Description


Details

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Robust calibration of imperfect mathematical models and prediction using experimental data

**Author(s)**

Mengyang Gu [aut, cre]

Maintainer: Mengyang Gu `<mgu6@jhu.edu>`

**References**


**See Also**

RobustGaSP

**Examples**

```r
## A simple example where the math model is not biased
##
## the reality
test_funct_eg1<-function(x){
    sin(pi/2*x)
}

## obtain 25 data from the reality plus a noise
set.seed(1)
```
## 10 data points are very small, one may want to add more data

```r
n=15
input=seq(0,4,4/(n-1))
input=as.matrix(input)

output=test_funct_eg1(input)+rnorm(length(input),mean=0,sd=0.2)

## plot input and output
#plot(input,output)
#num_obs=n=length(output)
```

## the math model
```r
math_model_eg1<-function(x,theta){
  sin(theta*x)
}
```

## fit the S-GaSP model for the discrepancy
```
## one can choose the discrepancy_type to GaSP, S-GaSP or no discrepancy
## p_theta is the number of parameters to calibrate and user needs to specify
## one may also want to change the number of posterior samples by change S and S_0
## one may change sd_proposal for the standard derivation of the proposal distribution
## one may also add a mean by setting X=... and have_trend=TRUE
model_sgasp=rCalibration(design=input, observations=output, p_theta=1, simul_type=1, math_model=math_model_eg1, theta_range=matrix(c(0,3),1,2), S=10000, S_0=2000, discrepancy_type='S-GaSP')
```

## posterior samples of calibration parameter and value
```
## the value is
plot(model_sgasp@post_sample[,1],type='l',xlab='num',ylab=expression(theta))
plot(model_sgasp@post_value,type='l',xlab='num',ylab='posterior value')
```

show(model_sgasp)

# Example: an example used in Susie Bayarri et. al. 2007 Technometrics paper

## reality
```r
test_funct_eg1<-function(x){
  3.5*exp(-1.7*x)+1.5
}
```

## math model
```r
math_model_eg1<-function(x,theta){
  5*exp(-x*theta)
}
```
## Noise Observations (sampled from reality + independent Gaussian noises)
Each has 3 replicates

```r
c(input=c(rep(.110,3),rep(.432,3),rep(.754,3),rep(1.077,3),rep(1.399,3),rep(1.721,3),
rep(2.043,3),rep(2.366,3),rep(2.688,3),rep(3.010,3)))
c(output=c(4.730,4.720,4.234,3.177,2.966,3.653,1.970,2.267,2.084,2.079,2.409,2.371,1.908,1.665,1.685,
1.773,1.608,1.922,1.370,1.661,1.757,1.868,1.505,1.638,1.390,1.275,1.679,1.461,1.157,1.530))
```

n_stack=length(output)/3

```r
output_stack=rep(0,n_stack)
input_stack=rep(0,n_stack)
for(j in 1:n_stack){
  output_stack[j]=mean(output[(j-1)*3+1:(3*j)])
  input_stack[j]=mean(input[(j-1)*3+1:(3*j)])
}
```

Output_stack=as.matrix(output_stack)
Input_stack=as.matrix(input_stack)

## Plot the Output and Stack

```r
plot(input_stack,output_stack,pch=16,col='red')
lines(input_stack,output_stack,pch=16,col='blue',type='p')
```

## Fit the Model with S-GaSP for the Discrepancy

```r
model_sgasp=r_calibration(design=input_stack, observations=output_stack, p_theta=1,simul_type=1,
  math_model=math_model_eg1,theta_range=matrix(c(0,10),1,2),S=10000,
  S_0=2000,discrepancy_type='S-GaSP')
```

# Posterior

```r
plot(model_sgasp@post_sample[,1],type='l',xlab='num',ylab=expression(theta))
show(model_sgasp)
```

### predict

**Prediction for the Robust Calibration Model**

#### Description

Function to make prediction on Robust Calibration models after the r_calibration class has been constructed.

#### Usage

```r
## S4 method for signature 'r_calibration'
predict(object, testing_input,X_testing=matrix(0,dim(testing_input)[1],1),
  n_thinning=10,
```

---

**predict**

**Prediction for the robust calibration model**

**Description**

Function to make prediction on Robust Calibration models after the r_calibration class has been constructed.

**Usage**

```r
## S4 method for signature 'r_calibration'
predict(object, testing_input,X_testing=matrix(0,dim(testing_input)[1],1),
  n_thinning=10,
```
 predict

 testing_output_weights=rep(1,dim(testing_input)[1]),
 interval_est=NULL,interval_data=F,
 math_model=NULL,...)

Arguments

 object an object of class `rcalibration`.
 testing_input a matrix containing the inputs where the `predict` is to perform prediction.
 x_testing a matrix of mean/trend for prediction.
 n_thinning number of points thinning the MCMC posterior samples.
 testing_output_weights
 the weight of testing outputs.
 interval_est a vector for the the posterior credible interval. If `interval_est` is NULL, we do not compute the posterior credible interval. It can be specified as a vector of values ranging from zero to one. E.g. if `interval_est=c(0.025, 0.975)`, the 95 posterior credible interval will be computed.
 interval_data a bool value to decide whether the experimental noise is included for computing the posterior credible interval.
 math_model a function for the math model to be calibrated.
 ... extra arguments to be passed to the function (not implemented yet).

Value

The returned value is a S4 Class `predictobj.rcalibration`.

Author(s)

Mengyang Gu [aut, cre]
Maintainer: Mengyang Gu <mgu6@jhu.edu>

References


Examples

#----------------------------------------------------------------------
# Example: an example used in Susie Bayarri et. al. 2007 Technometrics paper
#----------------------------------------------------------------------

## reality
test_funct_eg1<-function(x){
  3.5*exp(-1.7*x)+1.5
}

## math model
math_model_eg1<-function(x,theta){
  5*exp(-x*theta)
}

## noise observations (sampled from reality + independent Gaussian noises)
## each has 3 replicates
input=c(rep(.110,3),rep(.432,3),rep(.754,3),rep(1.077,3),rep(1.399,3),rep(1.721,3),
        rep(2.043,3),rep(2.366,3),rep(2.688,3),rep(3.010,3))
output=c(4.730,4.720,4.234,3.177,2.966,3.653,1.978,2.267,2.084,2.079,2.409,2.371,1.908,1.665,1.685,
        1.773,1.603,1.922,1.370,1.661,1.757,1.868,1.505,1.638,1.390,1.275,1.679,1.461,1.157,1.530)

## calculating the average or the stack data
n_stack=length(output)/3
output_stack=rep(0,n_stack)
input_stack=rep(0,n_stack)
for(j in 1:n_stack){
  output_stack[j]=mean(output[ ((j-1)*3+1):(3*j) ])
  input_stack[j]=mean(input[ ((j-1)*3+1):(3*j) ])
}
output_stack=as.matrix(output_stack)
input_stack=as.matrix(input_stack)
## plot the output and stack output
#plot(input,output,pch=16,col='red')
#lines(input_stack,output_stack,pch=16,col='blue',type='p')

## fit model using S-GaSP for the discrepancy
## one can change S and S_0 for the number of posterior and burn-in samples
## Normally you may need a larger number of posterior sample
## you can set S=50000 and S_0=5000
## one may also change the sd of the proposal distribution using sd_proposal
model_sgasp=rCalibration(design=input_stack,observations=output_stack, p_theta=1,simul_type=1,
                         math_model=math_model_eg1,theta_range=matrix(c(0,10),1,2),
                         S=10000,S_0=2000,discrepancy_type='S-GaSP')

# one can fit the GaSP model for discrepancy function by discrepancy_type='GaSP'
predict

# one can fit a model without the discrepancy function by discrepancy_type='no-discrepancy'

## posterior of the calibration parameter
plot(model_sgasp$post_sample[,1],type='l',xlab='num',ylab=expression(theta))
show(model_sgasp)

## test data set
testing_input=as.matrix(seq(0,6,0.02))

## perform prediction
prediction_sgasp=predict(model_sgasp,testing_input,math_model=math_model_eg1,
 interval_est=c(0.025,0.975),interval_data=TRUE,
 n_thinning =20 )

## real test output
testing_output=test_funct_eg1(testing_input)

## the prediction by S-GaSP
min_val=min(prediction_sgasp$mean,prediction_sgasp$interval,output,testing_output)
max_val=max(prediction_sgasp$mean,prediction_sgasp$interval,output,testing_output)

plot(testing_input,prediction_sgasp$mean,type='l',col='blue',xlab='x',ylab='y',
 ylim=c(min_val,max_val) )
lines(testing_input,prediction_sgasp$interval[,1],col='blue',lty=2)
lines(testing_input,prediction_sgasp$interval[,2],col='blue',lty=2)

lines(input,output,type='p')
lines(testing_input,prediction_sgasp$math_model_mean,col='blue',lty=3)

lines(testing_input,testing_output,type='l')
legend("topright", legend=c("reality", "predictive mean","95 percent posterior credible interval",
 "predictive mean of the math model"),
 col=c("black", "blue","blue","blue"), lty=c(1,1,2,3),cex=.6)

## MSE if the math model and discrepancy are used for prediction
mean((testing_output-prediction_sgasp$mean)^2)

## MSE if the math model is used for prediction
mean((testing_output-prediction_sgasp$math_model_mean)^2)

# the example with a mean structure

## now let's fit model with mean
model_sgasp_with_mean=rCalibration(design=input_stack,observations=output_stack,
p_theta=1,X=matrix(1,dim(input_stack)[1],1),
predict

```r
have_trend=TRUE,simul_type=1,
math_model=math_model_eg1,
theta_range=matrix(c(0,10),1,2),
S=10000,S_0=2000,
discrepancy_type='S-GaSP')

#posterior
#plot(model_sgasp_with_mean@post_sample[,1],type='l',xlab='num',ylab=expression(theta))
show(model_sgasp_with_mean)

## test data set
testing_input=as.matrix(seq(0,6,0.02))

## in this way we don't output the interval estimation
## one can have interval using interval_est=c(0.025,0.975),interval_data=TRUE
prediction_sgasp_with_mean=predict(model_sgasp_with_mean,testing_input,
X_testing=matrix(1,dim(testing_input)[1],1),
math_model=math_model_eg1,n_thinning = 20)

## plot for the S-GaSP
## for this example, with a mean structure, it fits much better
min_val=min(prediction_sgasp_with_mean@mean,output,testing_output)
max_val=max(prediction_sgasp_with_mean@mean,output,testing_output)
plot(testing_input,prediction_sgasp_with_mean@mean,type='l',col='blue',xlab='x',
     ylab='y',ylim=c(min_val,max_val) )
#lines(testing_input,prediction_sgasp_with_mean@interval[,1],col='blue',lty=2)
#lines(testing_input,prediction_sgasp_with_mean@interval[,2],col='blue',lty=2)
lines(input,output,type='p')
lines(testing_input,prediction_sgasp_with_mean@math_model_mean,col='blue',lty=3)
lines(testing_input,testing_output,type='l')
legend("topright", legend=c("reality", "predictive mean", "predictive mean of the math model"),
col=c("black", "blue","blue"), lty=c(1,1,3),cex=.6)

## MSE if the math model and discrepancy are used for prediction
mean((testing_output-prediction_sgasp_with_mean@mean)^2)

## MSE if the math model is used for prediction
mean((testing_output-prediction_sgasp_with_mean@math_model_mean)^2)

## Not run:
#--------------------------------------------------------------------------------------------------
#the example with the emulator
#--------------------------------------------------------------------------------------------------
n_design=80
```
design_simul<-matrix(runif(n_design*2),n_design,2)
#library(lhs)
#design_simul=maximinLHS(n=n_design,k=2)

design_simul[,1]=6*design_simul[,1]  ##the first one is the observed input x
design_simul[,2]=10*design_simul[,2]  ##the second one is the calibration parameter \theta
output_simul=math_model_eg1(design_simul[,1],design_simul[,2])

##this is a little slow compared with the previous model

model_sgasp_with_mean_emulator=r_calibration(design=input_stack, observations=output_stack,
      p_theta=1,simul_type=0,
      have_trend=T,X=matrix(1,dim(input_stack)[1],1),
      input_simul=design_simul, output_simul=output_simul,
      theta_range=matrix(c(0,10),1,2),
      S=10000,S_0=2000,discrepancy_type='S-GaSP')

##now the output is a list
show(model_sgasp_with_mean_emulator)

##here is the plot
plot(model_sgasp_with_mean_emulator@post_sample[,4],type='l',xlab='num',ylab=expression(theta))

##For this example, with a mean structure, it fits much better
min_val=min(prediction_sgasp_with_mean_emulator@mean/output,testing_output,
      prediction_sgasp_with_mean_emulator@math_model_mean)
max_val=max(prediction_sgasp_with_mean_emulator@mean/output,testing_output,
      prediction_sgasp_with_mean_emulator@math_model_mean)

plot((testing_input,prediction_sgasp_with_mean_emulator@mean,type='l',col='blue',xlab='x',
      ylab='y',ylim=c(min_val,max_val))
#lines(testing_input,prediction_sgasp_with_mean@interval[,1],col='blue',lty=2)
#lines(testing_input,prediction_sgasp_with_mean@interval[,2],col='blue',lty=2)

lines(input,output,type='p')
lines(testing_input,prediction_sgasp_with_mean_emulator@math_model_mean,col='blue',lty=3)

lines(testing_input,testing_output,type='l')

legend("topright", legend=c("reality", "predictive mean", "predictive mean of the math model"),
       col=c("black", "blue","blue"), lty=c(1,1,3),cex=.6)
predictobj.rcalibration-class

### Predictive results for the Robust Calibration class

#### Description

S4 class for prediction after Robust rcalibration with or without the specification of the discrepancy model.

#### Objects from the Class

Objects of this class are created and initialized with the function `predict` that computes the prediction and the uncertainty quantification.

#### Slots

- `mean`: object of class `vector`. The predictive mean at testing inputs combing the mathematical model and discrepancy function.
- `math_model_mean`: object of class `vector`. The predictive mean at testing inputs using only the mathematical model (and the trend if specified).
- `math_model_mean_no_trend`: object of class `vector`. The predictive mean at testing inputs using only the mathematical model without the trend.
- `interval`: object of class `matrix`. The upper and lower predictive credible interval. If `interval_data` is TRUE in the `predict.rcalibration`, the experimental noise is included for computing the predictive credible interval.

#### Author(s)

Mengyang Gu [aut, cre]
Maintainer: Mengyang Gu <mgu6@jhu.edu>
predictobj.rcalibration_MS-class

References


See Also

predict.rcalibration for more details about how to do prediction for a rcalibration object.

predictobj.rcalibration_MS-class

Predictive results for the Robust Calibration class

Description

S4 class for prediction after Robust rcalibration for multiple sources.

Objects from the Class

Objects of this class are created and initialized with the function predict_MS that computes the prediction and the uncertainty quantification.

Slots

mean: object of class list. Each element is a vector of the predictive mean at testing inputs combing the mathematical model and discrepancy function for each source.

math_model_mean: object of class list. Each element is a vector of the predictive mean at testing inputs using only the mathematical model (and the trend if specified).

math_model_mean_no_trend: object of class list. Each element is a vector of the predictive mean at testing inputs using only the mathematical model without the trend for each source.

interval: object of class list. Each element is a matrix of the upper and lower predictive credible interval. If interval_data is TRUE in the predict_MS, the experimental noise is included for computing the predictive credible interval.

Author(s)

Mengyang Gu [aut, cre]

Maintainer: Mengyang Gu <mgu6@jhu.edu>
**References**


**See Also**

predict_MS for more details about how to do prediction for a rcalibration_MS object.

---

**predict_MS**

**Prediction for the robust calibration model for multiple sources**

**Description**

Function to make prediction on Robust Calibration models after the rcalibration class has been constructed for multiple sources.

**Usage**

```r
## S4 method for signature 'rcalibration_MS'
predict_MS(object, testing_input,
            X_testing=as.list(rep(0,object@num_sources)),
            n_thinning=10, testing_output_weights=NULL,
            interval_est=NULL,
            interval_data=rep(F,length(testing_input)),
            math_model=NULL,...)
```

**Arguments**

- **object**
  an object of class rcalibration_MS.

- **testing_input**
  a list of matrices containing the inputs where the predict_MS is to perform prediction.

- **X_testing**
  a list of matrices of mean/trend for prediction if specified.

- **n_thinning**
  number of points thinning the MCMC posterior samples.

- **testing_output_weights**
  a list of vectors for the weight of testing outputs for multiple sources.

- **interval_est**
  a list of vectors for the posterior predictive credible interval for multiple sources.
  If interval_est is NULL, we do not compute the posterior credible interval. It can be specified as a vector of values ranging from zero to one. E.g.

- **interval_data**
  a vector of bool values to decide whether the experimental noise is included for computing the posterior credible interval.

- **math_model**
  a list of functions for the math model to be calibrated for multiple sources.

- **...**
  extra arguments to be passed to the function (not implemented yet).
Value

The returned value is a S4 CLass predictobj.rcalibration.

Author(s)

Mengyang Gu [aut, cre]
Maintainer: Mengyang Gu <mgu6@jhu.edu>

References


Examples

# An example for calibrating and predicting mathematical models for data from multiple sources

library(RobustCalibration)

## reality

test_funct<function(x){
  sin(pi*x/2)+2*cos(pi*x/2)
}

## math model from two sources

math_model_source_1<function(x,theta){
  sin(theta*x)
}
predict_MS

math_model_source_2 <- function(x, theta){
  cos(theta*x)
}

input1 = seq(0, 2/(10-1))
input2 = seq(0, 3/(10-1))
##
output1 = test_funct(input1) + rnorm(length(input1), sd = 0.01)
output2 = test_funct(input2) + rnorm(length(input2), sd = 0.02)

plot(input1, output1)
plot(input2, output2)

design = list()
design[[1]] = as.matrix(input1)
design[[2]] = as.matrix(input2)

observations = list()
observations[[1]] = output1
observations[[2]] = output2

p_theta = 1
theta_range = matrix(0, p_theta, 2)
theta_range[1, ] = c(0, 8)
simul_type = c(1, 1)

math_model = list()
math_model[[1]] = math_model_source_1
math_model[[2]] = math_model_source_2

## calibrating two mathematical models for these two sources
model_sgasp = rcalibration_MS(df_design = design, observations = observations, p_theta = 1,
simul_type = simul_type, math_model = math_model,
theta_range = theta_range,
S = 10000, S_0 = 2000,
discrepancy_type = rep('S-GaSP', length(design)))

# plot(model_sgasp@post_theta[, 1], type = 'l')
mean(model_sgasp@post_theta[, 1])

testing_input1 = seq(0, 2/(25-1))
testing_input2 = seq(0, 3/(25-1))
testing_input = list()
rcalibration

Setting up the robust Calibration model

Description

Setting up the Calibration model for estimating the parameters via MCMC with or without a discrepancy function.

Usage

rcalibration(design, observations, p_theta=NULL, X=matrix(0, dim(design)[1], 1), have_trend=FALSE, simul_type=0, input_simul=NULL, output_simul=NULL, simul_nug=FALSE, math_model=NULL, theta_range=NULL, sd_proposal=c(rep(0.05, p_theta), rep(0.25, dim(design)[2]), 0.25), s=10000, s_0=1000, discrepancy_type='S-GaSP', kernel_type='matern_5_2', tilde_lambda=1/2, a=1/2-dim(design)[2], b=1, alpha=rep(1.9, dim(design)[2]), output_weights=rep(1, dim(design)[1]))

Arguments

design  a matrix of observed inputs.
observations  a vector of experimental data.
p_theta  an integer about the number of parameters, which should be specified by the user.
X  a matrix of the mean/trend discrepancy between the reality and math model.
have_trend  a bool value meaning whether we assume a mean/trend discrepancy function.
**simul_type**

an integer about the math model/simulator. If the simul_type is 0, it means we use RobustGaSP R package to build an emulator for emulation. If the simul_type is 1, it means the function of the math model is given by the user. When simul_type is 2 or 3, the mathematical model is the geophysical model for Kilauea Volcano. If the simul_type is 2, it means it is for the ascending mode InSAR data; if the simul_type is 3, it means it is for the descending mode InSAR data.

**input_simul**

an D x (p_x+p_theta) matrix of design for emulating the math model. It is only useful if simul_type is 0, meaning that we emulate the output of the math model.

**output_simul**

a D dimensional vector of the math model runs on the design (input_simul). It is only useful if simul_type is 0, meaning that we emulate the output of the math model.

**simul_nug**

a bool value meaning whether we have a nugget for emulating the math model/simulator. If the math model is stochastic, we often need a nugget. If simul_Nug is TRUE, it means we have a nugget for the emulator. If simul_Nug is FALSE, it means we do not have a nugget for the emulator.

**math_model**

a function of the math model provided by the user. It is only useful if simul_type is 1, meaning that we know the math model and it can be computed fast. If the evaluation the math model is computationally slow, one should set simul_type to be 0 to emulate the math model.

**theta_range**

a p_theta x 2 matrix of the range of the calibration parameters. The first column is the lower bound and the second column is the upper bound. It should be specified by the user if the simul_type is 0.

**sd_proposal**

a vector of the standard deviation of the proposal distribution in MCMC.

**S**

an integer about about how many posterior samples to run.

**S_0**

an integer about about the number of burn-in samples.

**discrepancy_type**

characters about the type of the discrepancy. If it is 'no-discrepancy', it means no discrepancy function. If it is 'GaSP', it means the GaSP model for the discrepancy function. If it is 'S-GaSP', it means the S-GaSP model for the discrepancy function.

**kernel_type**

characters about the type of the discrepancy.type of kernel. matern_3_2 and matern_5_2 are Matern kernel with roughness parameter 3/2 and 5/2 respectively. pow_exp is power exponential kernel with roughness parameter alpha. If pow_exp is to be used, one needs to specify its roughness parameter alpha.

**tilde_lambda**

a numeric value about how close the math model to the reality in squared distance when the S-GaSP model is used for modeling the discrepancy.

**a**

a scalar of the prior parameter.

**b**

a scalar of the prior parameter.

**alpha**

a numeric parameter for the roughness in the kernel.

**output_weights**

a vector of the weights of the outputs.

**Value**

r_calibration returns an S4 object of class r_calibration (see r_calibration-class).
Author(s)
Mengyang Gu [aut, cre]
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References

Examples
library(RobustCalibration)

# an example with multiple local maximum of minimum in L2 loss

## the reality
test_funct_eg1=function(x){
  x*cos(3/2*x)+x
}

## obtain 25 data from the reality plus a noise
set.seed(1)
## 10 data points are very small, one may want to add more data
n=15
input=seq(0.5,5/(n-1))
input=as.matrix(input)
output=test_funct_eg1(input)+rnorm(length(input),mean=0,sd=0.1)
num_obs=n=length(output)

## the math model
### r calibration

\[
\text{math\_model\_eg1} <- \text{function}(x, \theta) \{
  \sin(\theta x) + x
\}
\]

---

```r
# fit the S-GaSP model for the discrepancy
# one can choose the discrepancy_type to GaSP, S-GaSP or no discrepancy
# p_theta is the number of parameters to calibrate and user needs to specify
# one may also want to change the number of posterior samples by change S and S_0
p_theta = 1
model_sgasp = rcalibration(design = input, observations = output, p_theta = p_theta, simul_type = 1,
                         math_model = math_model_eg1, theta_range = matrix(c(0, 3), 1, 2),
                         S = 10000, S_0 = 2000, discrepancy_type = 'S-GaSP')

# if the acceptance rate is too low or too high, one can adjust sd_proposal, e.g.
model_sgasp = rcalibration(design = input, observations = output, p_theta = 1, simul_type = 1,
                         sd_proposal = c(rep(0.02, p_theta), rep(0.2, dim(input)[2])), 0.2)
                         math_model = math_model_eg1, theta_range = matrix(c(0, 3), 1, 2),
                         S = 10000, S_0 = 2000, discrepancy_type = 'S-GaSP')

# posterior samples of calibration parameter and value
plot(model_sgasp@post_sample[, 1], type = 'l', xlab = 'num', ylab = expression(theta))
plot(model_sgasp@post_value, type = 'l', xlab = 'num', ylab = 'posterior value')

show(model_sgasp)

# one may want to fit a a model with an estimated baseline mean discrepancy by setting
# X = matrix(1, dim(input_stack)[1], 1), have_trend = TRUE

model_sgasp_with_mean = rcalibration(design = input, observations = output, p_theta = 1, simul_type = 1,
                      X = matrix(1, dim(input)[1], 1), have_trend = TRUE,
                      math_model = math_model_eg1, theta_range = matrix(c(0, 3), 1, 2),
                      S = 10000, S_0 = 2000, discrepancy_type = 'S-GaSP')

show(model_sgasp_with_mean)

# posterior samples of calibration parameter and value
plot(model_sgasp_with_mean@post_sample[, 1], type = 'l', xlab = 'num', ylab = expression(theta))
plot(model_sgasp_with_mean@post_value, type = 'l', xlab = 'num', ylab = 'posterior value')

## Not run:

# an example with multiple local maximum of minimum in L2 loss
# for combing the emulator

## the reality

\[
\text{test\_funct\_eg1} <- \text{function}(x) \{
  x * \cos(3/2*x) + x
\}
\]
### Obtain 20 Data from the Reality Plus a Noise

```r
set.seed(1)

n=20
input=seq(0,5,5/(n-1))
input=as.matrix(input)

output=test_funct_eg1(input)+rnorm(length(input),mean=0, sd=0.05)

num_obs=n=length(output)
```

### The Math Model

```r
math_model_eg1<-function(x,theta){
  sin(theta*x)+x
}
```

### Let's Build an Emulator for the Case if the Math Model is Too Slow

# Let's Say We Can Only Run the Math Model n_design Times

```r
n_design=80

design_simul=matrix(runif(n_design*2),n_design,2)
design_simul[,1]=5*design_simul[,1]  # The first one is the observed input x
design_simul[,2]=3*design_simul[,2]  # The second one is the calibration parameter

output_simul=math_model_eg1(design_simul[,1],design_simul[,2])
```

### This is a Little Slow Compared with the Previous Model

```r
model_sgasp_emulator=rcalibration(design=input, observations=output, p_theta=1, simul_type=0,
input_simul=design_simul, output_simul=output_simul,
theta_range=matrix(c(0,3),1,2),
S=10000,S_0=2000,discrepancy_type='S-GaSP')
```

### Now the Output Is a List

```r
show(model_sgasp_emulator)
```

### Here Is the Plot

```r
plot(model_sgasp_emulator@post_sample[,1],type='l',xlab='num',ylab=expression(theta))
plot(model_sgasp_emulator@post_value,type='l',xlab='num',ylab='posterior value')
```

### End (Not run)
**Description**

S4 class for Robust rcalibration with or without the specification of the discrepancy model.

**Objects from the Class**

Objects of this class are created and initialized with the function `rcalibration` that computes the calculations needed for setting up the calibration and prediction.

**Slots**

- `p_x`: Object of class `integer`. The dimension of the observed inputs.
- `p_theta`: Object of class `integer`. The calibration parameters.
- `num_obs`: Object of class `integer`. The number of experimental observations.
- `input`: Object of class `matrix` with dimension `n x p_x`. The design of experiments.
- `output`: Object of class `vector` with dimension `n x 1`. The vector of the experimental observations.
- `x`: Object of class `matrix` of with dimension `n x q`. The mean/trend discrepancy basis function.
- `have_trend`: Object of class `bool` to specify whether the mean/trend discrepancy is zero. "TRUE" means it has zero mean discrepancy and "FALSE" means the mean discrepancy is not zero.
- `q`: Object of class `integer`. The number of basis functions of the mean/trend discrepancy.
- `R0`: Object of class `list` of matrices where the j-th matrix is an absolute difference matrix of the j-th input vector.
- `kernel_type`: A character to specify the type of kernel to use.
- `alpha`: Object of class `vector`. Each element is the parameter for the roughness for each input coordinate in the kernel.
- `theta_range`: A matrix for the range of the calibration parameters.
- `tilde_lambda`: Object of class `numeric` about how close the math model to the reality in squared distance when the S-GaSP model is used for modeling the discrepancy.
- `S`: Object of class `integer` about how many posterior samples to run.
- `S_0`: Object of class `integer` about the number of burn-in samples.
- `prior_par`: Object of class `vector` about prior parameters.
- `output_weights`: Object of class `vector` about the weights of the experimental data.
- `sd_proposal`: Object of class `vector` about the standard deviation of the proposal distribution.
- `discrepancy_type`: Object of class `character` about the discrepancy. If it is 'no-discrepancy', it means no discrepancy function. If it is 'GaSP', it means the GaSP model for the discrepancy function. If it is 'S-GaSP', it means the S-GaSP model for the discrepancy function.
- `simul_type`: Object of class `integer` about the math model/simulator. If the simul_type is 0, it means we use RobustGaSP R package to build an emulator for emulation. If the simul_type is 1, it means the function of the math model is given by the user. When simul_type is 2 or 3, the mathematical model is the geophysical model for Kilauea Volcano. If the simul_type is 2, it means it is for the ascending mode InSAR data; if the simul_type is 3, it means it is for the descending mode InSAR data.
- `emulator`: An S4 class of rgasp from the RobustGaSP package.
post_sample: Object of class matrix for the posterior samples after burn-in.

post_value: Object of class vector for the posterior values after burn-in.

accept_s: Object of class vector for the number of proposed samples of the calibration parameters are accepted in MCMC. The first value is the number of proposed calibration parameters are accepted in MCMC. The second value is the number of proposed range and nugget parameters are accepted.

count_boundary: Object of class vector for the number of proposed samples of the calibration parameters are outside the range and they are rejected directly.

Methods

show Print the main slots of the object.

predict See predict.

predict_discrepancy_separable_2dim See predict_discrepancy_separable_Rdim.

Author(s)

Mengyang Gu [aut, cre]

Maintainer: Mengyang Gu <mgu6@jhu.edu>

References


See Also

rcalibration for more details about how to create a rcalibration object.
Usage

rcalibration_MS(list_of_design, list_of_observations, p_theta=NULL, index_theta=NULL, X=as.list(rep(0,length(list_of_design))), have_trend=rep(FALSE,length(list_of_design))), simul_type=rep(0, length(list_of_design)), input_simul=rep(NULL, length(list_of_design)), simul_nug=rep(FALSE,length(list_of_design)), math_model=NULL, theta_range=NULL, sd_proposal_theta=rep(0.05,p_theta), sd_proposal_cov_par=NULL, tilde.lambda=rep(1/2,length(list_of_design)), a=rep(NULL, length(list_of_design)), b=rep(NULL, length(list_of_design)), output_weights=rep(NULL, length(list_of_design)))

Arguments

design a list of observed inputs from multiple sources.
observations a list of experimental data from multiple sources.
index_theta a list of vectors for the index of calibration parameter contained in each source.
p_theta an integer about the number of parameters, which should be specified by the user.
X a list of matrices of the mean/trend discrepancy between the reality and math model for multiple sources.
have_trend a vector of bool value meaning whether we assume a mean/trend discrepancy function.
simul_type a vector of integer about the math model/simulator for multiple sources. If the simul_type is 0, it means we use RobustGaSP R package to build an emulator for emulation. If the simul_type is 1, it means the function of the math model is given by the user. When simul_type is 2 or 3, the mathematical model is the geophysical model for Kilauea Volcano. If the simul_type is 2, it means it is for the ascending mode InSAR data; if the simul_type is 3, it means it is for the descending mode InSAR data.
input_simul a list of matrices, each having dimension D x (p_x+p_theta) being the design for emulating the math model. It is only useful if the ith value of simul_type is 0 for the ith source, meaning that we emulate the output of the math model.
output_simul a list of vectors, each having dimension D x 1 being the math model outputs on the design (input_simul). It is only useful if the ith value of simul_type is 0 for the ith source, meaning that we emulate the output of the math model.
simul_nug a vectors of bool values meaning whether we have a nugget for emulating the math model/simulator for this source. If the math model is stochastic, we often need a nugget. If simul_nug is TRUE, it means we have a nugget for the
emulator. If simul_Nug is FALSE, it means we do not have a nugget for the emulator.

math_model a list of functions of the math models provided by the user for multiple sources. It is only useful if simul_type is 1, meaning that we know the math model and it can be computed fast. If the evaluation the math model is computationally slow, one should set simul_type to be 0 to emulate the math model.

theta_range a p_theta x 2 matrix of the range of the calibration parameters. The first column is the lower bound and the second column is the upper bound. It should be specified by the user if the simul_type is 0.

sd_proposal_theta a vector of the standard deviation of the proposal distribution for the calibration parameters in MCMC.

sd_proposal_cov_par a list of vectors of the standard deviation of the proposal distribution for range and nugget parameters in MCMC for each source.

S an integer about about how many posterior samples to run.

S_0 an integer about about the number of burn-in samples.

discrepancy_type a vector of characters about the type of the discrepancy for each source. If it is 'no-discrepancy', it means no discrepancy function. If it is 'GaSP', it means the GaSP model for the discrepancy function. If it is 'S-GaSP', it means the S-GaSP model for the discrepancy function.

kernel_type a vector of characters about the type of the discrepancy.type of kernel for each source. matern_3_2 and matern_5_2 are Matern kernel with roughness parameter 3/2 and 5/2 respectively. pow_exp is power exponential kernel with roughness parameter alpha. If pow_exp is to be used, one needs to specify its roughness parameter alpha.

tilde_lambda a vector numeric values about how close the math model to the reality in squared distance when the S-GaSP model is used for modeling the discrepancy for each source.

a a vector of the prior parameter for multiple sources.

b a vector of the prior parameter for multiple sources.

alpha a list of vectors of roughness parameters in the kernel for multiple sources.

output_weights a list of vectors of the weights of the outputs for multiple sources.

Value
rcalibration_MS returns an S4 object of class rcalibration_MS (see rcalibration_MS-class).

Author(s)
Mengyang Gu [aut, cre]
Maintainer: Mengyang Gu <mgu6@jhu.edu>
References


Examples

```r
# An example for calibrating mathematical models for data from multiple sources
libary(RobustCalibration)

test_funct<-function(x){
  sin(pi*x/2)+2*cos(pi*x/2)
}

math_model_source_1<-function(x,theta){
  sin(theta*x)
}

math_model_source_2<-function(x,theta){
  cos(theta*x)
}

input1=seq(0.2,2/(10-1))
input2=seq(0.3,3/(15-1))

output1=test_funct(input1)+rnorm(length(input1), sd=0.01)
output2=test_funct(input2)+rnorm(length(input2), sd=0.02)

plot(input1, output1)
plot(input2, output2)
```
r_calibration MS-class  Robust Calibration for multiple sources class

Description

S4 class for multiple sources Robust rcalibration with or without the specification of the discrepancy model.

Objects from the Class

Objects of this class are created and initialized with the function `r_calibration MS` that computes the prediction after calibrating the mathematical models from multiple sources.
**Slots**

- `num_sources`: Object of class integer. The number of sources.
- `p_x`: Object of class vector. Each element is the dimension of the observed inputs in each source.
- `p_theta`: Object of class integer. The number of calibration parameters.
- `num_obs`: Object of class vector. Each element is the number of experimental observations of each source.
- `index_theta`: Object of class list. The each element is a vector of the index of calibration parameters (theta) contained in each source.
- `input`: Object of class list. Each element is a matrix of the design of experiments in each source with dimension \( n_i \times p_{x,i} \) for \( i = 1, \ldots, \text{num\_sources} \).
- `output`: Object of class list. Each element is a vector of the experimental observations in each source with dimension \( n_i \times 1 \), for \( i = 1, \ldots, \text{num\_sources} \).
- `X`: Object of class list. Each element is a matrix of the mean/trend discrepancy basis function in each source with dimension \( n_i \times q_i \), for \( i = 1, \ldots, \text{num\_sources} \).
- `have_trend`: Object of class vector. Each element is a bool to specify whether the mean/trend discrepancy is zero in each source. "TRUE" means it has zero mean discrepancy and "FALSE" means the mean discrepancy is not zero.
- `q`: Object of class vector. Each element is integer of the number of basis functions of the mean/trend discrepancy in each source.
- `R0`: Object of class list. Each element is a list of matrices where the j-th matrix is an absolute difference matrix of the j-th input vector in each source.
- `kernel_type`: Object of class list. Each element is an object of character to specify the type of kernel to use in each source.
- `alpha`: Object of class vector. Each element is an object of vector about prior parameters.
- `theta_range`: A matrix for the range of the calibration parameters.
- `tilde_lambda`: Object of class vector. Each element is a numeric value about how close the math model to the reality in squared distance when the S-GaSP model is used for modeling the discrepancy in each source.
- `S`: Object of class integer about how many posterior samples to run.
- `S_0`: Object of class integer about the number of burn-in samples.
- `prior_par`: Object of class list. Each element is a vector about prior parameters.
- `output_weights`: Object of class list. Each element is a vector about the weights of the experimental data.
- `sd_proposal_theta`: Object of class vector about the standard deviation of the proposal distribution for the calibration parameters.
- `sd_proposal_cov_par`: Object of class list. Each element is a vector about the standard deviation of the proposal distribution for the calibration parameters in each source.
- `discrepancy_type`: Object of class vector. Each element is a character about the type of the discrepancy in each source. If it is 'no-discrepancy', it means no discrepancy function. If it is 'GaSP', it means the GaSP model for the discrepancy function. If it is 'S-GaSP', it means the S-GaSP model for the discrepancy function.
simul_type: Object of class vector. Each element is an integer about the math model/simulator. If the simul_type is 0, it means we use RobustGaSP R package to build an emulator for emulation. If the simul_type is 1, it means the function of the math model is given by the user. When simul_type is 2 or 3, the mathematical model is the geophysical model for Kilauea Volcano. If the simul_type is 2, it means it is for the ascending mode InSAR data; if the simul_type is 3, it means it is for the descending mode InSAR data.

emulator: Object of class list. Each element is an S4 class of rgasp from the RobustGaSP package in each source.

post_theta: Object of class matrix for the posterior samples of the calibration parameters after burn-in.

post_individual_par: Object of class list. Each element is a matrix for the posterior samples after burn-in in each source.

post_value: Object of class vector for the posterior values after burn-in.

accept_S_theta: Object of class numerical for the number of proposed samples of the calibration parameters are accepted in MCMC.

accept_S_beta: Object of class vector for the number of proposed samples of the range and nugget parameters in each source are accepted in MCMC.

count_boundary: Object of class vector for the number of proposed samples of the calibration parameters are outside the range and they are rejected directly.

Methods

predict_MS See predict_MS.

Author(s)

Mengyang Gu [aut, cre]

Maintainer: Mengyang Gu <mgu6@jhu.edu>

References


See Also

rcalibration_MS for more details about how to create a rcalibration_MS object.
Show an Robust Calibration object.

Description

Function to print the Robust Calibration model after the rcalibration class has been constructed.

Usage

```r
## S4 method for signature 'rcalibration'
show(object)
```

Arguments

- `object`: an object of class `rcalibration`.

Author(s)

Mengyang Gu [aut, cre]
Maintainer: Mengyang Gu <mgu6@jhu.edu>

References


Examples

```r
##-----------------------------------------------
#A simple example where the math model is not biased
##-----------------------------------------------
## the reality
test_funct_eg1<-function(x){
  sin(pi/2*x)
}

## obtain 15 data from the reality plus a noise
set.seed(1)
```
## 10 data points are very small, one may want to add more data

```r
n=15
input=seq(0,4,4/(n-1))
input=as.matrix(input)

output=test_funct_eg1(input)+rnorm(length(input),mean=0,sd=0.2)
```

## plot input and output

```r
#plot(input,output)
#num_obs=n=length(output)
```

## the math model

```r
math_model_eg1=function(x,theta){
  sin(theta*x)
}
```

## fit the S-GaSP model for the discrepancy

```r
# one can choose the discrepancy_type to GaSP, S-GaSP or no discrepancy
# p_theta is the number of parameters to calibrate and user needs to specify
# one may also want to change the number of posterior samples by change S and S_0
# one may change sd_proposal for the standard derivation of the proposal distribution
# one may also add a mean by setting X=... and have_trend=TRUE
model_sgasp=rCalibration(design=input, observations=output, p_theta=1, simul_type=1,
                         math_model=math_model_eg1, theta_range=matrix(c(0,3),1,2),
                         S=10000, S_0=2000, discrepancy_type='S-GaSP')
```

## posterior samples of calibration parameter and value

```r
# the value is
plot(model_sgasp$post_sample[,1],type='l',xlab='num',ylab=expression(theta))
plot(model_sgasp$post_value,type='l',xlab='num',ylab='posterior value')
```

```r
show(model_sgasp)
```

## an example with multiple local maximum of minimum in L2 loss

```r
## the reality
test_funct_eg1<-function(x){
  x*cos(3/2**x)+x
}
```

## obtain 15 data from the reality plus a noise

```r
set.seed(1)
n=15
input=seq(0,5,5/(n-1))
```
show

input = as.matrix(input)

output = test_funct_eg1(input) + rnorm(length(input), mean = 0, sd = 0.05)

num_obs = n = length(output)

## the math model
math_model_eg1 <- function(x, theta){
  sin(theta * x) + x
}

## fit the S-GaSP model for the discrepancy
model_sgasp = rcalibration(design = input, observations = output, p_theta = 1, simul_type = 1, math_model = math_model_eg1, theta_range = matrix(c(0, 1, 2), 1, 2), discrepancy_type = 'S-GaSP')

## posterior samples
plot(model_sgasp$post_sample[, 1], type = 'l', xlab = 'num', ylab = expression(theta))
show(model_sgasp)
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