

# Tensor-Train Kernel Learning for Gaussian Processes

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## Abstract

We propose a new kernel learning approach based on efficient low-rank tensor compression for Gaussian process (GP) regression. The central idea is to compose a low-rank function represented in a hierarchical tensor format with a GP covariance function. Compared to similar deep neural network architectures, this approach facilitates to learn significantly more expressive features at lower computational costs as illustrated in the examples. Additionally, over-fitting is avoided with this compositional model by taking advantage of its inherent regularisation properties. Estimates of the generalisation error are compared to five baseline models on three synthetic and six real-world data sets. The experimental results show that the incorporated tensor network enables a highly accurate GP regression with a comparatively low number of trainable parameters. The observed performance is clearly superior (usually by an order of magnitude in mean squared error) to all examined standard models, in particular to deep neural networks with more than 1000 times as many parameters.

**Keywords:** Tensor networks, Tensor-Train decomposition, Gaussian processes.

## 1. Introduction

This work introduces a kernel learning approach based on tensor network models for approximating high-dimensional functions. Specifically, we investigate the capabilities of tree tensor networks (Stoudenmire and Schwab, 2016; Oseledets, 2011) to learn expressive features for Gaussian process (GP) regression (Rasmussen and Williams, 2005) in order to enable uncertainty aware predictions. To achieve this, the tensor network parameters are treated as kernel hyper-parameters, which can be optimised through the marginal likelihood of the GP.

We also highlight tensor kernel learning as an approach for quantifying predictive uncertainty for tensor network models. Thereby, their applicability can be extended to high stakes domains like medical applications. For these domains, integration of prior knowledge into the modelling process and interpretability through rigorous mathematical foundations are important properties, which neural networks (arguably) do not provide (yet) although they exhibit similar (but theoretically larger) representational power.

A first approach for using tensor networks for kernel learning was proposed in (Konstantinidis et al., 2021). We add to that work in two important ways: first, we make use of a different type of low rank structure, namely Tensor-Trains (TTs) instead of the Canonical-Polyadic (CP) decomposition. Contrary to the latter, the TT format permits an efficient pre-training procedure so that the tensor network kernel starts with near optimal weights

before end-to-end training. Moreover, the TT format allows pre-training and gradient-based optimisation to be easily regularised w.r.t. the norm of an underlying function space. This prevents over-fitting even in complex architectures and speeds up convergence. As a second addition, we perform thorough experiments for all compared architectures in this work and are able to show that our TT kernel method beats all baselines on the considered real world tasks usually by at least one order of magnitude. These baselines include the architecture proposed in (Konstantinidis et al., 2021) as well as a deep neural network with about a hundred times as many parameters as our method.

The rest of the work is structured as follows. In section 2 we briefly review the related literature. Section 3 then introduces the concept of low-rank feature extractors using TTs, which will be combined with GPs in section 4 to define the Tensor-Train kernel learning method. In section 5 we detail the experimental setup on several synthetic and real world data sets and report the results. Finally, section 6 provides a summary of the work and some concluding remarks.

## 2. Related Work

Tensor network models for supervised learning have become increasingly popular in the machine learning community for the last five years. Some notable contributions are (Stoudenmire and Schwab, 2016; Novikov et al., 2017; Efthymiou et al., 2019; Cohen and Shashua, 2016; Glasser et al., 2020; Blagoveschensky and Phan, 2020; Grelier et al., 2019; Ali and Nouy, 2020a,b, 2021). It should be noted that the physics and numerical mathematics communities have driven research in this direction for a much longer time, resulting, e.g. in standard textbooks like (Hackbusch, 2012), a range of training methods (Holtz et al., 2011; White, 1992; Grasedyck and Krämer, 2019) or application of tensor networks to (uncertainty quantification of) partial differential equations (Bachmayr et al., 2016; Eigel et al., 2019; Dolgov et al., 2015; Eigel et al., 2017, 2018), optimal transport (Eigel et al., 2020) and control (Oster et al., 2020).

Uncertainty quantification for tensor networks and tensor regression can e.g. be found in (Kirstein et al., 2020; Hu et al., 2020; Hawkins and Zhang, 2019). In contrast to our work however, the authors treat the whole model as Bayesian rather than only using a Bayesian layer at the end of a multi-layer model architecture.

Another interesting work is (Izmailov et al., 2018), where tensor networks are used to compress a tensor-variate mean function of a sparse Gaussian process, in order to radically increase the possible number of inducing points.

In what follows, we treat kernel learning as the composition of standard kernels (i.e. a Matérn or squared-exponential kernel) with functions from a different model class (i.e. neural or tensor networks). This approach is mainly developed in (Fu Jie Huang and LeCun, 2006; Salakhutdinov and Hinton, 2007) and further extended, e.g. in (Wilson et al., 2016a,b; Damianou and Lawrence, 2013; Zhuang et al., 2011). The authors compose different kernel functions or use deep neural networks in conjunction with kernels to learn rich high level features, which improve performance compared to classical support vector machines and Gaussian processes.

In (Ober et al., 2021) the question of over-fitting in kernel learning with deep neural networks is posed. The authors show empirically that an optimisation of the marginal

likelihood does not provide enough regularisation for highly complex models with lots of parameters like deep neural networks. This is in contrast to the usual assumption that the marginal likelihood objective - as part of the Bayesian approach - should deliver an inherent complexity penalty sufficient to prevent over-fitting. At this point we conjecture that tensor networks are less prone to this kind of behaviour due to the inherent regularisation properties of a low-rank structure and the introduction of prior knowledge through the choice of basis functions for the tensor-product space.

Interestingly, (van Amersfoort et al., 2021) implicitly addresses the same problem as (Ober et al., 2021). As a remedy for over-fitting in their deep neural network model, the authors introduce further regularisation, which increases smoothness of the generated function approximation. This supports our claim that tensor networks can act as a stand-in for deep neural networks, especially when introduction of prior information is needed, since regularisation is much more intuitive and a direct feature of the model.

### 3. Low-Rank Tensor Networks

#### 3.1. Basis Representations in High-Dimensional Spaces

In order to construct a function approximator, we start with a set of basis functions  $\{P_{\alpha_i} : \mathbb{R} \rightarrow \mathbb{R}\}_{i=1}^d$ , with  $\alpha_i = 1, \dots, J_i$  and  $J_i \in \mathbb{N}$ . Under the assumption of a tensor product function space, we can represent the approximator as

$$f_W(x) = \sum_{\alpha_1=1}^{J_1} \cdots \sum_{\alpha_d=1}^{J_d} W_{\alpha_1, \dots, \alpha_d} \prod_{i=1}^d P_{\alpha_i}(x_i), \quad (1)$$

for  $x = (x_1, \dots, x_d) \in \mathcal{X}$ , where  $W$  is a coefficient tensor of order  $d$ , i.e.  $W \in \mathbb{R}^{J_1 \times \dots \times J_d}$ . While this means large expressive power, it poses a significant computational burden due to the exponential complexity growth in the dimensionality  $d$  (“curse of dimensionality”), which we alleviate by the introduction of a low-rank tensor format as described subsequently.

#### 3.2. Tensor Decomposition

The exponential full tensor complexity (regarding storage and algorithms) fortunately can be reduced significantly by exploiting an often encountered low-rank structure of  $f_W$  (see, e.g. (Cichocki et al., 2016)). For our experiments we choose the so-called Tensor-Train format (Oseledets, 2011), which is a popular linearised case of more general tree tensor formats (Grelier et al., 2019). Recalling expansion (1), a low-rank TT compression of the coefficient tensor  $W$  reads

$$W_{\alpha_1, \dots, \alpha_d} \approx \sum_{k_0=1}^{r_0} \cdots \sum_{k_d=1}^{r_d} \prod_{i=1}^d V_{k_{i-1}, \alpha_i, k_i}^{(i)}, \quad (2)$$

with  $r_0 = r_d = 1$ . The components  $V^{(i)} \in \mathbb{R}^{r_i \times J_i \times r_{i+1}}$ ,  $i = 1, \dots, d-1$ , are tensors of order three. The remaining components  $V^{(1)}, V^{(d)}$  are of order two. The upper summation bounds  $r_i$ ,  $i = 1, \dots, d-1$ , are called TT ranks.

The TT format exhibits a storage complexity of  $\mathcal{O}(\max(J_1, \dots, J_d)d \max(r_1, \dots, r_{d-1})^2)$ , which scales only linearly in the dimension  $d$ , hence avoiding the curse of dimensionality.

Imposing a low-rank structure on the coefficient tensor results in an embedded manifold of functions with low-rank tensors of rank at most  $\mathbf{r} = (r_1, \dots, r_{d-1})$ . In practice this can be interpreted as neglecting spurious features by controlling the rank parameters  $r_1, \dots, r_{d-1}$  in the sense of an implicit feature selection. Analytical bounds, which quantify this behaviour rigorously can be found in (Oseledets, 2011; Oseledets and Tyrtyshnikov, 2010).

Utilising a TT format for the coefficient tensor  $W$  yields

$$f_W(x) = \sum_{\alpha_1, \dots, \alpha_d} \sum_{k_0=1}^{r_0} \dots \sum_{k_d=1}^{r_d} \prod_{i=1}^d V_{k_{i-1}, \alpha_i, k_i}^{(i)} P_{\alpha_i}(x_i). \quad (3)$$

Ultimately, this model allows to perform practical computations in a very high dimensional space while staying trainable.

## 4. Tensor-Train Kernel Learning

As an application and extension of tensor network regression, Gaussian processes provide an immediate and principled way to equip an otherwise deterministic model with probabilistic features (see section A for a brief summary). Then again, tensor networks can also be beneficial to GP regression. Due to the inherent regularisation by the choice of basis functions and by imposing a low-rank structure, the problem of over-fitting as, e.g. encountered in standard deep kernel learning can be significantly reduced. Additionally, the low-rank structure performs implicit feature selection on the data, thus providing more expressive features for GP regression. Therefore, combining a low-rank tensor network model and GP should improve upon regular GP regression especially in the domain of high dimensional problems, where kernel evaluation is costly. Thus, the combination of a tensor network compression together with a GP model provides principled and flexible modelling options.

### 4.1. Tensor-Train Kernel Model

The general idea of our proposed method is to first propagate data through a TT function, then subsequently perform a GP regression on the resulting outputs (latents). Thus, the TT function serves as a feature extractor for the GP.

Concretely, we estimate an  $\mathbb{R}$ -valued random variable  $Y$  which depends on a  $d$ -dimensional  $\mathcal{X}$ -valued random vector  $X$ . Subsequently, we assume  $Y = \Phi(X)$  and our goal is to approximate  $\Phi$  by a GP  $g$  with TT feature extractor using samples of the pair  $(X, Y)$ . Therefore, we start by defining an i.i.d. data set  $D_N := \{x^{(n)}, y^{(n)}\}$  for  $n = 1, \dots, N$ .

To this end, let  $f_W: \mathcal{X} \rightarrow \mathcal{Z}$  denote a TT function with coefficient tensor  $W$ , mapping to the latent space  $\mathcal{Z}$ . This space need not be one-dimensional: multivariate latents can be achieved by adding a latent output dimension  $d_{\mathcal{Z}}$  to the first component  $V^{(1)}$  of the tensor train, transforming it to shape  $r_1 \times J_1 \times r_2 \times d_{\mathcal{Z}}$ . Now, assume the composite kernel

$$k(x, x' | \theta) := \hat{k}(f_W(x), f_W(x') | \theta),$$

where  $\hat{k}: \mathcal{Z} \times \mathcal{Z} \rightarrow \mathbb{R}$  is a base kernel with hyper-parameters  $\theta$ , e.g. a squared-exponential or linear kernel, and  $\boldsymbol{\theta} := \{W, \theta\}$  are the joint hyper-parameters. Now, we perform regression with the composite kernel, yielding the GP prior

$$g(x) \sim \mathcal{GP}(m(x), k(x, x' | \boldsymbol{\theta})). \quad (4)$$

With  $\mathbf{k}_i = k(x, x^{(i)} | \boldsymbol{\theta})$ ,  $K_{i,j} = k(x^{(i)}, x^{(j)} | \boldsymbol{\theta})$ ,  $\mathbf{y}_i = y^{(i)} - m(x^{(i)})$  this leads to the representation

$$m^\Phi(x) = m(x) + \mathbf{k}^\top K^{-1} \mathbf{y} \quad (5)$$

for the posterior GP’s mean function  $m^\Phi$  (for more details we refer again to [section A](#)).

With the described approach, we have defined a composition of a tensor network function and a GP, where the parameters of the TT are now treated as kernel hyper-parameters. Thus, we are able to jointly train our tensor network model and the GP using gradient-based optimisation. However, optimising the resulting composite hyper-parameters of the kernel globally by standard gradient descent neglects the multi-linear structure of the Tensor-Train. We thus prefer to pre-train the TT, using a regularised version of the Alternating Linear Scheme (ALS) ([Holtz et al., 2011](#)) (see [section B](#)).

## 4.2. Learning Tensor-Train Kernel Models

After pre-training the TT model with ALS, we resort to stochastic variational inference (VI) for training of a sparse GP with composite kernel  $k(x, x' | \boldsymbol{\theta})$ . For a thorough treatment of VI for GPs we further suggest ([Leibfried et al., 2021](#)).

The idea is to introduce a set of inducing point locations  $\{x_1, x_2, \dots, x_M\}$ , which are used as pseudo-training data in order to ensure a sparse GP. For VI we have the following optimisation problem

$$\begin{aligned} \max_{\boldsymbol{\theta}, \gamma} \sum_{n=1}^N \mathbb{E}_{q_{\boldsymbol{\theta}, \gamma}} \log p(y_n | g(x_n)) \\ - \text{KL}(q_{\boldsymbol{\theta}, \gamma}(u) || p_{\boldsymbol{\theta}, \gamma}(u)), \end{aligned} \quad (6)$$

where  $q_{\boldsymbol{\theta}, \gamma}$  is the variational density (depended on kernel hyper-parameters  $\boldsymbol{\theta}$  and variational parameters  $\gamma$ ) used to approximate the GP posterior with inducing variables  $u = g(x)$ , i.e. a point-wise evaluation of the GP at the inducing points. In VI related literature the stated objective is usually called evidence lower bound, ELBO for short.

Furthermore, we need the likelihood function  $p(y_n | g(x_n))$  and prior density  $p_{\boldsymbol{\theta}, \gamma}(u)$  which corresponds to equation 4. Due to the summation over data points, the objective is amenable to stochastic mini-batching. In our case of regression, the expectation can be evaluated in closed-form, assuming  $q_{\boldsymbol{\theta}, \gamma}$  is a Gaussian. Otherwise, we rely on Monte-Carlo integration.

In our experiments, we introduce the inducing points not on the data space  $\mathcal{X}$ , but on the (potentially lower dimensional) latent space  $\mathcal{Z}$  in order to reduce complexity. This leads to an overall asymptotic cost of

$\mathcal{O}(N(\max(J_1, \dots, J_d)d \max(r_1, \dots, r_{d-1})^2 + M^2))$  for  $M \ll N$  operations.

## 5. Numerical Experiments

Tensor-Train kernel learning (TTKL) and five baselines<sup>1</sup>, namely TT and sparse GP regression, a deep neural network (DNN), deep kernel learning (DKL) and the Canonical-Polyadic model from (Konstantinidis et al., 2021) (CPKL) are compared on three synthetic and six real-world data sets<sup>2</sup>. Moreover, we execute ablation experiments for TTKL.

As methodological tool for hyper-parameter optimisation of all models, we use a random search (Bergstra and Bengio, 2012) together with advanced early-stopping (Li et al., 2020). Additionally, we repeat model evaluation six times with randomly selected seeds and report the resulting mean and standard deviation.

Further information on the experiments, baselines, data sets and hyper-parameter optimisation is given in section C.

### 5.1. Tensor-Train Kernel Learning Set-up

The TT feature extractor uses orthonormal polynomials in a Sobolev space ( $H^1$ ) of order one up to a fixed degree  $J$  as basis functions  $P_{\alpha_i}$  which is also used to impose an  $H^1$ -regularisation during end-to-end training (see section B for details). Additionally, we constrain the tensor-train rank uniformly to a fixed value  $r$ . For pre-training, we use regularised ALS (Holtz et al., 2011), where we solve the underlying linear systems by LU factorisation. This yields improved initial values for subsequent stochastic gradient based optimisation of the tensor kernel model. For the base kernel  $\hat{k}$ , we use the standard radial basis function (RBF) kernel  $k_{RBF}$  (see section A). Subsequently, we use sparse variational inference (VI) (Leibfried et al., 2021) with a mean-field Gaussian density. Additionally, we assign individual initial learning rates for each TT component, the RBF kernel hyper-parameters and VI related parameters by specifying so-called parameter groups for PyTorch’s optimiser class. As optimisation method we choose ADAM (Kingma and Ba, 2015) with default hyper-parameters, except for the initial learning rates. In order to adjust the TT’s latent dimension from pre-training to end-to-end training, we simply duplicate the TT’s first component to match.

In the ablation experiments we compare the individual additions we made to the TTKL model to one with all additions removed (Vanilla). That is, we examine the influence of pre-training (Pre-Train),  $H^1$  regularisation ( $H^1$ ) and TT component specific optimisation (Opt).

### 5.2. Results

In Table 3 we present estimates of the generalisation error of TTKL and the various baselines. Additionally, we give a parameter count for each trained model on the specific data set. We also provide the probabilistic models’ log-likelihoods in Table 1, which by including uncertainty information, constitute a more “complete” metric. However, in our discussion, we will focus on the mean squared error, because it is available for probabilistic and deterministic models alike. Note that the results for DKL are different than those of (Wilson et al., 2016a), as we used a slightly different, richer architecture for the DNN (see C.3.2) and a standard

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1. All models were implemented with the PyTorch (Paszke et al., 2019) and GPyTorch (Gardner et al., 2018) frameworks. Distributed model selection and evaluation were facilitated by means of Ray (Moritz et al., 2018) and Tune (Liaw et al., 2018).

2. Available at [https://drive.google.com/open?id=OBxWe\\_IuTnMfcYXhdUNwRHBKTLU](https://drive.google.com/open?id=OBxWe_IuTnMfcYXhdUNwRHBKTLU)

kernel (squared exponential), which we also used for all other model classes. The change in DNN architecture is due to non-reproducible results of the original paper. For the sake of consistency, we used sparse GP regression on all data sets, irrespective of the size.

The TTKL model outperforms all baselines in terms of mean squared error on five of the six real data sets, on four even by an order of magnitude. On the synthetic data sets, it surpasses the GP, DNN, DKL and CPKL models, while only TT regression achieves better results. We attribute the TT regression’s strong performance to the structure of the synthetic data sets, which are readily approximated in a high dimensional polynomial space (Ali and Nouy, 2021). Moreover we can see that having a TT feature extractor improves upon a pure GP’s generalisation error. Additionally, we argue that better performance of TTKL compared to DKL corroborates our earlier conjecture about better regularisation properties and therefore a reduced risk of over-fitting in TTKL. Another point we stress is the immensely reduced number of trainable parameters of the TTKL model when compared to the neural network feature extractor of DKL. Furthermore, especially in the high dimensional setting, the use of a TT feature extractor for GP regression even reduces the amount of parameters when compared to ordinary sparse GP regression. Moreover, construction of the Euclidean distance-based kernel matrix is sped-up, due to the kernel’s evaluation on the lower dimensional latent space. Furthermore, we notice that CPKL fails in the high dimensional tasks. We attribute this to the exploding / vanishing gradient problem when training tensor network models purely by gradient descent. This behaviour underlines the importance of efficient and robust training methods for tensor based models such as ALS. With regard to the log-likelihood, TTKL outperforms the probabilistic baseline methods on all but one data set. This shows TTKL’s ability to explain the data much better than the compared models.

In Table 2 we give results of the ablation experiments carried out on the synthetic data sets. Specifically, we remove all extensions, i.e. pre-training,  $H^1$  regularisation and component specific optimisation, in order to receive a vanilla TTKL model. Subsequently, we only add one extension a time in order to investigate its effect on TTKL’s performance. From these results we conclude that training methods specific to the structure of low-rank functions used in pre-training provide the most benefit. Next in line for consistently increased performance is  $H^1$  based regularisation. Component specific gradient optimisation loses its benefit with increasing data dimension. Only on the low dimensional problem is the vanilla model able to achieve near comparable performance to extended models. We conclude that the interactions of extensions contribute the most to the performance of TTKL, especially in regard to increased feature dimension.

## 6. Conclusion

In this work we presented a new method for GP kernel learning in the form of low-rank functions parameterised by decomposed weight tensors. We derived the TTKL model and gave a short introduction into the learning method. Furthermore, we compared TTKL to five baseline models - namely TT and GP regression, a DNN, DKL and CPKL - on three synthetic and six real-world data sets. All models were subjected to extensive hyper-parameter optimisation and an average over six performance evaluations on an independent test set was reported together with the respective standard deviation. TTKL showed superior

Table 1: Log-likelihood (higher is better) and one standard deviation for all probabilistic models on three synthetic and six real-world data sets.

Data set	$N$	$d$	Test LL			
			TTKL (ours)	CPKL	DKL	GP
HighDimSin (Synthetic)	100 000	30	<b>2.46</b>	-1.94	$2.43 \times 10^{-2}$	-3.43
			$\pm 9.51 \times 10^{-1}$	$\pm 6.90 \times 10^{-4}$	$\pm 7.84 \times 10^{-2}$	$\pm 7.15 \times 10^{-2}$
Friedman (Synthetic)	100 000	5	<b>3.06</b>	2.50	$1.33 \times 10^{-1}$	1.53
			$\pm 3.05 \times 10^{-2}$	$\pm 1.28 \times 10^{-1}$	$\pm 6.66 \times 10^{-2}$	$\pm 9.32 \times 10^{-2}$
Grid (Synthetic)	65 536	2	<b>2.54</b>	1.88	1.34	
			$\pm 8.21 \times 10^{-2}$	$\pm 5.52 \times 10^{-1}$	$\pm 9.02 \times 10^{-2}$	$\pm$
Kegg (Real)	48 827	22	<b><math>6.40 \times 10^{-1}</math></b>	-1.74	$3.56 \times 10^{-1}$	$3.41 \times 10^{-1}$
			$\pm 5.11 \times 10^{-1}$	$\pm 2.72 \times 10^{-5}$	$\pm 1.11 \times 10^{-1}$	$\pm 1.38 \times 10^{-2}$
Skillcraft (Real)	3338	19	-1.53	$-7.20 \times 10^{-1}$	<b><math>-2.95 \times 10^{-1}</math></b>	$-3.05 \times 10^{-1}$
			$\pm 4.39 \times 10^{-1}$	$\pm 2.58 \times 10^{-1}$	$\pm 3.14 \times 10^{-2}$	$\pm 7.30 \times 10^{-3}$
Elevators (Real)	16 599	18	<b><math>8.17 \times 10^{-1}</math></b>	$-8.04 \times 10^{-2}$	$7.16 \times 10^{-2}$	$7.61 \times 10^{-2}$
			$\pm 3.51 \times 10^{-2}$	$\pm 8.30 \times 10^{-2}$	$\pm 1.18 \times 10^{-2}$	$\pm 1.80 \times 10^{-3}$
Housing (Real)	506	13	<b><math>-9.46 \times 10^{-1}</math></b>	-3.98	-3.54	-5.50
			$\pm 2.78 \times 10^{-1}$	$\pm 8.89 \times 10^{-2}$	$\pm 1.77 \times 10^{-1}$	$\pm 3.04 \times 10^{-1}$
Protein (Real)	45 730	9	<b><math>1.11 \times 10^{-1}</math></b>	-1.17	$-7.49 \times 10^{-1}$	-1.18
			$\pm 4.17 \times 10^{-2}$	$\pm 7.00 \times 10^{-5}$	$\pm 1.94 \times 10^{-1}$	$\pm 1.94 \times 10^{-2}$
Kin40K (Real)	40 000	8	<b>2.15</b>	$3.11 \times 10^{-1}$	$7.92 \times 10^{-1}$	$-2.78 \times 10^{-2}$
			$\pm 1.34 \times 10^{-1}$	$\pm 9.27 \times 10^{-1}$	$\pm 2.89 \times 10^{-2}$	$\pm 1.72 \times 10^{-3}$

performance when compared to the other baselines on real-world data sets. Only on synthetic data could better performance be achieved by the TT regression. Moreover, we conducted ablation experiments. These clearly suggested that best performance of TTKL can only be achieved with the interaction of all extensions proposed for our method.

More importantly, we gave evidence of the benefit of low-rank functions for kernel learning on regression problems when compared to deep neural networks. Additionally, we introduced means to extend ordinary low-rank tensor regression with uncertainty estimates for its prediction. We hence showed that TTKL is a viable alternative to DKL and a promising research area.

Table 2: Mean squared error and one standard deviation for ablation experiments (vanilla, extended and full TTKL model) on three synthetic data sets.

Data set	$N$	$d$	Test MSE				
			Vanilla	$H^1$	Pre-Train	Opt	Full
HighDimSin	100 000	30	2.84	$1.62 \times 10^{-2}$	$4.48 \times 10^{-5}$	2.89	$2.54 \times 10^{-4}$
			$\pm 1.04 \times 10^{-2}$	$\pm 4.75 \times 10^{-2}$	$\pm 1.92 \times 10^{-5}$	$\pm 7.92 \times 10^{-2}$	$\pm 3.75 \times 10^{-4}$
Friedman	100 000	5	1.92	$3.76 \times 10^{-2}$	$1.40 \times 10^{-2}$	$1.54 \times 10^{-2}$	$9.00 \times 10^{-6}$
			$\pm 2.84$	$\pm 8.59 \times 10^{-3}$	$\pm 1.71 \times 10^{-5}$	$\pm 1.91 \times 10^{-2}$	$\pm 9.42 \times 10^{-7}$
Grid	65 536	2	$8.39 \times 10^{-5}$	$4.58 \times 10^{-5}$	$5.15 \times 10^{-6}$	$1.15 \times 10^{-4}$	$4.99 \times 10^{-6}$
			$\pm 2.77 \times 10^{-5}$	$\pm 2.91 \times 10^{-5}$	$\pm 1.60 \times 10^{-6}$	$\pm 5.26 \times 10^{-5}$	$\pm 8.29 \times 10^{-7}$

Table 3: Mean squared error, one standard deviation and number of trainable parameters for all models on three synthetic and six real-world data sets.

Data set	$N$	$d$	Test MSE					
			Num Params					
			TTKL (ours)	CPKL	DKL	DNN	TT	GP
HighDimSin (Synthetic)	100 000	30	$2.54 \times 10^{-4}$	2.83	$2.77 \times 10^{-2}$	$3.34 \times 10^{-2}$	<b><math>2.69 \times 10^{-14}</math></b>	$2.11 \times 10^{-2}$
			$\pm 3.75 \times 10^{-4}$	$\pm 2.77 \times 10^{-4}$	$\pm 5.97 \times 10^{-3}$	$\pm 3.47 \times 10^{-3}$	$\pm 1.87 \times 10^{-15}$	$\pm 2.94 \times 10^{-4}$
			26 609	40 767	58 007 943	42 575 956	252 600	18 512
Friedman (Synthetic)	100 000	5	$9.00 \times 10^{-6}$	$1.20 \times 10^{-4}$	$1.14 \times 10^{-2}$	$1.23 \times 10^{-2}$	<b><math>2.34 \times 10^{-16}</math></b>	$1.73 \times 10^{-3}$
			$\pm 9.42 \times 10^{-7}$	$\pm 3.12 \times 10^{-5}$	$\pm 2.74 \times 10^{-3}$	$\pm 1.88 \times 10^{-3}$	$\pm 1.25 \times 10^{-16}$	$\pm 2.85 \times 10^{-4}$
			4308	3746	50 820 695	2 052 701	1020	2572
Grid (Synthetic)	65 536	2	$4.99 \times 10^{-6}$	$1.12 \times 10^{-3}$	$1.06 \times 10^{-3}$	$1.29 \times 10^{-3}$	<b><math>9.02 \times 10^{-8}</math></b>	$2.65 \times 10^{-6}$
			$\pm 8.29 \times 10^{-7}$	$\pm 1.48 \times 10^{-3}$	$\pm 3.36 \times 10^{-4}$	$\pm 1.97 \times 10^{-4}$	$\pm 7.66 \times 10^{-12}$	$\pm 4.61 \times 10^{-7}$
			2182	1394	2 695 770	2 614 358	52	1030
Kegg (Real)	48 827	22	<b><math>1.93 \times 10^{-3}</math></b>	1.89	$2.22 \times 10^{-2}$	$1.76 \times 10^{-2}$	$3.23 \times 10^{-2}$	$2.39 \times 10^{-2}$
			$\pm 3.85 \times 10^{-4}$	$\pm 2.66 \times 10^{-5}$	$\pm 1.60 \times 10^{-3}$	$\pm 8.90 \times 10^{-4}$	$\pm 2.78 \times 10^{-3}$	$\pm 2.52 \times 10^{-4}$
			1469	25 258	2 999 434	57 336 262	19 448	17 844
Skillcraft (Real)	3338	19	<b><math>3.61 \times 10^{-2}</math></b>	$1.51 \times 10^{-1}$	$8.18 \times 10^{-2}$	$1.14 \times 10^{-1}$	$3.18 \times 10^{-1}$	$9.78 \times 10^{-2}$
			$\pm 1.21 \times 10^{-2}$	$\pm 1.12 \times 10^{-5}$	$\pm 1.52 \times 10^{-3}$	$\pm 5.76 \times 10^{-2}$	$\pm 5.44 \times 10^{-2}$	$\pm 1.92 \times 10^{-3}$
			14 457	18 318	9 530 027	57 292 606	12 350	6 709
Elevators (Real)	16 599	18	$1.14 \times 10^{-2}$	$6.31 \times 10^{-2}$	$4.91 \times 10^{-2}$	$4.86 \times 10^{-2}$	<b><math>9.07 \times 10^{-3}</math></b>	$4.99 \times 10^{-2}$
			$\pm 1.64 \times 10^{-3}$	$\pm 2.18 \times 10^{-6}$	$\pm 3.82 \times 10^{-4}$	$\pm 3.23 \times 10^{-4}$	$\pm 1.74 \times 10^{-4}$	$\pm 2.15 \times 10^{-4}$
			2366	7400	12 513 644	14 920 122	52 200	7346
Housing (Real)	506	13	<b><math>1.62 \times 10^{-1}</math></b>	$8.17 \times 10^1$	$1.88 \times 10^1$	$2.82 \times 10^1$	$1.42 \times 10^1$	$3.83 \times 10^1$
			$\pm 6.75 \times 10^{-2}$	$\pm 4.76 \times 10^{-1}$	$\pm 1.22$	$\pm 1.16$	$\pm 1.54 \times 10^{-1}$	$\pm 2.62$
			2116	21 023	89 226 791	2 982 619	3770	1835
Protein (Real)	45 730	9	<b><math>5.18 \times 10^{-2}</math></b>	$6.04 \times 10^{-1}$	$4.00 \times 10^{-1}$	$2.06 \times 10^{-1}$	$6.70 \times 10^{-1}$	$6.00 \times 10^{-1}$
			$\pm 7.96 \times 10^{-3}$	$\pm 1.04 \times 10^{-5}$	$\pm 2.04 \times 10^{-1}$	$\pm 1.02 \times 10^{-2}$	$\pm 7.70 \times 10^{-2}$	$\pm 1.08 \times 10^{-3}$
			4754	18 974	7 557 785	26 561 968	1170	6185
Kin40K (Real)	40 000	8	<b><math>3.49 \times 10^{-4}</math></b>	$1.82 \times 10^{-2}$	$9.74 \times 10^{-3}$	$1.13 \times 10^{-2}$	$1.96 \times 10^{-3}$	$4.00 \times 10^{-2}$
			$\pm 1.84 \times 10^{-4}$	$\pm 1.78 \times 10^{-2}$	$\pm 5.09 \times 10^{-4}$	$\pm 3.55 \times 10^{-4}$	$\pm 6.43 \times 10^{-5}$	$\pm 4.62 \times 10^{-4}$
			9916	11 090	9 466 377	2 631 074	3600	7738

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## Appendix A. Gaussian Processes

We briefly recall the fundamentals of Gaussian process regression, following the notation used in the classical reference (Rasmussen and Williams, 2005).

Starting from the same point as in subsection 3.1, we assume random variables  $X, Y$  with realisations  $\mathbf{X} = [x^{(1)}, \dots, x^{(N)}]$  drawn according to  $\rho$  and  $\mathbf{y} = [y^{(1)}, \dots, y^{(N)}]$  with  $y^{(n)} = \Phi(x^{(n)})$  for all  $n = 1, \dots, N$ . The goal now is to approximate  $\Phi$  by  $f$  for which we assign a Gaussian process (GP) prior distribution  $f_0$  to  $f$ . A GP (prior)  $f_0$  is characterised by a mean function  $m_0: \mathcal{X} \rightarrow \mathbb{R}$  and a symmetric positive-definite function  $k_0: \mathcal{X} \times \mathcal{X} \rightarrow \mathbb{R}$ . Thus, we denote it by

$$f_0(x) \sim \mathcal{GP}(m_0(x), k_0(x, x')).$$

For any finite number of inputs, the corresponding function values of the GP have a joint Gaussian distribution

$$\begin{aligned} f_0(\mathbf{X}) &= [f_0(x^{(1)}), \dots, f_0(x^{(N)})] \\ &\sim \mathcal{N}(m_0(\mathbf{X}), k_0(\mathbf{X}, \mathbf{X})), \end{aligned}$$

where the mean vector  $m_0(\mathbf{X}) = [m_0(x^{(1)}), \dots, m_0(x^{(N)})]$  and covariance matrix  $k_0(\mathbf{X}, \mathbf{X}) = k_0(x^{(i)}, x^{(j)})$  for all  $i, j = 1, \dots, N$  are defined by the mean function and covariance function, respectively.

In order to predict at new points  $\mathbf{X}_* = (x_*^{(n+1)}, \dots, x_*^{(n+m)})$  we condition the GP on the data, yielding the posterior process

$$f_*(x) \sim \mathcal{GP}(m_*(x), k_*(x, x')).$$

In this case, we obtain updated mean and covariance functions  $m_*, k_*$  with corresponding mean vector  $m_*(\mathbf{X}_*) = m_0(\mathbf{X}_*) - k_0(\mathbf{X}_*, \mathbf{X})k_0(\mathbf{X}, \mathbf{X})^{-1}(m_0(\mathbf{X}) - \mathbf{y})$  and covariance matrix  $k_*(\mathbf{X}_*, \mathbf{X}_*) = k_0(\mathbf{X}_*, \mathbf{X}_*) - k_0(\mathbf{X}_*, \mathbf{X})k_0(\mathbf{X}, \mathbf{X})^{-1}k_0(\mathbf{X}, \mathbf{X}_*)$ .

Furthermore, assuming a mean-zero prior GP we have

$$m_*(x) = \sum_{n=1}^N c_n k_0(x, x^{(n)}) \quad \forall x \in \mathcal{X}, \quad (7)$$

with  $\mathbf{c} = [c_1, \dots, c_N]$  and  $\mathbf{c} = k_0(\mathbf{X}, \mathbf{X})^{-1}\mathbf{y}$ . This is the definition of a linear (kernel) predictor. Therefore, the prior covariance function identifies the function space of the posterior mean as a reproducing kernel Hilbert space, namely  $m_* \in \mathcal{K} := \text{span}\{k_0(\cdot, x), x \in \mathcal{X}\}$ .

For our purposes it is important to note that the covariance function  $k_0$  usually depends on some set of hyper-parameters  $\boldsymbol{\theta}$ , i.e.

$$k_0(x, x') = k_0(x, x' | \boldsymbol{\theta}).$$

For example, the common RBF kernel

$$k_{\text{RBF}}(x, x') = \sigma_f^2 \exp\left(-\frac{1}{2\ell} \|x - x'\|^2\right)$$

has the prior standard deviation  $\sigma_f$ , known as the signal variance, and the lengthscale  $\ell$ , which determines the rate of correlation decay with increasing distance between inputs, as hyper-parameters. Together with the noise variance, the set of hyper-parameters for this simple kernel is given by  $\boldsymbol{\theta} = \{\sigma_f, \ell, \sigma_n\}$ .

A GP is trained by optimising the kernel hyper-parameters. This can be done in a principled way by maximising the marginal likelihood of the targets  $\mathbf{y}$  conditioned on the data  $\mathbf{X}$  and the hyper-parameters  $\boldsymbol{\theta}$  w.r.t.  $\boldsymbol{\theta}$ :

$$\log p(\mathbf{y} | \mathbf{X}, \boldsymbol{\theta}) \propto -\mathbf{y}^\top K_\boldsymbol{\theta}^{-1} \mathbf{y} - \log |K_\boldsymbol{\theta}|.$$

Here,  $K_\boldsymbol{\theta}$  denotes the covariance matrix of the targets  $\mathbf{y}$  given the hyper-parameters  $\boldsymbol{\theta}$ , i.e.  $K_\boldsymbol{\theta} = k_0(\mathbf{X}, \mathbf{X})$ . The first term gives a measure of how well the chosen kernel represents the data, while the second term penalises complexity. Normally, this second term prevents over-fitting as the marginal likelihood naturally favors models of intermediate complexity. However, as shown in (van Amersfoort et al., 2021; Ober et al., 2021), when used together with deep neural networks the regularisation effect is not strong enough.

## Appendix B. Risk Minimisation with the Alternating Linear Scheme

Following (Grelier et al., 2019), we can formulate the risk minimisation problem from statistical learning as follows

$$\min_{f \in \mathcal{M}_r} L(f) := \min_{f \in \mathcal{M}_r} \int_{\mathcal{X}} \ell(f; x) \rho(dx), \quad (8)$$

where we choose  $\ell(f; x) = (\Phi(x) - f(x))^2$ . Furthermore, as hypothesis class we assume a manifold  $\mathcal{M}_r$  of functions with rank- $\mathbf{r}$  TT coefficient which is embedded into the infinite dimensional tensor product function space  $\mathcal{F}$ .

Due to the intractability of the integral, we compute the empirical risk for a set of realisations  $(x^{(n)}, y^{(n)} = \Phi(x^{(n)}))$  for  $n = 1, \dots, N$ , i.e.

$$L(f) \approx \frac{1}{N} \sum_{n=1}^N (y^{(n)} - f(x^{(n)}))^2. \quad (9)$$

To reduce over-fitting, we introduce regularisation, meaning that we minimise

$$\hat{L}(f) = \sum_{n=1}^N (y^{(n)} - f(x^{(n)}))^2 + \delta \|f\|_{\mathcal{F}}^2, \quad (10)$$

where  $\delta$  is a hyper-parameter and the type of regularisation is dependent on the function space  $\mathcal{F}$ . Assuming that the data is contained in a  $d$ -dimensional cube  $\Omega = (a, b)^d$ , we choose  $\mathcal{F} := H_{\text{mix}}^1(\Omega)$ , which is defined as the tensor product  $\bigotimes_{i=1}^d H^1((a, b))$ . Here,  $H^1((a, b))$  denotes the Sobolev space of square integrable functions on  $(a, b)$  whose weak derivative is again square integrable. By penalizing large derivative values in this way, we impose smoothness on the learned function. This regularization in mixed Sobolev Spaces for TTs has been used successfully already, e.g. in (Oster et al., 2022).

Each function  $f \in \mathcal{M}_r$  can be parametrised by a list of tensors  $(V^{(i)})_{i=1}^d$ . Furthermore, by the tensor structure and Parseval's identity we have  $\|f(\cdot; (V^{(i)})_i)\|_{\mathcal{F}}^2 = \|(V^{(i)})_i\|_F^2$ , where  $\|(V^{(i)})_i\|_F$  denotes the Frobenius-norm of the full tensor corresponding to  $(V^i)_i$ . Hence, we arrive at the finite dimensional minimisation problem

$$\begin{aligned} & \min_{(V^{(i)})_i} \hat{L}((V^{(i)})_i) = \\ & \min_{(V^{(i)})_i} \frac{1}{N} \sum_{n=1}^N (y^{(n)} - f(x^{(n)}; (V^{(i)})_i))^2 \\ & + \delta \|((V^{(i)})_i)\|_F^2, \end{aligned} \quad (11)$$

over all component tensors  $V^{(i)}$ . We can further exploit the multi-linear model structure by using an alternating optimisation method. The idea is to fix all but one component tensor  $V^{(j)}$  and sweep back and forth over the tensor network, sequentially performing

$$\min_{V^{(j)}} \frac{1}{N} \sum_{n=1}^N (y^{(n)} - f(x^{(n)}; (V^{(i)})_i))^2 + \delta \|V^{(j)}\|_F^2, \quad (12)$$

where  $V^{(i)}$  is fixed for all  $i \neq j$ . After the  $j$ -th component has been optimised it is orthogonalised using a QR-decomposition where the non-orthogonal part is shifted to the next component in order to ensure stability of the method. This procedure is known as the Alternating Linear Scheme (ALS) and was introduced in (Holtz et al., 2011). It is a method specific to the TT format, due to the latter’s linearised tree structure. Hence, we have relaxed the non-convex learning problem such that only a sequence of linear systems has to be solved. Additionally, regularisation is performed naturally by suitable choice of orthonormal basis functions and penalisation of the Frobenius-norm.

## Appendix C. Specifics for Numerical Experiments

### C.1. Data Sets

Three synthetic and six real-world data sets with differing feature dimensionality are used for evaluating model performance.

For the first data set called Grid we assume  $\mathcal{X} = [0, 1]^2$  and set the true function to  $\Phi(x) = \sum_{i=1}^2 \sin(2\pi i x_i) + \epsilon$  with  $\epsilon \sim \mathcal{N}(0, 0.1)$ . Furthermore, we discretise the domain by an equidistant grid with 256 vertices and evaluate  $\Phi$  on this grid, resulting in 65 536 total data points.

The second data set is known as Friedman data set (Friedman, 1991). Here, we assume  $\mathcal{X} = [0, 1]^5$  with the data generating measure  $\rho = \otimes_{i=1}^5 \mathcal{U}(0, 1)$  and target function  $\Phi(x) = 10 \sin(\pi x_1 x_2) + 20(x_3 - 0.5)^2 + 10x_4 + 5x_5$ . We draw an overall number of 100 000 data points from  $\rho$ .

The third data set represents a high dimensional sine function with a reasonable amplitude, hence the name HighDimSine. We let  $\mathcal{X} = [0, 1]^{30}$ . Thus, we set  $\rho = \otimes_{i=1}^{30} \mathcal{U}(0, 1)$  and sample 100 000 data points. This time the target function is given by  $\Phi(x) = \sum_{i=1}^{30} \sin(\pi x_i)$ .

Furthermore, we test on the six data sets from the UCI Machine Learning Repository. Namely, the Physicochemical Properties of Protein Tertiary Structure, KEGG Metabolic Relation Network (Directed), Kin40K, SkillCraft1 Master, Elevators and Housing data set.

In order to conduct experiments, we split each data set into 70% training, 15% validation and 15% test data, where we use the training set in conjunction with early stopping on the validation set.

### C.2. Tensor-Train Kernel Learning Hyper-parameters

The budgets for pre-training, hyper-parameter optimisation and test set evaluation are 20, 50 and 100 epochs respectively. For all instances we use early stopping on the validation set.

All hyper-parameters for Tensor-Train kernel learning and their respective search spaces are:

- Tensor-Train ranks:  $r \sim \mathcal{U}(2, 15)$ ,
- $H^1$  polynomial degree:  $J \sim \mathcal{U}(2, 14)$ ,
- ALS regularisation coefficient:  $\delta_1 = 10^b$ ,  $b \sim \mathcal{U}(\log(1 \times 10^{-10}), \log(1 \times 10^{-1}))$
- orthogonalisation of TT after pre-training with equal probability

- latents dimensionality:  $L \sim \mathcal{U}(1, d)$
- number of inducing points:  $M \sim \mathcal{U}(10, 1000)$
- TT regularisation during end-to-end training with equal probability
- TT regularisation coefficient during end-to-end training:  $\delta_2 = 10^b$ ,  $b \sim \mathcal{U}(\log(1 \times 10^{-10}), \log(1 \times 10^{-1}))$
- initial TT learning rate:  $\eta_1 = 10^b$ ,  $b \sim \mathcal{U}(\log(1 \times 10^{-5}), \log(1 \times 10^{-1}))$
- initial RBF hyper-parameters learning rate:  $\eta_2 = 10^b$ ,  $b \sim \mathcal{U}(\log(1 \times 10^{-4}), \log(1 \times 10^{-1}))$
- initial VI related hyper-parameters learning rate:  $\eta_3 = 10^b$ ,  $b \sim \mathcal{U}(\log(1 \times 10^{-3}), \log(1 \times 10^{-1}))$
- data batch size:  $S \sim \mathcal{U}(4, 1024)$

### C.3. Baselines

In this part, we present all baseline models together with their respective hyper-parameters and search spaces.

#### C.3.1. CANONICAL-POLYADIC KERNEL LEARNING

In order to take the results from (Konstantinidis et al., 2021) into consideration, we implement the Canonical-Polyadic model in PyTorch and use it as feature extractor. Similarly to the TTKL training, we also use different learning rates for core tensors, GP kernel hyper-parameters and VI related hyper-parameters. However, due to the structure of the Canonical-Polyadic model, pre-training with ALS is not possible, since the orthogonalisation procedure required for ALS cannot be performed. Therefore, we train by use of ADAM (Kingma and Ba, 2015), as suggested in (Konstantinidis et al., 2021). The budgets for hyper-parameters optimisation and test set evaluation are set to 50 and 1000, due to slower convergence. Early stopping is used on the validation set.

Hyper-parameters and respective search spaces used in the random search:

- Canonical-Polyadic rank:  $r \sim \mathcal{U}(2, 15)$ ,
- polynomial degree:  $J \sim \mathcal{U}(2, 20)$ ,
- latents dimensionality:  $L \sim \mathcal{U}(1, d)$
- number of inducing points:  $M \sim \mathcal{U}(10, 1000)$
- CP regularisation during end-to-end training with equal probability
- CP regularisation coefficient during end-to-end training:  $\delta_2 = 10^b$ ,  $b \sim \mathcal{U}(\log(1 \times 10^{-10}), \log(1 \times 10^{-1}))$
- initial CP learning rate:  $\eta_1 = 10^b$ ,  $b \sim \mathcal{U}(\log(1 \times 10^{-5}), \log(1 \times 10^{-1}))$

- initial RBF hyper-parameters learning rate:  $\eta_2 = 10^b$ ,  
 $b \sim \mathcal{U}(\log(1 \times 10^{-4}), \log(1 \times 10^{-1}))$
- initial VI related hyper-parameters learning rate:  $\eta_3 = 10^b$ ,  
 $b \sim \mathcal{U}(\log(1 \times 10^{-3}), \log(1 \times 10^{-1}))$
- data batch size:  $S \sim \mathcal{U}(4, 1024)$

### C.3.2. DEEP NEURAL NETWORK

To test against a deep learning model, we choose a variant of a pre-activation ResNet (including batch normalisation layers) (He et al., 2016; He et al., 2016) with trainable residual units to account for changes in dimensionality between layers. For all data sets, we assume 4 hidden layers to be sufficient. Furthermore, we use the ADAM optimisation algorithm (Kingma and Ba, 2015) with default hyper-parameters, except for the initial learning rate. The budgets for hyper-parameter optimisation and test set evaluation are 50 and 500 epochs respectively. For all instances we use early stopping on the validation set.

Hyper-parameters and respective search spaces used in the random search:

- output dimension of first and second hidden layer:  $h_{1,2} \sim \mathcal{U}(1000, 10\,000)$
- output dimension of third hidden layer:  $h_3 \sim \mathcal{U}(100, 1000)$
- output dimension of fourth hidden layer:  $h_4 \sim \mathcal{U}(10, 100)$
- hidden layer’s non-linearity is with equal probability either ReLU, Tanh or quadratic
- data batch size:  $S \sim \mathcal{U}(4, 1024)$
- initial learning rate:  $\gamma = 10^a$ ,  $a \sim \mathcal{U}(\log(1 \times 10^{-5}), \log(1 \times 10^{-1}))$
- regularisation is either applied or not with equal probability
- regularisation coefficient:  $\delta = 10^b$ ,  $b \sim \mathcal{U}(\log(1 \times 10^{-10}), \log(1 \times 10^{-1}))$

### C.3.3. TENSOR NETWORK

As a further baseline, we perform high dimensional tensor regression using the TT decomposition. The tensorised basis consists of polynomials of degree  $J$  in the Sobolev space  $H^1$ . We constrain the rank uniformly to  $r$  and train the model by ALS using LU factorisation to solve the resulting linear systems. Moreover, we regularise the problem by the  $H^1$  norm. The budgets for hyper-parameter optimisation and test set evaluation are 25 and 100 epochs respectively. For all instances we use early stopping on the validation set. Preliminary experiments have shown that stochastic training might also be possible. The investigation of this objective is deferred to future work.

Again, hyper-parameters and their search spaces are:

- Tensor-Train rank:  $r \sim \mathcal{U}(2, 15)$ ,
- $H^1$  polynomial degree:  $J \sim \mathcal{U}(2, 14)$ ,
- regularisation coefficient:  $\delta = 10^b$ ,  $b \sim \mathcal{U}(\log(1 \times 10^{-10}), \log(1 \times 10^{-1}))$ .

#### C.3.4. GAUSSIAN PROCESS

For the GP model we use the same training methods as for TTKL. The budgets for hyper-parameter optimisation and test set evaluation are 50 and 100 epochs respectively. For all instances we use early stopping on the validation set.

The GP hyper-parameters search spaces are:

- Number inducing points:  $M \sim \mathcal{U}(100, 1000)$
- initial learning rate:  $\gamma = 10^b$ ,  $b \sim \mathcal{U}(\log(1 \times 10^{-3}), \log(1 \times 10^{-1}))$
- Batch size:  $S \sim \mathcal{U}(4, 1024)$ .

#### C.3.5. DEEP KERNEL LEARNING

For deep kernel learning the same deep architecture (and hyper-parameters) is deployed as for the DNN. However, we specify an additional hyper-parameter. Namely, we optimise over the DNN's output dimension. That is, we specify the number of features used for the subsequent GP regression. We adjust the output dimension by simply duplicating the parameters of the last layer to fit the correct number of outputs.

Exactly as in the TKL case, we train by sparse variational inference (VI) (Leibfried et al., 2021) in conjunction with ADAM. Moreover, we specify separate learning rates for the DNN, RBF kernel hyper-parameters and VI related parameters. The budgets for hyper-parameter optimisation and test set evaluation are 50 and 500 epochs respectively. For all instances we use early stopping on the validation set.

Hyper-parameters and search spaces for DKL are:

- latents dimensionality:  $L \sim \mathcal{U}(1, d)$
- number of inducing points for sparse VI:  $M \sim \mathcal{U}(10, 1000)$
- initial learning rate GP:  $\gamma_1 = 10^a$ ,  $a \sim \mathcal{U}(\log(1 \times 10^{-3}), \log(1 \times 10^{-1}))$
- initial RBF hyper-parameters learning rate:  $\gamma_2 = 10^a$ ,  
 $a \sim \mathcal{U}(\log(1 \times 10^{-4}), \log(1 \times 10^{-1}))$
- initial DNN learning rate:  $\gamma_3 = 10^a$ ,  $a \sim \mathcal{U}(\log(1 \times 10^{-5}), \log(1 \times 10^{-1}))$
- data batch size:  $S \sim \mathcal{U}(4, 1024)$