RMoCap package documentation

July 26, 2018

July 20, 2016
Type Package
Title Package for processing and analysis motion capture (mocap) data.
Version 1.0.0
Author Tomasz Hachaj, Marek R. Ogiela
Maintainer Tomasz Hachaj <tomekhachaj@o2.pl></tomekhachaj@o2.pl>
Description With functions from this package you can: - load and save Biovision Hierarchy (BVH) files, - convert direct kinematic model to hierarchical kinematic model and vice versa, - plot interactive plots of motion capture (mocap) data, - align two mocap recordings, - while walking with translation and acceleration data is is possible to calculate correct body joints displacement,
average many motion capture recordings that presents the same activity and store it as a single one - perform comparision analysis of two motion capture recordings using advanced Dynamic Time Wapring - besed procedures and display results, This package has also many useful algebraic functions like Quaternion Markley averaging algorithms etc.
License GPL (>= 3)
Encoding UTF-8
LazyData true
Imports smoother, rgl, RSpincalc, subplex, signal, compiler
RoxygenNote 6.0.1
R topics documented:
aligninputandrefdata

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aligninputandrefdata This function translate inputdata so that it has common values of limbname Dx, Dy, Dz columns with refdata.

Description

This function translate inputdata so that it has common values of limbname Dx, Dy, Dz columns with refdata.

Usage

aligninputandrefdata(inputdata, refdata, limbname)

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Arguments

input data input motion capture data frame.
refdata reference motion capture data frame.

1 name of the column which is used to perform alignment.

Value

data frame aligned as it is described above.

Examples

#See example from analyze.mocap function

analyze.mocap

This function performs motion capture dat analysis based on Dynamic Time Warping.

Description

This procedure detects highest differences between reference and input mocap recording. Analysis goes as follows:

- Perform DTW on x1 and x2 from list at index 1. Plot DTW alignment This alignment will be used in aligning all other signals. Plot distance between x1 and x2 after alignment and find local maxima in this plot. In following analysis only maxima with relative value above treshold will be used.
- perform DTW alignment for each other element from the data.configuration list, however use the alignment function from first step. Maxima are detected in the same procedure as above, however we take into account only those of them, that are close enough to maxims from first step of analysis.

Usage

```
analyze.mocap(data.configuration, ref.d = NULL, in.d = NULL,
   extremumtreshold = 0.66, smoothSize = 0.1)
```

Arguments

data.configuration

a list containing configuration for the algorithm. Each element of the list is a list with following elements (all elements are obligatory):

- x1 reference signal for DTW (vectors list),
- x2 second signal for DTW (vectors list),
- FUN distance function for DTW, use euc.dist or euc.dist1d, however you can define any function that can opperates on x1 and x2,
- ylab label on Y axis of the results plot,
- legend part of the legend sting over plots,
- plotRGL name of the body joint for which DTW alignment of x1 and x2 will be drawn. It will be 3D rgl plot. If plotRGL = NULL plot will not be drawn.

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• skeleton - object of class mocap. It is used to get joints relations while plotting RGL plot. data frame with reference mocap data (default is ref.d=NULL). It is used to get joints relations while plotting RGL plot. data frame with reference mocap data (default is in.d=NULL). It is used to get joints relations while plotting RGL plot. extremumtreshold treshold from range [0,1], that is used to remove local extreme, which realtive value is below extremumtreshold (default value is extremumtreshold=0.66).

size of the gaussian smoothing window. Deafult value is smoothSize = 0.1. smoothSize

Value

a list containing data.configuration parameters plus algorihm results

data.configuration[[3]] <- list(x1 = x1[[1]],

Examples

ref.d

in.d

```
###########################
#analyze upper body data
##########################
data(right.arm.motion.1)
data(right.arm.motion.2)
refdata <- right.arm.motion.1$data.frame</pre>
inputdata <- right.arm.motion.2$data.frame</pre>
extremumtreshold <- 0.66
smoothSize <- 0.1</pre>
inputdataalignment <- rotatedata(inputdata, refdata, "LeftShoulder", "RightShoulder")</pre>
inputdataalignmentkinematic <- calculate.kinematic(inputdataalignment, bodypartname = "LeftShoulder")</pre>
refdatakinematic <- calculate.kinematic(refdata, bodypartname = "LeftShoulder")</pre>
inputdataalignmentkinematic <- aligninputandrefdata(inputdataalignmentkinematic, refdatakinematic, limbname
data.configuration <- list()</pre>
data.configuration[[1]] <- list(x1 = vector.to.list(refdatakinematic, "RightHand"),</pre>
   x2 = vector.to.list(inputdataalignmentkinematic, "RightHand"),
   FUN = euc.dist,
   ylab = "Distance [cm]",
   legend = "RightHand",
   plotRGL = "RightHand",
   skeleton = right.arm.motion.1)
data.configuration[[2]] <- list(x1 = vector.to.angles.list(refdatakinematic, "RightShoulder", "RightArm", "RightArm", "RightArm")</pre>
  x2 = vector.to.angles.list(inputdataalignmentkinematic, "RightShoulder", "RightArm", "RightForearm"),
   FUN = euc.dist1d,
   ylab = "Angle [rad]"
   legend = "Right elbow",
   plotRGL = NULL,
   skeleton = NULL)
x1 <- vector.to.angles.frame.list(refdatakinematic, "RightArm", "RightForearm", "RightShoulder", "LeftShoulde
x2 <- vector.to.angles.frame.list(inputdataalignmentkinematic, "RightArm", "RightForearm", "RightShoulder",
```

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```
x2 = x2[[1]],
   FUN = euc.dist1d,
   ylab = "Angle [rad]",
   legend = "X angle between RightArm and RightForearm",
   plotRGL = NULL,
   skeleton = NULL)
data.configuration[[4]] <- list(x1 = x1[[2]],
  x2 = x2\Gamma\Gamma277.
   FUN = euc.dist1d,
   ylab = "Angle [rad]",
   legend = "Y angle between RightArm and RightForearm",
   plotRGL = NULL,
   skeleton = NULL)
data.configuration[[5]] <- list(x1 = x1[[3]],
  x2 = x2[[3]],
   FUN = euc.dist1d,
   ylab = "Angle [rad]",
   legend = "Z angle between RightArm and RightForearm",
   plotRGL = NULL,
   skeleton = NULL)
res <- analyze.mocap(data.configuration,</pre>
refdatakinematic,
inputdataalignmentkinematic,
extremumtreshold,
smoothSize)
#analyze lower body data
############################
data(mawashi.geri.left.1)
data(mawashi.geri.left.2)
refdata <- mawashi.geri.left.1$data.frame</pre>
inputdata <- mawashi.geri.left.2$data.frame</pre>
extremumtreshold <- 0.66
smoothSize <- 0.1</pre>
inputdataalignment <- rotatedata(inputdata, refdata, "LeftThigh", "RightThigh")</pre>
inputdataalignmentkinematic <- calculate.kinematic(inputdataalignment, bodypartname = "RightFoot")</pre>
refdatakinematic <- calculate.kinematic(refdata, bodypartname = "RightFoot")</pre>
inputdataalignmentkinematic <- aligninputandrefdata(inputdataalignmentkinematic, refdatakinematic, limbname
data.configuration <- list()</pre>
data.configuration[[1]] <- list(x1 = vector.to.list(refdatakinematic, "LeftFoot"),</pre>
x2 = vector.to.list(inputdataalignmentkinematic, "LeftFoot"),
FUN = euc.dist,
ylab = "Distance [cm]",
 legend = "LeftFoot",
 plotRGL = "LeftFoot",
  skeleton = mawashi.geri.left.1)
x2 = vector.to.angles.list(inputdataalignmentkinematic, "LeftThigh", "LeftLeg", "LeftFoot"),
 FUN = euc.dist1d,
```

averaged.mocap-class

```
ylab = "Angle [rad]",
  legend = "Left knee",
  plotRGL = NULL)
x1 <- vector.to.angles.frame.list(refdatakinematic, "LeftThigh", "LeftLeg", "LeftThigh", "RightThigh")
x2 <- vector.to.angles.frame.list(inputdataalignmentkinematic, "LeftThigh", "LeftLeg", "LeftThigh", "RightThi
data.configuration[[3]] <- list(x1 = x1[[1]],</pre>
  x2 = x2[[1]],
  FUN = euc.dist1d,
  ylab = "Angle [rad]",
  legend = "X angle between LeftThigh and LeftLeg",
  plotRGL = NULL)
data.configuration[[4]] <- list(x1 = x1[[2]],
  x2 = x2[[2]],
  FUN = euc.dist1d,
  ylab = "Angle [rad]",
  legend = "Y angle between LeftThigh and LeftLeg",
  plotRGL = NULL)
data.configuration[[5]] <- list(x1 = x1[[3]],
  x2 = x2[[3]],
  FUN = euc.dist1d,
  ylab = "Angle [rad]",
  legend = "Z angle between LeftThigh and LeftLeg",
  plotRGL = NULL)
res <- analyze.mocap(data.configuration,</pre>
refdatakinematic,
inputdataalignmentkinematic,
extremumtreshold,
smoothSize)
```

averaged.mocap-class A class returned by mocap.averaging function.

Description

A class returned by mocap.averaging function.

Usage

```
see documentation of mocap.averaging.
```

Format

a list containing data frame (fullData) and data frame (norm.distance) with normalized distance optimized during averaging.

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Examples

```
data("mawashi.geri.right.list")
myList <- list()
for (a in 1:length(mawashi.geri.right.list))
{
    myList[[a]] <-mawashi.geri.right.list[[a]]$data.frame
}
res.data <- mocap.averagingCmp(myList, 2, eps = 0.000001)
plot(res.data)</pre>
```

avg.quaternion.markley

Quaternion Markley averaging algorithms.

Description

See: F. Landis Markley, Yang Cheng, John Lucas Crassidis, and Yaakov Oshman. "Averaging Quaternions", Journal of Guidance, Control, and Dynamics, Vol. 30, No. 4 (2007), pp. 1193-1197. https://doi.org/10.2514/1.28949

Usage

```
avg.quaternion.markley(Q)
```

Arguments

Q a data frame of quaternions (four dimensional vectors) to be averaged. Each row of data frame holds one quaternion.

Value

4D quaternion vector.

```
Q <- data.frame(c(0.9999986, 0.9999986, 0.9999986, 0.9999986, 0.9999986, 0.9999986), c(0.0008716584, 0.0008716584, 0.0009590162, 0.0009590162, 0.001046359, 0.001046359), c(0.0009608439, 0.001048034, 0.0008736689, 0.001048034, 0.0009608439, 0.0009590162, 0.001046359, 0.0008716584, 0.0008716584, 0.0009590162)) avg.quaternion.markley(Q)
```

8 calculate.kinematic

bvh.to.df	This function get direct kinematic from list generated with function read.bvh or read.mocap function and returns it in the form of data frame.

Description

This function does not perform any algebraic calculation - it just takes data from Dxyz and Rxyz columns. Function can additionally calculate second derivative of Dxyz.

Usage

```
bvh.to.df(skeleton, sd = TRUE)
```

Arguments

skeleton input hierarchical kinematic model.

sd does function should calculate second derivative? Default = TRUE.

Value

data frame with direct kinematic.

Examples

```
#an example BVH file
data("heian.nidan.bvh")
#write file to the disc
f <- file("heian.nidan.bvh")
writeChar(con = f, object = heian.nidan.bvh)
close(f)
#read hierarchical model stored in BVH file
heian.nidan <- read.bvh("heian.nidan.bvh")
df <- bvh.to.df(heian.nidan)</pre>
```

calculate.kinematic

This function corrects translation of the direct kinematic model.

Description

This heuristic method is especially usable while dealing with data acquired by IMU (Inertial measurement unit) sensors. There are two possible calls of this method. The first one utilize acceleration data of body joints (.ax, .ay and .az columns). If this data is avilable method can calculate displacement of long motion, during which an actor uses both left and right leg. If this call is used, one should left bodypartname as NULL. The second call is used when we are dealing with data without acceleration data and actor has one stationary limb. In this case bodypartname shoule have value of the limb which does not translate during motion.

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Usage

```
calculate.kinematic(dd, LeftFoot = "LeftFoot", RightFoot = "RightFoot",
    show.plot = FALSE, plot.title = "", bodypartname = NULL, dyEps = 5)
```

Arguments

dd data frame with motion capture data.

LeftFoot name of the column with data that holds coordinates of left foot (joint that touch

the ground). Default value is LeftFoot = "LeftFoot". Used only for the first type

of call.

RightFoot name of the column with data that holds coordinates of right foot (joint that

touch the ground). Default value is RightFoot = "RightFoot". Used only for the

first type of call.

show.plot if TRUE (default is show.plot = FALSE) plots results of an algorithm.

plot.title part of the title over plots.

bodypartname if not NULL value the stationary joint of the motion is column with bodypart-

name. Default value is bodypartname = NULL.

dyEps Treshold value for translation calculation (used only for the first type of call).

Default value is dyEps = 5.

Value

Data frame, which has Dxzy columns updated. All other columns are not updated.

```
#This example uses acceleration data to calculates correct displacement.
  #header of mocap file
  data("header.mocap")
  #data frame with displacements and acceleration data
  data("heian.shodan")
heian.shodan.corrected <- calculate.kinematic(heian.shodan, show.plot = "TRUE", plot.title = "Heian Shodan")
 original.bvh <- set.data.frame(header.mocap, heian.shodan)</pre>
  corrected.bvh <- set.data.frame(header.mocap, heian.shodan.corrected)</pre>
  #plot BVH, red is original data, green is corrected
 plot(original.bvh, frames.fraction = 0.1, my.color = "red", alpha = 0.1, spheres = FALSE)
plot(corrected.bvh, frames.fraction = 0.1, my.color = "green", alpha = 0.1, spheres = FALSE, append = TRUE)
  #writting BVH to disk
  write.bvh(original.bvh, "original.bvh")
  write.bvh(corrected.bvh, "corrected.bvh")
 #This example uses single body joint which coordinates should be constant during motion
  data(mawashi.geri.left.1)
plot(mawashi.geri.left.1, frames.fraction = 0.1, my.color = "red", alpha = 0.5, spheres = FALSE)
\verb|mawashi.geri.left.1$| data.frame <- calculate.kinematic(mawashi.geri.left.1$| data.frame, bodypartname = "Right of the content of the calculate.kinematic(mawashi.geri.left.1$| data.frame, bodypartname = "Right of the content of the calculate.kinematic(mawashi.geri.left.1$| data.frame, bodypartname = "Right of the content of the calculate.kinematic(mawashi.geri.left.1$| data.frame, bodypartname = "Right of the content of the calculate.kinematic(mawashi.geri.left.1$| data.frame, bodypartname = "Right of the content of the calculate.kinematic(mawashi.geri.left.1$| data.frame, bodypartname = "Right of the content of the calculate.kinematic(mawashi.geri.left.1$| data.frame, bodypartname = "Right of the content of the calculate.kinematic(mawashi.geri.left.1$| data.frame, bodypartname = "Right of the content of the calculate.kinematic(mawashi.geri.left.1$| data.frame, bodypartname = "Right of the calculate.kinematic(mawashi.geri.left.1$| data.frame, bodypartname = "Right of the calculate.kinematic(mawashi.geri.left.1$| data.frame = "Right of the calculate.kinematic(m
plot(mawashi.geri.left.1, frames.fraction = 0.1, my.color = "green", alpha = 0.5, spheres = FALSE, append = TR
```

10 df.to.bvh

df.to.bvh

This function recalculate direct to hierarchical kinematic model.

Description

Procedure implements iterative algebraic procedure with additional initial optimization, that is required to align root body joints. Optimization is done using simplex method. The rotation order in hierarchical model is automatically set to ZYX, even if input.skeleton has different order.

Usage

```
df.to.bvh(input.skeleton, df.to.save, plot.me = FALSE, frame.id = -1,
  debug.messages = FALSE)
```

Arguments

input.skeleton object of mocap class that defines hierarchical kinematic model.

df.to.save data frame with column names compatible with input.skeleton. Data that is used for calculation has to be placed in columns with names endind .Dx, .Dy and .Dz.

plot.me if TRUE plot steps of skeleton aligning of frame with index frame.id. Default value is plot.me = FALSE.

frame.id if frame.id > 0 and plot.me = TRUE plot steps of skeleton aligning of frame with index frame.id. Default value is frame.id = -1.

debug.messages print additional messages informing about calculation progress.

Value

object of class mocap.

Examples

}

points(df.to.save[,2],col="red", pch = 2)

```
data("header.mocap")
data("heian.yondan")

input.skeleton <- header.mocap

df.to.save <- heian.yondan[1:300,]
first.frame <- df.to.bvh(input.skeleton, df.to.save, plot.me = FALSE, debug.messages = TRUE)
write.bvh(first.frame, "e:\bvh in r\\gotowy_kod\\output\\heian.yondan.frames300.bvh")

plot(first.frame$skeleton$Joints[[1]]$Rxyz[,1], type = "1", col = "black", xlab = "sample", ylab = "angle (deglines(first.frame$skeleton$Joints[[1]]$Rxyz[,2], type = "1", col = "red")
lines(first.frame$skeleton$Joints[[1]]$Rxyz[,3], type = "1", col = "blue")
legend("bottomright", legend=c("X axis rotation", "Y axis rotation", "Z axis rotation"), col=c("black", "red" title("Hips rotation data")

plot(df.to.save[,2], ylab = "Displacement [cm]", xlab = "Time [10^-2 sec]", pch = 1)
for (a in 1:ncol(df.to.save))
{
    df.to.save[,a] <- jitter(df.to.save[,a], factor = 500)</pre>
```

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```
legend("bottomright", legend=c("Original", "Jitter"), col=c("black", "red"), pch = c(1,2))
title("Example channel of MoCap data")

first.frame <- df.to.bvh(input.skeleton, df.to.save, plot.me = FALSE, debug.messages = TRUE)

#plot rotation data
plot(first.frame$skeleton$Joints[[1]]$Rxyz[,1], type = "l", col = "black", xlab = "sample", ylab = "angle (deglines(first.frame$skeleton$Joints[[1]]$Rxyz[,2], type = "l", col = "red")
lines(first.frame$skeleton$Joints[[1]]$Rxyz[,3], type = "l", col = "blue")
legend("bottomright", legend=c("X axis rotation", "Y axis rotation", "Z axis rotation"), col=c("black", "red" title("Hips rotation data")

write.bvh(first.frame, "e:\\bvh in r\\gotowy_kod\\output\\jitter.heian.yondan.frames300.bvh")

df.to.save <- heian.yondan[1000:1001,]
foo <- df.to.bvh(input.skeleton, df.to.save, plot.me = TRUE, debug.messages = FALSE, frame.id = 1)</pre>
```

euc.dist

This function calulcates Euclidean distance between vectors x1 and x2.

Description

This function calulcates Euclidean distance between vectors x1 and x2.

Usage

```
euc.dist(x1, x2)
```

Arguments

x1 first numeric vector.

x2 second numeric vector.

Value

```
\operatorname{asqrt}(\operatorname{sum}((\operatorname{x} 1 - \operatorname{x} 2) \wedge 2)).
```

```
euc.dist(c(1,2,3,4),c(-5,0,-6,3))
```

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euc.dist1d

This function returns absolute value of difference between x1 and x2.

Description

This function returns absolute value of difference between x1 and x2.

Usage

```
euc.dist1d(x1, x2)
```

Arguments

x1 first numeric value.x2 second numeric value.

Value

```
abs(x1[1] - x2[1]).
```

Examples

```
euc.dist1d(1,-5)
```

generate.first.frame

This function generates object of mocap class with zero rotation data.

Description

This function generates object of mocap class with zero rotation data.

Usage

```
generate.first.frame(input.skeleton, Nframes = 1, FrameTime = 0.01)
```

Arguments

input.skeleton object of mocap class which defines hierarchical model.

Nframes number of frames to be generated.

FrameTime intervals between acquisitions.

Value

object of mocap class.

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Examples

```
data("heian.nidan.bvh")
f <- file("heian.nidan.bvh")
writeChar(con = f, object = heian.nidan.bvh)
close(f)
#read hierarchical model stored in hierarchical BVH file
heian.nidan <- read.mocap("heian.nidan.bvh")
first.frame <- generate.first.frame(heian.nidan)
plot(heian.nidan, frame = 1, alpha = 1, spheres = TRUE)
plot(first.frame, my.color = "red", frame = 1, alpha = 1, spheres = TRUE, append = TRUE)</pre>
```

header.mocap

An example object of class mocap that does not have any motion data.

Description

An example object of class mocap that does not have any motion data.

Usage

```
data("header.mocap")
```

Format

An object of mocap class.

heian.nidan.bvh

A raw byh file to be saved on disc (Shotokan Karate kata Heian Nidan).

Description

A raw bvh file to be saved on disc (Shotokan Karate kata Heian Nidan).

Usage

```
data(heian.nidan.bvh)
```

Format

A raw file.

```
data("heian.nidan.bvh")
f <- file("e:\\bvh in r\\gotowy_kod\\output\\heian.nidan.bvh")
writeChar(con = f, object = heian.nidan.bvh)
close(f)</pre>
```

heian.shodan	A data frame with mocap data (Shotokan Karate kata Heian Shodan). Also acceleration channels are included.

Description

A data frame with mocap data (Shotokan Karate kata Heian Shodan). Also acceleration channels are included.

Usage

```
data("heian.shodan")
```

Format

A data frame.

heian.yondan

A data frame with mocap data (Shotokan Karate kata Heian Yondan).

Description

A data frame with mocap data (Shotokan Karate kata Heian Yondan).

Usage

```
data("heian.yondan")
```

Format

A data frame.

hierarchical.to.direct.kinematic

Generates direct kinematic model from hierarchical kinematic model.

Description

This function makes calculations based on hierarchical kinematic data in input list (input.skeleton). It does not use Dxyz from input.skeleton.

Usage

```
hierarchical.to.direct.kinematic(input.skeleton)
```

Arguments

input.skeleton list in the same format as one generated with read.mocap function.

mawashi.geri.left.1

Value

data frame with direct kinematic model. Names of the columns are the same as in input.skeleton.

Examples

```
#an example BVH file
data("heian.nidan.bvh")
f <- file("heian.nidan.bvh")
writeChar(con = f, object = heian.nidan.bvh)
close(f)
#read hierarchical model stored in hierarchical BVH file
heian.nidan <- read.mocap("heian.nidan.bvh")
#plot kinematic data
plot(x = heian.nidan$data.frame$Hips.Dx, y = heian.nidan$data.frame$Hips.Dz, type = "l", ylab = "Displacement title("Hips displacement during motion")

#generate kinematic from hierarchical model - same results as above
df <- hierarchical.to.direct.kinematic(heian.nidan$skeleton)
plot(x = df$Hips.Dx, y = df$Hips.Dz, type = "l", ylab = "Displacement X [cm]", xlab = "Displacement Z [cm]")
title("Hips displacement during motion")</pre>
```

mawashi.geri.left.1 An object of class mocap that contains karate kick nawashi geri.

Description

An object of class mocap that contains karate kick nawashi geri.

Usage

```
data("mawashi.geri.left.1")
```

Format

An object of mocap class.

mawashi.geri.left.2 An object of class mocap that contains karate kick nawashi geri.

Description

An object of class mocap that contains karate kick nawashi geri.

Usage

```
data("mawashi.geri.left.2")
```

Format

An object of mocap class.

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```
mawashi.geri.right.list
```

A list of ten object of class mocap that contains karate kicks mawashi geri.

Description

A list of ten object of class mocap that contains karate kicks mawashi geri.

Usage

```
data("mawashi.geri.right.list")
```

Format

A list of objects of mocap class.

mocap-class

A class returned by read.mocap function.

Description

A class returned by read.mocap function.

Usage

```
read.mocap("file.name.bvh")
```

Format

a list containing:

- Joints list of joints,
- Time vector with time series,
- FrameTime value of time interval between samples
- Frame and samples count.

Each joint is a list that contains:

- Nestdepth level of joint in hierarchy,
- Name name of the joint,
- Parent id of the parent on the list, root joint has parent = -1,
- Offset 3D vector with offset from parent joint,
- Nchannels number of data channels (6 from root, 3 for other or 0 for end point),
- Order rotation order (accepted orders are XYZ, XZY, YXZ, YZX, ZXY or ZYX),
- Dxyz matrix with direct kinematic displacement (calculated from original data),
- RawDxyz matrix with direct kinematic displacement, present only in root joint,
- Rxyz matrix with rotation in degrees of hierarchical kinematic model
- Trans list of rotation translation matrices that are used to recalculates hierarchical to direct kinematic model.

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Examples

```
#an example BVH file
data("heian.nidan.bvh")
#write file to the disc
f <- file("heian.nidan.bvh")
writeChar(con = f, object = heian.nidan.bvh)
close(f)
#read hierarchical model stored in BVH file
heian.nidan <- read.mocap("heian.nidan.bvh")
summary(heian.nidan)</pre>
```

mocap.averaging

This function averages a list of motion capture recordings.

Description

Averaging is performed on each rotation channel of hierarchical model sepparetly with Dynami Time Warping barycenter averagin (DBA), see: François Petitjean, Alain Ketterlin, PierreGançarski, "A global averaging method for dynamic time warping, with applications to clustering", Pattern Recognition, Volume 44, Issue 3, March 2011, Pages 678-693, https://doi.org/10.1016/j.patcog.2010.09.013 Each rotation signal holds rotation represented as quaternion. Quaternion averging is performed with Quaternion Markley averaging algorithms, see: F. Landis Markley, Yang Cheng, John Lucas Crassidis, and Yaakov Oshman. "Averaging Quaternions", Journal of Guidance, Control, and Dynamics, Vol. 30, No. 4 (2007), pp. 1193-1197. https://doi.org/10.2514/1.28949 Results are smoothed with Weighted Quaternion Markley averaging algorithms using gaussian kernel.

Usage

```
mocap.averaging(myList, DBAIterationsCount = 50, eps = 1e-04,
    plot.me = TRUE)
```

Arguments

myList

list of mocap data frames. Algorithm uses columns with names that has .Rx, .Ry and .Rz names. Rotation should be represented by Euler angles in degrees.

DBAIterationsCount

maximal number of itereations of DBA algorithm (deafulat value is DBAItera-

tionsCount = 50).

eps treshold value for DBA - iteration stops when absolute value of difference be-

tween normalized DTW distances on this and previous iteration is less than eps

(default value is eps = 0.0001).

plot.me if TRUE, plots DTW distances for each averaged signal (deafulat value is plot.me

= 50).

Value

return object of class averaged.mocap.

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Examples

```
#load list of objects of mocap class
data("mawashi.geri.right.list")
myList <- list()
#assign data frames to list
for (a in 1:length(mawashi.geri.right.list))
{
    myList[[a]] <-mawashi.geri.right.list[[a]]$data.frame
}
#run compiled version of mocap.averaging function
res.data <- mocap.averagingCmp(myList, 50, eps = 0.000001)
plot(res.data)
#write results to disc as bvh file
skel <- set.data.frame(mawashi.geri.right.list[[1]], res.data$fullData)
write.bvh(path = "avg.mawashi.geri.right.bvh", skeleton.helper = skel)</pre>
```

myEV2Q

Calculates quaternion from axis angle.

Description

All other quternion function are impoted from package RSpincalc.

Usage

```
myEV2Q(axis, angle)
```

Arguments

```
axis 3D vector. angle in radians.
```

Value

4D quaternon vector.

```
\label{eq:myEV2Q} myEV2Q(vector.to.unit(c(1,2,3)),pi/4)
```

plot.averaged.mocap 19

plot.averaged.mocap

Plots normalized distance of all joints from averaged.mocap class.

Description

Plots normalized distance of all joints from averaged.mocap class.

Usage

```
## S3 method for class 'averaged.mocap'
plot(data.to.plot)
```

Arguments

data.to.plot object of class averaged.mocap.

Examples

```
data("mawashi.geri.right.list")
myList <- list()
for (a in 1:length(mawashi.geri.right.list))
{
    myList[[a]] <-mawashi.geri.right.list[[a]]$data.frame
}
res.data <- mocap.averagingCmp(myList, 2, eps = 0.000001)
plot(res.data)</pre>
```

plot.mocap

Overrides plot function to work with mocap class.

Description

This function uses rgl package for 3D visualizations. It creates interactive 3D plots that enables rotation and scalling.

Usage

```
## S3 method for class 'mocap'
plot(obj, frame = 0, my.color = "green",
   frames.fraction = 0.1, alpha = 0.05, spheres = FALSE, append = FALSE,
   print.text = FALSE)
```

Arguments

obj mocap object.

frame index of frame to be show. If default (frame = 0) all frames are drawn.

my.color color of the plot.

frames.fraction

if frame = 0 this parameter indicates what fraction of frames should be drawn (default frames.fraction = 0.1 means that 10% of frames will be ploted).

20 read.bvh

alpha value of alpha channel of the plot (0 is 100% transparency, 1 is no transparency,

default is alpha = 0.05).

spheres if TRUE, position of body joints will be marked as spheres (default is spheres =

FALSE).

append if FALSE (default is append = FALSE) a new plot is generated, if TRUE a plot

is drawn over open rgl window).

print.text if TRUE (default is append = FALSE) plots name of the joints.

Examples

```
data("right.arm.motion.1")
plot(right.arm.motion.1, frame = 1, my.color = "white", alpha = 1, spheres = TRUE)
plot(right.arm.motion.1, frames.fraction = 0.5, my.color = "white", alpha = 1, spheres = FALSE, print.text = TRUE)
```

read.bvh This function read file in BVH (Biovision Hierarchy) format and recal-

culates it to direct kinematic model.

Description

BVH file contains a hierarchical kinematic model definition (called a skeleton) and motion data. For more information see: "Motion Capture File Formats Explained" by M. Meredith S. Maddock, doi: 10.1.1.103.2097.

Usage

```
read.bvh(filepath)
```

Arguments

filepath A path to BVH file.

Value

an object of mocap class.

```
#an example BVH file
data("heian.nidan.bvh")
#write file to the disc
f <- file("heian.nidan.bvh")
writeChar(con = f, object = heian.nidan.bvh)
close(f)
#read hierarchical model stored in BVH file</pre>
```

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read.mocap

This function reads motion capture file on BVH format.

Description

It also caluclates direct kinematic from hierarchical kinematic. This function calls read.bvh and bvh.to.df to generate single object of mocap class. See documentation for those two functions.

Usage

```
read.mocap(filepath)
```

Arguments

```
filepath path to a file.
```

Value

object of mocap class.

Examples

```
#an example BVH file
data("heian.nidan.bvh")
f <- file("eian.nidan.bvh")
writeChar(con = f, object = heian.nidan.bvh)
close(f)
#read hierarchical model stored in hierarchical BVH file
heian.nidan <- read.mocap("heian.nidan.bvh")
summary(heian.nidan)</pre>
```

right.arm.motion.2

An object of class mocap that contains motion of right arm.

Description

An object of class mocap that contains motion of right arm.

Usage

```
data("right.arm.motion.2")
```

Format

An object of mocap class.

rotatedata

This function align to data frames that contains mocap data.

Description

Both data frames need to have two common column groups with names ending Dx and Dz. This function calulates vector vv = v1-v2 for both data frames (vv.m for mydata and vv.ref for reference-data) and rotates mydata around Y axis in order to minimize euclidean distane between vv.m and vv.ref. Minimization is performed with simplex method. After this procedure mydata face the same direction as referencedata. This procedure works correctly only if root joint of mocap is stationary.

Usage

```
rotatedata(mydata, referencedata, v1, v2)
```

Arguments

mydata input data frame with mocap data to be algined to referencedata.

referencedata reference data frame with mocap data.

v1 name of the first body joint.v2 name of the second body joint.

Value

mydata rotated by Y axis so that mydata and referencedata faces same direction.

Examples

```
data(mawashi.geri.left.1)
data(mawashi.geri.left.2)
refdata <- mawashi.geri.left.1$data.frame
inputdata <- mawashi.geri.left.2$data.frame
#after following function inputdata and refdata are alignined towards vector LeftThigh - RightThigh
inputdataalignment <- rotatedata(inputdata, refdata, "LeftThigh", "RightThigh")</pre>
```

rotation.matrix.between.vectors

This function returns n by n rotation matrix that align vector x onto y.

Description

Bth vector x and y has to be normalized and has to be same length.

Usage

```
rotation.matrix.between.vectors(x, y)
```

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Arguments

```
x vector to be rotated (has to be normalized).y reference vector (has to be normalized).
```

Value

n x n rotation matrix.

Examples

```
x <- vector.to.unit(c(1,0,0))
y <- vector.to.unit(c(0,1,0))
Rx2y <- rotation.matrix.between.vectors(x, y)
x %*% Rx2y
y
x <- vector.to.unit(c(5,0,2,4,6))
y <- vector.to.unit(c(0,1,0,8,-5))
Rx2y <- rotation.matrix.between.vectors(x, y)
x %*% Rx2y
y
x <- vector.to.unit(c(8,0,0))
y <- vector.to.unit(c(2,0,0))
Rx2y <- rotation.matrix.between.vectors(x, y)
x %*% Rx2y
y</pre>
```

second.derivative

Calculates second derivative

Description

Calculates second derivative using Central Difference Operator.

Usage

```
second.derivative(signal)
```

Arguments

signal

input vecotr.

Value

vector containing second derivative calculated from the input vector.

```
second.derivative(sin(seq(from=0, to=2*pi,by=pi/50)))
```

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set.data.frame

Assign data frame to object of mocap class.

Description

This function does not recalculate chierarchical kinematic model in skeleton.

Usage

```
set.data.frame(skel, df)
```

Arguments

skel object of mocap class.

df data frame with columns names compatible to hierarchical model definition in

skel object.

Value

object of mocap class with df assigned.

Examples

```
data("header.mocap")
data("heian.shodan")
print(header.mocap$skeleton$Frames)
original.bvh <- set.data.frame(header.mocap, heian.shodan)
print(original.bvh$skeleton$Frames)</pre>
```

summary.mocap

Plots information about number of frames, joints count and joints chierarchy of mocap class.

Description

Plots information about number of frames, joints count and joints chierarchy of mocap class.

Usage

```
## S3 method for class 'mocap'
summary(mocap.data)
```

Arguments

mocap.data an object of mocap class.

vector.angle 25

Examples

```
data("heian.nidan.bvh")
f <- file("e:\\bvh in r\\gotowy_kod\\output\\heian.nidan.bvh")
writeChar(con = f, object = heian.nidan.bvh)
close(f)
#read hierarchical model stored in hierarchical BVH file
heian.nidan <- read.mocap("e:\\bvh in r\\gotowy_kod\\output\\heian.nidan.bvh")
summary(heian.nidan)</pre>
```

vector.angle

This function calculates angle between two vectors on the plane designed by those vectors.

Description

The return value is in the range [0,pi].

Usage

```
vector.angle(a, b)
```

Arguments

a first vector.b second vector.

Value

angle between a and b (in radians).

Examples

```
vector.angle(c(1,2,3),c(4,5,6))
```

vector.cross

This function calculates cross product.

Description

Cross product is only defined for 3D vectors.

Usage

```
vector.cross(a, b)
```

Arguments

a first vector.b second vector.

26 vector.norm

Value

cross product.

Examples

```
vector.cross(c(1,2,3),c(3,4,5))
```

vector.dot

This function calculates dot product.

Description

This function calculates dot product.

Usage

```
vector.dot(a, b)
```

Arguments

a first vector.b second vector.

Value

dot product.

Examples

```
vector.dot(c(1,2,3,4),c(5,6,7,8))
```

vector.norm

This function returns length (norm) of the vector.

Description

This function returns length (norm) of the vector.

Usage

```
vector.norm(x)
```

Arguments

Χ

a vector.

Value

vector norm.

```
vector.norm(c(1,2,3))
```

```
vector.to.angles.frame.list
```

This function genereates list of angles between vector and coordinates frame.

Description

Vector is defined as v1=(f1-f2). The coordinate frame is:

- X=(xt1-xt2),
- $Z=X \times (0,1,0)$,
- Y=X x Z

All vectors are normalized.

Usage

```
vector.to.angles.frame.list(dataToCalculate, f1, f2, xt1, xt2)
```

Arguments

dataToCalculate

data frame with motion capture data.

- f1 name of the first joint of vector.
- f2 name of the second joint of vector.
- xt1 name of the first joint of horizontal axis of coordinate frame.
- xt2 name of the second joint of horizontal axis of coordinate frame.

Value

Results are list of vectors defined as follow: (angle(Y, v1), angle(Z, v1), angle(X, v1)).

Examples

```
data("heian.yondan")
vector.to.angles.frame.list(heian.yondan[1:3,], "RightArm", "RightForearm", "RightShoulder", "LeftShoulder")
```

 ${\tt vector.to.angles.list} \ \ \textit{This function generates list of angles}.$

Description

Angles are calculated on the plane between vectors v1=(f2-f1) and v2=(f2-f3).

Usage

```
vector.to.angles.list(dataToCalculate, f1, f2, f3)
```

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Arguments

dataToCalculate

data frame with motion capture data.

f1 name of the first joint.

f2 name of the second joint.

f3 name of the third joint.

Value

list of angles (in radians) defined as above.

Examples

```
data("heian.yondan")
vector.to.angles.list(heian.yondan[1:3,], "RightShoulder", "RightArm", "RightForearm")
```

vector.to.list

This function genereates list of vectors.

Description

Each list position equals c(ABC.Dx, ABC.Dy, ABC.Dz) where featuresName = ABC is a name of the column of data frame dataToCalculate.

Usage

```
vector.to.list(dataToCalculate, featuresName)
```

Arguments

 ${\tt dataToCalculate}$

data frame with motion capture data.

featuresName

column name of the motion capture feature. Column names should have names endings .Dx, .Dy, .Dz.

Value

list of vectors defined as above.

```
data("heian.yondan")
vector.to.list(heian.yondan[1:3,], "RightHand")
```

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vector.to.unit

This function performs vector normalizing.

Description

This function performs vector normalizing.

Usage

```
vector.to.unit(x)
```

Arguments

Χ

a vector.

Value

normalized vector x.

Examples

```
vector.to.unit(c(1,2,3,4,5))
```

wavg.quaternion.markley

Weighted quaternion Markley averaging algorithms.

Description

See: F. Landis Markley, Yang Cheng, John Lucas Crassidis, and Yaakov Oshman. "Averaging Quaternions", Journal of Guidance, Control, and Dynamics, Vol. 30, No. 4 (2007), pp. 1193-1197. https://doi.org/10.2514/1.28949

Usage

```
wavg.quaternion.markley(Q, weights)
```

Arguments

Q a data frame of quaternions (four dimensional vectors) to be averaged. Each row

of data frame holds one quaternion.

weights weights vector.

Value

4D quaternion vector.

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Examples

```
Q <- data.frame(c(0.9999986, 0.9999986, 0.9999986, 0.9999986, 0.9999986, 0.9999986), c(0.0008716584, 0.0008716584, 0.0009590162, 0.0009590162, 0.001046359, 0.001046359), c(0.0009608439, 0.001048034, 0.0008736689, 0.001048034, 0.0009608439, 0.0008736689), c(0.001046359, 0.0009590162, 0.001046359, 0.0008716584, 0.0008716584, 0.0009590162))  x <- seq(-2,2,length=6) \\ y <- dnorm(x,mean=0, sd=1) \\ y <- y / sum(y) \\ wavg.quaternion.markley(Q, y)
```

write.bvh

This function saves object of mocap class to the file.

Description

Function uses hierarchical model definition from hierarchy list however data of channels are taken from data frame.

Usage

```
write.bvh(skeleton.helper, path)
```

Arguments

```
\begin{tabular}{ll} skeleton.helper & mocap object to be save. \\ path & path to the file. \\ \end{tabular}
```

```
data("header.mocap")
data("heian.shodan")
heian.shodan.corrected <- calculate.kinematic(heian.shodan, show.plot = "TRUE", plot.title = "Heian Shodan")
original.bvh <- set.data.frame(header.mocap, heian.shodan)
corrected.bvh <- set.data.frame(header.mocap, heian.shodan.corrected)
#plotting BVH
plot(original.bvh, frames.fraction = 0.1, my.color = "red", alpha = 0.1, spheres = FALSE)
plot(corrected.bvh, frames.fraction = 0.1, my.color = "green", alpha = 0.1, spheres = FALSE, append = TRUE)
#writting BVH to disk
write.bvh(original.bvh, "original.bvh")
write.bvh(corrected.bvh, "corrected.bvh")</pre>
```

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