

# Package ‘BoneProfileR’

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**Type** Package

**Title** Tools to Study Bone Compactness

**Version** 2.3

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**Depends** imager, HelpersMG (>= 5.3-1), knitr, rmarkdown, R (>= 4.1), shiny

**Suggests** tiff, ijtiff, openxlsx, parallel, scatterplot3d

**Description** Bone Profiler is a scientific method and a software used to model bone section for paleontological and ecological studies. See Girondot and Laurin (2003) <[https://www.researchgate.net/publication/280021178\\_Bone\\_profiler\\_A\\_tool\\_to\\_quantify\\_model\\_and\\_statistically\\_compare\\_bone-section\\_compactness\\_profiles](https://www.researchgate.net/publication/280021178_Bone_profiler_A_tool_to_quantify_model_and_statistically_compare_bone-section_compactness_profiles)> and Gônet, Laurin and Girondot (2022) <<https://palaeo-electronica.org/content/2022/3590-bone-section-compactness-model>>.

**License** GPL-2

**LazyData** yes

**LazyLoad** yes

**Encoding** UTF-8

**RoxygenNote** 7.2.0

**NeedsCompilation** no

**Repository** CRAN

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## R topics documented:

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BoneProfileR-package    *A Model for Bone Compactness.*

---

## Description

A Model for Bone Compactness.

The lastest version of this package can always been installed using:

```
install.packages(c("imager", "tiff", "ijtiff", "HelpersMG", "knitr", "rmarkdown", "openxlsx", "shiny"))
install.packages("https://hebergement.universite-paris-saclay.fr/marcgirondot/CRAN/HelpersMG.tar.gz",
repos=NULL, type="source")
install.packages("https://hebergement.universite-paris-saclay.fr/marcgirondot/CRAN/BoneProfileR.tar.gz",
repos=NULL, type="source")
```

BoneProfileR uses a new results management software that is developed as part of the HelpersMG package. Using this results management system (RM), all the results are stored as part of the analyzed image.

This results management software has been developed to help users to maintain the results associated with the methodology used to obtain it. It is part of the large movement in science of replicative research.

An analysis is then stored with the image in a single file with the following information:  
name, timestamp, bg, fg, threshold, contour, centers, peripherie, compactness, array.compactness,  
cut.distance.center, cut.angle, used.centers, compactness.synthesis, partial, rotation.angle, global.compactness,  
optim, optimRadial

Several analyses can be stored within a single file.



## Details

A model for bone compactness.

Package:	BoneProfileR
Type:	Package
Version:	2.3 build 762
Date:	2022-07-14
License:	GPL (>= 2)
LazyLoad:	yes

## Author(s)

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

## References

Girondot M, Laurin M (2003) Bone Profiler: a tool to quantify, model, and statistically compare bone-section compactness profiles. Journal of Vertebrate Paleontology 23: 458-461

Laurin M, Girondot M, Loth M-M (2004) The evolution of long bone microstructure and lifestyle in lissamphibians. *Paleobiology* 30: 589-613

Gônet, Jordan, Jérémie Bardin, Marc Girondot, John R. Hutchinson, and Michel Laurin. 2022. The Reptilian Locomotor and Postural Diversity Seen through the Prism of Femoral Microanatomy: Paleobiological Implications for Some Permian and Mesozoic Reptiles. Submitted

Gônet, Jordan, Michel Laurin, and Marc Girondot. 2022. “Bone Profiler: The Next Step to Quantify, Model and Statistically Compare Bone Section Compactness Profiles.” *Paleontologica Electronica*. 25(1): a12

## Examples

```
## Not run:
# Not run:
library(BoneProfileR)
path_Hedgehog <- system.file("extdata", "Erinaceus_europaeus_fem_2-1_small.png",
                             package = "BoneProfileR")
bone <- BP_OpenImage(file=path_Hedgehog)
plot(bone, type="original")
bone <- BP_DetectBackground(bone=bone, analysis="logistic")
bone <- BP_DetectForeground(bone=bone, analysis="logistic")
bone <- BP_DetectCenters(bone=bone, analysis="logistic")
plot(bone, type="original")
plot(bone, type="mineralized")
plot(bone, type="unmineralized")
plot(bone, type="section")
plot(bone, type="colors")
plot(bone, type="3Dcolors")
bone <- BP_EstimateCompactness(bone, analysis="logistic", center="ontogenetic")
plot(bone, type="original")
plot(bone, type="mineralized")
plot(bone, type="observations")
bone <- BP_FitMLCompactness(bone, analysis="logistic")
plot(bone, type="model", analysis=1)
plot(bone, type="observations+model", analysis=1)
fittedpar <- BP_GetFittedParameters(bone, analysis="logistic")
bone <- BP_DuplicateAnalysis(bone, from="logistic", to="flexit")
bone <- BP_FitMLCompactness(bone,
                            fitted.parameters=c(fittedpar, K1=1, K2=1),
                            fixed.parameters=NULL, analysis="flexit")
compare_AIC(Logistic=BP_GetFittedParameters(bone, analysis="logistic", alloptim=TRUE),
            Flexit=BP_GetFittedParameters(bone, analysis="flexit", alloptim=TRUE))
# pdf(file = "Figure 2.pdf", width = 8, height = 10, pointsize = 12)
layout(1:2)
plot(bone, type="observations+model", analysis="logistic", restorePar=FALSE, mar=c(4, 4, 2, 5))
plot(bone, type="observations+model", analysis="flexit", restorePar=FALSE, mar=c(4, 4, 2, 5))
layout(1)
# dev.off()
out4p <- plot(bone, type="observations+model", analysis="logistic")
out6p <- plot(bone, type="observations+model", analysis="flexit")
bone <- BP_FitBayesianCompactness(bone, analysis="logistic")
plot(bone, type="observations+model", CI="MCMC")
bone <- BP_FitBayesianCompactness(bone, analysis="flexit")
```

```

plot(bone, type="observations+model", CI="MCMC", analysis="flexit")
plot(bone, type="mcmc", parameter="P",
      options.mcmc=list(xlim=c(0.55, 0.57), breaks=seq(from=0, to=1, by=0.001)))
plot(bone, type="mcmc", parameter="S",
      options.mcmc=list(xlim=c(0.02, 0.05), breaks=seq(from=0.02, to=.05, by=0.001)))
plot(bone, type="mcmc", parameter="Min",
      options.mcmc=list(xlim=c(0.05, 0.08), breaks=seq(from=0, to=1, by=0.001)))
plot(bone, type="mcmc", parameter="Max",
      options.mcmc=list(xlim=c(0.95, 0.97), breaks=seq(from=0, to=1, by=0.001)))
outMCMC <- RM_get(x = bone, RM = "RM", RMname = "logistic", valuename = "mcmc")
summary(outMCMC)
outMCMC <- RM_get(x = bone, RM = "RM", RMname = "flexit", valuename = "mcmc")
summary(outMCMC)
# pdf(file = "Figure 3.pdf", width = 8, height = 10, pointsize = 12)
layout(1:2)
plot(bone, type="mcmc", parameter="K1", analysis="flexit",
      options.mcmc=list(xlim=c(-1, 3), ylim=c(0,10),
                         breaks=seq(from=-1, to=3, by=0.001),
                         legend = FALSE, show.prior = FALSE, mar=c(4, 4, 1, 6)), restorePar=FALSE)
segments(x0=1, x1=1,
         y0=0, y1=10, lty=4, lwd=3)
text(x=ScalePreviousPlot(x=0.95, y=0.95)$x,
     y=ScalePreviousPlot(x=0.95, y=0.95)$y, labels="A", cex=3)
plot(bone, type="mcmc", parameter="K2", analysis="flexit",
      options.mcmc=list(xlim=c(-1, 3), ylim=c(0,10),
                         breaks=seq(from=-1, to=3, by=0.001),
                         legend = FALSE, show.prior = FALSE, mar=c(4, 4, 1, 6)), restorePar=FALSE)
segments(x0=1, x1=1,
         y0=0, y1=10, lty=4, lwd=3)
text(x=ScalePreviousPlot(x=0.95, y=0.95)$x,
     y=ScalePreviousPlot(x=0.95, y=0.95)$y, labels="B", cex=3)
# dev.off()

bone <- BP_FitMLRadialCompactness(bone, analysis = "flexit")
plot(bone, type="radial", radial.variable=c("P", "S"), analysis = "flexit")
plot(bone, type="radial", radial.variable=c("P", "S", "Min", "Max"), analysis = "flexit")
out <- RM_get(x=bone, RMname="flexit", valuename = "optimRadial")$synthesis
mean(out[, "P"]); sd(out[, "P"])
range(out[, "S"])
quantile(out[, "S"])
# pdf(file = "Figure 4.pdf", width=7, height = 9, pointsize = 12)
layout(1:2)
plot(bone, type="radial", radial.variable="P", analysis = "flexit", restorePar=FALSE)
text(x=ScalePreviousPlot(x=0.95, y=0.95)$x,
     y=ScalePreviousPlot(x=0.95, y=0.95)$y, labels="A", cex=3)
plot(bone, type="radial", radial.variable="S", analysis = "flexit", restorePar=FALSE)
text(x=ScalePreviousPlot(x=0.95, y=0.95)$x,
     y=ScalePreviousPlot(x=0.95, y=0.95)$y, labels="B", cex=3)
# dev.off()

## End(Not run)

```

---

BP                   *Run a shiny application to fit bone section*

---

**Description**

Run a shiny application to fit bone section

**Usage**

```
BP()
```

**Details**

BP runs a shiny application to fit bone section

**Value**

Nothing

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**Examples**

```
## Not run:  
# Not run:  
library(BoneProfileR)  
BP()  
  
## End(Not run)
```

---

BP\_AutoFit           *Fit model automatically*

---

**Description**

Open an image, fit a model and generate a report.

**Usage**

```
BP_AutoFit(  
  file = file.choose(),  
  xlsx = TRUE,  
  rotation.angle = 0,  
  center = "ontogenetic"  
)
```

**Arguments**

file	The file to be opened
xlsx	TRUE, FALSE or the name and path of the report
rotation.angle	The angle of rotation for analysis
center	Which center to be used.

**Details**

BP\_AutoFit fits model automatically

**Value**

Characteristics of an image with all the fit information

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**See Also**

Other BoneProfileR: [BP\\_ChooseBackground\(\)](#), [BP\\_ChooseCenter\(\)](#), [BP\\_ChooseForeground\(\)](#), [BP\\_DetectBackground\(\)](#), [BP\\_DetectCenters\(\)](#), [BP\\_DetectForeground\(\)](#), [BP\\_DuplicateAnalysis\(\)](#), [BP\\_EstimateCompactness\(\)](#), [BP\\_FitBayesianCompactness\(\)](#), [BP\\_FitMLCompactness\(\)](#), [BP\\_FitMLRadialCompactness\(\)](#), [BP\\_GetFittedParameters\(\)](#), [BP\\_ListAnalyses\(\)](#), [BP\\_LnLCompactness\(\)](#), [BP\\_OpenImage\(\)](#), [BP\\_Report\(\)](#), [Erinaceus\\_europaeus](#), [plot.BoneProfileR\(\)](#), [summary.BoneProfileR\(\)](#)

**Examples**

```
## Not run:
# Not run:
library(BoneProfileR)
path_Hedgehog <- system.file("extdata", "Erinaceus_europaeus_fem_2-1_small.png",
                             package = "BoneProfileR")
bone <- BP_AutoFit(file=path_Hedgehog, xlsx=TRUE)
# or to open a dialog box
bone <- BP_AutoFit()

## End(Not run)
```

BP\_ChooseBackground    *Let the user to choose the background color of an image*

**Description**

Let the user to choose the background color of an image.

**Usage**

```
BP_ChooseBackground(bone, analysis = 1)
```

**Arguments**

bone	The bone image to be used
analysis	The name or rank of analysis

**Details**

BP\_ChooseBackground lets the user to choose the background color of an image

**Value**

The original bone object with a new attribute for background color

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**See Also**

Other BoneProfileR: [BP\\_AutoFit\(\)](#), [BP\\_ChooseCenter\(\)](#), [BP\\_ChooseForeground\(\)](#), [BP\\_DetectBackground\(\)](#), [BP\\_DetectCenters\(\)](#), [BP\\_DetectForeground\(\)](#), [BP\\_DuplicateAnalysis\(\)](#), [BP\\_EstimateCompactness\(\)](#), [BP\\_FitBayesianCompactness\(\)](#), [BP\\_FitMLCompactness\(\)](#), [BP\\_FitMLRadialCompactness\(\)](#), [BP\\_GetFittedParameters\(\)](#), [BP\\_ListAnalyses\(\)](#), [BP\\_LnLCompactness\(\)](#), [BP\\_OpenImage\(\)](#), [BP\\_Report\(\)](#), [Erinaceus\\_europaeus](#), [plot.BoneProfileR\(\)](#), [summary.BoneProfileR\(\)](#)

**Examples**

```
## Not run:
# Not run:
path_Hedgehog <- system.file("extdata", "Erinaceus_europaeus_fem_2-1_small.tif",
                             package = "BoneProfileR")
bone <- BP_OpenImage(file=path_Hedgehog)
bone <- BP_ChooseBackground(bone=bone)
bone <- BP_ChooseForeground(bone=bone)
plot(bone)

## End(Not run)
```

---

BP\_ChooseCenter      *Let the user to choose the center of the bone*

---

## Description

Let the user to choose the center of the bone.

## Usage

```
BP_ChooseCenter(bone, analysis = 1)
```

## Arguments

bone	The bone image to be used
analysis	The name or rank of analysis

## Details

BP\_ChooseCenter lets the use to choose the center of the bone

## Value

The original bone object with a new attribute for center

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## See Also

Other BoneProfileR: [BP\\_AutoFit\(\)](#), [BP\\_ChooseBackground\(\)](#), [BP\\_ChooseForeground\(\)](#), [BP\\_DetectBackground\(\)](#), [BP\\_DetectCenters\(\)](#), [BP\\_DetectForeground\(\)](#), [BP\\_DuplicateAnalysis\(\)](#), [BP\\_EstimateCompactness\(\)](#), [BP\\_FitBayesianCompactness\(\)](#), [BP\\_FitMLCompactness\(\)](#), [BP\\_FitMLRadialCompactness\(\)](#), [BP\\_GetFittedParameters\(\)](#), [BP\\_ListAnalyses\(\)](#), [BP\\_LnLCompactness\(\)](#), [BP\\_OpenImage\(\)](#), [BP\\_Report\(\)](#), [Erinaceus\\_europaeus](#), [plot.BoneProfileR\(\)](#), [summary.BoneProfileR\(\)](#)

## Examples

```
## Not run:  
# Not run:  
library(BoneProfileR)  
path_Hedgehog <- system.file("extdata", "Erinaceus_europaeus_fem_2-1_small.png",  
                             package = "BoneProfileR")  
bone <- BP_OpenImage(file=path_Hedgehog)  
bone <- BP_DetectBackground(bone=bone)  
bone <- BP_DetectForeground(bone=bone)  
bone <- BP_ChooseCenter(bone=bone)  
# For partial section, only BP_ChooseCenter() must be used  
path_Dicynodon <- system.file("extdata", "Dicynodon_tibia_11.11.1.T_b_b-1.png",
```

```

    package = "BoneProfileR")
bone <- BP_OpenImage(file=path_Dicynodon)
bone <- BP_DetectBackground(bone=bone)
bone <- BP_DetectForeground(bone=bone)
bone <- BP_ChooseCenter(bone=bone)
bone <- BP_EstimateCompactness(bone, center="user", partial=TRUE)
bone <- BP_FitMLCompactness(bone, analysis=1)
plot(bone, type="observations+model")

## End(Not run)

```

**BP\_ChooseForeground**     *Let the user to choose the foreground color of an image*

## Description

Let the user to choose the foreground color of an image.

## Usage

```
BP_ChooseForeground(bone, analysis = 1)
```

## Arguments

bone	The bone image to be used
analysis	The name or rank of analysis

## Details

BP\_ChooseForeground let the user to choose the foreground color of an image

## Value

The original bone object with a new attribute for foreground color

## Author(s)

Marc Girondot <marc.girondot@u-psud.fr>

## See Also

Other BoneProfileR: [BP\\_AutoFit\(\)](#), [BP\\_ChooseBackground\(\)](#), [BP\\_ChooseCenter\(\)](#), [BP\\_DetectBackground\(\)](#), [BP\\_DetectCenters\(\)](#), [BP\\_DetectForeground\(\)](#), [BP\\_DuplicateAnalysis\(\)](#), [BP\\_EstimateCompactness\(\)](#), [BP\\_FitBayesianCompactness\(\)](#), [BP\\_FitMLCompactness\(\)](#), [BP\\_FitMLRadialCompactness\(\)](#), [BP\\_GetFittedParameters\(\)](#), [BP\\_ListAnalyses\(\)](#), [BP\\_LnLCompactness\(\)](#), [BP\\_OpenImage\(\)](#), [BP\\_Report\(\)](#), [Erinaceus\\_europaeus](#), [plot.BoneProfileR\(\)](#), [summary.BoneProfileR\(\)](#)

## Examples

```
## Not run:
# Not run:
bone <- BP_OpenImage()
bone <- BP_ChooseBackground(bone=bone)
bone <- BP_ChooseForeground(bone=bone)
plot(bone)

## End(Not run)
```

**BP\_DetectBackground**     *Detects the background color of an image*

## Description

Detects the background color of an image.

## Usage

```
BP_DetectBackground(bone, analysis = 1, show.plot = TRUE)
```

## Arguments

bone	The bone image to be used
analysis	The name or rank of analysis
show.plot	should plot is shown ?

## Details

BP\_DetectBackground detects the background color of an image

## Value

The original bone object with a new attribute for background color

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## See Also

Other BoneProfileR: [BP\\_AutoFit\(\)](#), [BP\\_ChooseBackground\(\)](#), [BP\\_ChooseCenter\(\)](#), [BP\\_ChOOSEforeground\(\)](#), [BP\\_DetectCenters\(\)](#), [BP\\_DetectForeground\(\)](#), [BP\\_DuplicateAnalysis\(\)](#), [BP\\_EstimateCompactness\(\)](#), [BP\\_FitBayesianCompactness\(\)](#), [BP\\_FitMLCompactness\(\)](#), [BP\\_FitMLRadialCompactness\(\)](#), [BP\\_GetFittedParameters\(\)](#), [BP\\_ListAnalyses\(\)](#), [BP\\_LnLCompactness\(\)](#), [BP\\_OpenImage\(\)](#), [BP\\_Report\(\)](#), [Erinaceus\\_europaeus](#), [plot.BoneProfileR\(\)](#), [summary.BoneProfileR\(\)](#)

## Examples

```
## Not run:
# Not run:
bone <- BP_OpenImage()
bone <- BP_DetectBackground(bone=bone)
bone <- BP_DetectForeground(bone=bone)
plot(bone)

## End(Not run)
```

**BP\_DetectCenters**      *Detect the centers of an image*

## Description

Detects the centers of an image. Note that this function must not be used with partial bone section.

## Usage

```
BP_DetectCenters(bone, analysis = 1, show.plot = TRUE)
```

## Arguments

bone	The bone image to be used
analysis	The name or rank of analysis
show.plot	should plot is shown ?

## Details

BP\_DetectCenters detects the centers of an image

## Value

The original bone object with a new attribute for centers

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## See Also

Other BoneProfileR: [BP\\_AutoFit\(\)](#), [BP\\_ChooseBackground\(\)](#), [BP\\_ChooseCenter\(\)](#), [BP\\_ChOOSEforeground\(\)](#), [BP\\_DetectBackground\(\)](#), [BP\\_DetectForeground\(\)](#), [BP\\_DuplicateAnalysis\(\)](#), [BP\\_EstimateCompactness\(\)](#), [BP\\_FitBayesianCompactness\(\)](#), [BP\\_FitMLCompactness\(\)](#), [BP\\_FitMLRadialCompactness\(\)](#), [BP\\_GetFittedParameters\(\)](#), [BP\\_ListAnalyses\(\)](#), [BP\\_LnLCompactness\(\)](#), [BP\\_OpenImage\(\)](#), [BP\\_Report\(\)](#), [Erinaceus\\_europaeus](#), [plot.BoneProfileR\(\)](#), [summary.BoneProfileR\(\)](#)

## Examples

```
## Not run:
# Not run:
library(BoneProfileR)
bone <- BP_OpenImage()
# or
path_Hedgehog <- system.file("extdata", "Erinaceus_europaeus_fem_2-1_small.png",
                             package = "BoneProfileR")
bone <- BP_OpenImage(file=path_Hedgehog)
bone <- BP_DetectBackground(bone=bone)
bone <- BP_DetectForeground(bone=bone)
bone <- BP_DetectCenters(bone=bone)
plot(bone, type="mineralized", show.grid=FALSE)
plot(bone, type="unmineralized", show.grid=FALSE)
plot(bone, type="section", show.grid=FALSE)

## End(Not run)
```

**BP\_DetectForeground**     *Detects the foreground color of an image*

## Description

Detects the foreground color of an image.

## Usage

```
BP_DetectForeground(bone, analysis = 1, show.plot = TRUE)
```

## Arguments

- bone              The bone image to be used
- analysis          The name or rank of analysis
- show.plot        should plot is shown ?

## Details

BP\_DetectForeground detects the foreground color of an image

## Value

The original bone object with a new attribute for foreground color

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

**See Also**

Other BoneProfileR: [BP\\_AutoFit\(\)](#), [BP\\_ChooseBackground\(\)](#), [BP\\_ChooseCenter\(\)](#), [BP\\_ChooseForeground\(\)](#), [BP\\_DetectBackground\(\)](#), [BP\\_DetectCenters\(\)](#), [BP\\_DuplicateAnalysis\(\)](#), [BP\\_EstimateCompactness\(\)](#), [BP\\_FitBayesianCompactness\(\)](#), [BP\\_FitMLCompactness\(\)](#), [BP\\_FitMLRadialCompactness\(\)](#), [BP\\_GetFittedParameters\(\)](#), [BP\\_ListAnalyses\(\)](#), [BP\\_LnLCompactness\(\)](#), [BP\\_OpenImage\(\)](#), [BP\\_Report\(\)](#), [Erinaceus\\_europaeus](#), [plot.BoneProfileR\(\)](#), [summary.BoneProfileR\(\)](#)

**Examples**

```
## Not run:
# Not run:
bone <- BP_OpenImage()
bone <- BP_DetectBackground(bone=bone)
bone <- BP_DetectForeground(bone=bone)
plot(bone)

## End(Not run)
```

**BP\_DuplicateAnalysis** *Duplicates an analysis stored in an object*

**Description**

Duplicates an analysis stored in an object.

**Usage**

```
BP_DuplicateAnalysis(bone, from = 1, to = 2)
```

**Arguments**

bone	The bone image to be used
from	The name or rank of analysis to be duplicated
to	The name or rank of analysis to be created

**Details**

`BP_DuplicateAnalysis` duplicates an analysis stored in an object

**Value**

The original bone object with a new analysis

**Author(s)**

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

## See Also

Other BoneProfileR: [BP\\_AutoFit\(\)](#), [BP\\_ChooseBackground\(\)](#), [BP\\_ChooseCenter\(\)](#), [BP\\_ChooseForeground\(\)](#), [BP\\_DetectBackground\(\)](#), [BP\\_DetectCenters\(\)](#), [BP\\_DetectForeground\(\)](#), [BP\\_EstimateCompactness\(\)](#), [BP\\_FitBayesianCompactness\(\)](#), [BP\\_FitMLCompactness\(\)](#), [BP\\_FitMLRadialCompactness\(\)](#), [BP\\_GetFittedParameters\(\)](#), [BP\\_ListAnalyses\(\)](#), [BP\\_LnLCompactness\(\)](#), [BP\\_OpenImage\(\)](#), [BP\\_Report\(\)](#), [Erinaceus\\_europaeus](#), [plot.BoneProfileR\(\)](#), [summary.BoneProfileR\(\)](#)

## Examples

```
## Not run:
# Not run:
library(BoneProfileR)
path_Hedgehog <- system.file("extdata", "Erinaceus_europaeus_fem_2-1_small.png",
                             package = "BoneProfileR")
bone <- BP_OpenImage(file=path_Hedgehog)
bone <- BP_DetectBackground(bone=bone, analysis="logistic")
bone <- BP_DetectForeground(bone=bone, analysis="logistic")
bone <- BP_DetectCenters(bone=bone, analysis="logistic")
bone <- BP_EstimateCompactness(bone, analysis="logistic")
bone <- BP_FitMLCompactness(bone, analysis="logistic")
plot(bone)
plot(bone, type="observations")
plot(bone, type="observations+model", analysis=1)
bone <- BP_DuplicateAnalysis(bone, from="logistic", to="flexit")

## End(Not run)
```

## BP\_EstimateCompactness

*Estimation of the compactness of a bone section*

### Description

Estimation of the compactness of a bone section.

The reference for radial estimation of compactness is the trigonometric circle for rotation.angle=0 in BP\_EstimateCompactness():

- The top of the section is located at -pi/2.
- The left of the section is located at -pi and +pi.
- The bottom of the section is located at pi/2.
- The right of the section is 0.

If rotation.angle is different from 0, the value of rotation.angle is added to the angle modulo 2.pi.

### Usage

```
BP_EstimateCompactness(
  bone,
  center = "ontogenetic",
  partial = FALSE,
```

```

cut.angle = 60,
cut.distance = 100,
rotation.angle = 0,
analysis = 1,
show.plot = TRUE
)

```

### Arguments

<code>bone</code>	The bone image to be used
<code>center</code>	Which center to be used: user, mineralized, unmineralized, section, ontogenetic
<code>partial</code>	Is the section partial?
<code>cut.angle</code>	Number of angles
<code>cut.distance</code>	Number of distances
<code>rotation.angle</code>	The angle of rotation for analysis
<code>analysis</code>	The name or rank of analysis
<code>show.plot</code>	should plot is shown ?

### Details

`BP_EstimateCompactness` estimates the compactness of a bone section

### Value

The original bone object with a new attribute for compactness

### Author(s)

Marc Girondot <marc.girondot@gmail.com>

### See Also

Other BoneProfileR: [BP\\_AutoFit\(\)](#), [BP\\_ChooseBackground\(\)](#), [BP\\_ChooseCenter\(\)](#), [BP\\_ChooseForeground\(\)](#), [BP\\_DetectBackground\(\)](#), [BP\\_DetectCenters\(\)](#), [BP\\_DetectForeground\(\)](#), [BP\\_DuplicateAnalysis\(\)](#), [BP\\_FitBayesianCompactness\(\)](#), [BP\\_FitMLCompactness\(\)](#), [BP\\_FitMLRadialCompactness\(\)](#), [BP\\_GetFittedParameters\(\)](#), [BP\\_ListAnalyses\(\)](#), [BP\\_LnLCompactness\(\)](#), [BP\\_OpenImage\(\)](#), [BP\\_Report\(\)](#), [Erinaceus\\_europaeus](#), [plot.BoneProfileR\(\)](#), [summary.BoneProfileR\(\)](#)

### Examples

```

## Not run:
# Not run:
library(BoneProfileR)
bone <- BP_OpenImage()
# or
path_Hedgehog <- system.file("extdata", "Erinaceus_europaeus_fem_2-1_small.png",
                             package = "BoneProfileR")
bone <- BP_OpenImage(file=path_Hedgehog)

```

```

bone <- BP_DetectBackground(bone=bone)
bone <- BP_DetectForeground(bone=bone)
bone <- BP_DetectCenters(bone=bone)
bone <- BP_EstimateCompactness(bone)
plot(bone, type="original", show.grid=FALSE)
plot(bone, type="mineralized", show.grid=FALSE)
plot(bone, type="unmineralized", show.grid=FALSE)
plot(bone, type="section", show.grid=FALSE)

## End(Not run)

```

**BP\_FitBayesianCompactness***Estimation of Bayesian model of a bone section***Description**

Estimation of Bayesian model of a bone section.

**Usage**

```

BP_FitBayesianCompactness(
  bone = stop("A result from BP_FitMLCompactness() must be provided"),
  priors = NULL,
  n.iter = 10000,
  n.chains = 1,
  n.adapt = 100,
  thin = 1,
  analysis = 1,
  silent = TRUE
)

```

**Arguments**

bone	The bone image to be used
priors	Priors
n.iter	Number of iterations
n.chains	Number of chains
n.adapt	Number of iteration to adapt
thin	Thin parameter for analysis
analysis	Name or rank of analysis
silent	Should some information must me shown ?

**Details**

BP\_FitBayesianCompactness estimates Bayesian model of a bone section

**Value**

The -Ln L

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**See Also**

Other BoneProfileR: [BP\\_AutoFit\(\)](#), [BP\\_ChooseBackground\(\)](#), [BP\\_ChooseCenter\(\)](#), [BP\\_ChOOSEforeground\(\)](#), [BP\\_DetectBackground\(\)](#), [BP\\_DetectCenters\(\)](#), [BP\\_DetectForeground\(\)](#), [BP\\_DuplicateAnalysis\(\)](#), [BP\\_EstimateCompactness\(\)](#), [BP\\_FitMLCompactness\(\)](#), [BP\\_FitMLRadialCompactness\(\)](#), [BP\\_GetFittedParameters\(\)](#), [BP\\_ListAnalyses\(\)](#), [BP\\_LnLCompactness\(\)](#), [BP\\_OpenImage\(\)](#), [BP\\_Report\(\)](#), [Erinaceus\\_europaeus](#), [plot.BoneProfileR\(\)](#), [summary.BoneProfileR\(\)](#)

**Examples**

```
## Not run:
# Not run:
library(BoneProfileR)
path_Hedgehog <- system.file("extdata", "Erinaceus_europaeus_fem_2-1_small.png",
                             package = "BoneProfileR")
bone <- BP_OpenImage(file=path_Hedgehog)
bone <- BP_DetectBackground(bone=bone, analysis="logistic")
bone <- BP_DetectForeground(bone=bone, analysis="logistic")
bone <- BP_DetectCenters(bone=bone, analysis="logistic")
bone <- BP_EstimateCompactness(bone, analysis="logistic")
bone <- BP_FitMLCompactness(bone, analysis="logistic")
plot(bone)
plot(bone, type="observations")
plot(bone, type="observations+model", analysis=1)
fittedpar <- BP_GetFittedParameters(bone, analysis="logistic")
bone <- BP_DuplicateAnalysis(bone, from="logistic", to="flexit")
bone <- BP_FitMLCompactness(bone,
                           fitted.parameters=c(fittedpar, K1=1, K2=1),
                           fixed.parameters=NULL, analysis="flexit")
compare_AIC(Logistic=BP_GetFittedParameters(bone, analysis="logistic", alloptim=TRUE),
            Flexit=BP_GetFittedParameters(bone, analysis="flexit", alloptim=TRUE))
out4p <- plot(bone, type="observations+model", analysis="logistic")
out6p <- plot(bone, type="observations+model", analysis="flexit")
bone <- BP_FitBayesianCompactness(bone, analysis="logistic")
plot(bone, type="observations+model", CI="MCMC", analysis="logistic")
bone <- BP_FitBayesianCompactness(bone, analysis="flexit")
plot(bone, type="observations+model", CI="MCMC", analysis="flexit")

## End(Not run)
```

---

BP\_FitMLCompactness     *Estimation of the likelihood of a bone section*

---

## Description

Estimation of the model of compactness of a bone section.

The two-steps analysis performs first a quasi-Newton method, then a Bayesian MCMC and finally again a quasi-Newton method. It generally ensures that global minimum is found. On the other hand, it doubles the time to complete.

## Usage

```
BP_FitMLCompactness(
  bone,
  fitted.parameters = c(P = 0.5, S = 0.05, Min = 0.001, Max = 0.999),
  priors = NULL,
  fixed.parameters = c(K1 = 1, K2 = 1),
  twosteps = TRUE,
  replicates.CI = 10000,
  analysis = 1,
  silent = FALSE
)
```

## Arguments

<code>bone</code>	The bone image to be used
<code>fitted.parameters</code>	Parameters of the model to be fitted
<code>priors</code>	Priors used for intermediate estimations
<code>fixed.parameters</code>	Fixed parameters of the model
<code>twosteps</code>	Does a 2-steps analysis be performed?
<code>replicates.CI</code>	Number of replicates to estimate confidence interval
<code>analysis</code>	Name or rank of analysis
<code>silent</code>	Should information be shown?

## Details

BP\_FitMLCompactness estimates likelihood of model of a bone section

## Value

The -Ln L

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**See Also**

Other BoneProfileR: [BP\\_AutoFit\(\)](#), [BP\\_ChooseBackground\(\)](#), [BP\\_ChooseCenter\(\)](#), [BP\\_ChOOSEforeground\(\)](#), [BP\\_DetectBackground\(\)](#), [BP\\_DetectCenters\(\)](#), [BP\\_DetectForeground\(\)](#), [BP\\_DuplicateAnalysis\(\)](#), [BP\\_EstimateCompactness\(\)](#), [BP\\_FitBayesianCompactness\(\)](#), [BP\\_FitMLRadialCompactness\(\)](#), [BP\\_GetFittedParameters\(\)](#), [BP\\_ListAnalyses\(\)](#), [BP\\_LnLCompactness\(\)](#), [BP\\_OpenImage\(\)](#), [BP\\_Report\(\)](#), [Erinaceus\\_europaeus](#), [plot.BoneProfileR\(\)](#), [summary.BoneProfileR\(\)](#)

**Examples**

```
## Not run:
# Not run:
library(BoneProfileR)
bone <- BP_OpenImage()
# or, to use the package imager to open a tiff image
bone <- BP_OpenImage(ijtiff=TRUE)
library(BoneProfileR)
path_Hedgehog <- system.file("extdata", "Erinaceus_europaeus_fem_2-1_small.png",
                             package = "BoneProfileR")
bone <- BP_OpenImage(file=path_Hedgehog)
bone <- BP_DetectBackground(bone=bone, analysis="logistic")
bone <- BP_DetectForeground(bone=bone, analysis="logistic")
bone <- BP_DetectCenters(bone=bone, analysis="logistic")
bone <- BP_EstimateCompactness(bone, analysis="logistic")
plot(bone, type="mineralized", show.grid=FALSE)
plot(bone, type="unmineralized", show.grid=FALSE)
plot(bone, type="section", show.grid=FALSE)
bone <- BP_FitMLCompactness(bone, analysis="logistic", twosteps=TRUE)
BP_GetFittedParameters(bone)
plot(bone)
plot(bone, type="observations")
plot(bone, type="observations+model", analysis=1)
bone <- BP_DuplicateAnalysis(bone, from="logistic", to="flexit")
fittedpar <- BP_GetFittedParameters(bone, analysis="logistic")
bone <- BP_DuplicateAnalysis(bone, from="logistic", to="flexit")
BP_ListAnalyses(bone)
bone <- BP_FitMLCompactness(bone,
                            fitted.parameters=c(fittedpar, K1=1, K2=1),
                            fixed.parameters=NULL, analysis="flexit", twosteps=TRUE)
compare_AIC(Logistic=BP_GetFittedParameters(bone, analysis="logistic", alloptim=TRUE),
            Flexit=BP_GetFittedParameters(bone, analysis="flexit", alloptim=TRUE))
out4p <- plot(bone, type="observations+model", analysis="logistic")
out6p <- plot(bone, type="observations+model", analysis="flexit")

## End(Not run)
```

---

BP\_FitMLRadialCompactness*Estimation of the likelihood of a bone section*

---

**Description**

Estimation of the compactness of a bone section using radial model.

If the fitted.parameters and fixed.parameters are NULL and the analysis includes a BP\_FitMLCompactness() result, the values of this result is used as a reference for fitted.parameters and fixed.parameters.

If no BP\_FitMLCompactness() result is available, it will use:

fitted.parameters=c(P=0.5, S=0.05, Min=-2, Max=5); fixed.parameters=c(K1=1, K2=1).

The reference for radial estimation of compactness is the trigonometric circle for rotation.angle=0 in BP\_EstimateCompactness():

- The top of the section is located at -pi/2.
- The left of the section is located at -pi and +pi.
- The bottom of the section is located at pi/2.
- The right of the section is 0.

If rotation.angle is different from 0, the value of rotation.angle is added to the angle modulo 2.pi.

The two-steps analysis performs first a quasi-Newton method, then a Bayesian MCMC and finally again a quasi-Newton method. It generally ensures that global minimum is found. On the other hand, it doubles the time to complete for each angle.

**Usage**

```
BP_FitMLRadialCompactness(
  bone,
  fitted.parameters = NULL,
  priors = NULL,
  fixed.parameters = NULL,
  analysis = 1,
  silent = FALSE,
  twosteps = TRUE
)
```

**Arguments**

bone	The bone image to be used
fitted.parameters	Parameters of the model to be fitted
priors	If twosteps is TRUE, tell what prior should be used.
fixed.parameters	Fixed parameters of the model
analysis	Name or rank of analysis
silent	Should the function displays some information?
twosteps	Should a 2-steps analysis be performed?

**Details**

*BP\_FitMLRadialCompactness* estimates likelihood of model of a bone section

**Value**

The -Ln L

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**See Also**

Other BoneProfileR: [BP\\_AutoFit\(\)](#), [BP\\_ChooseBackground\(\)](#), [BP\\_ChooseCenter\(\)](#), [BP\\_ChooseForeground\(\)](#), [BP\\_DetectBackground\(\)](#), [BP\\_DetectCenters\(\)](#), [BP\\_DetectForeground\(\)](#), [BP\\_DuplicateAnalysis\(\)](#), [BP\\_EstimateCompactness\(\)](#), [BP\\_FitBayesianCompactness\(\)](#), [BP\\_FitMLCompactness\(\)](#), [BP\\_GetFittedParameters\(\)](#), [BP\\_ListAnalyses\(\)](#), [BP\\_LnLCompactness\(\)](#), [BP\\_OpenImage\(\)](#), [BP\\_Report\(\)](#), [Erinaceus\\_europaeus](#), [plot.BoneProfileR\(\)](#), [summary.BoneProfileR\(\)](#)

**Examples**

```
## Not run:
# Not run
library(BoneProfileR)
path_Hedgehog <- system.file("extdata", "Erinaceus_europaeus_fem_2-1_small.png",
                             package = "BoneProfileR")
bone <- BP_OpenImage(file=path_Hedgehog)
# or
bone <- BP_OpenImage(ijtiff=TRUE)
bone <- BP_DetectBackground(bone=bone, analysis="logistic")
bone <- BP_DetectForeground(bone=bone, analysis="logistic")
bone <- BP_DetectCenters(bone=bone, analysis="logistic")
bone <- BP_EstimateCompactness(bone, analysis="logistic")
bone <- BP_EstimateCompactness(bone, analysis="logistic", cut.angle=30)
bone <- BP_FitMLCompactness(bone, analysis="logistic")
plot(bone)
plot(bone, type="observations")
plot(bone, type="observations+model", analysis=1)
fittedpar <- BP_GetFittedParameters(bone, analysis="logistic")
bone <- BP_DuplicateAnalysis(bone, from="logistic", to="flexit")
bone <- BP_FitMLCompactness(bone,
                           fitted.parameters=c(fittedpar, K1=1.01, K2=1.01),
                           fixed.parameters=NULL, analysis="flexit")
bone <- BP_FitBayesianCompactness(bone, analysis="flexit")
mcmc <- RM_get(bone, RMname = "flexit", value="mcmc")
fittedpar <- as.parameters(mcmc)
bone <- BP_FitMLCompactness(bone,
                           fitted.parameters=fittedpar,
                           fixed.parameters=NULL, analysis="flexit")
compare_AIC(Logistic=BP_GetFittedParameters(bone, analysis="logistic", alloptim=TRUE),
            Flexit=BP_GetFittedParameters(bone, analysis="flexit", alloptim=TRUE))
```

```

out4p <- plot(bone, type="observations+model", analysis="logistic")
out6p <- plot(bone, type="observations+model", analysis="flexit")
# The twosteps fit is more accurate but is around 100 times slower
bone <- BP_FitMLRadialCompactness(bone, analysis="logistic", twosteps=TRUE)
bone <- BP_FitMLRadialCompactness(bone, analysis="logistic", twosteps=FALSE)
plot(bone, type="observations", angle=0)
plot(bone, type="model", analysis="logistic", angle=0)
plot(bone, type="observations+model", angle=0)
plot(bone, type="observations+model", angle=pi)
plot(bone, type="radial", radial.variable=c("P", "S"), analysis="logistic")
plot(bone, type="radial", radial.variable=c("P", "S", "Min", "Max"), analysis="logistic")
plot(bone, type="radial", radial.variable=c("TRC"), analysis="logistic")
# Test using the change of orientation using default.angle from BP_EstimateCompactness():
bone <- BP_DuplicateAnalysis(bone, from="logistic", to="logistic_rotation_pi")
# With a pi rotation, the top moves to the bottom and the left moves to the right
bone <- BP_EstimateCompactness(bone, rotation.angle=pi, analysis="logistic_rotation_pi")
bone <- BP_FitMLRadialCompactness(bone, analysis="logistic_rotation_pi")
plot(bone, type="radial", radial.variable=c("P", "S"), analysis="logistic")
plot(bone, type="radial", radial.variable=c("P", "S"), analysis="logistic_rotation_pi")
BP_Report(bone=bone,
           analysis=1,
           docx=NULL,
           pdf=NULL,
           xlsx=file.path(getwd(), "report.xlsx"),
           author="Marc Girondot",
           title=attributes(bone)$name)

## End(Not run)

```

**BP\_GetFittedParameters***Return the fitted parameters***Description**

Return the fitted parameters.

**Usage**

```
BP_GetFittedParameters(bone, analysis = 1, alloptim = FALSE)
```

**Arguments**

<code>bone</code>	The bone image to be used
<code>analysis</code>	Name or rank of analysis
<code>alloptim</code>	If TRUE, return the complete object returned by optim

**Details**

`BP_GetFittedParameters` returns the fitted parameters

**Value**

The fitted parameters

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**See Also**

Other BoneProfileR: [BP\\_AutoFit\(\)](#), [BP\\_ChooseBackground\(\)](#), [BP\\_ChooseCenter\(\)](#), [BP\\_ChOOSEforeground\(\)](#), [BP\\_DetectBackground\(\)](#), [BP\\_DetectCenters\(\)](#), [BP\\_Detectforeground\(\)](#), [BP\\_DuplicateAnalysis\(\)](#), [BP\\_EstimateCompactness\(\)](#), [BP\\_FitBayesianCompactness\(\)](#), [BP\\_FitMLCompactness\(\)](#), [BP\\_FitMLRadialCompactness\(\)](#), [BP\\_ListAnalyses\(\)](#), [BP\\_LnLCompactness\(\)](#), [BP\\_OpenImage\(\)](#), [BP\\_Report\(\)](#), [Erinaceus\\_europaeus](#), [plot.BoneProfileR\(\)](#), [summary.BoneProfileR\(\)](#)

**Examples**

```
## Not run:
# Not run:
library(BoneProfileR)
path_Hedgehog <- system.file("extdata", "Erinaceus_europaeus_fem_2-1_small.png",
                             package = "BoneProfileR")
bone <- BP_OpenImage(file=path_Hedgehog)
bone <- BP_DetectBackground(bone=bone, analysis="logistic")
bone <- BP_Detectforeground(bone=bone, analysis="logistic")
bone <- BP_DetectCenters(bone=bone, analysis="logistic")
bone <- BP_EstimateCompactness(bone, analysis="logistic")
bone <- BP_FitMLCompactness(bone, analysis="logistic")
BP_GetFittedParameters(bone, analysis="logistic")

## End(Not run)
```

**BP\_ListAnalyses**

*List the analyses stored in an object*

**Description**

Get the analyses stored in an object.

**Usage**

```
BP_ListAnalyses(bone, silent = TRUE, max.level = FALSE)
```

**Arguments**

bone	The bone image to be used
silent	Should the results be shown ?
max.level	If TRUE, will return all list element of the objects

## Details

BP\_ListAnalyses lists the analyses stored in an object

## Value

The list of analyses

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## See Also

Other BoneProfileR: [BP\\_AutoFit\(\)](#), [BP\\_ChooseBackground\(\)](#), [BP\\_ChooseCenter\(\)](#), [BP\\_ChooseForeground\(\)](#), [BP\\_DetectBackground\(\)](#), [BP\\_DetectCenters\(\)](#), [BP\\_DetectForeground\(\)](#), [BP\\_DuplicateAnalysis\(\)](#), [BP\\_EstimateCompactness\(\)](#), [BP\\_FitBayesianCompactness\(\)](#), [BP\\_FitMLCompactness\(\)](#), [BP\\_FitMLRadialCompactness\(\)](#), [BP\\_GetFittedParameters\(\)](#), [BP\\_LnLCompactness\(\)](#), [BP\\_OpenImage\(\)](#), [BP\\_Report\(\)](#), [Erinaceus\\_europaeus](#), [plot.BoneProfileR\(\)](#), [summary.BoneProfileR\(\)](#)

## Examples

```
## Not run:
# Not run:
library(BoneProfileR)
path_Hedgehog <- system.file("extdata", "Erinaceus_europaeus_fem_2-1_small.png",
                             package = "BoneProfileR")
bone <- BP_OpenImage(file=path_Hedgehog)
bone <- BP_DetectBackground(bone=bone, analysis="logistic")
bone <- BP_DetectForeground(bone=bone, analysis="logistic")
bone <- BP_DetectCenters(bone=bone, analysis="logistic")
bone <- BP_EstimateCompactness(bone, analysis="logistic")
bone <- BP_FitMLCompactness(bone, analysis="logistic")
plot(bone)
plot(bone, type="observations")
plot(bone, type="observations+model", analysis=1)
bone <- BP_DuplicateAnalysis(bone, from="logistic", to="flexit")
BP_ListAnalyses(bone)

## End(Not run)
```

## Description

Estimation of the compactness of a bone section.

**Usage**

```
BP_LnLCompactness(
  par,
  bone = NULL,
  data_m = NULL,
  data_nm = NULL,
  distance.center = NULL,
  fixed.parameters = NULL,
  analysis = 1
)
```

**Arguments**

<code>par</code>	Parameters of the model
<code>bone</code>	The bone image to be used
<code>data_m</code>	Number of mineralized pixels
<code>data_nm</code>	Number of non-mineralized pixels
<code>distance.center</code>	Distances to the center
<code>fixed.parameters</code>	Fixed parameters of the model
<code>analysis</code>	Name or rank of analysis

**Details**

`BP_LnLCompactness` estimates likelihood of model of a bone section

**Value**

The -Ln L

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**See Also**

Other BoneProfileR: [BP\\_AutoFit\(\)](#), [BP\\_ChooseBackground\(\)](#), [BP\\_ChooseCenter\(\)](#), [BP\\_ChooseForeground\(\)](#), [BP\\_DetectBackground\(\)](#), [BP\\_DetectCenters\(\)](#), [BP\\_DetectForeground\(\)](#), [BP\\_DuplicateAnalysis\(\)](#), [BP\\_EstimateCompactness\(\)](#), [BP\\_FitBayesianCompactness\(\)](#), [BP\\_FitMLCompactness\(\)](#), [BP\\_FitMLRadialCompactness\(\)](#), [BP\\_GetFittedParameters\(\)](#), [BP\\_ListAnalyses\(\)](#), [BP\\_OpenImage\(\)](#), [BP\\_Report\(\)](#), [Erinaceus\\_europaeus](#), [plot.BoneProfileR\(\)](#), [summary.BoneProfileR\(\)](#)

## Examples

```
## Not run:
# Not run:
path_Hedgehog <- system.file("extdata", "Erinaceus_europaeus_fem_2-1_small.png",
                             package = "BoneProfileR")
bone <- BP_OpenImage(file=path_Hedgehog)
bone <- BP_DetectBackground(bone=bone)
bone <- BP_DetectForeground(bone=bone)
bone <- BP_DetectCenters(bone=bone)
bone <- BP_EstimateCompactness(bone)
plot(bone)

## End(Not run)
```

BP\_OpenImage

*Open an image*

## Description

Open an image.

## Usage

```
BP_OpenImage(file = file.choose(), name = NULL, ijtiff = FALSE)
```

## Arguments

file	The file to be opened
name	Name of this slice
ijtiff	Should the ijtiff must be used to read tiff image

## Details

BP\_OpenImage opens an image

## Value

Characteristics of an image

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## See Also

Other BoneProfileR: [BP\\_AutoFit\(\)](#), [BP\\_ChooseBackground\(\)](#), [BP\\_ChooseCenter\(\)](#), [BP\\_ChOOSEforeground\(\)](#), [BP\\_DetectBackground\(\)](#), [BP\\_DetectCenters\(\)](#), [BP\\_DetectForeground\(\)](#), [BP\\_DuplicateAnalysis\(\)](#), [BP\\_EstimateCompactness\(\)](#), [BP\\_FitBayesianCompactness\(\)](#), [BP\\_FitMLCompactness\(\)](#), [BP\\_FitMLRadialCompactness\(\)](#), [BP\\_GetFittedParameters\(\)](#), [BP\\_ListAnalyses\(\)](#), [BP\\_LnLCompactness\(\)](#), [BP\\_Report\(\)](#), [Erinaceus\\_europaeus](#), [plot.BoneProfileR\(\)](#), [summary.BoneProfileR\(\)](#)

## Examples

```
## Not run:
# Not run:
library(BoneProfileR)
path_Hedgehog <- system.file("extdata", "Erinaceus_europaeus_fem_2-1_small.png",
                             package = "BoneProfileR")
bone <- BP_OpenImage(file=path_Hedgehog)
plot(bone)
path_Hedgehog <- system.file("extdata", "Erinaceus_europaeus_fem_2-1_small.tif",
                             package = "BoneProfileR")
bone <- BP_OpenImage(file=path_Hedgehog)
plot(bone)
bone <- BP_OpenImage(file=path_Hedgehog, ijtiff=TRUE)
plot(bone)
# A partial section
path_Dicynodon <- system.file("extdata", "Dicynodon_tibia_11.11.1.T_b_b-1.png",
                               package = "BoneProfileR")
bone <- BP_OpenImage(file=path_Dicynodon)
plot(bone)
# To open a file with a dialog:
bone <- BP_OpenImage()

## End(Not run)
```

BP\_Report

*Generate a pdf report for the analyzed bone*

## Description

Generate a docx, xlsx, or pdf report.

## Usage

```
BP_Report(
  bone = stop("A bone section must be provided"),
  control.plot = list(message = NULL, show.centers = TRUE, show.colors = TRUE,
    show.grid = TRUE, CI = "ML", show.legend = TRUE),
  analysis = 1,
  docx = file.path(getwd(), "report.docx"),
  pdf = file.path(getwd(), "report.pdf"),
  xlsx = file.path(getwd(), "report.xlsx"),
  author = NULL,
  title = attributes(bone)$name
)
```

## Arguments

bone	The bone image
control.plot	A list with the parameters used for plot
analysis	Indicate analysis name or rank that you want report
docx	Name of Word file
pdf	Name of pdf file
xlsx	Name of Excel file
author	Name indicated in the report
title	Title of the report

## Details

BP\_Report save a pdf report for the analyzed bone

## Value

Nothing

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## See Also

Other BoneProfileR: [BP\\_AutoFit\(\)](#), [BP\\_ChooseBackground\(\)](#), [BP\\_ChooseCenter\(\)](#), [BP\\_ChooseForeground\(\)](#), [BP\\_DetectBackground\(\)](#), [BP\\_DetectCenters\(\)](#), [BP\\_DetectForeground\(\)](#), [BP\\_DuplicateAnalysis\(\)](#), [BP\\_EstimateCompactness\(\)](#), [BP\\_FitBayesianCompactness\(\)](#), [BP\\_FitMLCompactness\(\)](#), [BP\\_FitMLRadialCompactness\(\)](#), [BP\\_GetFittedParameters\(\)](#), [BP\\_ListAnalyses\(\)](#), [BP\\_LnLCompactness\(\)](#), [BP\\_OpenImage\(\)](#), [Erinaceus\\_europaeus\(\)](#), [plot.BoneProfileR\(\)](#), [summary.BoneProfileR\(\)](#)

## Examples

```
## Not run:
# Not run:
library(BoneProfileR)
path_Hedgehog <- system.file("extdata", "Erinaceus_europaeus_fem_2-1_small.png",
                             package = "BoneProfileR")
bone <- BP_OpenImage(file=path_Hedgehog)
bone <- BP_DetectBackground(bone=bone, analysis="logistic")
bone <- BP_DetectForeground(bone=bone, analysis="logistic")
bone <- BP_DetectCenters(bone=bone, analysis="logistic")
bone <- BP_EstimateCompactness(bone, analysis="logistic")
bone <- BP_FitMLCompactness(bone, analysis="logistic")
fittedpar <- BP_GetFittedParameters(bone, analysis="logistic")
bone <- BP_DuplicateAnalysis(bone, from="logistic", to="flexit")
bone <- BP_FitMLCompactness(bone,
                           fitted.parameters=c(fittedpar, K1=1, K2=1),
                           fixed.parameters=NULL, analysis="flexit")
```

```

compare_AIC(Logistic=BP_GetFittedParameters(bone, analysis="logistic", alloptim=TRUE),
            Flexit=BP_GetFittedParameters(bone, analysis="flexit", alloptim=TRUE))
bone <- BP_FitMLRadialCompactness(bone, analysis="logistic")
# Test using the change of orientation using default.angle from BP_EstimateCompactness():
bone <- BP_DuplicateAnalysis(bone, from="logistic", to="logistic_rotation_pi")
# With a pi rotation, the top moves to the bottom and the left moves to the right
bone <- BP_EstimateCompactness(bone, rotation.angle=pi, analysis="logistic_rotation_pi")
bone <- BP_FitMLRadialCompactness(bone, analysis="logistic_rotation_pi")
BP_Report(bone=bone,
          analysis=1,
          docx=NULL,
          pdf=NULL,
          xlsx=file.path(getwd(), "report.xlsx"),
          author="Marc Girondot",
          title=attributes(bone)$name)

BP_Report(bone=bone,
          analysis=1,
          docx=NULL,
          pdf=file.path(getwd(), "report.pdf"),
          xlsx=NULL,
          author="Marc Girondot",
          title=attributes(bone)$name)

BP_Report(bone=bone,
          analysis=1,
          docx=file.path(getwd(), "report.docx"),
          pdf=NULL,
          xlsx=NULL,
          author="Marc Girondot",
          title=attributes(bone)$name)

## End(Not run)

```

*Erinaceus\_europaeus      Example of hedgehog femur*

## Description

Example of hedgehog femur. A drawing produced by Michel Laurin.

## Usage

*Erinaceus\_europaeus*

## Format

A png image.

## Details

Example of hedgehog femur

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## See Also

Other BoneProfileR: [BP\\_AutoFit\(\)](#), [BP\\_ChooseBackground\(\)](#), [BP\\_ChooseCenter\(\)](#), [BP\\_ChOOSEForeground\(\)](#), [BP\\_DetectBackground\(\)](#), [BP\\_DetectCenters\(\)](#), [BP\\_DetectForeground\(\)](#), [BP\\_DuplicateAnalysis\(\)](#), [BP\\_EstimateCompactness\(\)](#), [BP\\_FitBayesianCompactness\(\)](#), [BP\\_FitMLCompactness\(\)](#), [BP\\_FitMLRadialCompactness\(\)](#), [BP\\_GetFittedParameters\(\)](#), [BP\\_ListAnalyses\(\)](#), [BP\\_LnLCompactness\(\)](#), [BP\\_OpenImage\(\)](#), [BP\\_Report\(\)](#), [plot.BoneProfileR\(\)](#), [summary.BoneProfileR\(\)](#)

## Examples

```
bone <- Erinaceus_europaeus  
plot(bone)
```

---

plot.BoneProfileR      *Plot a bone section*

---

## Description

Display a bone section.

type value can be:

Image plot: original, mineralized, unmineralized, section

Original is the original image, mineralized is the mineral interpretation of the section, unmineralized is the unmineralized interpretation of the section, section is the interpretation of the section.

colors show the histograms of pixel information with foreground and background colors if they are defined.

3Dcolors show the pixels colors in 3D

Global analysis: observations, model, observations+model

Radial analysis: radial

If angle is not null and a radial analysis exists, it will show the model for this angle.

mcmc: It will show the posterior distribution of parameter

## Usage

```
## S3 method for class 'BoneProfileR'  
plot(  
  x,  
  message = NULL,  
  type = "original",  
  angle = NULL,  
  show.centers = TRUE,
```

```

show.colors = TRUE,
show.grid = TRUE,
analysis = 1,
parameter.mcmc = "S",
options.mcmc = list(),
restorePar = TRUE,
mar = NULL,
angle.3D = 55,
CI = "ML",
radial.variable = "S",
show.legend = TRUE,
...
)

```

### Arguments

<code>x</code>	The bone image
<code>message</code>	The message to be displayed
<code>type</code>	The type of plot; see description
<code>angle</code>	Which angle model to show
<code>show.centers</code>	Should the centers be shown?
<code>show.colors</code>	Should the background and foreground colors be shown?
<code>show.grid</code>	Should the grid be shown?
<code>analysis</code>	Name or number of analysis to be plotted
<code>parameter.mcmc</code>	The posterior parameter to show for type = "mcmc"
<code>options.mcmc</code>	The option to plot type mcmc output
<code>restorePar</code>	If TRUE, restore the par parameter at the exit
<code>mar</code>	The margin for type being "model" or "observations"
<code>angle.3D</code>	The angle between x and y for 3Dcolors graph
<code>CI</code>	Which confidence interval should be plotted: MCMC or ML
<code>radial.variable</code>	Name of the radial variable to plot
<code>show.legend</code>	Should a legend be shown?
<code>...</code>	Not used

### Details

`plot.BoneProfileR` displays a bone section

### Value

Nothing

### Author(s)

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

## See Also

Other BoneProfileR: [BP\\_AutoFit\(\)](#), [BP\\_ChooseBackground\(\)](#), [BP\\_ChooseCenter\(\)](#), [BP\\_ChooseForeground\(\)](#), [BP\\_DetectBackground\(\)](#), [BP\\_DetectCenters\(\)](#), [BP\\_DetectForeground\(\)](#), [BP\\_DuplicateAnalysis\(\)](#), [BP\\_EstimateCompactness\(\)](#), [BP\\_FitBayesianCompactness\(\)](#), [BP\\_FitMLCompactness\(\)](#), [BP\\_FitMLRadialCompactness\(\)](#), [BP\\_GetFittedParameters\(\)](#), [BP\\_ListAnalyses\(\)](#), [BP\\_LnLCompactness\(\)](#), [BP\\_OpenImage\(\)](#), [BP\\_Report\(\)](#), [Erinaceus\\_europaeus](#), [summary.BoneProfileR\(\)](#)

## Examples

```
## Not run:
# Not run:
library(BoneProfileR)
bone <- BP_OpenImage()
# or
path_Hedgehog <- system.file("extdata", "Erinaceus_europaeus_fem_2-1_small.png",
                             package = "BoneProfileR")
bone <- BP_OpenImage(file=path_Hedgehog)
bone <- BP_DetectBackground(bone=bone, analysis="logistic")
bone <- BP_DetectForeground(bone=bone, analysis="logistic")
plot(bone, type="colors")
bone <- BP_DetectCenters(bone=bone, analysis="logistic")
plot(bone, type="3Dcolors")
bone <- BP_EstimateCompactness(bone, analysis="logistic", rotation.angle = 1)
bone <- BP_FitMLCompactness(bone, analysis="logistic")
plot(bone)
#
path_Hedgehog <- system.file("extdata", "Erinaceus_europaeus_fem_2-1_small.png",
                             package = "BoneProfileR")
bone <- BP_OpenImage(file=path_Hedgehog)
bone <- BP_DetectBackground(bone=bone, analysis="logistic")
bone <- BP_DetectForeground(bone=bone, analysis="logistic")
bone <- BP_DetectCenters(bone=bone, analysis="logistic")
bone <- BP_EstimateCompactness(bone, analysis="logistic")
bone <- BP_FitMLCompactness(bone, analysis="logistic")
plot(bone)
plot(bone, type="observations")
plot(bone, type="observations+model", analysis=1)
bone <- BP_DuplicateAnalysis(bone, from="logistic", to="flexit")
fittedpar <- BP_GetFittedParameters(bone, analysis="logistic")
bone <- BP_DuplicateAnalysis(bone, from="logistic", to="flexit")
bone <- BP_FitMLCompactness(bone,
                           fitted.parameters=c(fittedpar, K1=1, K2=1),
                           fixed.parameters=NULL, analysis="flexit")
compare_AIC(Logistic=BP_GetFittedParameters(bone, analysis="logistic", alloptim=TRUE),
            Flexit=BP_GetFittedParameters(bone, analysis="flexit", alloptim=TRUE))
out4p <- plot(bone, type="observations+model", analysis="logistic")
out6p <- plot(bone, type="observations+model", analysis="flexit")
bone <- BP_FitBayesianCompactness(bone, analysis="logistic")
plot(bone, type="observations+model", CI="MCMC")
bone <- BP_FitMLRadialCompactness(bone)
plot(bone, type="radial", radial.variable=c("P", "S"))
plot(bone, type="radial", radial.variable=c("P", "S", "Min", "Max"))
```

---

```
## End(Not run)
```

---

*summary.BoneProfileR Plot a bone section*

---

## Description

Display information of bone section

## Usage

```
## S3 method for class 'BoneProfileR'  
summary(object, max.level = FALSE, analysis = 1, ...)
```

## Arguments

object	The bone image
max.level	If TRUE, will return all list element of the objects
analysis	The analysis to report the global compacteness
...	Not used

## Details

*summary.BoneProfileR* displays a bone section

## Value

An invisible list with recorded information

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## See Also

Other BoneProfileR: [BP\\_AutoFit\(\)](#), [BP\\_ChooseBackground\(\)](#), [BP\\_ChooseCenter\(\)](#), [BP\\_ChooseForeground\(\)](#), [BP\\_DetectBackground\(\)](#), [BP\\_DetectCenters\(\)](#), [BP\\_DetectForeground\(\)](#), [BP\\_DuplicateAnalysis\(\)](#), [BP\\_EstimateCompactness\(\)](#), [BP\\_FitBayesianCompactness\(\)](#), [BP\\_FitMLCompactness\(\)](#), [BP\\_FitMLRadialCompactness\(\)](#), [BP\\_GetFittedParameters\(\)](#), [BP\\_ListAnalyses\(\)](#), [BP\\_LnLCompactness\(\)](#), [BP\\_OpenImage\(\)](#), [BP\\_Report\(\)](#), [Erinaceus\\_europaeus](#), [plot.BoneProfileR\(\)](#)

## Examples

```
## Not run:  
# Not run:  
library(BoneProfileR)  
bone <- BP_OpenImage()  
# or  
path_Hedgehog <- system.file("extdata", "Erinaceus_europaeus_fem_2-1_small.png",  
                             package = "BoneProfileR")  
bone <- BP_OpenImage(file=path_Hedgehog)  
summary(bone)  
  
## End(Not run)
```

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