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NeuroKit2: A Python Toolbox for Neurophysiological Signal Processing

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15 Abstract

NeuroKit2 is an open-source, community-driven and user-friendly Python package
dedicated to neurophysiological signal processing (e.g., ECG, EDA, EMG, ...). Its design
philosophy, which is centred on user-experience and a collaborative environment, makes it
accessible to both novice and advanced users. The package provides general functions
allowing for data processing and analysis in a few lines of code using validated pipelines,
as well as tools dedicated at specific processing steps, offering flexibility and fine-tuned
control for advanced users. NeuroKit2 aims at improving transparency and reproducibility
in neurophysiological research, as well as being a scaffolding for exploration and
innovation.

25 Keywords: Neurophysiology, Biosignals, Python, ECG, EDA, EMG

26 Word count:

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Cognitive neuroscience and psychology are increasingly relying on neurophysiological methods to assess brain and bodily activity. These approaches include electroencephalography 29 (EEG), electrocardiography (ECG), electromyography (EMG) and electrodermal activity 30 (EDA) signals. This trend was driven not only by theoretical motivations (e.g., the growth 31 of embodied or affective neuroscience; Kiverstein & Miller, 2015), but also by practical rea-32 sons including low monetary cost (especially compared with other imaging techniques, such 33 as MRI or MEG), high user convenience (e.g., portability, setup speed), and the increasing availability of recording devices (e.g., in smart watches; Yuehong, Zeng, Chen, & Fan, 35 2016). Together with the development of recording tools, advancements in the fields of signal processing and computational data science bolstered the emergence of new processing 37 algorithms (Clifton, Gibbons, Davies, & Tarassenko, 2012; Roy et al., 2019), in turn offering a myriad of novel methods for users to process and analyze neurophysiological signals.

- Unfortunately, because most of the algorithms are implemented as code, neurophysiological data processing remains a challenge for many researchers without a formal training or experience in programming. Moreover, many existing implementations are also limited to one type of signals (for instance, focused on ECG or EDA), which makes it inconvenient for researchers who might have to learn and concurrently rely on different software to process multimodal data.
- NeuroKit2 aims at addressing these challenges by offering a free, user-friendly, and comprehensive solution for neurophysiological data processing. It is an open-source Python
 package, developed in a collaborative environment that continues to welcome contributors
 from different countries and fields. Historically, NeuroKit2 is the re-forged successor NeuroKit1 (https://github.com/neuropsychology/NeuroKit.py), a PhD side project that ended
 up benefiting an important community of users (252 GitHub stars as of 03-05-2020). The
 new version takes on its best features and design choices, and re-implements them in a

professional and well-thought way. It aims at being 1) accessible and well-documented, 2) reliable and cutting-edge, and 3) flexible and efficient.

Being available for Python 3 (Van Rossum & Drake, 2009), one of the most popular program-55 ming languages, NeuroKit2's users benefit from an important amount of existing tutorials 56 and a large online community. The package is also relatively lightweight, using mainly standard dependencies (Virtanen et al., 2020) such as NumPy, pandas, SciPy, scikit-learn and MatplotLib (with an additional system of optional dependencies), enabling its use as 59 a dependency in other software. The package source code is available under a permissive license on GitHub (https://qithub.com/neuropsychology/NeuroKit). Its documentation, au-61 tomatically built and rendered from the code, is hosted at https://neurokit2.readthedocs.io/. Apart from guides for installation and contribution, and a description of the package's func-63 tions, the documentation also includes several "hands-on" examples and tutorials providing a walk-through on how to address specific issues (e.g., how to extract and visualize individual heartbeats, how to analyze event-related data). New examples can be easily added by users simply by uploading a Python notebook file (Kluyver et al., 2016) to the repository. This file will be automatically transformed into a webpage and displayed on the website, ensuring a transparent and evolutive documentation. Moreover, these examples can be used interactively via a cloud-based Binder environment (Jupyter et al., 2018), allowing users to try out the features directly in their browser. Finally, the accessibility for newcomers is reinforced by the issue tracker of GitHub, where users can create public issues to inquire for help. 73

The package aims at being reliable and trustworthy, including peer-reviewed processing pipelines and functions tested against existing implementations of established reference software such as *BioSPPy* (Carreiras et al., 2015), *hrv under review*, *PySiology* (Gabrieli, Azhari, & Esposito, 2019), *HeartPy* (Gent, Farah, Nes, & Arem, 2019), *systole* (Legrand & Allen, 2020) or *nolds* (Schölzel, 2020). The code itself includes a comprehensive test suite using continuous integration tools (e.g., ???) to ensure stability and prevent errors. Moreover,

⁸⁰ users are able to easily report any bugs and be notified of their fixes via the issue tracker.

Thanks to its collaborative and open development as well as its modular organization, Neu-

roKit2 can easily follow the latest developments and remain cutting-edge through its ability

to evolve, adapt, and integrate new methods as they are emerging.

Finally, we believe that the design philosophy of NeuroKit2 contributes to an efficient (i.e.,

allowing to achieve a lot with few functions) yet flexible (i.e., enabling fine control and

precision over what is done) user interface (API). We will illustrate these claims with two

examples of common use-cases (the analysis of event-related and resting state data), and will

88 conclude by discussing how NeuroKit2 contributes to neurophysiological research by raising

the standards for validity, reproducibility and accessibility.

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Design Philosophy

As stated above, NeuroKit2 aims at being accessible to beginners and, at the same time,

ning users to implement complex processing and analyses pipelines with very few functions.

offering a maximal level of control to experienced users. This is achieved by allowing begin-

while still enabling fine-tuned control and precision to more experienced users. In concrete

95 terms, this trade-off is allowed by the implementation of three abstract levels of functions.

₉₆ Low-level: Base Utilities for Signal Processing

The basic building blocks are functions to facilitate general signal processing, i.e., to do filtering, resampling, interpolating, peak detection, etc. These functions are signal-agnostic, and include a lot of tweakable parameters. For instance, one can change the filtering method, frequencies, order and such. Most of these functions are based on validated algorithms present in *scipy* (Virtanen et al., 2020). Examples of such functions include signal_filter(), signal_interpolate(), signal_resample(), signal_detrend(), and signal_findpeaks().

Mid-level: Neurophysiological Processing Steps

The signal processing utilities are then used by functions specific to the different types of 105 physiological signals (i.e., ECG, RSP, EDA, EMG, PPG). These functions aim at taking care 106 of specific steps of physiological data processing, such as cleaning, peak detection, phase 107 classification or rate computation. Critically, for each type of signals, the same function 108 names are called (in the form signaltype_functiongoal()) to achieve equivalent goals, e.g., 109 *_clean(), *_findpeaks(), *_process(), *_plot(), making it intuitive and consistent 110 across different modalities. 111 For example, the rsp clean() function uses signal filter() and signal detrend(), 112 with different possible sets of default parameters that can be switched via a "method" ar-113 gument (corresponding to different published or validated pipelines). For instance, setting 114 method="khodadad2018" will use the cleaning workflow described in Khodadad et al. (2018). 115 However, if a user wants to build its own custom cleaning pipeline, she or he can use the 116

cleaning function as a template, and directly tweak the parameters in the low-level signal

High-level Wrappers for Processing and Analysis

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processing operations.

These steps are then assembled in "master" functions, that are usually the entry point for 120 new users. For instance, the ecg process() function uses ecg clean(), ecg findpeaks(), 121 ecg rate(). A general processing pipeline can be selected via the method argument, that is 122 then propagated throughout its lower-level functions. Easily switching between processing pipelines allows for the comparison of different methods, and streamlines critical but timeconsuming steps in reproducible research, such as the validation of data preparation and 125 quality control (Quintana, Alvares, & Heathers, 2016). Finally, the package includes conve-126 nience meta-functions (e.g., bio process) that enable the combined processing of multiple 127 types of signals at once (e.g., bio_process(ecg=ecg_signal, eda=eda_signal)). 128

Performing a set of operations with sensible default parameters can be rewarding, especially for beginners, allowing them to perform cutting-edge processing or replication of research steps without requiring a programming expertise. Moreover, it contributes to the demystification of the usage of "pure" programming tools (as opposed to GUI-based software such as SPSS, Kubios, or Acqknowledge), providing a welcoming framework to further understand the frontend, backend and the in-betweens of physiological data processing. Importantly, more advanced users can again very easily build their own custom analysis pipeline by using the mid-level functions, allowing for a finer control over the processing parameters.

Overall, we believe that this code structure offers a calibrated trade-off between flexibility and user-friendliness, with functions that are easy to memorize and implement. We hope that it may further encourage researchers to become part of a supportive open-science community construed of many expertises - rather than relying on closed and proprietary software - to achieve their goals.

142 Example

In this section, we present two examples that illustrate the most common use-cases. The
first example is an event-related paradigm, in which the interest lies in the momentarily
short-term physiological changes related to specific stimuli, whereas the second shows how
to extract the characteristics (features) of physiological activity during a longer period of
time (not necessarily tied to a specific and sudden event). The example datasets are made
available with the package and can be downloaded using the data() function.

149 Event-related Paradigm

This example dataset contains ECG, RSP and EDA signals of one participant to whom were presented four emotional images (from the NAPS database; Marchewka, Żurawski, Jednoróg, & Grabowska, 2014), in a typical (albeit highly shortened) experimental psychology

paradigm.

Signals are 2.5 minutes long and are recorded at a frequency of 100Hz (note that the sampling rate is low for storage purposes and should be higher in actual recordings, see Quintana et al., 2016). It has 4 channels including three physiological signals, and one corresponding to the marking of events via a photosensor (which signal decreased when a stimulus appeared on the screen).

```
# Load the package
import neurokit2 as nk

# Download the example dataset
data = nk.data("bio_eventrelated_100hz")

# Visualize 10 seconds of data (on the same scale)
nk.signal_plot(data[900:1900], standardize=True)
```

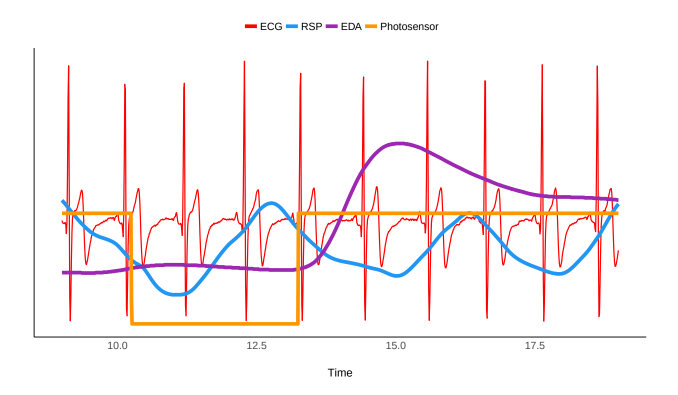


Figure 1. Subset of the dataset showing one event (in orange) and the other physiological signals.

```
# Extract event related features
results = nk.bio_analyze(epochs)

# Show subset of results
results[["Condition", "ECG_Rate_Mean", "RSP_Rate_Mean", "EDA_Peak_Amplitude"]]
```

Table 1
Subset of the outure related to event-related analysis characterizing the pattern of physiological changes related to specific stimuli.

Condition	ECG_Rate_Mean	RSP_Rate_Mean	EDA_Peak_Amplitude
Negative	-1.94	-0.22	None
Neutral	-4.36	1.57	None
Neutral	1.02	-0.30	None
Negative	-3.61	2.22	1.68

In this example, after loading the package and the example dataset, each physiological signal is processed using bio_process(). Data from the photosensor is processed separately with events_find(), that locates the stimuli onsets in the signal. Once we have preprocessed signals and the location of events, we can slice the data into segments corresponding to a time window (ranging from -0.1 to 4 seconds) around each stimulus with epochs_create(). Finally, relevant features are computed for each epoch (i.e., each stimulus) by providing them to bio_analyze().

The features include for example the changes in rates of ECG and RSP signals (e.g. maximum, minimum and mean rate after stimulus onset, and the time at which they occur),
and the peak characteristics of EDA signal (e.g., occurrence of skin conductance response
(SCR), and if SCR is present, its corresponding peak amplitude, time of peak, rise and recovery time). In addition, respiration and cardiac cycle phases are also extracted (i.e., the
respiration phase - inspiration/expiration - and cardiac phase - systole/diastole - occurring
at the onset of event).

This example shows the straightforward process of extracting features of physiological responses. This pipeline can easily scale up to group-level analysis by aggregating the average

of features across subjects. In addition to streamlining data analyses, NeuroKit2 aims to provide researchers an extensive suite of signal features, allowing for precise interpretations in terms of relationship between physiological activity and neurocognitive processes. In this example (see **Table 1**), exposure to negative stimuli, as compared to neutral stimuli, is related to stronger cardiac deceleration, higher skin conductance response, and accelerated breathing rate (note that this illustrative interpretation is purely descriptive).

181 Resting-state Features

The second dataset corresponds to 5 minutes physiological activity of a human participant at rest (eyes-closed in a seated position), under no specific set of instructions. It contains three channels (ECG, PPG and RSP) sampled at a frequency of 100Hz.

results[["ECG_Rate_Mean", "ECG_HRV_RMSSD", "RSP_Rate_Mean", "RSA_P2T_Mean"]]

Table 2
Subset of properties characterizing the physiological activity over a period of 5 minutes of resting-state.

ECG_Rate_Mean	ECG_HRV_RMSSD	RSP_Rate_Mean	RSA_P2T_Mean
86.42	4.28	15.86	0.01

In this example, the steps of the analysis are in fact identical to the previous example, 185 including loading the package, the dataset and processing the data. The difference is that 186 here there is no epoching, as we want to compute features related to the whole dataset. 187 Thus, we can directly pass the dataframe to bio analyze(), which will detect that these 188 are not epochs, and compute the appropriate features accordingly. These include for instance 189 the average heart and breathing rate, as well as indices of heart rate variability (HRV) and 190 respiratory sinus arrhythmia (RSA). 191 This example illustrates a second type of physiological analysis, that we could refer to as 192 interval-related (as opposed to event-related), where one is typically interested in computing 193 features of signal variability and activation patterns over a longer-term period or interval of 194 time (typically more than a few seconds, as is generally the case in event-related paradigms). 195 The simplicity of usage of NeuroKit2 allows for the fast creation of a standardized and 196

198 Discussion

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reproducible pipeline to describe physiological activity.

NeuroKit2 is a neurophysiological signal processing software accessible to people with very little knowledge of programming, due to its design choices focusing on user-experience and collaborative community. It is also a pragmatic answer to the broader need for transparent

and reproducible methods in neurophysiology. Its modular organization not only facilitates
the use of existing and validated processing pipelines, but is also an viable platform for
experimentation and innovation.

We expect future evolution to be mostly driven by the community and the advances in related fields. Possible directions include extending the support for other types of bodily signals (e.g., electrogastrography - EGG, electrooculography - EOG) and strengthening the efficiency of the code to obtain performance gains for large datasets. Further validation of the available processing pipelines could be made through the (re)analysis of public databases. In line with this objective, the support of standardized data structure formats (e.g. WFDB, BIDS, ...) could be extended.

In conclusion, we think NeuroKit2 provides useful tools not only for novice and senior 212 researchers, but amateurs and tech-enthusiasts interested in health or embodied aspects 213 of (neuro)psychology. Whether the data is produced by "smart health devices" or aca-214 demic research-grade equipment, the package foster reproducible science. Critically, it is a 215 community-oriented project. By increasing the autonomy of researchers and practitioners, 216 and by shortening the delay between data collection and results acquisition, NeuroKit2 could 217 be useful beyond fundamental research in neuroscience and psychology, including applica-218 tions such as biofeedback, personal physiological monitoring and sport research. 219

Conflict of Interest

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The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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