

# Package ‘Toothnroll’

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**Title** Dental Tissues Landmarking Measuring and Mapping

**Version** 1.11

**Description** Two- and three-dimensional morphometric maps of enamel and dentine thickness and multivariate analysis.

Volume calculation of dental materials.

Principal component analysis of thickness maps with associated morphometric map variations.

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Tooth2Dmap

*Tooth2Dmap***Description**

Create 2D morphometric maps of enamel/dentin thickness

**Usage**

```
Tooth2Dmap(
  tooth.shape,
  input,
  rem.out = FALSE,
  fac.out = 0.5,
  smooth = FALSE,
  scale = TRUE,
  smooth.iter = 5,
  gamMap = FALSE,
  nrow = 120,
  ncol = 80,
  gdl = 250,
  method = "equiangular",
  plot = TRUE,
  pal = blue2green2red(101),
  aspect = 0.6,
  labels = c("Li", "Mes", "Bu", "D", "Li"),
  ylab = "")
```

### Arguments

tooth.shape	list: output from the function ToothShape
input	list: output from the function ToothAlignment
rem.out	logical: if TRUE outliers will be removed
fac.out	numeric: parameter to set the threshold in outlier detection
smooth	logical: if TRUE a smoothing filter is applied
scale	logical: if TRUE the thickness matrix is scaled from 0 to 1
smooth.iter	numeric: number of smoothing iterations
gamMap	logical: if TRUE gam smoothing is applied
nrow	numeric: number of rows for gam smoothing matrix
ncol	numeric: number of columns for gam smoothing matrix
gdf	numeric: number of degree of freedom for gam smoothing matrix
method	character: if set on "equiangular" the dentine or enamel thickness is meant as the distance of the segment intersecting the external and internal outline starting from the centroid of the section. If set on "closest" the dentine or enamel thickness is calculated at each point as the closest distance between external and internal outlines
plot	logical: if TRUE the 2D morphometric map is plotted
pal	character vector: colors to be used in the map production
aspect	numeric: axis ratio for 2D morphometric map
labels	character vector: names for x labels in the morphometric map
ylab	character vector: label for y axis in the morphometric map

### Value

dataframe  
 dataframe for colormap production  
 2Dmap  
 thickness color map  
 gamoutput  
 output from GAM  
 data input  
 used to build the GAM map

### Author(s)

Antonio Profico; Mathilde Augoyard

### Examples

```

data("URI1_tooth")
require(morphomap)
Enamel<-URI1_tooth$mesh1
Dentin<-URI1_tooth$mesh2
Pulp<-URI1_tooth$mesh3
outline<-URI1_tooth$outline
  
```

```

set<-URI1_tooth$set
#Map of the crown
AlignMeshes<-ToothAlignment(mesh1=Enamel, mesh2=Dentin, mesh3=Pulp, set, outline, analyse = "c")
#Virtual sectioning dentine-pulp
External<-AlignMeshes$almesh1$mesh
Internal<-AlignMeshes$almesh2$mesh
#Define 16 cross-sections from the 30% to the 90% along the crown
Core<-ToothCore(External, Internal, num.points = 1000, num.sect = 16,
                  bio.len = AlignMeshes$length, start=0.3, end=0.9)
#Extract 25 equiangular semilandmarks from each cross-section (anticlockwise)
Shape<-ToothShape(Core, 25, sects_vector = NULL, cent.out = "E", direction="a")
Tooth2Dmap(Shape, AlignMeshes, rem.out = TRUE, scale=FALSE, smooth = FALSE, aspect = 0.5, gamMap = FALSE,
            nrow = 100, ncol = 100, gdl = 250, method="equiangular")

#Map of the root
AlignMeshes<-ToothAlignment(mesh1=Enamel, mesh2=Dentin, mesh3=Pulp, set, outline, analyse = "r")
#Virtual sectioning dentine-pulp
External<-AlignMeshes$almesh2$mesh
Internal<-AlignMeshes$almesh3$mesh
#Define 16 cross-sections from the 10% to the 50% along the root
Core<-ToothCore(External, Internal, num.points = 1000, num.sect = 16,
                  bio.len = AlignMeshes$length, start=0.1, end=0.5)
#Extract 25 equiangular semilandmarks from each cross-section (anticlockwise)
Shape<-ToothShape(Core, 25, sects_vector = NULL, cent.out = "E", direction="a")
Tooth2Dmap(Shape, AlignMeshes, rem.out = FALSE, scale=FALSE, smooth = FALSE, aspect = 0.5, gamMap = FALSE,
            nrow = 100, ncol = 100, gdl = 250, method="equiangular")

```

**ToothAlignment*****ToothAlignment*****Description**

Align dental meshes using as reference the cervical outline and five landmarks

**Usage**

```
ToothAlignment(mesh1, mesh2, mesh3 = NULL, set, outline, analyse = c("r", "c"))
```

**Arguments**

mesh1	3D mesh: dental mesh (enamel)
mesh2	3D mesh: dental mesh (dentin)
mesh3	3D mesh: dental mesh (dental pulp)
set	matrix: 5 landmarks acquired on the mesh (see details)
outline	matrix: set of coordinates along the cervical outline
analyse	character: "r" for root, "c" for crown

## Details

The function needs five landmarks to align the dental meshes. Usually landmarks from 1 to 4 define the Lingual, Mesial, Buccal and Distal margins. The fifth landmark defines the end of the z axis (biomechanical length). The centroid of the cervical outline defines the origin of axes.

## Value

- almesh1: mesh of the aligned mesh1
- almesh2: mesh of the aligned mesh2
- almesh3: mesh of the aligned mesh3
- alset: coordinates of the aligned landmark configuration
- length: biomechanical length of the root (see details)
- margins: coordinates of the landmarks in correspondence of the four margins
- margins\_sel: position of the margin along the outline
- aloutline: coordinates of the aligned cervical outline
- diamBL: bucco-lingual diameter
- diamMD: mesio-distal diameter

## Author(s)

Antonio Profico; Mathilde Augoyard

## Examples

```

data("URI1_tooth")
Enamel<-URI1_tooth$mesh1
Dentin<-URI1_tooth$mesh2
Pulp<-URI1_tooth$mesh3
outline<-URI1_tooth$outline
set<-URI1_tooth$set

#Example on the root
AlignMeshesR<-ToothAlignment(mesh1=Enamel,mesh2=Dentin,mesh3=Pulp,set,outline,analyse = "r")
require(rgl)
open3d()
shade3d(AlignMeshesR$almesh1$mesh,col="white",alpha=0.5)
shade3d(AlignMeshesR$almesh2$mesh,col="pink",alpha=0.5,add=TRUE)
shade3d(AlignMeshesR$almesh3$mesh,col="orange",alpha=0.5,add=TRUE)
spheres3d(AlignMeshesR$alset,radius=0.25)
spheres3d(AlignMeshesR$outline,radius=0.1,col="blue")
lines3d(AlignMeshesR$outline)
text3d(rbind(AlignMeshesR$outline[AlignMeshesR$margins_sel,],AlignMeshesR$alset[4,]),
texts=1:5,cex=4)
spheres3d(AlignMeshesR$alset[2,],radius=0.3,col="red")
arrow3d(colMeans(AlignMeshesR$aloutline),AlignMeshesR$alset[4,],lwd=3,col="green",
type="lines",s=1/10)
lines3d(rbind(AlignMeshesR$alset[2,],AlignMeshesR$alset[1,]),lwd=3,col="green")

```

```

axes3d()

#Example on the crown

AlignMeshesC<-ToothAlignment(mesh1=Enamel,mesh2=Dentin,mesh3=Pulp,set,outline,
analyse = "c")
require(rgl)
open3d()
shade3d(AlignMeshesC$almesh1$mesh,col="white",alpha=0.5)
shade3d(AlignMeshesC$almesh2$mesh,col="pink",alpha=0.5,add=TRUE)
shade3d(AlignMeshesC$almesh3$mesh,col="orange",alpha=0.5,add=TRUE)
spheres3d(AlignMeshesC$alset, radius=0.25)
spheres3d(AlignMeshesC$outline, radius=0.1, col="blue")
lines3d(AlignMeshesC$outline)
text3d(rbind(AlignMeshesC$outline[AlignMeshesC$margin_sel,],AlignMeshesC$alset[4,]),
texts=1:5,cex=4)
spheres3d(AlignMeshesC$alset[2,],radius=0.3,col="red")
arrow3d(colMeans(AlignMeshesC$outline),AlignMeshesC$alset[4,],lwd=3,col="green",
type="lines",s=1/10)
lines3d(rbind(AlignMeshesC$alset[2,],AlignMeshesC$alset[1,]),lwd=3,col="green")
lines3d(rbind(AlignMeshesC$alset[2,],AlignMeshesC$alset[1,]),lwd=3,col="green")

axes3d()

```

## Description

Tool to build 3D and 2D cross sections

## Usage

```

ToothCore(
  out.sur = out.sur,
  inn.sur = inn.sur,
  num.sect = 61,
  bio.len,
  clean_int_out_0 = TRUE,
  clean_int_out_I = TRUE,
  param1_out = 0.5,
  radius факт_out = 2.5,
  param1_inn = 0.5,
  radius факт_inn = 2.5,
  npovs = 100,
  num.points = 500,
  start = 0.2,
  end = 0.8,

```

```

    curved = FALSE,
    print.progress = TRUE
)

```

### Arguments

out.sur	object of class mesh3d
inn.sur	object of class mesh3d
num.sect	number of sections
bio.len	length of the selected region of interest along with extracting the digital section
clean_int_out_0	logical if TRUE the outer section will be cleaned by using spherical flipping
clean_int_out_I	logical if TRUE the inner section will be cleaned by using spherical flipping
param1_out	numeric parameter for clean_int_out_O spherical flipping operator (how much the section will be deformed)
radius факт_out	logical if TRUE the inner section will be cleaned by using spherical flipping
param1_inn	numeric parameter for clean_int_out_I spherical flipping operator (how much the section will be deformed)
radius факт_inn	logical if TRUE the inner section will be cleaned by using spherical flipping
npovs	numeric: number of points of view defined around the section
num.points	number of equiangular points to be defined on each section
start	percentage of the mechanical length from which the first section is defined
end	percentage of the mechanical length from which the last section is defined
curved	logical: if TRUE the cutting planes will follow the mesh curvature (beta version)
print.progress	logical: if TRUE a progress bar is printed to the screen

### Value

out3D	num.pointsx3xnum.sect array of the external outlines
inn3D	num.pointsx3xnum.sect array of the internal outlines
out3D	num.pointsx2xnum.sect array of the external outlines
inn3D	num.pointsx2xnum.sect array of the internal outlines
mech_length	mechanical length of the long bone
start	percentage of the mechanical length from which the first section is defined
end	percentage of the mechanical length from which the last section is defined

### Author(s)

Antonio Profico; Mathilde Augoyard

## Examples

```

data("URI1_tooth")
require(morphomap)
require(Morpho)
require(rgl)
Enamel<-URI1_tooth$mesh1
Dentin<-URI1_tooth$mesh2
Pulp<-URI1_tooth$mesh3
outline<-URI1_tooth$outline
set<-URI1_tooth$set

#Unrolling the crown
AlignMeshes<-ToothAlignment(mesh1=Enamel,mesh2=Dentin,mesh3=Pulp,set,outline,analyse = "c")
#Virtual sectioning dentine-pulp
External<-AlignMeshes$almesh1$mesh
Internal<-AlignMeshes$almesh2$mesh
#Define 16 cross-sections from the 30% to the 90% along the crown
Core<-ToothCore(External,Internal,num.points = 1000,num.sect =16,
                 bio.len = AlignMeshes$length,start=0.3,end=0.9)

plot3d(morphomapArray2matrix(Core$"out3D"),aspect=FALSE,xlab="x",ylab="y",zlab="z")
plot3d(morphomapArray2matrix(Core$"inn3D"),aspect=FALSE,add=TRUE)
wire3d(AlignMeshes$almesh2$mesh,col="white")
wire3d(AlignMeshes$almesh1$mesh,col="violet")

#Unrolling the root
AlignMeshes<-ToothAlignment(mesh1=Enamel,mesh2=Dentin,mesh3=Pulp,set,outline,analyse = "r")
#Virtual sectioning dentine-pulp
External<-AlignMeshes$almesh2$mesh
Internal<-AlignMeshes$almesh3$mesh
#Define 16 cross-sections from the 10% to the 50% along the root
Core<-ToothCore(External,Internal,num.points = 1000,num.sect =16,
                 bio.len = AlignMeshes$length,start=0.1,end=0.5)

plot3d(morphomapArray2matrix(Core$"out3D"),aspect=FALSE,xlab="x",ylab="y",zlab="z")
plot3d(morphomapArray2matrix(Core$"inn3D"),aspect=FALSE,add=TRUE)
wire3d(AlignMeshes$almesh3$mesh,col="red")
wire3d(AlignMeshes$almesh2$mesh,col="lightblue")
wire3d(AlignMeshes$almesh1$mesh,col="white")

```

## Description

Tool to build a data.frame suitable for morphometric maps

**Usage**

```
ToothDF(
  tooth.thickness,
  rem.out = TRUE,
  fac.out = 0.5,
  smooth = TRUE,
  scale = TRUE,
  smooth.iter = 5,
  method = "equiangular",
  labels = c("Li", "Mes", "Bu", "D", "Li"),
  relThick = FALSE
)
```

**Arguments**

tooth.thickness	list: tooth.Thickness object
rem.out	logical: if TRUE the outlier will be removed
fac.out	numeric: parameter to set the threshold in outliers detection
smooth	logical: if TRUE the smooth algorithm is applied
scale	logical: if TRUE the thickness matrix is scaled from 0 to 1
smooth.iter	numeric: number of smoothing iterations
method	character: if set on "equiangular" the dental thickness is meant as the distance of the segment intersecting the external and internal outline starting from the centroid of the section. If set on "closest" the dental thickness is calculated at each point as the closest distance between external and internal outlines
labels	character vector: names for x labels in the morphometric map
relThick	logical: if TRUE the dental thickness is converted into relative dental thickness

**Value**

XYZ data.frame for morphometric map

labels character vector for x labels in the morphometric map

**Author(s)**

Antonio Profico; Mathilde Augoyard

---

ToothExport

*ToothExport*

---

### Description

Export the output from ToothAlignment

### Usage

`ToothExport(input, id, file)`

### Arguments

input	list: output from ToothAlignment
id	character: label name
file	character: name the output file

### Value

no return value, called for side effects (see description)

### Author(s)

Antonio Profico; Mathilde Augoyard

---

ToothImport

*ToothImport*

---

### Description

Import the output from ToothAlignment

### Usage

`ToothImport(file)`

### Arguments

file	character: name of input file
------	-------------------------------

**Value**

Mesh1: mesh of the aligned mesh1  
Mesh2: mesh of the aligned mesh2  
Mesh3: mesh of the aligned mesh3  
B.Length: length of the region of interest  
Landmarks: landmark coordinates  
Outline: outline coordinates  
Margins: position of the margins along the outline  
diamBL: bucco-lingual diameter  
diamMD: mesio-distal diameter

**Author(s)**

Antonio Profico; Mathilde Augoyard

---

ToothPCA

*ToothPCA*

---

**Description**

Perform the Principal Component Analysis on a list of tooth.shape objects

**Usage**

```
ToothPCA(  
  mpShapeList,  
  gamMap = FALSE,  
  nrow = 120,  
  ncol = 80,  
  gdl = 250,  
  rem.out = TRUE,  
  scaleThick = FALSE,  
  relThick = FALSE,  
  fac.out = 1.5,  
  method = "equiangular",  
  scalePCA = TRUE  
)
```

**Arguments**

mpShapeList	list: tooth.shape objects
gamMap	logical: if TRUE gamMap spline method is applied
nrow	numeric: number of rows if gamMap is TRUE

ncol	numeric: number of columns if gamMap is TRUE
gdl	numeric: degree of freedom (if gamMap is TRUE)
rem.out	logical: if TRUE outliers are removed
scaleThick	logical: if TRUE thickness values are scaled from 0 to 1
relThick	logical: if TRUE the thickness values are scaled by the diameter from the centroid to the external outline
fac.out	numeric: threshold to define an outlier observation
method	character: "equiangular" or "closest" to define the thickness from evenly spaced or closest semilandmarks between the external and internal outline
scalePCA	logical: indicate whether the variables should be scaled to have unit variance

### Value

PCscores matrix of PC scores  
 PCs principal components  
 variance table of the explained Variance by the PCs  
 meanMap mean map  
 CorMaps maps of thickness used as input in the PCA

### Author(s)

Antonio Profico; Mathilde Augoyard

### Examples

```

### Example on the canine crown
data("UCcrown")
require(morphomap)
shapeList<-UCcrown
PCA<-ToothPCA(shapeList,gamMap = FALSE,scaleThick = TRUE,scalePCA = TRUE ,relThick = FALSE)

#gamMap set on TRUE
PCA<-ToothPCA(shapeList,gamMap = TRUE,scaleThick = TRUE,scalePCA = TRUE,relThick = FALSE)

otu<-substr(names(shapeList),1,2)
pchs <- ifelse(otu == "MH", 16, 17)
cols <- ifelse(otu == "MH", "orange", "darkblue")

plot(PCA$PCscores,col=cols,cex=1, pch = pchs,
      xlab=paste("PC1 (",round(PCA$Variance[1,2],2),"%)",),
      ylab=paste("PC2 (",round(PCA$Variance[2,2],2),"%)",),
      cex.lab=1,cex.axis=1)
title (main="UC (radicular dentine)", font.main= 1,adj = 0, cex.main = 1.2)
legend("topright", legend = c("MH", "NE"), col = c("orange", "darkblue"), pch = c(16,17), cex = 0.8)
abline(v=0,h=0,col="black",lwd=2,lty=3)

hpts1 <- chull(PCA$PCscores[which(otu=="MH"),1:2])
hpts1 <- c(hpts1, hpts1[1])

```

```

polygon(PCA$PCscores[which(otu=="MH")[hpts1],1:2 ], col = adjustcolor("orange", 0.3), border = NA)

hpts2 <- chull(PCA$PCscores[which(otu=="NE"),c(1:2)])
hpts2 <- c(hpts2, hpts2[1])
polygon(PCA$PCscores[which(otu=="NE")[hpts2], ], col = adjustcolor("darkblue", 0.3), border = NA)

PC1min<-ToothVariations(PCA,min(PCA$PCscores[,1]),PCA$PCs[,1],asp=0.5,meanmap = FALSE)
PC1max<-ToothVariations(PCA,max(PCA$PCscores[,1]),PCA$PCs[,1],asp=0.5,meanmap = FALSE)

### Example on the canine root
data("UCroot")
require(morphomap)
shapeList<-UCroot
PCA<-ToothPCA(shapeList,gamMap = FALSE,scaleThick = TRUE,scalePCA = TRUE,relThick = FALSE)
otu<-substr(names(shapeList),1,2)
pchs <- ifelse(otu == "MH", 16, 17)
cols <- ifelse(otu == "MH", "orange", "darkblue")

plot(PCA$PCscores,col=cols,cex=1, pch = pchs,
      xlab=paste("PC1 (",round(PCA$Variance[1,2],2),"%)",),
      ylab=paste("PC2 (",round(PCA$Variance[2,2],2),"%)",),
      cex.lab=1,cex.axis=1)
title (main="UC (radicular dentine)", font.main= 1,adj = 0, cex.main = 1.2)
legend("topright", legend = c("MH", "NE"), col = c("orange", "darkblue"), pch = c(16,17), cex = 0.8)
abline(v=0,h=0,col="black",lwd=2,lty=3)

hpts1 <- chull(PCA$PCscores[which(otu=="MH"),1:2])
hpts1 <- c(hpts1, hpts1[1])
polygon(PCA$PCscores[which(otu=="MH")[hpts1],1:2 ], col = adjustcolor("orange", 0.3), border = NA)

hpts2 <- chull(PCA$PCscores[which(otu=="NE"),c(1:2)])
hpts2 <- c(hpts2, hpts2[1])
polygon(PCA$PCscores[which(otu=="NE")[hpts2], ], col = adjustcolor("darkblue", 0.3), border = NA)

PC1min<-ToothVariations(PCA,min(PCA$PCscores[,1]),PCA$PCs[,1],asp=0.5,meanmap = FALSE)
PC1max<-ToothVariations(PCA,max(PCA$PCscores[,1]),PCA$PCs[,1],asp=0.5,meanmap = FALSE)

### Example on the central upper incisor (crown)
data("UI1crown")
require(morphomap)
shapeList<-UI1crown
PCA<-ToothPCA(shapeList,gamMap = FALSE,scaleThick = TRUE,scalePCA = TRUE ,relThick = FALSE)
otu<-substr(names(shapeList),1,2)
pchs <- ifelse(otu == "MH", 16, 17)
cols <- ifelse(otu == "MH", "orange", "darkblue")

plot(PCA$PCscores,col=cols,cex=1, pch = pchs,
      xlab=paste("PC1 (",round(PCA$Variance[1,2],2),"%)",),
      ylab=paste("PC2 (",round(PCA$Variance[2,2],2),"%)",),
      cex.lab=1,cex.axis=1)
title (main="UC (radicular dentine)", font.main= 1,adj = 0, cex.main = 1.2)
legend("topright", legend = c("MH", "NE"), col = c("orange", "darkblue"), pch = c(16,17), cex = 0.8)
abline(v=0,h=0,col="black",lwd=2,lty=3)

```

```

hpts1 <- chull(PCA$PCscores[which(otu=="MH"),1:2])
hpts1 <- c(hpts1, hpts1[1])
polygon(PCA$PCscores[which(otu=="MH")[hpts1],1:2 ], col = adjustcolor("orange", 0.3), border = NA)

hpts2 <- chull(PCA$PCscores[which(otu=="NE"),c(1:2)])
hpts2 <- c(hpts2, hpts2[1])
polygon(PCA$PCscores[which(otu=="NE")[hpts2], ], col = adjustcolor("darkblue", 0.3), border = NA)

PC1min<-ToothVariations(PCA,min(PCA$PCscores[,1]),PCA$PCs[,1],asp=0.5,meanmap = FALSE)
PC1max<-ToothVariations(PCA,max(PCA$PCscores[,1]),PCA$PCs[,1],asp=0.5,meanmap = FALSE)

### Example on the upper central incisor (root)
data("UI1root")
require(morphomap)
shapeList<-UI1root
PCA<-ToothPCA(shapeList,gamMap = FALSE,scaleThick = TRUE,scalePCA = TRUE,relThick = FALSE)
otu<-substr(names(UI1root),1,2)
pchs <- ifelse(otu == "MH", 16, 17)
cols <- ifelse(otu == "MH", "orange", "darkblue")

plot(PCA$PCscores,col=cols,cex=1, pch = pchs,
      xlab=paste("PC1 (",round(PCA$Variance[1,2],2),"%)","),
      ylab=paste("PC2 (",round(PCA$Variance[2,2],2),"%)","),
      cex.lab=1,cex.axis=1)
title (main="UC (radicular dentine)", font.main= 1,adj = 0, cex.main = 1.2)
legend("topright", legend = c("MH", "NE"), col = c("orange", "darkblue"), pch = c(16,17), cex = 0.8)
abline(v=0,h=0,col="black",lwd=2,lty=3)

hpts1 <- chull(PCA$PCscores[which(otu=="MH"),1:2])
hpts1 <- c(hpts1, hpts1[1])
polygon(PCA$PCscores[which(otu=="MH")[hpts1],1:2 ], col = adjustcolor("orange", 0.3), border = NA)

hpts2 <- chull(PCA$PCscores[which(otu=="NE"),c(1:2)])
hpts2 <- c(hpts2, hpts2[1])
polygon(PCA$PCscores[which(otu=="NE")[hpts2], ], col = adjustcolor("darkblue", 0.3), border = NA)

PC1min<-ToothVariations(PCA,min(PCA$PCscores[,1]),PCA$PCs[,1],asp=0.5,meanmap = FALSE)
PC1max<-ToothVariations(PCA,max(PCA$PCscores[,1]),PCA$PCs[,1],asp=0.5,meanmap = FALSE)

### Example on the lateral upper incisor (crown)
data("UI2crown")
require(morphomap)
shapeList<-UI2crown
PCA<-ToothPCA(shapeList,gamMap = FALSE,scaleThick = TRUE,scalePCA = TRUE ,relThick = FALSE)
otu<-substr(names(shapeList),1,2)
pchs <- ifelse(otu == "MH", 16, 17)
cols <- ifelse(otu == "MH", "orange", "darkblue")

plot(PCA$PCscores,col=cols,cex=1, pch = pchs,
      xlab=paste("PC1 (",round(PCA$Variance[1,2],2),"%)","),
      ylab=paste("PC2 (",round(PCA$Variance[2,2],2),"%)","),
      cex.lab=1,cex.axis=1)

```

```

title (main="UC (radicular dentine)", font.main= 1,adj = 0, cex.main = 1.2)
legend("topright", legend = c("MH", "NE"), col = c("orange", "darkblue"), pch = c(16,17), cex = 0.8)
abline(v=0,h=0,col="black",lwd=2,lty=3)

hpts1 <- chull(PCA$PCscores[which(otu=="MH"),1:2])
hpts1 <- c(hpts1, hpts1[1])
polygon(PCA$PCscores[which(otu=="MH")][hpts1],1:2 ], col = adjustcolor("orange", 0.3), border = NA)

hpts2 <- chull(PCA$PCscores[which(otu=="NE"),c(1:2)])
hpts2 <- c(hpts2, hpts2[1])
polygon(PCA$PCscores[which(otu=="NE")][hpts2], ], col = adjustcolor("darkblue", 0.3), border = NA)

PC1min<-ToothVariations(PCA,min(PCA$PCscores[,1]),PCA$PCs[,1],asp=0.5,meanmap = FALSE)
PC1max<-ToothVariations(PCA,max(PCA$PCscores[,1]),PCA$PCs[,1],asp=0.5,meanmap = FALSE)

### Example on the upper lateral incisor (root)
data("UI2root")
require(morphomap)
shapeList<-UI2root
PCA<-ToothPCA(shapeList,gamMap = FALSE,scaleThick = TRUE,scalePCA = TRUE,relThick = FALSE)
otu<-substr(names(UI2root),1,2)
pchs <- ifelse(otu == "MH", 16, 17)
cols <- ifelse(otu == "MH", "orange", "darkblue")

plot(PCA$PCscores,col=cols,cex=1, pch = pchs,
      xlab=paste("PC1 (",round(PCA$Variance[1,2],2),"%)"),
      ylab=paste("PC2 (",round(PCA$Variance[2,2],2),"%)"),
      cex.lab=1,cex.axis=1)
title (main="UC (radicular dentine)", font.main= 1,adj = 0, cex.main = 1.2)
legend("topright", legend = c("MH", "NE"), col = c("orange", "darkblue"), pch = c(16,17), cex = 0.8)
abline(v=0,h=0,col="black",lwd=2,lty=3)

hpts1 <- chull(PCA$PCscores[which(otu=="MH"),1:2])
hpts1 <- c(hpts1, hpts1[1])
polygon(PCA$PCscores[which(otu=="MH")][hpts1],1:2 ], col = adjustcolor("orange", 0.3), border = NA)

hpts2 <- chull(PCA$PCscores[which(otu=="NE"),c(1:2)])
hpts2 <- c(hpts2, hpts2[1])
polygon(PCA$PCscores[which(otu=="NE")][hpts2], ], col = adjustcolor("darkblue", 0.3), border = NA)

PC1min<-ToothVariations(PCA,min(PCA$PCscores[,1]),PCA$PCs[,1],asp=0.5,meanmap = FALSE)
PC1max<-ToothVariations(PCA,max(PCA$PCscores[,1]),PCA$PCs[,1],asp=0.5,meanmap = FALSE)

```

**Description**

Finds equiangular landmarks

**Usage**

```
ToothRegradius(mat, center, n, direction = c("c", "a"))
```

**Arguments**

mat	a kx2 matrix
center	coordinates of the center from which the calculation of regular radius started
n	number of points
direction	specify direction: "c"=clockwise; "a"=anticlockwise

**Value**

V2 position of landmarks equiangular spaced

**Author(s)**

Antonio Profico; Mathilde Augoyard

ToothShape

*ToothShape*

**Description**

Tool for the extraction of equiangular landmarks on the entire dental region of interest

**Usage**

```
ToothShape(
  tooth.core,
  num.land,
  sects_vector,
  cent.out = "E",
  delta = 0.1,
  direction = "c",
  out.sur = NULL,
  inn.sur = NULL
)
```

**Arguments**

tooth.core	list: tooth.core object
num.land	numeric: number of landmarks defining each section
sects_vector	numeric: number of sections
cent.out	how to define the center of each section. The method allowed are "CCA" (center of cortical area), "E" (barycenter of the external outline) and "I" (barycenter of the internal outline)

delta	pixel size used to calculate the CCA
direction	clockwise if "c", anticlockwise if "a"
out.sur	mesh: if provided, the external outlines will be projected back on the surface
inn.sur	mesh: if provided, the internal outlines will be projected back on the surface

**Value**

out3D num.pointsx3xnum.sect array in which the external outlines are stored  
 inn3D num.pointsx3xnum.sect array in which the internal outlines are stored  
 out2D num.pointsx2xnum.sect array in which the external outlines are stored  
 inn2D num.pointsx2xnum.sect array in which the internal outlines are stored  
 ALPM\_inn array with the coordinates of ALPM coordinates on the external outline  
 ALPM\_out array with the coordinates of ALPM coordinates on the internal outline  
 mech\_length length of the selected region of interest  
 start percentage of the mechanical length from which the first section is defined  
 end percentage of the mechanical length from which the last section is defined  
 centroids of the cross-sections

**Author(s)**

Antonio Profico; Mathilde Augoyard

**Examples**

```

data("URI1_tooth")
require(morphomap)
require(rgl)
Enamel<-URI1_tooth$mesh1
Dentin<-URI1_tooth$mesh2
Pulp<-URI1_tooth$mesh3
outline<-URI1_tooth$outline
set<-URI1_tooth$set

#Unrolling the crown
AlignMeshes<-ToothAlignment(mesh1=Enamel,mesh2=Dentin,mesh3=Pulp,set,outline,analyse = "c")
#Virtual sectioning dentine-pulp
External<-AlignMeshes$almesh1$mesh
Internal<-AlignMeshes$almesh2$mesh
#Define 16 cross-sections from the 30% to the 90% along the crown
Core<-ToothCore(External,Internal,num.points = 1000,num.sect =16,
                 bio.len = AlignMeshes$length,start=0.3,end=0.9)
Shape<-ToothShape(Core,num.land = 21,sects_vector = NULL,direction = "a")

plot3d(morphomapArray2matrix(Shape$"out3D"),type="s",radius = 0.1,aspect=FALSE,
       xlab="x",ylab="y",zlab="z")
plot3d(morphomapArray2matrix(Shape$"inn3D"),type="s",radius = 0.1,aspect=FALSE,

```

```

add=TRUE)
wire3d(AlignMeshes$almesh2$mesh,col="white")
wire3d(AlignMeshes$almesh1$mesh,col="violet")

#Unrolling the root
AlignMeshes<-ToothAlignment(mesh1=Enamel,mesh2=Dentin,mesh3=Pulp,set.outline,
analyse = "r")
#Virtual sectioning dentine-pulp
External<-AlignMeshes$almesh2$mesh
Internal<-AlignMeshes$almesh3$mesh
#Define 16 cross-sections from the 10% to the 50% along the root
Core<-ToothCore(External,Internal,num.points = 1000,num.sect =16,
                 bio.len = AlignMeshes$length,start=0.1,end=0.5)
Shape<-ToothShape(Core,num.land = 21,sects_vector = NULL,direction = "a")

plot3d(morphomapArray2matrix(Shape$"out3D"),type="s",radius = 0.1,aspect=FALSE,
xlab="x",ylab="y",zlab="z")
plot3d(morphomapArray2matrix(Shape$"inn3D"),type="s",radius = 0.1,aspect=FALSE,
add=TRUE)
wire3d(AlignMeshes$almesh3$mesh,col="red")
wire3d(AlignMeshes$almesh2$mesh,col="lightblue")
wire3d(AlignMeshes$almesh1$mesh,col="white")

```

ToothThickness

*ToothThickness*

## Description

Tool for the extraction of equiangular landmarks on the selected dental mesh

## Usage

```
ToothThickness(tooth.shape)
```

## Arguments

`tooth.shape`      list: tooth.shape object

## Value

`sect_thickness_eq` dental thickness (equiangular method)  
`sect_thickness_cp` dental thickness (closest point method)  
`sect_Rthickness_eq` relative dental thickness (equiangular method)  
`sect_Rthickness_cp` relative dental thickness (closest point method)  
`ALPM_thickness` dental thickness at ALPM quadrants  
`tooth.shape` tooth.shape object

**Author(s)**

Antonio Profico; Mathilde Augoyard

---

ToothVariations      *ToothVariations*

---

**Description**

Calculate and return morphometric map variation

**Usage**

```
ToothVariations(  
  PCA,  
  scores,  
  PC,  
  asp = 2,  
  pal = blue2green2red(101),  
  meanmap = TRUE  
)
```

**Arguments**

PCA	list: output from ToothPCA
scores	numeric: principal component value
PC	numeric: principal component eigenvalue
asp	numeric: x,y ratio of the morphometric map
pal	character vector: vector of colors
meanmap	logical: if TRUE the mean map corresponds to the real mean morphometric maps, otherwise the mean map is 0

**Value**

XYZ data.frame of morphometric map variation

**Author(s)**

Antonio Profico; Mathilde Augoyard

**ToothVolumes***ToothVolumes***Description**

Extract volumes from the object `ToothShape`

**Usage**

```
>ToothVolumes(
  ShapeExt,
  ShapeInn,
  col1 = "gray",
  col2 = "red",
  col3 = "green",
  alpha1 = 1,
  alpha2 = 1,
  alpha3 = 1,
  plot = FALSE
)
```

**Arguments**

<code>ShapeExt</code>	3D mesh: external mesh
<code>ShapeInn</code>	3D mesh: internal mesh
<code>col1</code>	color of the <code>ShapeExt</code>
<code>col2</code>	color of the <code>ShapeInn</code>
<code>col3</code>	color of the boolean operation between <code>ShapeExt</code> and <code>ShapeInn</code>
<code>alpha1</code>	value to set transparency of <code>col1</code>
<code>alpha2</code>	value to set transparency of <code>col1</code>
<code>alpha3</code>	value to set transparency of <code>col1</code>
<code>plot</code>	logical: if TRUE the volumes are shown

**Value**

`meshOut`: external selected mesh  
`meshInnT`: internal selected mesh  
`meshDiff`: differences between selected meshes  
`volumeT`: volume of the external mesh  
`volInn`: volume of the internal mesh  
`volDiff`: difference between `volumeT` and `volInn`

**Author(s)**

Antonio Profico; Mathilde Augoyard

**Examples**

```

data("URI1_tooth")
require(morphomap)
Enamel<-URI1_tooth$mesh1
Dentin<-URI1_tooth$mesh2
Pulp<-URI1_tooth$mesh3
outline<-URI1_tooth$outline
set<-URI1_tooth$set

#Unrolling the crown
AlignMeshes<-ToothAlignment(mesh1=Enamel,mesh2=Dentin,mesh3=Pulp,set,outline,analyse = "c")
#Virtual sectioning dentine-pulp
External<-AlignMeshes$almesh1$mesh
Internal<-AlignMeshes$almesh2$mesh
#Define 16 cross-sections from the 30% to the 90% along the crown
Core<-ToothCore(External,Internal,num.points = 1000,num.sect =16,
                 bio.len = AlignMeshes$length,start=0.3,end=0.9)
Shape<-ToothShape(Core,num.land = 100,sects_vector = NULL,direction = "a")
volumes<-ToothVolumes(Shape$"out3D",Shape$"inn3D",plot=TRUE)
unlist(volumes[4:6])

```

**Description**

list of ToothShape (modern human and neanderthal upper canines)

**Usage**

```
data(UCcrown)
```

**Author(s)**

Antonio Profico, Mathilde Augoyard

---

UCroot

*example dataset*

---

**Description**

list of ToothShape (modern human and neanderthal upper canines)

**Usage**

```
data(UCroot)
```

**Author(s)**

Antonio Profico, Mathilde Augoyard

---

UI1crown

*example dataset*

---

**Description**

list of ToothShape (modern human and neanderthal upper central incisors)

**Usage**

```
data(UI1crown)
```

**Author(s)**

Antonio Profico, Mathilde Augoyard

---

UI1root

*example dataset*

---

**Description**

list of ToothShape (modern human and neanderthal upper central incisors)

**Usage**

```
data(UI1root)
```

**Author(s)**

Antonio Profico, Mathilde Augoyard

---

UI2crown

*example dataset*

---

**Description**

list of ToothShape (modern human and neanderthal upper lateral incisors)

**Usage**

```
data(UI2crown)
```

**Author(s)**

Antonio Profico, Mathilde Augoyard

---

UI2root

*example dataset*

---

**Description**

list of ToothShape (modern human and neanderthal upper lateral incisors)

**Usage**

```
data(UI2root)
```

**Author(s)**

Antonio Profico, Mathilde Augoyard

---

URI1\_tooth

*example dataset*

---

**Description**

3D dental meshes of a modern human upper right central incisor

**Usage**

```
data(URI1_tooth)
```

**Author(s)**

Antonio Profico, Mathilde Augoyard

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