The GPML Toolbox version 3.5

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Abstract

The GPML toolbox is an Octave 3.2.x and Matlab 7.x implementation of inference and prediction in Gaussian process (GP) models. It implements algorithms discussed in Rasmussen & Williams: Gaussian Processes for Machine Learning, the MIT press, 2006 and Nickisch & Rasmussen: Approximations for Binary Gaussian Process Classification, JMLR, 2008.

The strength of the function lies in its flexibility, simplicity and extensibility. The function is flexible as firstly it allows specification of the properties of the GP through definition of mean function and covariance functions. Secondly, it allows specification of different inference procedures, such as e.g. exact inference and Expectation Propagation (EP). Thirdly it allows specification of likelihood functions e.g. Gaussian or Laplace (for regression) and e.g. cumulative Logistic (for classification). Simplicity is achieved through a single function and compact code. Extensibility is ensured by modular design allowing for easy addition of extension for the already fairly extensive libraries for inference methods, mean functions, covariance functions and likelihood functions.

This document is a technical manual for a developer containing many details. If you are not yet familiar with the GPML toolbox, the *user documentation* and examples therein are a better way to get started.

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1 Gaussian Process Training and Prediction

The gpml toolbox contains a single user function gp described in section 2. In addition there are a number of supporting structures and functions which the user needs to know about, as well as an internal convention for representing the posterior distribution, which may not be of direct interest to the casual user.

Inference Methods: An inference method is a function which computes the (approximate) posterior, the (approximate) negative log marginal likelihood and its partial derivatives w.r.t.. the hyperparameters, given a model specification (i.e., GP mean and covariance functions and a likelihood function) and a data set. Inference methods are discussed in section 3. New inference methods require a function providing the desired inference functionality and possibly extra functionality in the likelihood functions applicable.

Hyperparameters: The hyperparameters is a struct controlling the properties of the model, i.e.. the GP mean and covariance function and the likelihood function. The hyperparameters is a struct with the three fields mean, cov and lik, each of which is a vector. The number of elements in each field must agree with number of hyperparameters in the specification of the three functions they control (below). If a field is either empty or non-existent it represents zero hyperparameters. When working with FITC approximate inference, the inducing inputs xu can also be treated as hyperparameters for some common stationary covariances.

Hyperparameter Prior Distributions: When optimising the marginal likelihood w.r.t. hyperparameters, it is sometimes useful to softly constrain the hyperparameters by means of prior knowledge. A prior is a probability distribution over individual or a group of hyperparameters, section 7.

Likelihood Functions: The likelihood function specifies the form of the likelihood of the GP model and computes terms needed for prediction and inference. For inference, the required properties of the likelihood depend on the inference method, including properties necessary for hyperparameter learning, section 4.

Mean Functions: The mean function is a cell array specifying the GP mean. It computes the mean and its derivatives w.r.t.. the part of the hyperparameters pertaining to the mean. The cell array allows flexible specification and composition of mean functions, discussed in section 5. The default is the zero function.

Covariance Functions: The covariance function is a cell array specifying the GP covariance function. It computes the covariance and its derivatives w.r.t.. the part of the hyperparameters pertaining to the covariance function. The cell array allows flexible specification and composition of covariance functions, discussed in section 6.

Inference methods, see section 3, compute (among other things) an approximation to the posterior distribution of the latent variables f_i associated with the training cases, $i=1,\ldots,n$. This approximate posterior is assumed to be Gaussian, and is communicated via a struct post with the fields post.alpha, post.sW and post.L. Often, starting from the Gaussian prior $p(f) = \mathcal{N}(f|m,K)$ the approximate posterior admits the form

$$q(f|\mathcal{D}) = \mathcal{N}(f|\mu = m + K\alpha, V = (K^{-1} + W)^{-1}), \text{ where } W \text{ diagonal with } W_{ii} = s_i^2.$$
 (1)

In such cases, the entire posterior can be computed from the two vectors post.alpha and post.sW; the inference method may optionally also return $L = \text{chol}(\text{diag}(s) \times \text{diag}(s) + I)$. If on the other hand the posterior doesn't admit the above form, then post.L returns the matrix

 $L=-(K+W^{-1})^{-1}$ (and post.sW is unused). In addition, a sparse representation of the posterior may be used, in which case the non-zero elements of the post.alpha vector indicate the active entries.

2 The gp Function

39 %

The gp function is typically the only function the user would directly call.

It offers facilities for training the hyperparameters of a GP model as well as predictions at unseen inputs as detailed in the following help.

```
\langle gp \ function \ help \ 5b \rangle \equiv
5b
                                                                  (5a)
     1 % Gaussian Process inference and prediction. The gp function provides a
     2 % flexible framework for Bayesian inference and prediction with Gaussian
     3 % processes for scalar targets, i.e. both regression and binary
     4 % classification. The prior is Gaussian process, defined through specification
     5~\% of its mean and covariance function. The likelihood function is also
     6 % specified. Both the prior and the likelihood may have hyperparameters
     7 % associated with them.
     8 %
     9 % Two modes are possible: training or prediction: if no test cases are
     10 % supplied, then the negative log marginal likelihood and its partial
     11 % derivatives w.r.t. the hyperparameters is computed; this mode is used to fit
     12 % the hyperparameters. If test cases are given, then the test set predictive
    13 % probabilities are returned. Usage:
    14 %
    15 %
          training: [nlZ dnlZ
                                         ] = gp(hyp, inf, mean, cov, lik, x, y);
     16 % prediction: [ymu ys2 fmu fs2 ] = gp(hyp, inf, mean, cov, lik, x, y, xs);
     17 %
                 or: [ymu ys2 fmu fs2 lp] = gp(hyp, inf, mean, cov, lik, x, y, xs, ys);
    18 %
     19 % where:
    20 %
     21 %
                    struct of column vectors of mean/cov/lik hyperparameters
     22 %
                    function specifying the inference method
          inf
    23 %
          mean prior mean function
    24 % cov
                    prior covariance function
    25 %
                   likelihood function
          lik
    26 %
          X
                  n by D matrix of training inputs
           xs ns by D matrix of test inputs
ys column vector of
    27 %
                   column vector of length n of training targets
          У
    28 %
    29 %
                    column vector of length nn of test targets
    30 %
    31 %
           nlZ returned value of the negative log marginal likelihood
    32 %
           dnlZ
                    struct of column vectors of partial derivatives of the negative
    33 %
                        log marginal likelihood w.r.t. mean/cov/lik hyperparameters
     34 %
                    column vector (of length ns) of predictive output means
          ymu
    35 %
                    column vector (of length ns) of predictive output variances
           vs2
    36 %
                    column vector (of length ns) of predictive latent means
           fmu
     37 %
           fs2
                    column vector (of length ns) of predictive latent variances
    38 %
                    column vector (of length ns) of log predictive probabilities
```

```
40 %
                      struct representation of the (approximate) posterior
            post
     41 %
                      3rd output in training mode or 6th output in prediction mode
     42 %
                      can be reused in prediction mode gp(.., cov, lik, x, post, xs,..)
     43 %
     44 % See also infMethods.m, meanFunctions.m, covFunctions.m, likFunctions.m.
     45 %
     46 (gpml copyright 6a)
     \langle gpml\ copyright\ 6a \rangle \equiv
                                            (5b 10 11 18 21 23a 34 35 37 41 42 45)
6a
      1 % Copyright (c) by Carl Edward Rasmussen and Hannes Nickisch, 2015-07-13.
                                                  File automatically generated using noweb.
```

Depending on the number of input parameters, gp knows whether it is operated in training or in prediction mode. The highlevel structure of the code is as follows. After some initialisations, we perform inference and decide whether test set predictions are needed or only the result of the inference is demanded.

```
6b ⟨initializations 6b⟩≡ (5a)

1 ⟨minimalist usage 6c⟩

2 ⟨process input arguments 6d⟩

3 ⟨check hyperparameters 7a⟩
```

If the number of input arguments is incorrect, we echo a minimalist usage and return.

Set some useful default values for empty arguments, and convert inf and lik to function handles and mean and cov to cell arrays if necessary. Initialize variables.

```
\langle process\ input\ arguments\ 6d \rangle \equiv
6d
                                                                   (6b)
     1 if isempty(mean), mean = {@meanZero}; end
                                                                       % set default mean
     2 if ischar(mean) || isa(mean, 'function_handle'), mean = {mean}; end % make cell
     3 if isempty(cov), error('Covariance function cannot be empty'); end % no default
     4 if ischar(cov) || isa(cov, 'function_handle'), cov = {cov}; end
     5 cstr = cov{1}; if isa(cstr,'function_handle'), cstr = func2str(cstr); end
     6 if strcmp(cstr,'covFITC') && isfield(hyp,'xu'), cov{3} = hyp.xu; end %use hyp.xu
     7 if isempty(inf)
                                                           % set default inference method
         if strcmp(cstr,'covFITC'), inf = {@infFITC}; else inf = {@infExact}; end
     9 end
     10 if ischar(inf), inf = str2func(inf); end
                                                           % convert into function handle
     11 if ischar(inf) || isa(inf,'function_handle'), inf = {inf}; end
     12 istr = inf{1}; if isa(istr, 'function_handle'), istr = func2str(istr); end
     13 if strcmp(istr, 'infPrior')
         istr = inf{2}; if isa(istr,'function_handle'), istr = func2str(istr); end
     14
     15 end
    16 if strcmp(cstr,'covFITC')
                                                             % only infFITC* are possible
         if isempty(strfind(istr,'infFITC')==1)
     18
           error('Only infFITC* are possible inference algorithms')
     19
         end
                                       % only one possible class of inference algorithms
     21 if isempty(lik), lik = {@likGauss}; end
                                                                        % set default lik
     22 if ischar(lik) || isa(lik, 'function_handle'), lik = {lik}; end
     23 lstr = lik{1}; if isa(lstr,'function_handle'), lstr = func2str(lstr); end
```

Check that the sizes of the hyperparameters supplied in hyp match the sizes expected. The three parts hyp.mean, hyp.cov and hyp.lik are checked separately, and define empty entries if they don't exist.

Inference is performed by calling the desired inference method inf. In training mode, we accept a failure of the inference method (and issue a warning), since during hyperparameter learning, hyperparameters causing a numerical failure may be attempted, but the minimize function may gracefully recover from this. During prediction, failure of the inference method is an error.

```
7b
     \langle inference 7b \rangle \equiv
      1 try
                                                                % call the inference method
          \% issue a warning if a classification likelihood is used in conjunction with
          % labels different from +1 and -1
          if strcmp(lstr,'likErf') || strcmp(lstr,'likLogistic')
      5
            if ~isstruct(y)
      6
              uy = unique(y);
      7
              if any( uy~=+1 & uy~=-1 )
      8
                warning('You try classification with labels different from {+1,-1}')
      9
              end
     10
            end
     11
     12
          if nargin>7 % compute marginal likelihood and its derivatives only if needed
     13
            if isstruct(y)
     14
                                    % reuse a previously computed posterior approximation
              post = y;
     15
     16
              post = feval(inf{:}, hyp, mean, cov, lik, x, y);
     17
            end
     18
     19
            if nargout <= 1</pre>
     20
              [post nlZ] = feval(inf{:}, hyp, mean, cov, lik, x, y); dnlZ = {};
     21
     22
              [post nlZ dnlZ] = feval(inf{:}, hyp, mean, cov, lik, x, y);
     23
            end
     24
          end
     25 catch
          msgstr = lasterr;
```

We copy the already computed negative log marginal likelihood to the first output argument, and if desired report its partial derivatives w.r.t. the hyperparameters if running in inference mode.

Predictions are computed in a loop over small batches to avoid memory problems for very large test sets.

```
\langle compute\ test\ predictions\ 8a \rangle \equiv
8a
                                                             (5a)
     1 alpha = post.alpha; L = post.L; sW = post.sW;
     2 if issparse(alpha)
                               % handle things for sparse representations
     3 nz = alpha \sim 0;
                                                      % determine nonzero indices
       5 if issparse(sW), sW = full(sW(nz)); end
     6 else nz = true(size(alpha,1),1); end
                                                       % non-sparse representation
                                         % in case L is not provided, we compute it
     7 if isempty(L)
       K = feval(cov{:}, hyp.cov, x(nz,:));
     9 L = chol(eye(sum(nz))+sW*sW'.*K);
    11 %verify whether L contains valid Cholesky decomposition or something different
    12 Lchol = isnumeric(L) && all(all(tril(L,-1)==0)&diag(L)'>0&isreal(diag(L))');
    13 \text{ ns} = \text{size}(xs, 1);
                                                          % number of data points
    14 if strcmp(cstr,'covGrid'), xs = covGrid('idx2dat',cov{3},xs); end % expand xs
                                            % number of data points per mini batch
    15 nperbatch = 1000;
    16 \text{ nact} = 0;
                                     % number of already processed test data points
    17 ymu = zeros(ns,1); ys2 = ymu; fmu = ymu; fs2 = ymu; lp = ymu; % allocate mem
    18 while nact<ns
                              % process minibatches of test cases to save memory
    19 id = (nact+1):min(nact+nperbatch,ns);
                                                         % data points to process
    20 (make predictions 8b)
    21 nact = id(end); % set counter to index of last processed data point
    22 end
    23 if nargin<9
    24 varargout = {ymu, ys2, fmu, fs2, [], post}; % assign output arguments
    26 varargout = {ymu, ys2, fmu, fs2, lp, post};
```

In every iteration of the above loop, we compute the predictions for all test points of the batch.

```
8b
     \langle make\ predictions\ 8b \rangle \equiv
     1 kss = feval(cov{:}, hyp.cov, xs(id,:), 'diag');
                                                                     % self-variance
     2 if strcmp(cstr,'covFITC')
                                                                 % cross-covariances
         Ks = feval(cov{:}, hyp.cov, x, xs(id,:)); Ks = Ks(nz,:); % res indep. of x
     4 else
         Ks = feval(cov\{:\}, hyp.cov, x(nz,:), xs(id,:));
                                                          % avoid computation
     6 end
     7 ms = feval(mean{:}, hyp.mean, xs(id,:));
     8 N = size(alpha,2); % number of alphas (usually 1; more in case of sampling)
     9 Fmu = repmat(ms,1,N) + Ks'*full(alpha(nz,:)); % conditional mean fs|f
    10 fmu(id) = sum(Fmu, 2)/N;
                                                                  % predictive means
                   % L contains chol decomp => use Cholesky parameters (alpha,sW,L)
    11 if Lchol
    12 V = L'\(repmat(sW,1,length(id)).*Ks);
    13 fs2(id) = kss - sum(V.*V,1);
                                                              % predictive variances
                           % L is not triangular => use alternative parametrisation
    14 else
```

```
15 if isnumeric(L), LKs = L*Ks; else LKs = L(Ks); end % matrix or callback
16 fs2(id) = kss + sum(Ks.*LKs,1);
                                                        % predictive variances
17 \text{ end}
18 fs2(id) = max(fs2(id),0); % remove numerical noise i.e. negative variances
19 Fs2 = repmat(fs2(id),1,N); % we have multiple values in case of sampling
20 if nargin<9
21 [Lp, Ymu, Ys2] = feval(lik{:},hyp.lik,[], Fmu(:),Fs2(:));
22 else
23    Ys = repmat(ys(id),1,N);
24 [Lp, Ymu, Ys2] = feval(lik{:},hyp.lik,Ys(:),Fmu(:),Fs2(:));
26 lp(id) = sum(reshape(Lp, [],N),2)/N; % log probability; sample averaging
27 ymu(id) = sum(reshape(Ymu,[],N),2)/N;
                                                 % predictive mean ys|y and ..
28 ys2(id) = sum(reshape(Ys2,[],N),2)/N;
                                                                % .. variance
```

3 Inference Methods

Inference methods are responsible for computing the (approximate) posterior post, the (approximate) negative log marginal likelihood n1Z and its partial derivatives dn1Z w.r.t. the hyperparameters hyp. The arguments to the function are hyperparameters hyp, mean function mean, covariance function cov, likelihood function lik and training data x and y. Several inference methods are implemented and described this section.

```
10
     \langle infMethods.m \ 10 \rangle \equiv
     1 % Inference methods: Compute the (approximate) posterior for a Gaussian process.
     2 % Methods currently implemented include:
     3 %
     4 %
                            Exact inference (only possible with Gaussian likelihood)
           infExact
     5 %
           infLaplace
                            Laplace's Approximation
     6 %
                            Expectation Propagation
           infEP
     7 %
           infVB
                            Variational Bayes Approximation
     8 %
           infKL
                            Kullback-Leibler optimal Approximation
     9 %
    10 %
           infFITC
                            Large scale regression with approximate covariance matrix
    11 %
           infFITC_Laplace Large scale inference with approximate covariance matrix
    12 %
           infFITC_EP
                            Large scale inference with approximate covariance matrix
    13 %
    14 %
           infMCMC
                       Markov Chain Monte Carlo and Annealed Importance Sampling
    15 %
                       We offer two samplers.
    16 %
                         - hmc: Hybrid Monte Carlo
    17 %
                          - ess: Elliptical Slice Sampling
    18 %
                       No derivatives w.r.t. to hyperparameters are provided.
    19 %
    20 %
           infL00
                       Leave-One-Out predictive probability and Least-Squares Approxim.
    21 %
           infPrior
                       Perform inference with hyperparameter prior.
    22 %
    23 % The interface to the approximation methods is the following:
    24 %
    25 %
           function [post nlZ dnlZ] = inf..(hyp, cov, lik, x, y)
    26 %
    27 % where:
    28 %
    29 %
                    is a struct of hyperparameters
          hyp
    30 %
                    is the name of the covariance function (see covFunctions.m)
          COV
    31 %
         lik
                    is the name of the likelihood function (see likFunctions.m)
    32 %
                    is a n by D matrix of training inputs
    33 %
                    is a (column) vector (of size n) of targets
           У
    34 %
    35 %
                    is the returned value of the negative log marginal likelihood
          nlZ
    36 %
           dnlZ
                    is a (column) vector of partial derivatives of the negative
    37 %
                       log marginal likelihood w.r.t. each hyperparameter
    38 %
                    struct representation of the (approximate) posterior containing
    39 %
           alpha is a (sparse or full column vector) containing inv(K)*(mu-m),
    40 %
                       where K is the prior covariance matrix, m the prior mean,
    41 %
                       and mu the approx posterior mean
    42 %
                    is a (sparse or full column) vector containing diagonal of sqrt(W)
            sW
    43 %
                       the approximate posterior covariance matrix is inv(inv(K)+W)
    44 %
                    is a (sparse or full) matrix, L = chol(sW*K*sW+eye(n))
    45 %
    46 % Usually, the approximate posterior to be returned admits the form
    47 % N(mu=m+K*alpha, V=inv(inv(K)+W)), where alpha is a vector and W is diagonal;
    48 % if not, then L contains instead -inv(K+inv(W)), and sW is unused.
```

```
50 % For more information on the individual approximation methods and their
51 % implementations, see the separate inf??.m files. See also gp.m
52 %
53 (gpml copyright 6a)
```

Not all inference methods are compatible with all likelihood functions, e.g., exact inference is only possible with Gaussian likelihood. In order to perform inference, each method needs various properties of the likelihood functions, section 4.

3.1 Exact Inference

39

For Gaussian likelihoods, GP inference reduces to computing mean and covariance of a multivariate Gaussian which can be done exactly by simple matrix algebra. The program inf/infExact.m does exactly this. If it is called with a likelihood function other than the Gaussian, it issues an error. The Gaussian posterior $q(f|D) = \mathcal{N}(f|\mu, V)$ is exact.

```
\langle inf/infExact.m \ 11 \rangle \equiv
11
     1 function [post nlZ dnlZ] = infExact(hyp, mean, cov, lik, x, y)
     3 % Exact inference for a GP with Gaussian likelihood. Compute a parametrization
     4 % of the posterior, the negative log marginal likelihood and its derivatives
     5\ \% w.r.t. the hyperparameters. See also "help infMethods".
     7 (gpml copyright 6a)
     9 % See also INFMETHODS.M.
    11 if iscell(lik), likstr = lik{1}; else likstr = lik; end
    12 if ~ischar(likstr), likstr = func2str(likstr); end
    13 if ~strcmp(likstr,'likGauss')
                                                   % NOTE: no explicit call to likGauss
         error('Exact inference only possible with Gaussian likelihood');
    15 end
    16
    17 [n, D] = size(x);
    18 K = feval(cov{:}, hyp.cov, x);
                                                           % evaluate covariance matrix
    19 m = feval(mean{:}, hyp.mean, x);
                                                                  % evaluate mean vector
    20
    21 \text{ sn2} = \exp(2*\text{hyp.lik});
                                                           % noise variance of likGauss
    22 if sn2<1e-6
                                          % very tiny sn2 can lead to numerical trouble
    L = chol(K+sn2*eye(n)); sl =
                                      1; % Cholesky factor of covariance with noise
    24
       pL = -solve_chol(L,eye(n));
                                                                % L = -inv(K+inv(sW^2))
    26 L = chol(K/sn2+eye(n)); sl = sn2;
                                                                  % Cholesky factor of B
    27
         pL = L;
                                                           % L = chol(eye(n)+sW*sW'.*K)
    28 end
    29 alpha = solve_chol(L,y-m)/sl;
    31 post.alpha = alpha;
                                                     % return the posterior parameters
    32 post.sW = ones(n,1)/sqrt(sn2);
                                                      % sqrt of noise precision vector
    33 \text{ post.L} = pL;
    34
                                                  \mbox{\ensuremath{\mbox{\%}}} do we want the marginal likelihood?
    35 if nargout>1
    36  n1Z = (y-m)^* + alpha/2 + sum(log(diag(L))) + n*log(2*pi*s1)/2; % -log marg lik
    37
         if nargout>2
                                                               % do we want derivatives?
    38
                                                       % allocate space for derivatives
           dnlZ = hyp;
```

```
40     for i = 1:numel(hyp.cov)
41         dnlZ.cov(i) = sum(sum(Q.*feval(cov{:}, hyp.cov, x, [], i)))/2;
42     end
43     dnlZ.lik = sn2*trace(Q);
44     for i = 1:numel(hyp.mean)
45         dnlZ.mean(i) = -feval(mean{:}, hyp.mean, x, i)'*alpha;
46     end
47     end
48     end
```

3.2 Laplace's Approximation

For differentiable likelihoods, Laplace's approximation, approximates the posterior by a Gaussian centered at its mode and matching its curvature inf/infLaplace.m.

More concretely, the mean of the posterior $q(f|\mathcal{D}) = \mathcal{N}\big(f|\mu,V\big)$ is – defining $\ell_i(f_i) = \ln p(y_i|f_i)$ and $\ell(f) = \sum_{i=1}^n \ell_i(f_i)$ – given by

$$\mu = \arg\min_{\mathbf{f}} \phi(\mathbf{f}), \text{ where } \phi(\mathbf{f}) = \frac{1}{2} (\mathbf{f} - \mathbf{m})^{\top} \mathbf{K}^{-1} (\mathbf{f} - \mathbf{m}) - \ell(\mathbf{f}) \stackrel{c}{=} - \ln[p(\mathbf{f})p(\mathbf{y}|\mathbf{f})], \tag{2}$$

which we abbreviate by $\mu \leftarrow \mathcal{L}(\ell)$. The curvature $\frac{\partial^2 \varphi}{\partial f^{\top}} = K^{-1} + W$ with $W_{ii} = -\frac{\partial^2}{\partial f_i^2} \ln p(y_i|f_i)$ serves as precision for the Gaussian posterior approximation $V = (K^{-1} + W)^{-1}$ and the marginal likelihood $Z = \int p(f)p(y|f)df$ is approximated by $Z \approx Z_{LA} = \int \tilde{\varphi}(f)df$ where we use the 2nd order Taylor expansion at the mode μ given by $\tilde{\varphi}(f) = \varphi(\mu) + \frac{1}{2}(f - \mu)^{\top}V^{-1}(f - \mu) \approx \varphi(f)$.

Laplace's approximation needs derivatives up to third order for the mode fitting procedure (Newton method)

$$d_k = \frac{\partial^k}{\partial f^k} \log p(y|f), \quad k = 0, 1, 2, 3$$

and

$$d_k = \frac{\partial}{\partial \rho_i} \frac{\partial^k}{\partial f^k} \log p(y|f), \quad k = 0, 1, 2$$

evaluated at the latent location f and observed value y. The likelihood calls (see section 4)

• [d0, d1, d2, d3] = lik(hyp, y, f, [], 'infLaplace')

and

• [d0, d1, d2] = lik(hyp, y, f, [], 'infLaplace', i)

return exactly these values.

3.3 Expectation Propagation

The basic idea of Expectation Propagation (EP) as implemented in inf/infEP.m. is to replace the non-Gaussian likelihood terms $p(y_i|f_i)$ by Gaussian functions $t(f_i; \nu_i, \tau_i) = \exp(\nu_i f_i - \frac{1}{2}\tau_i f_i^2)$ and to adjust the natural parameters ν_i, τ_i such that the following identity holds:

$$\frac{1}{Z_{t,i}}\int f^kq_{-i}(f)\cdot t(f;\nu_i,\tau_i)df = \frac{1}{Z_{p,i}}\int f^kq_{-i}(f)\cdot p(y_i|f)df,\quad k=1,2$$

with the so-called cavity distributions $q_{-i}(f) = \mathcal{N}(f|m,K) \prod_{j \neq i} t(f_j; \nu_j, \tau_j) \propto \mathcal{N}(f|\mu,V)/t(f_i; \nu_i, \tau_i)$ equal to the posterior divided by the ith Gaussian approximation function and the two normalisers $Z_{t,i} = \int q_{-i}(f) \cdot t(f_i; \nu_i, \tau_i) df$ and $Z_{p,i} = \int q_{-i}(f) \cdot p(y_i|f_i) df$. The moment matching corresponds to minimising the following local KL-divergence

$$\nu_i, \tau_i = \arg\min_{\nu,\tau} KL[q_{-i}(f)p(y_i|f_i)/Z_{p,i}\|q_{-i}(f)t(f_i;\nu,\tau)/Z_{t,i}].$$

In order to apply the moment matching steps in a numerically safe way, EP requires the deriviatives of the expectations w.r.t. the Gaussian mean parameter μ

$$d_k = \frac{\partial^k}{\partial \mu^k} log \int p(y|f) \mathcal{N}(f|\mu, \sigma^2) df, \quad k = 0, 1, 2$$

and the ith likelihood hyperparameter ρ_i

$$d = \frac{\partial}{\partial \rho_i} \log \int p(y|f) \mathcal{N}(f|\mu, \sigma^2) df$$

which can be obtained by the likelihood calls (see section 4)

and

3.4 Kullback Leibler Divergence Minimisation

Another well known approach to approximate inference implemented inf/infKL.m in attempts to directly find the closest Gaussian $q(f|\mathcal{D}) = \mathcal{N}(f|\mu, V)$ to the exact posterior $p(f|\mathcal{D})$ w.r.t. to some proximity measure or equivalently to maximise a lower bound $Z(\mu, V)$ to the marginal likelihood Z as described in Nickisch & Rasmussen *Approximations for Binary Gaussian Process Classification*, JMLR, 2008. In particular, one minimises $KL(\mathcal{N}(f|\mu, V)||p(f|\mathcal{D}))$ which amounts to minimising $-\ln Z(\mu, V)$ as defined by:

$$\begin{split} -\ln \mathsf{Z} &= -\ln \int p(f)p(y|f)df = -\ln \int q(f|\mathcal{D})\frac{p(f)}{q(f|\mathcal{D})}p(y|f)df \\ &\overset{Jensen}{\leqslant} \int q(f|\mathcal{D})\ln \frac{q(f|\mathcal{D})}{p(f)}df - \int q(f|\mathcal{D})\ln p(y|f)df =: -\ln \mathsf{Z}(\mu, V) \\ &= \mathsf{KL}\left(\mathcal{N}(f|\mu, V)||\mathcal{N}(f|m, K)\right) - \sum_{i=1}^n \int \mathcal{N}(f_i|\mu_i, \nu_{i\,i})\ln p(y_i|f_i)df_i, \ \nu_{i\,i} = [V]_{i\,i} \\ &= \frac{1}{2}\left(\operatorname{tr}(VK^{-1} - I) - \ln |VK^{-1}|\right) + \frac{1}{2}(\mu - m)^\top K^{-1}(\mu - m) - \sum_{i=1}^n \ell^{KL}(\mu_i, \nu_{i\,i}) \right) \end{split}$$

where $\ell_v^{KL}(\mu_i) = \int \mathcal{N}(f_i|\mu_i,\nu_{i\,i})\ell_i(f_i)df_i$ is the convolution of the log likelihood ℓ_i with the Gaussian \mathcal{N} and $\mathbf{v} = dg(\mathbf{V})$. Equivalently, one can view ℓ^{KL} as a smoothed version of ℓ with univariate smoothing kernel \mathcal{N} .

From Challis & Barber Concave Gaussian Variational Approximations for Inference in Large Scale Bayesian Linear Models, AISTATS, 2011 we know that the mapping $(\mu, L) \mapsto -\ln Z(\mu, L^{\top}L)$ is

jointly convex whenever the likelihoods $f_i \mapsto \mathbb{P}(y_i|f_i)$ are log concave. In particular, this implies that every $(\mu_i, s_i) \mapsto -\ell^{KL}(\mu_i, s_i^2)$ is jointly convex.

We use an optimisation algorithm similar to EP (section 3.3) where we minimise the local KL-divergence the other way round μ_i , $s_i = \arg\min_{\mu,s} \text{KL}[\mathcal{N}(f|\mu,s^2)||q_{-i}(f)p(y_i|f_i)/Z_{p,i}]$. This view was brought forward by Tom Minka Convex Divergence measures and message passing, MSR-TR, 2005. The KL minimisation constitutes a jointly convex 2d optimisation problem solved by klmin using a scaled Newton approach which is included as a sub function in inf/infKL.m. The smoothed likelihood $\ell^{\text{KL}}(\mu_i,\nu_{ii})$ is implemented as a meta likelihood in likKL; it uses Gaussian-Hermite quadrature to compute the required integrals. Note that – as opposed to EP – Gaussian-Hermite quadrature is appropriate since we integrate against the $\ln \mathbb{P}(y_i|f_i)$ (which can be well approximated by a polynomial) instead of $\mathbb{P}(y_i|f_i)$ itself. The algorithm is – again unlike EP – provably convergent for log-concave likelihoods (e.g. likGauss, likLaplace, likSech2, likLogistic, likPoisson) since it can be regarded as coordinate descent with guaranteed decrease in the objective in every step. Due to the complex update computations, infKL can be quite slow although it has the same $\mathbb{O}(\mathfrak{n}^3)$ asymptotic complexity as EP and Laplace.

3.5 Variational Bayes

One can drive the bounding even further by means of local quadratic lower bounds to the log likelihood $\ell(f) = \ln p(y|f)$. Suppose that we use a super-Gaussian likelihood p(y|f) i.e. likelihoods that can be lower bounded by Gaussians of any width w (e.g. likLaplace, likT, likLogistic, likSech2). Formally, that means that there are $b, z \in \mathbb{R}$ such that

$$\rho(f) = \ln p(y|f-z) - bf$$

is symmetric and $\sqrt{f} \mapsto \rho(f)$ is a convex function for all $f \geqslant 0$. As a result, we obtain the following exact representation of the likelihood

$$\ell(f) = \ln p(y|f) = \max_{w>0} \left((b+wz)f - \frac{wf^2}{2} - \frac{1}{2}h(\gamma) \right),$$

which can be derived by convex duality and assuming the likelihoods to be super-Gaussian. Details can be found in papers by Palmer et al. *Variational EM Algorithms for Non-Gaussian Latent Variable Models*, NIPS, 2006 and Nickisch & Seeger Convex Variational Bayesian Inference for Large Scale Generalized Linear Models, ICML, 2009.

The bottom line is that we can treat the variational bounding as a sequence of Laplace approximations with the "variational Bayes" log likelihood

$$\ell^{VB}(f_{\mathfrak{i}}) = \ell(g_{\mathfrak{i}}) + b_{\mathfrak{i}}(f_{\mathfrak{i}} - g_{\mathfrak{i}}), \ \mathbf{g} = \text{sgn}(\mathbf{f} - \mathbf{z}) \odot \sqrt{(\mathbf{f} - \mathbf{z})^2 + \mathbf{v}} + \mathbf{z}$$

instead of the usual likelihood $\ell(f_i) = \ln p(y_i|f_i)$ i.e. we solve $\mu \leftarrow \mathcal{L}(\ell_v^{VB})$ instead of $\mu \leftarrow \mathcal{L}(\ell)$. See section 3.2. In the code of inf/infVB.m, the likelihood is implemented in the function likVB.

At the end, the optimal value of **W** can be obtained analytically via $w_i = |b_i - \ell'(g_i)|/|g_i - z_i|$.

For the minimisation in inf/infVB.m, we use a provably convergent double loop algorithm, where in the inner loop a nonlinear least squares problem (convex for log-concave likelihoods) is solved using inf/infLaplace.m such that $\mu \leftarrow \mathcal{L}(\ell_v^{VB})$ and in the outer loop, we compute $\mathbf{v} \leftarrow \text{dg}((\mathbf{K}^{-1} + \mathbf{W})^{-1})$. The only requirement to the likelihood function is that it returns the values z and b required by the bound which are delivered by the call (see section 4)

The negative marginal likelihood upper bound $-\ln Z_{VB}$ is obtained by integrating the prior times the exact representation of the likelihood

$$p(y|f) = \max_{\gamma > 0} q(y|f,\gamma), \ q(y|f,\gamma) = \mathcal{N}(f|\nu,\gamma) \exp\left(-\frac{h(\gamma)}{2} - \frac{\nu^2}{2\gamma}\right) \sqrt{2\pi\gamma}, \ \gamma = \frac{1}{w}, \ \nu = b\gamma + z$$

w.r.t. the latent variables f yielding

$$\begin{split} -\ln Z_{\mathrm{VB}} &= -\ln \int \mathcal{N}(\mathbf{f}|\mathbf{m},\mathbf{K}) \prod_{i=1}^{n} q_{i}(y_{i}|f_{i},\gamma_{i}) d\mathbf{f} \\ &= -\ln \mathcal{N}(\mathbf{m}|\mathbf{v},\mathbf{K}+\boldsymbol{\Gamma}) + \frac{1}{2} \left(h(\boldsymbol{\gamma}) - \mathbf{w}^{\top} \mathbf{v}^{2} - \mathbf{1}^{\top} \ln 2\pi \boldsymbol{\gamma}\right). \end{split}$$

3.6 FITC Approximations

One of the main problems with GP models is the high computational load for inference computations. In a setting with n training points x, exact inference with Gaussian likelihood requires $O(n^3)$ effort; approximations like Laplace of EP consist of a sequence of $O(n^3)$ operations.

There is a line of research with the goal to alleviate this burden by using approximate covariance functions \tilde{k} instead of k. A review is given by Candela and Rasmussen A Unifying View of Sparse Approximate Gaussian Process Regression, JMLR, 2005. One basic idea in those approximations is to work with a set of m inducing inputs u with a reduced computational load of $O(nm^2)$. In the following, we will provide a rough idea of the FITC approximation used in the toolbox. Let K denote the $n \times n$ covariance matrix between the training points x, K_u the $m \times n$ covariance matrix between the n training points and the m inducing points, and K_{uu} the $m \times m$ covariance matrix between the m inducing points. The FITC approximation to the covariance is given by

$$K \approx \tilde{K} = Q + G, \ G = \text{diag}(g), \ g = \text{diag}(K - Q), \ Q = K_u^\top Q_{uu}^{-1} K_u, \ Q_{uu} = K_{uu} + \sigma_{n_u}^2 I,$$

where σ_{n_u} is the noise from the inducing inputs. Note that \tilde{K} and K have the same diagonal elements diag(\tilde{K}) = diag(K); all off-diagonal elements are the same as for Q. Internally, the necessary covariance evaluations are performed by a meta covariance function cov/covFITC.m. The toolbox offers FITC versions for regression with Gaussian likelihood inf/infFITC.m, as well as for Laplace's approximation inf/infFITC_Laplace.m and expectation propagation inf/infFITC_EP.m.

The user can decide whether to treat the inducing inputs **u** as fixed or as hyperparameters. The latter is possible for a number of common stationary covariance functions (and also all three FITC inference methods inf/infFITC*.m) and allows to adjust the inducing inputs **u** w.r.t. the marginal likelihood. As detailed in the documentation of inf/infFITC*.m, **u** is treated as fixed if it is passed as the 2nd parameter of covFITC(cov,xu,...). If the hyperparameter structure hyp contains a field hyp.xu in inference method calls like infFITC*(hyp,...) or inference/prediction calls like gp(hyp,@infFITC*,...) the inducing inputs **u** are treated as hyperparameters and can be optimised.

3.7 Grid Approximations

Another way to bring down computational costs is to take advantage of (partial) grid structure present in the training points \mathbf{x} . For example, in geostatistics or image processing, the training data $\mathbf{x} \in \mathbb{R}^{n \times D}$ could be a complete 2d lattice of size $n_1 \times n_2$ as given by the axes $\mathbf{x}_1 \in \mathbb{R}^{n_1}$, $\mathbf{x}_2 \in \mathbb{R}^{n_2}$ so that $\mathbf{n} = \mathbf{n}_1 \cdot \mathbf{n}_2$, $\mathbf{D} = 2$ and $\mathbf{x} = [\text{vec}(\mathbf{x}_1 \mathbf{1}^\top), \text{vec}(\mathbf{1}\mathbf{x}_2^\top)]$. In general, a p-dimensional grid $\mathbf{x}_g \in \mathbb{R}^{N \times D}$ is specified by a set of axis matrices $\{\mathbf{x}_i \in \mathbb{R}^{n_i \times D_i}\}_{i=1..p}$ so that $\mathbf{N} = \prod_{i=1}^p n_i$ and $\mathbf{D} = \sum_{i=1}^p D_i$ where the axes do not need to be 1d nor do their components need to be sorted. As a consequence,

 \mathbf{x}_g represents a Cartesian product of its axes $\mathbf{x}_g = \mathbf{x}_1 \times \mathbf{x}_2 \times ... \times \mathbf{x}_p$. The cov/covGrid.m covariance function represents a Kronecker product covariance matrix

$$K=K_p\otimes ..\otimes K_2\otimes K_1$$

whose factorisation structure is given by the grid \mathbf{x}_g . The gain in computationial efficiency is due to the fact that matrix-vector product, determinant, inverse and eigenvalue computations decompose so that the overall cost of $\mathcal{O}(N^3)$ turns into $\mathcal{O}(\sum_{i=1}^p n_1^3)$. Internally, we use a meta covariance function cov/covGrid.m to represent the Kronecker covariance matrix and a Gaussian regression inference method inf/infGrid.m. We also support incomplete grids where n < N. A good starting point is Yunus Saatçi's PhD thesis Scalable Inference for Structured Gaussian Process Models, University of Cambridge, 2011. For incomplete grids, we use the extensions by Wilson et. al, Fast Kernel Learning for Multidimensional Pattern Extrapolation, NIPS, 2014 where conjugate gradients and a determinant approximations are used. See doc/demoGrid.m for an illustration. There is also an extension to non-Gaussian likelihoods by Seth Flaxman Fast Kronecker inference in Gaussian processes with non-Gaussian likelihoods.

4 Likelihood Functions

A likelihood function $p_{\rho}(y|f)$ (with hyperparameters ρ) is a conditional density $\int p_{\rho}(y|f)dy = 1$ defined for scalar latent function values f and outputs y. In the GPML toolbox, we use iid. likelihoods $p_{\rho}(y|f) = \prod_{i=1}^{n} p_{\rho}(y_i|f_i)$. The approximate inference engine does not explicitly distinguish between classification and regression likelihoods: it is fully generic in the likelihood allowing to use a single code in the inference step.

Likelihood functionality is needed both during inference and while predicting.

4.1 Prediction

A prediction at \mathbf{x}_* conditioned on the data $\mathcal{D}=(X,y)$ (as implemented in gp.m) consists of the predictive mean μ_{y_*} and variance $\sigma_{y_*}^2$ which are computed from the the latent marginal moments μ_{f_*} , $\sigma_{f_*}^2$ i.e. the Gaussian marginal approximation $\mathcal{N}(f_*|\mu_{f_*},\sigma_{f_*}^2)$ via

$$p(y_*|\mathcal{D}, \mathbf{x}_*) = \int p(y_*|f_*)p(f_*|\mathcal{D}, \mathbf{x}_*)df_* \approx \int p(y_*|f_*)\mathcal{N}(f_*|\mu_{f_*}, \sigma_{f_*}^2)df_*.$$
(3)

The moments are given by $\mu_{y_*} = \int y_* p(y_*|\mathcal{D}, \mathbf{x}_*) dy_*$ and $\sigma^2_{y_*} = \int (y_* - \mu_{y_*})^2 p(y_*|\mathcal{D}, \mathbf{x}_*) dy_*$. The likelihood call

• [lp,ymu,ys2] = lik(hyp, [], fmu, fs2)

does exactly this. Evaluation of the logarithm of $p_{y_*} = p(y_*|\mathcal{D}, \mathbf{x}_*)$ for values y_* can be done via

• [lp,ymu,ys2] = lik(hyp, y, fmu, fs2)

where 1p contains the number $\ln p_{u_*}$.

Using the moments of the likelihood $\mu(f_*) = \int y_* p(y_*|f_*) dy_*$ and $\sigma^2(f_*) = \int (y_* - \mu(f_*))^2 p(y_*|f_*) dy_*$ we obtain for the predictive moments the following (exact) expressions

$$\begin{array}{lcl} \mu_{y_*} & = & \int \mu(f_*) p(f_* | \mathcal{D}, \mathbf{x}_*) df_*, \text{ and} \\ \\ \sigma_{y_*}^2 & = & \int \left[\sigma^2(f_*) + (\mu(f_*) - \mu_{y_*})^2 \right] p(f_* | \mathcal{D}, \mathbf{x}_*) df_*. \end{array}$$

1. The binary case is simple since $y_* \in \{-1, +1\}$ and $1 = p_{y_*} + p_{-y_*}$. Using $\pi_* = p_{+1}$, we find

$$\begin{array}{lll} p_{y_*} & = & \begin{cases} \pi_* & y_* = +1 \\ 1 - \pi_* & y_* = -1 \end{cases} \\ \mu_{y_*} & = & \sum_{y_* = \pm 1} y_* p(y_* | \mathcal{D}, \mathbf{x}_*) = 2 \cdot \pi_* - 1 \in [-1, 1], \quad \text{and} \\ \sigma_{y_*}^2 & = & \sum_{y_* = \pm 1} (y_* - \mu_{y_*})^2 p(y_* | \mathcal{D}, \mathbf{x}_*) = 4 \cdot \pi_* (1 - \pi_*) \in [0, 1]. \end{array}$$

2. The continuous case for homoscedastic likelihoods depending on $r_* = y_* - f_*$ only and having noise variance $\sigma^2(f_*) = \sigma_n^2$ is also simple since the identity $p(y_*|f_*) = p(y_* - f_*|0)$ allows to substitute $y_* \leftarrow y_* + f_*$ yielding $\mu(f_*) = f_* + \int y_* p(y_*|0) dy_*$ and assuming $\int y_* p(y_*|0) dy_* = 0$ we arrive at

$$\begin{array}{rcl} \mu_{y_*} &=& \mu_{f_*}, \text{ and} \\ \sigma_{y_*}^2 &=& \sigma_{f_*}^2 + \sigma_n^2. \end{array}$$

3. The generalised linear model (GLM) case is also feasible. Evaluation of the predictive distribution is done by quadrature

$$p_{y_*} \ = \ \int p(y_*|f_*)p(f_*|\mathcal{D}, \mathbf{x}_*)df_* \approx \int p(y_*|f_*)\mathcal{N}(f_*|\mu_{f_*}, \sigma_{f_*}^2)df_*.$$

For GLMs the mean is given by $\mu(f_*) = g(f_*)$ and the variance is usually given by a simple function of the mean $\sigma^2(f_*) = \nu(g(f_*))$, hence we use Gaussian-Hermite quadrature with $\mathcal{N}(f_*|\mu_{f_*},\sigma_{f_*}^2) \approx p(f_*|\mathcal{D},\mathbf{x}_*)$ to compute

$$\begin{array}{lcl} \mu_{y_*} & = & \int g(f_*) p(f_* | \mathbb{D}, \mathbf{x}_*) df_*, \text{ and} \\ \\ \sigma_{y_*}^2 & = & \int \left[\nu(g(f_*)) + (g(f_*) - \mu_{y_*})^2 \right] p(f_* | \mathbb{D}, \mathbf{x}_*) df_* \neq \nu(\mu_{y_*}). \end{array}$$

4. Finally the warped Gaussian likelihood predictive distribution with strictly monotonically increasing warping function q is given by the expression

$$p(y_*|\mathcal{D}, \mathbf{x}_*) = g'(y_*)\mathcal{N}\left(g(y_*)|\mu_{f_*}, \sigma_n^2 + \sigma_{f_*}^2\right)$$

so that the predictive moments can be computed by Gaussian-Hermite quadrature.

In the following, we will detail how and which likelihood functions are implemented in the GPML toolbox. Further, we will mention dependencies between likelihoods and inference methods and provide some analytical expressions in addition to some likelihood implementations.

4.2 Interface

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The likelihoods are in fact the most challenging object in our implementation. Different inference algorithms require different aspects of the likelihood to be computed, therefore the interface is rather involved as detailed below.

```
\langle likFunctions.m \ 18 \rangle \equiv
 1 % likelihood functions are provided to be used by the gp.m function:
2 %
3 %
       likErf
                      (Error function, classification, probit regression)
4 %
      likLogistic
                                    classification, logit regression)
 5 %
                      (Uniform likelihood, classification)
      likUni
 6 %
 7 %
      likGauss
                      (Gaussian, regression)
8 %
      likGaussWarp
                       (Warped Gaussian, regression)
9 %
                       (Gumbel likelihood for extremal values)
      likGumbel
10 %
      likLaplace
                       (Laplacian or double exponential, regression)
11 %
      likSech2
                       (Sech-square, regression)
12 %
       likT
                       (Student's t, regression)
13 %
                       (Poisson regression, count data)
14 %
      likPoisson
                       (Nonnegative regression, positive data)
15 %
      likGamma
16 %
                       (Nonnegative regression, positive data)
      likExp
17 %
      likInvGauss
                       (Nonnegative regression, positive data)
                       (Beta regression, interval data)
18 %
       likBeta
19 %
20 %
                       (Mixture of individual covariance functions)
       likMix
21 %
22 % The likelihood functions have three possible modes, the mode being selected
```

```
23 % as follows (where "lik" stands for any likelihood function in "lik/lik*.m".):
24 %
25 % 1) With one or no input arguments:
                                          [REPORT NUMBER OF HYPERPARAMETERS]
27 %
       s = lik OR s = lik(hyp)
28 %
29 % The likelihood function returns a string telling how many hyperparameters it
30 % expects, using the convention that "D" is the dimension of the input space.
31 % For example, calling "likLogistic" returns the string '0'.
32 %
33 %
34 % 2) With three or four input arguments:
                                                                  [PREDICTION MODE]
35 %
36 %
       lp = lik(hyp, y, mu) OR [lp, ymu, ys2] = lik(hyp, y, mu, s2)
37 %
38 % This allows to evaluate the predictive distribution. Let p(y_*|f_*) be the
39 % likelihood of a test point and N(f_*|mu,s2) an approximation to the posterior
40 % marginal p(f_*|x_*,x,y) as returned by an inference method. The predictive
41 % distribution p(y_*|x_*,x,y) is approximated by.
42 %
      q(y_*) = \inf N(f_*|mu,s2) p(y_*|f_*) df_*
43 %
44 %
     lp = log(q(y)) for a particular value of y, if s2 is [] or 0, this
45 %
                       corresponds to log(p(y|mu))
46 %
                       the mean and variance of the predictive marginal q(y)
     ymu and ys2
47 %
                       note that these two numbers do not depend on a particular
48 %
                       value of y
49 % All vectors have the same size.
50 %
51 %
52 % 3) With five or six input arguments, the fifth being a string [INFERENCE MODE]
53 %
54 % [varargout] = lik(hyp, y, mu, s2, inf) OR
55 % [varargout] = lik(hyp, y, mu, s2, inf, i)
57 % There are three cases for inf, namely a) infLaplace, b) infEP and c) infVB.
58 % The last input i, refers to derivatives w.r.t. the ith hyperparameter.
60 % a1) [lp,dlp,d2lp,d3lp] = lik(hyp, y, f, [], 'infLaplace')
61 % lp, dlp, d2lp and d3lp correspond to derivatives of the log likelihood
62 % log(p(y|f)) w.r.t. to the latent location f.
63 % lp = log(p(y|f))
64 \% dlp = d log(p(y|f)) / df
65 \% d21p = d^2 log(p(y|f)) / df^2
66 \% d3lp = d^3 log(p(y|f)) / df^3
67 %
68 % a2) [lp_dhyp,dlp_dhyp,d2lp_dhyp] = lik(hyp, y, f, [], 'infLaplace', i)
69 % returns derivatives w.r.t. to the ith hyperparameter
     lp_dhyp = d log(p(y|f)) / (
                                          dhyp_i)
71 % dlp_dhyp = d^2 log(p(y|f)) / (df dhyp_i)
72 % d2lp_dhyp = d^3 log(p(y|f)) / (df^2 dhyp_i)
73 %
74 %
75 \% b1) [1Z,d1Z,d21Z] = lik(hyp, y, mu, s2, 'infEP')
76 % let Z = \inf p(y|f) N(f|mu,s2) df then
77 \% 1Z = log(Z)
78 \% dlZ = d log(Z) / dmu
79 \% d21Z = d^2 \log(Z) / dmu^2
80 %
```

```
81 % b2) [dlZhyp] = lik(hyp, y, mu, s2, 'infEP', i)
 82 % returns derivatives w.r.t. to the ith hyperparameter
 83 \% dlZhyp = d log(Z) / dhyp_i
 84 %
 85 %
 86 \% c1) [b,z] = lik(hyp, y, [], ga, 'infVB')
 87 % ga is the variance of a Gaussian lower bound to the likelihood p(y|f).
 88 % p(y|f) \neq exp(b*(f+z) - (f+z).^2/(2*ga) - h(ga)/2) \Rightarrow N(f|b*ga-z,ga)
 89 % The function returns the linear part b and z.
 90 %
 91 % Cumulative likelihoods are designed for binary classification. Therefore, they
 92 % only look at the sign of the targets y; zero values are treated as +1.
 93 %
 94 % Some examples for valid likelihood functions:
 95 % lik = @likLogistic;
 96 %
         lik = {'likMix',{'likUni',@likErf}}
 97 %
          lik = {@likPoisson, 'logistic'};
99 % See the help for the individual likelihood for the computations specific to
100 % each likelihood function.
101 %
102 (gpml copyright 6a)
```

4.3 Implemented Likelihood Functions

The following table enumerates all (currently) implemented likelihood functions that can be found at lik/lik<NAME>.m and their respective set of hyperparameters ρ .

lik <name></name>	regression $y_i \in \mathbb{R}$	$p_{\rho}(y_i f_i) =$	ρ =			
Gauss	Gaussian	$\mathcal{N}(y_i f_i,\sigma^2) = \frac{1}{\sqrt{2\pi}\sigma} \exp\left(-\frac{(y_i-f_i)^2}{2\sigma^2}\right)$	{ln σ}			
GaussWarp	Warped Gaussian	$\mathcal{N}(g_{\theta},(y_{i}) f_{i},\sigma^{2})g_{\theta}'(y_{i})$	$\{\theta_1,,\theta_{n_q},\ln\sigma\}$			
Gumbel	Gumbel	$\frac{\pi}{\sigma\sqrt{6}}\exp\left(-z_{i}-e^{-z_{i}}\right), z_{i}=\gamma+\frac{s\cdot\pi(y_{i}-f_{i})}{\sigma\sqrt{6}}, s =1$	{ln σ}			
Sech2	Sech-squared	$\left(\frac{\tau}{2\cosh^2(\tau(y_i-f_i))}, \tau = \frac{\pi}{2\sigma\sqrt{3}}\right)$	{ln σ}			
Laplace	Laplacian	$\frac{1}{2b} \exp\left(-\frac{ y_i-f_i }{b}\right), \ b=\frac{\sigma}{\sqrt{2}}$	{ln σ}			
Т	Student's t	$\frac{\Gamma(\frac{\nu+1}{2})}{\Gamma(\frac{\nu}{2})} \frac{1}{\sqrt{\nu\pi}\sigma} \left(1 + \frac{(y_i - f_i)^2}{\nu\sigma^2}\right)^{-\frac{\nu+1}{2}}$	$\{\ln(\nu-1), \ln\sigma\}$			
lik <name></name>	classification $y_i \in \{\pm 1\}$	$p_{\rho}(y_i f_i) =$	$\rho =$			
Erf	Error function	$\int_{-\infty}^{y_i f_i} \mathcal{N}(t) dt$	Ø			
Logistic	Logistic function	$\frac{1}{1+\exp(-y_i f_i)}$	Ø			
Uni	Label noise	$\frac{1}{2}$	Ø			
lik <name></name>	count data $y_i \in \mathbb{N}$	$p_{\rho}(y_i f_i) =$	ρ =			
Poisson	Poisson	$\mu^{y} \cdot \frac{e^{-\mu}}{y!}, \ \mu = e^{f} \text{ or } \mu = \log(1 + e^{f})$	Ø			
lik <name></name>	nonnegative data $y_i \in \mathbb{R}_+ \setminus \{0\}$	$p_{\rho}(y_i f_i) =$	ρ =			
Weibull	Weibull, $\gamma_1 = \Gamma(1 + 1/\kappa)$	$\kappa \gamma_1 / \mu (y \gamma_1 / \mu)^{\kappa - 1} \exp(-(y \gamma_1 / \mu)^{\kappa})$	{ln κ}			
Gamma	Gamma	$\frac{\kappa \gamma_1 / \mu (y \gamma_1 / \mu)^{\kappa - 1} \exp(-(y \gamma_1 / \mu)^{\kappa})}{\kappa \gamma_1 / \mu (y \gamma_1 / \mu)^{\kappa - 1} \exp(-(y \gamma_1 / \mu)^{\kappa})}$ $\frac{\alpha^{\alpha} y^{\alpha - 1}}{\Gamma(\alpha)} \mu^{-\alpha} \exp\left(-\frac{y \alpha}{\mu}\right)$	{ln α}			
Exp	Exponential	$\mu^{-1} \exp\left(-\frac{y}{\mu}\right)$	Ø			
InvGauss	Inverse Gaussian	$\sqrt{\frac{\lambda}{2\pi y^3}} \exp\left(-\frac{\lambda(y-\mu)^2}{2\mu^2 y}\right)$	$\{\ln\lambda\}$			
lik <name></name>	interval data $y_i \in [0, 1]$	$p_{\rho}(y_i f_i) =$	ρ =			
Beta	Beta	$\frac{\Gamma(\phi)}{\Gamma(\mu\phi)\Gamma((1-\mu)\phi)}y^{\mu\phi-1}(1-y)^{(1-\mu)\phi-1}$	{ln φ}			
Composite likelihood functions $[p_1(y_i f_i), p_1(y_i f_i),] \mapsto p_{\rho}(y_i f_i)$						
Mix	Mixture	$\sum_{j} \alpha_{j} p_{j}(y_{i} f_{i})$	$\{\ln \alpha_1, \ln \alpha_2,\}$			

4.4 Usage of Implemented Likelihood Functions

Some code examples taken from doc/usageLik.m illustrate how to use simple and composite likelihood functions to specify a GP model.

Syntactically, a likelihood function 1f is defined by

```
lk := 'func' | @func // simple
lf := {lk} | {param, lk} | {lk, {lk, .., lk}} // composite
```

i.e., it is either a string containing the name of a likelihood function, a pointer to a likelihood function or one of the former in combination with a cell array of likelihood functions and an additional list of parameters.

```
\langle doc/usageLik.m 21 \rangle \equiv
21
     1 % demonstrate usage of likelihood functions
     3 % See also likFunctions.m.
     4 %
     5 (gpml copyright 6a)
     6 clear all, close all
     7 n = 5; f = randn(n,1);
                                % create random latent function values
     9 % set up simple classification likelihood functions
     10 \text{ yc} = \text{sign}(f);
     11 1c0 = {'likErf'};
                             hypc0 = [];  % no hyperparameters are needed
     12 lc1 = {@likLogistic}; hypc1 = []; % also function handles are OK
                             hvpc2 = [];
     13 lc2 = {'likUni'};
     14 lc3 = {'likMix', {'likUni', @likErf}}; hypc3 = log([1;2]); %mixture
    1.5
    16\ \% set up simple regression likelihood functions
     17 \text{ yr} = f + randn(n,1)/20;
    18 \text{ sn} = 0.1;
                                                 % noise standard deviation
    19 lr0 = {'likGauss'}; hypr0 = log(sn);
    20 lr1 = {'likLaplace'}; hypr1 = log(sn);
    21 lr2 = {'likSech2'}; hypr2 = log(sn);
    22 \text{ nu} = 4;
                                             % number of degrees of freedom
    23 lr3 = {'likT'}; hypr3 = [log(nu-1); log(sn)];
    24 lr4 = {'likMix', {lr0, lr1}}; hypr4 = [log([1,2]); hypr0; hypr1];
    26 a = 1; % set up warped Gaussian with g(y) = y + a*sign(y).*y.^2
    27 1r5 = {'likGaussWarp',['poly2']}; hypr5 = log([a;sn]);
    28 lr6 = {'likGumbel', '+'}; hypr6 = log(sn);
     29
    30 % set up Poisson regression
    31 \text{ yp} = \text{fix}(abs(f)) + 1;
     32 lp0 = {@likPoisson, 'logistic'}; hypp0 = [];
     33 lp1 = {@likPoisson,'exp'};
                                    hypp1 = [];
    35 % set up other GLM likelihoods for positive or interval regression
     36 lg1 = {@likGamma, 'logistic'}; al = 2; hyp.lik = log(al);
    37 lg2 = {@likInvGauss,'exp'}; lam = 1.1; hyp.lik = log(lam);
    40
    41 % 0) specify the likelihood function
    42 lik = lc0; hyp = hypc0; y = yc;
    43 \% lik = lr4; hyp = hypr4; y = yr;
```

44 % lik = lp1; hyp = hypp1; y = yp;

```
45
46 % 1) query the number of parameters
47 feval(lik{:})
48
49 % 2) evaluate the likelihood function on f
50 exp(feval(lik{:},hyp,y,f))
51
52 % 3a) evaluate derivatives of the likelihood
53 [lp,dlp,d2lp,d3lp] = feval(lik{:}, hyp, y, f, [], 'infLaplace');
54
55 % 3b) compute Gaussian integrals w.r.t. likelihood
56 mu = f; s2 = rand(n,1);
57 [lZ,dlZ,d2lZ] = feval(lik{:}, hyp, y, mu, s2, 'infEP');
58
59 % 3c) obtain lower bound on likelihood
60 ga = rand(n,1);
61 [b,z] = feval(lik{:}, hyp, y, [], ga, 'infVB');
```

4.5 Compatibility Between Likelihoods and Inference Methods

The following table lists all possible combinations of likelihood function and inference methods.

Likelihood \ Inference	Exact FITC	EP FITC-EP	Laplace FITC-Laplace	VB	KL	MCMC	LOO	Type, Output Domain	Alternative Names
Entermood (Interence	Grid	1110 L1	Grid-Laplace	, ve late meme 100 Type, out		Type, Gutput Bomain	Thermative rannes		
Gaussian	√	✓	✓	√	√	✓	✓	regression, R	
Warped Gaussian		✓	✓	√	√	✓	✓	regression, R	
Gumbel			✓		√	✓	✓	regression, R	
Sech-squared		✓	✓	√	√	✓	✓	regression, R	logistic distribution
Laplacian		✓	✓	√	√	✓	√	regression, R	double exponential
Student's t			✓	√	√	✓	✓	regression, R	
Mixture		√	✓		√	√	√		mixing meta likelihood
Error function		✓	✓		✓	√	√	classification, {±1}	probit regression
Logistic function		✓	✓	√	√	✓	✓	classification, {±1}	logit regression
Uniform		✓	✓	√	✓	✓	✓	classification, {±1}	label noise
Weibull			✓			√	√	positive data, $\mathbb{R}_+ \setminus \{0\}$	nonnegative regression
Gamma			✓			√	√	positive data, $\mathbb{R}_+ \setminus \{0\}$	nonnegative regression
Exp			✓			√	√	positive data, $\mathbb{R}_+ \setminus \{0\}$	nonnegative regression
Inverse Gaussian			✓			√	√	positive data, $\mathbb{R}_+ \setminus \{0\}$	nonnegative regression
Poisson		(√)*	✓		√	✓	✓	count data, N	Poisson regression
Beta			√			✓	√	interval data, [0, 1]	beta regression

$(\checkmark)^*$ EP might not converge in some cases since quadrature is used.

Exact inference is only tractable for Gaussian likelihoods. Expectation propagation together with Student's t likelihood is inherently unstable due to non-log-concavity. Laplace's approximation for Laplace likelihoods is not sensible because at the mode the curvature and the gradient is undefined due to the non-differentiable peak of the Laplace distribution. Special care has been taken for the non-convex optimisation problem imposed by the combination Student's t likelihood and Laplace's approximation.

4.6 Gaussian Likelihood

The Gaussian likelihood is the simplest likelihood because the posterior distribution is not only Gaussian but can be computed analytically. In principle, the Gaussian likelihood would only be operated in conjunction with the exact inference method but we chose to provide compatibility with all other inference algorithms as well because it enables code testing and allows to switch between

different regression likelihoods very easily.

```
\langle lik/likGauss.m 23a \rangle \equiv
23a
       1 function [varargout] = likGauss(hyp, y, mu, s2, inf, i)
       3 % likGauss - Gaussian likelihood function for regression. The expression for the
       4 % likelihood is
             likGauss(t) = exp(-(t-y)^2/2*sn^2) / sqrt(2*pi*sn^2),
       6 \% where y is the mean and sn is the standard deviation.
       7 %
       8 % The hyperparameters are:
      9 %
      10 % hyp = [\log(sn)]
      11 %
      12 % Several modes are provided, for computing likelihoods, derivatives and moments
      13 % respectively, see likFunctions.m for the details. In general, care is taken
      14 % to avoid numerical issues when the arguments are extreme.
      15 %
      16 \(\langle gpml \) copyright 6a\(\rangle \)
      17 %
      18 % See also LIKFUNCTIONS.M.
      20 if nargin<3, varargout = {'1'}; return; end % report number of hyperparameters
      22 sn2 = exp(2*hyp);
      23
      24 if nargin<5
                                                       % prediction mode if inf is not present
      25
           (Prediction with Gaussian likelihood 23b)
      26 else
      27
           switch inf
      28 case 'infLaplace'
      29
            (Laplace's method with Gaussian likelihood 24a)
      30
         case 'infEP'
      31
             (EP inference with Gaussian likelihood 24b)
      32
           case 'infVB'
      33
             (Variational Bayes inference with Gaussian likelihood 24c)
      34
           end
      35 end
23b
      \langle Prediction \ with \ Gaussian \ likelihood \ 23b \rangle \equiv
                                                                        (23a)
      1 if isempty(y), y = zeros(size(mu)); end
       2 s2zero = 1; if nargin>3&&numel(s2)>0&&norm(s2)>eps, s2zero = 0; end % s2==0 ?
       3 if s2zero
                                                                             % log probability
       4 lp = -(y-mu).^2./sn2/2-log(2*pi*sn2)/2; s2 = 0;
       5 else
           lp = likGauss(hyp, y, mu, s2, 'infEP');
                                                                                   % prediction
       7 end
       8 \text{ ymu} = \{\}; \text{ ys2} = \{\};
       9 if nargout>1
      10 ymu = mu;
                                                                              % first y moment
      11
         if nargout>2
      12
            ys2 = s2 + sn2;
                                                                             % second y moment
      13
           end
      14 end
      15 varargout = {lp,ymu,ys2};
```

The Gaussian likelihood function has a single hyperparameter ρ , the log of the noise standard deviation σ_n .

4.6.1 Exact Inference

Exact inference doesn't require any specific likelihood related code; all computations are done directly by the inference method, section 3.1.

4.6.2 Laplace's Approximation

```
24a
      \langle Laplace's method with Gaussian likelihood 24a \rangle \equiv
                                                                   (23a)
      1 if nargin<6
                                                                   % no derivative mode
          if isempty(y), y=0; end
          ymmu = y-mu; dlp = {}; d2lp = {}; d3lp = {};
          lp = -ymmu.^2/(2*sn2) - log(2*pi*sn2)/2;
      5
          if nargout>1
            dlp = ymmu/sn2;
                                                   % dlp, derivative of log likelihood
      7
            if nargout>2
                                              % d2lp, 2nd derivative of log likelihood
      8
              d2lp = -ones(size(ymmu))/sn2;
      9
              if nargout>3
                                              % d3lp, 3rd derivative of log likelihood
      10
                d3lp = zeros(size(ymmu));
      11
               end
      12
            end
     13
          end
          varargout = {lp,dlp,d2lp,d3lp};
     15 else
                                                                      % derivative mode
          lp_dhyp = (y-mu).^2/sn2 - 1; % derivative of log likelihood w.r.t. hypers
     16
     17
          dlp_dhyp = 2*(mu-y)/sn2;
                                                                   % first derivative,
     18 d2lp_dhyp = 2*ones(size(mu))/sn2; % and also of the second mu derivative
      varargout = {lp_dhyp,dlp_dhyp,d2lp_dhyp};
      20 end
```

4.6.3 Expectation Propagation

```
\langle EP \text{ inference with Gaussian likelihood 24b} \rangle \equiv
24b
                                                                      (23a)
       1 if nargin<6
                                                                     % no derivative mode
         1Z = -(y-mu).^2./(sn2+s2)/2 - log(2*pi*(sn2+s2))/2; % log part function
          d1Z = {}; d21Z = {};
       3
          if nargout>1
       5
           dlZ = (y-mu)./(sn2+s2);
                                                           % 1st derivative w.r.t. mean
       6
            if nargout>2
       7
                                                           % 2nd derivative w.r.t. mean
              d21Z = -1./(sn2+s2);
       8
             end
      9
           end
      10
          varargout = {1Z,d1Z,d21Z};
                                                                        % derivative mode
           dlZhyp = ((y-mu).^2./(sn2+s2)-1) ./ (1+s2./sn2); % deriv. w.r.t. hyp.lik
           varargout = {dlZhyp};
      14 end
```

4.6.4 Variational Bayes

```
24c  \langle Variational Bayes inference with Gaussian likelihood 24c\rangle = (23a)
1 % variational lower site bound
2 % t(s) = exp(-(y-s)^2/2sn2)/sqrt(2*pi*sn2)
3 % the bound has the form: (b+z/ga)*f - f.^2/(2*ga) - h(ga)/2
4 n = numel(s2); b = zeros(n,1); y = y.*ones(n,1); z = y;
5 varargout = {b,z};
```

4.7 Warped Gaussian Likelihood

Starting from the likelihood p(y|f) we are sometimes facing the situation where the data $y \in \mathcal{Y} \subseteq \mathbb{R}$ is not distributed according to p(y|f) but some nonlinear transformation of the data g(y) = z so that $z \sim p(z|f)$. Here, the warping function $g: \mathcal{Y} \to \mathbb{R}$ needs to be strictly monotonically increasing i.e. g'(y) > 0. Formally, we start from the fact that p(z|f) integrates to one and use the derivative dz = g'(y)dy to substitute

$$\int p(z|f)dz = 1 = \int p_g(y|f)dy, \ p_g(y|f) = p(g(y)|f)g'(y)$$

where we have defined the log warped likelihood $\ln p_g(y|f) = \ln p(g(y)|f) + \ln g'(y)$. The interesting bit is that approximate inference methods such as infExact, infLaplace, infEP, infVB, infKL remain fully feasible; only prediction and derivatives become more involved. The usual GP inference is recovered by using the identity warping function $g: y \mapsto y$. The construction works in princple for any likelihood but our implementation in likGaussWarp is limited to the Gaussian likelihood.

Hyperparameter derivatives

Hyperparameter derivatives for infLaplace are obtained as follows

$$\begin{split} \frac{\partial}{\partial \theta} \ln \frac{\partial^k}{\partial f^k} p_g(y|f) &= \frac{\partial}{\partial \theta} \ln \frac{\partial^k}{\partial f} p(g(y)|f) + \frac{\partial}{\partial \theta} \frac{\partial^k}{\partial f^k} \ln g'(y), \ k = 0, 1, 2 \\ &= -\frac{\partial^{k+1}}{\partial f^{k+1}} \ln p(g(y)|f) \frac{\partial}{\partial \theta} g(y) + \frac{\partial}{\partial \theta} \frac{\partial^k}{\partial f^k} \ln g'(y). \end{split}$$

Similarly for infEP the derivatives are given by

$$\begin{split} \frac{\partial}{\partial \theta} \ln \int p_g(y|f) \mathcal{N}(f|\mu,\sigma^2) df &= \frac{\partial}{\partial \theta} \ln \int p(g(y)|f) \mathcal{N}(f|\mu,\sigma^2) df + \frac{\partial}{\partial \theta} \ln g'(y) \\ &= -\frac{\partial}{\partial \mu} \ln \int p(g(y)|f) \mathcal{N}(f|\mu,\sigma^2) df \frac{\partial}{\partial \theta} g(y) + \frac{\partial}{\partial \theta} \ln g'(y). \end{split}$$

This trick above works for any homoscedastic likelihood where $p(y|f) = p(y + \beta|f + \beta)$ such as likGauss, likLaplace, likSech2 and likT.

Predictive moments

As detailed in 4, the predictive distribution is – for Gaussian likelihood – given by

$$\begin{split} p(z_*|\mathcal{D},\mathbf{x}_*) &= \int p(z_*|f_*)p(f_*|\mathcal{D},\mathbf{x}_*)df_* = \int \mathcal{N}(z_*|f_*,\sigma_n^2)\mathcal{N}(f_*|\mu_{f_*},\sigma_{f_*}^2)df_* \\ &= \mathcal{N}(z_*|\mu_{f_*},\sigma_n^2+\sigma_{f_*}^2), \text{ where } z_* = g(y_*) \\ p(y_*|\mathcal{D},\mathbf{x}_*) &= g'(y_*)\mathcal{N}(g(y_*)|\mu_{f_*},\sigma_n^2+\sigma_{f_*}^2). \end{split}$$

Hence, the predictive moments are obtained by the 1d integrals

$$\begin{split} \mu_{y_*} &= \int y_* g'(y_*) \mathcal{N}(g(y_*) | \mu_{f_*}, \sigma_n^2 + \sigma_{f_*}^2) dy_* \\ &= \int g^{-1}(z_*) \mathcal{N}(z_* | \mu_{f_*}, \sigma_n^2 + \sigma_{f_*}^2) dz_*, \text{ and} \\ \sigma_{y_*}^2 &= \int (y_* - \mu_{y_*})^2 g'(y_*) \mathcal{N}(g(y_*) | \mu_{f_*}, \sigma_n^2 + \sigma_{f_*}^2) dy_* \\ &= \int (g^{-1}(z_*) - \mu_{y_*})^2 \mathcal{N}(z_* | \mu_{f_*}, \sigma_n^2 + \sigma_{f_*}^2) dz_*. \end{split}$$

4.8 Gumbel Likelihood

Distributions of extrema are well captured by the Gumbel distribution

$$p(y) = \frac{1}{\beta} \exp(-z - e^{-z}), z = s \frac{y - \eta}{\beta}, s \in \{\pm 1\}$$

with mean $\mu=\eta+\beta\gamma$ and variance $\sigma^2=\pi^2\beta^2/6$ where $\gamma=0.57721566490153$ denotes Euler–Mascheroni's constant. Skewness is approximately given by 1.1395s where s is a sign switching between left and right skewness and kurtosis is 12/5. The final expression for the Gumbel likelihood is

 $p(y|f) = \frac{\pi}{\sigma\sqrt{6}} \exp\left(-z - e^{-z}\right), \ z = \gamma + s \frac{\pi}{\sigma\sqrt{6}} (y - f), \ s \in \{\pm 1\}.$

4.9 Laplace Likelihood

Laplace's Approximation

The following derivatives are needed:

$$\begin{aligned} & \ln p(y|f) &= -\ln(2b) - \frac{|f - y|}{b} \\ & \frac{\partial \ln p}{\partial f} &= \frac{\text{sign}(f - y)}{b} \\ & \frac{\partial^2 \ln p}{(\partial f)^2} &= \frac{\partial^3 \ln p}{(\partial f)^3} = \frac{\partial^3 \ln p}{(\partial \ln \sigma_n)(\partial f)^2} = 0 \\ & \frac{\partial \ln p}{\partial \ln \sigma_n} &= \frac{|f - y|}{b} - 1 \end{aligned}$$

Expectation Propagation

Expectation propagation requires integration against a Gaussian measure for moment matching.

We need to evaluate $\ln Z = \ln \int \mathcal{L}(y|f,\sigma_n^2) \mathcal{N}(f|\mu,\sigma^2) df$ as well as the derivatives $\frac{\partial \ln Z}{\partial \mu}$ and $\frac{\partial^2 \ln Z}{\partial \mu^2}$ where $\mathcal{N}(f|\mu,\sigma^2) = \frac{1}{\sqrt{2\pi\sigma^2}} \exp\left(-\frac{(f-\mu)^2}{2\sigma^2}\right)$, $\mathcal{L}(y|f,\sigma_n^2) = \frac{1}{2b} \exp\left(-\frac{|y-f|}{b}\right)$, and $b = \frac{\sigma_n}{\sqrt{2}}$. As a first step, we reduce the number of parameters by means of the substitution $\tilde{f} = \frac{f-y}{\sigma_n}$ yielding

$$\begin{split} Z &= \int \mathcal{L}(y|f,\sigma_n^2) \mathcal{N}(f|\mu,\sigma^2) df \\ &= \frac{1}{\sqrt{2\pi}\sigma} \frac{\sqrt{2}}{2\sigma_n} \int exp\left(-\frac{(f-\mu)^2}{2\sigma^2}\right) exp\left(-\sqrt{2}\frac{|f-y|}{\sigma_n}\right) df \\ &= \frac{\sqrt{2}}{2\sigma\sqrt{2\pi}} \int exp\left(-\frac{(\sigma_n\tilde{f}+y-\mu)^2}{2\sigma^2}\right) exp\left(-\sqrt{2}|\tilde{f}|\right) d\tilde{f} \\ &= \frac{\sigma_n}{\sigma\sigma_n\sqrt{2\pi}} \int exp\left(-\frac{\sigma_n^2\left(\tilde{f}-\frac{\mu-y}{\sigma_n}\right)^2}{2\sigma^2}\right) \mathcal{L}(\tilde{f}|0,1) d\tilde{f} \\ &= \frac{1}{\sigma_n} \int \mathcal{L}(f|0,1) \mathcal{N}(f|\tilde{\mu},\tilde{\sigma}^2) df \\ ln Z &= ln \tilde{Z} - ln \, \sigma_n = ln \int \mathcal{L}(f|0,1) \mathcal{N}(f|\tilde{\mu},\tilde{\sigma}^2) df - ln \, \sigma_n \end{split}$$

with $\tilde{\mu} = \frac{\mu - y}{\sigma_n}$ and $\tilde{\sigma} = \frac{\sigma}{\sigma_n}$. Thus, we concentrate on the simpler quantity $\ln \tilde{Z}$.

$$\begin{split} \ln Z &= \ln \int \exp \left(-\frac{(f-\tilde{\mu})^2}{2\tilde{\sigma}^2} - \sqrt{2} |f| \right) df - \ln \tilde{\sigma} \sqrt{2\pi} - \ln \sqrt{2} \sigma_n \\ &= \ln \left[\int_{-\infty}^0 \exp \left(-\frac{(f-\tilde{\mu})^2}{2\tilde{\sigma}^2} + \sqrt{2} f \right) df + \int_0^\infty \exp \left(-\frac{(f-\tilde{\mu})^2}{2\tilde{\sigma}^2} - \sqrt{2} f \right) df \right] + C \\ &= \ln \left[\int_{-\infty}^0 \exp \left(-\frac{f^2 - 2(\tilde{\mu} + \tilde{\sigma}^2 \sqrt{2})f + \tilde{\mu}^2}{2\tilde{\sigma}^2} \right) df + \int_0^\infty \exp \left(-\frac{f^2 - 2(\tilde{\mu} - \tilde{\sigma}^2 \sqrt{2})f + \tilde{\mu}^2}{2\tilde{\sigma}^2} \right) df \right] + C \\ &= \ln \left[\exp \left(\frac{m_-^2}{2\tilde{\sigma}^2} \right) \int_{-\infty}^0 \exp \left(-\frac{(f-m_-)^2}{2\tilde{\sigma}^2} \right) df + \exp \left(\frac{m_+^2}{2\tilde{\sigma}^2} \right) \int_0^\infty \exp \left(-\frac{(f-m_+)^2}{2\tilde{\sigma}^2} \right) df \right] - \frac{\tilde{\mu}^2}{2\tilde{\sigma}^2} + C \\ &= \ln \left[\exp \left(\frac{m_-^2}{2\tilde{\sigma}^2} \right) \int_{-\infty}^0 \mathcal{N}(f|m_-, \tilde{\sigma}^2) df + \exp \left(\frac{m_+^2}{2\tilde{\sigma}^2} \right) \left(1 - \int_{-\infty}^0 \mathcal{N}(f|m_+, \tilde{\sigma}^2) df \right) \right] - \frac{\tilde{\mu}^2}{2\tilde{\sigma}^2} - \ln \sqrt{2} \sigma_n \\ &= \ln \left[\exp \left(\frac{m_-^2}{2\tilde{\sigma}^2} \right) \Phi \left(\frac{m_-}{\tilde{\sigma}} \right) - \exp \left(\frac{m_+^2}{2\tilde{\sigma}^2} \right) \Phi \left(\frac{m_+}{\tilde{\sigma}} \right) + \exp \left(\frac{m_+^2}{2\tilde{\sigma}^2} \right) \right] - \frac{\tilde{\mu}^2}{2\tilde{\sigma}^2} - \ln \sqrt{2} \sigma_n \end{split}$$

Here, $\Phi(z) = \int_{-\infty}^{z} \mathcal{N}(f|0,1) df$ denotes the cumulative Gaussian distribution. Finally, we have

$$\begin{split} \ln Z &= & \ln \left[\exp \left(- \sqrt{2} \tilde{\mu} \right) \Phi \left(\frac{m_-}{\tilde{\sigma}} \right) + \exp \left(\sqrt{2} \tilde{\mu} \right) \Phi \left(- \frac{m_+}{\tilde{\sigma}} \right) \right] + \tilde{\sigma}^2 - \ln \sqrt{2} \sigma_n \\ &= & \ln \left[\exp \left(\underbrace{\ln \Phi(-z_+) + \sqrt{2} \tilde{\mu}}_{\alpha_+} \right) + \exp \left(\underbrace{\ln \Phi(z_-) - \sqrt{2} \tilde{\mu}}_{\alpha_-} \right) \right] + \tilde{\sigma}^2 - \ln \sqrt{2} \sigma_n \\ &= & \ln (e^{\alpha_+} + e^{\alpha_-}) + \tilde{\sigma}^2 - \ln \sqrt{2} \sigma_n \end{split}$$

where $z_{+} = \frac{\tilde{\mu}}{\tilde{\sigma}} + \tilde{\sigma}\sqrt{2} = \frac{\mu - y}{\sigma} + \frac{\sigma}{\sigma_{n}}\sqrt{2}$, $z_{-} = \frac{\tilde{\mu}}{\tilde{\sigma}} - \tilde{\sigma}\sqrt{2} = \frac{\mu - y}{\sigma} - \frac{\sigma}{\sigma_{n}}\sqrt{2}$ and $\tilde{\mu} = \frac{\mu - y}{\sigma_{n}}$, $\tilde{\sigma} = \frac{\sigma}{\sigma_{n}}$. Now, using $\frac{d}{d\theta} \ln \Phi(z) = \frac{1}{\Phi(z)} \frac{d}{d\theta} \Phi(z) = \frac{N(z)}{\Phi(z)} \frac{dz}{d\theta}$ we tackle first derivative

$$\begin{split} \frac{\partial \ln Z}{\partial \mu} &= \frac{e^{\alpha_{+}} \frac{\partial \alpha_{+}}{\partial \mu} + e^{\alpha_{-}} \frac{\partial \alpha_{-}}{\partial \mu}}{e^{\alpha_{+}} + e^{\alpha_{-}}} \\ \frac{\partial \alpha_{+}}{\partial \mu} &= \frac{\partial}{\partial \mu} \ln \Phi(-z_{+}) + \frac{\sqrt{2}}{\sigma_{n}} \\ &= -\frac{\mathcal{N}(-z_{+})}{\sigma \Phi(-z_{+})} + \frac{\sqrt{2}}{\sigma_{n}} = -\frac{q_{+}}{\sigma} + \frac{\sqrt{2}}{\sigma_{n}} \\ \frac{\partial \alpha_{-}}{\partial \mu} &= \frac{\partial}{\partial \mu} \ln \Phi(z_{-}) - \frac{\sqrt{2}}{\sigma_{n}} \\ &= \frac{\mathcal{N}(z_{-})}{\sigma \Phi(z_{-})} - \frac{\sqrt{2}}{\sigma_{n}} = \frac{q_{-}}{\sigma} - \frac{\sqrt{2}}{\sigma_{n}} \\ \frac{\partial \alpha_{\pm}}{\partial \mu} &= \mp \frac{q_{\pm}}{\sigma} \pm \frac{\sqrt{2}}{\sigma_{n}}. \end{split}$$

as well as the second derivative

$$\begin{split} \frac{\partial^2 \ln Z}{\partial \mu^2} &= \frac{\frac{\partial}{\partial \mu} \left(e^{\alpha_+} \frac{\partial \alpha_+}{\partial \mu}\right) + \frac{\partial}{\partial \mu} \left(e^{\alpha_-} \frac{\partial \alpha_-}{\partial \mu}\right)}{e^{\alpha_+} + e^{\alpha_-}} - \left(\frac{\partial \ln Z}{\partial \mu}\right)^2 \\ \frac{\partial}{\partial \mu} \left(e^{\alpha_\pm} \frac{\partial \alpha_\pm}{\partial \mu}\right) &= e^{\alpha_\pm} \left[\left(\frac{\partial \alpha_\pm}{\partial \mu}\right)^2 + \frac{\partial^2 \alpha_\pm}{\partial \mu^2}\right] \\ \frac{\partial^2 \alpha_+}{\partial \mu^2} &= -\frac{1}{\sigma} \frac{\frac{\partial}{\partial \mu} \mathcal{N}(-z_+) \Phi(-z_+) - \frac{\partial}{\partial \mu} \Phi(-z_+) \mathcal{N}(-z_+)}{\Phi^2(-z_+)} \\ &= -\frac{1}{\sigma} \frac{\mathcal{N}(-z_+) \Phi(-z_+) \frac{\partial - z_+^2/2}{\partial \mu} - \mathcal{N}^2(-z_+) \frac{\partial - z_+}{\partial \mu}}{\Phi^2(-z_+)} \\ &= \frac{\mathcal{N}(-z_+)}{\sigma^2} \cdot \frac{\Phi(-z_+) z_+ - \mathcal{N}(-z_+)}{\Phi^2(-z_+)} = -\frac{q_+^2 - q_+ z_+}{\sigma^2} \\ \frac{\partial^2 \alpha_-}{\partial \mu^2} &= \frac{1}{\sigma} \frac{\frac{\partial}{\partial \mu} \mathcal{N}(z_-) \Phi(z_-) - \frac{\partial}{\partial \mu} \Phi(z_-) \mathcal{N}(z_-)}{\Phi^2(z_-)} \\ &= \frac{1}{\sigma} \frac{\mathcal{N}(z_-) \Phi(z_-) \frac{\partial - z_-^2/2}{\partial \mu} - \mathcal{N}^2(z_-) \frac{\partial z_-}{\partial \mu}}{\Phi^2(z_-)} \\ &= \frac{\mathcal{N}(z_-)}{\sigma^2} \cdot \frac{-\Phi(z_-) z_- - \mathcal{N}(z_-)}{\Phi^2(z_-)} = -\frac{q_-^2 + q_- z_-}{\sigma^2} \\ \frac{\partial^2 \alpha_\pm}{\partial \mu^2} &= -\frac{q_\pm^2 \mp q_\pm z_\pm}{\sigma^2} \end{split}$$

which can be simplified to

$$\frac{\partial^2 \ln Z}{\partial \mu^2} = \frac{e^{\alpha_+}b_+ + e^{\alpha_-}b_-}{e^{\alpha_+} + e^{\alpha_-}} - \left(\frac{\partial \ln Z}{\partial \mu}\right)^2$$

using

$$\begin{split} b_{\pm} &= \left(\frac{\partial \alpha_{\pm}}{\partial \mu}\right)^2 + \frac{\partial^2 \alpha_{\pm}}{\partial \mu^2} &= \left(\mp \frac{q_{\pm}}{\sigma} \pm \frac{\sqrt{2}}{\sigma_n}\right)^2 - \frac{q_{\pm}^2 \mp q_{\pm} z_{\pm}}{\sigma^2} \\ &= \left(\frac{q_{\pm}}{\sigma} - \frac{\sqrt{2}}{\sigma_n}\right)^2 - \frac{q_{\pm}^2}{\sigma^2} \pm \frac{q_{\pm} z_{\pm}}{\sigma^2} \\ &= \frac{2}{\sigma_n^2} - \left(\frac{\sqrt{8}}{\sigma \sigma_n} \mp \frac{z_{\pm}}{\sigma^2}\right) q_{\pm}. \end{split}$$

We also need

$$\frac{\partial \ln Z}{\partial \ln \sigma_n} = \frac{e^{\alpha_+} \frac{\partial \alpha_+}{\partial \ln \sigma_n} + e^{\alpha_-} \frac{\partial \alpha_-}{\partial \ln \sigma_n}}{e^{\alpha_+} + e^{\alpha_-}} - \frac{2\sigma^2}{\sigma_n^2} - 1.$$

Variational Bayes

We need $h(\gamma)$ and its derivatives as well as $\beta(\gamma)$:

$$h(\gamma) = \frac{2}{\sigma_n^2} \gamma + \ln(2\sigma_n^2) + y^2 \gamma^{-1}$$

$$h'(\gamma) = \frac{2}{\sigma_n^2} - y^2 \gamma^{-2}$$

$$h''(\gamma) = 2y^2 \gamma^{-3}$$

$$\beta(\gamma) = y \gamma^{-1}$$

4.10 Student's t Likelihood

The likelihood has two hyperparameters (both represented in the log domain to ensure positivity): the degrees of freedom ν and the scale σ_n with mean y (for $\nu > 1$) and variance $\frac{\nu}{\nu-2}\sigma_n^2$ (for $\nu > 2$).

$$p(y|f) = Z \cdot \left(1 + \frac{(f - y)^2}{\nu \sigma_n^2}\right)^{-\frac{\nu + 1}{2}}, \quad Z = \frac{\Gamma\left(\frac{\nu + 1}{2}\right)}{\Gamma\left(\frac{\nu}{2}\right)\sqrt{\nu \pi \sigma_n^2}}$$

Laplace's Approximation

For the mode fitting procedure, we need derivatives up to third order; the hyperparameter derivatives at the mode require some mixed derivatives. All in all, using r = y - f, we have

$$\begin{split} &\ln p(y|f) &= &\ln \Gamma \left(\frac{\nu+1}{2} \right) - \ln \Gamma \left(\frac{\nu}{2} \right) - \frac{1}{2} \ln \nu \pi \sigma_n^2 - \frac{\nu+1}{2} \ln \left(1 + \frac{r^2}{\nu \sigma_n^2} \right) \\ &\frac{\partial \ln p}{\partial f} &= &(\nu+1) \frac{r}{r^2 + \nu \sigma_n^2} \\ &\frac{\partial^2 \ln p}{(\partial f)^2} &= &(\nu+1) \frac{r^2 - \nu \sigma_n^2}{(r^2 + \nu \sigma_n^2)^2} \\ &\frac{\partial^3 \ln p}{(\partial f)^3} &= &2(\nu+1) \frac{r^3 - 3r\nu \sigma_n^2}{(r^2 + \nu \sigma_n^2)^3} \\ &\frac{\partial \ln p}{\partial \ln \nu} &= &\frac{\partial Z}{\partial \ln \nu} - \frac{\nu}{2} \ln \left(1 + \frac{r^2}{\nu \sigma_n^2} \right) + \frac{\nu+1}{2} \cdot \frac{r^2}{r^2 + \nu \sigma_n^2} \\ &\frac{\partial Z}{\partial \ln \nu} &= &\frac{\nu}{2} \frac{d \ln \Gamma \left(\frac{\nu+1}{2} \right)}{d \ln \nu} - \frac{\nu}{2} \frac{d \ln \Gamma \left(\frac{\nu}{2} \right)}{d \ln \nu} - \frac{1}{2} \\ &\frac{\partial^3 \ln p}{(\partial \ln \nu)(\partial f)^2} &= &\nu \frac{r^2(r^2 - 3(\nu+1)\sigma_n^2) + \nu \sigma_n^2}{(r^2 + \nu \sigma_n^2)^3} \\ &\frac{\partial \ln p}{\partial \ln \sigma_n} &= &(\nu+1) \frac{r^2}{r^2 + \nu \sigma_n^2} - 1 \\ &\frac{\partial^3 \ln p}{(\partial \ln \sigma_n)(\partial f)^2} &= &2\nu \sigma_n^2 (\nu+1) \frac{\nu \sigma_n^2 - 3r^2}{(r^2 + \nu \sigma_n^2)^3} \end{split}$$

4.11 Cumulative Logistic Likelihood

The likelihood has one hyperparameter (represented in the log domain), namely the standard deviation σ_n

$$p(y|f) = Z \cdot cosh^{-2} \left(\tau(f - y) \right), \ \tau = \frac{\pi}{2\sigma_n \sqrt{3}}, \ Z = \frac{\pi}{4\sigma_n \sqrt{3}}$$

Laplace's Approximation

The following derivatives are needed where $\phi(x) \equiv \ln(\cosh(x))$

$$\begin{array}{rcl} & \ln p(y|f) & = & \ln(\pi) - \ln(4\sigma_{\pi}\sqrt{3}) - 2\varphi\left(\tau(f-y)\right) \\ & \frac{\partial \ln p}{\partial f} & = & 2\tau\varphi'\left(\tau(f-y)\right) \\ & \frac{\partial^2 \ln p}{(\partial f)^2} & = & -2\tau^2\varphi''\left(\tau(f-y)\right) \\ & \frac{\partial^3 \ln p}{(\partial f)^3} & = & 2\tau^3\varphi'''\left(\tau(f-y)\right) \\ & \frac{\partial^3 \ln p}{(\partial \ln \sigma_{\pi})(\partial f)^2} & = & 2\tau^2\left(2\varphi''\left(\tau(f-y)\right) + \tau(f-y)\varphi'''\left(\tau(f-y)\right)\right) \\ & \frac{\partial \ln p}{\partial \ln \sigma_{\pi}} & = & 2\tau(f-y)\varphi'\left(\tau(f-y)\right) - 1 \end{array}$$

4.12 GLM Likelihoods: Poisson, Weibull, Gamma, Exponential, Inverse Gaussian and Beta

Data y from a space other than \mathbb{R} e.g. \mathbb{N} , \mathbb{R}_+ or [0,1] can be modeled using generalised linear model likelihoods p(y|f) where the expected value $\mathbb{E}[y] = \mu$ is related to the underlying Gaussian process f by means of an inverse link function $\mu = g(f)$. Typically, the likelihoods are from an exponential family, hence the variance $\mathbb{V}[y] = \nu(\mu)$, is a simple function of the mean μ as well as higher order moments such as skewness $\mathbb{S}[y] = s(\mu)$ and kurtosis $\mathbb{K}[y] = k(\mu)$.

Here, we directly specify the inverse link function $\mu = g(f)$ defining the mapping from the GP f to the mean intensity μ . For numerical reasons, we work with the log of the inverse link function $h(f) = \ln g(f)$ and use its derivatives h', h'' and h''' for subsequent computations. In the table below, we have summarised the GLM likelihood expressions, the moments, the range of their variables and the applicable inverse link functions.

Likelihood	$\rho =$	$\nu(\mu) =$	$s(\mu) =$	$k(\mu) =$	p(y f) =	y∈	$\mu \in$	Inverse Links
Poisson	Ø	μ	$1/\sqrt{\mu}$	1/μ	$\mu^{y} \exp(-\mu)/y!$	N	\mathbb{R}_{+}	exp, logistic
Weibull	{ln κ}	$\mu^2(\gamma_2/\gamma_1^2-1)$	$\frac{\gamma_3 - 3\gamma_1\gamma_2 + 2\gamma_1^3}{(\gamma_2 - \gamma_1^2)^{3/2}}$	$\frac{\gamma_4 - 4\gamma_1\gamma_3 + 12\gamma_1^2\gamma_2 - 3\gamma_2^2 - 6\gamma_1^4}{(\gamma_2 - \gamma_1^2)^2}$	$\kappa \gamma_1 / \mu (y \gamma_1 / \mu)^{\kappa - 1} \exp (-(y \gamma_1 / \mu)^{\kappa})$	$\mathbb{R}_+\backslash\{0\}$	$\mathbb{R}_+\backslash\{0\}$	exp, logistic
Gamma	$\{ln \ \alpha\}$	μ^2/α	$2/\sqrt{\alpha}$	6/α	$\frac{\alpha^{\alpha}y^{\alpha-1}}{\Gamma(\alpha)}\mu^{-\alpha}\exp\left(-\frac{y\alpha}{\mu}\right)$	$\mathbb{R}_+\backslash\{0\}$	$\mathbb{R}_+\backslash\{0\}$	exp, logistic
Exponential	Ø	μ^2	2	6	$\mu^{-1} \exp\left(-\frac{y}{\mu}\right)$	$\mathbb{R}_+\backslash\{0\}$	$\mathbb{R}_+\backslash\{0\}$	exp, logistic
Inv. Gauss	$\{ln\lambda\}$	μ^3/λ	$3\sqrt{\mu/\lambda}$	$15\mu/\lambda$	$\sqrt{\frac{\lambda}{2\pi y^3}} \exp\left(-\frac{\lambda(y-\mu)^2}{2\mu^2 y}\right)$	$\mathbb{R}_+\backslash\{0\}$	$\mathbb{R}_+\backslash\{0\}$	exp, logistic
Beta	{ln φ}	$\mu(1-\mu)/(1+\varphi)$	$\frac{(2-4\mu)(1+\phi)}{\sqrt{\nu(\mu)}(2+\phi)}$	$6\frac{(\phi+1)^2-\nu(\mu)(5\phi+6)}{\nu(\mu)(\phi+2)(\phi+3)}$	$\frac{\Gamma(\phi)}{\Gamma(\mu\phi)\Gamma((1-\mu)\phi)}y^{\mu\phi-1}(1-y)^{(1-\mu)\phi-1}$	[0, 1]	[0, 1]	expexp, logit

4.12.1 Inverse Link Functions

Possible inverse link functions and their properties (\cup convex, \cap concave, \uparrow monotone) are summarised below:

util/glm_invlink_*	$g(f) = \mu =$	$g:\mathbb{R} o$	g is	$h(f) = \ln \mu =$	h is
exp	e ^f	\mathbb{R}_+	∪,↑	f	∪,∩,↑
logistic	$ln(1+e^f)$	\mathbb{R}_{+}	∪,↑	$ln(ln(1+e^f))$	∩,↑
expexp	$\exp(-e^{-f})$	[0, 1]	↑	$-e^{-f}$	∪,↑
logit	$1/(1+e^{-f})$	[0, 1]	↑	$-\ln(1+e^{-f})$	∪,↑

Exponential inverse link: exp

For $g(f) = e^f$ things are simple since h(f) = f, h'(f) = 1 and h''(f) = h'''(f) = 0.

Logistic inverse link: logistic

For $g(f) = \ln(1 + e^f)$ the derivatives of h(f) are given by

$$\begin{array}{lcl} h(f) & = & \ln(\ln(1+e^f)) \\ h'(f) & = & \frac{1}{\ln(1+e^f)}s(-f), \ s(f) = \frac{1}{1+e^f}, \ s'(f) = \frac{-e^f}{(1+e^f)^2} = -s(-f)s(f) \\ h''(f) & = & \frac{1}{\ln(1+e^f)}\frac{e^{-f}}{(1+e^{-f})^2} - \frac{1}{\ln^2(1+e^f)}\frac{e^f}{1+e^f}\frac{1}{1+e^{-f}} \\ & = & h'(f)\left[s(f)-h'(f)\right] \\ h'''(f) & = & h''(f)\left[s(f)-h'(f)\right] + h'(f)\left[\frac{-e^f}{(1+e^f)^2}-h''(f)\right] \\ & = & h''(f)\left[s(f)-2h'(f)\right] - h'(f)s(f)s(-f). \end{array}$$

Note that $g(f) = e^{h(f)} = \ln(1 + e^f)$ is convex and $h(f) = \ln(\ln(1 + e^f))$ with

$$h''(f) = \frac{1}{\ln(1+e^f)} \left(1 - \frac{e^f}{\ln(1+e^f)}\right) \frac{1}{1+e^f} \frac{1}{1+e^{-f}} \leqslant 0$$

is concave since $e^f \ge \ln(1 + e^f)$ for all $f \in \mathbb{R}$.

Double negative exponential inverse link: expexp

For $g(f) = \exp(-e^{-f})$ the derivatives of h(f) are given by

$$h(f) = -e^{-f}$$
 $h'(f) = -h(f)$
 $h'''(f) = h(f)$
 $h''''(f) = -h(f)$

Logit regression inverse link: logit

For $g(f) = 1/(1+e^{-f})$ the derivatives of h(f) can be computed using the logistic inverse link function $h_{\ell}(f)$ since $h(f) = f - \exp(h_{\ell}(f))$

$$\begin{array}{lll} h(f) & = & f - e^{h_{\ell}(f)} \\ h'(f) & = & 1 - e^{h_{\ell}(f)} h'_{\ell}(f) \\ h''(f) & = & -e^{h_{\ell}(f)} [h'_{\ell}(f)^2 + h''_{\ell}(f)] = e^{h_{\ell}(f)} s_{\ell}(-f) s_{\ell}^2(f) \\ h'''(f) & = & -e^{h_{\ell}(f)} [h'_{\ell}(f)^3 + 3h''_{\ell}(f)h'_{\ell}(f) + h'''_{\ell}(f)] \end{array}$$

4.12.2 Poisson Likelihood

Count data $y \in \mathbb{N}^n$ can be modeled in the GP framework using the Poisson distribution $\mathfrak{p}(y) = \mu^y e^{-\mu}/y!$ with mean/variance $\mathbb{E}[y] = \mathbb{V}[y] = \mu$, skewness $\mathbb{S}[y] = 1/\sqrt{\mu}$ and kurtosis $\mathbb{K}[y] = 1/\mu$

leading to the likelihood

$$\begin{array}{rcl} p(y|f) & = & \mu^y \exp(-\mu)/y!, \; \mu = g(f) \\ \Leftrightarrow \ln p(y|f) & = & y \cdot \ln g(f) - g(f) - \ln \Gamma(y+1). \end{array}$$

For Laplace's method to work, we need the first three derivatives of the log likelihood $\ln p(y|f)$, where $h(f) = \ln g(f)$

Note that if $\ln \mu = h(f)$ is concave and $\mu = g(f)$ is convex then the Poisson likelihood p(y|f) is log-concave in f which is the case for both exp and logistic.

4.12.3 Weibull Likelihood

Nonnegative data $y \in \mathbb{R}_+$ such as time-to-failure can be modeled in the GP framework using the Weibull distribution $p(y) = \kappa/\lambda(y/\lambda)^{\kappa-1}e^{-(y/\lambda)^\kappa}$ with shape parameter $\kappa > 0$, scale parameter $\lambda > 0$, mean $\mathbb{E}[y] = \lambda \gamma_1 = \mu$ where $\gamma_j = \Gamma(1+j/\kappa)$, variance $\mathbb{V}[y] = \lambda^2 \gamma_2 - \mu^2 = \mu^2(\gamma_2/\gamma_1^2-1)$, skewness $\mathbb{S}[y] = (\gamma_3 - 3\gamma_1\gamma_2 + 2\gamma_1^3)/(\gamma_2 - \gamma_1^2)^{3/2}$ and kurtosis $\mathbb{K}[y] = (\gamma_4 - 4\gamma_1\gamma_3 + 12\gamma_1^2\gamma_2 - 3\gamma_2^2 - 6\gamma_1^4)/(\gamma_2 - \gamma_1^2)^2$. Using the substitution $\mu = \lambda \gamma_1 \Leftrightarrow 1/\lambda = \gamma_1/\mu$, we obtain

$$\begin{split} p(y|f) &= \gamma_1 \frac{\kappa}{\mu} \left(\gamma_1 \frac{y}{\mu} \right)^{\kappa-1} exp \left(- \left(\gamma_1 \frac{y}{\mu} \right)^{\kappa} \right), \; \mu = g(f) > 0 \\ \Leftrightarrow \ln p(y|f) &= \ln \left(\gamma_1 \frac{\kappa}{\mu} \right) + (\kappa - 1) \ln \left(\gamma_1 \frac{y}{\mu} \right) - \left(\gamma_1 \frac{y}{\mu} \right)^{\kappa}. \end{split}$$

Note that the Weibull likelihood p(y|f) is log-concave in f neither for the exp nor for the logistic inverse link.

4.12.4 Gamma Likelihood

Nonnegative data $y \in \mathbb{R}_+$ can be modeled in the GP framework using the Gamma distribution $p(y) = \theta^{-\alpha}/\Gamma(\alpha)y^{\alpha-1}e^{-y/\theta}$ with shape parameter $\alpha > 0$, scale parameter $\theta > 0$, mean $\mathbb{E}[y] = \alpha\theta = \mu$, variance $\mathbb{V}[y] = \alpha\theta^2 = \mu^2/\alpha$, skewness $\mathbb{S}[y] = 2/\sqrt{\alpha}$ and kurtosis $\mathbb{K}[y] = 6/\alpha$. Using the substitution $\mu = \alpha\theta \Leftrightarrow \alpha/\mu = 1/\theta$, we obtain

$$\begin{split} p(y|f) &= \frac{\alpha^{\alpha}y^{\alpha-1}}{\Gamma(\alpha)}\mu^{-\alpha}\exp\left(-\frac{y\alpha}{\mu}\right),\; \mu = g(f) > 0 \\ \Leftrightarrow \ln p(y|f) &= -\alpha\left(\ln \mu + \frac{y}{\mu}\right) - \ln Z_{\alpha}(y),\; \ln Z_{\alpha}(y) = \ln \Gamma(\alpha) - \alpha \ln \alpha + (1-\alpha) \ln y. \end{split}$$

Note that if $\ln \mu = h(f)$ was convex and $\mu = g(f)$ was concave then the Gamma likelihood p(y|f) would be log-concave in f which is not the case for both exp and logistic.

4.12.5 Exponential Likelihood

Nonnegative data $y \in \mathbb{R}_+$ can be modeled in the GP framework using the Exponential distribution $p(y) = \theta^{-1}e^{-y/\theta}$ with scale parameter $\theta > 0$, mean $\mathbb{E}[y] = \theta = \mu$, variance $\mathbb{V}[y] = \mu^2$, skewness $\mathbb{S}[y] = 2$ and kurtosis $\mathbb{K}[y] = 6$. We obtain

$$\begin{split} p(y|f) &= & \mu^{-1} \exp\left(-\frac{y}{\mu}\right), \; \mu = g(f) > 0 \\ \Leftrightarrow & \ln p(y|f) &= & -\ln \mu - \frac{y}{\mu}. \end{split}$$

Note that for exp (but not for logistic) the likelihood is log-concave. The exponential distribution corresponds to the Gamma distribution with $\alpha = 1$ and the Weibull distribution with $\kappa = 1$.

4.12.6 Inverse Gaussian Likelihood

Nonnegative data $y \in \mathbb{R}^n_+$ can be modeled in the GP framework using the Inverse Gaussian distribution $p(y) = \sqrt{\lambda/(2\pi y^3)} \exp(-\lambda(y-\mu)^2/(2\mu^2 y))$ with shape parameter $\lambda > 0$, mean parameter $\mu > 0$, mean $\mathbb{E}[y] = \mu$, variance $\mathbb{V}[y] = \mu^3/\lambda$, skewness $\mathbb{S}[y] = 3\sqrt{\mu/\lambda}$ and kurtosis $\mathbb{K}[y] = 15\mu/\lambda$. We obtain

$$\begin{split} p(y|f) &=& \sqrt{\frac{\lambda}{2\pi y^3}} \exp\left(-\frac{\lambda (y-\mu)^2}{2\mu^2 y}\right), \; \mu = g(f) > 0 \\ \Leftrightarrow & \ln p(y|f) &=& -\frac{\lambda (y-\mu)^2}{2\mu^2 y} - \ln Z_\alpha(y), \; \ln Z_\alpha(y) = -\frac{1}{2} (\ln \lambda - \ln 2\pi y^3). \end{split}$$

The inverse Gaussian likelihood is in general not log-concace in f for both exp and logistic.

4.12.7 Beta Likelihood

Interval data $y \in [0,1]^n$ can be modeled in the GP framework using the Beta distribution $p(y) = y^{\alpha-1}(1-y)^{\beta-1}/B(\alpha,\beta)$ with shape parameters $\alpha,\beta>0$, mean $\mathbb{E}[y]=\alpha/(\alpha+\beta)$ and variance $\mathbb{V}[y]=\alpha\beta/[(\alpha+\beta)^2(\alpha+\beta+1)]$ and $1/B(\alpha,\beta)=\Gamma(\alpha+\beta)/[\Gamma(\alpha)\Gamma(\beta)]$. Reparametrising using the mean parameter $\mu=\mathbb{E}[y]=\alpha/(\alpha+\beta)$, the shape parameter $\phi=\alpha+\beta$, the variance $\mathbb{V}[y]=\mu(1-\mu)/(1+\varphi)$ and hence

$$\begin{split} p(y|f) &=& \frac{\Gamma(\varphi)}{\Gamma(\mu\varphi)\Gamma((1-\mu)\varphi)} y^{\mu\varphi-1} (1-y)^{(1-\mu)\varphi-1}, \; \mu=g(f)>0 \\ \Leftrightarrow & \ln p(y|f) &=& \ln \Gamma(\varphi) - \ln \Gamma(\mu\varphi) - \ln \Gamma((1-\mu)\varphi) + (\mu\varphi-1) \ln y + ((1-\mu)\varphi-1) \ln (1-y). \end{split}$$

The Beta likelihood is in general not log-concace in f for both exp and logistic.

5 Mean Functions

A mean function $\mathfrak{m}_{\Phi}: \mathfrak{X} \to \mathbb{R}$ (with hyperparameters Φ) of a GP f is a scalar function defined over the whole domain \mathfrak{X} that computes the expected value $\mathfrak{m}(\mathbf{x}) = \mathbb{E}[f(\mathbf{x})]$ of f for the input \mathbf{x} .

5.1 Interface

In the GPML toolbox, a mean function $m: \mathcal{X} \to \mathbb{R}$ needs to implement evaluation $m = m_{\varphi}(X)$ and first derivatives $m_i = \frac{\partial}{\partial \varphi_i} m$ with respect to the components i of the parameter $\varphi \in \Phi$ as detailed below.

```
\langle meanFunctions.m 34 \rangle \equiv
34
      1 % mean functions to be use by Gaussian process functions. There are two
      2 % different kinds of mean functions: simple and composite:
      3 %
      4 % simple mean functions:
      5 %
      6 %
          meanZero - zero mean function
      7 % meanOne
                          - one mean function
      8 % meanConst
                          - constant mean function
     9 % meanLinear - linear mean function
10 % meanPoly - polynomial mean function
     11 % meanDiscrete - precomputed mean for discrete data
     12 % meanGP - predictive mean of another GP
     13 % meanGPexact - predictive mean of a regression GP
     14 \% meanNN - nearest neighbor mean function
     15 %
     16 % composite covariance functions (see explanation at the bottom):
     17 %
     18 % meanScale - scaled version of a mean function
19 % meanPow - power of a mean function
20 % meanProd - products of mean functions
21 % meanSum - sums of mean functions
     22 % meanMask
                          - mask some dimensions of the data
     23 % meanPref
                         - difference mean for preference learning
     25 % Naming convention: all mean functions are named "mean/mean*.m".
     26 %
     27 %
     28 % 1) With no or only a single input argument:
     29 %
     30 %
             s = meanNAME or s = meanNAME(hyp)
     31 %
     32 % The mean function returns a string s telling how many hyperparameters hyp it
     33\, % expects, using the convention that "D" is the dimension of the input space.
     34 % For example, calling "meanLinear" returns the string 'D'.
     35 %
     36 % 2) With two input arguments:
     37 %
     38 %
             m = meanNAME(hyp, x)
     39 %
     40 % The function computes and returns the mean vector where hyp are the
     41 % hyperparameters and x is an n by D matrix of cases, where D is the dimension
     42 % of the input space. The returned mean vector is of size n by 1.
```

44 % 3) With three input arguments:

```
45 %
46 % dm = meanNAME(hyp, x, i)
47 %
48 % The function computes and returns the n by 1 vector of partial derivatives
49 % of the mean vector w.r.t. hyp(i) i.e. hyperparameter number i.
50 %
51 % See also doc/usageMean.m.
52 %
53 \( \langle \text{pml copyright 6a} \rangle \)
```

5.2 Implemented Mean Functions

We offer simple and composite mean functions producing new mean functions m(x) from existing mean functions $\mu_j(x)$. All code files are named according to the pattern mean/mean<NAME>.m for simple identification. This modular specification allows to define affine mean functions $m(x) = c + a^{\top}x$ or polynomial mean functions $m(x) = (c + a^{\top}x)^2$. All currently available mean functions are summarised in the following table.

Simple mean functions $m(x)$							
<name></name>	Meaning	$m(\mathbf{x}) =$	ф				
Zero	mean vanishes always	0	Ø				
One	mean equals 1	1	Ø				
Const	mean equals a constant	С	$c\in\mathbb{R}$				
Linear	mean linearly depends on $\mathbf{x} \in \mathcal{X} \subseteq \mathbb{R}^D$	$\mathbf{a}^{T}\mathbf{x}$	$\mathbf{a} \in \mathbb{R}^{\mathrm{D}}$				
Poly	mean polynomially depends on $\mathbf{x} \in \mathcal{X} \subseteq \mathbb{R}^D$	$\sum_{\mathbf{d}} \mathbf{a}_{\mathbf{d}}^{T} \mathbf{x}^{\mathbf{d}}$	$\mathbf{a} \in \mathbb{R}^{\mathrm{D} imes \mathrm{d}}$				
Discrete	precomputed mean for discrete data $\mathbf{x} \in \mathfrak{X} \subseteq \mathbb{N}$	$\mu_{\mathbf{x}}$	$\mu \in \mathbb{R}^s$				
GP	predictive mean of another GP	$\int \mathbf{y} \cdot \mathbf{p}(\mathbf{y} \mathcal{D}, \mathbf{x}) d\mathbf{y}$	Ø				
GPexact	predictive mean of a regression GP	$\int \mathbf{y} \cdot \mathbf{p}(\mathbf{y} \mathcal{D}, \mathbf{x}) d\mathbf{y}$	ρ, ψ, σ_n				
NN	nearest neighbor for a set $(\mathbf{z_j}, \mathbf{m_j}) \in \mathcal{X} \times \mathbb{R}$	m_i , $i = arg min_j d(x, z_j)$	Ø				
Composite	mean functions $[\mu_1(\mathbf{x}), \mu_2(\mathbf{x}),] \mapsto \mathfrak{m}(\mathbf{x})$						
<name></name>	Meaning	$m(\mathbf{x}) =$	ф				
Scale	scale a mean	$\alpha\mu(\mathbf{x})$	$lpha \in \mathbb{R}$				
Sum	add up mean functions	$\sum_{j} \mu_{j}(\mathbf{x})$	Ø				
Prod	multiply mean functions	$\prod_{j} \mu_{j}(\mathbf{x})$	Ø				
Pow	raise a mean to a power	$\mu(\mathbf{x})^{d}$	Ø				
Mask	act on components $I \subseteq [1, 2,, D]$ of $\mathbf{x} \in \mathcal{X} \subseteq \mathbb{R}^D$ only	$\mu(\mathbf{x}_{\mathrm{I}})$	Ø				
Pref	preference learning mean $\mathbf{x} = [\mathbf{x}_1; \mathbf{x}_2], \ \mathbf{x}_i \subseteq \mathbb{R}^{D/2}$	$\mu(\mathbf{x}_1) - \mu(\mathbf{x}_2)$	Ø				

5.3 Usage of Implemented Mean Functions

Some code examples taken from doc/usageMean.m illustrate how to use simple and composite mean functions to specify a GP model.

Syntactically, a mean function mf is defined by

```
mn := 'func' | @func // simple
mf := {mn} | {mn, {param, mf}} | {mn, {mf, .., mf}} // composite
```

i.e., it is either a string containing the name of a mean function, a pointer to a mean function or one of the former in combination with a cell array of mean functions and an additional list of parameters.

 $\langle doc/usageMean.m \ 35 \rangle \equiv$

35

```
1 % demonstrate usage of mean functions
2 %
 3 % See also meanFunctions.m.
 4 %
 5 (gpml copyright 6a)
 6 clear all, close all
 7 n = 5; D = 2; x = randn(n,D);
                                    % create a random data set
9 % set up simple mean functions
12 mc = {@meanConst}; hypc = 2; % also function handles are possible
13 ml = {@meanLinear}; hypl = [2;3];
                                        % m(x) = 2*x1 + 3*x2
14 mp = {@meanPoly,2}; hypp = [1;1;2;3]; % m(x) = x1+x2+2*x1^2+3*x2^2
15 mn = {@meanNN,[1,0; 0,1],[0.9,0.5]}; hypn = []; % nearest neighbor
16 s = 12; hypd = randn(s,1);
                                     % discrete mean with 12 hypers
17 md = {'meanDiscrete',s};
18 \text{ hyp.cov} = [0;0]; \text{ hypg} = [];
                                                % GP predictive mean
19 xt = randn(2*n,D); yt = sign(xt(:,1)-xt(:,2));
                                                  % training data
20 mg = {@meanGP,hyp,@infEP,@meanZero,@covSEiso,@likErf,xt,yt};
21 hype = [0;0; log(0.1)]; % regression GP predictive mean
22 xt = randn(2*n,D); yt = xt(:,1).*xt(:,2);
                                                    % training data
23 me = {@meanGPexact,@meanZero,@covSEiso,xt,yt};
2.4
25 % set up composite mean functions
26 msc = {'meanScale', {m1}};
                                hypsc = [3; hyp1]; % scale by 3
27 msu = {'meanSum', {m0,mc,ml}}; hypsu = [hyp0; hypc; hyp1]; % sum
28 \text{ mpr} = \{0\text{meanProd}, \{\text{mc,ml}\}\};
                                29 mpo = {'meanPow',3,msu};
                                hyppo = hypsu;
                                                      % third power
30 mask = [false,true]; % mask excluding all but the 2nd component
31 mma = {'meanMask', mask, ml}; hypma = hypl(mask);
32 mpf = {@meanPref,ml};
                                hyppf = 2; % linear pref with slope
33
34 % 0) specify mean function
35 % mean = md; hyp = hypd; x = randi([1,s],n,1);
36 \% mean = mn; hyp = hypn;
37 % mean = mg; hyp = hypg;
38 mean = me; hyp = hype;
39 \% \text{ mean} = m0; \text{ hyp} = \text{hyp0};
40 % mean = msu; hyp = hypsu;
41 % mean = mpr; hyp = hyppr;
42 % mean = mpo; hyp = hyppo;
43 % mean = mpf; hyp = hyppf;
45 \% 1) query the number of parameters
46 feval(mean{:})
47
48 \% 2) evaluate the function on x
49 feval(mean{:},hyp,x)
51 % 3) compute the derivatives w.r.t. to hyperparameter i
52 i = 2; feval(mean{:},hyp,x,i)
```

6 Covariance Functions

A covariance function $k_{\psi}: \mathcal{X} \times \mathcal{X} \to \mathbb{R}$ (with hyperparameters ψ) of a GP f is a scalar function defined over the whole domain \mathcal{X}^2 that computes the covariance $k(\mathbf{x}, \mathbf{x}') = \mathbb{V}[f(\mathbf{x}), f(\mathbf{x}')] = \mathbb{E}[(f(\mathbf{x}) - m(\mathbf{x}))(f(\mathbf{x}') - m(\mathbf{x}'))]$ of f between the inputs \mathbf{x} and \mathbf{x}' .

6.1 Interface

Again, the interface is simple since only evaluation of the full covariance matrix $K = k_{\psi}(X)$ and its derivatives $K_i = \frac{\partial}{\partial \psi_i} K$ as well as cross terms $k_* = k_{\psi}(X, x_*)$ and $k_{**} = k_{\psi}(x_*, x_*)$ for prediction are required.

```
\langle covFunctions.m \ 37 \rangle \equiv
37
        1 % covariance functions to be use by Gaussian process functions. There are two
       2 % different kinds of covariance functions: simple and composite:
       3 %
       4 % simple covariance functions:
       5 % covConst - covariance for constant functions
       6 % covCos
                                 - sine periodic covariance function (1d) with unit period
       7 % covLIN - linear covariance function without parameters
8 % covLINard - linear covariance function with ARD
9 % covLINiso - linear covariance function
      10 % covLINone - linear covariance function with bias
      11 % covMaternard - Matern covariance function with nu=1/2, 3/2 or 5/2 with ARD
      12 \% covMaterniso - Matern covariance function with nu=1/2, 3/2 or 5/2
      13 % covNNone - neural network covariance function
      14 % covNoise - independent covariance function (i.e. white noise)
      15 % covPeriodic - smooth periodic covariance function (1d)
      16 % covPeriodicNoDC - as above but with zero DC component and properly scaled
      17 \% covPoly - polynomial covariance function
      18 % covPPard - piecewise polynomial covariance function (compact support)
19 % covPPiso - piecewise polynomial covariance function (compact support)
      20 % covRQard
                                - rational quadratic covariance function with ARD
      21 % covRQiso
                                - isotropic rational quadratic covariance function
      covRQiso - isotropic rational quadratic covariance function

covSEard - squared exponential covariance function with ARD

covSEiso - isotropic squared exponential covariance function

covSEisoU - same as above but without latent scale

covSEvlen - spatially varying lengthscale squared exponential

covSEfact - factor analysis squared exponential covariance function

covSM - spectral mixture covariance function
      28 % covGaborard - Gabor covariance function with ARD
      29 %
             covGaborsio - isotropic Gabor covariance function
      30 %
              covDiscrete - precomputed covariance for discrete data
      31 %
      32 % composite (meta) covariance functions (see explanation at the bottom):
      33 % covScale - scaled version of a covariance function
      34 % covProd
                                - products of covariance functions
      35 % covSum
                                - sums of covariance functions
      36 % covADD - additive covariance function
37 % covMask - mask some dimensions of the data
38 % covPERard - make ARD stationary covariance periodic
39 % covPERiso - make isotropic stationary covariance periodic
40 % covPref - difference covariance for preference learning
      41 %
      42 % special purpose (wrapper) covariance functions
```

- to be used in conjunction with infFITC* for large scale

```
44 %
                       inference problems; any covariance can be wrapped by
45 %
                       covFITC such that the FITC approximation is applicable
                     - to be used in conjunction with infGrid* for large scale
46 %
     covGrid
47 %
                       inference problems on grids resulting Kronecker structure
48 %
49 % Naming convention: all covariance functions are named "cov/cov*.m". A trailing
50~\% "iso" means isotropic, "ard" means Automatic Relevance Determination, and
51 % "one" means that the distance measure is parameterized by a single parameter.
52 %
 53 % The covariance functions are written according to a special convention where
54 % the exact behaviour depends on the number of input and output arguments
55 % passed to the function. If you want to add new covariance functions, you
56 % should follow this convention if you want them to work with the function gp.
 57 % There are four different ways of calling the covariance functions:
59 % 1) With no (or one) input argument(s):
60 %
61 %
      s = cov
62 %
63 % The covariance function returns a string s telling how many hyperparameters it
64 % expects, using the convention that "D" is the dimension of the input space.
65 % For example, calling "covRQard" returns the string '(D+2)'.
67 % 2) With two input arguments:
69 %
        K = cov(hyp, x) equivalent to K = cov(hyp, x, [])
70 %
 71 % The function computes and returns the covariance matrix where hyp are
72 % the hyperparameters and x is an n by D matrix of cases, where
73 % D is the dimension of the input space. The returned covariance matrix is of
74 % size n by n.
 75 %
76 % 3) With three input arguments:
 77 %
78 %
      Ks = cov(hyp, x, xs)
 79 %
        kss = cov(hyp, xs, 'diag')
80 %
81 % The function computes test set covariances; kss is a vector of self covariances
82 % for the test cases in xs (of length ns) and Ks is an (n by ns) matrix of cross
 83 % covariances between training cases x and test cases xs.
 85 % 4) With four input arguments:
86 %
 87 %
        dKi = cov(hyp, x, [], i)
88 %
        dKsi = cov(hyp, x, xs, i)
 89 %
        dkssi = cov(hyp, xs, 'diag', i)
90 %
91 % The function computes and returns the partial derivatives of the
92 % covariance matrices with respect to hyp(i), i.e. with
 93 % respect to the hyperparameter number i.
 94 %
 95 % Covariance functions can be specified in two ways: either as a string
96 % containing the name of the covariance function or using a cell array. For
97 % example:
98 %
99 % cov = 'covRQard';
100 % cov = {'covRQard'};
101 % cov = {@covRQard};
```

```
102 %
103 % are supported. Only the second and third form using the cell array can be used
104 % for specifying composite covariance functions, made up of several
105 % contributions. For example:
106 %
107 %
            cov = {'covScale', {'covRQiso'}};
108 %
           cov = {'covSum', {'covRQiso','covSEard','covNoise'}};
109 %
            cov = {'covProd',{'covRQiso','covSEard','covNoise'}};
110 %
            cov = {'covMask',{mask,'covSEiso'}}
111 % q=1; cov = {'covPPiso',q};
112 % d=3; cov = {'covPoly',d};
113 %
            cov = {'covADD',{[1,2],'covSEiso'}};
114 %
            cov = {@covFITC, {@covSEiso}, u}; where u are the inducing inputs
115 %
116 % specifies a covariance function which is the sum of three contributions. To
117 % find out how many hyperparameters this covariance function requires, we do:
118 %
119 %
      feval(cov{:})
120 %
121 % which returns the string '3+(D+1)+1' (i.e. the 'covRQiso' contribution uses
122\ \% 3 parameters, the 'covSEard' uses D+1 and 'covNoise' a single parameter).
123 %
124 % See also doc/usageCov.m.
125 %
126 (gpml copyright 6a)
```

6.2 Implemented Covariance Functions

Similarly to the mean functions, we provide a whole algebra of covariance functions $k: \mathcal{X} \times \mathcal{X} \to \mathbb{R}$ with the same generic name pattern cov/cov<NAME>.m as before.

Besides a long list of simple covariance functions, we also offer a variety of composite covariance functions as shown in the following table.

Simple covariar	nce functions k(x,x')		
<name></name>	Meaning	$k(\mathbf{x}, \mathbf{x}') =$	ψ
Zero	covariance vanishes always	0	ø
Eye	unit additive measurement noise	$\delta(\mathbf{x} - \mathbf{x}')$	Ø
Noise	additive measurement noise	$\sigma_f^2 \delta(\mathbf{x} - \mathbf{x}')$	ln σ _f
Const	covariance equals a constant	σ_{ϵ}^2	ln σ _f
LIN	linear, $\mathfrak{X} \subseteq \mathbb{R}^{D}$	$\mathbf{x}^{\dagger}\mathbf{x}'$	0
LINard	linear with diagonal weighting, $\mathfrak{X} \subseteq \mathbb{R}^{D}$	$\mathbf{x}^{T} \mathbf{\Lambda}^{-2} \mathbf{x}'$	$\{\ln \lambda_1,, \ln \lambda_D\}$
LINiso	linear with isotropic weighting, $\mathfrak{X} \subseteq \mathbb{R}^{D}$	$\mathbf{x}^{\top}\mathbf{x}'/\ell^2$	ln ℓ
LINone	linear with bias, $\mathfrak{X} \subseteq \mathbb{R}^{D}$	$(\mathbf{x}^{\top}\mathbf{x}'+1)/\ell^2$	In ℓ
Poly	polynomial covariance, $\mathfrak{X} \subseteq \mathbb{R}^{D}$	$\sigma_{\epsilon}^{2}(\mathbf{x}^{\top}\mathbf{x}'+\mathbf{c})^{d}$	$\{\ln c, \ln \sigma_f\}$
SEard	automatic relevence determination squared exponential, $\mathfrak{X} \subseteq \mathbb{R}^{D}$	$\sigma_f^2 \exp\left(-\frac{1}{2}(\mathbf{x} - \mathbf{x}')^{\top} \mathbf{\Lambda}^{-2}(\mathbf{x} - \mathbf{x}')\right)$	$\{\ln \lambda_1,, \ln \lambda_D, \ln \sigma_f\}$
SEiso	diagonal squared exponential, $\mathfrak{X} \subseteq \mathbb{R}^{D}$	$\sigma_f^2 \exp\left(-\frac{1}{2\ell^2}(\mathbf{x} - \mathbf{x}')^{\top}(\mathbf{x} - \mathbf{x}')\right)$	$\{\ln \ell, \ln \sigma_f\}$
SEisoU	squared exponential, $\mathfrak{X} \subseteq \mathbb{R}^{D}$	$\exp(-\frac{1}{2g^2}\mathbf{x}^{\top}\mathbf{x}')$	In £
SEvlen	spatially varying lengthscale squared exponential $\mathfrak{X}\subseteq\mathbb{R}^D$	$\sigma_f^2\left(\frac{a}{b}\right)^{\frac{3b}{2}}\exp\left(-\frac{\ \mathbf{x}-\mathbf{x}'\ ^2}{b}\right), a = 2\ell(\mathbf{x})\ell(\mathbf{x}'), b = \ell^2(\mathbf{x}) + \ell^2(\mathbf{x}')$	$\{\phi_{\ell}, \ln \sigma_f\}$
SEfact	factor analysis squared exponential $\mathfrak{X} \subseteq \mathbb{R}^{D}$	$\sigma_f^2 \exp\left(-\frac{1}{2}(\mathbf{x} - \mathbf{x}')^\top \mathbf{L}^\top \mathbf{L}(\mathbf{x} - \mathbf{x}')\right), \mathbf{L} \in \mathbb{R}^{d \times D}, d \leq D$	$\{L, \ln \sigma_f\}$
RQard	rational quadratic, $\mathfrak{X} \subseteq \mathbb{R}^{D}$	$ \frac{\sigma_f^2 \left(1 + \frac{1}{2\alpha} (\mathbf{x} - \mathbf{x}')^\top \mathbf{\Lambda}^{-2} (\mathbf{x} - \mathbf{x}')\right)^{-\alpha}}{\sigma_f^2 \left(1 + \frac{1}{2\alpha f^2} (\mathbf{x} - \mathbf{x}')^\top (\mathbf{x} - \mathbf{x}')\right)^{-\alpha}} $	$\{\ln \lambda_1,, \ln \lambda_D, \ln \sigma_f, \ln \alpha\}$
RQiso	rational quadratic, $\mathfrak{X} \subseteq \mathbb{R}^{D}$	$\sigma^{\frac{1}{2}}(1+\frac{1}{2}(\mathbf{x}-\mathbf{x}')^{\top}(\mathbf{x}-\mathbf{x}'))^{-\alpha}$	$\{\ln \ell, \ln \sigma_f, \ln \alpha\}$
Maternard	Matérn, $\mathfrak{X} \subseteq \mathbb{R}^{D}$, $f_{1}(t) = 1$, $f_{3}(t) = 1 + t$, $f_{5}(t) = f_{3}(t) + \frac{t^{2}}{3}$	$\frac{\sigma_f^2 f_d(\mathbf{r}_d) \exp(-\mathbf{r}_d), \mathbf{r}_d = \sqrt{d(\mathbf{x} - \mathbf{x}')^{\top} \mathbf{\Lambda}^{-2} (\mathbf{x} - \mathbf{x}')}}{\sigma_f^2 f_d(\mathbf{r}_d) \exp(-\mathbf{r}_d), \mathbf{r}_d = \sqrt{d(\mathbf{x} - \mathbf{x}')^{\top} \mathbf{\Lambda}^{-2} (\mathbf{x} - \mathbf{x}')}}$	$\{\ln \lambda_1,, \ln \lambda_D, \ln \sigma_f\}$
Materniso	Matérn, $\mathfrak{X} \subseteq \mathbb{R}^D$, $f_1(t) = 1$, $f_3(t) = 1 + t$, $f_5(t) = f_3(t) + \frac{t^2}{3}$	$\sigma_{\mathbf{f}}^{2} f_{\mathbf{d}}(\mathbf{r}_{\mathbf{d}}) \exp(-\mathbf{r}_{\mathbf{d}}), \ \mathbf{r}_{\mathbf{d}} = \sqrt{\frac{\mathbf{d}}{\ell^{2}} (\mathbf{x} - \mathbf{x}')^{\top} (\mathbf{x} - \mathbf{x}')}$	$\{\ln \ell, \ln \sigma_f\}$
NNone	neural net, $\mathcal{X} \subseteq \mathbb{R}^D$, $f(\mathbf{x}) = 1 + \mathbf{x}^\top \mathbf{\Lambda}^{-2} \mathbf{x}$	$\sigma_f^2 \sin^{-1} \left(\frac{\mathbf{x}^T \mathbf{A}^{-2} \mathbf{x}'}{\sqrt{f(\mathbf{x})f(\mathbf{x}')}} \right)$	$\{\ln \ell, \ln \sigma_f\}$
Periodic	periodic, $\mathfrak{X}\subseteq\mathbb{R}$	$\sigma_{\rm f}^2 \exp\left(-\frac{2}{\ell^2}\sin^2\left[\pi \ x - x'\ /p\right]\right)$	$\{\ln \ell, \ln p, \ln \sigma_f\}$
PeriodicNoDC	periodic, $\mathfrak{X} \subseteq \mathbb{R}$, rescaled and DC component removed	$\sigma_f^2 \frac{\kappa(x-x') - \frac{1}{\pi} \int_0^\pi \kappa(t) dt}{\kappa(0) - \frac{1}{\pi} \int_0^\pi \kappa(t) dt}, \kappa(t) = \exp\left(-\frac{2}{\ell^2} \sin^2[\pi t/p]\right)$	$\{\ln \ell, \ln p, \ln \sigma_f\}$
Cos	periodic cosine, $\mathfrak{X} \subseteq \mathbb{R}$	$\sigma_{\rm f}^2 \cos\left(\pi \left\ x - x'\right\ /p\right]\right)$	$\{\ln p, \ln \sigma_f\}$
PPard	compact support, piecewise polynomial $f_{\nu}(r)$, $\mathfrak{X} \subseteq \mathbb{R}^{D}$,	$\sigma_f^2 \max(0, 1 - r)^{j+\nu} \cdot f_{\nu}(r), r^2 = (\mathbf{x} - \mathbf{x}')^{\top} \mathbf{\Lambda}^{-2} (\mathbf{x} - \mathbf{x}')$	$\{\ln \lambda_1,, \ln \lambda_D, \ln \sigma_f\}$
PPiso	compact support, piecewise polynomial $f_{\nu}(r)$, $\mathfrak{X} \subseteq \mathbb{R}^{D}$,	$\sigma_f^2 \max(0, 1-r)^{j+\nu} \cdot f_{\nu}(r), r = \frac{\ \mathbf{x} - \mathbf{x}'\ }{\ell}, j = \left\lfloor \frac{D}{2} \right\rfloor + \nu + 1$	$\{\ln \ell, \ln \sigma_f\}$
SM	spectral mixture, $\mathfrak{X}\subseteq\mathbb{R}^D$, $\mathbf{w}\in\mathbb{R}_+^Q$, $\mathbf{M},\mathbf{V}\in\mathbb{R}_+^{D\times Q}$	$\mathbf{w}^{\top} \left(\prod_{d=1}^{D} \exp(-\frac{1}{2} \mathbf{v}_{d} \mathbf{r}^{2}) \odot \cos(\mathbf{m}_{d} \mathbf{r}) \right), \mathbf{r} = 2\pi \ \mathbf{x} - \mathbf{x}'\ $	$\{\ln w, \ln M, \ln V\}$
	spectral mixture, $\mathfrak{X} \subseteq \mathbb{R}^D$, $\mathbf{W} \in \mathbb{R}_+^{D \times Q}$, $\mathbf{M}, \mathbf{V} \in \mathbb{R}_+^{D \times Q}$	$\prod_{d=1}^{D} \mathbf{w}_{d}^{\top} \left(\exp(-\frac{1}{2} \mathbf{v}_{d} \tau^{2}) \odot \cos(\mathbf{m}_{d} \tau) \right), \ \tau = 2\pi \ \mathbf{x} - \mathbf{x}'\ $	{ln W, ln M, ln V}
Gaborard	anisotropic Gabor function, $\mathfrak{X} \subseteq \mathbb{R}^D$, $\lambda, p \in \mathbb{R}^D_+$	$\exp\left(-\sum_{d=1}^{D} \frac{t_{d}^{2}}{2\lambda_{d}^{2}}\right) \cos\left(2\pi \sum_{d=1}^{D} t_{d}/p_{d}\right), t_{d} = x_{d} - x_{d}'$	$\{\ln \lambda, \ln p\}$
Gaboriso	isotropic Gabor function, $\mathfrak{X} \subseteq \mathbb{R}^{D}$, $\ell, \mathfrak{p} \in \mathbb{R}_{+}$	$\exp(-\frac{\mathbf{t}^{T}\mathbf{t}}{2\ell^2})\cos(2\pi\mathbf{t}^{T}1/p), \mathbf{t} = \mathbf{x} - \mathbf{x}'$	{ln ℓ, ln p}
Discrete	precomputed covariance for discrete data $\mathbf{x} \in \mathfrak{X} \subseteq \mathbb{N}$	$k_{xx'}$ where $K = L^{T}L$ is Cholesky decomposition, $L \in \mathbb{R}^{s \times s}$	{L}
Composite cova	ariance functions $[\kappa_1(\mathbf{x}, \mathbf{x}'), \kappa_2(\mathbf{x}, \mathbf{x}'),] \mapsto k(\mathbf{x}, \mathbf{x}')$		
<name></name>	Meaning	$k(\mathbf{x}, \mathbf{x}') =$	ψ
Scale	scale a covariance	$\sigma_f^2 \kappa(\mathbf{x}, \mathbf{x}')$ and $\sigma_f(\mathbf{x}) \kappa(\mathbf{x}, \mathbf{x}') \sigma_f(\mathbf{x}')$	$\{\ln \sigma_f\}$
Sum	add up covariance functions	$\sum_{i} \kappa_{i}(\mathbf{x}, \mathbf{x}')$	Ø
Prod	multiply covariance functions	$\prod_{i} \kappa_{i}(\mathbf{x}, \mathbf{x}')$	Ø
PERard	turn ARD stationary into a periodic, $\mathfrak{X} \subseteq \mathbb{R}^{D}$	$\kappa(\mathbf{u}(\mathbf{x}), \mathbf{u}(\mathbf{x}'))$, $\mathbf{u}(\mathbf{x}) = [\sin \mathbf{x}_{\mathbf{p}}, \cos \mathbf{x}_{\mathbf{p}}]$, $\mathbf{x}_{\mathbf{p}} = 2\pi \operatorname{diag}(\mathbf{p}^{-1})\mathbf{x}$	{ln p ₁ ,,ln p _D }
PERiso	turn isotropic stationary into a periodic, $\mathfrak{X} \subseteq \mathbb{R}^D$	$\kappa(\mathbf{u}(\mathbf{x}), \mathbf{u}(\mathbf{x}'))$, $\mathbf{u}(\mathbf{x}) = [\sin \mathbf{x}_{p}, \cos \mathbf{x}_{p}]$, $\mathbf{x}_{p} = 2\pi \mathbf{x}/p$	lnp
Mask	act on components $I \subseteq [1, 2,, D]$ of $\mathbf{x} \in \mathcal{X} \subseteq \mathbb{R}^D$ only	$\kappa(\mathbf{x}_{\mathrm{I}},\mathbf{x}_{\mathrm{I}}')$	Ø
ADD	additive, $\mathfrak{X} \subseteq \mathbb{R}^{D}$, index degree set $\mathfrak{D} = \{1,, D\}$	$\sum_{d \in \mathcal{D}} \sigma_{f_d}^2 \sum_{ I =d} \prod_{i \in I} \kappa(x_i, x_i'; \psi_i)$	$\{\psi_1,, \psi_D, \ln \sigma_{f_1},, \ln \sigma_{f_{ \mathcal{D} }}\}$
Pref	preference learning covariance $\mathbf{x} = [\mathbf{x}_1; \mathbf{x}_2], \ \mathbf{x}_i \subseteq \mathbb{R}^{D/2}$	$\kappa(\mathbf{x}_1, \mathbf{x}_1') + \kappa(\mathbf{x}_2, \mathbf{x}_2') - \kappa(\mathbf{x}_1, \mathbf{x}_2') - \kappa(\mathbf{x}_2, \mathbf{x}_1')$	0

The spectral mixture covariance covSM was introduced by Wilson & Adams Gaussian Process Kernels for Pattern Discovery and Extrapolation, ICML, 2013.

The periodic covariance functions covPERiso and covPERard start from a stationary isotropic or ARD covariance function that depends on the data only through a distance $\mathbf{r}^2 = (\mathbf{x} - \mathbf{x}')^{\top} \mathbf{\Lambda}^{-2} (\mathbf{x} - \mathbf{x}')$ such as covMatern*, covPP*, covRQ*, covSE* where *=ard|iso and turn them into a periodic covariance function by embedding the data $\mathbf{x} \in \mathbb{R}^D$ into a periodic high-dimensional space $\mathbf{x}_p = \mathbf{u}(\mathbf{x}) \in \mathbb{R}^{2D}$ by a function $\mathbf{u}(\mathbf{x}) = 2\pi \text{diag}(\mathbf{p}^{-1})\mathbf{x}$.

The additive covariance function covADD starts from a one-dimensional covariance function $\kappa(x_i, x_i', \psi_i)$ acting on a single component $i \in [1, ..., D]$ of x. From that, we define covariance functions $\kappa_I(x_I, x_I) = \prod_{i \in I} \kappa(x_i, x_i', \psi_i)$ acting on vector-valued inputs x_I . The sums of exponential size can efficiently be computed using the Newton-Girard formulae. Samples functions drawn from a GP with additive covariance are additive functions. The number of interacting variables |I| is a measure of how complex the additive functions are.

6.3 Usage of Implemented Covariance Functions

Some code examples taken from doc/usageCov.m illustrate how to use simple and composite covariance functions to specify a GP model.

Syntactically, a covariance function cf is defined by

```
cv := 'func' | @func // simple
cf := {cv} | {cv, {param, cf}} | {cv, {cf, .., cf}} // composite
```

i.e., it is either a string containing the name of a covariance function, a pointer to a covariance function or one of the former in combination with a cell array of covariance functions and an additional list of parameters.

```
\langle doc/usageCov.m \ 41 \rangle \equiv
41
     1 % demonstrate usage of covariance functions
     3 % See also covFunctions.m.
     4 %
     5 (gpml copyright 6a)
     6 clear all, close all
     7 \text{ n} = 5; D = 3; x = \text{randn}(n,D); xs = \text{randn}(3,D); % create a data set
     9 % set up simple covariance functions
    10 cn = {'covNoise'}; sn = .1; hypn = log(sn); % one hyperparameter
    11 cc = {@covConst}; sf = 2; hypc = log(sf); % function handles OK
    12 \text{ ce} = \{@covEye\};
                                    hype = [];
    13 \text{ cl} = \{\text{@covLIN}\};
                                    hypl = []; % linear is parameter-free
    14 cla = {'covLINard'}; L = rand(D,1); hypla = log(L); % linear (ARD)
    16 clo = {@covLINone}; ell = .9; hyplo = log(ell); % linear with bias
    17 cp = \{(covPoly, 3)\}; c = 2; hypp = \log([c;sf]); % third order poly
    18 cga = {@covSEard}; hypga = log([L;sf]);
                                                    % Gaussian with ARD
    20 cgu = {'covSEisoU'}; hypgu = log(ell);  % isotropic Gauss no scale
    21 cra = {'covRQard'}; al = 2; hypra = log([L;sf;al]); % ration. quad.
    22 cri = {@covRQiso};
                                 hypri = log([ell;sf;al]);  % isotropic
    23 cma = {@covMaternard,5}; hypma = log([ell;sf]); % Matern class d=5
    24 cmi = {'covMaterniso',3}; hypmi = log([ell;sf]); % Matern class d=3
    25 cnn = {'covNNone'}; hypnn = log([L;sf]);
                                                         % neural network
    26 cpe = {'covPeriodic'}; p = 2; hyppe = log([ell;p;sf]);  % periodic
    27 cpn = {'covPeriodicNoDC'}; p = 2; hyppe = log([ell;p;sf]); % w/o DC
    28 cpc = {'covCos'}; p = 2; hypcpc = log([p;sf]);
                                                           % cosine cov
    29 cca = {'covPPard',3}; hypcc = hypgu;% compact support poly degree 3
    30 cci = {'covPPiso',2}; hypcc = hypgi; % compact support poly degree 2
    31 cgb = {@covGaboriso}; ell = 1; p = 1.2; hypgb=log([ell;p]); % Gabor
    32 Q = 2; w = ones(Q,1)/Q; m = rand(D,Q); v = rand(D,Q);
    33 csm = \{@covSM,Q\}; hypsm = log([w;m(:);v(:)]);
                                                     % Spectral Mixture
    34 cvl = {@covSEvlen, {@meanLinear}}; hypvl = [1;2;1; 0]; % var lenscal
    35 s = 12; cds = {@covDiscrete,s}; % discrete covariance function
    36 L = randn(s); L = chol(L'*L); L(1:(s+1):end) = log(diag(L));
    37 hypds = L(triu(true(s))); xd = randi([1,s],[n,1]); xsd = [1;3;6];
    38 cfa = \{\emptyset covSEfact, 2\}; hypfa = randn(D*2,1); % factor analysis
    40~\% set up composite i.e. meta covariance functions
    41 \csc = {\text{`covScale'}, \{cgu\}\}}; hypsc = [log(3); hypgu]; % scale by 9
    42 csu = {'covSum', {cn,cc,cl}}; hypsu = [hypn; hypc; hypl];
    43 cpr = {@covProd,{cc,cci}}; hyppr = [hypc; hypcc];
    44 mask = [0,1,0]; % binary mask excluding all but the 2nd component
```

```
45 cma = {'covMask', {mask, cgi{:}}}; hypma = hypgi;
46 % isotropic periodic rational quadratic
47 cpi = {'covPERiso',{@covRQiso}};
48 % periodic Matern with ARD
49 cpa = {'covPERard',{@covMaternard,3}};
50 % additive based on SEiso using unary and pairwise interactions
51 cad = {'covADD',{[1,2],'covSEiso'}};
52 % preference covariance with squared exponential base covariance
53 cpr = {'covPref', {'covSEiso'}}; hyppr = [0;0];
54 \text{ xp} = \text{randn}(n,2*D); \text{ xsp} = \text{randn}(3,2*D);
55
56 % 0) specify a covariance function
57 % cov = cma; hyp = hypma;
58 \% \text{ cov} = \text{cci; hyp} = \text{hypcc;}
59 \% \text{ cov} = \text{csm}; \text{hyp} = \text{hypsm};
60 cov = cds; hyp = hypds; x = xd; xs = xsd;
61 % cov = cfa; hyp = hypfa;
62 % cov = cvl; hyp = hypvl;
63 \% \text{ cov} = \text{cpr}; \text{ hyp} = \text{hyppr}; \text{ x} = \text{xp}; \text{ xs} = \text{xsp};
65 % 1) query the number of parameters
66 feval(cov{:})
68 \% 2) evaluate the function on x
69 feval(cov{:},hyp,x)
70
71 \% 3) evaluate the function on x and xs to get cross-terms
72 kss = feval(cov{:},hyp,xs,'diag')
73 Ks = feval(cov{:},hyp,x,xs)
75~\% 4) compute the derivatives w.r.t. to hyperparameter i
76 i = 1; feval(cov{:},hyp,x,[],i)
```

7 Hyperpriors

A hyperprior $p(\theta)$ with $\theta = [\rho, \varphi, \psi]$ is a joint probability distribution over the likelihood hyperparameters ρ , the mean hyperparameters φ and the covariance hyperparameters ψ . We concentrate on factorial priors $p(\theta) = \prod_j p_j(\theta_j)$. Hyperpriors can be used to regularise the optimisation of the hyperparameters via the marginal likelihood $Z(\theta)$ so that $p(\theta)Z(\theta)$ is maximised instead. As we wish to perform unconstrained optimisation, we require (mainly) smooth hyperpriors with infinite support.

7.1 Interface

In the GPML toolbox, a prior distribution $p(\theta)$ needs to implement the evaluation of the log density $\ln p(\theta)$ and its first derivative $\frac{\partial}{\partial \theta} \ln p(\theta)$. In addition, we require sampling capabilities i.e. the generation of $\theta \sim p(\theta)$.

```
42  ⟨priorDistributions.m 42⟩≡
   1 % prior distributions to be used for hyperparameters of Gaussian processes
   2 % using infPrior.
   3 % There are two different kinds of prior distributions: simple and composite:
   4 %
   5 % simple prior distributions:
```

```
6 %
7 % priorGauss
8 % priorLaplace
 7 %
                         - univariate Gaussian
                            - univariate Laplace
9 % priorT
                            - univariate Student's t
10 %
11 % priorSmoothBox1 - univariate interval (linear decay in log domain)
12 % priorSmoothBox2 - univariate interval (quadr. decay in log domain)
13 %
14 % priorGamma
                            - univariate Gamma, IR+
15 % priorWeibull
16 % priorInvGauss
                            - univariate Weibull, IR+
                            - univariate Inverse Gaussian, IR+
17 % priorLogNormal
                            - univariate Log-normal, IR+
18 %
     priorClamped or
19 %
                            - fix hyperparameter to its current value by setting
20 % priorDelta
                          derivatives to zero, no effect on marginal likelihood
21 %
22 % priorGaussMulti - multivariate Gauss
23 % priorLaplaceMulti - multivariate Laplace
24 % priorTMulti - multivariate Student's t
25 %
26 %
     priorClampedMulti or - fix hyperparameter to its current value by setting
27 %
     priorDeltaMulti derivatives to zero, no effect on marginal likelihood
28 %
29 \% composite prior distributions (see explanation at the bottom):
30 %
31 % priorMix
                             - nonnegative mixture of priors
32 %
     priorTransform
                            - prior on g(t) rather than t
33 %
34 % Naming convention: all prior distributions are named "prior/prior*.m".
35 %
36 %
37 % 1) With only a fixed input arguments:
39 %
        r = priorNAME(par1,par2,parN)
40 %
41 % The function returns a random sample from the distribution for e.g.
42 % random restarts, simulations or optimisation initialisation.
43 %
44 % 2) With one additional input arguments:
45 %
        [lp,dlp] = priorNAME(par1,par2,parN, t)
46 %
47 %
48 % The function returns the log density at location t along with its first
49 % derivative.
50 %
51 % See also doc/usagePrior.m, inf/infPrior.m.
52 %
53 (gpml copyright 6a)
```

7.2 Implemented Hyperpriors

All code files are named according to the pattern prior/prior<NAME>.m for simple identification. All currently available hyperpriors are summarised in the following table.

Simple hyperpriors $p(\theta)$						
Univariate hyperpriors defined over the whole reals with mean μ and variace σ^2						
<name></name>	Meaning	$p(\theta) =$	τ			
Gauss	normally distributed hyperparameter $\theta \in \mathbb{R}$	$\frac{1}{\sigma\sqrt{2\pi}}\exp\left(-\frac{(\theta-\mu)^2}{2\sigma^2}\right)$ $\frac{1}{2b}\exp\left(-\frac{(\theta-\mu)}{b}\right), b = \sigma/\sqrt{2}$	$\mu \in \mathbb{R}, \sigma^2 \in \mathbb{R}_+$			
Laplace	double exponentially hyperparameter $\theta \in \mathbb{R}$		$\mu \in \mathbb{R}, \sigma^2 \in \mathbb{R}_+$			
Т	Student's t distributed hyperparameter $\theta \in \mathbb{R}$	$\frac{\Gamma(\frac{\nu+1}{2})}{\Gamma(\frac{\nu}{2})}\frac{1}{\sqrt{(\nu-2)\pi\sigma}}\left(1+\frac{(\theta-\mu)^2}{(\nu-2)\sigma^2}\right)^{-\frac{\nu+1}{2}}$	$\mu \in \mathbb{R}, \sigma^2, u \in \mathbb{R}_+$			
Univariate hype	erpriors with effective bounded support but defin	ned over the whole real line				
SmoothBox1	interval hyperparameter $\theta \in \mathbb{R}$ i.e. $\theta \stackrel{\approx}{\in} [a,b]$	$\begin{vmatrix} \frac{1 - \exp(\eta(a - b))}{b - a} \cdot \frac{1}{1 + \exp(-\eta(\theta - a))} \cdot \frac{1}{1 + \exp(\eta(\theta - b))} \\ \mu = \frac{a + b}{2}, \sigma^2 = \frac{w^2}{1 - \exp(-2a)} \frac{1 + \pi^2/g^2}{12}, g = \frac{w\eta}{2}, w = a - b \end{vmatrix}$	$a\leqslant b\in\mathbb{R},\eta\in\mathbb{R}_{+}$			
SmoothBox2	localised hyperparameter $\theta \in \mathbb{R}$ i.e. $\theta \stackrel{\approx}{\in} [\mathfrak{a},\mathfrak{b}]$	$\begin{split} & \frac{1-\exp(\eta(a-b))}{b-a} \cdot \frac{1}{1+\exp(-\eta(\theta-a))} \cdot \frac{1}{1+\exp(\eta(\theta-b))} \\ & \mu = \frac{a+b}{2}, \ \sigma^2 = \frac{w^2}{1-\exp(-2g)} \frac{1+\pi^2/g^2}{12}, \ g = \frac{w\eta}{2}, \ w = a-b \\ & \frac{1}{(1/\eta+1)(b-a)} \begin{cases} \mathcal{N}(\theta a, \sigma^2_{ab}) & t < a \\ 1 & t \in [a,b], \ \sigma_{ab} = \frac{b-a}{\eta\sqrt{2\pi}} \end{cases} \\ & \mathcal{N}(\theta a, \sigma^2_{ab}) & b < t \end{cases} \\ & \mu = \frac{a+b}{2}, \ \sigma^2 = \frac{w^2}{4} \frac{\eta^3/3+\eta^2+4\eta/\pi+2/\pi}{\eta^3+\eta^2}, \ w = a-b \end{split}$	$a\leqslant b\in\mathbb{R},\eta\in\mathbb{R}_{+}$			
Univariate hype	erpriors supported only over the positive reals					
Gamma	Gamma hyperparameter $\theta \in \mathbb{R}_+$	$\frac{1}{\Gamma(k)t^k} \exp(\frac{\theta}{t})\theta^{k-1}$	$k \in \mathbb{R}_+, t \in \mathbb{R}_+$			
Weibull	Weibull hyperparameter $\theta \in \mathbb{R}_+$	$\frac{k}{\lambda} \left(\frac{\theta}{\lambda} \right)^{k-1} \exp \left(-(\frac{\theta}{\lambda})^k \right)$	$k \in \mathbb{R}_+, \lambda \in \mathbb{R}_+$			
InverseGauss	inverse Gaussian hyperparameter $\theta \in \mathbb{R}_+$	$\begin{array}{l} \frac{1}{\Gamma(k)t^k} \exp(\frac{\theta}{t}) \theta^{k-1} \\ \frac{k}{\lambda} \left(\frac{\theta}{\lambda}\right)^{k-1} \exp\left(-\left(\frac{\theta}{\lambda}\right)^k\right) \\ \frac{1}{2\pi\theta^3/\lambda} \exp\left(-\frac{\lambda(\theta-\mu)^2}{2\mu^2\theta}\right) \end{array}$	$k\in\mathbb{R}_+,\lambda\in\mathbb{R}_+$			
LogNormal	log-normal hyperparameter $\theta \in \mathbb{R}_+$	$\mathcal{N}(\theta \mu, \sigma^2) = \frac{1}{\theta \sigma \sqrt{2\pi}} \exp\left(-\frac{(\ln \theta - \mu)^2}{2\sigma^2}\right)$	$\mu \in \mathbb{R}, \sigma^2 \in \mathbb{R}_+$			
Multivariate hy	perpriors supported all over \mathbb{R}^D with mean μ a					
GaussMulti	multivariate normal distribution $\theta \in \mathbb{R}^D$	$ 2\pi\mathbf{\Sigma} ^{-\frac{1}{2}}\exp\left(-\frac{1}{2}(\mathbf{\theta}-\mathbf{\mu})^{\top}\mathbf{\Sigma}^{-1}(\mathbf{\theta}-\mathbf{\mu})\right)$	$\mu \in \mathbb{R}^{D}, \Sigma \in \mathbb{R}^{D \times D}$			
LaplaceMulti	multivariate Laplace distribution $\theta \in \mathbb{R}^D$	$ \sqrt{2}\Sigma ^{-\frac{1}{2}}\exp\left(-\sqrt{2}\left\ L^{-1}(\theta-\mu)\right\ _{1}\right), L^{\top}L=\Sigma$	$\mu \in \mathbb{R}^{D}, \Sigma \in \mathbb{R}^{D \times D}$			
TMulti	multivariate Student's t distribution $\boldsymbol{\theta} \in \mathbb{R}^D$	$ (\nu-2)\pi\pmb{\Sigma} ^{-\frac{1}{2}\frac{\Gamma(\frac{\nu+D}{2})}{\Gamma(\frac{\nu}{2})}}\left(1+\frac{(\theta-\mu)^{\top}\pmb{\Sigma}^{-1}(\theta-\mu)}{(\nu-2)}\right)^{-\frac{\nu+D}{2}}$	$\boldsymbol{\mu} \in \mathbb{R}^{D}, \boldsymbol{\Sigma} \in \mathbb{R}^{D \times D}, \boldsymbol{\nu} \in \mathbb{R}$			
Improper hyper	priors used to fix the value of a particular hyper	rparameter				
Delta	clamped hyperparameter $\theta = \theta_0 \in \mathbb{R}$	$\delta(\theta - \theta_0)$	0			
Clamped	amped					
DeltaMulti	clamped hyperparameter $\theta = \theta_0 \in \mathbb{R}^D$	$\delta(\theta - \theta_0)$	0			
ClampedMulti	ClampedMulti					
Composite hyperpriors $[\pi_1(\theta), \pi_2(\theta),] \mapsto p(\theta)$						
Transform	prior distribution on $g(\theta)$ instead of θ	$\pi(g(\theta))$	{g}			
Mix	mixture distribution	$\sum_{i} w_{i} \pi_{i}(\theta)$	{ w }			

The priorSmoothBox2 is a Gauss-uniform sandwich obtained by complementing a uniform distribution on [a,b] with two Gaussian halves at each side. The parameter η balances the probability mass between the constituents so that $\eta/(\eta+1)$ is used for the box and $1/(\eta+1)$ for the Gaussian sides. Its brother priorSmoothBox1 is the product of two sigmoidal functions.

The priorDelta or equivalently priorClamped can be used to exclude some hyperparameters from the optimisation. Their values are clamped to θ_0 and the derivative vanishes. There are also multivariate counterparts priorDeltaMulti and priorClampedMulti.

7.3 Usage of Implemented Hyperpriors

Some code examples taken from doc/usagePrior.m illustrate how to use univariate, multivariate and composite priors on hyperparameters. Syntactically, a hyperprior hp is defined by

```
hp := {pr} | {pr, {param, hp}} | {pr, {hp, ..., hp}} // composite
```

i.e., it is either a string containing the name of a hyperprior function, a pointer to a hyperprior function or one of the former in combination with a cell array of hyperprior functions and an additional list of parameters. Furthermore, we have multivariate hyperprior variants and 2 (equivalent) predefined hyperpriors allowing to exclude variables from optimisation.

```
45
     \langle doc/usagePrior.m \ 45 \rangle \equiv
     1 % demonstrate usage of prior distributions
     2 %
     3 % See also priorDistributions.m.
     5 (gpml copyright 6a)
     6 clear all, close all
     8 % 1) specify some priors
     9 % a) univariate priors
    10 \text{ mu} = 1.0; s2 = 0.01^2; nu = 3;
    11 pg = {@priorGauss,mu,s2};
                                                         % Gaussian prior
    12 pl = {'priorLaplace', mu, s2};
                                                          % Laplace prior
    13 pt = {@priorT, mu, s2, nu};
                                                      % Student's t prior
    14 p1 = {@priorSmoothBox1,0,3,15}; % smooth box constraints lin decay
    15 p2 = {@priorSmoothBox2,0,2,15}; % smooth box constraints qua decay
    16 pd = {'priorDelta'}; % fix value of prior exclude from optimisation
    17 pc = {@priorClamped};
                                                    % equivalent to above
    18 \text{ lam} = 1.05; k = 2.5;
    19 pw = {@priorWeibull,lam,k};
                                                          % Weibull prior
    20
    21 % b) meta priors
    23 g = 0exp; dg = 0exp; ig = 0log;
    25
    26 % c) multivariate priors
    27 \text{ m} = [1;2]; V = [2,1;1,2];
    28 pG = {@priorGaussMulti,m,V};
                                                      % 2d Gaussian prior
    29 pD = {'priorDeltaMulti'};  % fix value of prior exclude from optim
    30 pC = {@priorClampedMulti};
                                                    % equivalent to above
    31
    32 % 2) evaluation
    33 % pri = pt; hp = randn(1,3);
    34 \% pri = pmx; hp = randn(1,3);
    35 % pri = ptr; hp = randn(1,3);
    36 \text{ pri} = pG; \quad hp = randn(2,3);
    37
    38 % a) draw a sample from the prior
    39 feval(pri{:})
    41 \% b) evaluate prior and derivative if requires
    42 [lp,dlp] = feval(pri{:},hp)
    44 % 3) comprehensive example
    45 \times = (0:0.1:10); y = 2*x + randn(size(x)); % generate training data
    46 mean = {@meanSum,{@meanConst,@meanLinear}}; % specify mean function
    47 cov = {@covSEiso}; lik = {@likGauss}; % specify covariance and lik
    48 hyp.cov = [log(1);log(1.2)]; hyp.lik = log(0.9); hyp.mean = [2;3];
    49 par = {mean,cov,lik,x,y}; mfun = @minimize; % input for GP function
    50
```

```
51 % a) plain marginal likelihood optimisation (maximum likelihood)
52 im = @infExact;
                                                   % inference method
53 hyp_plain = feval(mfun, hyp, @gp, -10, im, par{:});
                                                           % optimise
55~\% b) regularised optimisation (maximum a posteriori) with 1d priors
56 prior.mean = {pg;pc}; % Gaussian prior for first, clamp second par
57 prior.cov = {p1;[]}; % box prior for first, nothing for second par
58 im = {@infPrior,@infExact,prior};
                                          % inference method
59 hyp_p1 = feval(mfun, hyp, @gp, -10, im, par{:});
                                                           % optimise
60
61~\% c) regularised optimisation (maximum a posteriori) with Nd priors
62 prior = [];
                                                % clear the structure
63 % multivariate Student's t prior on the first and second mean hyper
64 prior.multi{1} = {@priorTMulti,[mu;mu],diag([s2,s2]),nu,...
65
                   struct('mean',[1,2])};
                                                   % use hyper struct
66 % Equivalent shortcut (same mu and s2 for all dimensions)
67 prior.multi{1} = {@priorTMulti,mu,s2,nu,struct('mean',[1,2])};
68 % multivariate Gaussian prior jointly on 1st and 3rd hyper
69 prior.multi{2} = {@priorGaussMulti,[mu;mu],diag([s2,s2]),...
                    [1,3]};
                                         % use unwrapped hyper vector
71 % Equivalent shortcut (same mu and s2 for all dimensions)
72 prior.multi{2} = {@priorGaussMulti,mu,s2,[1,3]};
                                                   % inference method
73 im = {@infPrior,@infExact,prior};
74 hyp_pN = feval(mfun, hyp, @gp, -10, im, par{:});
                                                           % optimise
76 [unwrap(hyp), unwrap(hyp_plain), unwrap(hyp_p1), unwrap(hyp_pN)]
```