Package 'JQL'

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Title Jump Q-Learning for Individualized Interval-Valued Dose Rule
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Description We provide tools to estimate the individualized intervalvalued dose rule (I2DR) that maximizes the expected beneficial clinical outcome for each individual and returns an optimal interval-valued dose, by using the jump Q-learning (JQL) method. The jump Q-learning method directly models the conditional mean of the response given the dose level and the baseline covariates via jump penalized least squares regression under the framework of Q learning. We develop a searching algorithm by dynamic programming in order to find the optimal I2DR with the time complexity O(n2) and spatial complexity O(n). To alleviate the effects of misspecification of the Q-function, a residual jump Q-learning is further proposed to estimate the optimal I2DR. The outcome of interest includes the best partition of the entire dosage of interest, the regression coefficients of each partition, and the value function under the estimated I2DR as well as the Waldtype confidence interval of value function constructed through the Bootstrap.
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R topics documented:
find.I2DR 2 opt.dose 3 tune.JQL 4 tune.RJQL 5

2 find.I2DR

find.I2DR	Estimating the Individualized Interval-valued Dose Rule via (Resid-
	ual) $Jump$ Q - $learning$.

Description

This function estimates the optimal Individualized Interval-valued Dose Rule (I2DR), and calculates a Wald-type confidence interval for the value function under the estimated optimal I2DR via Bootstrap.

Usage

```
find.I2DR(Y,A,X,cm=6,method='JQL',Gamma.list=seq(from=1,to=20,by=2)/5, Lambda.list=seq(from=1,to=20,by=2)/5,RF_A.list=c(0,0.25,0.5,0.75,1), folds_num=5,alpha=0.95,nboots=500)
```

Arguments

Υ	The patient's associated response/outcome, the larger the better by convention.
A	The dose level received by each patient, should be continuous.
X	The patient's baseline covariates, could be a matrix, including continous or discrete covariates.
cm	The constent cm in m=n/cm, where m is the number of total subinterval that diverges with sample size n. The default value is 6.
method	Two methods are available, Jump Q-learning ('JQL') and Residual Jump Q-learning ('RJQL'). The default method is 'JQL'.
Gamma.list	The candidate tuning paramter space for c1 in penalty term gamma=c1 $\log(n)/n$. The default value is $seq(from=1,to=20,by=2)/5$. If the length of Gamma.list is 1, then the tuning process will be skipped.
Lambda.list	The candidate tuning paramter space for c2 in penalty term lambda= $c2 \log(n)/n$. The default value is $seq(from=1,to=20,by=2)/5$. If the length of Lambda.list is 1, then the tuning process will be skipped.
RF_A.list	The candidate tuning paramter space for A in fitted $E(Y A=a,X)$ by Random Forest Regression for method 'RJQL' only. The default value is $c(0,0.25,0.5,0.75,1)$. If the length of RF_A.list is 1, then the tuning process will be skipped.
folds_num	The number of the folds in the cross-validation process. The default value is 5.
alpha	The Confidence level. The default level is 0.95.
nboots	The number of Bootstrap. The default number is 500.

opt.dose 3

Value

An object of class "I2DR" which is a list with components:

Partition A partition of the entire dose range.

Beta The regression coefficients for each partition.

Value The estimated value function under our proposed I2DR.

low_bd The lower bound of the confidence interval.

up_bd The upper bound of the confidence interval.

method The method used to find the I2DR.

References

Jump Q-learning for Individualized Interval-valued Dose Rule.

Examples

opt.dose

Optimal Interval-valued Dose under the Individualized Interval-valued Dose Rule via (Residual) Jump Q-learning.

Description

This function assigns each individual to one of the subintervals of the entire dosage according to his/her baseline covariates under the estimated I2DR.

Usage

```
opt.dose(X,I2DR)
```

Arguments

X The patient's baseline covariates, coule be a matrix, including continous or dis-

crete covariates.

I2DR The Individualized Interval-valued Dose Rule found by the function "JQL" or

"RJQL".

Value

opt.dose The optimal Interval-valued dosage for each individual.

4 tune.JQL

References

Jump Q-learning for Individualized Interval-valued Dose Rule.

Examples

```
 \begin{array}{l} n=50 \\ d=4 \\ x=matrix(runif(n*(d-1),-1,1),nrow=n,ncol=d-1) \\ a=runif(n,0,1) \\ y=(1+x[,1])*(a>=0&a<0.35)+(x[,1]-x[,2])*(a>=0.35&a<0.65)+(1-x[,2])*(a>=0.65&a<=1)+rnorm(n,0,1) \\ rule=find.I2DR(Y=y,A=a,X=x) \\ n0=10 \\ xnew=matrix(runif(n0*(d-1),-1,1),nrow=n0,ncol=d-1) \\ opt.dose(X=xnew,I2DR=rule) \\ \end{array}
```

tune.JQL

Tuning function via k-fold cross vaidation for Jump Q-learning.

Description

This function uses the cross-validation to train the best tuning parameters lambda_n and gamma_n for Jump Q-learning.

Usage

```
tune.JQL(sample,cm=6,Gamma.list=seq(from=1,to=20,by=2)/5,
Lambda.list=seq(from=1,to=20,by=2)/5,folds_num=5)
```

Arguments

sample	The training dataset (Y,A,X), where Y is the patient's associated response/outcome, A is the dose level received by each patient, and X is the patient's baseline covariates.
cm	The constent cm in m=n/cm, where m is the number of total subinterval that diverges with sample size n. The default value is 6.
Gamma.list	The candidate tuning paramter space for c1 in penalty term gamma=c1 $\log(n)/n$. The default value is $seq(from=1,to=20,by=2)/5$.
Lambda.list	The candidate tuning paramter space for c2 in penalty term lambda=c2 log(n)/n. The default value is seq(from=1,to=20,by=2)/5.
folds_num	The number of the folds in the cross-validation process. The default value is 5.
Lambda.list	The candidate tuning paramter space for c1 in penalty term gamma=c1 log(n)/n. The default value is seq(from=1,to=20,by=2)/5. The candidate tuning paramter space for c2 in penalty term lambda=c2 log(n)/n. The default value is seq(from=1,to=20,by=2)/5.

Value

best_gamma	The best tuning parameter gamma by minimuming the least square loss function.
best_lambda	The best tuning parameter lambda by minimuming the least square loss function.

tune.RJQL 5

References

Jump Q-learning for Individualized Interval-valued Dose Rule.

Examples

```
 \begin{array}{l} n=50 \\ d=4 \\ x=matrix(runif(n*(d-1),-1,1),nrow=n,ncol=d-1) \\ a=runif(n,0,1) \\ y=(1+x[,1])*(a>=0&a<0.35)+(x[,1]-x[,2])*(a>=0.35&a<0.65)+(1-x[,2])*(a>=0.65&a<=1)+rnorm(n,0,1) \\ sample=data.frame(y=y,a=a,x=x) \\ tune.JQL(sample) \\ \end{array}
```

tune.RJQL

Tuning function via k-fold cross vaidation for Residual Jump Q-learning.

Description

This function uses the cross-validation to train the best tuning parameters lambda_n and gamma_n for Residual Jump Q-learning.

Usage

```
tune.RJQL(sample,cm=6,Gamma.list=seq(from=1,to=20,by=2)/5,\\ Lambda.list=seq(from=1,to=20,by=2)/5,RF_A.list=c(0,0.25,0.5,0.75,1),folds_num=5)
```

Arguments

sample	The training dataset (Y,A,X) , where Y is the patient's associated response/outcome, A is the dose level received by each patient, and X is the patient's baseline covariates.
CM	The constent cm in m=n/cm, where m is the number of total subinterval that diverges with sample size n. The default value is 6.
Gamma.list	The candidate tuning paramter space for c1 in penalty term gamma=c1 log(n)/n. The default value is seq(from=1,to=20,by=2)/5.
Lambda.list	The candidate tuning paramter space for c2 in penalty term lambda=c2 log(n)/n. The default value is seq(from=1,to=20,by=2)/5.
RF_A.list	The candidate tuning paramter space for A in fitted $E(Y A=a,X)$ by Random Forest Regression for method 'RJQL' only. The default value is $c(0,0.25,0.5,0.75,1)$.
folds_num	The number of the folds in the cross-validation process. The default value is 5.

Value

best_gamma	The best tuning parameter gamma by minimuming the least square loss function.
best_lambda	The best tuning parameter lambda by minimuming the least square loss function.
best_a	The best tuning parameter a to fit random forest by minimuming the least square
	loss function.

6 tune.RJQL

References

Jump Q-learning for Individualized Interval-valued Dose Rule.

Examples

```
 \begin{array}{l} n=50 \\ d=4 \\ x=matrix(runif(n*(d-1),-1,1),nrow=n,ncol=d-1) \\ a=runif(n,0,1) \\ y=(1+x[,1])*(a>=0&a<0.35)+(x[,1]-x[,2])*(a>=0.35&a<0.65)+(1-x[,2])*(a>=0.65&a<=1)+rnorm(n,0,1) \\ sample=data.frame(y=y,a=a,x=x) \\ tune.RJQL(sample) \\ \end{array}
```

Index

```
find.I2DR, 2
opt.dose, 3
tune.JQL, 4
tune.RJQL, 5
```