Elucidating regulators of astrocytic Ca\(^{2+}\) signaling via multi-threshold event detection (MTED)

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Abstract
Recent achievements in indicator optimization and imaging techniques promote the advancement of functional imaging to decipher complex signaling processes in living cells, such as Ca\(^{2+}\) activity patterns. Astrocytes are important regulators of the brain network and well known for their highly complex morphology and spontaneous Ca\(^{2+}\) activity. However, the astrocyte community is lacking standardized methods to analyze and interpret Ca\(^{2+}\) activity recordings, hindering global comparisons. Here, we present a biophysically-based analytical concept for deciphering the complex spatio-temporal changes of Ca\(^{2+}\) biosensor fluorescence for understanding the underlying signaling mechanisms. We developed a pixel-based multi-threshold event detection (MTED) analysis of multidimensional data, which accounts for signal strength as an additional signaling dimension and provides the experimenter with a comprehensive toolbox for a differentiated and in-depth characterization of fluorescence signals. MTED was validated by analyzing astrocytic Ca\(^{2+}\) activity across Ca\(^{2+}\) indicators, imaging setups, and model systems from primary cell culture to awake, head-fixed mice. We identified extended Ca\(^{2+}\) activity at 25°C compared to 37°C physiological body temperature and dissected how neuronal activity shapes long-lasting astrocytic Ca\(^{2+}\) activity. Our MTED strategy, as a parameter-free approach, is easily transferrable to other fluorescent indicators and biosensors and embraces the additional dimensionality of signaling activity strength. It will also advance the definition of standardized procedures and parameters to improve comparability of research data and reports.

KEYWORDS
astrocyte, biosensor, Ca\(^{2+}\), GCaMP, MTED

JEL CLASSIFICATION
C00; I29
1 | INTRODUCTION

Fluorescence microscopy is a method used across many scientific disciplines. In biology, it allows researchers to visualize cellular processes in time and space largely preserving the integrity of the sample. Recent developments in novel microscopy techniques, biosensors, and indicators push functional imaging to its limits. A persistent major challenge is to adequately analyze multidimensional data. For example, Ca$^{2+}$ is an important signaling molecule and, because of its large concentration gradients, a well-established readout for cellular activity. In some cell types, such as astrocytes, Ca$^{2+}$ activity patterns can be very complex. Astrocytes are versatile glial cells participating in the maintenance of numerous brain functions, including the regulation of ion homeostasis, neurotransmitter clearance, energy supply, and blood flow (Sofroniew & Vinters, 2010). They are a crucial element in the brain architecture and display very distinct morphological characteristics that vary between brain regions and sub-regional networks (Chai et al., 2017). With their fine processes, astrocytes contact synapses. As active partners at the tripartite synapse, they release gliotransmitters that regulate synaptic activity (Araque et al., 2014; Bonansco et al., 2011; Henneberger et al., 2010; Panatier et al., 2011). In return, the release of a variety of neurotransmitters has been shown to modulate the cytosolic Ca$^{2+}$ concentration ([Ca$^{2+}$] i) in astrocytes (Porter & Mccarthy, 1997). Variations in astrocyte [Ca$^{2+}$] have been proposed to represent a unique manner of cellular signaling, but revealing the exact functions and regulatory mechanisms remains a challenge (Semyanov et al., 2020; Volterra & Meldolesi, 2005). Ca$^{2+}$ signals in astrocytes are not solely stimulus-dependent, but also occur spontaneously (Nett et al., 2002). They can also propagate and fuse, thus displaying very distinct morphological characteristics that vary between cell types.

Ca$^{2+}$ activity in astrocytes appears as local and very distinct fluctuations but could also spread into distant regions or engage the whole cell. Therefore, the spatial extent and amplitude of Ca$^{2+}$ events are additional measures and standard evaluation approaches used in other applications cannot simply be transferred to astrocytic Ca$^{2+}$ activity (Volterra et al., 2014).

A standard approach to analyzing Ca$^{2+}$ activity is fluorescence microscopy using Ca$^{2+}$ sensitive fluorescent dyes (Meldolesi, 2004). They generally come in two flavors: organic dyes, such as Oregon Green 488 BAPTA-1 (OGB-1), and genetically encoded Ca$^{2+}$ indicators (GECIs), such as GCaMPs, which both have their specific advantages and disadvantages (Chen et al., 2013; Semyanov et al., 2020). Both are easy to use in single channel recordings, which simplifies the readout and subsequent data processing (Horikawa, 2015; Pérez Koldenкова & Nagai, 2013). Various tools are available to extract relative changes in [Ca$^{2+}$] i, which mainly analyze the frequency and amplitude of events (Venugopal et al., 2019) based on predefined or automatically detected active regions (Agarwal et al., 2017; Castro et al., 2011). Recently, alternative approaches have been presented that also analyze the directionality of propagating events and the origin of Ca$^{2+}$ events, which is predominantly located in the fine peripheral processes (Wang et al., 2019; Wu et al., 2014, 2019). This has shifted the focus from major Ca$^{2+}$ changes in the somatic region to the periphery. However, these regions provide only a weak fluorescent signal due to their low volume. Therefore, when the detection method is based on identifying signals above noise level, events in these areas can escape detection due to a low signal-to-noise ratio.

We developed a strategy to determine the fluorescence signal of Ca$^{2+}$ indicators at a basal $[\text{Ca}^{2+}]_{o}$, which was the basis for calculating $(F-F_0)/F_0$, expressed as $\Delta F/F_0$, as a measure of relative changes in $[\text{Ca}^{2+}]_{i}$ in a pixel-based manner. We then applied an automated multi-threshold event detection (MTED) algorithm to quantify the intrinsic Ca$^{2+}$ activity of astrocytes in primary hippocampal cultures, organotypic slice cultures, and cortical astrocytes in vivo. This allows to combine the information of amplitude and spatial extend over time to investigate the magnitude of Ca$^{2+}$ events as a measure of signaling strength. Next, we tested MTED in two straightforward experimental settings. We could demonstrate substantially different Ca$^{2+}$ activity dynamics of cultured astrocytes at room temperature (RT) compared to 37°C, which resembled the activity measured in vivo. We could further reveal that neuronal activity favors the generation of long-lasting $[\text{Ca}^{2+}]_{i}$ elevations in astrocytes. Our flexible MTED algorithm, exploited here successfully for Ca$^{2+}$ indicators and astrocyte physiology across preparations in 2D over time, is easily applicable to investigate functional dynamics using various fluorescence-based indicators and biosensors and can be easily extended to 3D over time and to other cell types.

2 | MATERIAL AND METHODS

2.1 | Animals for in vitro Ca$^{2+}$ imaging

For all in vitro experiments, wildtype animals of both genders from strain C57BL/6J were used. Animals were housed and cared for in accordance to directive 2010/63/EU. Mice were kept in a 14 h light and 10 h dark cycle with lights on starting at 7 am. Animals had ad libitum access to food and water and were kept under standard conditions at 22 ± 2°C RT with 55% ± 5% humidity. Mice were killed by decapitation, and all experiments were conducted according to the recommendations of the European commission and Lower Saxony’s “Landesamt für Verbraucherschutz und Lebensmittelsicherheit” (authorization number 2018/179).

2.2 | Animals for in vivo Ca$^{2+}$ imaging

Mice were maintained in the animal facility of the Center for Integrative Physiology and Molecular Medicine (IIPMM, University of Saarland). Astrocyte-specific knock-in GLAST-CreERT2 mice (Slc1a3tm1 [cre/ERT2]Mgoe, MGI:3830051) (Mori et al., 2006) were crossbred to Rosa26 reporter mice with GCaMP3 expression (Pt(ROSA)26Sortm1 [Cre]/ERT2)Mgoe, MGI: 5659933) (Paukert et al., 2014). Imaging sessions were performed at 8–10 weeks of age. Mouse administration was managed via the PyRAT database (Python based Relational Animal Tracking) from Scionics Computer Innovation GmbH (Dresden, Germany). Animals were kept and bred in strict accordance with the...
recommendations to European and German guidelines for the welfare of experimental animals. Animal experiments were approved by the Saarland state’s “Landesamt für Gesundheit und Verbraucherschutz” in Saarbrücken/Germany (license numbers: 71/2013 and 36/2016).

2.3 | Primary hippocampal astrocyte cultures

Primary astrocyte cell cultures were prepared according to a previously described protocol (Wu et al., 2014) with slight modifications: Hippocampi were isolated from brains of neonatal mice between P1-3 and cells were seeded after dissociation at a density of 5 x 10^4 cells per 12 mm glass coverslip for microscopy in 500 μL plating medium (49 mL MEM, 1 mL B-27 supplement, 500 μL sodium pyruvate, 500 μL L-Glutamine, 50 μL Penicillin–Streptomycin; all Thermo Fisher Scientific Inc., Waltham). On DIV3 the entire plating medium was replaced with 1 mL maintenance medium (49 mL Neurobasal-A, 1 mL B-27 supplement, 500 μL L-Glutamine, 50 μL Penicillin–Streptomycin; all Thermo Fisher Scientific Inc., Waltham). On DIV11, ½ of the medium was exchanged with prewarmed maintenance medium prior to infection of the cells with of 0.1 μL AAV-mGFAP-GCaMP6s (3.7 x 10^8 vg/μL) and AAV-mGFAP-tdTomato (1 x 10^7 vg/μL). Astrocytes were maintained at 37°C in a humidified incubator in a 5% CO2 atmosphere used for experiments between DIV14-17. Cells were transferred to a prewarmed recording chamber for microscopy and kept in a balanced salt solution (BSS), which was adjusted to pH 7.4 and 290 mOsm with glucose, containing 115 mM NaCl, 5.4 mM KCl, 1 mM MgCl2, 2 mM CaCl2 and 20 mM HEPES.

2.4 | Organotypic slice cultures

Organotypic slice cultures were prepared after an adapted protocol from Kobe et al. (Kobe et al., 2012). Briefly, mice were decapitated at P6 under sterile conditions and the isolated hippocampus was placed in ice-cold oxygenized slice medium in a 60 mm dish for 30 min. 350 μm thick slices were prepared with McIlwain Tissue Chopper (Mickle, Surrey, UK) and separated with a needle to select 2-4 slices with complete hippocampal structures. Selected slices were transferred onto Millicell filter inserts (#PICM03050, Merck, Darmstadt, Germany) at pH 7.3). Excess liquid around the slice was removed and cells were subsequently infected by application of 0.2 μL AAV-mGFAP-GCaMP6s (3.7 x 10^8 vg/μL) into the medium. Slices were kept in a humidified atmosphere (5% CO2, 37°C) with ½ of the medium being exchanged on DIV2, DIV4 and DIV6. Ca2+ imaging was conducted at DIV5-7.

2.5 | Cranial window surgery for in vivo two-photon imaging

During surgical procedures, animals were kept on a heating pad and eyes were covered with Bepanthen ointment (Bayer, Leverkusen, Germany). Anesthesia was induced with a mixture of 5% isoflurane, 47.5% O2 (0.6 L/min) and 47.5% N2O (0.4 L/min) and maintained with 2% isoflurane (Harvard Apparatus anesthetic vaporizer, Harvard, Holliston). A standard craniotomy (Cupido et al., 2014) of 3 mm in diameter was performed over the somatosensory cortex (2 mm posterior and 1.5 mm lateral to bregma). The craniotomy was sealed with a glass coverslip (Glawarenfabrik Karl Hecht, Sondheim, Germany; #1.5 thickness code) and fixed with dental cement (ReliX®, 3 M ESPE, Seefeld, Germany). Subsequently, a metal holder for head restraining (5 mm diameter) was applied and fixed to the skull with dental cement. After surgery, the animals were kept on the heating pad until complete recovery. Post-operative treatment consisted of buprenorphine (3 mg/kg, s.c.) and dexamethasone (0.2 mg/kg, i.p.), for three consecutive days. Recovery was assessed by body weight and mouse grimace scale. After five to seven days the first imaging session was performed.

In preparation for Ca2+ imaging, animals were habituated before the first imaging session according to adapted protocols without water restriction from Guo et al. (Guo et al., 2014) and Kislin et al. (Kislin et al., 2014) The animals were head-fixed with a custom-designed head-restrainer, 3D-printed using stainless steel. During imaging, anesthesia was applied using a custom-made, magnetically attachable anesthesia mask. Each field of view (FOV) was imaged twice: first in anesthetized, then in awake state. During imaging in anesthetized state, isoflurane concentration was kept at 1.5%, and flow of O2 and N2O was set to 0.6 and 0.4 L/min, respectively. Before awake state imaging, isoflurane and other gases were switched off and it was verified that the animals were fully awake. The selected FOVs for Ca2+ imaging were located in the somatosensory cortex, 80-100 μm beneath the dura. Each FOV was recorded for 5 min to investigate the Ca2+ signals. The total duration of one imaging session ranged between 30 and 60 min per animal. After imaging, animals were kept on a heating pad at 37°C until they recovered completely, additionally Fresubin ( Fresenius Kabi, Bad Homburg, Germany) was provided ad libitum.

2.6 | Reagents

Tetrodotoxin citrate (TTX; #Asc-055; Ascent Scientific, Princeton, NJ) was used at a concentration of 10 nM to block neuronal activity and was applied several minutes prior to imaging. The Ca2+-ATPase inhibitor Cyclopiazonic acid (CPA; #120300, Abcam, Cambridge, UK) was applied at a concentration of 10 μM at least 10 min before the measurements.

2.7 | Microscopy

Ca2+ imaging in vitro and in situ was conducted on an upright Andor Spinning Disk microscope (Oxford Instruments, Belfast, Northern Ireland) equipped with a CSU-X1 (Yokogawa, Musashino, Japan) using filter cubes (537/26 nm) for full frame imaging of GCaMP6s or split filter cubes (609/54) for simultaneous imaging of GCaMP6s and tdTomato. Cells were recorded for 10 min with 5 frames/s using
excitation wavelength 488 nm (GCaMP6s) and 561 nm (tdTomato). The temperature of the BSS for measurement was controlled by a custom-built heating device and additionally supervised with an external thermometer. To achieve thermal stability and avoid artifacts during recordings the objective, stage and chamber were all heated to the desired temperature.

In vivo Ca\(^{2+}\) imaging was conducted in anaesthetized (1.5% iso-flurane) and awake head-fixed mice through a cortical cranial window in the prefrontal cortex using two-photon excitation laser scanning microscopy (TPE-LSM). The custom-built microscope was equipped with a resonant scanner (RESSCAN-MOM, Sutter instrument, Novato) and a 20x water-immersion objective (W Plan-Apochromat 20x/1.0 DIC D = 0.17; Zeiss, Jena, Germany). Images were acquired with a frame rate of 30 Hz and a 10 Hz frame-averaging factor, resulting in an effective acquisition rate of 3 Hz. To minimize photo-damage, the excitation laser power was kept at a minimum for a sufficient signal-to-noise ratio (<40 mW at 60 ns pixel dwell time). Laser wavelength was set to 890 nm (Chameleon Ultra II, Ti:Sapphire Laser; Coherent, Santa Clara). The emitted light was detected by a photomultiplier tube (R6357; Hamamatsu, Hamamatsu, Japan) and pre-amplified (DHPCA-100, Femto, Berlin, Germany). Digitizer (NI-5734) and control hardware (NI-6341) was housed in a PXIe (1082) chassis, connected to a control-PC via a high bandwidth PXIe-PCIe8398 interface (NI, Austin). Scanning and image acquisition were controlled by ScanImage (SI 5.6R1) (Pologruto et al., 2003).

2.8 | Data processing

Data were processed using Matlab. In vitro data obtained by confocal spinning disk microscopy were denoised using VBM3D (Maggioni et al., 2012) and low intensity TPE-LSM data by SURE-LET. An automated data offset control was obtained by analyzing the data intensity histogram. The $F_0$ calculation is based on ‘moving window’ filter functions, where the mean and variance of $F$ are used as weighting functions for $F$ to identify low intensities. The ‘moving window’ sizes for filtering are input parameters, dependent on acquisition settings and the temporal profile of expected signal changes. The MTED algorithm uses $F_0$ or $F_R$, the fluorescence signal of an equally expressed reference fluorescent protein (FP), and applied various Ca\(^{2+}\) thresholds to $\Delta F/F_0$ or $\Delta F/F_R$, respectively. To reject false-positive pixels, we applied an appropriate $F_0$ threshold to eliminate background signals, such as readout noise. Next, basic morphological gray-scale operations (opening and closing) were performed to close holes and remove small isolated peaks, followed by Gaussian smoothing of the Ca\(^{2+}\) signals. This was used as a weighting function. Groups of Ca\(^{2+}\)-positive regions were then identified based on their spatio-temporal connectivity. Small groups not exhibiting a minimum size and duration were rejected. The time dependence of a detected event was stored for visualization and further analysis. The Ca\(^{2+}\) event detection was repeated for various Ca\(^{2+}\) thresholds. For practical reasons, we use logarithmic-like spaced threshold levels, such as [0.2, 0.5, 1, 2, 5, 10] fold change in $F$.

2.9 | Statistical analysis

Data were analyzed for statistical significance using GraphPad Prism 8. The individual statistical tests are indicated in the respective figure legends.

2.10 | Data and materials availability

The Matlab code developed in this study is available from the corresponding author upon request.

3 | RESULTS

3.1 | Visualization of Ca\(^{2+}\) transients in astrocytes

For developing a strategy to accurately characterize Ca\(^{2+}\) activity, we used primary cultures of mouse hippocampal astrocytes expressing the Ca\(^{2+}\) indicator GCaMP6s. These cells exhibit extensive endogenous Ca\(^{2+}\) activity with varying amplitudes and magnitudes, changing directionality and regional patterns (Figure 1, Movie S1). The GCaMP6s fluorescence signal $F$ is sensitive to changes in Ca\(^{2+}\) levels, but also depends on the indicator concentration itself and the cellular structure, which determines the number of molecules in the focal volume. Figure 1 shows an example of Ca\(^{2+}\) changes in cultured hippocampal astrocytes. From the GCaMP6s fluorescence signal $F$ (Figure 1(a) and Supplementary Figure S1(a)), which scales with both the Ca\(^{2+}\) and GCaMP concentration, it is not possible to deduce the changes in [Ca\(^{2+}\)] directly. The time series shown in Figure 1(b) contains various regions presenting similar brightness at some points over time (regions of interest [ROIs] 1–4). While the intensity of ROI 4 stays constant over time, $F$ in ROIs 1–3 varies and can be identified as changes in Ca\(^{2+}\) levels. To obtain a quantity, which is independent from the indicator concentration and the number of indicator molecules in the focal volume, we developed an automated pixel-based calculation of $F_0$ derived from the indicator signal $F$ (Supplementary Figure S2 and Movie S2). $F_0$ deduced from the temporal behavior of $F$, represents the indicator signal at lowest [Ca\(^{2+}\)]. The resulting $\Delta F/F_0$ provides an indicator concentration-independent readout which reflects relative changes in [Ca\(^{2+}\)] (Figure 1(c),(e), Movie S1). Alternatively, cytosolic co-expression of a Ca\(^{2+}\)-insensitive FP (e.g., tdTomato) as a reference probe ($F_R$) can be used to account for pixel and time dependent differences in the number of indicator molecules in the focal volume (compare Supplementary Figure S1 and Movie S1). A common strategy of detecting Ca\(^{2+}\) events from $F$ is based on identifying signals above noise level (often a factor times the standard deviation of $F$). This can be a source of biases because only very pronounced changes in Ca\(^{2+}\) exceed the noise level at small, peripheral structures, where $F$ is small as well (see Supplementary Figure S2). We can further visualize the spatiotemporal increases and decreases in cytosolic Ca\(^{2+}\) ($d(\Delta F/F_0)/dt$), depicted as red and blue regions, respectively, in Figure 1(d) and Movie S1(d).
Comparison of the overview images in Figure 1(a),(f), where the maximum projection of a 15 s time window is shown as intensity $F$ and as false color $\Delta F/F_0$, respectively, underlines the benefit of the ratiometric concept. Bright regions, like some cell somata in Figure 1(a), are characterized by low changes in Ca\textsuperscript{2+}, whereas others exhibit high Ca\textsuperscript{2+} activity.

### 3.2 Multi-threshold event detection

The analysis of Ca\textsuperscript{2+} activity in astrocytes should provide detailed information on event duration, size, and magnitude (see Figure 1, Movie S1). A fundamental question is: which changes in Ca\textsuperscript{2+} can be called a Ca\textsuperscript{2+} event? Furthermore, differences in the characteristics of Ca\textsuperscript{2+} events need to be identified and should be reported by an event detection algorithm. Such an algorithm should also capture the spatial extent of the events without prior definition of static ROIs. Furthermore, the detection algorithm should not require experiment-specific settings. Therefore, we developed MTED, a pixel-based, dynamic event detection algorithm relying on multiple thresholds, which computes the abovementioned variables (Figure 2(a),(b), Movie S3). Condition-specific adjustments of parameters are not required.

Through Monte Carlo simulations, we found that the accuracy in identifying a pixel above a specific Ca\textsuperscript{2+} threshold can be far below 95% confidence and varies with $F_0$ (see Supplementary Figure S2). We selected a set of six thresholds (0.2, 0.5, 1, 2, 5, 10) expressed in terms of $\Delta F/F_0$, where a Ca\textsuperscript{2+} event reaching a threshold of 0.2 or 1 does not necessarily reflect a 20% or double increase in basal Ca\textsuperscript{2+} concentration, respectively (see Semyanov et al., 2020). In conjunction with proximity relationships of neighboring pixels (xyt), MTED identifies groups of Ca\textsuperscript{2+}-positive pixels for a set of Ca\textsuperscript{2+} thresholds, thereby allowing for dynamic region changes. The output for the sequence shown in Figure 1(c) is shown in Figure 2(c) and Movie S3. Here, the six Ca\textsuperscript{2+} thresholds applied are depicted as contour lines with different colors, which show the respective spatial signal dynamics. A statistical analysis of the complete dataset is shown in Figure 2(d) as frequency plots for all analyzed Ca\textsuperscript{2+} thresholds exemplifying the maximum size of the events, their duration, and the maximum distance of the event center traveled over time (see Supplementary Figure S3 for more statistical parameters). The frequency plots shown in Figure 2(d) can be accumulated for different experimental conditions and be compared as shown below. Overall, the MTED algorithm provides a comprehensive analysis of spatiotemporally complex fluorescence changes.

### 3.3 The temperature-dependence of Ca\textsuperscript{2+} event characteristics

We next applied the analysis to two physiologically meaningful experimental scenarios. Investigations with cultured astrocytes are often carried out at RT (i.e., 22°C) rather than at the physiological body temperature (i.e., 37°C). When investigating Ca\textsuperscript{2+} event characteristics of the same astrocytic culture at different environmental temperatures, we observed substantial differences in the Ca\textsuperscript{2+} activity.
patterns (Figure 3, Movie S4). Figure 3(a) illustrates the Ca\(^{2+}\) events detected within the zoomed-in region shown in Figure 1(b)–(e) for 25, 34, and 37°C. At 25°C, both events (ROI 1 and 2 from Figure 1) slowly increased, reaching large areas and high amplitudes. At 34°C, events in nearby regions were much shorter and occupied much smaller areas. At 37°C, the detected events that originated from the same initial region were spatially restricted and shorter. Such temperature-dependent changes become more obvious when illustrated in 3D (Figure 3(b), Movie S5). Statistical analysis of the maximum event size, event duration, and maximum distance the area center traveled within the event revealed that at 25°C strong Ca\(^{2+}\) changes up to 10-fold were detected, spanning over the complete parameter range for the
maximal size and distance of Ca\(^{2+}\) events (Figure 3(c), Supplementary Figure S3, Movie S5). At 34°C, high-magnitude Ca\(^{2+}\) events were still detected but overall smaller magnitudes became more frequent. At 37°C, high Ca\(^{2+}\) levels were not reached any longer and small-magnitude Ca\(^{2+}\) events dominated. Event duration and maximum propagation distance also display a clear temperature dependence such that Ca\(^{2+}\) events become shorter and propagated less at higher temperature.

**FIGURE 3**  Astrocytic Ca\(^{2+}\) event characteristics are shaped by ambient temperature. (a) Comparison of Ca\(^{2+}\) activity detected in the same cell at 37°C, 34°C, and 25°C for a sequence of 14 s color-coded for various thresholds. Data were captured in a time sequence of 10 min at 25°C, the sample heated to 34°C, and a sequence acquired from the same region. The measurement was repeated at 37°C and the sample cooled back to 34°C and then to 25°C (see Movie S4). Scale bar = 20 μm. (b) 3D contour plots of detected events, with darker colors representing higher changes in Ca\(^{2+}\), illustrating the change in Ca\(^{2+}\) activity with environmental temperatures. (c) 2D histograms as a statistical presentation of selected parameters. At 37°C, the detected events are very small in space, transient in time, and do not reach high magnitudes. These characteristics change with cooler environments, reaching longer lasting, spatially more extensive, and comparably high magnitude events during measurement at 25°C.
3.4 | Suppression of cytosolic Ca\(^{2+}\) clearance reproduces the low temperature pattern

To understand the mechanisms underlying the observed temperature-dependent differences in the Ca\(^{2+}\) patterns, we compared the maximum increase and decrease in the Ca\(^{2+}\) signal, \(\max_t (dF/F_0)/dt\) and \(\min_t (dF/F_0)/dt\), respectively, for each detected Ca\(^{2+}\) event (Figure 4(a)). At 37°C, the signal increases were very local with low amplitude. In contrast, regions with positive changes in Ca\(^{2+}\) were spread over wider areas and reached greater amplitudes at lower temperatures. Similarly, the decrease in Ca\(^{2+}\) overlaying centers of maximal increase was punctual at 37°C but spread over wider regions at lower temperatures. Overall, negative Ca\(^{2+}\) changes were substantially greater at 34°C than 25°C. The rate of positive and negative Ca\(^{2+}\) changes was similar at 37°C, the positive Ca\(^{2+}\) changes (i.e., spreading of events) overcame the negative changes (i.e., depletion of events) at 25°C (Figure 4(b)).

Next, we used this analysis to investigate what underlies the profoundly different Ca\(^{2+}\) characteristics. The decrease in Ca\(^{2+}\) levels reflects its removal from the cytosol, for instance into the endoplasmic reticulum (ER). The latter depends to a significant degree on Ca\(^{2+}\) pumps localized in the ER membrane (sarco/ER Ca\(^{2+}\) ATPases; SERCA pumps). Therefore, their inhibition should strongly affect Ca\(^{2+}\) activity patterns. Pharmacological inhibition of SERCA pumps by cyclopiazonic acid (CPA, 10 μM) strongly prolonged Ca\(^{2+}\) transients. The results obtained from these experiments suggest that a temperature-dependent slowdown of Ca\(^{2+}\) extrusion from the cytosol could explain the differences in the observed Ca\(^{2+}\) activity patterns (Figure 4(c),(d)). Notably, lowering the temperature only partially slows down ATP-driven Ca\(^{2+}\) uptake processes, whereas CPA blocks ATP-driven Ca\(^{2+}\) uptake to a greater extent. As in Figure 4(b), analyses of the ratio of maximal positive to negative temporal Ca\(^{2+}\) changes were performed for each pixel of an event (Supplementary Figure S4). At low temperatures, the uptake is substantially blocked homogeneously throughout the cells. Overall, the MTED analysis presented here successfully captured complex signaling patterns and enabled us to compare them between experiments.

3.5 | Event characteristics in situ

We next analyzed astrocytic Ca\(^{2+}\) events in organotypic slice cultures from the mouse hippocampus. In such preparations, the morphological

![Figure 4](image-url)
FIGURE 5  Temperature phenotype of Ca\(^{2+}\) activity is not restricted to primary astrocyte cultures. (a),(b) Average \(\Delta F/F_0\) (left images) and the fraction of active time, which is the relative time for each pixel above a Ca\(^{2+}\) threshold of 0.2 (right images), detected in organotypic hippocampal slice cultures at 37°C (a) and 25°C (b). Comparably higher \(\Delta F/F_0\) with spatially more extended and longer lasting events was detected at 25°C, similar to primary astrocyte cultures. Scale bars = 20 μm. (c) 2D histograms of the selected parameters maximum size, duration, and traveled distance of events characteristic for organotypic slice cultures incubated at 37°C, and 25°C incubation temperature depict similar changes in features as observed in primary cultures (n = 6 for 37°C and n = 5 for 25°C from N = 3 independent cultures). (d) Averaged 2D histogram of control measurements in organotypic slice cultures showing prolonged event duration at 37°C compared to primary astrocyte cultures (red framed area, left panel). Blocking the neuronal activity with TTX reduces the occurrence of long-lasting Ca\(^{2+}\) events and exhibits similar 2D histogram features as primary astrocyte cultures at 37°C. (e) Statistical evaluation of Ca\(^{2+}\) activity with duration >90 s (red framed area in (d)) as a function of the Ca\(^{2+}\) threshold applied in the MTED algorithm. Data show mean and SEM. Two-way ANOVA with Sidak’s multiple comparisons post-hoc test (n = 10 recordings from N = 3 experiment days in (d) and (e)).
organization of the hippocampus is mostly preserved, and the maturation of different cell types, network connections, and receptor/channel expression are closer to the situation in vivo (Gähwiler et al., 1997; Gogolla et al., 2006). Figure 5(a),(b) show the average ΔF/F₀ and the fraction of active time detected for the same preparation at 37 and 25°C. Notably, in organotypic slices, we obtained a similar temperature-dependence of astrocyte Ca²⁺ characteristics as in primary astrocyte cultures. At 37°C, the maximal Ca²⁺ levels were lower and events shorter and spatially more restricted, whereas at 25°C, astrocytes exhibited long-lasting Ca²⁺ events of lower frequency (Figure 5(c)). Detailed analysis of Ca²⁺ event duration in primary astrocyte cultures and organotypic hippocampal slices at 37°C revealed an additional set of long-lasting high-magnitude events in the latter preparations (areas in open red boxes, compare Figure 5(c) to Figure 3(c)). As the main difference between these two preparations is the absence of neurons in primary astrocytic cultures, we hypothesized that long-lasting Ca²⁺ events may be a consequence of neuronal activity. Supporting this hypothesis, we observed a similar Ca²⁺ activity pattern in astrocyte-neuron co-cultures (Supplementary Figure S5). To verify the impact of neurons in the hippocampal slice preparations, we blocked neuronal activity by applying tetrodotoxin (TTX; Figure 5(d)). This treatment led to a pronounced reduction in long-lasting Ca²⁺ events in both organotypic preparations and astrocyte-neuron co-cultures, shifting Ca²⁺ activity to an event pattern observed in primary hippocampal astrocyte cultures at 37°C (Figure 5(e), Supplementary Figure S5). These findings demonstrate that the MTED algorithm can identify the impact of neuronal activity on astrocytic Ca²⁺ activity.

3.6 | Event detection in recordings obtained in vivo

To further investigate the versatility of the MTED approach, we evaluated astrocytic Ca²⁺ activity in vivo, for which the recorded fluorescence signal F is commonly much smaller, resulting in more challenging quantification of Ca²⁺ signals. To this end, we implanted a cortical cranial window into transgenic mice with astrocytic expression of GCaMP3 (Figure 6(a)). We set the excitation power between 30 and 40 mW to avoid light-induced Ca²⁺ activity and tissue damage, rarely detecting more than one photon per pixel and time point. Subsequently, we recorded and compared the Ca²⁺ activity patterns of the same cortical region in anesthetized and awake mice (Figure 6). Figure 6(b) depicts the average ΔF/F₀ obtained, which coincides with the fraction of active time and shows overall low activity in the cortex of an anesthetized mouse. When the volatile anesthetic isoflurane was removed and mice woke up, the Ca²⁺ activity in the same region was drastically increased (Figure 6(b), Supplementary Figure S6 and Movie S6). Our analysis revealed that, compared to anesthetized conditions, awake mice had higher magnitude Ca²⁺ events and increased frequency of long-lasting events (Figure 6(c), Supplementary...
Figure S6(c)–(e)). Figure 6(d) displays the differences in the Ca\textsuperscript{2+} activity pattern between anesthetized and awake conditions. Changes in frequency are most prominent at threshold levels 0.5 and 1 and for the subset of 1.5–7 s lasting events, which was supported by the statistical analysis (Figure 6(e)).

Overall, the Ca\textsuperscript{2+} activity in cortical astrocytes observed by in vivo imaging showed similar features in the Ca\textsuperscript{2+} pattern as obtained from cultured astrocytes measured at 37°C. The datasets further identify and define the impact of neuronal activity on a selective subset of Ca\textsuperscript{2+} characteristics of astrocytes (Figure 6(c) and Figure 5(d)). Importantly, the only necessary parameters for MTED of Ca\textsuperscript{2+} signals were the thresholds of \( F_0 \) and Ca\textsuperscript{2+}. More importantly, our approach robustly detected and comprehensively characterized Ca\textsuperscript{2+} event patterns not only in vitro and in situ but also in vivo.

4 | DISCUSSION

4.1 | Establishing a new event detection algorithm for analyzing astrocytic Ca\textsuperscript{2+} signals

With the development of fluorescent indicators of biological activities and metabolites, it has become possible to visualize structural activity and function on a subcellular level. Using GCaMP indicators to monitor the Ca\textsuperscript{2+} signals in astrocytes revealed the complexity of these processes, which cannot be described as on or off activity (i.e., below or above a particular threshold). Ca\textsuperscript{2+} event characterization in terms of magnitude, duration, propagation, regional connectivity, and correct event origin provides valuable information about this signaling pathway and is fundamental for understanding the functional role of Ca\textsuperscript{2+} activity in astrocytes.

During the past years, analysis of astrocyte Ca\textsuperscript{2+} activity has been continuously improved. Two examples of algorithms are CaS-CaDe and AQuA (Agarwal et al., 2017; Wang et al., 2019), which use statistics-based data standardization techniques to estimate and remove the per-pixel fluorescence baseline. Such standardization, however, results in constant values, which does not permit to adapt to fluctuation in baseline fluorescence within the temporal evolution of a pixel. In contrast, MTED’s \( F_0 \) estimation is able to adapt to these fluctuations and can thus compensate changes in fluorescence at basal levels of Ca\textsuperscript{2+} for the time-course of each individual pixel.

In addition, CaS-CaDe uses classical, static ROIs to represent Ca\textsuperscript{2+} events. These are then analyzed and classified using a Support-vector machine (SVM), as a machine learning approach. AQuA extends the concept of static ROIs to so-called dynamic events, possessing the ability to adapt to morphological changes of a Ca\textsuperscript{2+} event on a per-frame basis. In both cases, detected ROIs are then subjected to further analysis.

In the present study, we further improve the flexibility of analysis by developing the MTED algorithm, a novel event detection strategy for time-lapse data obtained by fluorescence microscopy. It is pixel-based and uses multiple thresholds. We chose the latter because signal-to-noise-based strategies for detection of events are prone to underestimating activity in regions with low fluorescence intensity (e.g., low indicator concentrations). This can lead to missed events because the signaling amplitudes stay below a single threshold. The multi-threshold approach developed here overcomes this problem. We applied our analysis to data of astrocytic Ca\textsuperscript{2+} signaling as a test case. It enabled us to quantitatively analyze multiple aspects of Ca\textsuperscript{2+} event patterns that evolve in time and space. Indeed, the approach maintained its robustness when weak signals from small astrocytic processes in more intact preparations were analyzed. Characterization of such weak signals is important because those processes are in close proximity with the extracellular matrix, surrounding cells, and synapses, and also represent the origin of the majority of Ca\textsuperscript{2+} events in astrocytes (Bindocci et al., 2017; Wu et al., 2019). This robustness is also important for other fluorescent indicators that do not display the large fluorescence changes of modern Ca\textsuperscript{2+} indicators, such as many FRET-based sensors.

For all indicators, the relationship between indicator fluorescence and ligand concentration requires careful assessment. In many cases, this relationship is non-linear, and it needs careful consideration to what extent this affects the interpretation of the results. In our study, we used a common measure of fluorescence intensity and its changes (\( \Delta F/F_0 \)). While this quantification has its drawbacks, see for instance (Semyanov et al., 2020), it is straight-forward and abundantly used throughout biomedical research and therefore suitable for testing the analytical approach and its versatility. Since the MTED algorithm is oblivious to the type of data it is working on and does not make specific assumption about it, it can be fed with more quantitative data (e.g., \([\text{Ca}^{2+}]\) after calibration of the imaging setup). Furthermore, it is not limited to Ca\textsuperscript{2+} signaling and could be applied to various fluorescent indicator families and readouts.

In this study we used only a fraction of parameters provided by MTED (compare Figure S3) which are already rather complex due to their multi-dimensionality. Further reduction of complexity can be achieved by applying dimensionality-reducing approaches such as principal component analysis, as recently suggested by Vaidyanathan et. al (Vaidyanathan et al., 2021). Depending on the scientific question those data could be also further examined with cluster analysis strategies such as k-Means.

4.2 | Temperature-dependence of Ca\textsuperscript{2+} signaling in astrocytes

Comparing the Ca\textsuperscript{2+} event detection strategies, some predominately describe only fast and regionally restricted event spots, from which they conclude that discussing frequency patterns in restricted active regions is sufficient for the characterization of astrocytic Ca\textsuperscript{2+} activity (Agarwal et al., 2017; Wang et al., 2017). Other approaches identify a substantial fraction of long-lasting events with regional growth and overlapping patterns, which require the analysis of dynamic regions (Wang et al., 2019; Wu et al., 2019). Here, we observed both types; when examining Ca\textsuperscript{2+} activity at physiological temperature (i.e., 37°C),
we observed the first, whereas imaging the same region at RT led to observation of the latter scenario. Our MTED algorithm sufficiently describes both states of this reversible behavior. We revealed that changes in Ca²⁺ event patterns rely on changes in Ca²⁺ clearance from the cytosol, which is temperature-dependent (Arrhenius, 1889; Komin et al., 2015; Schipper et al., 2014). A relatively straight-forward explanation is that when the Ca²⁺ uptake is slowed down, Ca²⁺ diffuses to larger volumes, accumulates, and initiates Ca²⁺-induced Ca²⁺ release leading to high magnitude Ca²⁺ transients. Blocking the Ca²⁺ uptake by CPA at 37°C indeed switched the Ca²⁺ activity profile to the patterns obtained at 25°C. Remarkably, minor deviations from the species-dependent physiological temperature, such as cooling to 34°C (a temperature typically used in patch-clamp experiments), cause prominent aberrations in the characteristics of Ca²⁺ activity.

Our observation of the temperature-dependent Ca²⁺ activity has broad consequences for interpreting published findings, as well as for future experimental designs. Deviations between astrocytic Ca²⁺ behavior obtained in vitro, in situ, and in vivo may be partly explained by different environmental conditions: 25°C for primary cultures, 34°C for organotypic slices, and 37°C for in vivo measurements. More importantly, Ca²⁺ activity patterns obtained in ex vivo experiments at 37°C were similar to those in vivo. The dissection of the temperature dependence of astrocytic Ca²⁺ signaling demonstrates that our analytical approach can be used to compare experimental results across preparations and experimental conditions.

4.3 Cellular environment and state of consciousness shape Ca²⁺ signals in astrocytes

As a second test scenario, we explored how astrocytic Ca²⁺ event patterns are shaped by neuronal activity. We found, for instance, that the occurrence of long-lasting events can be reduced by blocking action potentials in organotypic slices and mixed cultures of astrocytes and neurons with the application of TTX. This adds to previous studies on how neuronal activity modifies astrocytic Ca²⁺ signaling (Bazargani & Attwell, 2016), by identifying the subset of Ca²⁺ events that is impacted. The obtained results suggest that the connectivity between co-cultured neurons and astrocytes is not as dense as in organotypic slice cultures (compare Figure 5(d),(e) and Supplementary Figure S5). An advantage of the presented analysis is the simpler visualization and identification of groups of events that depend on a specific experimental condition, in our example the presence of neurons/neuronal activity. Furthermore, the location and time of this special group of long-lasting Ca²⁺ events can be identified and could potentially be correlated with nearby neuronal structures. More generally, this approach may help with associating spatiotemporal patterns and properties of cellular events reported by fluorescence with biologically relevant mechanisms and conditions.

As a first step toward the latter, we analyzed Ca²⁺ transients of anesthetized and awake animals. Doing so confirmed that Ca²⁺ activity in astrocytes is reduced under anesthetized conditions (Bojarskaite et al., 2020; Thrane et al., 2012; Vaidyanathan et al., 2021). Interestingly, this is partly because of a reduction of pronounced Ca²⁺ events with large magnitude whereas the frequency of small magnitude events did not significantly change: an important distinction and refinement. This is a third example of how the MTED analysis can extract important additional information from time-lapse fluorescence microscopy, a type of data set ubiquitous in neurobiology and beyond.

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CONFLICT OF INTERESTS

The authors declare no competing interests.

AUTHOR CONTRIBUTIONS

Conceptualization: Andre Zeug, Volodymyr Cherkas, Gebhard Stopper. Methodology: Andre Zeug, Volodymyr Cherkas, Gebhard Stopper. Investigation: Franziska E. Müller, Volodymyr Cherkas, Gebhard Stopper, Laura Stopper, Laura Caudal, Laura Caudal. Visualization: Franziska E. Müller, Andre Zeug. Supervision: Evgeni G. Ponimaskin, Frank Kirchhoff, Christian Henneberger. Writing–original draft: Franziska E. Müller, Andre Zeug. Writing–review & editing: Franziska E. Müller, Volodymyr Cherkas, Gebhard Stopper, Laura Caudal, Laura Stopper, Frank Kirchhoff, Christian Henneberger, Evgeni G. Ponimaskin, Andre Zeug. This manuscript is part of the PhD thesis of Franziska E. Müller.

DATA AVAILABILITY STATEMENT

The Matlab code developed in this study is available from the corresponding author upon request.

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REFERENCES

Agarwal, A., Wu, P.-H., Hughes, E. G., Fukaya, M., Tischfield, M. A., Langseth, A. J., Wirtz, D., & Bergles, D. E. (2017). Transient Opening of the Mitochondrial Permeability Transition Pore Induces Micromodular Calcium Transients in Astrocyte Processes. Neuron, 93(3), 587–605. e7. https://doi.org/10.1016/j.neuron.2016.12.034
Araque, A., Carmignoto, G., Haydon, P. G., Oliet, S. H. R., Robitaille, R., & Volterra, A. (2014). Gliotransmitters Travel in Time and Space. Neuron, 81(4), 728–739. https://doi.org/10.1016/j.neuron.2014.02.007
sleep depth and duration via separate GPCR pathways. eLife, 10, e63329. https://doi.org/10.7554/eLife.63329
Venugopal, S., Srinivasan, R., & Khakh, B. S. (2019). GECIquant: Semi-automated Detection and Quantification of Astrocyte Intracellular Ca2+ Signals Monitored with GCaMP6f. In M. De Pittà & H. Berry (Eds.), Computational Glioscience (S. 455–470). Springer International Publishing. https://doi.org/10.1007/978-3-030-00817-8_17
Volterra, A., Liaudet, N., & Savtchouk, I. (2014). Astrocyte Ca2+ signalling: An unexpected complexity. Nature Reviews Neuroscience, 15(5), 327–335. https://doi.org/10.1038/nrn3725
Volterra, A., & Meldolesi, J. (2005). Astrocytes, from brain glue to communication elements: The revolution continues. Nature Reviews Neuroscience, 6(8), 626–640. https://doi.org/10.1038/nrn1722
Wang, Y., DelRosso, N. V., Vaidyanathan, T. V., Cahill, M. K., Reitman, M. E., Pittolo, S., Mi, X., Yu, G., & Poskanzer, K. E. (2019). Accurate quantification of astrocyte and neurotransmitter fluorescence dynamics for single-cell and population-level physiology. Nature Neuroscience, 22(11), 1936–1944. https://doi.org/10.1038/s41593-019-0492-2
Wang, Y., Shi, G., Miller, D. J., Wang, Y., Wang, C., Brousard, G., Wang, Y., Tian, L., & Yu, G. (2017). Automated Functional Analysis of Astrocytes from Chronic Time-Lapse Calcium Imaging Data. Frontiers in Neuroinformatics, 11, 1–23. https://doi.org/10.3389/fninf.2017.00048
Wu, Y.-W., Gordleeva, S., Tang, X., Shih, P.-Y., Dembitskaya, Y., & Semyanov, A. (2019). Morphological profile determines the frequency of spontaneous calcium events in astrocytic processes. Glia, 67(2), 246–262. https://doi.org/10.1002/glia.23537
Wu, Y.-W., Tang, X., Arizono, M., Bannai, H., Shih, P.-Y., Dembitskaya, Y., Kazantsev, V., Tanaka, M., Itohara, S., Mikoshiba, K., & Semyanov, A. (2014). Spatiotemporal calcium dynamics in single astrocytes and its modulation by neuronal activity. Cell Calcium, 55(2), 119–129. https://doi.org/10.1016/j.ceca.2013.12.006

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