pyPhotometry: Open source Python based hardware and software for fiber photometry data acquisition

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Fiber photometry is the process of recording bulk neural activity by measuring fluorescence changes in activity sensitive indicators such as GCaMP through an optical fiber. We present a system of open source hardware and software for fiber photometry data acquisition consisting of a compact, low cost, data acquisition board built around the Micropython microcontroller, and a cross platform graphical user interface (GUI) for controlling acquisition and visualising signals. The system can acquire two analog and two digital signals, and control two external LEDs via built in LED drivers. Time-division multiplexed illumination allows independent readout of fluorescence evoked by different excitation wavelengths from a single photoreceiver signal. Validation experiments indicate this approach offers better signal to noise for a given average excitation light intensity than sinusoidally-modulated illumination.

pyPhotometry is substantially cheaper than commercial hardware filling the same role, and we anticipate, as an open source and comparatively simple tool, it will be easily adaptable and therefore of broad interest to a wide range of users.

Fiber photometry has emerged as an important tool for behavioural neuroscience, which allows for the measurement of neuronal activity from genetically-defined neuronal populations or axonal projections in superficial or deep brain structures in behaving animals¹–¹¹. The rapid development of novel genetically encoded fluorescent indicators for intracellular calcium¹²–¹⁶ and chloride¹⁷, neurotransmitter release¹⁸–²⁰, and membrane voltage²¹–²³ suggest that experimental applications for fiber photometry will continue to grow rapidly.

In fiber photometry experiments, excitation light of one or more wavelengths is transmitted to the brain structure of interest through an optic fiber where it excites one or more fluorescent indicators. Light emitted by the indicators is transmitted back through the optic fiber, separated from the illumination light and fluorescence from other indicators using optical filters, and converted to electrical signals by high sensitivity photoreceivers or cameras. These electrical signals are digitised and constitute the data generated by the experiment. Excitation light of different wavelengths may be differentially modulated to allow the fluorescence evoked by each to be demultiplexed from a single photodetector signal, for example to independently measure fluorescence evoked from the genetically encoded calcium indicator GCaMP by 470 nm and 405 nm excitation light. As GCaMP is approximately isosbestic at 405 nm (i.e. fluorescence is independent of calcium concentration) this gives a calcium sensitive signal and a calcium insensitive signal that can be used to control for movement artefacts⁵.

Turn-key commercial systems exist for fiber photometry data acquisition that are convenient but expensive. Alternatively acquisition can be controlled using generic hardware such as that from National Instruments, but this requires a substantial investment of time to setup as well as proprietary hardware and software licences. We sought to develop an open source acquisition system that offered the convenience of commercial hardware at low cost. As well as democratising access to new experimental methods, open source hardware can help to improve reproducibility as the entire signal acquisition and processing pipeline is open, and facilitate new applications as researchers can modify tools themselves²⁴–²⁷.

pyPhotometry is a system of hardware and software consisting of an acquisition board and graphical user interface (GUI). The system implements the following functionality: (1) Digitisation of 2 analog voltage signals

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(at 15-bit resolution) and 2 digital signals. (2) Two constant current LED driver circuits with a 0–100 mA output. (3) Control of data acquisition and online visualisation of signals via the GUI. (4) Streaming of acquired data to disk in a compact binary format. (5) Time-division multiplexed illumination to prevent crosstalk between fluorescence signals and bleed-through of ambient light, with online demultiplexing and visualisation. (6) User documentation at https://pyphotometry.readthedocs.io.

We report the system design and rationale, validation experiments characterising system performance, and data showing its application to recording calcium signals from VTA dopamine neurons.

Results

Acquisition board. The acquisition board is built around the Micropython Pyboard, an ARM Cortex microcontroller that is programmed in Python (Fig. 1). Programming the firmware in a high level language allows it to be simple and compact (<200 lines of code), facilitating rapid development. The acquisition board has 2 analog and 2 digital inputs, each a BNC connector.

Analog signals are acquired using the Pyboard's analog to digital converters (ADCs). These are 12-bit ADCs with a 0–3.3 V range. Oversampling is used to increase the resolution to 15-bit, i.e. for each sample the ADC is read 64 times and the values averaged to give an extra 3 bits of resolution. This gives a resolution of ~10⁻⁴ volts per division of the digitised signal.

The digital inputs are read at the same sampling rate as the analog signals and their intended use is to enable post hoc synchronisation of the photometry data with other systems such as behavioural hardware. We typically use one digital input to receive synchronisation pulses from the behavioural hardware and the other to receive a TTL each time a behavioural event of particular interest (e.g. reward delivery) occurs. This allows the average response to the event of interest to be visualized online (see below), and the response to other behavioural events to be calculated offline by using the sync pulse times to align behavioural and photometry data. We typically use sync pulses with random inter-pulse intervals as this creates a unique and unambiguous match between pulse times recorded on different systems (see https://pycontrol.readthedocs.io/en/latest/user-guide/synchronisation/). The digital inputs connect directly to pins on the microcontroller so in principle could be used as outputs, e.g. for triggering closed loop stimulation, though this is not currently supported by the firmware.

The LED driver circuits are voltage controlled current sinks implemented using an op-amp and MOSFET, adapted from Figure 200 of 30. Their outputs are M8 connectors compatibles with commonly-used connectorised LEDs from Doric Lenses or Thorlabs. The LED driver circuit is linear over a 1–100 mA current range and responds to control voltage transients in ~1 uS (see electrical characterisation).

The acquisition board draws power from the Pyboard's USB connection which is also used to stream data to the computer. Capacitors on the 5 V rail improve transient response and smooth the load presented to the power supply, with a current limiting IC to restrict the inrush current when the board is powered up.

Assembly of the acquisition board requires only standard through-hole and surface-mount soldering techniques. To make a complete photometry system the acquisition board must be paired with LEDs, photoreceivers, filter cubes and other optical components. A complete parts list for the setup we use for green/red two colour experiments (e.g. GCaMP/TdTomato) is provided in Table 1. Assembled acquisition boards can be purchased from the Open Ephys store for €350 (http://www.open-ephys.org/store/pyphotometry).
**Data format.** pyPhotometry can save data either as binary data files with a .ppd file extension or as comma separated value files with a .csv file extension.

The binary file format is designed to be straightforward to import into analysis programs while generating files that are no larger than necessary. The user guide provides detailed information on importing .ppd files, and functions for importing .ppd files into Python and Matlab are provided in the tools folder of the code repository (https://bitbucket.org/takam/pyphotometry/src/default/tools/). Files consist of header information and data. The header is a JSON object that contains the subject ID, start date and time, and acquisition parameters. JSON is a lightweight data interchange format designed to be readable by humans and easy for machines to parse, and is supported by most programming languages. The remainder of the data file contains the analog and digital signals encoded as binary data. Each 2 byte chunk encodes one 15-bit analog signal sample and one digital signal sample, resulting in very compact data files – e.g. a 1 hour recording at 130 Hz sampling rate results in a ~1.9 MB file.

The comma separated value files generated by pyPhotometry are text files where each line contains one sample from each of the two analog inputs and two digital inputs, separated by commas. When data is stored as .csv file, acquisition settings are saved as a JSON object in a separate text file with a .json file extension. Saving data as .csv files results in files sizes approximately 4 times larger than binary files.

**Time-division illumination.** Photometry experiments often employ sinusoidal modulation of excitation light combined with lock-in amplification - i.e. multiplication of the photoreceiver signal by a sinusoidal reference signal synchronised with the excitation light, followed by low pass filtering. This has two advantages over continuous illumination: Firstly, the recovered signal is sensitive only to inputs at the modulation frequency, and thus insensitive to bleed through of ambient light and other noise sources well separated in frequency. As many noise sources have more power at low frequencies, lock in amplification may achieve better signal-to-noise than simply low pass filtering (which also reduces noise bandwidth but does so about 0 Hz). Secondly, sinusoidal

| Name | Quantity | Part | Vendor |
|------|----------|------|--------|
| Photoreciever | 2 | Newport 2151 with lensed FC adapter | Doric |
| 465 nm LED | 1 | CLED_465 | Doric |
| 560 nm LED | 1 | CLED_560 | Doric |
| Minicube | 1 | FMCS_E1(450–490)_F1(550–580)_E2(660–680)_S | Doric |
| Fiber patchcord: LED - Minicube | 2 | MFP_220/220/LWMJ-0.48_0.3 m_FCM-FCM | Doric |
| Fiber patchcord: Minicube - Photoreciever | 2 | MFP_600/630/LWMJ-0.48_0.3 m_FCM-FCM | Doric |
| Fiber patchcord: Rotary-sample | 1 | MFP_200/220/900-0.48_1 m_FCM-FCM | Doric |
| Pigtailed rotary joint | 1 | FRI_1 x 1 PT_200/230/LWMJ-0.48_1 m_FCM-0.2 m_FCM | Doric |
| FC-FC adapter | 1 | ADAPTER_FC style 2 | Doric |
| Optical breadboard 150 × 300 mm metric | 1 | MB1530F/M | Thorlabs |
| M6 - M3 thread adapter (for mounting acquisition board) | 4 | MSA6/M | Thorlabs |
| Clamp for minicube | 1 | CL3/M-P5 Compact Variable Height Clamp, M6 Tapped | Thorlabs |
| Clamp for photodetector | 2 | MSC2 Clamping Fork for 1/4" 20 (M6) Cap Screw | Thorlabs |
| Pillar for photodetector | 2 | TRP14/M Ø12 mm Pedestal Pillar Post | Thorlabs |
| m6 screw 12 mm (for mounting LEDs and photodetectors) | 4 | M6 x 12 mm Socket Cap Screw | Thorlabs |
| USB A - micro B cable 1.8 m | 1 | 121–3251 | RS |
| BNC cable 0.3 m: Photoreciever to acquisition board | 2 | 886–0706 | RS |
| M3 screw 10 mm (for mounting acquisition board) | 4 | 660–4636 | RS |
| M3 spacer 3 mm (for mounting acquisition board) | 4 | 161–3676 | RS |
| M6 × 45 mm (for minicube clamp) | 2 | 468–0133 | RS |

**Table 1.** Optical components for red/green system.
modulation at different frequencies can be applied to different wavelengths of excitation light, and the fluorescence evoked by each independently measured by lock in amplification at the appropriate frequency – a form of frequency-division multiplexing. This is typically used to independently measure the fluorescence evoked in GCaMP by excitation light at 470 nm and 405 nm.

pyPhotometry uses a different approach to achieve these ends, based on time-division rather than frequency-division principles. Independent readout of fluorescence evoked by different excitation light wavelengths is achieved by alternately switching on LEDs 1 and 2, and acquiring samples of signal 1 when LED 1 is on and of signal 2 when LED 2 is on (Fig. 3c). Additionally, baseline subtraction is used to render measurements insensitive to ambient light levels and other low frequency noise sources; for each sample the ADC is read twice; once with both LEDs off to obtain a baseline measurement, and again with the respective LED on. The baseline is subtracted from the sample such that only the difference in light intensity between the LED on and off conditions influences the signal. The acquisition sequence is:

1. Turn LEDs 1 and 2 off, read signal 1 baseline.
2. Turn LED 1 on, read signal 1 sample, subtract baseline and send sample to GUI.
3. Turn LEDs 1 and 2 off, read signal 2 baseline.
4. Turn LED 2 on, read signal 2 sample, subtract baseline and send sample to GUI.

Figure 2. Graphical user interface.
and returned to baseline 2.5 ms after the light was turned off. The transient response of the photoreceiver voltage faster timescale, ~1
is determined by the photoreceiver’s bandwidth, as both the LED currents and fluorescent lifetimes are on a much
emission spectra to be exploited to minimise crosstalk.

different excitation and emission spectra (e.g. GCaMP/tdTomato), allowing separation of both the excitation and
i.e. from two separate photoreceivers. This acquisition mode is designed to be used with two fluorophores with
Validation of time-division illumination.

This acquisition sequence is used for two acquisition modes, which differ with respect to the analog inputs
read to generate signals 1 and 2. The first mode, termed ‘1 colour time-division’, uses the same analog input (i.e.
a single photoreceiver) for signals 1 and 2. This mode is designed for independently recording the fluorescence evoked at a single emission wavelength by two different excitation wavelengths, for example for measuring the fluorescence from GCaMP due to illumination at 470 and 405 nm.

The second mode termed ‘2 colour time-division’ reads signals 1 and 2 from analog inputs 1 and 2 respectively,
we independently recording the fluorescence evoked at a single emission wavelength by two different excitation wavelengths, for example for measuring the fluorescence from GCaMP due to illumination at 470 and 405 nm.

Validation of time-division illumination. To determine the timing of the acquisition sequence for
time-division illumination we characterised the time course of photoreceiver voltage signals in response to excitation light transients (Fig. 3a,b). The photoreceiver response peaked 0.76 ms after the onset of the excitation light and returned to baseline 2.5 ms after the light was turned off. The transient response of the photoreceiver voltage is determined by the photoreceiver's bandwidth, as both the LED currents and fluorescent lifetimes are on a much faster timescale, ~1 μs for the LED current (Fig. 4c), and ~1 ns for the fluorescence lifetime of DiI and GCaMP.

Using a photoreceiver with a faster transient response would therefore allow time division illumination with a
faster sampling rate.

Based on these timings we implemented the acquisition sequence shown in Fig. 3c. Light pulses from each
LED were 0.75 ms in duration and occurred at 130 Hz, with the two LEDs in anti-phase. Acquisition from the
ADCs used 64x oversampling at a 256 KHz oversampling rate such that reading a sample took 0.25 ms. The baseline
for each signal was read immediately before the respective LED was turned on, and the signal sample was read immediately before the LED was turned off.

We assessed the linearity of measurements using this time-division acquisition sequence by measuring the acquired signal as a function of LED current for in-phase illumination of the fluorescent target - i.e. using signal 1 timings for acquisition and LED 1 timings for the light pulses (Fig. 3c). In-phase illumination produced a linear relationship between LED current and acquired signal (Fig. 3d). We assessed crosstalk between the two signals by
measuring the acquired signal as a function LED current for anti-phase illumination – i.e. using signal 1 timings for acquisition but LED 2 timings for the light pulses. The acquired signal was 0 independent of the intensity of out of phase illumination (Fig. 3c) indicating no detectable crosstalk. Finally we measured the acquired signal as a function of LED current for continuous illumination – i.e. using signal 1 timings for acquisition with the LED on continuously. Again, the acquired signal was 0 independent of LED current, indicating that baseline subtraction successfully removed sensitivity to steady state light intensity.

We assessed how the time-division illumination used by pyPhotometry compared with the more commonly used sinusoidal modulation in terms of (i) bandwidth (the frequency range of signals that can be measured) and (ii) signal to noise.

The bandwidth achievable using sinusoidal illumination with lock-in amplification is determined by the integration time needed for the modulations of different signals to be orthogonal. Over short integration times the modulations will not be orthogonal and readout will be contaminated by noise due to varying overlap between the target signal's modulation and the modulation of the other signal. We numerically evaluated the strength of this overlap noise as a function of window duration for two signals of equal amplitude modulated at 211 and 531 Hz (as used in 5). Overlap noise decreased smoothly with integration time (Fig. 3e) such that the required integration time depended on the noise level deemed acceptable. To achieve an overlap noise standard deviation <10⁻³ of the target signal size required a widow duration of ~10 ms.

Using time division illumination we can acquire samples from two channels at 130 Hz, corresponding to a time window of 7.7 ms per sample. The time division illumination used by pyPhotometry therefore achieves comparable signal bandwidth to that achieved by...
the frequency-division methods used in the literature. Both approaches achieve a bandwidth substantially larger than that of GCaMP6f.

Intrinsic noise due to e.g. thermal fluctuations in the photoreceivers, may differentially affect signals acquired using time-division illumination with baseline subtraction and sinusoidal illumination with lock-in amplification. We assessed this experimentally by measuring the coefficient of variation of signal recorded using both approaches from a fluorescent target (DiI solution) as a function of average illumination intensity (Fig. 3f). For time-division illumination the pyPhotometry acquisition board was used to acquire signals and control the LED. For sinusoidal illumination a National Instruments USB-6212 BNC board was used to acquire signals and generate the sinusoidal modulation signal, and a Doric Lenses LED driver was used to drive the LED. The optical setup was otherwise identical for the two approaches (see methods). Signals acquired using both methods were low pass filtered at 20 Hz to ensure their bandwidth was equivalent.

Noise was substantially lower for signals acquired using time-division illumination, such that a given signal to noise level was achieved at approximately 50% of the average LED current required with sinusoidal illumination. This likely reflects the fact that each LED is on for only 10% of the time during time-division acquisition, such that the illumination light intensity when the signal is measured is 10x higher than the average illumination intensity.

Electrical characterisation. We performed electrical measurement to characterise the accuracy of the LED driver outputs and the analog inputs.

Linearity of the LED driver was assessed by measuring the LED current as a function of the value written to the Pyboard DAC. LED current was linear in the DAC value across the full range of values (Fig. 4a). The standard deviation of LED currents across the 4 driver circuits tested was <1% of the mean current over a range of 1–100 mA mean current (Fig. 4b). The transient response of the LED driver was assessed by measuring the current waveform in response to a 1 ms control voltage pulse (Fig. 4c). Rise and fall times of the LED current were of order 1 μs.

To assess the accuracy of voltage measurement using the Pyboard ADCs we used the GUI to read both ADCs at 1 KHz while we presented constant voltage inputs. We tested 2 ADCs on each of 4 Pyboards for a total of 8 analog inputs. The measured voltage was very close to linear across the measured voltage range (Fig. 4d), with variation in measured voltage across inputs <1% of the mean across the range tested (Fig. 4e). We also assessed the extent to which individual inputs deviated from linearity by fitting a least squares linear fit to the measurements from each input separately and plotting the mean and standard deviation of the residuals across inputs (Fig. 4f). Deviation from linearity was <1 mV across the measured input voltage range and had a characteristic shape across inputs. We measured the noise amplitude of the measured signal. Noise amplitude increased with input voltage but was <0.06% of the measured voltage across the range of voltages tested (Fig. 4g).

These results confirm the acquisition board is able to control LED current with high accuracy and temporal precision, and acquire analog signals with good linearity and low noise.

Dopamine neuron recordings. Having characterised the performance of the hardware we tested its application to recording neuronal activity in vivo. Calcium transients were recorded from GCaMP6f expressing VTA dopamine neurons in response to unpredictable reward delivery, which is known to activate a large proportion of these neurons. For these recordings water restricted mice nose-poked in a reward port where 6 μL water rewards became available on a random interval 20 s schedule.

Signals were recorded from cell bodies in VTA using the ‘2 colour time-division’ acquisition mode to record a GCaMP signal and a movement control signal from co-expressed tdTomato (Fig. 5a,b). Similar to previous studies5,7,8, reward delivery consistently produced a marked, fast transient increase in GCaMP6f fluorescence, while the tdTomato signal was unaffected.

We also recorded signals from dopamine axons in nucleus accumbens (NAc) using the ‘1 colour time-division’ acquisition mode to record a calcium sensitive fluorescence signal evoked from GCaMP by 465 nm light and a calcium insensitive signal evoked by 405 nm isosbestic illumination (Fig. 5c,d). Unpredictable reward delivery again produced sharp calcium transients in the calcium sensitive signal while the isosbestic signal was unaffected.

These recordings demonstrate the ability of the system to acquire measures of bulk activity from cell bodies and axons in deep-brain structures in behaving animals.

Discussion We have developed an open source acquisition board and GUI for fiber photometry data acquisition. The system is compact, convenient, and cheap, and we have characterised its performance in detail.

The system can use time-division illumination to independently measure fluorescence evoked by different excitation wavelengths, combined with baseline subtraction to render measurements insensitive to ambient light. We compared the bandwidth and signal to noise performance of this approach to the more commonly used sinusoidal modulation and lock-in amplification. We found that time-division illumination offered comparable bandwidth to sinusoidal modulation schemes used in the literature, but better signal to noise as a function of illumination light intensity, allowing approximately 50% lower light intensity for a given noise level. Reducing illumination intensity is desirable as it reduces photo-bleaching of fluorophores.

Components of the system lend themselves to adaptation for other applications. The LED driver circuit may be useful in applications where a linear voltage controlled current source with fast transient response is required. The circuit can be modified to handle higher currents by using a MOSFET with higher power dissipation and changing some resistor values. For instance, we use a modified version of the circuit as an analog LED driver for optogenetics.
The GUI code may be useful as a template for applications which require a Python based GUI for data acquisition and plotting. Some pyPhotometry GUI code is shared by pyControl (https://pycontrol.readthedocs.io), a system of open source hardware and software for controlling behavioural neuroscience experiments, also built around the Micropython microcontroller.

While writing this manuscript we became aware of another open source project for fiber photometry called PhotometryBox34. This supports generation of sinusoidally modulated control signals (sent to external LED drivers), online demodulation of signals for visualisation (though for analysis demodulation is performed offline), and recording of signals to disk. PhotometryBox uses a microcontroller for generating sinusoidal signals and online demodulation, and a National Instruments board for recording data. The principal differences between the systems are: pyPhotometry has built in LED drivers and is USB bus powered; PhotometryBox must record at higher sampling rates (5 KHz) as demodulation is performed offline, resulting in larger data files. pyPhotometry is controlled via a dedicated GUI custom-written for photometry while PhotometryBox uses generic electrophysiology acquisition software (WinEDR) combined with physical controls. Ultimately both systems provide open source photometry data acquisition at a fraction of the cost of commercial systems. We expect this proliferation of open source tools to both reduce research costs and facilitate the development of novel applications for photometry.

Methods

Open source repositories. Design files for the system are at https://bitbucket.org/takam/pyphotometry and https://bitbucket.org/takam/pyphotometry_hardware. User documentation is at https://pyphotometry.readthedocs.io.

Electrical calibration experiments. A Picoscope 2204A USB oscilloscope was used for all electrical measurements and as a signal generator for testing the analog inputs.

LED current was measured via the voltage across the 4.7 ohm current sense resistors that form part of the driver circuit, using the provided connection points on the board. LED current was measured at 8 DAC values spaced from 500 to 4000 (the DAC takes values from 0 to 4095 in 12 bit mode), and a further 10 values spaced from 10 to 100 to cover the low current range. A single linear fit was made to the full set of points.

Figure 5. VTA dopamine neuron recordings. (a) Fluorescent signals acquired in VTA from GCaMP6f (green) and tdTomato (red) expressed in dopamine neurons, recorded using the ‘2 colour time division’ acquisition mode. Black triangles indicate times of unpredictable reward delivery. (b) VTA response aligned on reward delivery, line shows mean and shaded area shows standard error. (c) Signals acquired from GCaMP6f expressing dopamine neuron terminals in nucleus accumbens using the ‘1 colour time division’ acquisition mode with 465 nm illumination (green) and 405 nm isosbestic illumination (purple). (d) NAc terminal response aligned on reward delivery.
**Time-division illumination.** Experiments characterising the time-division illumination mode and comparing it with sinusoidally modulated illumination used a Doric Lenses CLED 560 nM LED for illumination, a Doric Lenses FCMS5 minicube to separate excitation and emission light (excitation filter 555–570 nM, emission filter 580–680 nM), and a Newport 2151 photoreceiver in DC coupled mode to detect the emitted light. The fluorescent target was a solution of Dil in ethanol, shielded from ambient light and coupled to the minicube via 200 um core 0.48 NA optical fiber.

To evaluate signal to noise for time-division illumination (Fig. 3f) we used the pyPhotometry acquisition board and GUI to both control the LED and acquire signal. For sinusoidal illumination the LED was controlled using a Doric Lenses LEDD_2 driver in low power analog modulation mode. The sinusoidal voltage signal used to modulate the LED was generated by a National Instruments USB-6212-BNC board controlled by WinEDR software (John Dempster, University of Strathclyde). The modulation amplitude was equal to the average value, i.e. the modulation depth was 100%. The USB-6212 was used to record both the LED modulation signal and the photoreceiver signal at 10 KHz. Lock in amplification was performed offline using the following steps: (1) Bandpass filtering the recorded modulation and photoreceiver signals around the modulation frequency. (2) Applying a time lag to the modulation signal to phase align it with the photoreceiver signal. (3) Multiplying the lagged modulation with the photoreceiver signal. (4) Low pass filtering the product at 20 Hz.

All filtering steps used 4th order zero phase filters, implemented by filtering in the forward and reverse directions using a 2nd order Butterworth filter. The same 20 Hz low pass filtering was applied to signals acquired using time-division illumination to ensure noise comparison was made on signals of equivalent bandwidth.

Figure 3e used numerical simulation to assess the bandwidth achievable with the sinusoidally modulated illumination and lock-in amplification used in1. Consider using lock-in amplification to measure the amplitude of sinusoidal signal $a(t)$ in the presence of sinusoidal signal $b(t)$ at a different frequency. Lock-in amplification works by multiplying the input with a sinusoidal reference signal synchronised to the target signal, followed by low pass filtering. The lock-in output at time $T$ is:

$$D(T) = \int_{-\infty}^{\infty} F(t-T)r(t)(a(t) + b(t))dt$$

where $r(t)$ is the sinusoidal reference signal and $F(t-T)$ is the impulse response of the low pass filter.

The quantity we plot in Fig. 3e which we term normalised overlap noise is:

$$\frac{\text{standard deviation} \int_{-\infty}^{T} F(t-T)r(t)b(t)dt}{\text{expectation} \int_{-\infty}^{\infty} F(t-T)r(t)a(t)dt}$$

where the expectation and standard deviation are over measurement time-point $T$. The numerator is the standard deviation of noise on the lock-in output due to overlap between the sinusoidal input $b(t)$ and the reference signal $r(t)$. The denominator is the expected contribution of the target signal $a(t)$ to the lock-in output.

We evaluated this quantity numerically for sinusoids of equal amplitude at 211 and 531 Hz. A Dolph-Chebyshev window with 100 dB side lobe attenuation was used as the filter impulse response function $F(t-T)$. We report the overlap noise as a function of the width of this integration window. This window function was chosen because it performed best out of all those tested.

**Dopamine recordings.** Mice were anaesthetised with isoflurane (3% induction, 0.5–1% maintenance), treated with buprenorphine (0.1 mg/kg) and meloxicam (5 mg/kg), and placed in a stereotactic frame. The skull was exposed and holes drilled to allow virus injection and implantation of fiber optic cannula. Mice were given addi-

For VTA cell body recordings (Figs. 5a,b), GCaMP and tdTomato were expressed in VTA dopamine neurons using AAV1-Syn-Flex-GCaMP6f-WPRE-SV40 (titer 6.2 × 10^{13}) and AAV1-CAG-Flex-tdTomato-WPRE-bGH (titer 3.1 × 10^{13}) viruses (Penn Vector Core) in male B6.SJL-Slc6a3tm1.1(cre)Bknm/f mice. The viruses were mixed and diluted in a ratio of 20% GCaMP6f, 10% tdTomato, 70% saline. 500 nL per hemisphere of the diluted virus was injected at 1 nL/second at AP: −3.3, ML: ±0.4, DV: −4.5 mm relative to bregma. Recordings were made through a 200 um 0.53 NA fiber optic cannula implanted at AP: −3.3, ML: ±0.4, DV: −4.3 mm relative to bregma.

For the NAc terminal recordings (Figs. 5c,d) GCaMP was expressed in VTA dopamine neurons using AAV1-Syn-Flex-GCaMP6f-WPRE-SV40 (titer 6.2 × 10^{13}). The virus was diluted 1:10 with saline and 1 μL was injected per hemisphere at AP: −3.3, ML: ±0.4, DV: −4.2 mm relative to bregma. Optic fiber cannula were implanted at AP: 1.4, ML 0.8, DV: 4.1 relative to bregma.

Prior to recording, mice were put on a water restriction schedule where on training days they received 0.5–1.5 mL water from rewards received in the task and on non-training days 1 hour of unrestricted access in their home cage. Mice maintained a typical body weight of >90% pre-restriction levels. Experiments were carried out in accordance with the Oxford University animal use guidelines and performed under UK Home Office Project Licence P6F11BC25.

VTA cell body recordings were acquired using the ‘2 colour time division’ acquisition mode using 470 and 560 nm wavelength LEDs respectively for the GCaMP and tdTomato excitation light and 500–540 and 600–680 nm emission filters for the GCaMP and tdTomato signals. NAc terminal recordings were acquired using the ‘1 colour time division’ acquisition mode using 470 and 405 nm wavelength LEDs respectively for the GCaMP and isosbestic signals.
Acquired signals were bandpass filtered between 0.01 and 20 Hz using a fourth order zero phase filter. The full set of optical components used is listed Table 1 as well as in the hardware repository. The Newport photo receivers were used in DC coupled mode for all recordings.

Data Availability
All data and analysis code used in the study are in the file ‘manuscript code and data.zip’ in the downloads section of the code repository: https://bitbucket.org/takam/pyphotometry/downloads/.

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Author Contributions
T.A. designed and built the hardware and software, designed experiments, analysed data and wrote the manuscript. M.E.W. designed experiments and edited the manuscript.

Additional Information
Competing Interests: T.A. has a consulting contract with Open Ephys Production Site who sell assembled pyPhotometry acquisition boards. M.E.W. has no competing interests.

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