

Package: fpod (via r-universe)

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Title Read and Process 'FPOD' and 'CPOD' Data

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Description Read 'FPOD' and 'CPOD' data into 'R' directly from the 'FPOD' data files (i.e. .CP1, .CP3, .FP1 and .FP3 files). The 'FPOD' data files contain binary data, so they can't trivially be read into 'R' using the usual approach, e.g. fread() or read.csv(). This package decodes the binary data and imports all the data in one go (i.e. header/metadata, clicks, 'KERNO' classifications, environmental data and pseudo-WAV data). It is then trivial to aggregate data as you please, e.g. detection-positive-minutes per time block. The advantage of handling data processing in 'R' is a long topic, but suffice it to say that it 1) simplifies things (many fewer steps, as different vars have to be exported in multiple goes in the official 'FPOD' app), and more importantly, 2) makes data processing transparent and reproducible. References: Pirotta et al. 2014 <doi:10.1111/1365-2435.12146>.

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LinkingTo Rcpp

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Suggests knitr, mixtools, rmarkdown, testthat (>= 3.0.0)

Depends R (>= 3.5)

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VignetteBuilder knitr

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BugReports <https://github.com/supermoan/fpod/issues>

Repository <https://supermoan.r-universe.dev>

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fp_example	<i>Get path to fpod example</i>
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Description

Get path to fpod example

Usage

```
fp_example(path = NULL)
```

Arguments

path Name of the file. If NULL, all example files are listed.

Value

A character vector, containing the path(s) that matched, or the empty string, "", if none matched.

Examples

```
# get a list all example files
fp_example()

# get the path for this particular file
fp_example("gullars_period1.FP3")
```

fp_find_buzzes	<i>Finds harbor porpoise feeding buzzes</i>
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Description

This function uses one of two methods to classify NBHF clicks into one of two classes: feeding buzz or non feeding buzz.

Usage

```
fp_find_buzzes(x, method = "clicks")
```

Arguments

x	a data.table where each row is a click, as the "clicks" element in the list object returned by <code>fp_read()</code> . Each row must minimally have a POSIXct column time, with nanosecond precision.
method	the method to use to find feeding buzzes - "clicks" or "trains". See details.

Details

Note that the so-called "feeding buzzes" are usually considered to represent a combination of feeding buzzes and social calls. Even so, the classifications resulting from both of these methods are commonly used as a proxy of foraging activity.

The two available methods are:

- `clicks` method: any inter-click-interval (ICI) less than 20 milliseconds is considered a NBHF feeding buzz.
- `trains` method: buzzes are identified using a mixture Gaussian model, with the number of components $k=3$. All clicks associated with the first component are considered a NBHF feeding buzz. This method requires the package `mixtools` to work.

Value

An integer vector of the same length as `nrow(x)`, where the values indicate that that click can be considered a feeding buzz (value = 1) or not (value = 0).

References

Pirotta, E., Thompson, P.M., Miller, P.I., Brookes, K.L., Cheney, B., Barton, T.R., Graham, I.M. and Lusseau, D. (2014), Scale-dependent foraging ecology of a marine top predator modelled using passive acoustic data. *Funct Ecol*, 28: 206-217. <https://doi.org/10.1111/1365-2435.12146>

Examples

```
# first read some FPOD data
fn <- fp_example("gullars_period1.FP3")
dat <- fp_read(fn)

# extract porpoise clicks of quality Hi and Mod
nbhf <- dat$clicks[species == "NBHF" & quality_level >= 2]

# then add a 'feeding buzz' column to the clicks data.table
nbhf$buzz <- fp_find_buzzes(nbhf, method = "clicks")
```

fp_plot

*Plot waveform and spectrum for a click***Description**

This function constructs and plots the waveform and spectrum for a click for which extra information has been recorded (IPI and SPL values for up to 21 cycles). By default, FPODs collect extra data for about 1% of detected clicks.

Usage

```
fp_plot(x, click_no = NULL, legend.pos = "topleft")
```

Arguments

x	A list object, as returned from <code>fp_read()</code> .
click_no	integer. The click number. If NULL, then all clicks are plotted, in sequential order, one at a time.
legend.pos	character/logical. If character; a single keyword specifying the legend position. This parameter is passed as x when calling <code>legend()</code> . See <code>legend()</code> for details. If legend.pos is logical and FALSE, then the legend is suppressed.

Details

According to the [FPOD software guide](#), extra data are stored for a small percentage of clicks (about 1%). The clicks to record are selected by a "real-time train detection algorithm" running on the FPOD. The information stored is the amplitude and timing (at 250 nanosecond resolution) of the peak of the soundwave. This allows for the reconstruction of the waveform by fitting sine waves to the points stored.

Note that there's an option in the FPOD app to configure the FPOD to record all clicks rather than just a small sample..

Value

Invisibly returns a numeric vector with the frequency values used for plotting

See Also[fp_read\(\)](#)**Examples**

```
# read a FP3 file
fn <- fp_example("gullars_period1.FP3")
dat <- fp_read(fn)

# get the click number of the first five clicks for which we have WAV data.
first5 <- head(unique(dat$wav$click_no))

# plot one of them. Here, we plot number three
fp_plot(dat, first5[3])

# If the legend covers the curve, we can move it somewhere else.
fp_plot(dat, first5[3], legend.pos = "topright")

# If we only want to plot clicks that meet certain criteria, we have to do that
# manually. For example, we can plot 9 random NBHF clicks:
nbhf_clicks <- dat$clicks[species == "NBHF" & has_wav == TRUE, click_no]
old.mfrow = par()$mfrow
par(mfrow = c(3,3)) # we'll do 3 by 3 panels
for (i in sample(nbf_clicks, size = 9)) {
  fp_plot(dat, click_no = i, legend.pos = FALSE)
}
par(mfrow = old.mfrow) # reset graphics device to whatever it was before
```

fp_read

*Read FPOD data***Description**

This function reads an FPOD or CPOD data file (FP1, FP3, CP1, CP3) into R.

Usage

```
fp_read(file, tz = "", simplify = TRUE, amp = "extended")
```

Arguments

file	a character string. The path to the FPOD (or CPOD) data file.
tz	a character string. The time zone specification to be used for calculating dates. Passed unchanged to as.POSIXct() .
simplify	logical. If TRUE, simplifies the clicks data.table by stripping away some columns, such as <code>clk_ipi_range</code> , <code>ipi_pre_max</code> , <code>amp_reversals</code> , <code>duration</code> , and <code>has_wav</code> .
amp	a character string. With <code>amp="extended"</code> , higher values are extrapolated from the duration of clipping and the IPI. For any other values of <code>amp</code> , the compressed SPL values recorded by the FPOD are used directly.

Details

The clicks data.frame contains the following columns:

- pod: the ID number of the pod
- time: The time and date of the click, at microsecond resolution. Note that R might only display dates and times to a second precision, but any date or time calculations will use the full precision.
- minute: minutes elapsed, since starting the FPOD
- microsec: microseconds elapsed, since the start of the minute.
- click_no: an ID number that uniquely identifies the click.
- train_id: an ID number from the KERNO classifier, reset for each minute.
- species: the species classification from the KERNO classifier
- quality_level: the quality level of the classification, 0 (?/echo), 1 (Lo), 2 (Mod) or 3 (Hi).
- echo: TRUE if the KERNO classifier thinks this click might be an echo of another click that has already been classified.
- ncyc: number of cycles in click. This is a proxy for the click duration.
- pkat: the number of the cycle with the highest amplitude
- clk_ipi_range: range of inter-peak-intervals (IPIs) among cycles in click
- ipi_pre_max: the IPI before pkat, in 250 nanosecond units
- ipi_at_max: the IPI at pkat, in 250 nanosecond units
- khz: The frequency (in kHz) of the peak cycle.
- amp_at_max: the peak amplitude (SPL) of the loudest cycle
- amp_reversals: the number of amplitude reversals
- duration: click duration
- has_wav: TRUE if there is a pseudo-WAV recorded for this click.

Value

A list, with one or more of the following data.frames (or data.tables, if available):

- header: a list with pod name, coordinates, starting time, stopping time, user notes, etc.
- clicks: A data.frame (or data.table) with data about each click. See details.
- wav (only FPx files): pseudo-wav data - inter-peak-intervals (in 250 ns units) and raw amplitudes for a subset of clicks
- env: misc data, angle from vertical (in degrees), ambient temperature (in deg C), battery voltage per stack (in units of volts), which battery column is in use, and the pod on/off state.

See Also

[fp_find_buzzes\(\)](#), [fp_summarize\(\)](#)

Examples

```
# read a FP3 file
fn <- fp_example("gullars_period1.FP3")
dat <- fp_read(fn)

# show misc. information (pod number, deployment date, etc.)
dat$header

# show battery levels and recorded temperatures for each minute
dat$env

# tally up the number of clicks in each species category
table(dat$clicks$species)
```

fp_summarize

Calculates minute-resolution summaries of clicks

Description

Calculates minute-resolution summaries of clicks

Usage

```
fp_summarize(x)
```

Arguments

x data.table where each row is a click, as the "clicks" element in the list object returned by `fp_read()`. Each row must minimally have a POSIXct column time. The return value of `fp_read()` is also accepted with a warning, provided that the clicks data.table is present .

Value

A data.table with three or four columns:

- **time**: POSIXct timestamp of the start of the 1-minute time chunk, in YYYY-mm-dd HH:MM format
- **dpm**: detection-positive-minutes, 1 if at least one click is registered during the time chunk; 0 otherwise.
- **bpm**: buzz-positive-minutes, 1 if at least one feeding buzz is registered during the time chunk, 0 otherwise. Note that bpm is only available if there is a 'buzz' column in the clicks data.table, e.g. from first calling `fp_find_buzzes()`

See Also

`fp_find_buzzes()`, `lubridate::floor_date()`

Examples

```
# first read some FPOD data
fn <- fp_example("gullars_period1.FP3")
dat <- fp_read(fn)

# extract porpoise clicks of quality Hi and Mod
nbhf <- dat$clicks[species == "NBHF" & quality_level >= 2]

# get a simple summary, with timestamp, degC and dpm.
dpm <- fp_summarize(nbhf)

# If we also wanted buzz-positive-minutes, we could do this first.
nbhf$buzz <- fp_find_buzzes(nbhf, method = "clicks")

# This time, when we call fp_summarize, it should detect that there's a
# buzz column, and automatically calculate feeding-positive-minutes as well.
dpm <- fp_summarize(nbhf)

# Once we have DPMs, we can easily aggregate these into coarser time chunks
dpm_per_hour <- dpm[, .(dpm = sum(dpm)),
  .(date = as.POSIXct(trunc(time, unit = "hours")))] # per hour
dpm_per_day <- dpm[, .(dpm = sum(dpm)),
  .(date = as.Date(time))] # per day
dpm_per_month <- dpm[, .(dpm = sum(dpm)),
  .(date = as.POSIXct(trunc(time, unit = "months")))] # per month
```

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