attribute this property of DKL methods, even outside the few-shot regime where they have been trained — this same property does not hold true for CNP. We attribute this property of DKL methods to their use of kernel methods. In fact, their role in adaptation and generalization increases as we move away from the few-shot training regime.

![Figure 3: Average MSE performance on the meta-test during active learning. The width of the shaded regions denotes the uncertainty over five runs for the sinusoidal collection. No uncertainty is shown for the real-world tasks as they were too time consuming.](image)

### Table 1: Average MSE on Sinusoids

<table>
<thead>
<tr>
<th>m model</th>
<th>5</th>
<th>10</th>
<th>20</th>
</tr>
</thead>
<tbody>
<tr>
<td>BMAML</td>
<td>2.042</td>
<td>1.371</td>
<td>0.844</td>
</tr>
<tr>
<td>CNP</td>
<td>1.616</td>
<td>0.392</td>
<td>0.117</td>
</tr>
<tr>
<td>Learned Basis</td>
<td>3.587</td>
<td>0.800</td>
<td>0.127</td>
</tr>
<tr>
<td>MAML</td>
<td>2.896</td>
<td>1.634</td>
<td>0.901</td>
</tr>
<tr>
<td>ADKL-GP</td>
<td>1.178</td>
<td>0.084</td>
<td>0.007</td>
</tr>
<tr>
<td>ADKL-KRR</td>
<td>0.867</td>
<td>0.061</td>
<td>0.005</td>
</tr>
<tr>
<td>ProtoMAML</td>
<td>2.044</td>
<td>1.369</td>
<td>0.846</td>
</tr>
<tr>
<td>FSDKL(R2D2)</td>
<td>1.002</td>
<td>0.073</td>
<td>0.009</td>
</tr>
</tbody>
</table>

### Table 2: Average MSE on Binding

<table>
<thead>
<tr>
<th>m model</th>
<th>5</th>
<th>10</th>
<th>20</th>
</tr>
</thead>
<tbody>
<tr>
<td>BMAML</td>
<td>0.061</td>
<td>0.059</td>
<td>0.057</td>
</tr>
<tr>
<td>CNP</td>
<td>0.064</td>
<td>0.062</td>
<td>0.061</td>
</tr>
<tr>
<td>Learned Basis</td>
<td>0.063</td>
<td>0.060</td>
<td>0.059</td>
</tr>
<tr>
<td>MAML</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>ADKL-GP</td>
<td>0.064</td>
<td>0.056</td>
<td>0.051</td>
</tr>
<tr>
<td>ADKL-KRR</td>
<td>0.063</td>
<td>0.054</td>
<td>0.051</td>
</tr>
<tr>
<td>ProtoMAML</td>
<td>0.061</td>
<td>0.059</td>
<td>0.065</td>
</tr>
<tr>
<td>FSDKL(R2D2)</td>
<td>0.060</td>
<td>0.060</td>
<td>0.055</td>
</tr>
</tbody>
</table>

### Table 3: Average MSE on Antibacterial

<table>
<thead>
<tr>
<th>m model</th>
<th>5</th>
<th>10</th>
<th>20</th>
</tr>
</thead>
<tbody>
<tr>
<td>BMAML</td>
<td>0.067</td>
<td>0.060</td>
<td>0.060</td>
</tr>
<tr>
<td>CNP</td>
<td>0.070</td>
<td>0.069</td>
<td>0.068</td>
</tr>
<tr>
<td>Learned Basis</td>
<td>0.068</td>
<td>0.065</td>
<td>0.093</td>
</tr>
<tr>
<td>MAML</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>ADKL-GP</td>
<td>0.068</td>
<td>0.064</td>
<td>0.060</td>
</tr>
<tr>
<td>ADKL-KRR</td>
<td>0.068</td>
<td>0.059</td>
<td>0.058</td>
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<tr>
<td>ProtoMAML</td>
<td>0.065</td>
<td>0.063</td>
<td>0.070</td>
</tr>
<tr>
<td>FSDKL(R2D2)</td>
<td>0.066</td>
<td>0.064</td>
<td>0.063</td>
</tr>
</tbody>
</table>

6.3 ABLATION EXPERIMENTS

In our final set of experiments, we more closely evaluate the impact of the task encoder and the pseudo-inputs on the generalization during meta-testing. We do so by training and evaluating ADKL on Sinusoids with different hyperparameter combinations. Figs. 4a to 4d show the relative improvements (negative values) or setbacks (positive values) in the meta-test MSE compared to different baselines (but the joint impact of $\gamma_{task}$ and $\gamma_{pseudo}$ is only discussed in Appendix A.3).
First, Fig. 4a compares $\gamma_{\text{task}} \in \{0.01, 0.1\}$ relative to $\gamma_{\text{task}} = 0$ and consequently demonstrates that regularizing the task encoder by maximizing the mutual information between the support set and the query set significantly improves the generalization performance. This conclusion holds for all support set sizes tested, as shown in Appendix A.1. Combined with the results from Section 6.1, this figure shows the importance of good task encoders for generalization in few-shot learning and how using the regularization term that we introduced is a step forward in that direction.

Then, Fig. 4c measures the relative differences between $\gamma_{\text{pseudo}} \in \{0.01, 0.1\}$ and $\gamma_{\text{pseudo}} = 0$ for different values of hyperparameter combinations. It shows that improving the kernel map evaluations using pseudo-input representations can significantly help with the generalization performance of ADKL. This conclusion also holds for all values tested for $|D_{\text{trn}}|$ (see Appendix A.2). However, the improvements were more consistent for smaller support sets, which is not surprising as improving the kernel map estimations in these cases is more critical.

Finally, Figs. 4b and 4d illustrate for ADKL-GP and ADKL-KRR, and different sizes of support sets, how the number of pseudo-representations (i.e, $|U|$) affects performance. The values for each cell are relative performance using $|U| \in \{20, 50\}$ versus $|U| = 0$ and have been averaged over different hyperparameters and $\gamma_{\text{pseudo}}$. In general, we can confirm that increasing the number of pseudo-representations increases the estimates of the kernel maps and improves generalization. However, the improvements are more prominent with KRR in comparison to GP, which may be due to the fact that GP attributes a part of the modelling noise to the kernel evaluations, leading to more constraints on the optimization of the pseudo-representation parameters.

7 CONCLUSION

In this work, we investigate the modelling of biological assays using few-shot learning methods. We propose a new framework, ADKL, that stores meta-knowledge in kernel functions and adapts to new tasks using KRR or GP. Our experiments provide evidence that the additional adaptation capacity at test-time provided by these methods increases generalization when modelling bio-assay and on 1D sinusoidal regression tasks. In a Bayesian setup, they better estimate predictive uncertainty, increasing their utility in real-world applications such as drug discovery. Finally, by making our bio-assay task collections publicly available, we hope that the community will leverage them to propose FSR algorithms that are ready to be deployed under real-world constraints, with the ultimate aim of accurately predicting key molecular properties early in the drug discovery pipeline.
REFERENCES


Appendices

A Regularization Impact

A.1 Task Regularization

Table 4 presents the hyperparameter combinations used in the experiments to assess the impact of the trade-off parameter $\gamma_{\text{task}}$. We report the MSE performance obtained on the meta-test for each combination. To make reading this table easier, we also repeat the Fig. 5 showing the improvement of the MSE relative to $\gamma_{\text{task}} = 0$ (no regularization).

Table 4: Effect of using task regularization (parameter $\gamma_{\text{task}}$) on the MSE performance

<table>
<thead>
<tr>
<th>Algorithm</th>
<th>K</th>
<th>$\gamma_{\text{pseudo}}$</th>
<th>$\gamma_{\text{task}}$</th>
<th>0.00</th>
<th>0.01</th>
<th>0.10</th>
</tr>
</thead>
<tbody>
<tr>
<td>ADKL-KRR</td>
<td>20</td>
<td>0.01</td>
<td>1</td>
<td>0.0585</td>
<td>0.0327</td>
<td><strong>0.0289</strong></td>
</tr>
<tr>
<td></td>
<td>10</td>
<td>0.00</td>
<td>2</td>
<td>0.4051</td>
<td><strong>0.2944</strong></td>
<td>0.3671</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>0.10</td>
<td>0.4363</td>
<td>0.2964</td>
<td><strong>0.2882</strong></td>
</tr>
<tr>
<td>ADKL-GP</td>
<td>5</td>
<td>0.10</td>
<td>4</td>
<td>2.4920</td>
<td><strong>2.2511</strong></td>
<td>2.2994</td>
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<tr>
<td>ADKL-KRR</td>
<td>20</td>
<td>0.00</td>
<td>5</td>
<td>0.0574</td>
<td>0.0305</td>
<td><strong>0.0302</strong></td>
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<tr>
<td>ADKL-GP</td>
<td>5</td>
<td>0.01</td>
<td>6</td>
<td>2.5611</td>
<td>2.1511</td>
<td><strong>2.2112</strong></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>0.01</td>
<td>3.2933</td>
<td><strong>2.7663</strong></td>
<td>3.0971</td>
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<tr>
<td></td>
<td>10</td>
<td>0.01</td>
<td>8</td>
<td>0.7675</td>
<td>0.7105</td>
<td><strong>0.4352</strong></td>
</tr>
<tr>
<td></td>
<td>20</td>
<td>0.00</td>
<td>9</td>
<td>1.201</td>
<td>0.0873</td>
<td><strong>0.0646</strong></td>
</tr>
<tr>
<td>ADKL-KRR</td>
<td>20</td>
<td>0.10</td>
<td>10</td>
<td>0.0575</td>
<td>0.0447</td>
<td><strong>0.0273</strong></td>
</tr>
</tbody>
</table>

Figure 5: Relative improvement of the MSE depending on the $\gamma_{\text{task}}$ parameter

For a more in-depth analysis, we show below the similar tables and figures for different values of $K$ (5, 10 and 20). These results confirm that regularizing the task encoder is helpful for any value of $K$, even though the impact seems to become much more important as $K$ increases (observe that the maximum improvement in each figure increases with $K$).

For $K = 5$

![Image](image5.png)

For $K = 10$

![Image](image10.png)
For $K = 20$

Table 5 presents the hyperparameter combinations used in the experiments to assess the impact of the trade-off parameter $\gamma_{\text{pseudo}}$, which governs the penalty applied to the divergence between the distribution of learned pseudo-representations and the distribution of actual representations. We also repeat in Fig. 6, the relative improvement of MSE compared to $\gamma_{\text{pseudo}} = 0$ as shown in the main text.

Table 5: Effect of the pseudo-examples regularization (parameter $\gamma_{\text{pseudo}}$) on the MSE performance

<table>
<thead>
<tr>
<th>Algorithm</th>
<th>$\gamma_{\text{task}}$</th>
<th>$\gamma_{\text{pseudo}}$</th>
<th>0.00</th>
<th>0.01</th>
<th>0.10</th>
</tr>
</thead>
<tbody>
<tr>
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<td>1</td>
<td>0.6079</td>
<td>0.4352</td>
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<tr>
<td></td>
<td>20</td>
<td>0.01</td>
<td>2</td>
<td>0.0873</td>
<td>0.0761</td>
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<tr>
<td>ADKL-KRR</td>
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<td>0.00</td>
<td>3</td>
<td>0.0526</td>
<td>0.0375</td>
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<tr>
<td>ADKL-GP</td>
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<td>0.10</td>
<td>4</td>
<td>2.2801</td>
<td>2.2112</td>
</tr>
<tr>
<td>ADKL-KRR</td>
<td>20</td>
<td>0.01</td>
<td>5</td>
<td>0.0535</td>
<td>0.0325</td>
</tr>
<tr>
<td>ADKL-GP</td>
<td>5</td>
<td>0.01</td>
<td>6</td>
<td>2.9466</td>
<td>2.7663</td>
</tr>
<tr>
<td></td>
<td>20</td>
<td>0.10</td>
<td>7</td>
<td>0.1147</td>
<td>0.1144</td>
</tr>
<tr>
<td></td>
<td>5</td>
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<td>8</td>
<td>0.1201</td>
<td>0.0958</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>0.01</td>
<td>9</td>
<td>3.1136</td>
<td>2.1511</td>
</tr>
<tr>
<td></td>
<td>0.00</td>
<td>10</td>
<td>2.8528</td>
<td>2.5611</td>
<td>2.4920</td>
</tr>
</tbody>
</table>

Figure 6: Relative improvement of the MSE depending on the $\gamma_{\text{task}}$ Parameter

Once again, for a more in-depth analysis, we show below the same format of tables and figures for different values of $K$, confirming again that regularizing using the pseudo-representation can be very helpful for any value of $K$. It is worth noticing here that the improvement gain is more consistent for
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$K = 5$ compared to $K \in \{10, 20\}$, supporting the fact that improving kernel maps evaluations using pseudo-representations is critical as size of the support set decreases.

For $K = 5$

\begin{tabular}{|l|c|c|c|}
\hline
algorithm & $\gamma_{\text{pseudo}}$ & $\gamma_{\text{task}}$ & 0.00 & 0.01 & 0.10 \\
\hline
ADKL-GP & 0.01 & 2.9466 & 2.7663 & 2.7121 \\
ADKL-KRR & 0.00 & 1.7123 & 1.6344 & 1.6868 \\
ADKL-GP & 0.00 & 1.1951 & 1.1655 & 1.1658 \\
ADKL-KRR & 0.00 & 2.8528 & 2.5661 & 2.4920 \\
ADKL-GP & 0.10 & 1.1998 & 1.1416 & 1.1442 \\
ADKL-KRR & 0.01 & 1.2129 & 1.1611 & 1.1716 \\
ADKL-GP & 0.10 & 1.2808 & 1.1974 & 1.2173 \\
ADKL-KRR & 0.01 & 1.7079 & 1.6655 & 1.6532 \\
\hline
\end{tabular}

For $K = 10$

\begin{tabular}{|l|c|c|c|}
\hline
algorithm & $\gamma_{\text{pseudo}}$ & $\gamma_{\text{task}}$ & 0.00 & 0.01 & 0.10 \\
\hline
ADKL-GP & 0.01 & 0.7329 & 0.7907 & 0.6294 \\
ADKL-KRR & 0.10 & 0.7479 & 0.7800 & 0.7663 \\
ADKL-GP & 0.00 & 0.3170 & 0.3070 & 0.3038 \\
ADKL-KRR & 0.00 & 0.3671 & 0.3628 & 0.2882 \\
ADKL-GP & 0.00 & 0.2967 & 0.2888 & 0.2893 \\
ADKL-KRR & 0.01 & 0.6556 & 0.7105 & 0.6577 \\
ADKL-GP & 0.10 & 0.7145 & 0.6758 & 0.7326 \\
ADKL-KRR & 0.10 & 0.6079 & 0.4352 & 0.5244 \\
ADKL-KRR & 0.10 & 0.2395 & 0.2299 & 0.2326 \\
\hline
\end{tabular}

For $K = 20$

\begin{tabular}{|l|c|c|c|}
\hline
algorithm & $\gamma_{\text{pseudo}}$ & $\gamma_{\text{task}}$ & 0.00 & 0.01 & 0.10 \\
\hline
ADKL-GP & 0.00 & 0.1201 & 0.0958 & 0.0940 \\
ADKL-KRR & 0.00 & 0.0794 & 0.1069 & 0.0702 \\
ADKL-GP & 0.00 & 0.0305 & 0.0327 & 0.0447 \\
ADKL-KRR & 0.01 & 0.0302 & 0.0289 & 0.0273 \\
ADKL-GP & 0.00 & 0.0574 & 0.0585 & 0.0575 \\
ADKL-KRR & 0.10 & 0.1147 & 0.1144 & 0.0870 \\
ADKL-GP & 0.01 & 0.0535 & 0.0325 & 0.0325 \\
ADKL-KRR & 0.10 & 0.0430 & 0.0414 & 0.0395 \\
\hline
\end{tabular}

Overall, the effect of the regularization is beneficial, even though we witness a few pathological cases.

A.3 Joint Impact of $\gamma_{\text{task}}$ and $\gamma_{\text{pseudo}}$

Since both $\gamma_{\text{task}}$ and $\gamma_{\text{pseudo}}$, have a high impact on the training and the generalization performance, we need to assess the relationship between the two. Fig. 7 shows, for different values of $K$, the relative improvement of the test MSE compared to the case where no regularization is done, i.e. $\gamma_{\text{task}} = 0$ and $\gamma_{\text{pseudo}} = 0$. Overall, one can see that higher is better in both dimensions but there seems to be a sweet spot on the grid for each value of $K$ and therefore we can only advise the user to cross-validate on those hyperparameters.

Overall, the effect of the regularization is beneficial, even though we witness a few pathological cases.

Figure 7: Average relative improvement of the MSE and joint impact of $\gamma_{\text{task}}$ and $\gamma_{\text{pseudo}}$. 

15
Prediction curves on the Sinusoids collection

Figure 8 presents a visualization of the results obtained by each model on three tasks taken randomly from the meta-test set. We provide the model with ten examples from an unseen task consisting of a slightly noisy sine function (shown in blue), and present in orange the predictions made by the network based on these ten examples.