

Package ‘BoneDensityMapping’

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Type Package

Title Maps Bone Densities from CT Scans to Surface Models

Version 0.1.3

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Description Allows local bone density estimates to be derived from CT data and mapped to 3D bone models in a reproducible manner. Processing can be performed at the individual bone or group level. Also includes tools for visualizing the bone density estimates. Example methods are described in Telfer et al., (2021) <[doi:10.1002/jor.24792](https://doi.org/10.1002/jor.24792)>, Telfer et al., (2021) <[doi:10.1016/j.jse.2021.05.011](https://doi.org/10.1016/j.jse.2021.05.011)>.

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VignetteBuilder knitr

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bone_scan_check	<i>Check if surface model is fully contained within scan volume</i>
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Description

Check if surface model is fully contained within scan volume

Usage

```
bone_scan_check(surface_mesh, nifti, return_limits = FALSE)
```

Arguments

surface_mesh	mesh object (class mesh3d) or numeric matrix/dataframe of vertex coordinates (cols: X, Y, Z)
nifti	NIfTI image object representing CT scan.
return_limits	Logical. If TRUE returns a summary of the bounding boxes of the scan and mesh

Value

If any vertices lie outside the scan volume, it raises an error.

Author(s)

Scott Telfer <scott.telfer@gmail.com>

Examples

```
# Download CT scan
url <- "https://github.com/Telfer/BoneDensityMapping/releases/download/v1.0.2/test_CT_hip.nii.gz"
scan_filepath <- tempfile(fileext = ".nii.gz")
download.file(url, scan_filepath, mode = "wb")
nifti <- import_scan(scan_filepath)
url2 <- "https://github.com/Telfer/BoneDensityMapping/releases/download/v1.0.2/test_CT_femur.stl"
bone_filepath <- tempfile(fileext = ".stl")
download.file(url2, bone_filepath, mode = "wb")
surface_mesh <- import_mesh(bone_filepath)
bone_scan_check(surface_mesh, nifti, return_limits = TRUE)
```

color_bar	<i>Produce stand alone color bar</i>
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Description

Produce stand alone color bar

Usage

```
color_bar(  
  colors,  
  mini,  
  maxi,  
  orientation = "vertical",  
  breaks,  
  title = "",  
  text_size = 11,  
  plot = TRUE  
)
```

Arguments

colors	String
mini	Numeric
maxi	Numeric
orientation	"horizontal" or "vertical"
breaks	Numeric vector
title	String
text_size	Numeric
plot	Logical

Value

ggplot object

Examples

```
colors <- c("darkblue", "blue", "lightblue", "green", "yellow", "red", "pink")
color_bar(colors, 0, 2000, breaks = c(0, 500, 1000, 1500, 2000))
```

color_mapping *maps numeric values to a color*

Description

maps numeric values to a color

Usage

```
color_mapping(x, maxi, mini, color_sel)
```

Arguments

x	Vector.
maxi	Numeric. Maximum value. Defaults to maximum value in vector. Can be useful to set this manually if there are outliers in the scan data
mini	Numeric. Minimum value. Defaults to minimum value in vector. Can be useful to set this manually if there are outliers in the scan data
color_sel	Vector. Colors to use for map. Defaults to a scale of "grey", "blue", "green", "yellow", "orange", "red", "pink".

Value

Vector of hex color values same length as x

Author(s)

Scott Telfer <scott.telfer@gmail.com>

Examples

```
# Download CT scan
url <- "https://github.com/Telfer/BoneDensityMapping/releases/download/v1.0.2/test_CT_hip.nii.gz"
scan_filepath <- tempfile(fileext = ".nii.gz")
download.file(url, scan_filepath, mode = "wb")
nifti <- import_scan(scan_filepath)
url2 <- "https://github.com/Telfer/BoneDensityMapping/releases/download/v1.0.2/test_CT_femur.stl"
bone_filepath <- tempfile(fileext = ".stl")
download.file(url2, bone_filepath, mode = "wb")
surface_mesh <- import_mesh(bone_filepath)
mat_peak <- voxel_point_intersect(surface_mesh, nifti, ct_eqn = "linear",
                                ct_params = c(68.4, 1.106),)
colors <- color_mapping(mat_peak)
```

color_mesh	<i>Takes a density vector mapped to standardized coordinates and maps it to a surface mesh for visualization.</i>
------------	---

Description

Takes a density vector mapped to standardized coordinates and maps it to a surface mesh for visualization.

Usage

```
color_mesh(  
  surface_mesh,  
  template_pts,  
  density_vector,  
  maxi = NULL,  
  mini = NULL,  
  export_path,  
  color_sel  
)
```

Arguments

surface_mesh	Mesh object
template_pts	Matrix
density_vector	Vector
maxi	Numeric
mini	Numeric
export_path	Character
color_sel	String

Value

mesh3d object with added color dimension

Author(s)

Scott Telfer <scott.telfer@gmail.com>

Examples

```
# Download CT scan  
url <- "https://github.com/Telfer/BoneDensityMapping/releases/download/v1.0.1/test_CT_hip.nii.gz"  
scan_filepath <- tempfile(fileext = ".nii.gz")  
download.file(url, scan_filepath, mode = "wb")  
nifti <- import_scan(scan_filepath)
```

```
url2 <- "https://github.com/Telfer/BoneDensityMapping/releases/download/v1.0.2/test_CT_femur.stl"
bone_filepath <- tempfile(fileext = ".stl")
download.file(url2, bone_filepath, mode = "wb")
surface_mesh <- import_mesh(bone_filepath)
landmark_path <- system.file("extdata", "test_femur.mrk.json",
                             package = "BoneDensityMapping")
landmarks <- import_lmks(landmark_path)
mapped_coords <- surface_points_template(surface_mesh, landmarks,
                                       no_surface_sliders = 1000)
dens <- voxel_point_intersect(mapped_coords, nifti,
                              ct_eqn = "linear",
                              ct_params = c(68.4, 1.106))
colored_mesh <- color_mesh(surface_mesh, mapped_coords, dens)
```

ct_calibration

Sigma beta CT calculations

Description

Sigma beta CT calculations

Usage

```
ct_calibration(ct_nos, calibration_type, params)
```

Arguments

ct_nos Numeric vector. CT numbers from scan
calibration_type String. Currently only linear supported.
params Numeric vector. beta and sigma values for calibration eqn

Value

Vector with estimated density values in mg/cm³

Author(s)

Scott Telfer <scott.telfer@gmail.com>

fill_bone_points	<i>Fills bone with orthogonally spaced points for internal analysis</i>
------------------	---

Description

Fills bone with orthogonally spaced points for internal analysis

Usage

```
fill_bone_points(surface_mesh, spacing)
```

Arguments

surface_mesh	Mesh object
spacing	Numeric

Value

Matrix with internal point coordinates

Examples

```
url <- "https://github.com/Telfer/BoneDensityMapping/releases/download/v1.0.2/test_CT_femur.stl"
bone_filepath <- tempfile(fileext = ".stl")
download.file(url, bone_filepath, mode = "wb")
surface_mesh <- import_mesh(bone_filepath)
internal_fill <- fill_bone_points(surface_mesh, 2)
```

import_lmks	<i>import landmark coordinates</i>
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Description

import landmark coordinates

Usage

```
import_lmks(landmark_path, x = 1, y = 1, z = 1)
```

Arguments

landmark_path	String. File path to landmark data. .json or .fcsv format
x	Integer Value to apply to convert mesh i.e. -1 will mirror x coords
y	Integer Value to apply to convert mesh i.e. -1 will mirror y coords
z	Integer Value to apply to convert mesh i.e. -1 will mirror z coords

Value

dataframe. Columns are landmark name, x, y, and z coordinates

Author(s)

Scott Telfer <scott.telfer@gmail.com>

Examples

```
landmark_path <- system.file("extdata", "test_femur.mrk.json",  
                             package = "BoneDensityMapping")  
landmarks <- import_lmks(landmark_path)
```

import_mesh

import surface mesh

Description

import surface mesh

Usage

```
import_mesh(surface_mesh_filepath)
```

Arguments

surface_mesh_filepath
String. File path to bone models. .stl or .ply

Value

mesh object

Author(s)

Scott Telfer <scott.telfer@gmail.com>

Examples

```
# Download bone model  
url <- "https://github.com/Telfer/BoneDensityMapping/releases/download/v1.0.2/test_CT_femur.stl"  
bone_filepath <- tempfile(fileext = ".stl")  
download.file(url, bone_filepath, mode = "wb")  
import_mesh(bone_filepath)
```

import_scan	<i>import CT scan</i>
-------------	-----------------------

Description

import CT scan

Usage

```
import_scan(scan_filepath)
```

Arguments

scan_filepath String. File path to CT scan data. Should be .nii or .nrrd

Value

scan object

Author(s)

Scott Telfer <scott.telfer@gmail.com>

Examples

```
# Download CT scan
url <- "https://github.com/Telfer/BoneDensityMapping/releases/download/v1.0.2/test_CT_hip.nii.gz"
scan_filepath <- tempfile(fileext = ".nii.gz")
download.file(url, scan_filepath, mode = "wb")
import_scan(scan_filepath)
```

landmark_check	<i>Check landmarks are close to the mesh</i>
----------------	--

Description

Check landmarks are close to the mesh

Usage

```
landmark_check(surface_mesh, landmarks, threshold = 1)
```

Arguments

surface_mesh mesh object
 landmarks Dataframe. Columns are landmark name, x, y, and z coords
 threshold Numeric. Distance landmark can be from surface without warning being thrown

Value

String. Returns a message warning that landmarks are not on bone surface

Author(s)

Scott Telfer <scott.telfer@gmail.com>

Examples

```
# Download bone model
url <- "https://github.com/Telfer/BoneDensityMapping/releases/download/v1.0.2/test_CT_femur.stl"
bone_filepath <- tempfile(fileext = ".stl")
download.file(url, bone_filepath, mode = "wb")
surface_mesh <- import_mesh(bone_filepath)
landmark_path <- system.file("extdata", "test_femur.fcsv",
                             package = "BoneDensityMapping")
landmarks <- import_lmks(landmark_path)
landmark_check(surface_mesh, landmarks, threshold = 1.0)
```

plot_cross_section_bone

Plot Cross-Sectional Bone Visualization in 3D

Description

Visualizes a 3D cross-section of a bone using surface mesh and internal density (fill) points. Clips the surface mesh at a given axis and value, and overlays a 2D projection of internal density.

Usage

```
plot_cross_section_bone(
  surface_mesh,
  surface_colors = NULL,
  fill_coords,
  fill_colors,
  slice_axis,
  slice_val,
  slice_thickness = 1,
  IncludeSurface = FALSE,
  title = "Bone Cross-Section",
  userMat = NULL,
  legend = TRUE,
  legend_color_sel = NULL,
  legend_maxi = NULL,
  legend_mini = NULL
)
```

Arguments

surface_mesh	A 'mesh3d' object representing the outer surface of the bone.
surface_colors	Optional. A vector of colors for each vertex of the surface mesh. If NULL, uses mesh's own material colors.
fill_coords	A numeric matrix of internal fill point coordinates.
fill_colors	A vector of colors corresponding to fill points.
slice_axis	Character. 'x', 'y', or 'z'. Axis along which to slice.
slice_val	Numeric (0 to 1). Relative slice location along selected axis.
slice_thickness	Numeric. Width of the slice (default = 1).
IncludeSurface	Logical. Whether to include the clipped surface mesh.
title	Character. Title for the plot.
userMat	Optional. A 4x4 matrix controlling view orientation.
legend	Logical. Optional color bar.
legend_color_sel	Optional character with color gradient
legend_maxi	Numeric. Maximum bone density.
legend_mini	Numeric. Minimum bone density.

Value

Generates an 'rgl' plot

Examples

```
# Download CT scan
url <- "https://github.com/Telfer/BoneDensityMapping/releases/download/v1.0.1/test_CT_hip.nii.gz"
scan_filepath <- tempfile(fileext = ".nii.gz")
download.file(url, scan_filepath, mode = "wb")
nifti <- import_scan(scan_filepath)
url2 <- "https://github.com/Telfer/BoneDensityMapping/releases/download/v1.0.2/test_CT_femur.stl"
bone_filepath <- tempfile(fileext = ".stl")
download.file(url2, bone_filepath, mode = "wb")
surface_mesh <- import_mesh(bone_filepath)
landmark_path <- system.file("extdata", "test_femur.mrk.json",
                             package = "BoneDensityMapping")
landmarks <- import_lmks(landmark_path)
mapped_coords <- surface_points_template(surface_mesh, landmarks,
                                       no_surface_sliders = 100)
mat_peak <- voxel_point_intersect(mapped_coords, nifti)
colored_mesh <- color_mesh(surface_mesh, mapped_coords, mat_peak)
internal_fill <- fill_bone_points(surface_mesh, 3)
internal_density <- voxel_point_intersect(internal_fill, nifti,
                                          ct_eqn = "linear",
                                          ct_params = c(68.4, 1.106))
internal_colors <- color_mapping(internal_density)
```

```
plot_cross_section_bone(colored_mesh, surface_colors = NULL,
                        internal_fill, internal_colors, slice_axis = 'x',
                        slice_val = 0.5)
```

plot_mesh

plot mesh

Description

plot mesh

Usage

```
plot_mesh(
  surface_mesh,
  density_color = NULL,
  title = NULL,
  legend = TRUE,
  legend_color_sel = NULL,
  legend_maxi = 2000,
  legend_mini = 0,
  userMat = NULL
)
```

Arguments

surface_mesh	Mesh object
density_color	Vector. Colors mapped from density values.
title	String. Plot title.
legend	Logical. Optional color bar.
legend_color_sel	Optional character with color gradient
legend_maxi	Numeric. Maximum bone density.
legend_mini	Numeric. Minimum bone density.
userMat	Optional matrix. Controls graph orientation.

Value

plot of mesh with color

Author(s)

Scott Telfer <scott.telfer@gmail.com>

Examples

```

# Download CT scan
url <- "https://github.com/Telfer/BoneDensityMapping/releases/download/v1.0.1/test_CT_hip.nii.gz"
scan_filepath <- tempfile(fileext = ".nii.gz")
download.file(url, scan_filepath, mode = "wb")
nifti <- import_scan(scan_filepath)
url2 <- "https://github.com/Telfer/BoneDensityMapping/releases/download/v1.0.2/test_CT_femur.stl"
bone_filepath <- tempfile(fileext = ".stl")
download.file(url2, bone_filepath, mode = "wb")
surface_mesh <- import_mesh(bone_filepath)
vertices <- t(surface_mesh$vb)[, c(1:3)]
landmark_path <- system.file("extdata", "test_femur.mrk.json",
                             package = "BoneDensityMapping")
landmarks <- import_lmks(landmark_path)
mapped_coords <- surface_points_template(surface_mesh, landmarks,
                                         no_surface_sliders = 5000)
mat_peak <- surface_normal_intersect(surface_mesh, mapped_coords,
                                     normal_dist = 3.0, nifti, rev_y=FALSE)
color_mesh <- color_mesh(surface_mesh, mapped_coords, mat_peak, maxi = 2000,
                          mini = 0)
plot <- plot_mesh(color_mesh)

```

rm_local_sig	<i>local significance</i>
--------------	---------------------------

Description

local significance

Usage

```
rm_local_sig(vertices, sig_vals, changes, sig_level = 0.05, dist)
```

Arguments

vertices	Matrix
sig_vals	Numeric vector
changes	Numeric vector
sig_level	Numeric. Default 0.05
dist	Numeric. Distance to check for vertices

Value

Numeric vector

Author(s)

Scott Telfer <scott.telfer@gmail.com>

surface_normal_intersect

Find material properties of bone at surface point using surface normal

Description

Find material properties of bone at surface point using surface normal

Usage

```
surface_normal_intersect(
  surface_mesh,
  mapped_coords = NULL,
  normal_dist = 3,
  nifti,
  ct_eqn = NULL,
  ct_params = NULL,
  rev_x = FALSE,
  rev_y = FALSE,
  rev_z = FALSE,
  check_in_vol = FALSE
)
```

Arguments

surface_mesh	Mesh object
mapped_coords	Data frame. 3D coords of remapped surface points. If NULL, surface_mesh vertices will be used
normal_dist	Numeric. Distance surface normal should penetrate surface
nifti	Nifti CT scan image
ct_eqn	String. Equation to use for density calibration. Currently "linear" supported.
ct_params	Numeric vector. Calibration parameters for density calculation. For linear, first value is beta coefficient (y intercept), second value is sigma coefficient (gradient)
rev_x	Logical. Reverses x voxel coordinates
rev_y	Logical. Reverses y voxel coordinates
rev_z	Logical. Reverses z voxel coordinates
check_in_vol	Logical. Include check that model is in scans volume and print dimensions

Value

Vector. Vector with value for each point on surface

Author(s)

Scott Telfer <scott.telfer@gmail.com>

Examples

```
# Download CT scan
url <- "https://github.com/Telfer/BoneDensityMapping/releases/download/v1.0.2/test_CT_hip.nii.gz"
scan_filepath <- tempfile(fileext = ".nii.gz")
download.file(url, scan_filepath, mode = "wb")
nifti <- import_scan(scan_filepath)
url2 <- "https://github.com/Telfer/BoneDensityMapping/releases/download/v1.0.2/test_CT_femur.stl"
bone_filepath <- tempfile(fileext = ".stl")
download.file(url2, bone_filepath, mode = "wb")
surface_mesh <- import_mesh(bone_filepath)
landmark_path <- system.file("extdata", "test_femur.mrk.json",
                             package = "BoneDensityMapping")
landmarks <- import_lmks(landmark_path)
mapped_coords <- surface_points_template(surface_mesh, landmarks,
                                         no_surface_sliders = 1000)
mat_peak <- surface_normal_intersect(surface_mesh, normal_dist = 3.0,
                                     nifti = nifti, ct_eqn = "linear",
                                     ct_params = c(68.4, 1.106))
```

surface_points_new *New mapped surface points from template*

Description

New mapped surface points from template

Usage

```
surface_points_new(
  surface_mesh,
  landmarks,
  template,
  mirror = FALSE,
  plot_check = FALSE
)
```

Arguments

surface_mesh	List. Mesh data imported via ply_import function
landmarks	Data frame. Contains 3D coords of landmarks
template	Data frame. 3D coords of remapped surface points
mirror	Logical or character. Set to "x", "y", or "z" to mirror the mesh and landmarks across that axis before remapping.

`plot_check` Logical. If TRUE, generates a 3D plot showing the mirrored mesh, mirrored landmarks, remapped surface points, and original template points to visually verify correct orientation and laterality.

Value

Data frame. 3D coords of remapped surface points

Author(s)

Scott Telfer <scott.telfer@gmail.com> Adapted from geomorph

Examples

```
url <- "https://github.com/Telfer/BoneDensityMapping/releases/download/v1.0.2/SCAP001.stl"
bone_filepath <- tempfile(fileext = ".stl")
download.file(url, bone_filepath, mode = "wb")
scap_001_mesh <- import_mesh(bone_filepath)
landmark_path <- system.file("extdata", "SCAP001_landmarks.fcsv",
                             package = "BoneDensityMapping")
scap_001_lmk <- import_lmks(landmark_path)
template_coords <- surface_points_template(scap_001_mesh, scap_001_lmk,
                                           1000)

url2 <- "https://github.com/Telfer/BoneDensityMapping/releases/download/v1.0.2/SCAP002.stl"
bone_filepath <- tempfile(fileext = ".stl")
download.file(url2, bone_filepath, mode = "wb")
scap_002_mesh <- import_mesh(bone_filepath)
landmark_path <- system.file("extdata", "SCAP002_landmarks.fcsv",
                             package = "BoneDensityMapping")
scap_002_lmk <- import_lmks(landmark_path)
scap_002_remapped <- surface_points_new(scap_002_mesh, scap_002_lmk,
                                       template_coords, mirror = "x",
                                       plot_check = FALSE)
```

`surface_points_template`

Redefine surface points. Adds additional surface points ("sliders") that are spatially distributed across the mesh surface. Adapted from geomorph

Description

Redefine surface points. Adds additional surface points ("sliders") that are spatially distributed across the mesh surface. Adapted from geomorph

Usage

```
surface_points_template(surface_mesh, landmarks, no_surface_sliders)
```


Arguments

surface_mesh Mesh object
landmarks Data frame with landmark coordinates (columns: ID, x, y, z)
no_surface_sliders
 Numeric. No. of surface points to generate

Value

Data frame. 3D coordinates for the combined set of original landmarks and the new surface points

Author(s)

Scott Telfer <scott.telfer@gmail.com>

Examples

```
url <- "https://github.com/Telfer/BoneDensityMapping/releases/download/v1.0.2/test_CT_femur.stl"
bone_filepath <- tempfile(fileext = ".stl")
download.file(url, bone_filepath, mode = "wb")
surface_mesh <- import_mesh(bone_filepath)
landmark_path <- system.file("extdata", "test_femur.mrk.json",
                             package = "BoneDensityMapping")
landmarks <- import_lmks(landmark_path)
mapped_coords <- surface_points_template(surface_mesh, landmarks, 1000)
```

voxel_point_intersect *Finds material properties of bone at any point*

Description

Finds material properties of bone at any point

Usage

```
voxel_point_intersect(  
  vertex_coords,  
  nifti,  
  ct_eqn = NULL,  
  ct_params = NULL,  
  rev_x = FALSE,  
  rev_y = FALSE,  
  rev_z = FALSE,  
  check_in_vol = FALSE  
)
```


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