# gprege Quick Guide

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#### 1 Abstract

The *gprege* package implements our methodology of Gaussian process regression models for the analysis of microarray time series. The package can be used to filter quiet genes and quantify differential expression in time-series expression ratios or for ranking candidate targets of a trascription factor. The purpose of this quick guide is to present a few examples of using gprege.

# 2 Citing gprege

Citing gprege in publications will involve citing the methodology paper [Kalaitzis and Lawrence, 2011] that the software is based on as well as citing the software package itself.

# 3 Introductory example analysis - TP63 microarray data

In this section we introduce the main functions of the *gprege* package by repeating some of the analysis from the BMC Bioinformatics paper [Kalaitzis and Lawrence, 2011].

#### 3.1 Installing the *gprege* package

The recommended way to install *gprege* is to use the biocLite function available from the bioconductor website. Installing in this way should ensure that all appropriate dependencies are met.

- > source("http://www.bioconductor.org/biocLite.R")
- > biocLite("gprege")

Otherwise, to manually install the gprege software, unpack the software and run

R CMD INSTALL gprege

To load the package start R and run

> library(gprege)

## 3.2 Loading the data

For convenience, a fragment of the preprocessed data is provided in this package and can be loaded using

> data(FragmentDellaGattaData)

The full dataset is available at http://staffwww.dcs.shef.ac.uk/people/A. Kalaitzis/DellaGattaData.RData and can be loaded using

- > # Download full Della Gatta dataset.
- > load(url('http://staffwww.dcs.shef.ac.uk/people/A.Kalaitzis/DellaGattaData.RData'))

Type ?DellaGattaData for more details on the dataset. If you need to preprocess some gene expression time-series from scratch and the dataset originates from Affymetrix arrays, we highly recommend processing it with rma or mmgmos from the affy and puma packages respectively. mmgmos extracts error bars on the expression measurements directly from the array data to allow judging the reliability of individual measurements. A detailed guide on how to perform these steps can be found in the tiqre Quick Guide.

#### 3.3 Interactive mode

Let us now examine one by one, in an interactive fashion, a few profiles of the full dataset (not available in the package). We start by downloading it.

- > # Download full Della Gatta dataset.
- > con <- url('http://staffwww.dcs.shef.ac.uk/people/A.Kalaitzis/DellaGattaData.RData')
- > load(con); close.connection(con)
- > # Timepoints / GP inputs.
- > tTrue = matrix(seq(0,240,by=20), ncol=1)
- > gpregeOptions <- list()</pre>

These gene expression profiles correspond to the top ranks as direct targets suggested by TSNI [Della Gatta et al., 2008].

- > # Set index range so that only a top few targets suggested by TSNI be explored.
- > gpregeOptions\$indexRange <- which(DGatta\_labels\_byTSNItop100)[1:2]</pre>

If the download fails, use the following instead.

- > # Load installed fragment-dataset.
- > data(FragmentDellaGattaData)
- > # Fragment dataset is comprised of top-ranked targets suggested by TSNI.
- > # Explore only the first few.
- > gpregeOptions\$indexRange <- 1:2</pre>

The option gpregeOptions\$explore<-TRUE makes gprege wait for a keystroke to proceed to the next profile.

```
# Explore individual profiles in interactive mode.
>
    gpregeOptions$explore <- TRUE</pre>
    # Exhaustive plot resolution of the LML function.
   gpregeOptions$exhaustPlotRes <- 30</pre>
    # Exhaustive plot contour levels.
    gpregeOptions$exhaustPlotLevels <- 10</pre>
    # Exhaustive plot maximum lengthscale.
    gpregeOptions$exhaustPlotMaxWidth <- 100</pre>
    # Noisy ground truth labels: which genes are in the top 786 ranks of the TSNI ranking.
   gpregeOptions$labels <- DGatta_labels_byTSNItop100</pre>
   # SCG optimisation: maximum number of iterations.
    gpregeOptions$iters <- 100</pre>
    # SCG optimisation: no messages.
    gpregeOptions$display <- FALSE</pre>
    # Matrix of different hyperparameter configurations as rows:
                             percent-signal-variance percent-noise-variance].
    # [inverse-lengthscale
   gpregeOptions$inithypers <-</pre>
      matrix( c( 1/1000,
                                   1e-3,
                                                0.999,
                   1/30.
                                0.999,
                                               1e-3,
                   1/80,
                                2/3, 1/3
    ), ncol=3, byrow=TRUE)
```

gprege will generate a report and a few plots for each explored profile. The first line contains the index of the profile in exprs\_tp63\_RMA and its ground truth (according to TSNI). The hyperparameters of the optimised log-marginal likelihood (LML) follow and then the initialisation (indicated by '<-') from which it came. Then a few statistics; the observed standard deviation of the time-series and the sum of the explained std.deviation (by the GP) and noise (Gaussian) std.deviation. Finally the function exhaustivePlot will reveal, through an exhaustive search in the hyperparemeter space<sup>1</sup>, the approximate true maximum LML and corresponding hyperparameters.

> gpregeOutput<-gprege(data=exprs\_tp63\_RMA,inputs=tTrue,gpregeOptions=gpregeOptions)

Profile 249		Label: 1
Length-scale 84.73909	Signal 0.22901	Noise 0.03297
Init.le 1000 30	LML 2.94548585 9.87342036	Best

 $<sup>^1 {\</sup>tt gpregeOptions\$exhaustPlotMaxWidth~defines~the~limit~of~the~length scale~search~range}.$ 

80 16.69472737 <--

Total st.dev. Estim.sig + noise

0.200493 0.261986

Log-ratio (max(LML[2:end]) - LML[1])

13.74924

======= EXHAUSTIVE LML SEARCH ==========

Length-scale Signal Noise 72.68966 0.16579 0.03470

Max LML Estim. sig + noise

16.32234561 0.200493

ENTER to continue

------

Profile 370 Label: 1

Length-scale Signal Noise 185.38847 0.37661 0.10492

Init.le LML Best

 1000
 0.24660390

 30
 2.47031897

 80
 6.37127488

80 6.37127488 <--

Total st.dev. Estim.sig + noise

0.246752 0.481527

Log-ratio (max(LML[2:end]) - LML[1])

6.124671

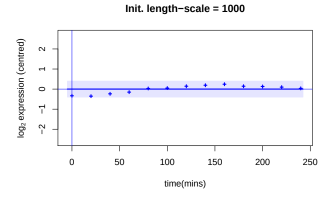
====== EXHAUSTIVE LML SEARCH ===========

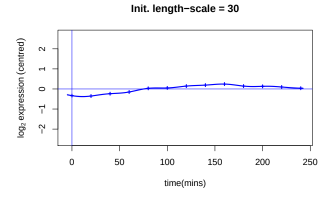
Length-scale Signal Noise 62.44828 0.15310 0.09366

Max LML Estim. sig + noise

5.08007151 0.246752

ENTER to continue





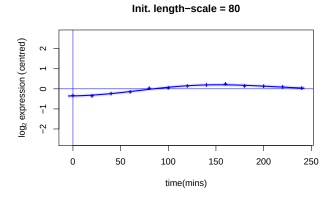
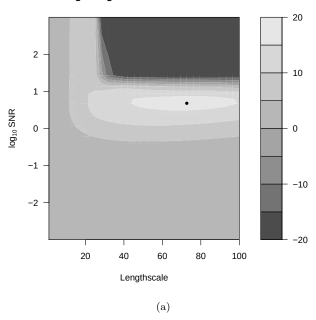


Figure 1: GP fit with different initialisations on profile #249.

### Log-marginal likelihood function



# max LML length-scale = 72.7

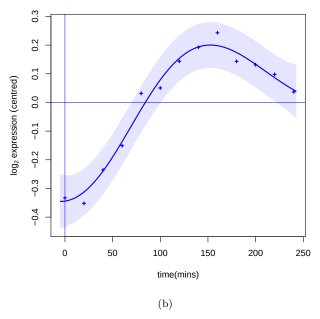
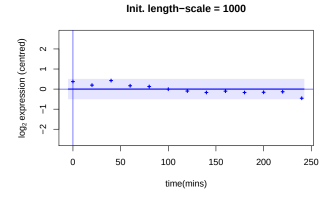
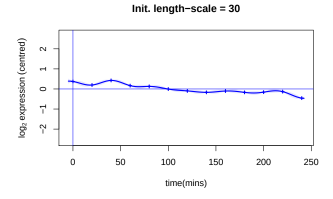


Figure 2: Profile #249: (a) Log-marginal likelihood (LML) contour. (b) GP fit with maximum LML hyperparameters from the exhaustive search.





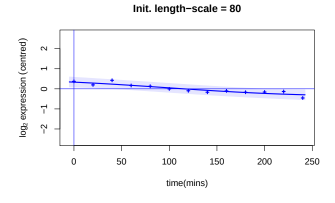
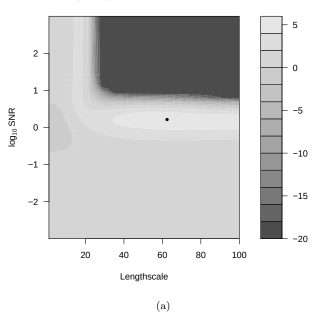


Figure 3: GP fit with different initialisations on profile #370.

### Log-marginal likelihood function



# max LML length-scale = 62.4

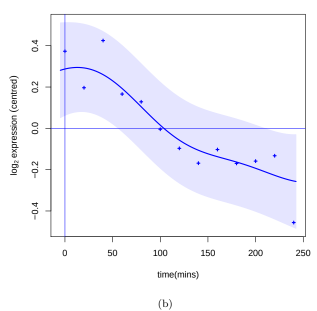


Figure 4: Profile #370: (a) Log-marginal likelihood (LML) contour. (b) GP fit with maximum LML hyperparameters from the exhaustive search.

## 3.4 Comparing against BATS

Now we demonstrate compareROC, a facility for comparing the performance of gprege on a dataset with some other method (see figures below). In this example, we reproduce the main result presented in [Kalaitzis and Lawrence, 2011] <sup>2</sup>. Specifically, we apply standard Gaussian process regression and BATS [Angelini et al., 2007] on experimental gene expression data, where only the top 100 ranks of TSNI were labelled as truly differentially expressed in the noisy ground truth. From the output of each model, a ranking of differential expression is produced and assessed with ROC curves to quantify how well in accordance to the noisy ground truth each method performs. For convenience, gprege was already run on the full DellaGatta dataset and the resulting rank metrics are stored in gpregeOutput\$rankingScores (see demTp63Gp1(fulldataset=TRUE)).

```
# Load fragment dataset.
   data(FragmentDellaGattaData)
   # Download BATS rankings (Angelini, 2007)
   # Case 1: Delta error prior, case 2: Inverse Gamma error prior,
    # case 3: Double Exponential error prior.
   BATSranking = matrix(0, length(DGatta_labels_byTSNItop100), 3)
    for (i in 1:3) {
      # Read gene numbers.
      tmp <- read.table(url(paste(</pre>
        'http://arxiv.org/src/1106.4333v1/anc/DGdat_p63_case',
        i,'_GL.txt',sep='')), skip=1)
     genenumbers <- as.numeric(lapply( as.character(tmp[,2]),</pre>
                                         function(x) x=substr(x,2,nchar(x))))
     # Sort rankings by gene numbers.
     BATSranking[,i] <- tmp[sort(genenumbers, index.return=TRUE)$ix, 4]
    }
+
```

<sup>&</sup>lt;sup>2</sup>Results on experimental data are slightly better because more initialisation points were used in optimising the likelihood wrt the kernel hypeparameters (gpregeOptions\$inithypers), and the optimisation with the best log-marginal likelihood was always used in the end.

#### [1] 0.876575

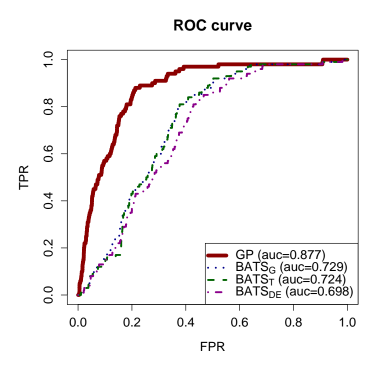


Figure 5: ROC comparison on experimental data from [Della Gatta et al., 2008]. One curve for the GP method and three for BATS, using a different noise model (subscript 'G' for Gaussian, 'T' for Student's-t and 'DE' for double exponential marginal distributions of error), followed by the area under the corresponding curve (AUC).

## 3.5 Ranking the targets

Finally, the bulk ranking for differential expression is done with the full dataset, no interactive mode, and a ROC comparison in the end. Total computation time is approximately 45 minutes on a desktop running Ubuntu 10.04 with a dual-core CPU at 2.8 GHz and 3.2 GiB of memory.

```
# Download full Della Gatta dataset.
con <- url('http://staffwww.dcs.shef.ac.uk/people/A.Kalaitzis/DellaGattaData.RData')</pre>
load(con); close.connection(con)
# Timepoints / GP inputs.
tTrue = matrix(seq(0,240,by=20), ncol=1)
gpregeOptions <- list()</pre>
# Explore individual profiles in interactive mode.
gpregeOptions$explore <- FALSE</pre>
# Exhaustive plot resolution of the LML function.
gpregeOptions$exhaustPlotRes <- 30</pre>
# Exhaustive plot contour levels.
gpregeOptions$exhaustPlotLevels <- 10</pre>
# Exhaustive plot maximum lengthscale.
gpregeOptions$exhaustPlotMaxWidth <- 100</pre>
# Noisy ground truth labels: which genes are in the top 786 ranks of the TSNI ranking.
gpregeOptions$labels <- DGatta_labels_byTSNItop100</pre>
# SCG optimisation: maximum number of iterations.
gpregeOptions$iters <- 100</pre>
# SCG optimisation: no messages.
gpregeOptions$display <- FALSE</pre>
# Matrix of different hyperparameter configurations as rows:
# [inverse-lengthscale
                         percent-signal-variance percent-noise-variance].
gpregeOptions$inithypers <-</pre>
  matrix( c( 1/1000, 1e-3,
                                      0.999,
               1/8,
                           0.999,
                                          1e-3,
               1/80,
                            2/3, 1/3
), ncol=3, byrow=TRUE)
gpregeOutput<-gprege(data=exprs_tp63_RMA,inputs=tTrue,gpregeOptions=gpregeOptions)</pre>
```

#### References

- C. Angelini, D. De Canditiis, M. Mutarelli, and M. Pensky. A Bayesian approach to estimation and testing in time-course microarray experiments. *Stat Appl Genet Mol Biol*, 6:24, 2007.
- G. Della Gatta, M. Bansal, A. Ambesi-Impiombato, D. Antonini, C. Missero, and D. di Bernardo. Direct targets of the TRP63 transcription factor revealed by a combination of gene expression profiling and reverse engineering. *Genome research*, 18(6):939, 2008.
- Alfredo A. Kalaitzis and Neil D. Lawrence. A simple approach to ranking differentially expressed gene expression time courses through gaussian process regression. *BMC Bioinformatics*, 12(180), 2011. doi: 10.1186/1471-2105-12-180.