

#### hu-neuro-pipeline

A Python-based and R-compatible pipeline for processing single trial EEG data

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# The Frömer et al. (2018) pipeline



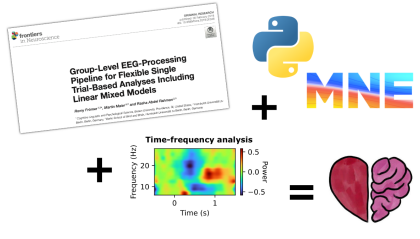
# The Frömer et al. (2018) pipeline



- Allows single trial analysis of ERP amplitudes
  - Random effects for items (Bürki et al., 2018)
  - Trial and item level covariates (Volpert-Esmond et al., 2021)
  - Continuous predictor variables
  - Unbalanced designs
  - Brain-behavior associations

# Python implementation





#### Python, I choose you!



 $\textbf{Blog post:} \ \ \mathsf{https:} / / \mathsf{dominiquemakowski.github.io/post/2020-05-22-r\_or\_python$ 

Online course: https://swcarpentry.github.io/python-novice-inflammation

#### MNE-Python



- Versatile
  - EEG, MEG, ECoG, fNIRS
  - Preprocessing, statistics, time-frequency analysis, visualization, machine learning, connectivity, source localization, . . .
- Open source
  - > 300 contributors on GitHub (January 2023)
  - Funded by NIH, NSF, ERC, Google, Amazon, ...
  - Code review, automated tests, user forum, office hours, ...

### Python implementation



- No MATLAB required
- No Python skills required can be called from R
- New features:
  - Time-frequency analysis
  - Fully automatic ocular correction (ICA)
  - Automatic bad channel detection.
  - Automatic missing trial detection
- Code standards + version control (https://github.com/alexenge/hu-neuro-pipeline/)

# Python implementation



Read raw da	ata <					
*						
(Downsample)						
Read channel locations		(Auto-detect bad channels)				
		det				
(Interpolate bad channels)		eg .				
Re-reference to average		bad o				
thar						
Ocular correction (BESA/ICA)						
Frequency filter						
rrequency n	iter					
Segment to ep	nochs					
+						
Read + match	log file					
Reject bad ep	ochs					
Compute single	Compute	ъ	-	- 77		
trial ampl.	evokeds	Participant 01	Participant 02	Participant 03		
trial diripii	evolues.	g:	<u>₽</u> .	₽.		
(QC report)		Ē.	ä	Ħ		
		2	8	8	1	
- t	-					
Combine trials	Combine evokeds		(Cluster-based			S.
			permutation			ę
Linear mixed model	Grand average		test)			Group pipeline
						a in

#### Installation



#### For Python users:

```
# Install via the command line from the Python Packaging Index (PyPI) python3 \mbox{-m} pip install hu-neuro-pipeline
```

#### For R users:

```
# Install reticulate for interfacing with Python from R
install.packages("reticulate")

# Install Python (Miniconda distribution)
reticulate::install_miniconda()

# Install the actual package from PyPI
reticulate::py_install("hu-neuro-pipeline", pip = TRUE, python_version = "3.8")
```

# General usage



```
# Import the Python package
pipeline <- reticulate::import("pipeline")

# Run the pipeline
res <- pipeline$group_pipeline(...)</pre>
```

## Minimal example



For details, see Frömer et al. (2018), https://doi.org/10.3389/fnins.2018.00048

### Minimal example



```
# Import the Python package
pipeline <- reticulate::import("pipeline")</pre>
# Run the pipeline
res <- pipeline$group pipeline(
  # Input/output paths
 vhdr_files = "data/raw",
 log_files = "data/log",
 output_dir = "output",
  # Preprocessing options
 besa_files = "data/cali",
  # Epoching options
 triggers = c(201:208, 211:218).
 components = list(
   "name" = list("N2", "P3b"),
   "tmin" = list(0.25, 0.4),
    "tmax" = list(0.35, 0.55),
    "roi" = list(
      c("FC1", "FC2", "C1", "C2", "Cz"),
     c("CP3", "CP1", "CPz", "CP2", "CP4", "P3", "Pz", "P4", "P03", "P0z", "P04")
 ).
  # Averaging options
 average_by = c("n_b", "DeviantPosRL", "n_b/DeviantPosRL")
```



```
# Input/output paths
vhdr_files = "data/raw",
log_files = "data/log",
output_dir = "output",
```

- Directory or list of raw EEG files (.vhdr)
- Directory or list of behavioral log files (.txt/.tsv/.csv)
- Output directory



```
# Preprocessing options
besa_files = "data/cali",
```

- Directory path or list of BESA files (.matrix)
- Default bandpass filter (0.1–40 Hz)



```
# Epoching options
triggers = c(201:208, 211:218),
components = list(
    "name" = list("N2", "P3b"),
    "tmin" = list(0.25, 0.4),
    "tmax" = list(0.35, 0.56),
    "roi" = list(
        c("FG1", "FC2", "C1", "C2", "C2"),
        c("CP3", "CP1", "CP2", "CP4", "P3", "P2", "P4", "P03", "P02", "P04")
    )
),
```

- List of numerical EEG triggers
- List of ERP component definitions:
  - name: Column names for each component
  - tmin + tmax: Onset and offset times (in s)
  - roi: List of channel names for each component



```
# Averaging options
average_by = c("n_b", "DeviantPosRL", "n_b/DeviantPosRL")
```

 List of column names (for main effects) and combinations of column names (for interaction effects, separated by "/")

#### More pipeline inputs



- Downsampling (downsample\_sfreq)
- Interpolate bad channels (bad\_channels)
- Frequency filter (highpass\_freq, lowpass\_freq)
- Epoch duration (epochs\_tmin, epochs\_tmax)
- Baseline duration (baseline)
- Skip log file rows (skip\_log\_rows, skip\_log\_conditions)
- Threshold for artifact rejection (reject\_peak\_to\_peak)
- . . .



#### Extract directly from the pipeline run:

```
trials <- res[[1]] # Single trial data frame
evokeds <- res[[2]] # Evokeds data frame
config <- res[[3]] # List of pipeline options
```

#### Or read from the output directory:

```
library(tidyverse)
trials <- read_csv("output/trials.csv")
evokeds <- read_csv("output/ave.csv")
config <- jsonlite::read_json("output/config.json")</pre>
```

See https://github.com/alexenge/hu-neuro-pipeline/blob/main/docs/outputs.md

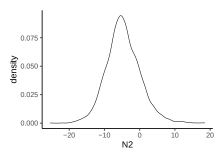


```
# Single trial data frame
print(trials)
## # A tibble: 3.840 x 32
     participa~1 VPNum~2 version wdh lfdNr n b Stand~3 Deviant Objek~4 BedCo~5
     <chr>
                            <dbl> <dbl> <dbl> <chr> <chr> <chr>
                                                                     <dbl> <chr>
##
                    <db1>
## 1.05
                                           1 norm~ objekt~ objekt~
                                                                         8 gngf
                                         2 blurr objekt~ objekt~
## 2.05
                                                                        10 un
## 3 05
                                    1 3 blurr objekt~ objekt~ 10 un
## 4 05
                               1 1 4 norm~ objekt~ objekt~ 3 un
                               1 1 5 norm~ objekt~ objekt~ 15 unuf
## 5.05
                               1 1 6 norm- objekt- objekt- 7 unuf
1 1 7 blurr objekt- objekt- 2 un
1 1 8 norm- objekt- objekt- 16 gngf
## 6 05
## 7 05
## 8.05
## 9 05
                                         9 norm~ objekt~ objekt~
## 10 05
                                          10 norm~ objekt~ objekt~
                                                                        11 un
## # ... with 3.830 more rows, 22 more variables: BedCode neu <chr>, bot <dbl>,
## #
      DeviantPosRL <chr>, DeviantPosNR <dbl>, BedCodeRL <chr>, kev <dbl>,
      ErrorCode <dbl>, RT <dbl>, Pos1 <chr>, Pos2 <chr>, Pos3 <chr>, Pos4 <chr>,
## #
## #
      Pos5 <chr>, Pos6 <chr>, Pos7 <chr>, Pos8 <chr>, Pos9 <chr>, Pos10 <chr>,
## #
      Pos11 <chr>, Pos12 <chr>, N2 <dbl>, P3b <dbl>, and abbreviated variable
      names 1: participant id, 2: VPNummer, 3: Standard, 4: Objektpaar,
## #
      5: BedCode_alt
## #
```



```
# Single trial N2 mean amplitudes
ggplot(trials, aes(x = N2)) +
  geom_density() +
  theme_classic(base_size = 30)
```

## Warning: Removed 7 rows containing non-finite values ('stat\_density()').

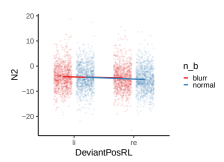




```
# Linear mixed-effects model
form <- N2 ~ n_b * DeviantPosRL + (1 | participant_id)
mod <- lme4::lmer(form, trials)
summary(mod)
## Linear mixed model fit by REML ['lmerMod']
## Formula: N2 ~ n_b * DeviantPosRL + (1 | participant_id)
##
     Data: trials
##
## REML criterion at convergence: 22696.1
##
## Scaled residuals:
##
      Min
               10 Median
                                     Max
## -4.6856 -0.6130 -0.0002 0.6148 5.1121
##
## Random effects:
## Groups
                             Variance Std.Dev.
                  Name
## participant_id (Intercept) 2.287 1.512
## Residual
                             21.780 4.667
## Number of obs: 3833, groups: participant_id, 2
##
## Fixed effects:
                           Estimate Std. Error t value
##
## (Intercept)
                          -4.2285 1.0800 -3.915
## n_bnormal
                          -0.1796 0.2132 -0.842
                          -0.4587 0.2132 -2.151
## DeviantPosRLre
## n bnormal:DeviantPosRLre -0.4732 0.3015 -1.569
```



```
# Single trial N2 mean amplitudes by condition
ggplot(trials, aes(x = DeviantPosRL, y = N2, color = n_b, group = n_b)) +
geom_point(position = position_jitterdodge(0.3), alpha = 0.1) +
stat_summary(
    geom = "line",
    size = 2.,
    position = position_dodge(0.75)
) +
theme_classic(base_size = 30)
```





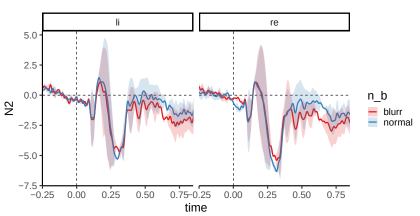
```
# Evokeds by participant and condition
print(evokeds)
```

```
## # A tibble: 16,000 x 70
##
     particip~1 avera~2 n b Devia~3 time
                                              Fp1
                                                    Fpz Fp2 AF7 AF3 AFz
     <chr>
##
                <chr>
                       <chr> <chr>
                                      <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <
##
  1 05
               n b
                       norm~ <NA>
                                  -0.5 -0.751 -1.49 -1.35 -1.14 -1.94 -1.86
## 2.05
               n b
                                  -0.498 -0.779 -1.46 -1.30 -1.16 -2.04 -1.93
                       norm~ <NA>
## 3 05
                       norm~ <NA>
                                  -0.496 -0.809 -1.43 -1.25 -1.15 -2.11 -2.00
               n_b
## 4 05
               n b
                      norm~ <NA> -0.494 -0.836 -1.40 -1.21 -1.13 -2.14 -2.06
## 5 05
               n b
                     norm~ <NA> -0.492 -0.859 -1.39 -1.17 -1.10 -2.14 -2.10
## 6 05
                     norm~ <NA> -0.49 -0.876 -1.39 -1.15 -1.05 -2.10 -2.12
               n_b
                n_b norm~ <NA> -0.488 -0.886 -1.42 -1.14 -1.01 -2.06 -2.12
## 7 05
## 8 05
                n b
                       norm~ <NA> -0.486 -0.893 -1.46 -1.15 -0.976 -2.00 -2.10
## 9 05
                       norm~ <NA> -0.484 -0.896 -1.51 -1.17 -0.952 -1.95 -2.09
                n_b
                       norm~ <NA> -0.482 -0.898 -1.56 -1.19 -0.942 -1.90 -2.07
## 10 05
                n_b
## # ... with 15.990 more rows, 59 more variables: AF4 <dbl>, AF8 <dbl>, F9 <dbl>,
## #
      F7 <dbl>, F5 <dbl>, F3 <dbl>, F2 <dbl>, F4 <dbl>, F6 <dbl>, F8 <dbl>,
## #
      F10 <dbl>, FT7 <dbl>, FC5 <dbl>, FC3 <dbl>, FC1 <dbl>, FC2 <dbl>,
## #
      FC4 <dbl>, FC6 <dbl>, FT8 <dbl>, T7 <dbl>, C5 <dbl>, C3 <dbl>, C1 <dbl>,
## #
      Cz <dbl>, C2 <dbl>, C4 <dbl>, C6 <dbl>, T8 <dbl>, TP9 <dbl>, TP7 <dbl>,
      CP5 <dbl>, CP3 <dbl>, CP1 <dbl>, CPz <dbl>, CP2 <dbl>, CP4 <dbl>,
## #
      CP6 <dbl>, TP8 <dbl>, TP10 <dbl>, P7 <dbl>, P5 <dbl>, P3 <dbl>, ...
## #
```



```
# Evokeds by participant/condition
evokeds %>%
 filter(average by == "n b/DeviantPosRL") %>%
 Rmisc::summarySEwithin(
   measurevar = "N2",
   withinvars = c("time", "n_b", "DeviantPosRL"),
   idvar = "participant id"
 ) %>%
 mutate(time = as.numeric(levels(time))[time]) %>%
 ggplot(aes(
   x = time,
   v = N2
   vmin = N2 - se.
   vmax = N2 + se.
    color = n_b,
   fill = n b
 )) +
 facet_wrap(~DeviantPosRL) +
 geom hline(vintercept = 0, linetype = "dashed") +
 geom vline(xintercept = 0, linetype = "dashed") +
  geom_line(size = 1) +
 geom_ribbon(color = NA, alpha = 0.2) +
  coord cartesian(xlim = c(-0.2, 0.8)) +
 theme_classic(base_size = 20)
```







## # List of pipeline options names(config)

```
[1] "whdr files"
                                "log files"
                                                       "output dir"
## [4] "clean dir"
                                "epochs dir"
                                                       "report dir"
                                "downsample_sfreq"
                                                       "veog_channels"
## [7] "to_df"
## [10] "heog channels"
                                "montage"
                                                       "bad channels"
## [13] "besa_files"
                                "ica method"
                                                       "ica_n_components"
## [16] "highpass_freq"
                                                       "triggers"
                                "lowpass_freq"
## [19] "triggers column"
                                "epochs tmin"
                                                       "epochs tmax"
## [22] "baseline"
                                "skip_log_rows"
                                                       "skip log conditions"
                                "components"
                                                       "average_by"
## [25] "reject_peak_to_peak"
## [28] "perform_tfr"
                                "tfr_subtract_evoked"
                                                       "tfr_freqs"
## [31] "tfr_cycles"
                                "tfr mode"
                                                       "tfr baseline"
## [34] "tfr_components"
                                "perm_contrasts"
                                                       "perm_tmin"
## [37] "perm_tmax"
                                "perm_channels"
                                                       "perm_fmin"
## [40] "perm fmax"
                                "n_jobs"
                                                       "auto rejected epochs"
```

# Number of rejected epochs per participant
lengths(config\$auto\_rejected\_epochs)

```
## 05 07
## 7 0
```

### More pipeline outputs



- Cleaned continuous data (clean\_dir)
- Epoched data (epochs\_dir)
- Automated QC reports (reports\_dir)

#### Cluster-based permutation tests



```
# Permutation test input
perm_contrasts = list(
 c("blurr", "normal"),
 c("blurr/re", "blurr/li"),
 c("normal/re", "normal/li")
# Permutation test outputs
clusters <- read csv("output/clusters.csv") # or clusters <- res[[4]]
print(na.omit(clusters))
## # A tibble: 5.748 x 6
     contrast
                   time channel t_obs cluster p_val
     <chr>
                    <dbl> <chr> <dbl> <chr>
                                                <dh1>
## 1 blurr - normal 0
                        AF3
                                -18.1 neg 282
## 2 blurr - normal 0 FT7
                                   52.8 pos_1
## 3 blurr - normal 0 C6
                                  -30.0 neg_281
## 4 blurr - normal 0 POz
                                   15.3 pos_96
## 5 blurr - normal 0.002 F10
                                  -24.5 neg_280
## 6 blurr - normal 0.002 FT7
                                   20.1 pos_1
## 7 blurr - normal 0.002 FC3
                                   17.5 pos 95
## 8 blurr - normal 0.002 FC1
                                   22.1 pos 95
## 9 blurr - normal 0.002 C6
                                 -154. neg_281
## 10 blurr - normal 0.002 CPz
                                   16.6 pos 94
## # ... with 5.738 more rows
```

#### Artifact correction



- Multiple source eye correction (MSEC)
  - Requires .matrix files from BESA

#### Artifact correction



#### • Independent component analysis (ICA)

- Different algorithms available (e.g., ica\_method = "fastica")
- Can specify initial number of principal components with ica\_n\_components
- Automatic detection + exclusion of eye movement components based on correlation with HEOG and VEOG (see https://mne.tools/ stable/generated/mne.preprocessing.ICA.html#mne.preprocessing.ICA.find\_bads\_eog)
- Verify in QC reports
- Other selection methods (manual selection, ICLabel) not yet implemented

#### Artifact rejection



- Per-channel peak-to-peak amplitude threshold via reject\_peak\_to\_peak (default: 200.0)
- In addition to or instead of BESA or ICA

## Repairing bad channels



- Pass participant-specific vectors of bad channel labels
  - E.g., bad\_channels = list("05" = c("C3", "P7"),
    ...)
- Uses spherical spline interpolation
- Experimental: Automatic bad channel detection (bad\_channels = "auto")
  - Based on channel SDs across epochs

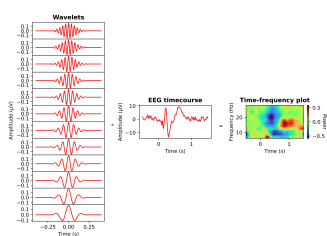
#### Detecting missing epochs



- Requires log file column (triggers\_column) with the EEG trigger for every trial
- Pipeline magically detects and deletes log file trials with missing EEG

#### Time-frequency analysis





#### Time-frequency analysis

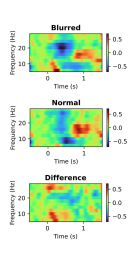


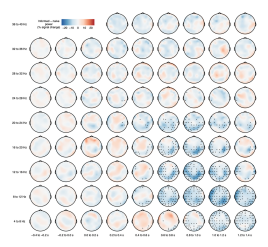
```
# Time-frequency analysis options
perform_tfr = TRUE,
tfr_components = list(
    "name" = list("alpha"),
    "tmin" = list(0.0), "tmax" = list(0.2),
    "fmin" = list(8.0), "fmax" = list(14.0),
    "roi" = list(c("PO9", "PO7", "PO3", "PO2", "PO4", "PO8", "PO10", "O1", "O2", "O2"))
)
```

- tfr\_components extracts single trial power values
- Additional options:
  - Morlet frequencies (tfr\_freqs, default 4, 5, 6, ..., 40 Hz)
  - Morlet no. of cycles (tfr\_cycles, default 2, 2.5, 3, ..., 20)
  - Baseline window (tfr\_baseline, default -450 ms to -50 ms)
  - Baseline method (tfr\_method, default percent signal change)

## Time-frequency analysis







#### **Plans**



- Proper documentation / preprint
- Unit tests
- RIDE correction for speech artifacts
- Mixed models with pymer4 (?)
- Better permutation tests (Frossard & Renaud, 2021, 2022)
- BIDS interface
- Your ideas + contributions?

## **Thanks**



#### References



- Bürki, A., Frossard, J., & Renaud, O. (2018). Accounting for stimulus and participant effects in event-related potential analyses to increase the replicability of studies. Journal of Neuroscience Methods, 309, 218-227. https://doi.org/10.1016/j.jneumeth.2018.09.016
- Frömer, R., Maier, M., & Abdel Rahman, R. (2018). Group-level EEG-processing pipeline for flexible single trial-based analyses including linear mixed models. Frontiers in Neuroscience, 12, 48. https://doi.org/10.3389/fnins.2018.00048
- Frossard, J., & Renaud, O. (2021). Permutation tests for regression, ANOVA, and comparison of signals: The permuco package. Journal of Statistical Software, 99, 1-32. https://doi.org/10.18637/jss.v099.i15
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- Volpert-Esmond, H. I., Page-Gould, E., & Bartholow, B. D. (2021). Using multilevel models for the analysis of event-related potentials. International Journal of Psychophysiology, 162, 145–156. https://doi.org/10.1016/j.ijpsycho.2021.02.006