

# Package ‘aqp’

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**Depends** R (>= 2.15.0), methods

**Imports** plyr, reshape, grid, lattice, cluster, sp, Hmisc, stringr,  
scales, plotrix, RColorBrewer, MASS

**Suggests** colorspace, maptools, foreign, ape, soilDB, latticeExtra,  
digest, maps

**Description** A collection of algorithms related to modeling of soil resources, soil classification, soil profile aggregation, and visualization.

**License** GPL (>= 2)

**Repository** CRAN

**URL** <http://aqp.r-forge.r-project.org/>

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aqp-package

*Algorithms for Quantitative Pedology*

---

## Description

The aqp (Algorithms for Quantitative Pedology) package for R was developed to address some of the difficulties associated with processing soils information, specifically related to visualization, aggregation, and classification of soil profile data. This package is based on a mix of S3/S4 functions and classes, and most functions use basic dataframes as input, where rows represent soil horizons and columns define properties of those horizons. Common to most functions are the requirements that horizon boundaries are defined as depth from 0, and that profiles are uniquely defined by an id column. The aqp package defines an S4 class, "SoilProfileCollection", for storage of profile-level metadata, as well as summary, print, and plotting methods that have been customized for common tasks related to soils data.

Demos: `demo(aqp)`  
`demo(slope_effect_hz_thickness)`  
 Extended Examples: [AQP-related blog posts](#)  
[Introduction to SoilProfileCollection objects](#)  
[Soil profile dissimilarity demo](#)  
[Vertical vs. perpendicular horizon depth measurement demo](#)

**Author(s)**

Dylan E. Beaudette <debeaudette@ucdavis.edu>

**References**

<http://casoilresource.lawr.ucdavis.edu/>

**See Also**

[ca630](#), [sp1](#), [sp2](#), [sp3](#), [sp4](#), [sp5](#)

---

 addBracket

---

*Add Depth Brackets*


---

**Description**

Add depth-wise brackets to an existing plot of a SoilProfileCollection object.

**Usage**

```
addBracket(top, bottom, idx=NULL, tick.length = 0.05,
  arrow.length = 0.05, offset = -0.3, missing.bottom.depth = 25,
  ...)
```

**Arguments**

<code>top</code>	numeric vector of bracket top depths
<code>bottom</code>	numeric vector of bracket bottom depths
<code>idx</code>	optional integer index, associating bracket with profile
<code>tick.length</code>	length of bracket "tick" mark
<code>arrow.length</code>	length of arrowhead
<code>offset</code>	numeric, length of left-hand offset from each profile
<code>missing.bottom.depth</code>	distance (in depth units) to extend brackets that are missing a lower depth
<code>...</code>	further arguments passed on to segments or arrows

**Details**

The optional argument `idx` can be used to manually specify which profile a given bracket will be associated with. When `idx` is `NULL`, an integer sequence associated with plotting order (via `plotSPC`) is used. See examples below.

**Note**

This is a ‘low-level’ plotting function: you must first plot a `SoilProfileCollection` object before using this function.

**Author(s)**

D.E. Beaudette

**See Also**

[plotSPC](#)

**Examples**

```
# sample data
data(sp1)

# add color vector
sp1$soil_color <- with(sp1, munsell2rgb(hue, value, chroma))

# promote to SoilProfileCollection
depths(sp1) <- id ~ top + bottom

# plot profiles
plot(sp1)

# extract top/bottom depths associated with all A horizons
f <- function(i) {
  h <- horizons(i)
  idx <- grep('^A', h$name)
  c(min(h$top[idx]), max(h$bottom[idx], na.rm=TRUE))
}

# apply function to sp1, result is a list
a <- profileApply(sp1, f, simplify=FALSE)
# convert list into matrix
a <- do.call('rbind', a)

# plot
plot(sp1)
# annotate with brackets
# note that plotting order is derived from the call to `plot(sp1)`
addBracket(a[, 1], a[, 2], col='red')

# more interesting example using diagnostic horizons
```

```
if(require(soilDB)) {  
  # load some sample data with diagnostic horizons  
  data(loafercreek)  
  
  # extract first 15 profiles  
  x <- loafercreek[1:15, ]  
  s <- site(x)  
  
  # plot  
  par(mar=c(0,0,0,0))  
  plot(x, name='hzname', id.style='top')  
  
  # add brackets describing the argillic horizon  
  addDiagnosticBracket(x, 'argillic horizon', col='red')  
  # add brackets describing paralithic contact  
  addDiagnosticBracket(x, 'paralithic contact', col='blue')  
}
```

---

addVolumeFraction      *Symbolize Volume Fraction on a Soil Profile Collection Plot*

---

## Description

Symbolize volume fraction on an existing soil profile collection plot.

## Usage

```
addVolumeFraction(x, colname, res = 10, cex.min = 0.1,  
cex.max = 0.5, pch = 1, col = "black")
```

## Arguments

x	a SoilProfileCollection object
colname	character vector of length 1, naming the column containing volume fraction data (horizon-level attribute)
res	integer, resolution of the grid used to symbolize volume fraction
cex.min	minimum symbol size
cex.max	maximum symbol size
pch	plotting character
col	color of the symbol

## Details

This function can only be called after plotting a SoilProfileCollection object.

**Note**

Details associated with a call to `plot.SoilProfileCollection` are automatically accounted for within this function: e.g. `plot.order`, `width`, etc.

**Author(s)**

D.E. Beaudette

**See Also**

[plotSPC](#)

**Examples**

```
# sample data
data(loafercreek, package='soilDB')

# subset first 10 profiles
s <- loafercreek[1:10, ]

# replace 0% frags with NA
s$total_frags_pct[which(s$total_frags_pct == 0)] <- NA

# plot in random order, note that annotations follow
par(mar=c(0, 0, 3, 0))
plot(s, color='total_frags_pct', plot.order=sample(1:length(s)))
addVolumeFraction(s, 'total_frags_pct', pch=1)

par(mar=c(0, 0, 0, 0))
plot(s, max.depth=100, name='total_frags_pct', cex.name=1, axis.line.offset=-4.25)
addVolumeFraction(s, 'total_frags_pct', pch=1)
```

---

amarillo

*Amarillo Soils*

---

**Description**

This sample dataset contains laboratory and published soil survey (i.e. generalized) data on the Amarillo soil series.

**Usage**

```
data(amarillo)
```

**Format**

The format is: List of 2 \$ lab:'data.frame': 323 obs. of 40 variables: ..\$ ID : int [1:323] 1 2 3 4 5 6 7 8 9 10 ... ..\$ user\_pedon\_id : chr [1:323] "53TX305059" "53TX305059" "53TX305059" "53TX305059" ... ..\$ user\_site\_id : chr [1:323] "53TX305059" "53TX305059" "53TX305059" "53TX305059" ... ..\$ horizontal\_datum\_name : chr [1:323] "" "" "" "" ... ..\$ latitude\_direction : chr [1:323] "north" "north" "north" "north" ... ..\$ latitude\_degrees : int [1:323] 33 33 33 33 33 33 33 33 33 33 ... ..\$ latitude\_minutes : int [1:323] 10 10 10 10 10 10 11 11 11 11 ... ..\$ latitude\_seconds : num [1:323] 30 30 30 30 30 30 5 5 5 5 ... ..\$ longitude\_direction : chr [1:323] "west" "west" "west" "west" ... ..\$ longitude\_degrees : int [1:323] 101 101 101 101 101 101 101 101 101 101 ... ..\$ longitude\_minutes : int [1:323] 47 47 47 47 47 47 46 46 46 46 ... ..\$ longitude\_seconds : num [1:323] 40 40 40 40 40 40 48 48 48 48 ... ..\$ latitude\_std\_decimal\_degrees : num [1:323] 33.2 33.2 33.2 33.2 33.2 ... ..\$ longitude\_std\_decimal\_degrees: num [1:323] -102 -102 -102 -102 -102 ... ..\$ taxon\_name : chr [1:323] "Amarillo" "Amarillo" "Amarillo" "Amarillo" ... ..\$ class\_type : chr [1:323] "correlated" "correlated" "correlated" "correlated" ... ..\$ natural\_key : chr [1:323] "40A34174" "40A34175" "40A34176" "40A34177" ... ..\$ hzn\_desgn : chr [1:323] "Ap" "B1" "Bt1" "Bt2" ... ..\$ hzn\_top : int [1:323] 0 18 51 86 137 190 0 20 51 76 ... ..\$ hzn\_bot : int [1:323] 18 51 86 137 190 236 20 51 76 102 ... ..\$ CEC : num [1:323] NA NA NA NA NA NA NA NA NA NA NA ... ..\$ ECEC : logi [1:323] NA NA NA NA NA NA NA ... ..\$ ph\_h2o : num [1:323] 7.5 7.2 7.5 8 8.4 8.4 NA NA NA NA ... ..\$ ph\_cacl2 : num [1:323] NA NA NA NA NA NA NA NA NA NA NA ... ..\$ caco3 : int [1:323] NA NA NA 8 53 38 NA NA NA NA ... ..\$ c\_gyp12 : int [1:323] NA NA NA NA NA NA NA NA NA NA NA ... ..\$ EC : num [1:323] NA NA NA NA NA NA NA NA NA NA NA ... ..\$ SAR : int [1:323] NA NA NA NA NA NA NA NA NA NA NA ... ..\$ Total.Clay : num [1:323] 17.9 19.5 25.2 29.2 37.8 32.6 10.8 26.8 25.5 30.8 ... ..\$ Total.Silt : num [1:323] 9.4 9.4 14.7 18.7 26.8 25.5 10.5 13.6 16.6 15.3 ... ..\$ Total.Sand : num [1:323] 72.7 71.1 60.1 52.1 35.4 41.9 78.7 59.6 57.9 53.9 ... ..\$ dB.33.bar : num [1:323] NA NA NA NA NA NA NA NA NA 1.2 NA ... ..\$ dB.15.bar : num [1:323] NA NA NA NA NA NA NA NA NA NA NA ... ..\$ LE : num [1:323] NA NA NA NA NA NA NA NA NA NA NA ... ..\$ Water.Ret..2mm.33.bar : num [1:323] NA NA NA NA NA NA NA NA NA NA NA ... ..\$ Water.Ret.clod.33.bar : num [1:323] NA NA NA NA NA NA NA NA NA 36.6 NA ... ..\$ Water.Ret..2mm.15.bar : num [1:323] NA NA NA NA NA NA NA NA NA NA NA ... ..\$ OC\_lab : num [1:323] NA NA NA NA NA NA NA NA NA NA NA ... ..\$ OM : num [1:323] NA NA NA NA NA NA NA NA NA NA NA ... ..\$ model\_desg : chr [1:323] "Ap" "B" "Bt" "Bt" ... \$ dmu:'data.frame': 383 obs. of 88 variables: ..\$ ID : int [1:383] 8177 8178 8179 8180 8181 8182 8183 8184 8185 8186 ... ..\$ State : chr [1:383] "NM" "NM" "NM" "NM" ... ..\$ Database : chr [1:383] "SSURGO" "SSURGO" "SSURGO" "SSURGO" ... ..\$ Area.Symbol : chr [1:383] "NM021" "NM021" "NM021" "NM021" ... ..\$ Area.Name : chr [1:383] "Harding County, New Mexico" "Harding County, New Mexico" "Harding County, New Mexico" "Harding County, New Mexico" ... ..\$ mukey : int [1:383] 376313 376313 376313 376379 376379 376379 376380 376380 376380 376405 ... ..\$ Mapunit.Symbol : chr [1:383] "AM" "AM" "AM" "SR" ... ..\$ Component.Name : chr [1:383] "Amarillo" "Amarillo" "Amarillo" "Amarillo" ... ..\$ SIR....obsolete: chr [1:383] "TX0130" "TX0130" "TX0130" "TX0130" ... ..\$ cokey : int [1:383] 504646 504646 504646 504918 504918 504918 504923 504923 504923 505030 ... ..\$ RV : int [1:383] 85 85 85 40 40 40 40 40 40 85 ... ..\$ Local.Phase : chr [1:383] "" "" "" "" ... ..\$ chkey : int [1:383] 1113073 1113074 1113075 1113731 1113732 1113733 1113743 1113744 1113745 1114027 ... ..\$ Designation : chr [1:383] "H1" "H2" "H3" "H1" ... ..\$ top : int [1:383] 0 13 122 0 20 122 0 13 122 0 ... ..\$ bott : int [1:383] 13 122 165 20 122 152 13 122 152 20 ... ..\$ CECL : num [1:383] 5 8 8 3 8 8 3 8 8 15 ... ..\$ CECR : num [1:383] 10 14 14 6.5 14 14 6.5 14 14 20 ... ..\$ S : int [1:383] 3 3 3 3 3 3 3 3 3 3 ... ..\$ CECH : num [1:383] 15 20 20 10 20 20 10 20 20 25 ... ..\$ cec\_mean : num [1:383] 10 14 14 6.5 14 14 6.5 14 14 20 ... ..\$ ECECL : num [1:383] NA NA NA

NA NA NA NA NA NA NA NA ... ..\$ ECECR : num [1:383] NA NA NA NA NA NA NA NA NA NA NA NA  
 ... ..\$ S1 : int [1:383] NA NA NA NA NA NA NA NA NA NA NA NA ... ..\$ ECECH : num [1:383] NA NA  
 NA NA NA NA NA NA NA NA NA NA ... ..\$ PHWL : num [1:383] 6.6 7.4 7.9 6.6 7.4 7.9 6.6 7.4 7.9 6.6  
 ... ..\$ PHWR : num [1:383] 7.2 7.9 8.2 7.2 7.9 8.2 7.2 7.9 8.2 7 ... ..\$ PHWH : num [1:383] 7.8 8.4  
 8.4 7.8 8.4 8.4 7.8 8.4 8.4 7.3 ... ..\$ PHWM : num [1:383] 7.2 7.9 8.15 7.2 7.9 8.15 7.2 7.9 8.15 6.95  
 ... ..\$ PHCL : num [1:383] NA NA NA NA NA NA NA NA NA NA NA NA ... ..\$ PHCR : num [1:383] NA  
 NA NA NA NA NA NA NA NA NA NA ... ..\$ PHCH : num [1:383] NA NA NA NA NA NA NA NA NA NA  
 NA NA ... ..\$ CACOL : int [1:383] 0 0 5 0 0 5 0 0 5 0 ... ..\$ CACOR : int [1:383] 3 3 7 3 3 7 3 5 7 0  
 ... ..\$ CACOH : int [1:383] 5 5 10 5 5 10 5 7 10 1 ... ..\$ CACOM : num [1:383] 2.5 2.5 7.5 2.5 2.5  
 7.5 2.5 3.5 7.5 0.5 ... ..\$ GYPL : int [1:383] 0 0 0 0 0 0 0 0 0 0 ... ..\$ GYPR : int [1:383] 0 0 0 0 0 0  
 0 0 0 0 ... ..\$ GYPH : int [1:383] 0 0 0 0 0 0 0 0 1 ... ..\$ GYPM : num [1:383] 0 0 0 0 0 0 0 0 0.5  
 ... ..\$ ECL : int [1:383] 0 0 0 0 0 0 0 0 0 ... ..\$ ECR : num [1:383] 1 1 1 1 1 1 1 1 0 ... ..\$ ECH  
 : num [1:383] 2 2 2 2 2 2 2 2 2 ... ..\$ ECM : num [1:383] 1 1 1 1 1 1 1 1 1 ... ..\$ SARL : int  
 [1:383] 0 0 0 0 0 0 0 0 0 ... ..\$ SARR : int [1:383] 0 0 0 1 1 1 0 0 0 ... ..\$ SARH : int [1:383] 1 1  
 1 2 2 2 1 1 1 2 ... ..\$ SARM : num [1:383] 0.5 0.5 0.5 1 1 1 0.5 0.5 0.5 1 ... ..\$ CLAYL : int [1:383]  
 10 20 20 5 20 20 5 20 20 13 ... ..\$ CLAYR : num [1:383] 14 27.5 27.5 10 27.5 27.5 10 27.5 27.5  
 21.5 ... ..\$ CLAYH : int [1:383] 18 35 35 15 35 35 15 35 35 30 ... ..\$ CLAYM : num [1:383] 14  
 27.5 27.5 10 27.5 27.5 10 27.5 27.5 21.5 ... ..\$ SILL : num [1:383] NA NA NA NA NA NA NA NA NA  
 NA 30 ... ..\$ SILR : num [1:383] 16.4 17.4 17.4 9.2 37.8 37.8 9.2 37.8 37.8 37.1 ... ..\$ SILH : num  
 [1:383] NA NA NA NA NA NA NA NA NA NA 45 ... ..\$ SILM : num [1:383] NA NA NA NA NA NA NA NA NA  
 NA NA NA 37.5 ... ..\$ SANL : num [1:383] NA NA NA NA NA NA NA NA NA NA 35 ... ..\$ SANR  
 : num [1:383] 69.6 55.1 55.1 80.8 34.7 34.7 80.8 34.7 34.7 41.4 ... ..\$ SANH : int [1:383] NA NA  
 NA NA NA NA NA NA NA NA 50 ... ..\$ SANM : num [1:383] NA NA NA NA NA NA NA NA NA NA  
 42.5 ... ..\$ S2 : int [1:383] 1 1 1 1 1 1 1 1 1 ... ..\$ DB3L : num [1:383] 1.35 1.3 1.4 1.4 1.3 1.4  
 1.4 1.3 1.4 1.3 ... ..\$ DB3R : num [1:383] 1.48 1.48 1.6 1.5 1.48 1.6 1.5 1.48 1.6 1.43 ... ..\$ DB3H  
 : num [1:383] 1.6 1.65 1.8 1.6 1.65 1.8 1.6 1.65 1.8 1.55 ... ..\$ DB3M : num [1:383] 1.47 1.47 1.6  
 1.5 1.47 1.6 1.5 1.47 1.6 1.42 ... ..\$ S3 : int [1:383] 3 3 3 3 3 3 3 3 3 ... ..\$ KSATL : num [1:383]  
 14.11 4.23 4.23 14.11 4.23 ... ..\$ KSATR : num [1:383] 28.23 9.17 9.17 28.23 9.17 ... ..\$ KSATH  
 : num [1:383] 42.3 14.1 14.1 42.3 14.1 ... ..\$ KSATM : num [1:383] 28.22 9.17 9.17 28.22 9.17 ...  
 ..\$ AWCL : num [1:383] 0.11 0.14 0.1 0.06 0.14 0.1 0.06 0.14 0.1 0.12 ... ..\$ AWCR : num [1:383]  
 0.13 0.16 0.13 0.08 0.16 0.13 0.08 0.16 0.13 0.15 ... ..\$ AWCH : num [1:383] 0.15 0.18 0.15 0.1  
 0.18 0.15 0.1 0.18 0.15 0.18 ... ..\$ AWCM : num [1:383] 0.13 0.16 0.12 0.08 0.16 0.12 0.08 0.16  
 0.12 0.15 ... ..\$ LEPL : num [1:383] 0 0 0 0 0 0 0 0 0 ... ..\$ LEPR : num [1:383] 1.5 1.5 1.5 1.5  
 1.5 1.5 1.5 1.5 1.5 1.5 ... ..\$ LEPH : num [1:383] 2.9 2.9 2.9 2.9 2.9 2.9 2.9 2.9 2.9 ... ..\$ LEPM  
 : num [1:383] 1.45 1.45 1.45 1.45 1.45 1.45 1.45 1.45 1.45 ... ..\$ OML : num [1:383] 0.5 0.2  
 0.1 0.5 0.2 0.1 0.5 0.2 0.1 1 ... ..\$ OMR : num [1:383] 0.75 0.3 0.2 0.75 0.3 0.2 0.75 0.3 0.2 1.5 ...  
 ..\$ OMH : num [1:383] 1 0.5 0.3 1 0.5 0.3 1 0.5 0.3 2 ... ..\$ OMM : num [1:383] 0.75 0.35 0.2 0.75  
 0.35 0.2 0.75 0.35 0.2 1.5 ... ..\$ DB15L : logi [1:383] NA NA NA NA NA NA NA NA ... ..\$ DB15R : num  
 [1:383] NA NA NA NA NA NA NA NA NA NA NA NA ... ..\$ DB15H : logi [1:383] NA NA NA NA NA NA  
 NA NA ... ..\$ DB15M : int [1:383] 0 0 0 0 0 0 0 0 0 ... ..\$ S4 : int [1:383] NA NA NA NA NA NA NA  
 NA NA NA NA NA ... ..\$ model\_desg : chr [1:383] "" "" "" "" "" ...

**Source**

USDA-NRCSS SSURGO database, and the KSSL database; c/o Skye Wills.

**Examples**

data(amarillo)



```
## maybe str(amarillo) ; plot(amarillo) ...
```

---

 ca630

*Soil Data from the Central Sierra Nevada Region of California*


---

## Description

Site and laboratory data from soils sampled in the central Sierra Nevada Region of California.

## Usage

```
data(ca630)
```

## Format

List containing:

\$site : A data frame containing site information.

user\_site\_id national user site id

mlra the MLRA

county the county

ssa soil survey area

lon longitude, WGS84

lat latitude, WGS84

pedon\_key national soil profile id

user\_pedon\_id local soil profile id

cntrl\_depth\_to\_top control section top depth (cm)

cntrl\_depth\_to\_bot control section bottom depth (cm)

sampled\_taxon\_name soil series name

\$lab : A data frame containing horizon information.

pedon\_key national soil profile id

layer\_key national horizon id

layer\_sequence horizon sequence number

hzn\_top horizon top (cm)

hzn\_bot horizon bottom (cm)

hzn\_desgn horizon name

texture\_description USDA soil texture

nh4\_sum\_bases sum of bases extracted by ammonium acetate (pH 7)

ex\_acid exchangeable acidity [method ?]

CEC8.2 cation exchange capacity by sum of cations method (pH 8.2)

CEC7 cation exchange capacity by ammonium acetate (pH 7)

bs\_8.2 base saturation by sum of cations method (pH 8.2)

bs\_7 base saturation by ammonium acetate (pH 7)

## Details

These data were extracted from the NSSL database. 'ca630' is a list composed of site and lab data, each stored as dataframes. These data are modeled by a 1:many (site:lab) relation, with the 'pedon\_id' acting as the primary key in the 'site' table and as the foreign key in the 'lab' table.

## Source

<http://ssldata.nrcs.usda.gov/>

## Examples

```
library(plyr)
library(lattice)
library(Hmisc)
library(maps)
library(sp)

# check the data out:
data(ca630)
str(ca630)

# note that pedon_key is the link between the two tables

# make a copy of the horizon data
ca <- ca630$lab

# promote to a SoilProfileCollection class object
depths(ca) <- pedon_key ~ hzn_top + hzn_bot

# add site data, based on pedon_key
site(ca) <- ca630$site

# ID data missing coordinates: '|' is a logical OR
(missing.coords.idx <- which(is.na(ca$lat) | is.na(ca$lon)))

# remove missing coordinates by safely subsetting
if(length(missing.coords.idx) > 0)
ca <- ca[-missing.coords.idx, ]

# register spatial data
coordinates(ca) <- ~ lon + lat

# assign a coordinate reference system
proj4string(ca) <- '+proj=longlat +datum=NAD83'

# check the result
print(ca)

# map the data (several ways to do this, here is a simple way)
map(database='county', region='california')
points(coordinates(ca), col='red', cex=0.5)
```

```

# aggregate %BS 7 for all profiles into 1 cm slices
a <- slab(ca, fm= ~ bs_7)

# plot median & IQR by 1 cm slice
xyplot(
top ~ p.q50, data=a, lower=a$p.q25, upper=a$p.q75,
ylim=c(160,-5), alpha=0.5, scales=list(alternating=1, y=list(tick.num=7)),
panel=panel.depth_function, prepanel=prepanel.depth_function,
ylab='Depth (cm)', xlab='Base Saturation at pH 7',
par.settings=list(superpose.line=list(col='black', lwd=2))
)

# aggregate %BS at pH 8.2 for all profiles by MLRA, along 1 cm slices
# note that mlra is stored in @site
a <- slab(ca, mlra ~ bs_8.2)

# keep only MLRA 18 and 22
a <- subset(a, subset=mlra %in% c('18', '22'))

# plot median & IQR by 1 cm slice, using different colors for each MLRA
xyplot(
top ~ p.q50, groups=mlra , data=a, lower=a$p.q25, upper=a$p.q75,
ylim=c(160,-5), alpha=0.5, scales=list(y=list(tick.num=7, alternating=3), x=list(alternating=1)),
panel=panel.depth_function, prepanel=prepanel.depth_function,
ylab='Depth (cm)', xlab='Base Saturation at pH 8.2',
par.settings=list(superpose.line=list(col=c('black','blue'), lty=c(1,2), lwd=2)),
auto.key=list(columns=2, title='MLRA', points=FALSE, lines=TRUE)
)

# safely compute hz-thickness weighted mean CEC (pH 7)
# using data.frame objects
head(lab.agg.cec_7 <- ddply(ca630$lab, .(pedon_key),
.fun=summarise, CEC_7=wtd.mean(bs_7, weights=hzn_bot-hzn_top)))

# extract a SPDF with horizon data along a slice at 25 cm
s.25 <- slice(ca, fm=25 ~ bs_7 + CEC7 + ex_acid)
splot(s.25, zcol=c('bs_7','CEC7','ex_acid'))

# note that the ordering is preserved:
all.equal(s.25$pedon_key, profile_id(ca))

# extract a data.frame with horizon data at 10, 20, and 50 cm
s.multiple <- slice(ca, fm=c(10,20,50) ~ bs_7 + CEC7 + ex_acid)

# Extract the 2nd horizon from all profiles as SPDF
ca.2 <- ca[, 2]

# subset profiles 1 through 10
ca.1.to.10 <- ca[1:10, ]

# basic plot method: profile plot
plot(ca.1.to.10, name='hzn_desgn')

```

---

estimateSoilDepth      *Estimate Soil Depth*

---

### Description

Estimate the soil depth of a single profile within a SoilProfileCollection object.

### Usage

```
estimateSoilDepth(f, name = "hzname", top = "hzdept", bottom = "hzdepb", p = 'Cr|R|Cd',
  no.contact.depth = NULL, no.contact.assigned = NULL)
```

### Arguments

f	A SoilProfileCollection object of length 1, e.g. a single profile
name	the name of the column that contains horizon designations
top	the name of the column that contains horizon top depths
bottom	the name of the column that contains horizon bottom depths
p	a REGEX pattern for determining contact with bedrock
no.contact.depth	in the absence of contact with bedrock, a depth at which we can assume a standard depth
no.contact.assigned	assumed standard depth

### Value

a single integer, the soil depth

### Author(s)

D.E. Beaudette and J.M. Skovlin

### See Also

[getSoilDepthClass](#), [profileApply](#)

### Examples

```
data(sp1)
depths(sp1) <- id ~ top + bottom

# apply to each profile in a collection, and save as site-level attribute
sp1$depth <- profileApply(sp1, estimateSoilDepth, name='name', top='top', bottom='bottom')

## Not run:
# sample data
```

```

data(gopheridge, package='soilDB')

# run on a single profile
estimateSoilDepth(gopheridge[1, ])

# apply to an entire collection
profileApply(gopheridge, estimateSoilDepth)

## End(Not run)

```

---

evalGenHZ

*Evaluate Generalized Horizon Labels*


---

## Description

Data-driven evaluation of generalized horizon labels using nMDS and silhouette width.

## Usage

```

evalGenHZ(obj, genhz, vars, non.matching.code = "not-used",
stand = TRUE, trace = FALSE, metric = "euclidean")

```

## Arguments

obj	a SoilProfileCollection object
genhz	name of horizon-level attribute containing generalized horizon labels
vars	character vector of horizon-level attributes to include in the evaluation
non.matching.code	code used to represent horizons not assigned a generalized horizon label
stand	standardize variables before computing distance matrix (default = TRUE), passed to <a href="#">daisy</a>
trace	verbose output from passed to <a href="#">isoMDS</a> , (default = FALSE)
metric	distance metric, passed to <a href="#">daisy</a>

## Details

Non-metric multidimensional scaling is performed via [isoMDS](#). The input distance matrix is generated by [daisy](#) using (complete cases of) horizon-level attributes from obj as named in vars.

Silhouette widths are computed via [silhouette](#). The input distance matrix is generated by [daisy](#) using (complete cases of) horizon-level attributes from obj as named in vars. Note that observations with genhz labels specified in non.matching.code are removed filtered before calculation of the distance matrix.

**Value**

a list is returned containing:

**horizons** c('mds.1', 'mds.2', 'sil.width', 'neighbor')

**stats** mean and standard deviation of vars, computed by generalized horizon label

**dist** the distance matrix as passed to [isoMDS](#)

**Author(s)**

D.E. Beaudette

**See Also**

[get.ml.hz](#)

---

f.noise

*Example Objective Function for Full-Pattern Matching*

---

**Description**

Basic objective function that can be used as a starting point for developing XRD full-pattern matching strategies. [details pending...]

**Usage**

```
f.noise(inits, pure.patterns, sample.pattern, eps.total = 0.05)
```

**Arguments**

inits	vector of initial guesses for mineral fractions, last item is a noise component
pure.patterns	a matrix of XRD patterns of pure samples, resampled to the same twotheta resolution and rescaled according to an external standard
sample.pattern	the unknown or composite pattern, aligned to the same twotheta axis as the pure patterns and rescaled to an external standard
eps.total	precision of comparisons; currently not used

**Details**

This is similar to the work of Chipera and Bish (2002), using the methods described in (Bish, 1994). If the flexibility of a custom objective function is not required, the linear model framework should be sufficient for pattern fitting. GLS should be used if realistic standard errors are needed.

**Value**

the sum of absolute differences between the unknown pattern and combination of pure patterns for the current set of mixture proportions

**Author(s)**

Dylan E. Beaudette

**References**

Chipera, S.J., & Bish, D.L. (2002) FULLPAT: A full-pattern quantitative analysis program for X-ray powder diffraction using measured and calculated patterns. *J. Applied Crystallography*, 35, 744-749.

Bish, D. 1994. Quantitative Methods in Soil Mineralogy, in *Quantitative X-Ray Diffraction Analysis of Soil*. Amonette, J. & Zelazny, L. (ed.) Soil Science Society of America, pp 267-295.

**See Also**

[resample.twotheta](#)

**Examples**

```
# sample data
data(rruff.sample)

# get number of measurements
n <- nrow(rruff.sample)

# number of components
n.components <- 6

# mineral fractions, normally we don't know these
w <- c(0.346, 0.232, 0.153, 0.096, 0.049, 0.065)

# make synthetic combined pattern
# scale the pure substances by the known proportions
rruff.sample$synthetic_pat <- apply(sweep(rruff.sample[,2:7], 2, w, '*'), 1, sum)

# add 1 more substance that will be unknown to the fitting process
rruff.sample$synthetic_pat <- rruff.sample$synthetic_pat +
(1 - sum(w)) * rruff.sample[,8]

# try adding some nasty noise
# rruff.sample$synthetic_pat <- apply(sweep(rruff.sample[,2:7], 2, w, '*'), 1, sum) +
# runif(n, min=0, max=100)

# look at components and combined pattern
par(mfcol=c(7,1), mar=c(0,0,0,0))
plot(1:n, rruff.sample$synthetic_pat, type='l', axes=FALSE)
legend('topright', bty='n', legend='combined pattern', cex=2)
for(i in 2:7)
{
plot(1:n, rruff.sample[, i], type='l', axes=FALSE)
```

```

legend('topright', bty='n',
legend=paste(names(rruff.sample)[i], '(', w[i-1], ')', sep=''), cex=2)
}

## fit pattern mixtures with a linear model
l <- lm(synthetic_pat ~ nontronite + montmorillonite + clinochlore
+ antigorite + chamosite + hematite, data=rruff.sample)

summary(l)

par(mfcol=c(2,1), mar=c(0,3,0,0))
plot(1:n, rruff.sample$synthetic_pat, type='l', lwd=2, lty=2, axes=FALSE,
xlab='', ylab='')
lines(1:n, predict(l), col=2)
axis(2, cex.axis=0.75, las=2)
legend('topright', legend=c('original','fitted'), col=c(1,2), lty=c(2,1),
lwd=c(2,1), bty='n', cex=1.25)

plot(1:n, resid(l), type='l', axes=FALSE, xlab='', ylab='', col='blue')
abline(h=0, col=grey(0.5), lty=2)
axis(2, cex.axis=0.75, las=2)
legend('topright', legend=c('residuals'), bty='n', cex=1.25)

## fitting by minimizing an objective function (not run)

# SANN is a slower algorithm, sometimes gives strange results
# default Nelder-Mead is most robust
# CG is fastest --> 2.5 minutes max
# component proportions (fractions), and noise component (intensity units)
# initial guesses may affect the stability / time of the fit

## this takes a while to run
# # synthetic pattern
# o <- optim(par=c(0.1, 0.1, 0.1, 0.1, 0.1, 0.1, 0.1), f.noise,
# method='CG', pure.patterns=rruff.sample[,2:7],
# sample.pattern=rruff.sample$synthetic_pat)
#
#
# # estimated mixture proportions
# o$par
#
# # compare with starting proportions
# rbind(o$par[1:n.components], w)
#
# # if we had an unknown pattern we were trying to match, compare fitted here
# # compute R value 0.1 - 0.2 considered good
# # sum(D^2) / sum(s)

```



```

# # o$value / sum(rruff.sample$sample)
#
# # plot estimated mixture vs sample
# # combine pure substances
# pure.mixture <- apply(sweep(rruff.sample[, 2:7], 2, o$par[1:n.components], '*'), 1, sum)
#
# # add in noise
# noise.component <- o$par[n.components+1]
# est.pattern <- pure.mixture + noise.component
#
#
# # plot results
# par(mfcol=c(2,1), mar=c(0,3,0,0))
# plot(1:n, rruff.sample$synthetic_pat, type='l', lwd=2, lty=2, axes=FALSE,
# xlab='', ylab='')
# lines(1:n, est.pattern, col=2)
# lines(1:n, rep(noise.component, n), col=3)
# axis(2, cex.axis=0.75, las=2)
# legend('topright', legend=c('original', 'fitted', 'noise'), col=c(1,2,3), lty=c(2,1,1),
# lwd=c(2,1,1), bty='n', cex=1.25)
#
# plot(1:n, rruff.sample$synthetic_pat - est.pattern, type='l', axes=FALSE,
# xlab='', ylab='')
# abline(h=0, col=grey(0.5), lty=2)
# axis(2, cex.axis=0.75, las=2)
# legend('topright', legend=c('difference'), bty='n', cex=1.25)
#

```

---

generalize.hz

*Generalize Horizon Names*


---

## Description

Generalize a vector of horizon names, based on new classes, and REGEX patterns.

## Usage

```
generalize.hz(x, new, pat, non.matching.code)
```

## Arguments

x	a character vector of horizon names
new	a character vector of new horizon classes
pat	a character vector of REGEX, same length as x
non.matching.code	text used to describe any horizon not matching any item in pat

**Value**

factor of the same length as x

**Author(s)**

Dylan E. Beaudette

**References**

<http://casoilresource.lawr.ucdavis.edu/>

**Examples**

```
data(sp1)

# check original distribution of hz designations
table(sp1$name)

# generalize
sp1$genhz <- generalize.hz(sp1$name,
  new=c('O', 'A', 'B', 'C', 'R'),
  pat=c('O', '^A', '^B', 'C', 'R'))

# see how we did / what we missed
table(sp1$genhz, sp1$name)
```

---

get.ml.hz

*Determine ML Horizon Boundaries*

---

**Description**

This function accepts input from `slab()` along with a vector of horizon names, and returns a data.frame of the most likely horizon boundaries.

**Usage**

```
get.ml.hz(x, o.names)
```

**Arguments**

x	output from <a href="#">slab</a>
o.names	an optional character vector of horizon designations that will be used in the final table

**Details**

This function is expecting that x is a data.frame generated by [slab](#). If x was not generated by `slab`, then o.names is required.

**Value**

A dataframe with the following columns:

hz	horizon names
top	top boundary
bottom	bottom boundary
confidence	integrated probability / ML horizon thickness, rounded to the nearest integer
pseudo.brier	A "pseudo" Brier Score for a multi-class prediction, where the most-likely horizon label is treated as the "correct" outcome. Details on the calculation for traditional Brier Scores here: <a href="http://en.wikipedia.org/wiki/Brier_score#Original_definition_by_Brier">http://en.wikipedia.org/wiki/Brier_score#Original_definition_by_Brier</a> .

**Author(s)**

D.E. Beaudette

**See Also**

[slab](#)

**Examples**

```
data(sp1)
depths(sp1) <- id ~ top + bottom

# normalize horizon names: result is a factor
sp1$name <- generalize.hz(sp1$name,
  new=c('O', 'A', 'B', 'C'),
  pat=c('O', '^A', '^B', 'C'))

# compute slice-wise probability so that it sums to contributing fraction, from 0-150
a <- slab(sp1, fm= ~ name, cpm=1, slab.structure=0:150)

# generate table of ML horization
get.ml.hz(a, o.names=c('O', 'A', 'B', 'C'))
```

---

getSoilDepthClass      *Generate Soil Depth Class Matrix*

---

**Description**

Generate a boolean matrix of soil depth classes from a SoilProfileCollection object.

**Usage**

```
getSoilDepthClass(f, depth.classes = c(very.shallow = 25,
  shallow = 50, mod.deep = 100, deep = 150, very.deep = 1000), ...)
```

**Arguments**

f a SoilProfileCollection object  
depth.classes a named vector of classes and depth breaks  
... arguments passed to [estimateSoilDepth](#)

**Value**

a data.frame containing soil depth and depth class for each profile, see examples

**Author(s)**

D.E. Beaudette and J.M. Skovlin

**See Also**

[estimateSoilDepth](#)

**Examples**

```
data(sp1)
depths(sp1) <- id ~ top + bottom

# generate depth-class matrix
sdc <- getSoilDepthClass(sp1, name='name', top='top', bottom='bottom')

# inspect
head(sdc)

# join back into sp1 as site-level data
site(sp1) <- sdc

## Not run:
# sample data
data(gopheridge, package='soilDB')

getSoilDepthClass(gopheridge)

## End(Not run)
```

---

hzDistinctnessCodeToOffset

*Convert Horizon Distinctness Codes*

---

**Description**

This function accepts a vector of horizon distinctness codes, a look-up vector of codes, and a corresponding vector of vertical offsets. The defaults are based on USDA-NCSS field methods.

**Usage**

```
hzDistinctnessCodeToOffset(x, codes = c("A", "C", "G", "D"), offset = c(0.5, 1.5, 5, 10))
```

**Arguments**

x	vector of distinctness codes to be converted to offsets
codes	vector of unique distinctness codes
offset	vector of offsets associated with distinctness codes

**Details**

Missing data (NA) or codes that are not defined in the 'codes' argument are returned as 0 offsets.

**Value**

a vector of vertical offsets, with the same length as the number of distinctness codes passed to the function

**Author(s)**

D.E. Beaudette

**References**

<http://soils.usda.gov/technical/fieldbook/>

**See Also**

[plotSPC](#)

**Examples**

```
data(sp1)
hzDistinctnessCodeToOffset(sp1$bound_distinct)
```

---

missingDataGrid

*Missing Data Grid*

---

**Description**

Generate a levelplot of missing data from a SoilProfileCollection object.

**Usage**

```
missingDataGrid(s, max_depth, vars, filter.column = NULL,
  filter.regex = NULL, cols = NULL, ...)
```

**Arguments**

<code>s</code>	a SoilProfilecollection object
<code>max_depth</code>	integer specifying the max depth of analysis
<code>vars</code>	character vector of column names over which to evaluate missing data
<code>filter.column</code>	a character string naming the column to apply the filter REGEX to
<code>filter.regex</code>	a character string with a regular expression used to filter horizon data OUT of the analysis
<code>cols</code>	a vector of colors
<code>...</code>	additional arguments passed on to <code>levelplot</code>

**Details**

This function evaluates a ‘missing data fraction‘ based on slice-wise evaluation of named variables in a SoilProfileCollection object.

**Value**

A data.frame describing the percentage of missing data by variable.

**Note**

A lattice graphic is printed to the active output device.

**Author(s)**

D.E. Beaudette

**See Also**

[slice](#)

**Examples**

```
## visualizing missing data
# 10 random profiles
require(ply)
s <- ldply(1:10, random_profile)

# randomly sprinkle some missing data
s[sample(nrow(s), 5), 'p1'] <- NA
s[sample(nrow(s), 5), 'p2'] <- NA
s[sample(nrow(s), 5), 'p3'] <- NA

# set all p4 and p5 attributes of `soil 1` to NA
s[which(s$id == '1'), 'p5'] <- NA
s[which(s$id == '1'), 'p4'] <- NA

# upgrade to SPC
depths(s) <- id ~ top + bottom
```

```
# plot missing data via slicing + levelplot
missingDataGrid(s, max_depth=100, vars=c('p1', 'p2', 'p3', 'p4', 'p5'),
main='Missing Data Fraction')
```

---

munsell

*Munsell-RGB Lookup Table for Common Soil Colors*

---

## Description

A lookup table of interpolated Munsell color chips for common soil colors.

## Usage

```
data(munsell)
```

## Format

A data frame with 8825 observations on the following 6 variables.

hue Munsell Hue, upper case

value Munsell Value

chroma Munsell Chroma

r red value (0-1)

g green value (0-1)

b blue value (0-1)

## Details

See `munsell2rgb` for conversion examples. Note that this table does not currently have entries for values of 2.5– common in most soil color books. These chips should be added in the next major release of `aqp`.

## Source

Color chip XYZ values: <http://www.cis.rit.edu/mcsl/online/munsell.php>

## References

<http://www.brucelindbloom.com/index.html?ColorCalcHelp.html> Color conversion equations

<http://casoilresource.lawr.ucdavis.edu/drupal/node/201> Methods used to generate this table

## Examples

```
data(munsell)
```

---

`munsell2rgb`*Convert Munsell Notation to and from RGB color coordinates*

---

### Description

Color conversion based on a look-up table of common soil colors.

### Usage

```
munsell2rgb(the_hue, the_value, the_chroma, alpha=1,
maxColorValue=1, return_triplets=FALSE)
rgb2munsell(color)
```

### Arguments

<code>the_hue</code>	a vector of one or more more hues, upper-case
<code>the_value</code>	a vector of one or more values
<code>the_chroma</code>	a vector of one or more chromas
<code>alpha</code>	alpha channel value (for transparency effects)
<code>maxColorValue</code>	maximum RGB color value (see <a href="#">rgb</a> )
<code>return_triplets</code>	should the function return raw RGB triplets instead of an R color
<code>color</code>	a <code>data.frame</code> or <code>matrix</code> object containing color-space coordinates: [R, G, B]

### Details

These functions generalizes to vectorized usage, as long as the length of each argument is the same. Both functions will pad output with NA if there are any NA present in the inputs.

### Value

For Munsell to RGB conversion, a vector of R colors is returned that is the same length as the input data. If `return_triplets` is TRUE, then a dataframe (of sample length as input) of r,g,b values is returned.

For RGB to Munsell conversion, a dataframe (NA-padded) of hue, value, chroma, and Euclidean distance to nearest matching color is returned.

### Warning

As of `plyr` 1.6 (CRAN), there are cases when this function will fail (<https://github.com/hadley/plyr/issues/43>). The next version of `plyr` should address this problem. Until then, be sure not to pass in Munsell (hue, value, chroma) as factors.



**Note**

Care should be taken when using the resulting RGB values; they are close to their Munsell counterparts, but will vary based on your monitor and ambient lighting conditions. Also, the value used for `maxColorValue` will affect the brightness of the colors. The default value (1) will usually give acceptable results, but can be adjusted to force the colors closer to what the user thinks they should look like.

**Author(s)**

Dylan E. Beaudette

**References**

<http://casoilresource.lawr.ucdavis.edu/drupal/node/201> <http://www.brucelindbloom.com/index.html?ColorCalcHelp.html> <http://www.cis.rit.edu/mcsl/online/munsell.php>

**Examples**

```
# Munsell to RGB triplets:
# function is vectorized as long as arguments are the same length
color <- munsell2rgb(the_hue=c('10YR', '2.5YR'), the_value=c(3, 5),
the_chroma=c(5, 6), return_triplets=TRUE)

# RGB triplets to closest Munsell color (in RGB space)
# function is vectorized
rgb2munsell(color)

# basic example: no factors!
d <- expand.grid(hue='10YR', value=2:8, chroma=1:8, stringsAsFactors=FALSE)
d$color <- with(d, munsell2rgb(hue, value, chroma))

# similar to the 10YR color book page
plot(value ~ chroma, data=d, col=d$color, pch=15, cex=3)

# multiple pages of hue:
hues <- c('2.5YR', '5YR', '7.5YR', '10YR')
d <- expand.grid(hue=hues, value=2:8, chroma=seq(2,8,by=2), stringsAsFactors=FALSE)
d$color <- with(d, munsell2rgb(hue, value, chroma))

# plot: note that we are setting panel order from red-->yellow
library(lattice)
xyplot(value ~ factor(chroma) | factor(hue, levels=hues),
main="Common Soil Colors", layout=c(4,1), scales=list(alternating=1),
strip=strip.custom(bg=grey(0.85)),
data=d, as.table=TRUE, subscripts=TRUE, xlab='Chroma', ylab='Value',
panel=function(x, y, subscripts, ...)
{
panel.xyplot(x, y, pch=15, cex=4, col=d$color[subscripts])
}
```

```

}
)

# try again, this time annotate with LAB coordinates:
if(require(colorspace)) {
  d.rgb <- with(d, munsell2rgb(hue, value, chroma, return_triplets=TRUE))
  d.lab <- as(with(d.rgb, RGB(r,g,b)), 'LAB')
  d <- data.frame(d, d.lab@coords)

  xyplot(value ~ factor(chroma) | factor(hue, levels=hues),
    main="Common Soil Colors - Annotated with LAB Coordinates", layout=c(4,1),
    scales=list(alternating=1), strip=strip.custom(bg=grey(0.85)),
    data=d, as.table=TRUE, subscripts=TRUE, xlab='Chroma', ylab='Value',
    panel=function(x, y, subscripts, ...) {
      panel.xyplot(x, y, pch=15, cex=7, col=d$color[subscripts])
      lab.text <- with(d[subscripts, ], paste(round(L), round(A), round(B), sep='\n'))
      panel.text(x, y, labels=lab.text, cex=0.75, col='white', font=2)
    }
  )

  # also demonstrate A ~ hue for each slice of chroma
  xyplot(A ~ factor(hue, levels=hues) | factor(value), groups=chroma, data=d,
    scales=list(alternating=1), strip=strip.custom(bg=grey(0.85)),
    main="A-coordinate vs. Munsell Hue", sub='panels are Munsell value, colors are Munsell chroma',
    xlab='Munsell Hue', ylab='A-coordinate', pch=16,
    type='b', as.table=TRUE, auto.key=list(lines=TRUE, points=FALSE, columns=4))
}

# soils example
data(sp1)

# convert colors
sp1$soil_color <- with(sp1, munsell2rgb(hue, value, chroma))

# simple plot, may need to tweak gamma-correction...
image(matrix(1:nrow(sp1)), axes=FALSE, col=sp1$soil_color, main='Soil Colors')

# convert into a more useful color space
# you will need the colorspace package for this to work
if(require(colorspace)) {
  # keep RGB triplets from conversion
  sp1.rgb <- with(sp1, munsell2rgb(hue, value, chroma, return_triplets=TRUE))

  # convert into LAB color space
  sp1.lab <- as(with(sp1.rgb, RGB(r,g,b)), 'LAB')
  plot(sp1.lab)
}

```

---

panel.depth\_function *Lattice Panel Function for Soil Profiles*

---

### Description

Panel function for plotting grouped soil property data, along with upper and lower estimates of uncertainty.

### Usage

```
panel.depth_function(x, y, id, upper = NA, lower = NA,
  subscripts = NULL, groups = NULL, sync.colors=FALSE, cf=NA,
  cf.col=NA, cf.interval=20, ...)
```

### Arguments

x	x values (generated by calling lattice function)
y	y values (generated by calling lattice function)
id	vector of id labels, same length as x and y—only required when plotting segments (see Details section)
upper	vector of upper confidence envelope values
lower	vector of lower confidence envelope values
subscripts	paneling indices (generated by calling lattice function)
groups	grouping data (generated by calling lattice function)
sync.colors	optionally sync the fill color within the region bounded by (lower–upper) with the line colors
cf	optionally annotate contributing fraction data at regular depth intervals see <a href="#">slab</a>
cf.col	optionall color for contributing fraction values, typically used to override the line color
cf.interval	number of depth units to space printed contributing fraction values
...	further arguments to lower-level lattice plotting functions, see below

### Details

This function can be used to replace `panel.superpose` when plotting depth function data. When requested, contributing fraction data are printed using colors the same color as corresponding depth function lines unless a single color value is given via `cf.col`.

### Author(s)

Dylan E. Beaudette

### References

<http://casoilresource.lawr.ucdavis.edu/>

**See Also**

[sp1](#), [slice](#), [slab](#)

**Examples**

```
library(lattice)
data(sp1)

# 1. plotting mechanism for step-functions derived from soil profile data
xyplot(cbind(top,bottom) ~ prop, data=sp1,id=sp1$id,
panel=panel.depth_function, ylim=c(250,-10),
scales=list(y=list(tick.number=10)), xlab='Property',
ylab='Depth (cm)', main='panel.depth_function() demo'
)

# 1.1 include groups argument to leverage lattice styling framework
sp1$group <- factor(sp1$group, labels=c('Group 1', 'Group2'))

xyplot(cbind(top,bottom) ~ prop, groups=group, data=sp1, id=sp1$id,
panel=panel.depth_function, ylim=c(250,-10),
scales=list(y=list(tick.number=10)), xlab='Property',
ylab='Depth (cm)', main='panel.depth_function() demo',
auto.key=list(columns=2, points=FALSE, lines=TRUE),
par.settings=list(superpose.line=list(col=c('Orange', 'RoyalBlue'))))
)
```

---

plot\_distance\_graph    *Between Individual Distance Plot*

---

**Description**

Plot pair-wise distances between individuals as line segments.

**Usage**

```
plot_distance_graph(D, idx = 1:dim(as.matrix((D)))[1])
```

**Arguments**

D                    distance matrix, should be of class 'dist' or compatible class  
idx                    an integer sequence defining which individuals should be compared

**Details**

By default all individuals are plotting on the same axis. When there are more than about 10 individuals, the plot can become quite messy. See examples below for ideas.

**Value**

No value is returned.

**Author(s)**

Dylan E Beaudette

**References**

<http://casoilresource.lawr.ucdavis.edu/>

**See Also**

[sp2](#), [profile\\_compare](#)

**Examples**

```
data(sp2)

d <- profile_compare(sp2, vars=c('prop', 'field_ph', 'hue', 'value'),
max_d=100, k=0.01, sample_interval=5)

par(mfcol=c(3,1), mar=c(2.5,4.5,1,1))
plot_distance_graph(d, idx=1:6)
plot_distance_graph(d, idx=7:12)
plot_distance_graph(d, idx=12:18)
```

---

profileApply-methods *Apply a function to soil profiles within a SoilProfileCollection object.*

---

**Description**

Apply a function to soil profiles within a SoilProfileCollection object, each iteration has access to a SoilProfileCollection object.

**Usage**

```
# method for SoilProfileCollection objects
profileApply(object, FUN, simplify=TRUE, ...)
```

**Arguments**

object	a SoilProfileCollection
FUN	a function to be applied to each profile within the collection
simplify	logical, should the result be simplified to a vector? see examples
...	further arguments passed to FUN

**Value**

When `simplify` is `TRUE`, a vector of length `nrow(object)` (horizon data) or of length `length(object)` (site data). When `simplify` is `FALSE`, a list is returned.

**Methods**

```
signature(object = "SoilProfileCollection")
```

**See Also**

[slab](#), [estimateSoilDepth](#)

**Examples**

```
data(sp1)
depths(sp1) <- id ~ top + bottom

# estimate soil depth using horizon designations
profileApply(sp1, estimateSoilDepth, name='name', top='top', bottom='bottom')

# scale properties within each profile
# scaled = (x - mean(x)) / sd(x)
sp1$d <- profileApply(sp1, FUN=function(x) round(scale(x$prop), 2))
plot(sp1, name='d')

# compute depth-wise differencing by profile
# note that our function expects that the column 'prop' exists
f <- function(x) { c(x$prop[1], diff(x$prop)) }
sp1$d <- profileApply(sp1, FUN=f)
plot(sp1, name='d')

# compute depth-wise cumulative sum by profile
# note the use of an anonymous function
sp1$d <- profileApply(sp1, FUN=function(x) cumsum(x$prop))
plot(sp1, name='d')

# compute profile-means, and save to @site
# there must be some data in @site for this to work
site(sp1) <- ~ group
sp1$mean_prop <- profileApply(sp1, FUN=function(x) mean(x$prop, na.rm=TRUE))

# re-plot using ranks defined by computed summaries (in @site)
plot(sp1, plot.order=rank(sp1$mean_prop))

## iterate over profiles, subsetting horizon data

# example data
data(sp1)
```

```

# promote to SoilProfileCollection
depths(sp1) <- id ~ top + bottom
site(sp1) <- ~ group

# make some fake site data related to a depth of some importance
sp1$dep <- profileApply(sp1, function(i) {round(rnorm(n=1, mean=mean(i$top)))})

# custom function for subsetting horizon data, by profile
# keep horizons with lower boundary < site-level attribute 'dep'
fun <- function(i) {
# extract horizons
h <- horizons(i)
# make an expression to subset horizons
exp <- paste('bottom < ', i$dep, sep='')
# subset horizons, and write-back into current SPC
horizons(i) <- subset(h, subset=eval(parse(text=exp)))
# return modified SPC
return(i)
}

# list of modified SoilProfileCollection objects
l <- profileApply(sp1, fun, simplify=FALSE)

# re-combine list of SoilProfileCollection objects into a single SoilProfileCollection
sp1.sub <- do.call(rbind, l)

# graphically check
par(mfrow=c(2,1), mar=c(0,0,1,0))
plot(sp1)
points(1:length(sp1), sp1$dep, col='red', pch=7)
plot(sp1.sub)

##
## helper functions: these must be modified to suit your own data
##

# compute the weighted-mean of some property within a given diagnostic horizon
# note that this requires conditional eval of data that may contain NA
# see ?slab for details on the syntax
# note that function expects certain columns within 'x'
f.diag.wt.prop <- function(x, d.hz, prop) {
# extract diagnostic horizon data
d <- diagnostic_hz(x)
# subset to the requested diagnostic hz
d <- d[d$diag_kind == d.hz, ]
# if missing return NA
if(nrow(d) == 0)
return(NA)

# extract depths and check for missing

```

```

sv <- c(d$featdept, d$featdepb)
if(any(is.na(sv)))
return(NA)

# create formula from named property
fm <- as.formula(paste('~', prop))
# return just the (weighted) mean, accessed from @horizons
s <- slab(x, fm, seg_vect=sv)$p.mean
return(s)
}

# conditional eval of thickness of some diagnostic feature or horizon
# will return a vector of length(x), you can save to @site
f.diag.thickness <- function(x, d.hz) {
# extract diagnostic horizon data
d <- diagnostic_hz(x)
# subset to the requested diagnostic hz
d <- d[d$diag_kind == d.hz, ]
# if missing return NA
if(nrow(d) == 0)
return(NA)

# compute thickness
thick <- d$featdepb - d$featdept
return(thick)
}

# conditional eval of property within particle size control section
# makes assumptions about the SPC that is passed-in
f.psc.prop <- function(x, prop) {
# these are accessed from @site
sv <- c(x$psctopdepth, x$pscbotdepth)
# test for missing PCS data
if(any(is.na(sv)))
return(NA)

# this should never happen... unless someone made a mistake
# check to make sure that the lower PSC boundary is shallower than the depth
if(sv[2] > max(x))
return(NA)

# create formula from named property
fm <- as.formula(paste('~', prop))
# return just the (weighted) mean, accessed from @horizons
s <- slab(x, fm, seg_vect=sv)$p.mean
return(s)
}

```



---

 profile\_compare-methods

*Numerical Soil Profile Comparison*


---

**Description**

Performs a numerical comparison of soil profiles using named properties, based on a weighted, summed, depth-segment-aligned dissimilarity calculation. If `s` is a [SoilProfileCollection](#), site-level variables (2 or more) can also be used. The site-level and horizon-level dissimilarity matrices are then re-scaled and averaged.

**Usage**

```
pc(s, vars, max_d, k, filter=NULL, sample_interval=NA,
  replace_na=TRUE, add_soil_flag=TRUE,
  return_depth_distances=FALSE, strict_hz_eval=FALSE,
  progress='none', plot.depth.matrix=FALSE, rescale.result=FALSE,
  verbose=FALSE)
```

**Arguments**

<code>s</code>	a dataframe with at least 2 columns of soil properties, and an 'id' column for each profile. horizon depths must be integers and self-consistent
<code>vars</code>	A vector with named properties that will be used in the comparison. These are typically column names describing horizon-level attributes (2 or more), but can also contain site-level attributes (2 or more) if <code>s</code> is a <a href="#">SoilProfileCollection</a> .
<code>max_d</code>	depth-slices up to this depth are considered in the comparison
<code>k</code>	a depth-weighting coefficient, use '0' for no depth-weighting (see examples below)
<code>filter</code>	an index used to determine which horizons (rows) are included in the analysis
<code>sample_interval</code>	use every n-th depth slice instead of every depth slice, useful for working with > 1000 profiles at a time
<code>replace_na</code>	if TRUE, missing data are replaced by maximum dissimilarity (TRUE)
<code>add_soil_flag</code>	The algorithm will generate a 'soil'/'non-soil' matrix for use when comparing soil profiles with large differences in depth (TRUE). See details section below.
<code>return_depth_distances</code>	return intermediate, depth-wise dissimilarity results (FALSE)
<code>strict_hz_eval</code>	should horizons be strictly checked for internal self-consistency? (FALSE)
<code>progress</code>	'none' (default): argument passed to <code>ddply</code> and related functions, see <a href="#">create_progress_bar</a> for all possible options; 'text' is usually fine.
<code>plot.depth.matrix</code>	should a plot of the 'soil'/'non-soil' matrix be returned (FALSE)
<code>rescale.result</code>	should the result be rescaled by dividing by <code>max(D)</code> (FALSE)
<code>verbose</code>	extra debug output (FALSE)

## Details

Variability in soil depth can interfere significantly with the calculation of between-profile dissimilarity—what is the numerical “distance” (or dissimilarity) between a slice of soil from profile A and the corresponding, but missing, slice from a shallower profile B? Gower’s distance metric would yield a NULL distance, despite the fact that intuition suggests otherwise: shallower soils should be more dissimilar from deeper soils. For example, when a 25 cm deep profile is compared with a 50 cm deep profile, numerical distances are only accumulated for the first 25 cm of soil (distances from 26 - 50 cm are NULL). When summed, the total distance between these profiles will generally be less than the distance between two profiles of equal depth. Our algorithm has an option (setting `replace_na=TRUE`) to replace NULL distances with the maximum distance between any pair of profiles for the current depth slice. In this way, the numerical distance between a slice of soil and a corresponding slice of non-soil reflects the fact that these two materials should be treated very differently (i.e. maximum dissimilarity).

This alternative calculation of dissimilarities between soil and non-soil slices solves the problem of comparing shallow profiles with deeper profiles. However, it can result in a new problem: distances calculated between two shallow profiles will be erroneously inflated beyond the extent of either profile’s depth. Our algorithm has an additional option (setting `add_soil_flag=TRUE`) that will preserve NULL distances between slices when both slices represent non-soil material. With this option enabled, shallow profiles will only accumulate mutual dissimilarity to the depth of the deeper profile.

Note that when the `add_soil_flag` option is enabled (default), slices are classified as ‘soil’ down to the maximum depth to which at least one of variables used in the dissimilarity calculation is not NA. This will cause problems when profiles within a collection contain all NAs within the columns used to determine dissimilarity. An approach for identifying and removing these kind of profiles is presented in the examples section below.

A notice is issued if there are any NA values within the matrix used for distance calculations, as these values are optionally replaced by the max dissimilarity.

Our approach builds on the work of (Moore, 1972) and the previously mentioned depth-slicing algorithm.

## Value

A dissimilarity matrix object of class ‘dissimilarity, dist’, optionally scaled by `max(D)`.

## Methods

- `data = "SoilProfileCollection"` see [SoilProfileCollection](#)
- `data = "data.frame"` see [profile\\_compare](#)

## Author(s)

Dylan E. Beaudette

## References

1. <http://casoilresource.lawr.ucdavis.edu/>
2. Moore, A.; Russell, J. & Ward, W. Numerical analysis of soils: A comparison of three soil profile models with field classification. *Journal of Soil Science*, 1972, 23, 194-209.

**See Also**[slice, daisy](#)**Examples**

```
## 1. check out the influence depth-weight coef:
require(lattice)
z <- rep(1:100,4)
k <- rep(c(0,0.1,0.05,0.01), each=100)
w <- 100*exp(-k*z)

xyplot(z ~ w, groups=k, ylim=c(105,-5), xlim=c(-5,105), type='l',
       ylab='Depth', xlab='Weighting Factor',
       auto.key=list(columns=4, lines=TRUE, points=FALSE, title="k", cex=0.8, size=3),
       panel=function(...) {
         panel.grid(h=-1,v=-1)
         panel.superpose(...)
       }
)

## 2. basic implementation, requires at least two properties
# implementation for a data.frame class object
data(sp1)
d <- profile_compare(sp1, vars=c('prop','group'), max_d=100, k=0.01,
plot.depth.matrix=TRUE)

# upgrade to SoilProfileCollection
depths(sp1) <- id ~ top + bottom
op <- par(mfrow=c(1,2))
# perform comparison on SoilProfileCollection object
# compare soil/non-soil matrix plot
d <- profile_compare(sp1, vars=c('prop','group'), max_d=100, k=0.01,
plot.depth.matrix=TRUE)

# plot profile collection
plot(sp1)
# annotate max depth of profile comparison
abline(h=100, col='red', lty=2)
par(op)

# more soil properties
data(sp2)
d.1 <- profile_compare(sp2, vars=c('prop','field_ph','hue','value'),
max_d=100, k=0.01, plot.depth.matrix=TRUE)

# add some missing data:
sp2$prop[1:2] <- NA
d.2 <- profile_compare(sp2, vars=c('prop','field_ph','hue','value'),
max_d=100, k=0.01, plot.depth.matrix=TRUE)

# note small changes in D:
```

```

cor(d.1, d.2)

## 3. identify profiles within a collection that contain all NAs
require(ply)
s <- ldply(1:10, random_profile)
depths(s) <- id ~ top + bottom

# replace first profile's data with NA
na.required <- nrow(s[1, ])
s$p1[1:na.required] <- NA
s$p2[1:na.required] <- NA

# attempt profile comparison: this won't work, throws an error
# d <- profile_compare(s, vars=c('p1','p2'), max_d=100, k=0)

# check for soils that are missing all clay / total RF data
f.check.NA <- function(i) length(which(is.na(i$p1) | is.na(i$p2))) / nrow(i) == 1
missing.too.much.data.idx <- which(profileApply(s, f.check.NA))

# remove bad profiles and try again: works
s.no.na <- profile_compare(s[-missing.too.much.data.idx, ], vars=c('p1','p2'),
max_d=100, k=0, plot.depth.matrix=TRUE)

## 4. better plotting of dendrograms with ape package:
require(ape)
require(cluster)
h <- diana(d)
p <- as.phylo(as.hclust(h))
plot(ladderize(p), cex=0.75, label.offset=1)
tiplabels(col=cutree(h, 3), pch=15)

## 5. other uses of the dissimilarity matrix
require(MASS)
# Sammon Mapping: doesn't like '0' values in dissimilarity matrix
d.sam <- sammon(d)

# simple plot
dev.off() ; dev.new()
plot(d.sam$points, type = "n", xlim=range(d.sam$points[,1] * 1.5))
text(d.sam$points, labels=row.names(as.data.frame(d.sam$points)),
cex=0.75, col=cutree(h, 3))

## 6. try out the 'sample_interval' argument
# compute using successively larger sampling intervals
data(sp3)
d <- profile_compare(sp3, vars=c('clay','cec','ph'),
max_d=100, k=0.01)
d.2 <- profile_compare(sp3, vars=c('clay','cec','ph'),
max_d=100, k=0.01, sample_interval=2)
d.10 <- profile_compare(sp3, vars=c('clay','cec','ph'),
max_d=100, k=0.01, sample_interval=10)

```

```
d.20 <- profile_compare(sp3, vars=c('clay','cec','ph'),
max_d=100, k=0.01, sample_interval=20)

# check the results via hclust / dendrograms
oldpar <- par(mfcol=c(1,4), mar=c(2,1,2,2))
plot(as.dendrogram(hclust(d)), horiz=TRUE, main='Every Depth Slice')
plot(as.dendrogram(hclust(d.2)), horiz=TRUE, main='Every 2nd Depth Slice')
plot(as.dendrogram(hclust(d.10)), horiz=TRUE, main='Every 10th Depth Slice')
plot(as.dendrogram(hclust(d.20)), horiz=TRUE, main='Every 20th Depth Slice')
par(oldpar)
```

---

random\_profile

*Random Profile*


---

## Description

Generate a random soil profile according to set criteria, with correlated depth trends.

## Usage

```
random_profile(id, n = c(3, 4, 5, 6), min_thick = 5,
max_thick = 30, n_prop = 5, exact = FALSE, method = 'random_walk',
HzDistinctSim=FALSE, ...)
```

## Arguments

id	a character or numeric id used for this profile
n	vector of possible number of horizons, or the exact number of horizons (see below)
min_thick	minimum thickness criteria for a simulated horizon
max_thick	maximum thickness criteria for a simulated horizon
n_prop	number of simulated soil properties (columns in the returned dataframe)
exact	should the exact number of requested horizons be generated? (defaults to FALSE)
method	named method used to synthesize depth function ('random_walk' or 'LPP'), see details
HzDistinctSim	optionally simulate horizon boundary distinctness codes
...	additional parameters passed-in to the LPP (.lpp) function

## Details

The random walk method produces profiles with considerable variation between horizons and is based on values from the normal distribution seeded with means and standard deviations drawn from the uniform distribution of [0, 10].

The logistic power peak (LPP) function can be used to generate random soil property depth functions that are sharply peaked. LPP parameters can be hard-coded using the optional arguments:

"lpp.a", "lpp.b", "lpp.u", "lpp.d", "lpp.e". Amplitude of the peak is controlled by ("lpp.a + "lpp.b"), depth of the peak by "lpp.u", and abruptness by "lpp.d" and "lpp.e". Further description of the method is outlined in (Brenton et al, 2011). Simulated horizon distinctness codes are based on the USDA-NCSS field description methods (<http://soils.usda.gov/technical/fieldbook/>). Simulated distinctness codes are constrained according to horizon thickness, i.e. a gradual boundary (+/- 5cm) will not be simulated for horizons that are thinner than 3x this vertical distance

### Value

A dataframe with the simulated profile.

### Note

See examples for ideas on simulating several profiles at once.

### Author(s)

Dylan E. Beaudette

### References

Myers, D. B.; Kitchen, N. R.; Sudduth, K. A.; Miles, R. J.; Sadler, E. J. & Grunwald, S. Peak functions for modeling high resolution soil profile data Geoderma, 2011, 166, 74-83.

### See Also

[profile\\_compare](#), [hzDistinctnessCodeToOffset](#)

### Examples

```
# generate 10 random profiles with default settings:
require(plyr)
d <- ldply(1:10, random_profile)

# add a fake color
d$soil_color <- 'white'

# promote to SoilProfileCollection and plot
depths(d) <- id ~ top + bottom
plot(d)

# simulate horizon boundary distinctness codes:
d <- ldply(1:10, random_profile, HzDistinctSim=TRUE)
d$soil_color <- grey(0.85)
depths(d) <- id ~ top + bottom
d$HzD <- hzDistinctnessCodeToOffset(d$HzDistinctCode)
plot(d, hz.distinctness.offset='HzD')

# depth functions are generated using the LPP function
```

```

opar <- par(mfrow=c(2,1), mar=c(0,0,3,0))
# generate data
d <- ldply(1:10, random_profile, n=c(6, 7, 8), n_prop=1, method='LPP')

# promote to SPC and plot
depths(d) <- id ~ top + bottom
plot(d, color='p1')

# do this again, this time set all of the LPP parameters
d <- ldply(1:10, random_profile, n=c(6, 7, 8), n_prop=1, method='LPP',
lpp.a=5, lpp.b=10, lpp.d=5, lpp.e=5, lpp.u=25)

depths(d) <- id ~ top + bottom
plot(d, color='p1')

# reset plotting defaults
par(opar)

# try plotting the LPP-derived simulated data
# aggregated over all profiles
a <- slab(d, fm= ~ p1)
a$mid <- with(a, (top + bottom) / 2)

library(lattice)
(p1 <- xyplot(mid ~ p.q50, data=a,
lower=a$p.q25, upper=a$p.q75, ylim=c(150,-5), alpha=0.5,
panel=panel.depth_function, prepanel=prepanel.depth_function,
cf=a$contributing_fraction, xlab='Simulated Data', ylab='Depth',
main='LPP(a=5, b=10, d=5, e=5, u=25)',
par.settings=list(superpose.line=list(col='black', lwd=2))
))

# optionally add original data as step-functions
if(require(latticeExtra)) {
  h <- horizons(d)
  p1 + as.layer(xyplot(top ~ p1, groups=id, data=h,
horizontal=TRUE, type='S',
par.settings=list(superpose.line=list(col='blue', lwd=1, lty=2))))
}

```

---

resample.twotheta

*Resample an XRD Pattern*


---

## Description

Resample an XRD pattern along a user-defined twotheta resolution via local spline interpolation.

**Usage**

```
resample.twotheta(twotheta, x, tt.min = min(twotheta),  
tt.max = max(twotheta), new.res = 0.02)
```

**Arguments**

twotheta	a vector of twotheta value
x	a vector of diffraction intensities corresponding with twotheta values
tt.min	new minimum twotheta value, defaults to current minimum
tt.max	new maximum twotheta value, defaults to current maximum
new.res	new twotheta resolution, defaults to 0.02

**Details**

Sometimes XRD patterns are collected at different resolutions, or at a resolution that is too great for full pattern matching. This function can be used to resample patterns to a consistent twotheta resolution, or to decimate massive patterns.

**Value**

A dataframe with the following columns

twotheta	new sequence of twotheta values
x	resampled diffraction intensities

**Author(s)**

Dylan E Beaudette

**References**

<http://casoilresource.lawr.ucdavis.edu/>

**See Also**

[rruff.sample](#)

**Examples**

```
data(rruff.sample)  
  
# resample single pattern  
nontronite.resamp <- with(rruff.sample,  
resample.twotheta(twotheta, nontronite, new.res=0.02) )  
  
# plot original vs. resampled pattern  
plot(nontronite ~ twotheta, data=rruff.sample, type='l', col='grey')  
lines(nontronite.resamp, col='blue')
```



---

rruff.sample                      *Sample XRD Patterns*

---

**Description**

Several sample XRD patterns from the RRUFF project site.

**Usage**

```
data(rruff.sample)
```

**Format**

A data frame with 3000 observations on the following 8 variables.

twotheta twotheta values

nontronite XRD pattern for nontronite

montmorillonite XRD pattern for montmorillonite

clinochlore XRD pattern for clinochlore

antigorite XRD pattern for antigorite

chamosite XRD pattern for chamosite

hematite XRD pattern for hematite

goethite XRD pattern for goethite

**Source**

<http://rruff.info/>

**References**

<http://rruff.info/>

**Examples**

```
data(rruff.sample)

# plot all patterns
matplot(rruff.sample, type='l', lty=1)
```

---

sim *Simulate Soil Profiles*

---

### Description

Simulate a collection of soil profiles based on the horization of a single soil profile.

### Usage

```
sim(x, n=1, iterations=25, hz.sd=2, min.thick=2)
```

### Arguments

x	a SoilProfileCollection object containing a single profile from which to draw simulated data
n	the number of requested simulations
iterations	sampling iterations used to determine each horizon thickness
hz.sd	standard deviation used to simulate horizon thickness, can be a vector but must divide evenly into the number of horizons found in x
min.thick	mininum horizon thickness allowed in simulation results

### Details

This function generates a collection of simulated soil profiles based on the horizon thickness data associated with a single "template" profile. Simulation is based on sampling from a family of gaussian distribution with means defined by the "template" profile and standard deviation defined by the user.

### Value

A SoilProfileCollection object with n simulated profiles, each containing the same number of horizons and same data as x.

### Author(s)

D. E. Beaudette

### See Also

[random\\_profile](#)

**Examples**

```

# load sample data and convert into SoilProfileCollection
data(sp3)
depths(sp3) <- id ~ top + bottom

# select a profile to use as the basis for simulation
s <- sp3[3, ]

# reset horizon names
s$name <- paste('H', seq_along(s$name), sep='')

# simulate 25 new profiles, using 's' and function defaults
sim.1 <- sim(s, n=25)

# simulate 25 new profiles using 's' and variable SD for each horizon
sim.2 <- sim(s, n=25, hz.sd=c(1, 2, 5, 5, 5, 10, 2))

# plot
par(mfrow=c(2,1), mar=c(0, 0, 0, 0))
plot(sim.1)
mtext('SD = 2', side=2, line=-1.5, font=2, cex=0.75)
plot(sim.2)
mtext('SD = c(1, 2, 5, 5, 5, 10, 2)', side=2, line=-1.5, font=2, cex=0.75)

# aggregate horization of simulated data
# note: set class_prob_mode=2 as profiles were not defined to a constant depth
sim.2$name <- factor(sim.2$name)
a <- slab(sim.2, ~ name, class_prob_mode=2)

# convert to long format for plotting simplicity
library(reshape)
a.long <- melt(a, id.vars=c('top','bottom'), measure.vars=levels(sim.2$name))

# plot horizon probabilities derived from simulated data
# dashed lines are the original horizon boundaries
library(lattice)
xyplot(top ~ value, groups=variable, data=a.long, subset=value > 0,
ylim=c(100, -5), type=c('l','g'), asp=1.5,
ylab='Depth (cm)', xlab='Probability',
auto.key=list(columns=4, lines=TRUE, points=FALSE),
panel=function(...) {
  panel.xyplot(...)
  panel.abline(h=s$stop, lty=2, lwd=2)
})

```

## Description

Aggregate soil properties along user-defined ‘slabs’, and optionally within groups.

## Usage

```
# method for SoilProfileCollection objects
slab(object, fm, slab.structure=1, strict=FALSE,
      slab.fun=.slab.fun.numeric.default, cpm=1, weights=NULL, ...)
```

## Arguments

<code>object</code>	a <code>SoilProfileCollection</code>
<code>fm</code>	A formula: either ‘groups ~ var1 + var2 + var3’ where named variables are aggregated within ‘groups’ OR where named variables are aggregated across the entire collection ‘ ~ var1 + var2 + var3’. If ‘groups’ is a factor it must not contain NA.
<code>slab.structure</code>	A user-defined slab thickness (defined by an integer), or user-defined structure (numeric vector). See details below.
<code>strict</code>	logical: should horizons be strictly checked for self-consistency?
<code>slab.fun</code>	Function used to process each ‘slab’ of data, ideally returning a vector with names attribute. Defaults to a wrapper function around <code>hdquantile</code> . See details.
<code>cpm</code>	Strategy for normalizing slice-wise probabilities, dividing by either: number of profiles with data at the current slice ( <code>cpm=1</code> ), or by the number of profiles in the collection ( <code>cpm=2</code> ). Mode 2 values will always sum to the contributing fraction, while mode 1 values will always sum to 1.
<code>weights</code>	Column name containing weights. NOT YET IMPLEMENTED
<code>...</code>	further arguments passed to <code>slab.fun</code>

## Details

Multiple continuous variables OR a single categorical (factor) variable can be aggregated within a call to `slab`. Basic error checking is performed to make sure that top and bottom horizon boundaries make sense. User-defined aggregate functions (`slab.fun`) should return a named vector of results. A new, named column will appear in the results of `slab` for every named element of a vector returned by `slab.fun`. See examples below for a simple example of a slab function that computes mean, mean-1SD and mean+1SD. The default slab function wraps `hdquantile` from the `Hmisc` package, which requires at least 2 observations per chunk. Note that if ‘group’ is a factor it must not contain NAs.

Execution time scales linearly (slower) with the total number of profiles in `object`, and exponentially (faster) as the number of profiles / group is increased. `slab()` and `slice()` are much faster and require less memory if input data are either numeric or character.

There are several possible ways to define slabs, using `slab.structure`:

**a single integer** e.g. 10: data are aggregated over a regular sequence of 10-unit thickness slabs

**a vector of 2 integers** e.g. `c(50, 60)`: data are aggregated over depths spanning 50–60 units

**a vector of 3 or more integers** e.g. `c(0, 5, 10, 50, 100)`: data are aggregated over the depths spanning 0–5, 5–10, 10–50, 50–100 units

**Value**

Output is returned in long format, such that slice-wise aggregates are returned once for each combination of grouping level (optional), variable described in the `fm` argument, and depth-wise 'slab'.

Aggregation of numeric variables, using the default slab function:

**variable** The names of variables included in the call to `slab`.

**groupname** The name of the grouping variable when provided, otherwise a fake grouping variable named 'all.profiles'.

**p.q5** The slice-wise 5th percentile.

**p.q25** The slice-wise 25th percentile

**p.q50** The slice-wise 50th percentile (median)

**p.q75** The slice-wise 75th percentile

**p.q95** The slice-wise 95th percentile

**top** The slab top boundary.

**bottom** The slab bottom boundary.

**contributing\_fraction** The fraction of profiles contributing to the aggregate value, ranges from  $1/n_{\text{profiles}}$  to 1.

When a single factor variable is used, slice-wise probabilities for each level of that factor are returned as:

**variable** The names of variables included in the call to `slab`.

**groupname** The name of the grouping variable when provided, otherwise a fake grouping variable named 'all.profiles'.

**A** The slice-wise probability of level A

**B** The slice-wise probability of level B

...

**n** The slice-wise probability of level n

**top** The slab top boundary.

**bottom** The slab bottom boundary.

**contributing\_fraction** The fraction of profiles contributing to the aggregate value, ranges from  $1/n_{\text{profiles}}$  to 1.

**Methods**

**data = "SoilProfileCollection"** Typical usage, where input is a [SoilProfileCollection](#).

**Note**

Arguments to `slab` have changed with aqp 1.5 (2012-12-29) as part of a code clean-up and optimization. Calculation of weighted-summaries was broken in aqp 1.2-6 (2012-06-26), and removed as of aqp 1.5 (2012-12-29). `slab` replaced the previously defined `soil.slot.multiple` function as of aqp 0.98-8.58 (2011-12-21).

**Author(s)**

D.E. Beaudette

**References**

D.E. Beaudette, P. Roudier, A.T. O'Geen, Algorithms for quantitative pedology: A toolkit for soil scientists, *Computers & Geosciences*, Volume 52, March 2013, Pages 258-268, 10.1016/j.cageo.2012.10.020.

Harrell FE, Davis CE (1982): A new distribution-free quantile estimator. *Biometrika* 69:635-640.

**See Also**

[slice](#), [hdquantile](#)

**Examples**

```
##
## basic examples
##
library(lattice)
library(grid)

# load sample data, upgrade to SoilProfileCollection
data(sp1)
depths(sp1) <- id ~ top + bottom

# aggregate entire collection with two different segment sizes
a <- slab(sp1, fm = ~ prop)
b <- slab(sp1, fm = ~ prop, slab.structure=5)

# check output
str(a)

# stack into long format
ab <- make.groups(a, b)
ab$which <- factor(ab$which, levels=c('a','b'),
labels=c('1-cm Interval', '5-cm Interval'))

# plot median and IQR
# custom plotting function for uncertainty viz.
xyplot(top ~ p.q50 | which, data=ab, ylab='Depth',
xlab='median bounded by 25th and 75th percentiles',
lower=ab$p.q25, upper=ab$p.q75, ylim=c(250,-5),
panel=panel.depth_function,
prepanel=prepanel.depth_function,
cf=ab$contributing_fraction,
layout=c(2,1), scales=list(x=list(alternating=1))
)

##
## categorical variable example
```

```

##
library(reshape)

# normalize horizon names: result is a factor
sp1$name <- generalize.hz(sp1$name,
new=c('0','A','B','C'),
pat=c('0', '^A', '^B', 'C'))

# compute slice-wise probability so that it sums to contributing fraction, from 0-150
a <- slab(sp1, fm= ~ name, cpm=1, slab.structure=0:150)

# reshape into long format for plotting
a.long <- melt(a, id.vars=c('top','bottom'), measure.vars=c('0','A','B','C'))

# plot horizon type proportions using panels
xyplot(top ~ value | variable, data=a.long, subset=value > 0,
ylim=c(150, -5), type=c('S','g'), horizontal=TRUE, layout=c(4,1), col=1 )

# again, this time using groups
xyplot(top ~ value, data=a.long, groups=variable, subset=value > 0,
ylim=c(150, -5), type=c('S','g'), horizontal=TRUE, asp=2)

# adjust probability to size of collection, from 0-150
a.1 <- slab(sp1, fm= ~ name, cpm=2, slab.structure=0:150)

# reshape into long format for plotting
a.1.long <- melt(a.1, id.vars=c('top','bottom'), measure.vars=c('0','A','B','C'))

# group mode 1 and mode 2 data
g <- make.groups(mode_1=a.long, mode_2=a.1.long)

# plot horizon type proportions
xyplot(top ~ value | variable, groups=which, data=g, subset=value > 0,
ylim=c(240, -5), type=c('S','g'), horizontal=TRUE, layout=c(4,1),
auto.key=list(lines=TRUE, points=FALSE, columns=2),
par.settings=list(superpose.line=list(col=c(1,2))))

# apply slice-wise evaluation of max probability, and assign ML-horizon at each slice
(gen.hz.ml <- get.ml.hz(a, c('0','A','B','C'))

## Not run:
##
## multivariate examples
##
data(sp3)

# add new grouping factor
sp3$group <- 'group 1'
sp3$group[as.numeric(sp3$id) > 5] <- 'group 2'
sp3$group <- factor(sp3$group)

# upgrade to SPC

```

```

depths(sp3) <- id ~ top + bottom
site(sp3) <- ~ group

# custom 'slab' function, returning mean +/- 1SD
mean.and.sd <- function(values) {
  m <- mean(values, na.rm=TRUE)
  s <- sd(values, na.rm=TRUE)
  upper <- m + s
  lower <- m - s
  res <- c(mean=m, lower=lower, upper=upper)
  return(res)
}

# aggregate several variables at once, within 'group'
a <- slab(sp3, fm=group ~ L + A + B, slab.fun=mean.and.sd)

# check the results:
# note that 'group' is the column containing group labels
library(lattice)
xyplot(
  top ~ mean | variable, data=a, groups=group,
  lower=a$lower, upper=a$upper, sync.colors=TRUE, alpha=0.5,
  cf=a$contributing_fraction,
  ylim=c(125,-5), layout=c(3,1), scales=list(x=list(relation='free')),
  par.settings=list(superpose.line=list(lwd=2, col=c('RoyalBlue', 'Orange2'))),
  panel=panel.depth_function,
  prepanel=prepanel.depth_function,
  auto.key=list(columns=2, lines=TRUE, points=FALSE)
)

# compare a single profile to the group-level aggregate values
a.1 <- slab(sp3[1, ], fm=group ~ L + A + B, slab.fun=mean.and.sd)

# manually update the group column
a.1$group <- 'profile 1'

# combine into a single data.frame:
g <- rbind(a, a.1)

# plot with customized line styles
xyplot(
  top ~ mean | variable, data=g, groups=group, subscripts=TRUE,
  lower=a$lower, upper=a$upper, ylim=c(125,-5),
  layout=c(3,1), scales=list(x=list(relation='free')),
  panel=panel.depth_function,
  prepanel=prepanel.depth_function,
  sync.colors=TRUE, alpha=0.25,
  par.settings=list(superpose.line=list(col=c('orange', 'royalblue', 'black'),
    lwd=2, lty=c(1,1,2))),
  auto.key=list(columns=3, lines=TRUE, points=FALSE)
)

```



```

## convert mean value for each variable into long format
library(reshape)

# note that depths are no longer in order
a.wide <- cast(a, group + top + bottom ~ variable, value=c('mean'))

## again, this time for a user-defined slab from 40-60 cm
a <- slab(sp3, fm=group ~ L + A + B, slab.structure=c(40,60), slab.fun=mean.and.sd)

# now we have weighted average properties (within the defined slab)
# for each variable, and each group
(a.wide <- cast(a, group + top + bottom ~ variable, value=c('mean'))))

## this time, compute the weighted mean of selected properties, by profile ID
a <- slab(sp3, fm= id ~ L + A + B, slab.structure=c(40,60), slab.fun=mean.and.sd)
(a.wide <- cast(a, id + top + bottom ~ variable, value=c('mean'))))

## aggregate the entire collection, using default slab function (hdquantile)
## note the missing left-hand side of the formula
a <- slab(sp3, fm= ~ L + A + B)

## weighted-aggregation -- NOT YET IMPLEMENTED --
# load sample data, upgrade to SoilProfileCollection
data(sp1)
depths(sp1) <- id ~ top + bottom

# generate pretend weights as site-level attribute
set.seed(10101)
sp1$site.wts <- runif(n=length(sp1), min=20, max=100)

## End(Not run)

```

---

slice-methods

*Slicing of SoilProfilecollection Objects*

---

## Description

Slicing of SoilProfilecollection Objects

## Usage

```

# method for SoilProfileCollection objects
slice(object, fm, top.down=TRUE, just.the.data=FALSE, strict=TRUE)

```

**Arguments**

object	a SoilProfileCollection
fm	A formula: either 'integer.vector ~ var1 + var2 + var3' where named variables are sliced according to 'integer.vector' OR where all variables are sliced according to 'integer.vector' 'integer.vector ~.'
top.down	Logical, should slices be defined from the top-down? The default is usually what you want.
just.the.data	Logical, return just the sliced data or a new SoilProfileCollection object.
strict	Logical, should the horizonation be strictly checked for self-consistency?

**Value**

Either a new SoilProfileCollection with data sliced according to fm, or a data.frame.

**Methods**

**data = "SoilProfileCollection"** Typical usage, where input is a [SoilProfileCollection](#).

**Note**

slab() and slice() are much faster and require less memory if input data are either numeric or character.

**Author(s)**

D.E. Beaudette

**References**

D.E. Beaudette, P. Roudier, A.T. O'Geen, Algorithms for quantitative pedology: A toolkit for soil scientists, Computers & Geosciences, Volume 52, March 2013, Pages 258-268, 10.1016/j.cageo.2012.10.020.

**See Also**

[slab](#)

**Examples**

```
# simulate some data, IDs are 1:20
library(plyr)
d <- ldply(1:20, random_profile)

# init SoilProfilecollection object
depths(d) <- id ~ top + bottom
head(horizons(d))

# generate single slice at 10 cm
# output is a SoilProfilecollection object
s <- slice(d, 10 ~ name + p1 + p2 + p3)
```

```

# generate single slice at 10 cm, output data.frame
s <- slice(d, 10 ~ name + p1 + p2 + p3, just.the.data=TRUE)

# generate integer slices from 0 - 25 cm
s <- slice(d, 0:25 ~ name + p1 + p2 + p3)
plot(s)

# generate slices from 0 - 10 cm, for all variables
s <- slice(d, 0:10 ~ .)
print(s)

# note that pct missing is computed for each slice,
# if all vars are missing, then NA is returned
d$p1[1:10] <- NA
s <- slice(d, 10 ~ ., just.the.data=TRUE)
print(s)

##
## check sliced data
##

# test that mean of 1 cm slices property is equal to the
# hz-thickness weighted mean value of that property
data(sp1)
depths(sp1) <- id ~ top + bottom

# get the first profile
sp1.sub <- sp1[which(profile_id(sp1) == 'P009'), ]

# compute hz-thickness wt. mean
hz.wt.mean <- with(horizons(sp1.sub),
sum((bottom - top) * prop) / sum(bottom - top)
)

# hopefully the same value, calculated via slice()
s <- slice(sp1.sub, 0:max(sp1.sub) ~ prop)
hz.slice.mean <- mean(s$prop, na.rm=TRUE)

# same?
if(!all.equal(hz.slice.mean, hz.wt.mean))
stop('there is a bug in slice() !!!')

```

---

SoilProfileCollection-class

*SoilProfileCollection Class*


---

**Description**

Basic class for storing soil profile collections, associated site data, and metadata.

**Objects from the Class**

Objects can be created by calls of the form `new("SoilProfileCollection", ...)`.

**Slots**

**idcol:** Object of class "character" the name of the column used to uniquely identify profiles

**depthcols:** Object of class "character" with the names of columns containing the horizon top and bottom boundaries

**metadata:** Object of class "data.frame" with collection-level metadata, having a single row, and user-defined columns

**horizons:** Object of class "data.frame" with 1 or more rows per profile

**site:** Object of class "data.frame" with 1 row per profile

**sp:** Object of class "SpatialPoints" with 1 row per profile

**diagnostic:** Object of class "data.frame" with 0 or more rows per profile

**Methods**

**\$** signature(x = "SoilProfileCollection"): ...

**\$<-** signature(x = "SoilProfileCollection"): ...

**[** signature(x = "SoilProfileCollection", i = "ANY", j = "missing"): ...

**[[** signature(x = "SoilProfileCollection", i = "ANY", j = "missing"): ...

**[[<-** signature(x = "SoilProfileCollection", i = "ANY", j = "missing"): ...

**coordinates<-** signature(object = "SoilProfileCollection"): ...

**horizonDepths** signature(object = "SoilProfileCollection"): ...

**horizons** signature(object = "SoilProfileCollection"): ...

**horizons<-** signature(object = "SoilProfileCollection"): ...

**idname** signature(object = "SoilProfileCollection"): ...

**names** signature(x = "SoilProfileCollection"): ...

**length** signature(x = "SoilProfileCollection"): ...

**max** signature(x = "SoilProfileCollection"): ...

**metadata** signature(object = "SoilProfileCollection"): ...

**metadata<-** signature(object = "SoilProfileCollection"): ...

**min** signature(x = "SoilProfileCollection"): ...

**profile\_id** signature(object = "SoilProfileCollection"): ...

**profile\_plot** signature(object = "SoilProfileCollection"): ...

**show** signature(object = "SoilProfileCollection"): ...

**site** signature(object = "SoilProfileCollection"): ...

```

site<- signature(object = "SoilProfileCollection"): ...
slab signature(data = "SoilProfileCollection"): ...
units signature(object = "SoilProfileCollection"): ...
units<- signature(object = "SoilProfileCollection"): ...
unique signature(object = "SoilProfileCollection"), vars = "character": ...

```

### Author(s)

Pierre Roudier and Dylan E. Beaudette

### Examples

```

## use the digest library to detect duplicate data
data(sp1)

# make a copy, make new IDs, and stack
s.1 <- sp1
s.2 <- sp1
s.2$id <- paste(s.2$id, '-copy', sep='')
s <- rbind(s.1, s.2)
depths(s) <- id ~ top + bottom

# digests are computed from horizon-level data only
# horizon boundaries and 'prop'
# result is an index of unique profiles
if(require(digest)) {
  u <- unique(s, vars=c('top', 'bottom', 'prop'))
}

# compare with and without dupes:
# note subsetting of SoilProfileCollection
cbind(dupes=length(s), no.dupes=length(s[u, ]))

## 3. concatenate SoilProfileCollection objects
require(plyr)
d <- ldply(1:10, random_profile)

# promote to SoilProfileCollection and plot
depths(d) <- id ~ top + bottom
plot(d)

# split into new SoilProfileCollection objects by index
d.1 <- d[1, ]
d.2 <- d[2, ]
d.345 <- d[3:5, ]

# recombine, note that profiles are sorted according to ID
d.new <- rbind(d.345, d.1, d.2)
plot(d.new)

```

```
## Not run:
# these next examples should throw an error
# insert a missing horizon boundary
data(sp1)
sp1$top[1] <- NA
depths(sp1) <- id ~ top + bottom

## End(Not run)
```

---

## SoilProfileCollection-plotting-methods

### *Profile Plot*

---

#### **Description**

Generate a simple diagram of a soil profile, with annotated horizon names.

#### **Usage**

```
plotSPC(x, color='soil_color', width=0.2, name=NULL, label=idname(x),
alt.label=NULL, alt.label.col='black', cex.names=0.5,
cex.depth.axis=cex.names, cex.id=cex.names+(0.2*cex.names),
print.id=TRUE, id.style='auto', plot.order=1:length(x), add=FALSE,
scaling.factor=1, y.offset=0, n=length(x),
max.depth=max(x), n.depth.ticks=5, shrink=FALSE,
shrink.cutoff=3, abbr=FALSE, abbr.cutoff=5, divide.hz=TRUE,
hz.distinctness.offset=NULL, hz.distinctness.offset.col='black',
hz.distinctness.offset.lty=2, axis.line.offset=-2.5,
density=NULL, col.label=color,
col.palette = rev(brewer.pal(10, 'Spectral')), lwd=1, lty=1, ...)
```

#### **Arguments**

<code>x</code>	a SoilProfileCollection object
<code>color</code>	the name of the column containing R-compatible color descriptions, or a column containing numeric data; see details
<code>width</code>	scaling of profile widths
<code>name</code>	the name of the (horizon-level) attribute containing horizon designation labels
<code>label</code>	the name of the (site-level) attribute used to identify profiles in the plot
<code>alt.label</code>	the name of a (site-level) attribute used for secondary annotation
<code>alt.label.col</code>	color used when printing secondary annotation
<code>cex.names</code>	character scaling applied to horizon names
<code>cex.depth.axis</code>	character scaling applied to depth scale
<code>cex.id</code>	character scaling applied to profile id
<code>print.id</code>	should the profile id be printed above each profile? (TRUE)

<code>id.style</code>	profile ID printing style: 'auto' (default) = simple heuristic used to select from: 'top' = centered above each profile, 'side' = 'along the top-left edge of profiles'
<code>plot.order</code>	a vector describing the order in which individual SoilProfile objects from the parent should be plotted
<code>add</code>	add to an existing figure
<code>scaling.factor</code>	vertical scaling of the profile heights
<code>y.offset</code>	vertical offset for top of profiles
<code>n</code>	integer describing amount of space along x-axis to allocate, defaults to <code>length(x)</code>
<code>max.depth</code>	suggested lower depth boundary of plot
<code>n.depth.ticks</code>	suggested number of ticks in depth scale
<code>shrink</code>	should long horizon names be shrunk by 80% ?
<code>shrink.cutoff</code>	character length defining long horizon names
<code>abbr</code>	should the profile ID be abbreviated?
<code>abbr.cutoff</code>	suggested minimum length for abbreviated IDs
<code>divide.hz</code>	should horizons be divided with a thin black line? (default is TRUE)
<code>hz.distinctness.offset</code>	column name containing vertical offsets used to depict horizon boundary distinctness (same units as profiles)
<code>hz.distinctness.offset.col</code>	color used to encode horizon distinctness (default is 'black')
<code>hz.distinctness.offset.lty</code>	line style used to encode horizon distinctness (default is 2)
<code>axis.line.offset</code>	horizontal offset applied to depth axis (default is -2.5)
<code>density</code>	fill density used for horizon color shading, either a single integer or a column name containing integer values (default is NULL, no shading)
<code>col.label</code>	text printed above the color-coded legend
<code>col.palette</code>	color palette used to plot numeric data
<code>lwd</code>	single numeric value: line width multiplier
<code>lty</code>	single integer: line style
<code>...</code>	other arguments passed into lower level plotting functions

### Details

Depth limits (`max.depth`) and number of depth ticks (`n.depth.ticks`) are *suggestions* to the `pretty` function. You may have to tinker with both parameters to get what you want.

The 'side' `id.style` is useful when plotting a large collection of profiles, and/or, when profile IDs are long.

If the column containing horizon designations is not specified (the `name` argument), a column (presumed to contain horizon designation labels) is guessed based on regular expression matching of the pattern 'name'— this usually works, but it is best to manually specify the name of the column containing horizon designations.

The color argument can either name a column containing R-compatible colors, possibly created via [munsell2rgb](#), or column containing numeric values. In the second case, numeric values are converted into colors and displayed along with a simple legend above the plot. Note that this functionality makes several assumptions about plot geometry and is most useful in an interactive setting.

### Value

A new plot of soil profiles is generated, or optionally added to an existing plot.

### Methods

```
signature(x = "SoilProfileCollection")
```

### Author(s)

Dylan E. Beaudette

### References

<http://casoilresource.lawr.ucdavis.edu/>

### See Also

[SoilProfileCollection-class](#), [pretty](#), [hzDistinctnessCodeToOffset](#), [addBracket](#)

### Examples

```
data(sp1)

# usually best to adjust margins
par(mar=c(0,0,3,0))

# add color vector
sp1$soil_color <- with(sp1, munsell2rgb(hue, value, chroma))

# promote to SoilProfileCollection
depths(sp1) <- id ~ top + bottom

# plot profiles
plot(sp1, id.style='side')

# title, note line argument:
title('Sample Data 1', line=1, cex.main=0.75)

# plot profiles without horizon-line divisions
plot(sp1, divide.hz=FALSE)

# add dashed lines illustrating horizon boundary distinctness
sp1$hzD <- hzDistinctnessCodeToOffset(sp1$bound_distinct)
plot(sp1, hz.distinctness.offset='hzD')
```



```

# plot horizon color according to some property
data(sp4)
depths(sp4) <- id ~ top + bottom
plot(sp4, color='clay')

# another example
data(sp2)
depths(sp2) <- id ~ top + bottom
site(sp2) <- ~ surface

# label with site-level attribute: `surface`
plot(sp2, label='surface', plot.order=order(sp2$surface))

```

---

sp1

*Soil Profile Data Example 1*


---

### Description

Soil profile data from Pinnacles National Monument, CA.

### Usage

```
data(sp1)
```

### Format

A data frame with 60 observations on the following 21 variables.

```

group a numeric vector
id a character vector
top a numeric vector
bottom a numeric vector
bound_distinct a character vector
bound_topography a character vector
name a character vector
texture a character vector
prop a numeric vector
structure_grade a character vector
structure_size a character vector
structure_type a character vector
stickiness a character vector
plasticity a character vector
field_ph a numeric vector
hue a character vector
value a numeric vector
chroma a numeric vector

```

## References

<http://casoilresource.lawr.ucdavis.edu/>

## Examples

```
data(sp1)
# convert colors from Munsell to hex-encoded RGB
sp1$soil_color <- with(sp1, munsell2rgb(hue, value, chroma))

# promote to SoilProfileCollection
depths(sp1) <- id ~ top + bottom
site(sp1) <- ~ group

# re-sample each profile into 1 cm (thick) depth slices
# for the variables 'prop', 'name', 'soil_color'
# result is a SoilProfileCollection object
s <- slice(sp1, 0:25 ~ prop + name + soil_color)

# plot, note slices
plot(s)

# aggregate all profiles along 1 cm depth slices,
# using data from column 'prop'
s1 <- slab(sp1, fm= ~ prop)

# check median & IQR
library(lattice)
xyplot(top ~ p.q50 + p.q25 + p.q75,
data=s1, type='S', horizontal=TRUE, col=1, lty=c(1,2,2),
panel=panel.superpose, ylim=c(110,-5), asp=2)
```

---

sp2

*Honcut Creek Soil Profile Data*

---

## Description

A collection of 18 soil profiles, consisting of select soil morphologic attributes, associated with a stratigraphic study conducted near Honcut Creek, California.

## Usage

```
data(sp2)
```

**Format**

A data frame with 154 observations on the following 21 variables.

id profile id  
surface dated surface  
top horizon top in cm  
bottom horizon bottom in cm  
bound\_distinct horizon lower boundary distinctness class  
bound\_topography horizon lower boundary topography class  
name horizon name  
texture USDA soil texture class  
prop field-estimated clay content  
structure\_grade soil structure grade  
structure\_size soil structure size  
structure\_type soil structure type  
stickiness stickiness  
plasticity plasticity  
field\_ph field-measured pH  
hue Munsell hue  
value Munsell value  
chroma Munsell chroma  
r RGB red component  
g RGB green component  
b RGB blue component  
soil\_color R-friendly encoding of soil color

**Author(s)**

Dylan E. Beaudette

**Source**

Busacca, Alan J.; Singer, Michael J.; Verosub, Kenneth L. 1989. Late Cenozoic stratigraphy of the Feather and Yuba rivers area, California, with a section on soil development in mixed alluvium at Honcut Creek. USGS Bulletin 1590-G.

**References**

<http://casoilresource.lawr.ucdavis.edu/>

**Examples**

```

data(sp2)

# convert into SoilProfileCollection object
depths(sp2) <- id ~ top + bottom

# transfer site-level data
site(sp2) <- ~ surface

# generate a new plotting order, based on the dated surface each soil was described on
p.order <- order(sp2$surface)

# plot
par(mar=c(1,0,3,0))
plot(sp2, plot.order=p.order)

# setup multi-figure output
par(mfrow=c(2,1), mar=c(0,0,1,0))

# truncate plot to 200 cm depth
plot(sp2, plot.order=p.order, max.depth=200)
abline(h=200, lty=2, lwd=2)

# compute numerical distances between profiles
# based on select horizon-level properties, to a depth of 200 cm
d <- profile_compare(sp2, vars=c('prop','field_ph','hue'),
max_d=200, k=0, sample_interval=5, rescale.result=TRUE)

# plot dendrogram with ape package:
require(ape) ; require(cluster)
h <- diana(d)
p <- as.phylo(as.hclust(h))
plot(p, cex=0.75, label.offset=0.01, font=1, direct='down', srt=90, adj=0.5, y.lim=c(-0.125, 0.5))

# add in the dated surface type via color
tiplabels(col=as.numeric(sp2$surface), pch=15)

# based on distance matrix values, YMMV
legend('topleft', legend=levels(sp2$surface), col=1:6, pch=15, bty='n', bg='white', cex=0.75)

```

---

sp3

*Soil Profile Data Example 3*


---

**Description**

Soil samples from 10 soil profiles, taken from the Sierra Foothill Region of California.

**Usage**

```
data(sp3)
```

**Format**

A data frame with 46 observations on the following 15 variables.

id soil id  
 top horizon upper boundary (cm)  
 bottom horizon lower boundary (cm)  
 clay clay content  
 cec CEC by amonium acetate at pH 7  
 ph pH in 1:1 water-soil mixture  
 tc total carbon percent  
 hue Munsell hue (dry)  
 value Munsell value (dry)  
 chroma Munsell chroma (dry)  
 mid horizon midpoint (cm)  
 ln\_tc natural log of total carbon percent  
 L color: l-coordinate, CIE-LAB colorspace (dry)  
 A color: a-coordinate, CIE-LAB colorspace (dry)  
 B color: b-coordinate, CIE-LAB colorspace (dry)  
**name** horizon name  
**soil\_color** horizon color

**Details**

These data were collected to support research funded by the Kearney Foundation of Soil Science.

**References**

<http://casoilresource.lawr.ucdavis.edu/>

**Examples**

```
## this example investigates the concept of a "median profile"

# required packages
library(ape)
data(sp3)

# generate a RGB version of soil colors
# and convert to HSV for aggregation
sp3$h <- NA ; sp3$s <- NA ; sp3$v <- NA
sp3.rgb <- with(sp3, munsell2rgb(hue, value, chroma, return_triplets=TRUE))
sp3[, c('h','s','v')] <- t(with(sp3.rgb, rgb2hsv(r, g, b, maxColorValue=1)))

# promote to SoilProfileCollection
depths(sp3) <- id ~ top + bottom
```

```

# aggregate across entire collection
a <- slab(sp3, fm= ~ clay + cec + ph + h + s + v, slab.structure=10)

# check
str(a)

# convert back to wide format
library(reshape)
a.wide.q25 <- cast(a, top + bottom ~ variable, value=c('p.q25'))
a.wide.q50 <- cast(a, top + bottom ~ variable, value=c('p.q50'))
a.wide.q75 <- cast(a, top + bottom ~ variable, value=c('p.q75'))

# add a new id for the 25th, 50th, and 75th percentile pedons
a.wide.q25$id <- 'Q25'
a.wide.q50$id <- 'Q50'
a.wide.q75$id <- 'Q75'

# combine original data with "mean profile"
vars <- c('top', 'bottom', 'id', 'clay', 'cec', 'ph', 'h', 's', 'v')
# make data.frame version of sp3
sp3.df <- as(sp3, 'data.frame')
sp3.grouped <- rbind(
sp3.df[, vars], a.wide.q25[, vars], a.wide.q50[, vars], a.wide.q75[, vars]
)

# re-constitute the soil color from HSV triplets
# convert HSV back to standard R colors
sp3.grouped$soil_color <- with(sp3.grouped, hsv(h, s, v))

# give each horizon a name
sp3.grouped$name <- paste(round(sp3.grouped$clay), '/',
round(sp3.grouped$cec), '/', round(sp3.grouped$ph,1))

## perform comparison, and convert to phylo class object
## D is rescaled to [0,]
d <- profile_compare(sp3.grouped, vars=c('clay', 'cec', 'ph'), max_d=100,
k=0.01, replace_na=TRUE, add_soil_flag=TRUE, rescale.result=TRUE)

require(cluster)
h <- agnes(d, method='ward')
p <- ladderize(as.phylo(as.hclust(h)))

# look at distance plot-- just the median profile
plot_distance_graph(d, 12)

# similarity relative to median profile (profile #12)
round(1 - (as.matrix(d)[12, ] / max(as.matrix(d)[12, ])), 2)

## make dendrogram + soil profiles

```

```

# first promote to SoilProfileCollection
depths(sp3.grouped) <- id ~ top + bottom

# setup plot: note that D has a scale of [0,1]
par(mar=c(1,1,1,1))
p.plot <- plot(p, cex=0.8, label.offset=3, direction='up', y.lim=c(2,0),
x.lim=c(1.25,length(sp3.grouped)+1), show.tip.label=FALSE)

# get the last plot geometry
lastPP <- get("last_plot.phylo", envir = .PlotPhyloEnv)

# the original labels, and new (indexed) order of pedons in dendrogram
d.labels <- attr(d, 'Labels')

new_order <- sapply(1:lastPP$Ntip,
function(i) which(as.integer(lastPP$xx[1:lastPP$Ntip]) == i))

# plot the profiles, in the ordering defined by the dendrogram
# with a couple fudge factors to make them fit
plot(sp3.grouped, color="soil_color", plot.order=new_order,
scaling.factor=0.01, width=0.1, cex.names=0.5,
y.offset=max(lastPP$yy)+0.1, add=TRUE)

```

---

sp4

*Soil Chemical Data from Serpentinic Soils of California*


---

### Description

Soil Chemical Data from Serpentinic Soils of California

### Usage

```
data(sp4)
```

### Format

A data frame with 30 observations on the following 13 variables.

```

id site name
name horizon designation
top horizon top boundary in cm
bottom horizon bottom boundary in cm
K exchangeable K in c mol/kg
Mg exchangeable Mg in cmol/kg
Ca exchangeable Ca in cmol/kg
CEC_7 cation exchange capacity (NH4OAc at pH 7)
ex_Ca_to_Mg extractable Ca:Mg ratio

```

```
sand sand content by weight percentage
silt silt content by weight percentage
clay clay content by weight percentage
CF >2mm fraction by volume percentage
```

### Details

Selected soil physical and chemical data from (McGahan et al., 2009).

### Source

<https://www.soils.org/publications/sssaj/articles/73/6/2087>

### References

McGahan, D.G., Southard, R.J., Claassen, V.P. 2009. Plant-Available Calcium Varies Widely in Soils on Serpentinite Landscapes. *Soil Sci. Soc. Am. J.* 73: 2087-2095.

### Examples

```
# setup environment
library(aqp)

# load sample data set, a simple data.frame object with horizon-level data from 10 profiles
data(sp4)
str(sp4)

# optionally read about it...
# ?sp4

# upgrade to SoilProfileCollection
# 'id' is the name of the column containing the profile ID
# 'top' is the name of the column containing horizon upper boundaries
# 'bottom' is the name of the column containing horizon lower boundaries
depths(sp4) <- id ~ top + bottom

# check it out
class(sp4) # class name
str(sp4) # internal structure

# inspect object properties
idname(sp4) # self-explanatory
horizonDepths(sp4) # self-explanatory

# you can change these:
depth_units(sp4) # defaults to 'cm'
metadata(sp4) # not much to start with

# alter the depth unit metadata
depth_units(sp4) <- 'inches' # units are really 'cm'
```



```

# more generic interface for adjusting metadata
md <- metadata(sp4) # save original metadata

# add columns
md$describer <- 'DGM'
md$date <- as.Date('2009-01-01')
md$citation <- 'McGahan, D.G., Southard, R.J., Claassen, V.P.
2009. Plant-Available Calcium Varies Widely in Soils
on Serpentinite Landscapes. Soil Sci. Soc. Am. J. 73: 2087-2095.'

# re-assign
metadata(sp4) <- md
depth_units(sp4) <- 'cm' # fix depth units, back to 'cm'

# further inspection with common function overloads
length(sp4) # number of profiles in the collection
nrow(sp4) # number of horizons in the collection
names(sp4) # column names
min(sp4) # shallowest profile depth in collection
max(sp4) # deepest profile depth in collection

# extraction of soil profile components
profile_id(sp4) # vector of profile IDs
horizons(sp4) # horizon data

# extraction of specific horizon attributes
sp4$clay # vector of clay content

# subsetting SoilProfileCollection objects
sp4[1, ] # first profile in the collection
sp4[, 1] # first horizon from each profile

# basic plot method, highly customizable: see manual page ?plotSPC
plot(sp4)
# inspect plotting area, very simple to overlay graphical elements
abline(v=1:length(sp4), lty=3, col='blue')
# profiles are centered at integers, from 1 to length(obj)
axis(1, line=-1.5, at=1:10, cex.axis=0.75, font=4, col='blue', lwd=2)
# y-axis is based on profile depths
axis(2, line=-1, at=pretty(1:max(sp4)), cex.axis=0.75, font=4, las=1, col='blue', lwd=2)

# symbolize soil properties via color
par(mar=c(0,0,4,0))
plot(sp4, color='clay')
plot(sp4, color='CF')

# apply a function to each profile, returning a single value per profile,
# in the same order as profile_id(sp4)
soil.depths <- profileApply(sp4, max) # recall that max() gives the depth of a soil profile

# check that the order is correct
all.equal(names(soil.depths), profile_id(sp4))

```

```

# a vector of values that is the same length as the number of profiles
# can be stored into site-level data
sp4$depth <- soil.depths
# check: looks good
max(sp4[1, ]) == sp4$depth[1]

# extract site-level data
site(sp4) # as a data.frame
sp4$depth # specific columns as a vector

# use site-level data to alter plotting order
new.order <- order(sp4$depth) # the result is an index of rank
par(mar=c(0,0,0,0))
plot(sp4, plot.order=new.order)

# deconstruct SoilProfileCollection into a data.frame, with horizon+site data
as(sp4, 'data.frame')

```

---

sp5

---

*Sample Soil Database #5*


---

## Description

296 Soil Profiles from the La Rochelle region of France (F. Carre and Girard, 2002)

## Usage

```
data(sp5)
```

## Format

```

Formal class 'SoilProfileCollection' [package "aqp"] with 6 slots
 ..@ idcol      : chr "soil"
 ..@ depthcols: chr [1:2] "top" "bottom"
 ..@ metadata  :'data.frame': 1 obs. of  1 variable:
 .. ..$ depth_units: chr "cm"
 ..@ horizons  :'data.frame': 1539 obs. of  17 variables:
 .. ..$ soil      : soil ID
 .. ..$ sand      : sand
 .. ..$ silt      : silt
 .. ..$ clay      : clay
 .. ..$ R25       : RGB r-coordinate
 .. ..$ G25       : RGB g-coordinate
 .. ..$ B25       : RGB b-coordinate
 .. ..$ pH        : pH
 .. ..$ EC        : EC
 .. ..$ CaC03    : CaC03 content
 .. ..$ C         : C content

```

```

.. ..$ Ca      : Ca
.. ..$ Mg      : Mg
.. ..$ Na      : Na
.. ..$ top     : horizon top boundary (cm)
.. ..$ bottom  : horizon bottom boundary (cm)
.. ..$ soil_color: soil color in r-friendly format
..@ site      : 'data.frame': 296 obs. of 1 variable:
.. ..$ soil: chr [1:296] "soil1" "soil10" "soil100" "soil101" ...
..@ sp       : Formal class 'SpatialPoints' [package "sp"] with 3 slots
.. .. ..@ coords : num [1, 1] 0
.. .. ..@ bbox   : logi [1, 1] NA
.. .. ..@ proj4string: Formal class 'CRS' [package "sp"] with 1 slots
.. .. .. ..@ projargs: chr NA

```

## Details

These data are c/o F. Carre (Florence.CARRE@ineris.fr).

## Source

296 Soil Profiles from the La Rochelle region of France (F. Carre and Girard, 2002). These data can be found on the OSACA project page (<http://eussoils.jrc.ec.europa.eu/projects/OSACA/>).

## References

F. Carre, M.C. Girard. 2002. Quantitative mapping of soil types based on regression kriging of taxonomic distances with landform and land cover attributes. *Geoderma*. 110: 241–263.

## Examples

```

library(scales)
data(sp5)
par(mar=c(1,1,1,1))
# plot a random sampling of profiles
s <- sample(1:length(sp5), size=25)
plot(sp5[s, ], divide.hz=FALSE)

# plot the first 100 profiles, as 4 rows of 25, hard-coding the max depth
layout(matrix(c(1,2,3,4), ncol=1), height=c(0.25,0.25,0.25,0.25))
plot(sp5[1:25, ], max.depth=300)
plot(sp5[26:50, ], max.depth=300)
plot(sp5[51:75, ], max.depth=300)
plot(sp5[76:100, ], max.depth=300)

# 4x1 matrix of plotting areas
layout(matrix(c(1,2,3,4), ncol=1), height=c(0.25,0.25,0.25,0.25))

# plot profiles, with points added to the mid-points of randomly selected horizons
sub <- sp5[1:25, ]
plot(sub, max.depth=300) ; mtext('Set 1', 2, line=-0.5, font=2)

```

```

y.p <- profileApply(sub, function(x) {
  s <- sample(1:nrow(x), 1)
  h <- horizons(x); with(h[s,], (top+bottom)/2)
})
points(1:25, y.p, bg='white', pch=21)

# plot profiles, with arrows pointing to profile bottoms
sub <- sp5[26:50, ]
plot(sub, max.depth=300); mtext('Set 2', 2, line=-0.5, font=2)
y.a <- profileApply(sub, function(x) max(x))
arrows(1:25, y.a-50, 1:25, y.a, len=0.1, col='white')

# plot profiles, with points connected by lines: ideally reflecting some kind of measured data
sub <- sp5[51:75, ]
plot(sub, max.depth=300); mtext('Set 3', 2, line=-0.5, font=2)
y.p <- 20*(sin(1:25) + 2*cos(1:25) + 5)
points(1:25, y.p, bg='white', pch=21)
lines(1:25, y.p, lty=2)

# plot profiles, with polygons connecting horizons with max clay content (+/-) 10 cm
sub <- sp5[76:100, ]
y.clay.max <- profileApply(sub, function(x) {
  i <- which.max(x$clay)
  h <- horizons(x)
  with(h[i, ], (top+bottom)/2)
})

plot(sub, max.depth=300); mtext('Set 4', 2, line=-0.5, font=2)
polygon(c(1:25, 25:1), c(y.clay.max-10, rev(y.clay.max+10)),
border='black', col=rgb(0,0,0.8, alpha=0.25))
points(1:25, y.clay.max, pch=21, bg='white')

# close plot
dev.off()

# plotting parameters
yo <- 100 # y-offset
sf <- 0.65 # scaling factor
# plot profile sketches
par(mar=c(0,0,0,0))
plot(sp5[1:25, ], max.depth=300, y.offset=yo, scaling.factor=sf)
# optionally add describe plotting area above profiles with lines
# abline(h=c(0,90,100, (300*sf)+yo), lty=2)
# simulate an environmental variable associated with profiles (elevation, etc.)
r <- vector(mode='numeric', length=25)
r[1] <- -50 ; for(i in 2:25) {r[i] <- r[i-1] + rnorm(mean=-1, sd=25, n=1)}
# rescale
r <- rescale(r, to=c(80, 0))
# illustrate gradient with points/lines/arrows
lines(1:25, r)
points(1:25, r, pch=16)
arrows(1:25, r, 1:25, 95, len=0.1)

```

```

# add scale for simulated gradient
axis(2, at=pretty(0:80), labels=rev(pretty(0:80)), line=-1, cex.axis=0.75, las=2)
# depict a secondary environmental gradient with polygons (water table depth, etc.)
polygon(c(1:25, 25:1), c((100-r)+150, rep((300*sf)+yo, times=25)),
border='black', col=rgb(0,0,0.8, alpha=0.25))

##
# sample 25 profiles from the collection
s <- sp5[sample(1:length(sp5), size=25), ]
# compute pair-wise dissimilarity
d <- profile_compare(s, vars=c('R25', 'pH', 'clay', 'EC'), k=0,
replace_na=TRUE, add_soil_flag=TRUE, max_d=300)
# keep only the dissimilarity between profile 1 and all others
d.1 <- as.matrix(d)[1, ]
# rescale dissimilarities
d.1 <- rescale(d.1, to=c(80, 0))
# sort in ascending order
d.1.order <- rev(order(d.1))
# plotting parameters
yo <- 100 # y-offset
sf <- 0.65 # scaling factor
# plot sketches
par(mar=c(0,0,0,0))
plot(s, max.depth=300, y.offset=yo, scaling.factor=sf, plot.order=d.1.order)
# add dissimilarity values with lines/points
lines(1:25, d.1[d.1.order])
points(1:25, d.1[d.1.order], pch=16)
# link dissimilarity values with profile sketches via arrows
arrows(1:25, d.1[d.1.order], 1:25, 95, len=0.1)
# add an axis for the dissimilarity scale
axis(2, at=pretty(0:80), labels=rev(pretty(0:80)), line=-1, cex.axis=0.75, las=2)

```

---

SPC-utils

*Getters, Setters, and Utility Methods for SoilProfileCollection Objects*


---

## Description

Getters, Setters, and Utility Methods for SoilProfileCollection Objects

## Methods

```
signature(object = "SoilProfileCollection")
```

## Author(s)

Dylan E. Beaudette

**References**

<http://casoilresource.lawr.ucdavis.edu/>

**Examples**

```

data(sp1)

## init SoilProfileCollection objects from data.frame
depths(sp1) <- id ~ top + bottom

## depth units
(du <- depth_units(sp1))
depth_units(sp1) <- 'in'
depth_units(sp1) <- du

## get/set metadata on SoilProfileCollection objects
# this is a 1-row data.frame
m <- metadata(sp1)
m$sampler <- 'Dylan'
metadata(sp1) <- m

## extract horizon data from SoilProfileCollection objects as data.frame
h <- horizons(sp1)

# also replace horizon data in SoilProfileCollection objects
# original order and length must be preserved!
horizons(sp1) <- h

# get number of horizons
nrow(sp1)

## getting site-level data
site(sp1)

## setting site-level data
# site-level data from horizon-level data (stored in @horizons)
site(sp1) <- ~ group

# make some fake site data, and append from data.frame
# a matching ID column must be present in both @site and new data
# note that IDs should all be character class
d <- data.frame(id=profile_id(sp1), p=runif(n=length(sp1)), stringsAsFactors=FALSE)
site(sp1) <- d

```

---

subsetProfiles-methods

*Subset SoilProfileCollection Objects.*

---

**Description**

This function is used to subset SoilProfileCollection objects using either site-level or horizon-level attributes, or both.

**Details**

The `s` argument supplies a fully-quoted search criteria for matching via site-level attributes. The `h` argument supplies a fully-quoted search criteria for matching via horizon-level attributes. All horizons associated with a single horizon-level match (i.e. out of several, only a single horizon matches the search criteria) are returned. See examples for usage.

**Value**

A SoilProfileCollection class object.

**Methods**

```
signature(object = "SoilProfileCollection", s = 'character', h = 'character', ...)
```

**See Also**

[profileApply](#), [site](#), [horizons](#)

**Examples**

```
data(sp1)
depths(sp1) <- id ~ top + bottom
site(sp1) <- ~ group

# save par settings, and setup plot for 3 columns
op <- par(mar=c(1,1,8,1), mfc=c(1,3))

# subset sp1 via site-level attributes
# note quoting style
plot(group.1 <- subsetProfiles(sp1, s="group == '1'"))

# subset sp1 via horizon-level attributes
# note quoting style
plot(coarse.soils <- subsetProfiles(sp1, h="texture == 'LS'"))

# re-combine subsets, note that duplicates are removed
g <- rbind(group.1, coarse.soils)
plot(g)

# reset plot area
par(op)

# subset sp1 via horizon and site-level attributes
plot(tiny.set <- subsetProfiles(sp1, s="group == 2", h='prop < 8'))
```

```
## other ways to subset SoilProfileCollection objects, via index
# more interesting sample data
data(sp2)
depths(sp2) <- id ~ top + bottom
site(sp2) <- ~ surface

# subset by integer index, note that this does not re-order the profiles
plot(sp2[1:5, ])

# generate an integer index via pattern-matching
idx <- grep('modesto', sp2$surface, ignore.case=TRUE)
plot(sp2[idx, ])

# generate in index via profileApply:
# subset those profiles where: min(ph) < 5.6
idx <- which(profileApply(sp2, function(i) min(i$field_ph, na.rm=TRUE) < 5.6))
plot(sp2[idx, ])
```

---

test\_hz\_logic

*Test Horizon Logic*


---

## Description

Simple tests for horizon logic, based on a simple data.frame of ordered horizons.

## Usage

```
test_hz_logic(i, topcol, bottomcol, strict = FALSE)
```

## Arguments

<code>i</code>	a data.frame associated with a single soil profile, ordered by depth
<code>topcol</code>	character, giving the name of the column in <code>i</code> that describes horizon top depth
<code>bottomcol</code>	character, giving the name of the column in <code>i</code> that describes horizon bottom depth
<code>strict</code>	logical, should continuity tests be performed– i.e. for non-contiguous horizon boundaries

## Details

By default, this function tests for NA and overlapping horizons. If any either are encountered, FALSE is returned.

## Value

logical: TRUE → pass, FALSE → fail



**Author(s)**

D.E. Beaudette

**References**<http://casoilresource.lawr.ucdavis.edu/>**See Also**[depths<-](#)**Examples**

```
## simple example: just one profile
data(sp1)
depths(sp1) <- id ~ top + bottom
s <- horizons(sp1[1, ])

## check
# fails due to missing hz boundary
s$bottom[6] <- NA # missing horizon boundary, common on bottom-most hz
test_hz_logic(s, 'top', 'bottom', strict=FALSE)

# fails due to inconsistent hz boundary
s$bottom[3] <- 30 # inconsistent hz boundary
test_hz_logic(s, 'top', 'bottom', strict=TRUE)

## filtering bad data
## Not run:
# missing bottom horizons
x$hzn_bot[!is.na(x$hzn_top) & is.na(x$hzn_bot)] <- x$hzn_top[is.na(x$hzn_top) & is.na(x$hzn_bot)]

# remove 0 horizons where top > bottom
bad.0.hz.idx <- which(x$hzn_top > x$hzn_bot)
if(length(bad.0.hz.idx) > 0)
x <- x[-bad.0.hz.idx, ]

## End(Not run)

## checking for bad data: do this before promoting to SoilProfileCollection object
library(plyr)
data(sp1)

# horizon logic can be tested via data.frame, at 2 levels of scrutiny:
ddply(sp1, 'id', test_hz_logic, topcol='top', bottomcol='bottom', strict=FALSE)
ddply(sp1, 'id', test_hz_logic, topcol='top', bottomcol='bottom', strict=TRUE)
```

---

```
texture.triangle.low.rv.high
```

*Soil Texture Low-RV-High as Defined by Quantiles*

---

### Description

This function accepts soil texture components (sand, silt, and clay percentages) and plots a soil texture triangle with a "representative value" (point) and low-high region (polygon) defined by quantiles. Marginal quantiles of sand, silt, and clay are used to define the boundary of a low-high region that encloses a several likely soil texture classes based on the values in ssc. The default settings place the RV symbol at the texture defined by marginal medians of sand, silt, and clay. The default low-high region is defined by the 5th and 95th marginal percentiles of sand, silt, and clay.

### Usage

```
texture.triangle.low.rv.high(ssc, p = c(0.05, 0.5, 0.95),
  method = c("approx", "closest"), n.closest = 3, delta = 1,
  pop.rv.col = "red", range.col = "RoyalBlue", range.alpha = 75)
```

### Arguments

ssc	a matrix-like object with columns: 'sand', 'silt', 'clay', values are percentages that should add to 100.
p	percentiles defining 'low', 'representative value', and 'high'
method	'approximate' uses marginal quantiles of [sand, silt, clay], 'closest' uses those data points that are closest to the marginal quantiles
n.closest	for method='closest', the number of observations used to define the low-high region
delta	step-size used to form low-high region via 'approx' method
pop.rv.col	the symbol color used to denote the population representative value on the texture triangle
range.col	color of the polygon enclosing the low-high region
range.alpha	transparency of the low-high range polygon (0-255)

### Value

A high-level plot as generated by `soil.texture`.

### Author(s)

D.E. Beaudette

### See Also

[triax.points](#), [soil.texture](#)

**Examples**

```
# sample data
data(loafercreek, package='soilDB')

# extract sand, silt, clay proportions
x <- na.omit(data.frame(sand=loafercreek$sand, silt=loafercreek$silt, clay=loafercreek$clay))

# test out the function
texture.triangle.low.rv.high(x, p=c(0.05, 0.5, 0.95))
texture.triangle.low.rv.high(x, p=c(0.25, 0.5, 0.75), range.col='darkgreen')

texture.triangle.low.rv.high(x, p=c(0.05, 0.5, 0.95), method='closest')
texture.triangle.low.rv.high(x, p=c(0.25, 0.5, 0.75), method='closest', range.col='darkgreen')
```

---

unroll

*Unroll Genetic Horizons*


---

**Description**

Generate a discretized vector of genetic horizons along a user-defined pattern.

**Usage**

```
unroll(top, bottom, prop, max_depth, bottom_padding_value = NA, strict=FALSE)
```

**Arguments**

top	vector of upper horizon boundaries, must be an integer
bottom	vector of lower horizon boundaries, must be an integer
prop	vector of some property to be "unrolled" over a regular sequence
max_depth	maximum depth to which missing data is padded with NA
bottom_padding_value	value to use when padding missing data
strict	should horizons be strictly checked for self-consistency? defaults to FALSE

**Details**

This function is used internally by several higher-level components of the aqp package. Basic error checking is performed to make sure that bottom and top horizon boundaries make sense. Note that the horizons should be sorted according to depth before using this function. The `max_depth` argument is used to specify the maximum depth of profiles within a collection, so that data from any profile shallower than this depth is padded with NA.

**Value**

a vector of "unrolled" property values

**Author(s)**

Dylan E. Beaudette

**References**

<http://casoilresource.lawr.ucdavis.edu/>

**Examples**

```
data(sp1)

# subset a single soil profile:
sp1.1 <- subset(sp1, subset=id == 'P001')

# demonstrate how this function works
x <- with(sp1.1, unroll(top, bottom, prop, max_depth=50))
plot(x, 1:length(x), ylim=c(90,0), type='b', cex=0.5)
```

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