

# Package ‘support’

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**Type** Package

**Title** Support Points

**Version** 0.1.7

**Description** The functions `sp()` and `sp_seq()` compute the support points in Mak and Joseph (2018) <[DOI:10.1214/17-AOS1629](https://doi.org/10.1214/17-AOS1629)>. Support points can be used as a representative sample of a desired distribution, or a representative reduction of a big dataset (e.g., an “optimal” thinning of Markov-chain Monte Carlo sample chains). This work was supported by USARO grant W911NF-14-1-0024 and NSF DMS grant 1712642.

**License** GPL (>= 2)

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## Contents

support-package . . . . .	2
e_dist . . . . .	2
mcmc_franke . . . . .	4
sp . . . . .	5
sp_seq . . . . .	8

<b>Index</b>	<b>12</b>
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 support-package

*Support Points*


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### Description

The 'support' package provides functions for computing support points.

### Details

Package: support  
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 Date: 2019-07-15  
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The 'support' package provides the functions `sp()` and `sp_seq()` for computing the support points in Mak and Joseph (2018) <DOI:10.1214/17-AOS1629>. Support points can be used as a representative sample of a desired distribution, or a representative reduction of a big dataset (e.g., an "optimal" thinning of Markov-chain Monte Carlo sample chains). This work was supported by USARO grant W911NF-14-1-0024 and NSF DMS grant 1712642.

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### References

Mak, S. and Joseph, V. R. (2018). Support points. *Annals of Statistics*, 46(6A):2562-2592.

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 e\_dist

*Computes the energy distance of a point set*


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### Description

`e_dist` computes the energy distance between points  $D$  and a target distribution (or big dataset)  $F$ . The cross-term  $E[||X - X'||]$ ,  $X, X' \sim F$  is *NOT* computed in `e_dist` for computational efficiency, since this is not needed for optimizing  $D$ . The target distribution or big dataset can be set using `dist.str` or `dist.samp`, respectively.

### Usage

```
e_dist(D, dist.str=NA, dist.param=vector("list", ncol(D)),
       nsamp=1e6, dist.samp=NA)
```

**Arguments**

D	An $n \times p$ point set.
dist.str	A $p$ -length string vector for target distribution (assuming independence). Current options include uniform, normal, exponential, gamma, lognormal, student-t, weibull, cauchy and beta. Exactly one of dist.str or dist.samp should be NA.
dist.param	A $p$ -length list for desired distribution parameters in dist.str. The following parameters are supported: <ul style="list-style-type: none"> <li>• <b>Uniform:</b> Minimum, maximum;</li> <li>• <b>Normal:</b> Mean, standard deviation;</li> <li>• <b>Exponential:</b> Rate parameter;</li> <li>• <b>Gamma:</b> Shape parameter, scale parameter;</li> <li>• <b>Lognormal:</b> Log-mean, Log-standard deviation;</li> <li>• <b>Student-t:</b> Degree-of-freedom;</li> <li>• <b>Weibull:</b> Shape parameter, scale parameter;</li> <li>• <b>Cauchy:</b> Location parameter, scale parameter;</li> <li>• <b>Beta:</b> Shape parameter 1, shape parameter 2.</li> </ul>
nsamp	Number of samples to draw from dist.str for comparison.
dist.samp	An $N \times p$ matrix for the target big dataset (e.g., MCMC chain). Exactly one of dist.str or dist.samp should be NA.

**References**

Szekely, G. J. and Rizzo, M. L. (2013). Energy statistics: A class of statistics based on distances. *Journal of Statistical Planning and Inference*, 143(8):1249-1272.

**Examples**

```
## Not run:
#####
# Generate 25 SPs for the 2-d i.i.d. N(0,1) distribution
#####
n <- 25 #number of points
p <- 2 #dimension
D <- sp(n,p,dist.str=rep("normal",p))
Drnd <- matrix(rnorm(n*p),ncol=p)
e_dist(D$sp,dist.str=rep("normal",p)) #smaller
e_dist(Drnd,dist.str=rep("normal",p))

#####
# Support points for big data reduction: Franke's function
#####
# library(MHadaptive) # Package archived, but you can use your favorite MCMC sampler

# #Generate MCMC samples
# li_func <- franke2d #Desired log-posterior
# ini <- c(0.5,0.5) #Initial point for MCMC
```

```

# NN <- 1e5 #Number of MCMC samples desired
# burnin <- NN/2 #Number of burn-in runs
# mcmc_franke <- Metro_Hastings(li_func, pars=ini, prop_sigma=0.05*diag(2),
#                               iterations=NN, burn_in=burnin)
data(mcmc_franke) # Loading MCMC sample from data

#Use modified Franke's function as posterior
franke2d <- function(xx){
  if ((xx[1]>1)|| (xx[1]<0)|| (xx[2]>1)|| (xx[2]<0)){
    return(-Inf)
  }
  else{
    x1 <- xx[1]
    x2 <- xx[2]

    term1 <- 0.75 * exp(-(9*x1-2)^2/4 - (9*x2-2)^2/4)
    term2 <- 0.75 * exp(-(9*x1+1)^2/49 - (9*x2+1)/10)
    term3 <- 0.5 * exp(-(9*x1-7)^2/4 - (9*x2-3)^2/4)
    term4 <- -0.2 * exp(-(9*x1-4)^2 - (9*x2-7)^2)

    y <- term1 + term2 + term3 + term4
    return(2*log(y))
  }
}

#Generate ncur SPs
ncur <- 50
D <- sp(ncur,2,dist.samp=mcmc_franke$trace)$sp
Drnd <- mcmc_franke$trace[sample(1:nrow(mcmc_franke$trace),n,F),]
e_dist(D,dist.samp=mcmc_franke$trace) #smaller
e_dist(Drnd,dist.samp=mcmc_franke$trace)

## End(Not run)

```

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mcmc\_franke

*MCMC sample from the modified Franke function*


---

### Description

MCMC sample from the modified Franke function using the package MHadaptive (which is currently archived).

### Usage

```
data(mcmc_franke)
```

## Description

sp is the main function for computing the support points in Mak and Joseph (2018). Current options include support points on standard distributions (specified via `dist.str`) or support points for reducing big data (specified via `dist.samp`). For big data reduction, weights on each data point can be specified via `wts`.

## Usage

```
sp(n, p, ini=NA,
   dist.str=NA, dist.param=vector("list",p),
   dist.samp=NA, scale.flg=TRUE, wts=NA, bd=NA,
   num.subsamp=ifelse(any(is.na(dist.samp)),
   max(10000,10*n),min(10000,nrow(dist.samp))),
   rnd.flg=ifelse(any(is.na(dist.samp)),
   TRUE,ifelse(num.subsamp<=10000,FALSE,TRUE)),
   iter.max=max(250,iter.min), iter.min=50,
   tol=1e-10, par.flg=TRUE, n0=n*p)
```

## Arguments

n	Number of support points.
p	Dimension of sample space.
ini	An $n \times p$ matrix for initialization.
dist.str	A $p$ -length string vector for desired distribution (assuming independence). Current options include uniform, normal, exponential, gamma, lognormal, student-t, weibull, cauchy and beta. Exactly one of <code>dist.str</code> or <code>dist.samp</code> should be NA.
dist.param	A $p$ -length list for desired distribution parameters in <code>dist.str</code> . The following parameters are supported: <ul style="list-style-type: none"> <li>• <b>Uniform:</b> Minimum, maximum;</li> <li>• <b>Normal:</b> Mean, standard deviation;</li> <li>• <b>Exponential:</b> Rate parameter;</li> <li>• <b>Gamma:</b> Shape parameter, scale parameter;</li> <li>• <b>Lognormal:</b> Log-mean, Log-standard deviation;</li> <li>• <b>Student-t:</b> Degree-of-freedom;</li> <li>• <b>Weibull:</b> Shape parameter, scale parameter;</li> <li>• <b>Cauchy:</b> Location parameter, scale parameter;</li> <li>• <b>Beta:</b> Shape parameter 1, shape parameter 2.</li> </ul>
dist.samp	An $N \times p$ matrix for the big dataset (e.g., MCMC chain) to be reduced. Exactly one of <code>dist.str</code> or <code>dist.samp</code> should be NA.

scale.flg	Should the big data <code>dist.samp</code> be normalized to unit variance before processing?
wts	Weights on each data point in <code>dist.samp</code> . Uniform weights are assigned if NA.
bd	A $p \times 2$ matrix for the lower and upper bounds of each variable.
num.subsamp	Batch size for resampling. For distributions, the default is $\max(10000, 10 \times n)$ . For data reduction, the default is $\min(10000, \text{nrow}(\text{dist.samp}))$ .
rnd.flg	Should the big data be randomly subsampled?
iter.max	Maximum iterations for optimization.
iter.min	Minimum iterations for optimization.
tol	Error tolerance for optimization.
par.flg	Should parallelization be used?
n0	Momentum parameter for optimization.

### Value

sp	An $n \times p$ matrix for support points.
ini	An $n \times p$ matrix for initial points.

### References

Mak, S. and Joseph, V. R. (2018). Support points. *Annals of Statistics*, 46(6A):2562-2592.

### Examples

```
## Not run:
#####
# Support points on distributions
#####

#Generate 25 SPs for the 2-d i.i.d. N(0,1) distribution
n <- 25 #number of points
p <- 2 #dimension
D <- sp(n,p,dist.str=rep("normal",p))

x1 <- seq(-3.5,3.5,length.out=100) #Plot contours
x2 <- seq(-3.5,3.5,length.out=100)
z <- exp(-outer(x1^2,x2^2,FUN="+")/2)
contour.default(x=x1,y=x2,z=z,drawlabels=FALSE,nlevels=10)
points(D$sp,pch=16,cex=1.25,col="red")

#####
# Generate 50 SPs for the 2-d i.i.d. Beta(2,4) distribution
#####
n <- 50
p <- 2
dist.param <- vector("list",p)
for (l in 1:p){
  dist.param[[l]] <- c(2,4)
}
```

```

}
D <- sp(n,p,dist.str=rep("beta",p),dist.param=dist.param)

x1 <- seq(0,1,length.out=100) #Plot contours
x2 <- seq(0,1,length.out=100)
z <- matrix(NA,nrow=100,ncol=100)
for (i in 1:100){
  for (j in 1:100){
    z[i,j] <- dbeta(x1[i],2,4) * dbeta(x2[j],2,4)
  }
}
contour.default(x=x1,y=x2,z=z,drawlabels=FALSE,nlevels=10 )
points(D$sp,pch=16,cex=1.25,col="red")

#####
# Generate 100 SPs for the 3-d i.i.d. Exp(1) distribution
#####
n <- 100
p <- 3
D <- sp(n,p,dist.str=rep("exponential",p))
pairs(D$sp,xlim=c(0,5),ylim=c(0,5),pch=16)

#####
# Support points for big data reduction: Franke's function
#####

#Use modified Franke's function as posterior
franke2d <- function(xx){
  if ((xx[1]>1)|| (xx[1]<0)|| (xx[2]>1)|| (xx[2]<0)){
    return(-Inf)
  }
  else{
    x1 <- xx[1]
    x2 <- xx[2]

    term1 <- 0.75 * exp(-(9*x1-2)^2/4 - (9*x2-2)^2/4)
    term2 <- 0.75 * exp(-(9*x1+1)^2/49 - (9*x2+1)/10)
    term3 <- 0.5 * exp(-(9*x1-7)^2/4 - (9*x2-3)^2/4)
    term4 <- -0.2 * exp(-(9*x1-4)^2 - (9*x2-7)^2)

    y <- term1 + term2 + term3 + term4
    return(2*log(y))
  }
}

# library(MHadaptive) # Package archived, but you can use your favorite MCMC sampler

# #Generate MCMC samples
# li_func <- franke2d #Desired log-posterior
# ini <- c(0.5,0.5) #Initial point for MCMC
# NN <- 1e5 #Number of MCMC samples desired
# burnin <- NN/2 #Number of burn-in runs
# mcmc_franke <- Metro_Hastings(li_func, pars=ini, prop_sigma=0.05*diag(2),

```

```

#               iterations=NN, burn_in=burnin)
data(mcmc_franke) # Loading MCMC sample from data

#Compute n SPs
n <- 100
D <- sp(n,2,dist.samp=mcmc_franke$trace)

#Plot SPs
oldpar <- par(mfrow=c(1,2))
x1 <- seq(0,1,length.out=100) #contours
x2 <- seq(0,1,length.out=100)
z <- matrix(NA,nrow=100,ncol=100)
for (i in 1:100){
  for (j in 1:100){
    z[i,j] <- franke2d(c(x1[i],x2[j]))
  }
}
plot(mcmc_franke$trace,pch=4,col="gray",cex=0.75,
     xlab="",ylab="",xlim=c(0,1),ylim=c(0,1)) #big data
points(D$sp,pch=16,cex=1.25,col="red")
contour.default(x=x1,y=x2,z=z,drawlabels=TRUE,nlevels=10) #contour
points(D$sp,pch=16,cex=1.25,col="red")
par(oldpar)

## End(Not run)

```

---

sp\_seq

*Computing (batch) sequential support points using difference-of-convex programming*

---

## Description

sp\_seq computes (batch) sequential support points to add onto a current point set D. Current options include sequential support points on standard distributions (specified via dist.str) or sequential support points for reducing big data (specified via dist.samp).

## Usage

```

sp_seq(D, nseq, ini=NA, num.rep=1,
       dist.str=NA, dist.param=vector("list",p),
       dist.samp=NA, scale.flg=TRUE, bd=NA,
       num.subsamp=ifelse(any(is.na(dist.samp)),
                           max(10000,10*(nseq+nrow(D))),
                           min(10000,nrow(dist.samp))),
       iter.max=max(200,iter.min), iter.min=50,
       tol=1e-10, par.flg=TRUE)

```



**Arguments**

D	An $n \times p$ matrix for the current point set.
nseq	Number of support points to add to D.
ini	An $nseq \times p$ matrix for initialization.
num.rep	Number of random restarts for optimization.
dist.str	A $p$ -length string vector for desired distribution (assuming independence). Current options include uniform, normal, exponential, gamma, lognormal, student-t, weibull, cauchy and beta. Exactly one of <code>dist.str</code> or <code>dist.samp</code> should be NA.
dist.param	A $p$ -length list for desired distribution parameters in <code>dist.str</code> . The following parameters are supported: <ul style="list-style-type: none"> <li>• <b>Uniform:</b> Minimum, maximum;</li> <li>• <b>Normal:</b> Mean, standard deviation;</li> <li>• <b>Exponential:</b> Rate parameter;</li> <li>• <b>Gamma:</b> Shape parameter, scale parameter;</li> <li>• <b>Lognormal:</b> Log-mean, Log-standard deviation;</li> <li>• <b>Student-t:</b> Degree-of-freedom;</li> <li>• <b>Weibull:</b> Shape parameter, scale parameter;</li> <li>• <b>Cauchy:</b> Location parameter, scale parameter;</li> <li>• <b>Beta:</b> Shape parameter 1, shape parameter 2.</li> </ul>
dist.samp	An $N \times p$ matrix for the big dataset (e.g., MCMC chain) to be reduced. Exactly one of <code>dist.str</code> or <code>dist.samp</code> should be NA.
scale.flg	Should the big data <code>dist.samp</code> be normalized to unit variance before processing?
bd	A $p \times 2$ matrix for the lower and upper bounds of each variable.
num.subsamp	Batch size for resampling. For distributions, the default is $\max(10000, 10*n)$ . For data reduction, the default is $\min(10000, \text{nrow}(\text{dist.samp}))$ .
iter.max	Maximum iterations for optimization.
iter.min	Minimum iterations for optimization.
tol	Error tolerance for optimization.
par.flg	Should parallelization be used?

**Value**

D	An $n \times p$ matrix for the current point set.
seq	An $nseq \times p$ matrix for the additional nseq support points.

**References**

Mak, S. and Joseph, V. R. (2018). Support points. *Annals of Statistics*, 46(6A):2562-2592.

## Examples

```

## Not run:

#####
# Generate 50 SPs for the 2-d i.i.d. N(0,1) distribution
#####
ncur <- 50
cur.sp <- sp(ncur,2,dist.str=rep("normal",2))$sp

#Add 50 sequential SPs
nseq <- 50
seq.sp <- sp_seq(cur.sp,nseq,dist.str=rep("normal",2))$seq

x1 <- seq(-3.5,3.5,length.out=100) #Plot contours
x2 <- seq(-3.5,3.5,length.out=100)
z <- exp(-outer(x1^2,x2^2,FUN="+")/2)
contour.default(x=x1,y=x2,z=z,drawlabels=FALSE,nlevels=10)
points(cur.sp,pch=4,cex=1.25,col="black",lwd=2) # (current in black)
points(seq.sp,pch=16,cex=1.25,col="red") # (new SPs in red)

#####
# Support points for big data reduction: Franke distribution
#####

#Use modified Franke's function as posterior
franke2d <- function(xx){
  if ((xx[1]>1)|| (xx[1]<0)|| (xx[2]>1)|| (xx[2]<0)){
    return(-Inf)
  }
  else{
    x1 <- xx[1]
    x2 <- xx[2]

    term1 <- 0.75 * exp(-(9*x1-2)^2/4 - (9*x2-2)^2/4)
    term2 <- 0.75 * exp(-(9*x1+1)^2/49 - (9*x2+1)/10)
    term3 <- 0.5 * exp(-(9*x1-7)^2/4 - (9*x2-3)^2/4)
    term4 <- -0.2 * exp(-(9*x1-4)^2 - (9*x2-7)^2)

    y <- term1 + term2 + term3 + term4
    return(2*log(y))
  }
}

# library(MHadaptive) # Package archived, but you can use your favorite MCMC sampler
#
# #Generate MCMC samples
# li_func <- franke2d #Desired log-posterior
# ini <- c(0.5,0.5) #Initial point for MCMC
# NN <- 1e5 #Number of MCMC samples desired
# burnin <- NN/2 #Number of burn-in runs
# mcmc_franke <- Metro_Hastings(li_func, pars=ini, prop_sigma=0.05*diag(2),

```

```
#                               iterations=NN, burn_in=burnin)
data(mcmc_franke) # Loading MCMC sample from data

#Generate ncur SPs
ncur <- 50
cur.sp <- sp(ncur,2,dist.samp=mcmc_franke$trace)$sp

#Add nseq sequential SPs
nseq <- 50
seq.sp <- sp_seq(cur.sp,nseq,dist.samp=mcmc_franke$trace)$seq

#Plot SPs
par(mfrow=c(1,2))
x1 <- seq(0,1,length.out=100) #contours
x2 <- seq(0,1,length.out=100)
z <- matrix(NA,nrow=100,ncol=100)
for (i in 1:100){
  for (j in 1:100){
    z[i,j] <- franke2d(c(x1[i],x2[j]))
  }
}
plot(mcmc_franke$trace,pch=4,col="gray",cex=0.75,
      xlab="",ylab="",xlim=c(0,1),ylim=c(0,1)) #big data
points(cur.sp,pch=4,cex=1.25,col="black",lwd=2) # (current in black)
points(seq.sp,pch=16,cex=1.25,col="red") # (new SPs in red)
contour.default(x=x1,y=x2,z=z,
                drawlabels=TRUE,nlevels=10) #contour
points(cur.sp,pch=4,cex=1.25,col="black",lwd=2) # (current in black)
points(seq.sp,pch=16,cex=1.25,col="red") # (new SPs in red)

## End(Not run)
```

# Index

\* **package**

support-package, [2](#)

e\_dist, [2](#)

mcmc\_franke, [4](#)

sp, [5](#)

sp\_seq, [8](#)

support (support-package), [2](#)

support-package, [2](#)