funGp: An R Package for Gaussian Process
Regression with Scalar and Functional Inputs

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Abstract

This article introduces funGp, an R package which handles regression problems involving multiple scalar and/or functional inputs, and a scalar output, through the Gaussian process model. This is particularly of interest for the design and analysis of computer experiments with expensive-to-evaluate numerical codes that take as inputs regularly sampled time series. Rather than imposing any particular parametric input-output relationship in advance (e.g., linear, polynomial), Gaussian process models extract this information directly from the data. The package offers built-in dimension reduction, which helps to simplify the representation of the functional inputs and obtain lighter models. It also implements an ant colony based optimization algorithm which supports the calibration of multiple structural characteristics of the model such as the state of each input (active or inactive) and the type of kernel function, while seeking for greater prediction power. The implemented methods are tested and applied to a real case in the domain of marine flooding.

Keywords: Gaussian process, metamodeling, functional inputs, computer experiments, R.

1. Introduction

Gaussian process (GP) models (Sacks, Welch, Mitchell, and Wynn 1989; Oakley and O’Hagan 2002) are one of the most popular regression methods thanks to the great flexibility they offer in the representation of complex non-linear input-output relationships, their high prediction
power, interpretability, and ability to provide both an interpolation of the data and an uncertainty quantification in the unexplored regions, e.g., Marrel, Iooss, Van Dorpe, and Volkova (2008). The GP model was first developed in spatial statistics under the name of kriging (Cressie 1990) and rapidly gained attention in the machine learning community where it was classified as a kernel machine (Rasmussen and Williams 2006). Over the years, the GP model has become popular in a wide range of applications such as nuclear safety (Bachoc, Ammar, and Martinez 2016), wind power generation (Mori and Kurata 2008), vehicle design and navigation (Chen, Dai, Wang, and Liu 2014), modeling and prediction of natural hazards (Rohmer and Idier 2012; Liu and Guillas 2017), among several others.

Here, a special focus will be given to the use of GP models as metamodeling techniques in the field of design and analysis of computer experiments, where the regression models are used as surrogates of expensive-to-run numerical models, see e.g., Yuan and Nian (2018) and Iooss and Marrel (2019). Indeed, in recent years, the use of computer codes for the study of complex systems has become a widespread practice to simulate systems at a lower resource expense owing to lack of observations, in particular regarding the input-output conditions for some natural processes (e.g., volcanic activity, earthquakes, flooding, etc.). Despite the increasing computer power over the last years, computer codes for environmental and industrial applications are often too time-consuming for direct application (e.g., for uncertainty quantification or fast prediction within an early warning system). A possible option to overcome this computational burden is to set up quick-to-evaluate mathematical emulators of those numerical codes (Forrester, Sobester, and Keane 2008; Simpson, Poplinski, Koch, and Allen 2001), based on a limited collection of model runs within the generic framework of design of computer experiments (Santner, Williams, Notz, and Williams 2003); such emulators being often called surrogate models or metamodels.

As shown in Østergård, Jensen, and Maagaard (2018) through comparison in 13 different metamodeling instances, GP metamodels are often able to outperform other popular methods, such as neural networks (Fonseca, Navarresse, and Moynihan 2003), multivariate adaptive regression splines (Friedman 1991) and support vector regression (Drucker, Burges, Kaufman, Smola, and Vapnik 1997), in terms of prediction accuracy.

In regression, a large amount of research works have addressed the case of scalar-valued inputs and outputs, e.g., (Forrester et al. 2008; Simpson et al. 2001; Santner et al. 2003). However, industrial and environmental applications often deal with complex inputs or outputs, which can be spatial, temporal, spatio-temporal i.e., functional in a more general way. As a motivating real case example, the coastal flood early warning system (FEWS) developed at Gâvres, France (Idier et al. 2021) predicts scalar output indicators of flooding (like total flooded area, maximum volume of water entering the territory, maximum water depth at a given location, etc.), based on inputs which are multiple scalar and time-varying maritime conditions (e.g., mean sea level, tide, atmospheric storm surge, and wave conditions), i.e., multiple time series (that are here sampled regularly over time). This FEWS is based on an hydrodynamic numerical model, but, to reach computation times compatible with short term forecast, metamodels had to be developed taking into account these inputs and providing the listed outputs.

Although GPs for scalar-valued inputs and outputs have been studied for almost 30 years, solving the problem of metamodeling in the functional framework (as described for the FEWS example) is still a much less developed research area (Muehlenstaedt, Fruth, and Roustant 2017). In principle one could use any GP implementation to fit a model with
functional inputs by representing each function as a vector, and then using each element of it as an individual scalar input of the model. However, this might lead to challenging and long hyperparameter optimization processes due to the large number of resulting decision variables. Furthermore, this approach disregards the functional nature of the inputs. To fill these gaps, this paper presents the package \texttt{funGp} (Betancourt, Bachoc, Klein, and Rohmer 2024) for the \textit{R} programming environment (\textit{R} Core Team 2023), developed on GitHub (https://github.com/djbetancourt-gh/funGp) and available from the Comprehensive \textit{R} Archive Network at https://CRAN.R-project.org/package=funGp. The package provides the construction of GP regression models of a single scalar output, dealing at the same time with multiple functional and scalar inputs by restricting on the situations where the functional inputs are regularly sampled time series (time series are considered for convenience of exposition; the \texttt{funGp} package can apply to other types of one-dimensional inputs as well). The \texttt{funGp} package offers dimension reduction (DR) methods which allow the projection of functional inputs onto a space of lower dimension while preserving the main statistical or geometric properties of each variable (Ramsay and Silverman 2007; Nanty, Helbert, Marrel, Péro, and Prieur 2016). Then, \texttt{funGp} provides the standard functionalities of GP models such as prediction, computation of confidence intervals, and conditional simulation. The second main contribution of \texttt{funGp} is an automatic procedure for the selection of the optimal structural parameters: (i) The state of each input (active or inactive), (ii) the DR method (e.g., PCA, B-splines), (iii) projection dimension and (iv) distance function used for each functional input, and (v) the type of kernel for the model (e.g., Gaussian, Matérn 5/2).

Table 1 provides a comparison between the functionalities of 24 classical open-source packages that either implement GP regression or support scalar-on-function regression with the GP or other types of models. This list includes the benchmark packages listed in the state of the art in functional data analysis in \textit{R} presented by Febrero-Bande and Oviedo de la Fuente (2012) and later updated in the website https://rpubs.com/moviedo/fda_usc_introduction, and also in the state of the art in software for GP regression presented in Roustant, Ginsbourger, and Deville (2012). To our best knowledge, and as we can see from our classification, \texttt{funGp} is the first open-source software that offers GP regression with scalar and functional inputs together with an automatic optimization method to set up the structural parameters of the model. We also see from this classification that many packages implementing GP regression allow for modeling observation noise. As discussed further in Section 2, the current conceptual choice of \texttt{funGp} is not to implement an observation noise model, since the intended application field is given by computer experiments.

The remainder of this paper is structured as follows. In Section 2 we provide the necessary statistical background. Then, in Section 3 we explain the four fundamental functionalities of \texttt{funGp}: Creation and diagnostic of GP regression models (Section 3.1), prediction (Section 3.2), simulation (Section 3.3) and model updating (Section 3.4). In Section 4 we show how the structural parameters of a \texttt{funGp} model can be set up manually. Then, in Section 5 we explain how to delegate the automatic structural configuration of the model to \texttt{funGp}. In Section 6 we show how the procedures of model construction and structural configuration can be sped up by means of parallelized implementations available in \texttt{funGp}. Section 7 applies \texttt{funGp} functionalities to a real case in the domain of marine flooding. Finally, Section 8 concludes. A compact overview of the package including a list of its main programming functions is provided in Appendix A.
funGp: Gaussian Process Regression with Scalar and Functional Inputs in R

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<th>Software/tool</th>
<th>Ref.</th>
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2. Statistical background

2.1. Structure of the regression problem

The funGp package considers the approximation of a real-valued function \( F_{sys} \) by means of a GP regression model. Throughout the article, we discriminate between three possible settings.
regarding the inputs of $\mathcal{F}_{\text{sys}}$: 

(a) When the function is $x \mapsto \mathcal{F}_{\text{sys}}(x)$, with $x = (x^{(1)}, \ldots, x^{(d_s)})^\top$ and $x^{(k)} \in \mathbb{R}$ for $k = 1, \ldots, d_s$, we say that there are $d_s$ scalar inputs, we call $x^{(k)}$ for $k = 1, \ldots, d_s$ a scalar input, and we call $x$ a vector of scalar inputs. For simplicity, we may also refer to $x$ as scalar inputs.

(b) When the function is $f \mapsto \mathcal{F}_{\text{sys}}(f)$, with $f = (f^{(1)}, \ldots, f^{(d_f)})^\top$ and $f^{(k)} : \mathcal{T}_k \subset \mathbb{R} \to \mathbb{R}$ for $k = 1, \ldots, d_f$, we say that there are $d_f$ functional inputs, we call $f^{(k)}$ for $k = 1, \ldots, d_f$ a functional input, and we call $f$ a vector of functional inputs. For simplicity, we may also refer to $f$ as functional inputs. We also denote by $\mathcal{F}$ the set of all $(f^{(1)}, \ldots, f^{(d_f)})^\top$ with $f^{(k)} : \mathcal{T}_k \subset \mathbb{R} \to \mathbb{R}$ for $k = 1, \ldots, d_f$.

(c) When the function is $(x, f) \mapsto \mathcal{F}_{\text{sys}}(x, f)$, we say that there are $d_s$ scalar inputs and $d_f$ functional inputs, and we use the same vocabulary as before for $x$ and $f$. In addition, we may refer to this setting as the hybrid-input case.

### 2.2. Gaussian process regression

Let us consider in the rest of Section 2 the hybrid-input case, of which the scalar- and functional-input ones are special cases. The goal is to build an approximate model for $\mathcal{F}_{\text{sys}}$ using a learning set $D = \{(x_1, f_1), \mathcal{F}_{\text{sys}}(x_1, f_1), \ldots, (x_n, f_n), \mathcal{F}_{\text{sys}}(x_n, f_n)\}$. As in Rasmussen and Williams (2006), we may call training points the elements of $D$. The GP model treats the fixed function $\mathcal{F}_{\text{sys}}$ as a realization of a Gaussian (random) process $\xi$, specified by its mean and covariance functions $\mu$ and $C$. The GP model is thus a Bayesian framework over functions. Recall that a Gaussian process $\xi : (x, f) \mapsto \xi(x, f)$ is defined by the fact that all vectors of the form $(\xi(x_1, f_1), \ldots, \xi(x_r, f_r))$ are Gaussian vectors, for any $r$-tuple of input points. Here $\xi$ is assumed to be centered ($\mu = 0$) and it is thus specified by its covariance function $C$:

$$\xi(\cdot) \sim \mathcal{GP}(0, C(\cdot, \cdot)), \quad (1)$$

where the kernel $C((x, f), (\tilde{x}, \tilde{f})) = \text{COV}(\xi(x, f), \xi(\tilde{x}, \tilde{f}))$ evaluates the covariance for any pair of input vectors $(x, f), (\tilde{x}, \tilde{f}) \in \mathbb{R}^{d_s} \times \mathcal{F}$. For instance, if $(x, f) = (\tilde{x}, \tilde{f})$, then $C((x, f), (\tilde{x}, \tilde{f}))$ is large and corresponds to perfect correlation. Otherwise, the covariance tends to zero as the distance between the two inputs increases. Since funGp currently does not model observation noise (nor local discontinuities similar to a nugget model in geostatistics, see Chiles and Delfiner 2009), the only admissible covariance functions here are those which are continuous at the diagonal.

One of the main benefits of GP models lies in the tractability of posterior distributions, which constitute the core mechanism to perform prediction and simulation. In the GP model, predicting the value of $\mathcal{F}_{\text{sys}}$ at a set of unobserved test points amounts to conditioning $\xi$ on the learning set $D$. Given $\xi(x_1, f_1) = \mathcal{F}_{\text{sys}}(x_1, f_1), \ldots, \xi(x_n, f_n) = \mathcal{F}_{\text{sys}}(x_n, f_n)$, $\xi$ is a GP with posterior mean values at $(X_*, F_*) = (x_{s1}, \ldots, x_{sn_s}, f_{s1}, \ldots, f_{sn_s})$ given by the vector:

$$C((X_*, F_*), (X, F)) C((X, F), (X, F))^{-1} y \quad (2)$$
and posterior covariance matrix at the same \( n_s \) points given by

\[
C((\mathbf{x}, \mathbf{F}_s), (\mathbf{x}, \mathbf{F}_s)) - C((\mathbf{x}, \mathbf{F}_s), (\mathbf{X}, \mathbf{F})) C((\mathbf{X}, \mathbf{F}), (\mathbf{X}, \mathbf{F}))^{-1} C((\mathbf{X}, \mathbf{F}), (\mathbf{x}, \mathbf{F}_s)).
\]

(3)

In the expressions above, \( \mathbf{y} = (\mathcal{F}_{\text{sys}}(x_1, f_1), \ldots, \mathcal{F}_{\text{sys}}(x_n, f_n))^\top \), \( C((\mathbf{X}, \mathbf{F}_s), (\mathbf{x}, \mathbf{F}_s)) \) is the \( n \times n_s \) prior covariance matrix between the observed values and the values to be predicted, \( C((\mathbf{x}, \mathbf{F}_s), (\mathbf{X}, \mathbf{F})) = C((\mathbf{X}, \mathbf{F}_s), (\mathbf{X}, \mathbf{F}))^\top \), \( C((\mathbf{X}, \mathbf{F}), (\mathbf{X}, \mathbf{F})) \) is the \( n \times n \) prior covariance matrix of the observed values and \( C((\mathbf{X}, \mathbf{F}_s), (\mathbf{x}, \mathbf{F}_s)) \) is the \( n_s \times n_s \) prior covariance matrix of the values to be predicted. These formulas can be found for instance in Rasmussen and Williams (2006). We emphasize that the posterior is given by noiseless observations \( \xi(x_1, f_1) = \mathcal{F}_{\text{sys}}(x_1, f_1), \ldots, \xi(x_n, f_n) = \mathcal{F}_{\text{sys}}(x_n, f_n) \), although incorporating an observation noise is common in statistical models (see also Table 1). The reason for treating the noiseless case here is that the main application field of \texttt{funGp} is constituted by computer experiments, where the noiseless setting is predominant.

The covariance function \( C \) is determined by the structural parameters listed in Section 1 (the state of each scalar and functional input, the DR method, projection dimension and distance function used for each functional input, and the type of kernel function for the model). For a given choice of these structural parameters, \( C \) is decomposed as a variance \( \sigma^2 \) times the tensor product of a correlation function for the scalar inputs and a correlation function for the functional inputs:

\[
C((x, f), (\bar{x}, \bar{f})) = \sigma^2 R(x - \bar{x}; \theta_s) R(f - \bar{f}; \theta_f),
\]

where \( \theta_s \) and \( \theta_f \) are the length scale correlation parameters, on which more details are provided in Section 4.4. Since we address the noiseless setting as discussed above, there is no noise variance parameter to estimate. In the covariance expression, the product between a correlation function for the scalar inputs and another for the functional inputs means that we consider separability between the scalar and functional inputs. Allowing also for more general covariances, non-separable between the scalar and functional inputs, could be an option for future developments, even though separability is relatively standard in GP modeling for computer experiments (Roustant et al. 2012; Sacks et al. 1989), and we obtain very satisfying prediction performances in the flooding real case of Section 7.

For the correlation function \( R \), the three possibilities in \texttt{funGp} are the Gaussian, Matérn 5/2 and Matérn 3/2 kernels, which are detailed in Section 4.1. For any choice of the structural parameters, the covariance parameters \( \sigma^2, \theta_s \) and \( \theta_f \) (a.k.a. hyperparameters) are selected by optimization of the likelihood computed on the learning set. The likelihood expression is

\[
\frac{1}{(2\pi)^{n/2}\det(C((\mathbf{X}, \mathbf{F}), (\mathbf{X}, \mathbf{F})))^{1/2}} \exp\left(-\mathbf{y}^\top C((\mathbf{X}, \mathbf{F}), (\mathbf{X}, \mathbf{F}))^{-1}\mathbf{y}\right).
\]

In \texttt{funGp}, it is numerically optimized by the L-BFGS-B algorithm (Byrd, Lu, Nocedal, and Zhu 1995) with the \texttt{optim()} R function (R Core Team 2023), allowing for constraints on the optimization variables. We remark that Leave-One-Out based selection of the hyperparameters is also possible (Bachoc 2013; Rasmussen and Williams 2006; Zhang and Wang 2010), but is not implemented in \texttt{funGp} as of now. Complementary information on our modeling approach can also be found in Betancourt, Bachoc, Klein, Idier, Pedreros, and Rohmer (2020b).
2.3. Optimization of the structural parameters

As explained above, a core feature of funGp is the possibility to optimize the structural parameters of the regression model to achieve better predictive performance. So far, for GP models, most research has exclusively focused on variable selection, meaning the optimization of the state of the input variables (Marrel et al. 2008; Lee and Park 2017; Ben Salem, Bachoc, Roustant, Gamboa, and Tomaso 2019). This task is typically performed through sensitivity analysis, which seeks to apportion the variation in the output variable to variations in the input variables (Saltelli, Tarantola, Campolongo, and Ratto 2004). By contrast, funGp treats the structural configuration of the model as a combinatorial optimization task, where each decision variable is linked to one structural parameter and the predictive power of the model is used as the objective function. By now, funGp offers structural optimization based on the well known ant colony optimization (ACO) algorithm (Dorigo and Gambardella 1997), which has been recognized as one of the most successful research lines in the area of swarm intelligence (Bonabeau, Dorigo, and Theraulaz 1999) and is always listed among the preferred metaheuristic optimization techniques (Blum 2005). We validated the ability of our ACO-based algorithm to find high performing model configurations in more than a dozen instances. The results obtained in various of those tests can be found in Chapters 3–5 of Betancourt (2020). We describe the functioning of this algorithm more in depth in Section 5.

3. Base funGp functionalities

In this section, we start by explaining the four fundamental procedures available in funGp: (i) Creation and diagnostic of regression models, (ii) prediction of the output at unobserved input points, (iii) simulation of trajectories from the underlying GP linked to any funGp model, and (iv) update of an existing model. The workflow for each of these four functionalities is illustrated through a follow-along example which considers the following analytic black-box function to regress:

\[ G : [0,1]^2 \times F \rightarrow \mathbb{R}, \]
\[ (x, f) \mapsto x^{(1)} + 2x^{(2)} + 4 \int_0^1 tf^{(1)}(t) \, dt + \int_0^1 f^{(2)}(t) \, dt, \]

with \( x = (x^{(1)}, x^{(2)})^\top \) the scalar inputs, \( f = (f^{(1)}, f^{(2)})^\top \) the functional inputs, and \( F \) the set of pairs of continuous functions from \([0,1]\) to \( \mathbb{R} \). This function corresponds to the first analytic example in Muehlenstaedt et al. (2017), and is accessible in funGp through \texttt{fgp BB3}(). By construction, function \( G \) has the first scalar and second functional input of the same importance, whereas the second scalar input and the first functional input are comparably influential but are more important than the first scalar and the second functional input. All the examples of the paper were executed on a 3.1 Ghz Intel Core i5 iMac personal computer with 16 GB RAM and 6 cores.

3.1. Create a funGp model

Let us start by creating a model. To do so, we must first put the input and output data in a suitable format. The scalar inputs should be provided as a matrix or data.frame. The functional inputs should be provided as a list of matrices, one per functional input. When the data set contains inputs of the form \( x_1, f_1, \ldots, x_n, f_n \) (see Section 2), the scalar input
matrix is $n \times d_s$, with line $\ell$ containing the vector $x_\ell = (x_\ell^{(1)}, \ldots, x_\ell^{(d_s)})$. The functional input list is of size $d_f$. In this list, the matrix corresponding to the $j$-th functional input has size $n \times N^{(j)}$ and its line $\ell$ contains the vector $(f_\ell^{(j)}(t_1^{(j)}), \ldots, f_\ell^{(j)}(t_{N^{(j)}}^{(j)}))$, where $t_1^{(j)}, \ldots, t_{N^{(j)}}^{(j)} \in \mathcal{T}_j$ are the time instants at which the input function $f_\ell^{(j)}$ is measured. Note that the input functions can thus also be thought of as time series inputs. Note also that the time instants should be equispaced, as integrals (see Section 4.4) related to the $j$-th functional input are numerically implemented as averages over the $N^{(j)}$ time instants. See also the discussion at the end of Section 4.

The output should be provided as an array or single-column matrix. Here, we will use synthetic data based on the analytic case defined above, which involves two scalar inputs and two functional inputs. To generate the scalar input data, we set up a factorial design over $[0,1]^2$. For the functional inputs, we assume that $f^{(1)}$ is measured at 10 time instants, $f^{(2)}$ at 22, and then we sample all the values of each function randomly from the standard uniform $U(0,1)$. We use an arbitrary number of 25 training points for this example.

```r
R> library("funGp")
R> set.seed(100)
R> n.tr <- 25
R> sIn <- expand.grid(x1 = seq(0, 1, length = sqrt(n.tr)),
+ x2 = seq(0, 1, length = sqrt(n.tr)))
R> fIn <- list(f1 = matrix(runif(n.tr * 10), ncol = 10),
+ f2 = matrix(runif(n.tr * 22), ncol = 22))
R> sOut <- fgp_BB3(sIn, fIn, n.tr)
R> m1 <- fgpm(sIn = sIn, fIn = fIn, sOut = sOut)
** Presampling...
** Optimising hyperparameters...

** Hyperparameters done!
```

The optimization of hyperparameters in `funGp` is handled by the `optim()` function from the R `stats` package (R Core Team 2023), which by default performs minimization. In the output above, "final value" refers to the negative logarithm of the likelihood of Section 2.2 after its numerical minimization. We refer to Section 7.3 for more information on the convergence monitoring of the optimization. Here the estimated covariance parameters are one variance, two scalar length scales and two functional length scales (see Section 4.4). They can be retrieved from the `kern` slot of `m1` (in that order) as follows.

```r
R> c(m1@kern@varHyp, m1@kern@s_lsHyps, m1@kern@f_lsHyps)
[1] 1.640431 2.000000 2.000000 2.580354 3.037035
```
It also should be noted that the function \texttt{fgpm()} does not take any input of type \texttt{formula}, although it provides a fitted regression model. This is in contrast, for instance, with the creation of linear models with the function \texttt{lm()} in R. The reason for this difference is that linear models are parametric and require the user to pre-specify regression functions that are linearly combined for prediction, for instance monomial functions of the inputs. In contrast, centered GPs provide a non-parametric regression model, where the predictions are fully data-driven and not restricted to combinations of user-specified functions. Note that if non-centered GPs were considered, it would be possible to select the mean functions as linear combinations of pre-specified functions, which could be done with a \texttt{formula} input to the function \texttt{fgpm()}. This would correspond to using universal Kriging equations instead of simple Kriging ones (see for instance Roustant \textit{et al.} 2012) in Section 2. In this case, the predictions would still be non-parametric, but with a parametric complementary component. This may be a topic of future extension of the \texttt{funGp} package.

The output of \texttt{fgpm()} is an object of class "\texttt{fgpm}" representing the fitted model. The next examples in Section 3 will reuse this model to illustrate other functionalities of the package. A first diagnostic procedure of the model can be made by calling

\texttt{R> plot(m1)}

which will display a calibration plot based on the leave-one-out (LOO) predictions, as shown in Figure 1. For a design with $n_{tr}$ points, LOO consists in building the model $n_{tr}$ times, each using a different set of $n_{tr} - 1$ points for training and computing the prediction at the ignored point. The plot will also display the LOO cross-validated squared correlation coefficient $Q^{2}_{\text{loocv}}$ as a measure of the external prediction capability of the model:

$$Q^{2}_{\text{loocv}} := 1 - \frac{\sum_{i=1}^{n_{tr}} (y_i - \hat{y}_{i,-i})^2}{\sum_{i=1}^{n_{tr}} (y_i - \bar{y})^2},$$

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{model_diagnostic.png}
\caption{Model diagnostic by leave-one-out cross-validation.}
\end{figure}
funGp: Gaussian Process Regression with Scalar and Functional Inputs in R

with \((y_i)_{i=1,\ldots,n_t}\) the vector of observed output values \((y_i = F_{sys}(x_i, f_i))\) in the hybrid-input case), \(\bar{y}\) the average of that vector and \(\hat{y}_{i-\bar{y}}\) the LOO prediction of \(y_i\). Since the classical LOO procedure might be time-expensive, *funGp* implements the virtual LOO formulas (Dubrule 1983) that require a single model training.

The main features of the model are printed by means of the `summary()` method:

```R
> summary(m1)

Gaussian Process Model____________________________________
* Scalar inputs: 2
* Functional inputs: 2

<table>
<thead>
<tr>
<th>Input</th>
<th>Orig. dim</th>
<th>Proj. dim</th>
<th>Basis</th>
<th>Distance</th>
</tr>
</thead>
<tbody>
<tr>
<td>F1</td>
<td>10</td>
<td>3</td>
<td>B-splines</td>
<td>L2_bygroup</td>
</tr>
<tr>
<td>F2</td>
<td>22</td>
<td>3</td>
<td>B-splines</td>
<td>L2_bygroup</td>
</tr>
</tbody>
</table>

* Total data points: 25
* Trained with: 25
* Kernel type: matern5_2
* Convergence: 0
* NegLogLik: 2.8411
* Hyperparameters:
  -> variance: 1.6404
  -> length-scale:
    ls(X1): 2.0000
    ls(X2): 2.0000
    ls(F1): 2.5804
    ls(F2): 3.0370

__________________________________________________________
```

The field Proj. dim relates to the possibility of performing DR (dimension reduction) on the functional inputs. Recall from the introduction that DR allows the projection of a functional input of dimension \(q\) onto a space of lower dimension \(p\) while preserving the main statistical or geometric properties of the variable. This process typically leads to \(p \ll q\), which improves the tractability and processing speed of the model. By default, the `fgpm()` function sets \(p = 3\) for every functional input. However, the user is allowed to pick custom projection dimensions and also not to project some inputs. The projection method used for each input is indicated under the field Basis; at this point, the implemented options are B-splines and PCA. Other methods might be integrated in the near future. The details on the choice of the projection method and dimension are discussed in Sections 4.2 and 4.3, respectively.

3.2. Predict using a funGp model

Now let us use our model to make predictions. To do so, we must prepare the input data corresponding to the coordinates at which the output is to be estimated. The inputs should have the same format as used for creating the model with the `fgpm()` function in Section 3.1.
The scalar inputs should be provided as a matrix or data.frame and the functional inputs should be provided as a list of matrices, one per functional input. This time, each row of an input matrix must correspond to a prediction point. For the example, we generate prediction points in a similar way as the training points, i.e., the scalar inputs from a factorial design over $[0,1]^2$ and the functional values randomly sampled from the $U(0,1)$ distribution.

```R
n.pr <- 100
sIn.pr <- as.matrix(expand.grid(x1 = seq(0, 1, length = sqrt(n.pr)),
  + x2 = seq(0, 1, length = sqrt(n.pr))))
fIn.pr <- list(f1 = matrix(runif(n.pr * 10), ncol = 10),
  + f2 = matrix(runif(n.pr * 22), ncol = 22))
```

Predictions can be requested by passing a funGp model and the prediction points to the predict() function. For instance, we can use the model m1 defined in Section 3.1.

```R
m1.preds <- predict(m1, sIn.pr = sIn.pr, fIn.pr = fIn.pr)
t(data.frame(row.names = names(m1.preds), Length = lengths(m1.preds)))
```

<table>
<thead>
<tr>
<th>mean</th>
<th>sd</th>
<th>lower95</th>
<th>upper95</th>
</tr>
</thead>
<tbody>
<tr>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
</tr>
</tbody>
</table>

The output of predict() is mainly a list containing the estimated mean, standard deviation, and limits of the 95% confidence intervals for the output at the prediction points. These confidence intervals are meant to contain the values of the Gaussian process $\xi$ which realization is the black box function $F_{sys}$. Their radii are equal to an appropriate Gaussian quantile times the posterior standard deviation presented in Section 2, with the covariance parameters fixed to their maximum likelihood estimates. Hence these intervals do not take the uncertainty on the covariance parameters into account, which is common for GP models that do not use Bayesian priors on these covariance parameters. Note that we consider a statistical model with no observation noise on $F_{sys}$, so that the confidence intervals are also predictive intervals, in the sense that they also contain the value of a new observation with a certain probability, provided that the GP model is adequate. In practice, the estimated mean of a GP model is used as the prediction of the output, while the standard deviation is often interpreted as a measure of the local error of the prediction. Actually the object returned by the predict() method on a funGp object has S3 class "predict.fgpm" inheriting from "list". This allows to use the plot() method for this S3 class.

```R
plot(m1.preds)
```

The resulting plot shows the increasingly sorted mean and corresponding confidence intervals as displayed in Figure 2. This plot can be used as a diagnostic tool for the hyperparameter optimization process. In this regard, overly wide confidence intervals for a large proportion of prediction points or missing confidence intervals for points out of the training set, as shown in Figure 3, might indicate a far-from-optimal solution.

The plot() method can also be used to compare predictions against true output values. For that aim, the vector of “true” output values at the prediction points should be passed in the optional argument sOut.pr, along with the object returned by predict().

```R
plot(m1.preds)
```
funGp: Gaussian Process Regression with Scalar and Functional Inputs in R

Sorted predictions

Figure 2: Predictions of a funGp model sorted in increasing order.

Figure 3: Signs of difficulties during the hyperparameter optimization. Left: the confidence intervals are too wide. Right: some confidence intervals are too thin.

R> sOut.pr <- fgp_BB3(sIn.pr, fIn.pr, n.pr)
R> plot(m1.preds, sOut.pr = sOut.pr)

In this case, a calibration plot will be placed on top of a sorted-output plot as shown in Figure 4. The calibration plot will display the predictive squared correlation coefficient $Q^2_{\text{total}}$, corresponding to the classical coefficient of determination $R^2$ for a test sample (Nilsson, De Jong, and Smilde 1997). The ordering in the sorted-output plot will be lead by the true output vector to facilitate the comparison of different models fitting the same data.

By default the predict() function in funGp returns so-called light predictions, which include the predicted mean, standard deviation and limits of the 95% confidence intervals. Some users might be interested in full predictions, which also include the unconditional training-prediction cross-covariance matrix $K_{tp}$ and the unconditional prediction auto-covariance matrix $K_{pp}$, which can be used for verification purposes (see Appendix B). To make full predictions, it suffices to set detail = "full" when calling predict(), as shown below.

R> m1.preds <- predict(m1, sIn.pr = sIn.pr, fIn.pr = fIn.pr, detail = "full")
R> t(data.frame(row.names = names(m1.preds), Length = lengths(m1.preds)))

<p>| | | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>mean</td>
<td>sd</td>
<td>K.tp</td>
<td>K.pp</td>
<td>lower95</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>100</td>
</tr>
<tr>
<td>Length</td>
<td>100</td>
<td>2500</td>
<td>10000</td>
<td>100</td>
</tr>
</tbody>
</table>
3.3. Simulate from a funGp model

Conditional simulations in funGp are requested through `simulate()`, similarly as predictions are requested through `predict()`. The scalar inputs should be provided as a matrix or `data.frame` and the functional inputs should be provided as a list of matrices, one per functional input. Each row of an input matrix is interpreted as a point at which to provide simulations. By default, `simulate()` will perform so-called light simulations, thus returning a \( n_{\text{rep}} \times n_{\text{sm}} \) matrix, with \( n_{\text{rep}} \) replications at \( n_{\text{sm}} \) input points. For this example we use the model \( m1 \) created in Section 3.1. In addition, we take the scalar input values from a factorial design over \([0, 1]\) and the functional values randomly from \( U(0, 1) \).

\begin{verbatim}
R> n.sm <- 100
R> sIn.sm <- as.matrix(expand.grid(x1 = seq(0, 1, length = sqrt(n.sm)),
  +   x2 = seq(0, 1, length = sqrt(n.sm))))
R> fIn.sm <- list(f1 = matrix(runif(n.sm * 10), ncol = 10),
  +   f2 = matrix(runif(n.sm * 22), ncol = 22))
R> m1.sims_l <- simulate(m1, nsim = 10, sIn.sm = sIn.sm, fIn.sm = fIn.sm)
\end{verbatim}

Simulations in funGp are plotted by the `plot()` method for the S3 class "simulate.fgpm". In contrast to prediction plots, simulation plots do not have the output sorted in increasing order, but instead in the order given by the simulation coordinates specified by the user. This plot is illustrated in Figure 5.

\begin{verbatim}
R> plot(m1.sims_l)
\end{verbatim}
funGp: Gaussian Process Regression with Scalar and Functional Inputs in R

Simulations from a funGp model

Figure 5: Simulations from a funGp model.

Figure 6: Simulations from a funGp model with confidence intervals. The confidence intervals are the same as provided by the predict() function.

If requested, simulate() will return a list containing the simulated output, predicted mean, standard deviation and limits of the 95% confidence intervals at the specified input coordinates. These confidence intervals are the same as provided by the predict() function. This corresponds to a full simulation, available through the option detail = "full".

R> m1.sims_f <- simulate(m1, nsim = 10, sIn.sm = sIn.sm, fIn.sm = fIn.sm, +   detail = "full")
R> t(data.frame(row.names = names(m1.sims_f), Length = lengths(m1.sims_f)))

<table>
<thead>
<tr>
<th>sims mean sd lower95 upper95</th>
</tr>
</thead>
<tbody>
<tr>
<td>Length 1000 100 100 100 100</td>
</tr>
</tbody>
</table>

Full simulations can be plotted similarly as light ones by means of the plot() function.

R> plot(m1.sims_f)
By default, the full simulation plot will include the predicted mean and limits of the confidence intervals (Figure 6). A light plot with just the simulated output values can also be requested for full simulations by setting the argument `detail = "light"` in the `plot()` call.

### 3.4. Update of a funGp model

Models created by `funGp` can be easily modified at any time by means of the `update()` function, which incorporates the following nine updating operations:

1. Deletion of data points.
2. Substitution of data points.
3. Addition of data points.
4. Substitution of the variance hyperparameter.
5. Substitution of the vector of scalar length-scale hyperparameters.

Through `update()`, `funGp` exploits the work done for the construction of the original model, making the updating process much faster than building a new model from zero. The request of updates is illustrated below through examples based on the model `m1` created in Section 3.1.

- **Deletion and addition of data points:**
  
  Here two points are deleted.
  ```r
  R> ind.dl <- sample(1:m1@n.tot, 2)
  R> m1up <- update(m1, ind.dl = ind.dl)
  
  --------------
  Update summary
  --------------
  * Complete tasks:
    - data deletion
  
  Here five points are added.
  ```
  ```r
  R> n.nw <- 5
  R> sIn.nw <- matrix(runif(n.nw * m1@ds), nrow = n.nw)
  R> fIn.nw <- list(f1 = matrix(runif(n.nw * 10), ncol = 10),
  +   f2 = matrix(runif(n.nw * 22), ncol = 22))
  R> sOut.nw <- fgp_BB3(sIn.nw, fIn.nw, n.nw)
  R> m1up <- update(m1, sIn.nw = sIn.nw, fIn.nw = fIn.nw,
  +   sOut.nw = sOut.nw, trace = FALSE)
  ```
Above, \texttt{trace = FALSE} enables to turn off the previous display summarizing the update. A \texttt{trace} argument is also used similarly below for the functions \texttt{fgpm()} and \texttt{fgpm\_factory()}. A more extensive study shows us that the computation time cost is largely decreased when using the \texttt{update()} function. With 25 additional points, it is reduced by 50\%, with 100, it is reduced by nearly 80\%.

- **Substitution of data points:**

Here, we generate the substitute input data, then we generate substitute output data, then we generate indices for substitution, and finally we update the scalar and functional inputs and the output.

\begin{verbatim}
R> n.sb <- 2
R> sIn.sb <- matrix(runif(n.sb * m1@ds), nrow = n.sb)
R> fIn.sb <- list(f1 = matrix(runif(n.sb * 10), ncol = 10),
+                f2 = matrix(runif(n.sb * 22), ncol = 22))
R> sOut.sb <- fgp_BB3(sIn.sb, fIn.sb, n.sb)
R> ind.sb <- sample(1:(m1@n.tot), n.sb)
R> m1up <- update(m1, sIn.sb = sIn.sb, fIn.sb = fIn.sb,
+                sOut.sb = sOut.sb, ind.sb = ind.sb, trace = FALSE)
\end{verbatim}

Substituting points only from some of the data structures is also possible. We illustrate it below with first the scalar inputs and second the output.

\begin{verbatim}
R> m1up1 <- update(m1, sIn.sb = sIn.sb, ind.sb = ind.sb, trace = FALSE)
R> m1up2 <- update(m1, sOut.sb = sOut.sb, ind.sb = ind.sb, trace = FALSE)
\end{verbatim}

- **Substitution of hyperparameters:**

Here we define the variance and the scalar and functional length-scales for substitution and then we perform the update.

\begin{verbatim}
R> var.sb <- 3
R> ls_s.sb <- c(2.44, 1.15)
R> ls_f.sb <- c(5.83, 4.12)
R> m1up <- update(m1, var.sb = var.sb, ls_s.sb = ls_s.sb,
+                ls_f.sb = ls_f.sb, trace = FALSE)
\end{verbatim}

Substituting only one of the three data structures is possible as well. We illustrate it below with first the variance and second the functional length-scales.

\begin{verbatim}
R> m1up <- update(m1, var.sb = var.sb, trace = FALSE)
R> m1up <- update(m1, ls_f.sb = ls_f.sb, trace = FALSE)
\end{verbatim}

- **Re-estimation of hyperparameters:**

Here we re-estimate successively the variance, the scalar length-scales, all the length-scales and all the hyperparameters.

\begin{verbatim}
R> m1up <- update(m1, var.re = TRUE, trace = FALSE)
R> m1up <- update(m1, ls_s.re = TRUE, trace = FALSE,
+                control.optim = list(trace = FALSE))
\end{verbatim}
Above, `control.optim = list(trace = FALSE)` enables to turn off the information display regarding the covariance parameter optimization. This is used in various other instances in the sequel.

When re-estimating hyperparameters, it is possible to use the current set of estimates as starting point of the new likelihood optimization procedure, by setting the `extend` argument to `TRUE`. We illustrate this next, when re-estimating all the hyperparameters. The benefit of this is further illustrated in Section 7.3.

Finally, it is possible to request multiple update tasks in a single call to `update()`. When doing so, tasks will be performed in the following order: data deletion/substitution → data addition → hyperparameter substitution/re-estimation.

### 4. Customization of a `funGp` model

Without delving too deep in the technical details, this section covers the choice of a `funGp` model through its so-called structural parameters. This set of categorical features (e.g., type of kernel, projection method) can be given different values in order to produce alternative models departing from the same input-output data. The interested reader is referred to Betancourt et al. (2020b) for a formal and more detailed explanation of the underlying theory.

#### 4.1. Kernel family

The selection of a suitable kernel function is something that naturally comes to mind when working with GP models. At this point, `funGp` offers the possibility to choose among the Gaussian, Matérn 5/2 and Matérn 3/2 kernels. This selection can be specified in the `fgpm()` call through the argument `kertype`, which might take a value among "gauss", "matern5_2" and "matern3_2". See for instance the example below for the Gaussian kernel.

```r
m1 <- fgpm(sIn = sIn, fIn = fIn, sOut = sOut, kerType = "gauss", 
           control.optim = list(trace = FALSE))
```

By default, `fgpm()` uses the Matérn 5/2 function, popular in the machine learning and spatial statistics community (Rasmussen and Williams 2006; Stein 1999).
Each kernel is applied to a distance \( t \geq 0 \) (the distances used are described in Section 4.4 and use length scale coefficients for importance weighting). The expressions for the Gaussian, Matérn 5/2 and Matérn 3/2 kernels are given below in that order:

\[
e^{-t^2/2}, \quad \left(1 + \sqrt{5}t + \frac{5}{3}t^2\right)e^{-\sqrt{5}t}, \quad \left(1 + \sqrt{3}t\right)e^{-\sqrt{3}t}.
\]

4.2. Projection method

In earlier sections we mentioned DR, the process of reducing the dimension of the data structures in such a way and extent that the model becomes significantly more tractable while most of its prediction power is retained. A common DR approach used on functional inputs is to project each input onto a space of lower dimension. This method requires the constitution of the basis functions defining the projection space for each input, which may come from diverse families including B-splines (De Boor 1978), PCA (Jolliffe 2002) and PLS (Papaioannou, Ehre, and Straub 2019), among the most popular ones. The B-splines and PCA bases are currently implemented in funGp for the projection of functional inputs, while the PLS ones are not as of now. This option is accessible in fgpm() through the f_basType argument, which can be set to "B-splines" or "PCA". When multiple functional inputs are handled, a custom basis can be selected for each of them. This is done by passing an array with the selection for each input:

```r
R> m1 <- fgpm(sIn = sIn, fIn = fIn, sOut = sOut,
+ f_basType = c("B-splines", "PCA"), trace = FALSE,
+ control.optim = list(trace = FALSE))
```

If multiple functional inputs are provided, but a single f_basType value is specified, that selection is used for all the inputs. By default, all functional inputs use a B-splines basis. Note that PCA is performed on the vectorized functional inputs and thus no functional basis needs to be specified. We refer to the end of Section 4 for additional discussions on using PCA on the vectorized functional inputs, as opposed to a purely functional PCA in funGp.

4.3. Projection dimension

This parameter is complementary to the projection method discussed above. Ideally, the projection dimension must be set substantially lower than the original one, but not so low that too much information is lost. The selection for each input can be specified in the fgpm() call through the f_pdims argument. Possible values are all the integer numbers from 0 to the original dimension of each input, with 0 denoting no projection. If multiple functional inputs are used, an array can be used to specify the selection for each. Below is an example where the first input is not projected while the second one is projected onto a space of dimension 7.

```r
R> m1 <- fgpm(sIn = sIn, fIn = fIn, sOut = sOut, f_pdims = c(0, 7),
+ trace = FALSE, control.optim = list(trace = FALSE))
```

If multiple functional inputs are passed, but a single f_pdims value is specified, that selection is used for all the inputs. By default, all functional inputs are projected to dimension 3.
4.4. Distance for functions

Kernel-based regression models require the computation of the distance between input coordinates in order to estimate the behavior of the output. This is the case of GP models, which use such distances to compute the correlation between pairs of observations. A set of scaling factors called length-scale coefficients are normally used to quantify the rate of change of the output in terms of each input. For scenarios with only scalar inputs, the typical is to use one length-scale parameter per input, which yields the distance

$$\|x - \tilde{x}\|_{L^2, \theta_s} := \sqrt{\sum_{k=1}^{d_s} \left\| x^{(k)} - \tilde{x}^{(k)} \right\|^2 / \left( \theta_s^{(k)} \right)^2},$$

with $\| \cdot \|$ the $L^2$ norm for scalars (i.e., the absolute value), and $\theta_s = (\theta_s^{(1)}, \ldots, \theta_s^{(d_s)})^\top$ the vector of length-scale parameters for the scalar inputs. This distance corresponds to the concept of geometric anisotropy in spatial statistics (Allard, Senoussi, and Porcu 2016; Chiles and Delfiner 2009).

When dealing with functional inputs, the norm $\| \cdot \|$ must be replaced by a suitable norm for functions. Two options are currently implemented in funGp, both based on a projection of each functional input of the form:

$$\Pi^{(k)}(f^{(k)})(t) = \sum_{r=1}^{p^{(k)}} \alpha^{(k)}_r B_r^{(k)}(t),$$

with $f^{(k)}$ a curve of the $k$-th functional input, with values $f^{(k)}(t), t \in T_k$, $B_r^{(k)}$ the $r$-th basis function used for its projection, $\alpha^{(k)}_r$ the corresponding projection coefficient, and $p^{(k)}$ the projection dimension. Note that $p^{(k)}$ is typically significantly smaller than the number of time instants $N^{(k)}$ (see Section 3) and that when $p^{(k)} = N^{(k)}$, the function $f^{(k)}$ and its projection coincide at the time instants. Note that with the PCA projection method, the basis functions $B_1^{(k)}, \ldots, B_{N^{(k)}}^{(k)}$ are taken from the $N^{(k)}$ ordered principal component functions, and thus for instance $B_1^{(k)}$ does not depend on the choice of $p^{(k)}$. Recall that actually vectorial PCA is performed on the vectors of function values at the equispaced time instants, and thus the basis functions $B_1^{(k)}, \ldots, B_{N^{(k)}}^{(k)}$ are represented by their vectors of values at the time instants. In contrast, with the B-spline basis functions, for each value of $p^{(k)}$, there is a specific set of $p^{(k)}$ basis functions (selected to cover the entire domain $T_k$ approximately uniformly) and thus in particular $B_1^{(k)}$ changes when $p^{(k)}$ is changed, in general.

The first type of distance (that can also be thought of as a semi-norm) implemented for functions considers each curve as a whole element:

$$\|f - \tilde{f}\|_{G, \theta_f} := \sqrt{\sum_{k=1}^{d_f} \int_{T_k} \left( \sum_{r=1}^{p^{(k)}} \left( \alpha^{(k)}_r - \tilde{\alpha}^{(k)}_r \right) B_r^{(k)}(t) \right)^2 dt / \left( \theta_f^{(k)} \right)^2},$$

with $f$ and $\tilde{f}$ two functional input points, $T_k \subset \mathbb{R}$ the domain of $f^{(k)}$, and $\theta_f = \left( \theta_f^{(1)}, \ldots, \theta_f^{(d_f)} \right)^\top$ the vector of length-scale parameters for the functional inputs. This distance is identified in
the package as L2_bygroup, since it uses a single length-scale parameter for the group of projection terms linked to one functional input. Note that when \( p^{(k)} = N^{(k)} \) as discussed above, the integral above is simply the square L2 distance \( \int [f^{(k)}(t) - \tilde{f}^{(k)}(t)]^2 dt \), thus making (6) similar to (4). Note also that the above integral is numerically handled as an average over the \( N^{(k)} \) time instants.

The second type of distance implemented only considers the projection coefficients, and disregards the basis functions:

\[
\|f - \tilde{f}\|_{f, \hat{\theta}_f} := \sqrt{\sum_{k=1}^{d_f} \sum_{r=1}^{p^{(k)}} \left( \alpha_r^{(k)} - \tilde{\alpha}_r^{(k)} \right)^2 / \left( \hat{\theta}_f^{(k)} \right)^2},
\]

(7)

where \( \hat{\theta}_f = (\hat{\theta}_{f,r}^{(k)})_{1 \leq r \leq p^{(k)}, 1 \leq k \leq d_f} \) denotes the vector of functional length-scale coefficients. This distance is identified in the package as L2_byindex since it involves one length-scale parameter per projection index. Using this distance corresponds to a common approach, which is to first perform the projection and then use each projection coefficient as an individual scalar input of the model. Remark that if for each \( k \) the basis functions \( B^{(k)}_1, \ldots, B^{(k)}_{p^{(k)}} \) are orthonormal and \( \theta_f^{(k)} = \hat{\theta}_f^{(k)} = \cdots = \hat{\theta}_f^{(k)} \), then (6) and (7) coincide. Both types of distances have existing counterparts in the literature, see for instance Muehlenstaedt et al. (2017).

Below we show some examples on how to specify the distance for each functional input in the fgpm() call. For these examples we still consider two functional inputs, \( f^{(1)} \) and \( f^{(2)} \), with original dimensions 10 and 22, respectively:

R> n.tr <- 25
R> sIn <- expand.grid(x1 = seq(0, 1, length = sqrt(n.tr)),
  + x2 = seq(0, 1, length = sqrt(n.tr)))
R> fIn <- list(f1 = matrix(runif(n.tr * 10), ncol = 10),
  + f2 = matrix(runif(n.tr * 22), ncol = 22))
R> sOut <- fgp_BB3(sIn, fIn, n.tr)

For the first example, \( f^{(1)} \) uses the L2_byindex distance and no projection, which yields 10 length-scale coefficients. On the other hand, \( f^{(2)} \) uses the L2_bygroup distance, which yields a single length-scale coefficient regardless of the projection dimension.

R> m1 <- fgpm(sIn = sIn, fIn = fIn, sOut = sOut, f_pdims = c(0, 5),
  + f_disType = c("L2_byindex", "L2_bygroup"), trace = FALSE,
  + control.optim = list(trace = FALSE))

Now, in a second example, we use the L2_bygroup distance for both inputs, which yields 2 length-scale coefficients, regardless of the projection dimensions.

R> m1 <- fgpm(sIn = sIn, fIn = fIn, sOut = sOut, f_pdims = c(0, 5),
  + f_disType = "L2_bygroup", trace = FALSE,
  + control.optim = list(trace = FALSE))

Finally, we use the L2_byindex distance for both inputs, we use no projection for \( f^{(1)} \) and projection dimension 5 for \( f^{(2)} \). This yields a total of 10 + 5 = 15 length-scale coefficients.
If no projection is requested for some input, both distances use the original data instead of the projection coefficients, and (6) uses the identity as the matrix of discretized basis functions.

Let us conclude by discussing the assumption of having equispaced time instances for the functional inputs. In \texttt{funGp}, the distances for functional inputs, in particular in (6), rely on integrals that are numerically computed as sums, which corresponds to representing functional inputs with equispaced time instances. Many functional inputs of the application fields of \texttt{funGp}, in particular in computer experiments, are indeed represented with equispaced time instances. We remark that it is also possible to pre-process functional inputs with non-equispaced time instances to represent them with equispaced ones, for instance by constructing a functional interpolation and evaluating it at new instants. Note also that using pseudo-inputs (Snelson and Ghahramani 2005) can also be an option to obtain equispaced time instants.

Another line of development that would allow for inputs with non-equispaced, and input-point-varying, time instants, would be to perform function decomposition at the functional rather than vectorial level. More precisely, vectorial PCA would become a fully functional PCA, where each functional input point (with its specific set of time instants) would be represented as a rich decomposition into basis functions, and vectorial PCA would be performed on the obtained coefficients, see Ramsay and Silverman (2007). This would of course allow for performing predictions for new functional inputs with different time instants. It is a prospect of extension for the \texttt{funGp} package.

Finally, remark that having equispaced time instants does not allow us to benefit from Kronecker structures, as can be the case in different settings with GPs (Wilson and Nickisch 2015), since if there are \( n \) input points each represented by \( N \) time instants, we still have only \( n \) scalar observations, while there would be \( nN \) scalar observations in settings where Kronecker structures would be valuable.

5. Automatic structural configuration

The possibility to set up the structural parameters of the model naturally comes with the non-trivial question of how to choose a good structural configuration, namely: (i) the state of each input (active or inactive), (ii) the DR method (e.g., PCA, B-splines), (iii) projection dimension, (iv) distance function used for each functional input, and (v) the type of kernel for the model (e.g., Gaussian, Matérn 5/2). In this section we introduce the \texttt{funGp} model factory, a boosting feature that enables the automatic selection of structural parameters seeking for optimal predictive performance. This tool is accessible through the \texttt{fgpm}\_\texttt{factory()} function, which enables the smart exploration of the solution space composed of all the structural parameter configurations available under \texttt{fgpm()}. Remark that Bayesian methodological alternatives to our automatic selection exist in the literature (Garnett 2022, although for GPs these alternatives are not provided in a unified package, as for \texttt{funGp}, recall the discussion of Table 1). We provide some discussion in Section 8.
Figure 7: Decision network used by the ACO-Gp heuristic in funGp. Dist., Dim., and Bas., stand for the distance, projection dimension and projection method (basis family) used for each functional input. The illustration considers \(d_s\) scalar inputs, \(d_f\) functional inputs, \(n_d\) types of distances, \(n_b\) projection methods and \(n_c\) kernel functions. The projection dimension 0 denotes no projection. Any end-to-end path over the network provides a candidate model.

5.1. Ant colony based model selection

At this point, funGp performs heuristic optimization of structural parameters supported on ACO-Gp, the ant colony based algorithm presented in Betancourt, Bachoc, Klein, and Gamboa (2020a). In ACO-Gp, virtual ants go through a decision network, selecting at each node the value for one structural parameter of the model (Figure 7). Each end-to-end path in the decision network provides a structural configuration, that is a selection of levels of kernel function, projection basis and distance type, listed in Sections 4.1, 4.2 and 4.4, as well as the state of each scalar and functional input in the model (active or inactive).

The ACO-Gp algorithm is summarized as follows. Each edge in the decision network (a choice of the next node, see Figure 7) is initialized with a quantity (load) of pheromone. At each iteration, there is a population of individual ants, each of them going independently through the decision network. For each ant, each next edge is selected stochastically, an edge with higher pheromone load being selected with higher probability. Once all the ants have completed their network paths, each of these paths yields a quantitative model quality criterion (\(Q^2_{\text{loocv}}\) or \(Q^2_{\text{hout}}\), see below). To complete the iteration, the pheromone load of each edge is updated. The pheromone loads of edges that belong to ant paths with better quality criteria are increased more. At the end of all the iterations, the ACO-Gp algorithm thus provides the list of all the structural configurations corresponding to each ant paths, associated with their quality criteria. The goal of this algorithm is that some of the configurations in the list have exceptionally good quality criteria. The calculations of the edge probabilities and pheromone updates can be tuned with parameters (see below), that enable to select a tradeoff between exploration (the edge probabilities are less dependent on the pheromone loads and the pheromone loads are more similar across edges) and exploitation (the converse).

Note that both ACO-Gp and its implementation within funGp are general enough to allow an easy extension to additional levels of kernel function, projection basis and distance type, to those listed in Sections 4.1, 4.2 and 4.4, in future versions of the package.
5.2. Getting started with the funGp model factory

The examples in this section are based on the analytic black-box function $H$ from $[0, 1]^5 \times \mathcal{F}$ to $\mathbb{R}$, with $\mathcal{F}$ the set of pairs of continuous functions $f = (f^{(1)}, f^{(2)})\top$ from $[0, 1]$ to $\mathbb{R}$, defined by

$$H(x, f) = a + \left[ b_0 + b_1 x^{(1)} + b_2 x^{(1)2} + b_3 x^{(2)} + b_4 x^{(3)} \right]^2 + c x^{(2)2} x^{(5)3} \cos x^{(1)} + d \sin x^{(4)} + e \sin x^{(4)} \times \int_0^1 (1 - t) f^{(1)}(t) dt + \left[ h_0 + h_1 x^{(1)} x^{(5)} \right] \times \int_0^1 t f^{(2)}(t) dt,$$

with $a = 10$, $b_0 = -6$, $b_1 = 5/\pi$, $b_2 = -5/4\pi^2$, $b_3 = 1$, $b_4 = 4$, $c = 10(1 - 1/8\pi)$, $d = -280\pi$, $e = 840\pi$, $h_0 = 240\pi$ and $h_1 = 3\pi$.

This function is adapted from the second analytic example of Muehlenstaedt et al. (2017). It is accessible in funGp through the black-box function `fgp_BB7()`. Here we generate the scalar and functional input values in a similar way to how we did in the previous sections:

```r
R> set.seed(100)
R> n.tr <- 32
R> sIn <- expand.grid(x1 = seq(0, 1, length = n.tr^(1/5)),
+                     x2 = seq(0, 1, length = n.tr^(1/5)),
+                     x3 = seq(0, 1, length = n.tr^(1/5)),
+                     x4 = seq(0, 1, length = n.tr^(1/5))
+                     x5 = seq(0, 1, length = n.tr^(1/5)))
R> fIn <- list(f1 = matrix(runif(n.tr * 10), ncol = 10),
+              f2 = matrix(runif(n.tr * 22), ncol = 22))
R> sOut <- fgp_BB7(sIn, fIn, n.tr)
```

**Getting started.** In our first example, we make a basic call to the model factory using its default arguments:

```r
R> xm <- fgpm_factory(sIn = sIn, fIn = fIn, sOut = sOut)
```

** Initializing decision network...**

** Optimizing structural parameters...**

** Ants are done ;)**

The output of `fgpm_factory()` is an object of class "Xfgpm" containing the optimized functional model (i.e., an object of class "fgpm" as the one delivered by `fgpm()`), a statistical indicator of its prediction power, the selected structural configuration, a record of all the evaluated models, among various other types of information. For now, we focus on the @model slot storing the optimized regression model. Let us first compare this optimized model with the one produced by `fgpm()` using its default arguments. This can be done by means of the `plot()` function introduced in Section 3.1, which receives an `fgpm` object as argument. We first plot the optimized model and then create and plot a model with default `fgpm` structural configuration.
funGp: Gaussian Process Regression with Scalar and Functional Inputs in R

(a) Model produced by \texttt{fgpm\_factory()} using its default arguments.

(b) Model produced by \texttt{fgpm()} using its default arguments.

Figure 8: Calibration plot for optimized and unoptimized structural configurations.

\[
\begin{align*}
R & \text{> plot(xm@model, main = "")} \\
R & \text{> m1 <- fgpm(sIn = sIn, fIn = fIn, sOut = sOut,} \\
& \text{+ trace = FALSE, control.optim = list(trace = FALSE))} \\
R & \text{> plot(m1, main = "")}
\end{align*}
\]

The outputs of the two \texttt{plot()} calls are displayed in Figures 8a and 8b, respectively. As can be seen, right away, just by calling \texttt{fgpm\_factory()} with its default arguments, we were able to find a model of greater quality than the one obtained with the default \texttt{fgpm()} arguments. Some key points in the light of this first result are:

- While the default structural parameters implemented in \texttt{fgpm()} usually provide reasonable predictions, they are not tailored to the data set at hand. In that view, the purpose of having \texttt{fgpm\_factory()} in the package is to be able to generate high quality structural configurations tailored to any data set that \texttt{fgpm()} could handle.

- The superiority of the model delivered by \texttt{fgpm\_factory()} is exclusively fostered by the optimization of the structural parameter configuration, and is unrelated to the mechanism for the optimization of the hyperparameters. Each model evaluated by \texttt{fgpm\_factory()} is internally created by a call to \texttt{fgpm()}. Thus, the same mechanism of hyperparameter optimization is used by both functions.

Let us move on with the exploration of the \texttt{Xfgpm} object delivered by \texttt{fgpm\_factory()} using the \texttt{plot()} method.

\[
\begin{align*}
R & \text{> plot(xm, which = "diag")} \\
R & \text{> plot(xm, which = "evol")}
\end{align*}
\]

The \texttt{plot()} method has an argument \texttt{which} allowing to choose between a diagnostic plot and an evolution plot. With the default choice \texttt{which = "diag"} the plot provides information on the absolute and relative quality of the selected model (Figure 9a). The choice \texttt{which = "evol"} illustrates the evolution in quality of the explored models along the optimization (Figure 9b). In the evolution plot, the points below zero usually correspond to models whose hyperparameters were difficult to optimize. This happens sporadically during the log-likelihood optimization for GPs due to the non-linearity of the objective function, and is not an issue that affects exclusively the funGp package.
Model diagnostic by leave-one-out cross-validation

Q2loocv = 0.939

Fitness of explored models

Sel. model
All models
fitness < 0

Figure 9: Figures available for the Xfgpm model delivered by fgpm_factory().

The quality of the selected model generally increases with the number of iterations completed by the algorithm. By default, the model factory performs 15 iterations. This quantity can be customized through the setup argument of the function, defined as a list. The way to do it is by including an element named n.iter in the setup list and giving it the value of the required number of iterations.

R> set.seed(100)
R> xm <- fgpm_factory(sIn = sIn, fIn = fIn, sOut = sOut,
+   setup = list(n.iter = 30), trace = FALSE)

Figure 10 displays the output of the plot() call on the model obtained with 30 iterations.
In this case, the algorithm explored a larger number of model configurations and was able to find a model of higher quality than the one obtained with the default 15 iterations (Figure 9). In the examples above, \texttt{fgpm\_factory()} optimized the model structure for $Q_{\text{loocv}}^2$ (see Section 3). Validating against external observations is also possible. This type of optimization can be requested by specifying, through the \texttt{ind.vl} argument, the indices of the data that should be used for validation. For instance, suppose that we have the same data as in the previous example, but we want to use about 85% of the points for training and the remaining ones for validation.

\texttt{R> ind.vl <- sample(seq\_len(n.tr), 5)}
\texttt{R> xm <- fgpm\_factory(sIn = sIn, fIn = fIn, sOut = sOut, ind.vl = ind.vl, + trace = FALSE)}

With this call, the model factory trains each model using all the data except for the points specified by \texttt{ind.vl}. Once built, each model is used to predict the output at the points ignored during training, and the predictive squared correlation coefficient $Q_{\text{hout}}^2$ (Nilsson \textit{et al.} 1997) is computed. This procedure ensures homogeneity in the comparison, since all the models use the same training and validation sets. To account for the sampling noise, it is possible to define multiple pairs of training-validation sets. This option is requested to the factory by passing a \texttt{matrix} instead of an \texttt{array} through the argument \texttt{ind.vl}. Such a \texttt{matrix} should have a vector of validation indices per column, and as many columns as replicates. Next, there are 30 replicates with approximately 15% of points for validation.

\texttt{R> ind.vl <- replicate(30, sample(seq\_len(n.tr), 5))}
\texttt{R> xm <- fgpm\_factory(sIn = sIn, fIn = fIn, sOut = sOut, ind.vl = ind.vl)}

Note that the calibration plot produced by \texttt{plot()} with \texttt{which = "diag"} will always report the $Q_{\text{loocv}}^2$ statistic, regardless of whether this or the $Q_{\text{hout}}^2$ is used for the optimization of the
structural parameters. In contrast, the evolution plot produced by this same function will display the statistic used during the optimization. When validation indices are provided, the optimized model stored in the \texttt{Xfgpm} object is one trained with as many points as left once the specified validation points are removed. When multiple validation sets are specified, the optimal model is selected by first identifying the structural configuration with largest average $Q^2_{\text{hout}}$ over all the replicates, and then picking among the corresponding models the one with largest individual $Q^2_{\text{hout}}$.

5.3. Setting up the parameters of the ACO algorithm

Our model selection algorithm relies on a set of parameters typical of any ant colony based method (recall the description of the method in Section 5.1). Those parameters control the number of individuals and iterations, degree of exploration, rate of convergence, as well as the learning mechanism in the algorithm. The full list of parameters implemented in the \texttt{fgpm\_factory()} function are listed below; those marked with an asterisk (*) are explained more thoroughly in Betancourt \textit{et al.} (2020a).

**Initial pheromone load.** At the beginning of the algorithm, each edge in the decision network is assigned a quantity indicating its desirability for the ants.

- **tao0**: Initial pheromone load on links pointing out to the selection of a distance type, a projection basis or a kernel type. Default: 0.1.
- **dop.s**: Factor to control how likely it is to activate a scalar input. It operates on a relation of the type $A = \text{dop.s} \times I$, where $A$ is the initial pheromone load of links pointing out to the activation of scalar inputs and $I$ is the initial pheromone load of links pointing out to their inactivation (which is set equal to $\text{tao0}$). Default: 1.
- **dop.f**: Analogous to dop.s for functional inputs. Default: 1.
- **delta.f** and **dispr.f**: Shape parameters for the regularization function that determines the initial pheromone values on the links connecting the \texttt{L2\_byindex} distance with the projection dimension*. Default: 2 and 1.4, respectively.

**Local pheromone update.** As the ants traverse the decision network, the pheromone load of the used links is reduced to foment diversification in the structural configurations generated.

- **rho.l**: Pheromone evaporation rate*. Default: 0.1.

**Global pheromone update.** The algorithm works by iterations, each involving a given number of ants. After each iteration, an update process is performed to increase the pheromone load in the links included in the best configurations evaluated so far.

- **n.ibest**: The algorithm always reinforces the links of the best \texttt{n.ibest} ants of the current iteration; how many ants should be used? Default: 1.
- **u.gbest**: Should the links of the best ant *over all the iterations so far* be reinforced during the global pheromone update as well? Default: `FALSE`.

- **rho.g**: Learning reinforcement rate*. Default: `0.1`.

**Population factors.** The extent of the search is primarily driven by the number of iterations and ants per iteration.

- **n.iter**: Number of iterations. Each iteration involves exploration, local pheromone update, constitution of model configurations, evaluation of performance in prediction, and system feedback through global pheromone update (recall Section 5.1). Default: 15.

- **n.pop**: Number of ants (structural configurations) per iteration. Default: 10.

**Bias strength.** Ants use one of two rules to select their next node at each step. The first rule leads the ant through the link with larger pheromone load; the second rule works based on probabilities which are proportional to the pheromone load on the feasible links. The ants will randomly chose one of the two rules at each time.

- **q0**: At each step, each ant will opt for rule 1 with probability `q0*`. Default: 0.95. For larger number of input variables, we recommend to slightly reduce it to e.g., 0.90. This might promote the evaluation of each input in at least some few models.

The default values of ACO-Gp parameters in `funGp` are based on the setup used by Dorigo and Gambardella (1997) in the introductory paper of the ant colony system. Similar values have proved to deliver a good performance for ACO-based algorithms in a variety of applications (see e.g., Li, Soleimani, and Zohal 2019, or Singh, Singh, Kumar, and Biswas 2020). All the parameters listed above can be customized in a `fgpm_factory` call through the argument `setup`, which should be a list. Below is an example using arbitrary setup values.

```R
R> mysup <- list(tao0 = 0.15, dop.s = 1.2, dop.f = 1.3, delta.f = 4, +    displ.f = 1.1, rho.1 = 0.2, u.gbest = TRUE, n.ibest = 2, +    rho.g = 0.08, n.iter = 30, n.pop = 12, q0 = 0.85) R> xm <- fgpm_factory(sIn = sIn, fIn = fIn, sOut = sOut, setup = mysup, +    trace = FALSE)
```

### 5.4. Defining the solution space

By default, `fgpm_factory()` considers feasible all possible combinations of: inputs state, distance type, projection dimension, basis family, and kernel family. However, the user is allowed to modify the solution space by imposing a system of constraints. This is achieved through the `ctrains` argument, which should be provided as a list. Below is an example considering the following list of constraints:

- Keep both scalar inputs always active.

- Keep \( f^{(2)} \) always active.
• Only use the L2_byindex distance for $f(2)$.

• Input $f(2)$ should be projected onto a space of dimension 4.

• Input $f(1)$ should be projected onto a space of dimension $\leq 5$.

• Only use the B-splines projection method for $f(1)$.

• Only test the Matérn 5/2 and Gaussian kernels.

```r
R> myctr <- list(s_keepOn = c(1, 2), f_keepOn = c(2),
+               f_disTypes = list("2" = c("L2_byindex")),
+               f_fixDims = matrix(c(2, 4), ncol = 1),
+               f_maxDims = matrix(c(1,5), ncol = 1),
+               f_basTypes = list("1" = c("B-splines")),
+               kerTypes = c("matern5_2", "gauss"))
R> xm <- fgpm_factory(sIn = sIn, fIn = fIn, sOut = sOut, ctraints = myctr)
```

The consistency of the explored configurations with the specified constraints can be verified through the `@log.success@sols` slot of the `Xfgpm` object returned by `fgpm_factory()`. This slot provides a `data.frame` with all the configurations successfully built and tested. Their corresponding prediction quality indicators can be obtained from the `@log.success@fitness` slot of the `Xfgpm` object (either $Q^2_{loocv}$ or $Q^2_{hout}$, depending on the optimized statistic). This is summarized by the `summary()` function as follows.

```r
R> summary(xm)
```

### 5.5. Time based stopping condition

In practice, it might be difficult to know in advance how many iterations will be enough to find a good configuration. Hence, `fgpm_factory()` offers an alternative stopping condition based on processing time. This feature is accessible in `fgpm_factory()` through the `time.lim` argument, which should be provided in seconds.

```r
R> mysup <- list(n.iter = 2000)
R> xm <- fgpm_factory(sIn = sIn, fIn = fIn, sOut = sOut, setup = mysup,
+                     time.lim = 60)
```

** Initializing decision network...**

** Optimizing structural parameters...**

** Time limit reached, exploration stopped after 60.00 seconds.**

** Ants are done ;)**
The number of iterations was set large enough to avoid a premature stop of the algorithm. Once the time limit is reached, the algorithm will attempt to stop as soon as possible, however, an ongoing model training process (i.e., the hyperparameter optimization) will never be interrupted. Thus, the actual processing time will often slightly exceed the specified budget.

5.6. Further exploring the Xfgpm object

We close this section by discussing in more detail the information contained in the Xfgpm object delivered by fgpm_factory(). Below we provide a list of the slots included in this object.

Optimal structural configuration.

- @model: Object of class "fgpm" associated to the selected model configuration.
- @structure: data.frame indicating the selected structural configuration.
- @stat and @fitness: Type ($Q^2_{\text{loocv}}$ or $Q^2_{\text{hout}}$) and value of the performance statistic used for the optimization of the structural parameters.

Record of explored models.

- @log.success: Object of class "antsLog" with the structure, function call and performance statistic of all the models successfully built and tested during the optimization.
- @log.crashes: Object of class "antsLog" with the structure and function calls of all the models whose fgpm() function call crashed.

Exploration extent.

- @n.solspce: Number of feasible structural configurations.
- @n.explored: Number of structural configurations successfully built and tested during the optimization.

Further information.

- @details: A list containing: (i) the set of heuristic parameters used; and (ii) the series of fitness vectors over the iterations of the heuristic.
- @factoryCall: A reminder of the expression used in the fgpm_factory() call.

The @log.success slot opens the possibility for a variety of interesting analyses that could be performed after the structural configuration. Based on the @log.success@sols component, for instance, which contains a data.frame with the structural parameters used in each explored model, one could seek to identify patterns leading to high quality models. The @log.success@args component contains analogous information, but in a format that allows the easy reconstruction of any explored model. We illustrate this possibility with the following example. We start from the structural optimization using 32 training points.
R> set.seed(100)
R> n.tr <- 32
R> sIn <- expand.grid(x1 = seq(0, 1, length = n.tr^(1/5)),
+ x2 = seq(0, 1, length = n.tr^(1/5)),
+ x3 = seq(0, 1, length = n.tr^(1/5)),
+ x4 = seq(0, 1, length = n.tr^(1/5)),
+ x5 = seq(0, 1, length = n.tr^(1/5)))
R> fIn <- list(f1 = matrix(runif(n.tr * 10), ncol = 10),
+ f2 = matrix(runif(n.tr * 22), ncol = 22))
R> sOut <- fgp_BB7(sIn, fIn, n.tr)
R> xm <- fgpm_factory(sIn = sIn, fIn = fIn, sOut = sOut, trace = FALSE)

Let us now rebuild the three best models using a larger number of training points. The modelDef() function returns a parsed R code that can be evaluated to rebuild the model corresponding to an index given by the ind formal argument. This code is to be evaluated in an environment containing the required objects sIn, fIn and sOut. By default, eval() uses the global environment.

R> n.tr <- 243
R> sIn <- expand.grid(x1 = seq(0, 1, length = n.tr^(1/5)),
+ x2 = seq(0, 1, length = n.tr^(1/5)),
+ x3 = seq(0, 1, length = n.tr^(1/5)),
+ x4 = seq(0, 1, length = n.tr^(1/5)),
+ x5 = seq(0, 1, length = n.tr^(1/5)))
R> fIn <- list(f1 = matrix(runif(n.tr * 10), ncol = 10),
+ f2 = matrix(runif(n.tr * 22), ncol = 22))
R> sOut <- fgp_BB7(sIn, fIn, n.tr)
R> m1 <- eval(modelDef(xm, ind = 1, trace = FALSE,
+ control.optim = list(trace = FALSE)))
R> m2 <- eval(modelDef(xm, ind = 2, trace = FALSE,
+ control.optim = list(trace = FALSE)))
R> m3 <- eval(modelDef(xm, ind = 3, trace = FALSE,
+ control.optim = list(trace = FALSE)))

Remark that m1, m2 and m3 are rebuilt from a different data set from the original data set used to obtain them when building xm. In the case where these three models were rebuilt using the same data set as when building xm, the estimated hyperparameters could still take different values than when building xm, since the hyperparameter optimization procedures use random initializations.

The three models m1, m2 and m3 can easily be used to make predictions thanks to the get_active_in() function, which automatically removes the inactive inputs from prediction or simulation data structures based on the fgpm() arguments of the model. To illustrate this, we start by stacking the three rebuilt models and their arguments into two lists.

R> modStack <- list(m1, m2, m3)
R> argStack <- xm@log.success@args[1:3]

Then, we generate some input points for prediction.
funGp: Gaussian Process Regression with Scalar and Functional Inputs in R

Figure 11: Predictions with models based on the best three structural configurations.

R> n.pr <- 32
R> sIn.pr <- expand.grid(x1 = seq(0, 1, length = n.pr^(1/5)),
+   x2 = seq(0, 1, length = n.pr^(1/5)),
+   x3 = seq(0, 1, length = n.pr^(1/5)),
+   x4 = seq(0, 1, length = n.pr^(1/5)),
+   x5 = seq(0, 1, length = n.pr^(1/5)))
R> fIn.pr <- list(f1 = matrix(runif(n.pr * 10), ncol = 10),
+   f2 = matrix(runif(n.pr * 22), ncol = 22))

Finally, we perform predictions based on the three models with the aid of `get_active_in()`.

R> preds <- do.call(cbind, Map(function(model, args) {
+   active <- get_active_in(sIn = sIn.pr, fIn = fIn.pr, args)
+   predict(model, sIn.pr = active$sIn.on, fIn.pr = active$fIn.on)$mean
+ }, modStack, argStack))

The resulting predictions are displayed in Figure 11.

6. Parallelization in funGp

Both, the `fgpm()` and `fgpm_factory()` functions have been equipped with the ability to exploit the availability of parallel environments. Now we explain how to access this feature.

6.1. Parallelized hyperparameter optimization

In funGp, the calibration of the hyperparameters of the model is made by likelihood maximization. For GPs, this corresponds to a nonlinear optimization problem, sometimes strongly affected by the selection of the starting points. A common way to deal with this issue is to
start the optimization multiple times from different points, which prevents the stagnation at local optima. This option is accessible in \texttt{fgpm()} through the argument \texttt{n.starts}, which should be assigned an integer value corresponding to the number of starting points to use. Below is an example using 10 starting points.

\begin{verbatim}
\texttt{R> m1 <- fgpm(sIn = sIn, fIn = fIn, sOut = sOut, n.starts = 10, +   control.optim = list(trace = FALSE))}

** Presampling...**

** Optimising hyperparameters...**

** Parallel backend register not found. Multistart optimizations done in sequence.**

** Hyperparameters done!**
\end{verbatim}

Since each starting point triggers an independent optimization process, the requested task can be performed in parallel. To do so, the user must define a parallel processing cluster and then pass it to \texttt{fgpm()} through the \texttt{par.clust} argument. As a good practice, the cluster must be stopped right after finishing the requested task in order to prevent memory issues.

\begin{verbatim}
\texttt{R> cl <- parallel::makeCluster(3)}
\texttt{R> m1 <- fgpm(sIn = sIn, fIn = fIn, sOut = sOut, n.starts = 10, +   par.clust = cl, control.optim = list(trace = FALSE))}

** Presampling...**

** Optimising hyperparameters...**

** Parallel backend register found. Multistart optimizations done in parallel.**

** Hyperparameters done!**
\end{verbatim}

\texttt{R> parallel::stopCluster(cl)}

\subsection*{6.2. Parallelized structural optimization}

During a call to \texttt{fgpm\_factory()}, each time all the ants of one iteration complete a feasible path, the corresponding models are built and evaluated in an independent fashion. Thus, this task can be performed in parallel. The way to request this to \texttt{fgpm\_factory()} is identical to how it is requested to \texttt{fgpm()}. The user must define a parallel processing cluster and then pass it to \texttt{fgpm\_factory()} through the \texttt{par.clust} argument, as shown below.

\begin{verbatim}
\texttt{R> cl <- parallel::makeCluster(3)}
\texttt{R> xm.par <- fgpm\_factory(sIn = sIn, fIn = fIn, sOut = sOut, par.clust = cl)}
\texttt{R> parallel::stopCluster(cl)}
\end{verbatim}
7. Flooding real case application

7.1. The case study

Here we apply the functionalities of \texttt{funGp} to a real case that is related to the problem of setting up a rapid FEWS for coastal flooding at Gâvres, France (Idier et al. 2021). The modeling of coastal flooding at the proper resolution (metric) to predict floods in urban areas and account precisely for processes such as wave overtopping is so time consuming (i.e., the computation time is hardly smaller than the real time) that forecasting efforts are greatly hindered. To overcome this difficulty, one solution relies on the construction of metamodels by following the approach described by Betancourt et al. (2020b).

We focus here on the prediction of the maximum value of the flooded area over time (denoted $Y$). See histogram in Figure 12. Below, $df$ is the number of functional inputs.

\begin{verbatim}
R> load("FloodingCase.rda")
R> df <- length(Xf)
R> n <- nrow(Xs)
R> hist(Y, xlab = "Flooded Area (m^2)", main = "Model outputs")
\end{verbatim}

Seven offshore meteo-oceanic forcing conditions depicted in Figure 13 are considered, namely

- the tide;
- the surge;
- the significant wave height, denoted $H_{sx}$ and $H_{sy}$ (projected onto the Cartesian x- and y-axis by means of the wave direction);
- the peak period of waves;
- the wind speed at 10m height, denoted $U_{x}$ and $U_{y}$ (projected onto the Cartesian x- and y-axis by means of the wind direction).

![Figure 12: Histogram of flooded area (m^2).](image)

\begin{itemize}
  \item ...
\end{itemize}
Each time series is sampled every 10 minutes (37 time steps) over the time interval (HT − 3 hours to HT + 3 hours), with HT referring to high tide.

To ease the analysis, the time-varying offshore meteo-oceanic forcings have been decomposed as $A_0 + f_n(t)$ where $A_0$ is a constant value over time (corresponding to the tide peak, i.e., the maximum value of the tide over the considered time interval, and to the temporal average for the other variables), and $f_n(t)$ is a time-varying signal after removal of $A_0$.
In total the inputs of the GP model include seven \( f_n(t) \) time series and eight scalar inputs (the seven \( A_0 \) scalar inputs as well as one additional scalar input corresponding to the mean sea level \( msl \)). Note that the seven functional inputs are in pairwise correspondence with seven of the scalar inputs, from the above decomposition. This raises the question of whether using covariance structures that are non-separable between functional and scalar inputs could be relevant (recall from Section 2.2 that we consider separability). In fact, our prediction performances below are already very satisfying (\( Q^2 \) values above 0.95) with a separable structure between functional and scalar inputs. The exploration of a non-separable structure is thus left open for future work.

The random selection of the inputs is based on the procedure described by Idier et al. (2021, Section 2.5.2). A total number of 174 numerical experiments were performed. To showcase funGp functionalities, we use 100 numerical experiments for the training and 74 for the predictions.

### 7.2. Fitting GP models

In this example, we set up a GP model with default parameters i.e., a "matern5_2" kernel, a "L2_bygroup" distance to be used for each functional coordinates within the covariance function of the GP, and a projection using B-splines basis functions onto a 3-dimensional space.

```r
R> set.seed(100)
R> id.train <- sample(1:n, 100, replace = FALSE)
R> id.test <- (1:n)[-id.train]
R> Xs.train <- Xs[id.train, ]
R> Xs.test <- Xs[id.test, ]
R> Xf.train <- Xf.test <- list()
R> for (i in 1:df) {
+   Xf.train[[i]] <- Xf[[i]][id.train, ]
+   Xf.test[[i]] <- Xf[[i]][id.test, ]
+ }
R> Y.train <- Y[id.train]
R> Y.test <- Y[id.test]
R> m1 <- fgpm(sIn = Xs.train, fIn = Xf.train, sOut = Y.train,
+     trace = FALSE, control.optim = list(trace = FALSE))
```

We first check the predictive capability of the constructed GP model using leave-one-out cross-validation. Figure 14 compares the observed and predicted \( Y \) values and shows a very satisfactory agreement with a \( Q^2_{loocv} > 0.95 \).

```r
R> plot(m1)
```

An alternative option is to check the predictive capability by means of a dataset of independent model results (Figure 15). The predictions can be performed by using the kriging mean plotted together with the prediction intervals at 0.95.

```r
R> m1.preds <- predict(m1, sIn.pr = Xs.test, fIn.pr = Xf.test)
R> plot(m1.preds)
```
Due to the many assumptions (related to the account for 8 scalar and 7 functional inputs) to be made to parameterize the GP model, we apply the ACO-Gp algorithm (with default parameters) to automatically select the structural parameters depending on the predictive performance. The best five solutions provide $Q^2_{loocv}$ values ranging from 0.967 to 0.969, see Figure 16. The next line of code calls the model factory. The lines after it then plot the best solution and provide a summary of the 5 best ones.

```r
R> xm <- fgpm_factory(sIn = Xs.train, fIn = Xf.train, sOut = Y.train,
  + trace = FALSE)
R> plot(xm, which = "diag")
```

![Figure 14: Leave-one-out cross validation diagnostic for the flooding application case.](image1)

![Figure 15: Predictions on independent samples for the flooding application case.](image2)
**funGp**: Gaussian Process Regression with Scalar and Functional Inputs in *R*

Figure 16: Best solutions resulting from the ACO-Gp algorithm applied to the flooding case.

```r
R> summary(xm, n = 5)

State of inputs

X1 X2 X3 X4 X5 X6 X7 X8 F1 F2 F3 F4 F5 F6 F7 Kern Q2
1  x  x  x  x  x  x  x  x  x  x  x  x  x  x  gauss  0.970
2  x  x  x  x  x  x  x  x  x  x  x  x  x  x  gauss  0.968
3  x  x  x  x  x  x  x  x  x  x  x  x  x  x  gauss  0.968
4  x  x  x  x  x  x  x  x  x  x  x  x  x  x  gauss  0.968
5  x  x  x  x  x  x  x  x  x  x  x  x  x  x  gauss  0.966

Details for functional inputs

F1 D_F1 Dim_F1 Bas_F1 F2 D_F2 Dim_F2 Bas_F2 F3 D_F3 Dim_F3 Bas_F3
1  x  grp  23  PCA x idx  4  PCA x idx  3  Bspl
2  x  grp  23  PCA x idx  2  Bspl x idx  3  Bspl
3  --  --  --  --  --  --  --  x  idx  3  Bspl
4  x  grp  23  PCA x idx  3  Bspl x idx  3  Bspl
5  --  --  --  --  --  --  --  x  idx  3  Bspl

F4 D_F4 Dim_F4 Bas_F4 F5 D_F5 Dim_F5 Bas_F5 F6 D_F6 Dim_F6 Bas_F6
1  --  --  --  --  --  --  --  --  x  idx  2  Bspl
2  --  --  --  --  --  --  --  --  x  idx  2  Bspl
3  --  --  --  --  --  --  --  --  x  idx  2  Bspl
4  --  --  --  --  --  --  --  --  x  idx  2  Bspl
5  --  --  --  --  --  --  --  --  x  idx  2  Bspl

F7 D_F7 Dim_F7 Bas_F7 Kern Q2
1  x  grp  33  PCA gauss  0.970
2  x  grp  33  PCA gauss  0.968
3  x  grp  33  PCA gauss  0.968
4  x  grp  33  PCA gauss  0.968
5  x  grp  33  PCA gauss  0.966
```
7.3. Cautionary note on convergence for covariance parameter estimation

The first fitted GP model of Section 7.2 uses the default number 100 of iteration limit. Details on its optimization process can be obtained by removing both occurrences of `trace = FALSE`.

```R
m2 <- fgpm(sIn = Xs.train, fIn = Xf.train, sOut = Y.train)
```

** Presampling...

** Optimising hyperparameters...

iter 10 value 1178.686098
iter 20 value 1169.785515
iter 30 value 1169.358788
iter 40 value 1168.584676
iter 50 value 1168.117087
iter 60 value 1167.926415
iter 70 value 1167.854377
iter 80 value 1167.782569
iter 90 value 1167.236280
iter 100 value 1167.049624
final value 1167.048204
stopped after 101 iterations

The function value is the negated log-likelihood

** Hyperparameters done!

We can also display a convergence code, which is directly provided by the `optim()` function.

```R
m2@convergence
```

[1] 1

Here the code 1 means that the iteration limit has been reached, which was also pointed out when `m2` was created with the sentence `stopped after 101 iterations`.

In this case, even though the prediction performance of `m1` (created with the same iteration limit as `m2`) was satisfying in Section 7.2, the optimization process did not completely converge. To see this, we can refit the GP model with a larger iteration limit.

```R
m3 <- fgpm(sIn = Xs.train, fIn = Xf.train, sOut = Y.train,
+     trace = TRUE, control.optim = list(trace = TRUE, maxit = 500))
```

** Presampling...

** Optimising hyperparameters...
funGp: Gaussian Process Regression with Scalar and Functional Inputs in R

iter 10 value 1174.852660
iter 20 value 1173.392364
iter 30 value 1172.160197
iter 40 value 1171.341091
iter 50 value 1170.522756
iter 60 value 1169.983111
iter 70 value 1169.712522
iter 80 value 1167.982626
iter 90 value 1167.150706
iter 100 value 1167.089213
iter 110 value 1167.085926
iter 120 value 1167.083327
iter 130 value 1167.077044
iter 140 value 1166.886057
iter 150 value 1166.884338
iter 160 value 1166.883324
iter 170 value 1166.883100
iter 180 value 1166.882840
iter 190 value 1166.862427
iter 200 value 1166.855279
iter 210 value 1166.849313
iter 220 value 1166.846073
final value 1166.845922
converged

The function value is the negated log-likelihood

** Hyperparameters done!

Now it is displayed that the optimization process did converge. Furthermore, the final negated log-likelihood is smaller (better) than before. The convergence can also be confirmed by displaying the convergence code.

R> m3@convergence

[1] 0

Here the value 0 is obtained, indicating successful completion of the call to the optim() function. Hence, the default iteration limit may be too small to reach convergence, in complex settings. This can be monitored simply from the display given when calling fgpm, or from the convergence slot of a fgpm object. The user has the possibility to increase this iteration limit, by using the control.optim argument.

We conclude by illustrating the benefit of the extend argument of the update function, discussed in Section 3.4. First, let us re-estimate the hyperparameters of the fgpm model m1 of Section 7.2, without benefiting from extend = TRUE, that is, by starting the likelihood optimization from scratch.
\begin{verbatim}
R> m4 <- update(m1, var.re = TRUE, ls_s.re = TRUE, ls_f.re = TRUE,
+     extend = FALSE, trace = FALSE)

iter  10 value 1192.597779
iter  20 value 1184.610629
iter  30 value 1176.694606
iter  40 value 1176.082474
iter  50 value 1174.242786
iter  60 value 1169.484368
iter  70 value 1168.365643
iter  80 value 1167.828297
iter  90 value 1167.823696
final value 1167.621387
stopped after 101 iterations

The function value is the negated log-likelihood

As for the creation of m2 above, since we use the default number 100 of iteration limit, the
iteration limit is reached before convergence. Next, using extend = TRUE, we can restart the
likelihood optimization from the non-converged set of hyperparameter estimates of m1.

R> m5 <- update(m1, var.re = TRUE, ls_s.re = TRUE, ls_f.re = TRUE,
+     extend = TRUE, trace = FALSE)

iter  10 value 1166.974505
iter  20 value 1166.964709
iter  30 value 1166.961533
iter  40 value 1166.913197
iter  50 value 1166.906878
iter  60 value 1166.883106
iter  70 value 1166.845932
final value 1166.845921
converged

The function value is the negated log-likelihood

We see that about 70 additional iterations are sufficient for convergence (as opposed to when
starting the likelihood optimization from scratch, in which case 100 iterations are not sufficient
for convergence).

8. Closing discussion

This article introduced the R package \texttt{funGp}, which allows the treatment of regression prob-
lens involving multiple scalar and/or functional inputs, and a scalar output, through the
fairly general GP model. In addition to being one of the few packages providing functional-
input regression under this type of model, \texttt{funGp} features the ability to automatically set
the structural parameters of the model for prediction performance. So far, this is a distinctive characteristic rarely found in current open-source regression software. The package is also equipped with multiple plotting functions that enable the quick inspection of the outputs delivered by its main functions: \texttt{fgpm()}, \texttt{fgpm\_factory()}, \texttt{predict()} and \texttt{simulate()}. Moreover, parallelization has been incorporated as an efficiency booster for the procedures of model construction and structural optimization. The robustness and stability of \texttt{funGp} has been validated on numerous analytic examples and on the real life early warning application.

We envisage the extension of the package in various directions, including the addition of complementary levels of the structural parameters already available (e.g., other projection methods, distance functions and kernel functions), incorporation of additional structural parameters (e.g., type of transformation function for the output) and implementation of alternative structural optimization methods (e.g., brute-force search and Genetic Algorithms). Our code implementations were made with these potential extensions in mind and thus, the package is architected under the principle of modularity for easy extension. Further potential extensions, both in terms of methodology and implementation, are possible. It would be valuable to allow for functional outputs and for multi-dimensional functional inputs, for instance two-dimensional maps numerically given by matrices. For multi-dimensional functional inputs, tensor product splines may be considered. As discussed in Section 3, an extension of the \texttt{funGp} package from simple to universal Kriging could be investigated, hence allowing to include a trend in the GP model. Note also that the automatic structural configuration procedure of Section 5 currently aims at returning a “best” structural configuration. Since close-to-optimal predictive performances could be achieved by very different structural configurations, an interesting extension of the \texttt{funGp} package could be to carry out Bayesian model averaging of multiple structural configurations, in the spirit of Zhang and Taflanidis (2019). Sections 4.5 and 4.6 of Garnett (2022) also discuss Bayesian priors on structural configurations for GPs, together with efficient methods for exploring the posterior. In relation with this, note that the Bayesian framework on the covariance function could enable to have credible intervals that take the uncertainty on the covariance parameters into account. Another extension perspective of the \texttt{funGp} package is to allow for covariance computations for functional inputs based on $L^2$ norms applied directly to the functions rather than their projections, and weighted by functional correlation lengths. This would provide new distances, sharing commonalities with (6), but where $\theta_f^{(k)}$ would be a function. The \texttt{funGp} package is available from GitHub (https://github.com/djbetancourt-gh/funGp) and CRAN (https://CRAN.R-project.org/package=funGp). We encourage the community to make contributions in the proposed topics or any other found relevant.

Acknowledgments

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References


A. funGp at a glance

The funGp package allows the easy construction of GP regression models with special treatment of functional inputs, an aspect scarcely treated by other R packages. The package is constituted by two major modules which are described in the paragraphs that follow.

Regression. This module is responsible for the core functionalities of the package: (i) Creation and diagnostic of GP regression models, (ii) prediction at unobserved input coordinates, (iii) conditional simulation, and (iv) model updating. These procedures are implemented in the five functions listed in Table 2. All the listed functions are set up with default arguments that allow an effortless first interaction with the package. However, high degree of customization is enabled to let the users get greater control over the outputs and extend the scope of the analysis as they become familiar with the overall functioning of the package. As an example, the \texttt{fgpm()} function allows the customization of the so-called structural parameters and the definition of custom hyperparameters, among other features.

<table>
<thead>
<tr>
<th>Function</th>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>\texttt{fgpm()}</td>
<td>Function</td>
<td>Creation of GP regression models. Outputs an object of S4 class &quot;\texttt{fgpm}&quot; representing the fitted model</td>
</tr>
<tr>
<td>\texttt{predict()}</td>
<td>Method</td>
<td>Prediction at new input points. Delivers the estimated output values and corresponding uncertainty at specified points</td>
</tr>
<tr>
<td>\texttt{simulate()}</td>
<td>Method</td>
<td>Conditional simulation. Delivers the simulated output values and corresponding uncertainty at specified points</td>
</tr>
<tr>
<td>\texttt{update()}</td>
<td>Method</td>
<td>Efficient model update. Allows to add, delete or substitute data, and also to replace or re-estimate hyperparameters</td>
</tr>
<tr>
<td>\texttt{plot()}</td>
<td>Method</td>
<td>Diagnostic plot for GP model; receives an \texttt{fgpm} object as main argument</td>
</tr>
</tbody>
</table>

Table 2: Base funGp functionalities related to the class "\texttt{fgpm}".

Structural optimization. The set of functions listed in Table 2 could constitute by themselves a regression package. However, funGp goes further and incorporates the automatic calibration of structural parameters of the model, a functionality widely neglected by other regression tools. Structural optimization enables the efficient exploration of the space of possible model configurations, seeking for optimal predictive performance. Diagnostic plots are provided for a quick evaluation of the selected model and the evolution of the optimization. The main procedures of this module are implemented in the three functions listed in Table 3.

B. Manual prediction

As explained in Section 3.2, GP predictions are accessible in funGp through the \texttt{predict()} method. However, as a verification/inspection procedure, it is also possible to recreate the different data structures necessary for making the predictions manually based on the equations (2) and (3) of Gaussian conditioning. The recovery of these data structures is explained below, departing from an hypothetical model \texttt{m1} created with \texttt{fgpm()}. 

Table 3: Complementary `funGp` functionalities related to the class "Xfgpm" enabling structural optimization.

- Vector of observed output values \( y \): this vector is stored at the `@sOut` slot of the `fgpm` object associated with the model.

  ```R
  R> y <- m1@sOut
  ```

- Unconditional \( n \times n \) training-prediction cross-covariance matrix \( C((X, F), (X, F)) \) and \( n \times n \) prediction auto-covariance matrix \( C((X, F_*), (X, F_*)) \): these matrices can be obtained from a full prediction generated by `predict()`, as shown below.

  ```R
  R> preds <- predict(m1, sIn.pr = sIn.pr, fIn.pr = fIn.pr, detail = "full")
  R> K.tp <- preds$K.tp
  R> K.pp <- preds$K.pp
  ```

- Unconditional \( n \times n \) training auto-covariance matrix \( C((X, F), (X, F)) \): this matrix can be regenerated from the lower triangular matrix of its LU decomposition, stored at the `@preMats$L` slot of the model.

  ```R
  R> K.tt <- tcrossprod(m1@preMats$L)
  ```

Once the \( y \), \( K.tp \), \( K.pp \) and \( K.tt \) data structures have been recovered, the predicted output values and corresponding confidence intervals can be reproduced based on Gaussian conditioning as follows (with the two first lines providing the predictive means and variances and the next two providing the limits of the 95% confidence intervals).

  ```R
  R> y.pr <- t(K.tp) %*% solve(K.tt) %*% y
  R> v.pr <- diag(K.pp - t(K.tp) %*% solve(K.tt) %*% K.tp)
  R> ll <- y.pr - 1.96 * sqrt(v.pr)
  R> ul <- y.pr + 1.96 * sqrt(v.pr)
  ```

We remark that this way of computing GP predictions is computationally inefficient, and is only recommended for purposes of verification and examination of the involved data structures. Predictions and simulations performed through the `predict()` and `simulate()` functions in `funGp` are highly optimized for computational efficiency.
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