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Reinforcing interdisciplinary collaborations
to unravel the astrocyte “Calcium Code”

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Abstract

In this review article, we present the major insights from and challenges faced in the acquisition, analysis and modeling of astrocyte calcium activity, aiming at bridging the gap between those fields to crack the complex astrocyte “Calcium Code”. We then propose strategies to reinforce interdisciplinary collaborative projects to unravel astrocyte function in health and disease.

Keywords: astrocyte, glia, calcium, interdisciplinary

Introduction

Astrocytes, the most abundant non-neuronal cells of the nervous system, are essential to brain function, from synaptogenesis and neurotransmission to higher brain functions such as memory and learning [1]. Those functions of astrocytes are altered in various brain diseases such as epilepsy, brain tumours, neurodegenerative diseases, Down syndrome, major depressive
disorder and schizophrenia [1]. Astrocytes notably respond to stimuli with transient elevations in cytosolic calcium concentration, referred to as calcium signals. Those calcium signals are essential to brain function and are altered in various brain diseases [1, 2]. Importantly, astrocyte calcium signals can trigger the release of molecules referred to as gliotransmitters that modulate neuronal communication at synapses (for recent reviews on gliotransmission and the associated controversies, see [3, 4]). Better understanding astrocyte physiology and the communication between astrocytes and other cells of the central nervous system thus relies on our ability to make sense of those calcium signals. Astrocyte calcium signals are characterized by diverse spatial (from microdomains to signals spreading within astrocyte networks) and temporal characteristics (from hundreds of milliseconds to tens of seconds) [2]. The majority of those signals occur in fine astrocyte compartments (50-200 nm), referred to as processes, that account for as much as 80 % of the volume of an astrocyte, yet cannot be resolved by diffraction-limited light microscopy [2, 5] (see Fig. 1). This strongly hinders our ability to characterize astrocyte calcium activity, from the molecular pathways involved to the quantification of the spatio-temporal properties of the signals. Consequently, the functions of the various signals observed, referred to as the astrocyte “Calcium Code”, remain unclear. Better characterizing astrocyte activity concomitantly with the activity of other brain cells will be essential to unravel the roles of astrocyte calcium signals in brain function [6]. Please refer to the review [2] for more details on the current challenges associated with the study of calcium signals in astrocytes.

Fig. 1 Confocal image of an astrocyte expressing GCaMP6f (maximum intensity projection over time) that shows its different structural compartments and their size.
In this review article, we highlight the importance of reinforcing interdisciplinary collaborations to crack the astrocyte “Calcium Code”, with a focus on the characterization of the properties of astrocyte calcium signals. We present the major insights from and challenges faced in data acquisition, analysis and modeling of astrocyte calcium activity and propose strategies to facilitate and strengthen collaborations between these fields, which are essential to unravel the functions of astrocyte calcium signals in health and disease.

**Acquisition of astrocyte calcium signals**

Data acquisition is the first step to characterize astrocyte calcium activity. In this section, we present a brief overview of the tools that are available for imaging astrocyte calcium signals, both in slices and *in vivo*. We further highlight the insights, challenges and perspectives associated with measuring calcium signals in astrocytes.

**Imaging tools for astrocyte calcium acquisition**

The development of calcium indicators, which change their fluorescence properties when binding to calcium ions, allowed neuroscientists to study astrocyte calcium activity. Numerous indicators exist, characterized by diverse kinetics and diffusion properties, so that they should be chosen carefully. In the early days, chemical calcium dyes, such as Fluo-4 or Oregon Green BAPTA, were commonly used [7–12]. One of the main caveats of these chemical sensors is the low signal to background noise ratio of the resulting signals, which only allows visualizing calcium signals in the soma and the main thick branches of astrocytes (see Fig. 1), unless loaded through a patch-clamp recording pipette and visualized with high-resolution microscopy [13–15]. More recently, the development of genetically-encoded calcium indicators (GECIs) [16] has considerably improved our understanding of astrocyte calcium dynamics. Various GECIs have been developed in the last years that can be imaged by different tools, for precise or wide imaging at cellular or subcellular levels [17–21]. GECIs have several advantages compared to classical calcium dyes. First, they can be easily targeted to be expressed specifically in astrocytes. Moreover, they provide a higher signal to background noise ratio compared to classical calcium dyes and diffuse better into the fine processes. Additionally, GECIs can be expressed in live organisms, thus allowing *in vivo* calcium imaging in anesthetized [21–24], awake head-fixed [17, 25–27] or freely-moving mice during consecutive behavioral sessions [25, 28, 29]. While many GECIs have been designed in the last years for neurons, only a few are available to target astrocytes specifically (reviewed in [30]). These GECIs have different spectral, temporal and spatial properties that make them suitable for specific experimental applications [31]. Importantly, they yield calcium signals with different spatio-temporal properties that may not be comparable and may be difficult to analyze with certain software (see section Analysis of astrocyte...
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calcium signals).

Astrocytes display most of their activity in their fine processes. The majority of those signals are spatially-restricted, forming so-called microdomains, and display strikingly diverse spatio-temporal properties [32]. Understanding the physiological relevance of those calcium signals requires powerful imaging techniques that can be used in combination with complementary methods to manipulate astrocyte and neuronal activity, such as electrophysiology, optogenetics, pharmacology and behavioral tests. Notably, because of the small size of astrocyte processes, high-resolution microscopy is needed to obtain a thorough view of the astrocyte calcium activity. Both confocal and two-photon microscopy are good options for imaging astrocyte calcium activity because these setups are generally compatible with other techniques, allowing for the study of calcium signals at the tripartite synapse level in slices and in vivo in anesthetized [21–23, 27] or awake head-fixed mice [17, 24–26]. Light sheet fluorescence microscopy (LSFM) and Lattice LSFM are novel imaging techniques that allow fast 3D scanning with low phototoxicity and a resolution comparable to confocal microscopy [33, 34]. Therefore, those techniques are excellent imaging options for experiments in brain slices. Please refer to Table 1 for an overview of optical resolution, phototoxicity/photobleaching, and compatibility of the different imaging techniques. High-resolution microscopy allows recording calcium signals at a high acquisition speed (in the order of ms) but its spatial resolution is limited by diffraction (x-y: 0.2-0.3 µm and z: 0.5 µm at best) and, thus, does not allow visualizing fine processes in detail. Recent studies have used super-resolution microscopy such as stimulated emission depletion (STED) and stochastic optical reconstruction microscopy (STORM) to study astrocyte morphology at the tripartite synapse level in live tissue [35–37]. STED microscopy has revealed that the complex spongiform morphology of astrocyte processes contains functionally-isolated nanostructures that are characterized by spatially-restricted calcium signals [37]. Currently, because of their low acquisition speed and high laser intensity, which induces high photobleaching of calcium sensors, super-resolution microscopy techniques cannot be used to perform calcium imaging. Thus, in the aforementioned study, calcium signals were acquired using high-resolution microscopy and were then mapped onto super-resolution structural images. Super-resolution imaging requires powerful computational tools, both for acquisition and analysis, which are not broadly available in the experimental community (in terms of knowledge, software and hardware) and emphasizes the need to establish collaborations between experimental and computer scientists.

The need for interdisciplinary approaches

It is important to keep in mind that experimental approaches have inherent limitations. First, calcium indicators are calcium buffers. Therefore, calcium indicators compete with calcium binding sites in the cell, altering calcium signals and the normal functioning of the cell. Second, the spatial and temporal
Characteristics of the measured signals are constrained by the imaging technique as well as the kinetics of the calcium indicator used. It is thus possible that some faster or smaller calcium signals than those currently reported exist in astrocytes that cannot be detected by the calcium imaging tools that are currently available. Importantly, this effect can be amplified during 3D scanning for calcium signals that are faster than the z-scanning time. Lastly, experimental manipulations, such as using a knock-out mouse line or bath applying drugs, can have unexpected off-site effects that can impact the results, making it difficult to extract causal relationships between the experimental manipulation and the obtained results. Collaborative work with computational scientists is essential to build mechanistic models to go beyond those limitations. For example, models have been essential to characterize the effect of the concentration, kinetics and diffusion coefficient of calcium buffers, such as calcium indicators, on calcium dynamics [38, 39]. Models can thus be used to predict the free calcium signals that would occur in the absence of indicators. Further, models can measure in silico calcium signals at very high spatial and temporal resolution (depending on the method used, see section Modeling astrocyte calcium signals), thus predicting the range of calcium signals that could not be resolved experimentally.

Analysis of astrocyte calcium signals

In the quest of characterizing astrocyte calcium signals, the key role of analysis is to provide tools to experimentalists and modelers to process their data, of increasing size and complexity. Statistical as well as advanced computational image analysis tools are thus needed. In this section, we focus on the analysis of calcium images, which is meant to quantify what is observed, i.e., to extract meaningful information or measurements from images. In particular, we emphasize the importance of developing computational image analysis tools dedicated to the quantification of astrocyte calcium signals, and the challenges to get there.

Image analysis to characterize astrocyte calcium signals

Decoding the astrocyte “Calcium Code” involves the characterization of the spatio-temporal dynamics of astrocyte calcium signals. Computational image analysis tools aim at accurately detecting all astrocyte calcium signals in a sequence of microscopy images and, for each signal, extracting its dynamical and spatial features, such as its amplitude, its duration, its trajectory, its propagation speed, the location from which it originates and its volume. Various information, such as the number of calcium signals in a specified region or cell, their frequency at a position, and the different types of signals induced by a stimulus, can be deduced from those measurements. From an image analysis point of view, reaching this ideal of output information requires preprocessing steps (e.g., denoising, deconvolution, motion
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Table 1 Overview of the main calcium imaging techniques used to study astrocyte calcium signals. * The ability to perform 3D fast scanning depends on the scanning method that the microscope uses, which varies depending on its hardware settings. ** Photobleaching and phototoxicity can be high at the focal plane with two-photon microscopy because it uses high intensity lasers, but it is low if the whole sample is considered (see [40]). *** Note that Light sheet fluorescence microscopy (LSFM) and Lattice LSFM cannot be used in vivo in postnatal murine models but can be used in vivo in embryos.

| Imaging method     | Optical resolution | Photobleaching & phototoxicity | Preparation | Compatible with fast 3D scanning * | Compatible with other techniques |
|--------------------|--------------------|---------------------------------|-------------|-----------------------------------|-----------------------------------|
| Wide-field         | Soma & main branches | High                            | In vitro & in vivo (anesthetised & head-fixed) | No                                | Electrophysiology, pharmacology, wide-field photostimulation |
| Confocal           | Soma, main branches & fine processes | High                            | In vitro & in vivo (anesthetised & head-fixed) | No                                | Electrophysiology, pharmacology, localized photostimulation |
| Two-photon         | Soma, main branches & fine processes | ** Low                           | In vitro & in vivo (anesthetised & head-fixed) | Yes, depending on the microscope | Electrophysiology, pharmacology, localized photostimulation |
| LSFM               | Soma, main branches & fine processes | Low                             | *** In vitro | Yes, faster than two-photon         | Electrophysiology, pharmacology |
| Lattice LSFM       | Soma, main branches & fine processes | Very low                        | *** In vitro | Yes, faster than LSFM               | Electrophysiology, pharmacology |
| Fiber photometry   | Population         | High                            | In vivo (freely behaving)         | No                                | Electrophysiology, wide-field photostimulation |
| Miniscopes         | Soma               | High                            | In vivo (freely behaving)         | No                                | Wide-field photostimulation |

correction) as well as the detection, the segmentation and the quantification
of the calcium signals in a sequence of microscopy images, which is very chal-
 lenging due to the complex nature of these signals. First, calcium signals are
 characterized by various durations (from milliseconds to tens of seconds), fre-
 quencies and signal-to-noise ratios. Second, their spatial spreads vary from
 microdomains to signals that propagate within the astrocyte in regions of
 various sizes and shapes. Third, they can overlap in space and time [41]. As
 most signals occur in fine astrocyte processes that cannot be fully resolved by
 diffraction-limited light microscopy techniques, image analysis methods cannot
 rely on morphological criteria to detect calcium signals, which also complex-
 ifies their quantification. In addition, the developed image analysis methods
 should ideally operate across data with different spatial scales, taken in vivo
 or in vitro, and acquired with different imaging techniques.

Lack of computational image analysis tools adapted to
the complexity and diversity of the data

Recently, several image analysis algorithms have been developed to quantify
astrocyte calcium signals in 2D+time microscopy images. Among them, we
 can cite GECIquant [42], CaSCaDe [17], FASP [43], AQuA [44] and, more
 recently, Begonia [45] and Astral [46]. Most of these methods are ROI-based
 approaches (ROI: region of interest), meaning that calcium signals are ana-
 lyzed through fixed spatial boundaries in the image. As the spatial spread of
 calcium signals can vary over time and become larger than or get out of the
 ROI, those approaches can lead to inaccurate or partial detection of the sig-
 nals. To solve this issue, event-based algorithms have been developed, such as
 AQuA [44]. For more details about these algorithms (e.g., analysis approach
 and outputs), please refer to the dedicated section in the review article from
 [47].

The aforementioned analysis tools have considerably improved the detec-
 tion and characterization of astrocyte calcium signals. Yet, their use can be
 restricted, either because they are not adapted to the diversity of acquisition
 modes and calcium indicators (see section Acquisition of astrocyte calcium
 signals) or because they are not open-source or not user-friendly [48]. This can
 constrain some neuroscientists to implement “in-house” analysis pipelines,
 which is time-consuming and less reproducible, or to use tools that were ini-
 tially developed for neuronal calcium imaging analysis, such as CaImAn [49],
 Suite2P [50] and LC_Pro [51]. This latter approach is not optimal as astrocytes
differ from neurons in many ways. For example, they have a different morphol-
 ogy. Further, notably because astrocytes are not polarized cells, their calcium
 signals display different spatio-temporal properties than the ones of neurons.

The continuous scientific and technical advances in calcium imaging will
 always call for new and adapted image analysis algorithms. Until now, most of
 the quantification of calcium signals has been performed on 2D+time fluores-
 cence microscopy data. The recent emergence of 3D+time imaging techniques
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gives access to new and major structural and dynamical information, such as
the number of calcium signals occurring in an entire astrocyte volume, their
synchronization, their trajectory and the location from which they originate
[5]. To the best of our knowledge, there is currently no image analysis tool to
detect, segment and quantify astrocyte calcium signals in 3D+time microscopy
images.

Challenges hindering the development of 3D+time image
analysis tools

The main reason why the quantification of astrocyte calcium signals has been
so far restricted to 2D+time images is because of the trade-off between tem-
poral and spatial resolution in microscopy techniques. The access to a refined
3D imaging of the dynamical behavior of calcium signals in astrocytes is quite
recent, owing to the emergence of microscopes enabling a high 3D spatio-
temporal resolution with low phototoxicity (e.g., lattice light sheet fluorescence
microscopy (LSFM) [33, 34] and of genetically encoded calcium indicators
(GECIs) [16]. Despite these scientific and technical advances, the development
of 3D+time image analysis tools tailored for the astrocyte calcium activ-
ity is not straightforward and calls for new quantitative analysis algorithms
with new constraints and challenges. First, a key challenge in the develop-
ment of 3D+time image analysis tools is the memory and computational costs
required to process large 3D+time data. To give the reader an idea, the equiv-
alent of one hour of acquisition of Lattice LSFM data represents about 1 To
of data. Importantly, the developed analysis tools should be accessible and
thus ideally be able to run on standard desktop computers. To tackle this
challenge, ingenious solutions for image processing are needed such as using
data-dimensionality reduction techniques. Second, and more critical, reliable
and large amounts of labeled data are not available. Such datasets are cru-
cial to evaluate image analysis tools and to train data-driven tools, which are
increasingly common with the emergence of deep learning in biological image
analysis [52]. Manually annotating 3D time-lapse images is a tedious task –
mainly because of the complex visualization in 4D space – that cannot be per-
formed reliably. There is a significant intra- and inter-experimenter variability.
There is currently a major lack of annotations of astrocyte calcium activity
images. Note that this is also true for other datasets of 3D images in live tis-
sue [53]. For all of those reasons, a common and promising approach is to use
realistic synthetic datasets with known ground-truths (i.e., all morphological
and dynamical properties are known and controlled) to train and quantita-
tively assess the performance of analysis software. This highlights the need for
developing models and simulators that are able to mimic real image sequences.
Need for public realistic synthetic datasets: join the forces!

To solve the difficulty to reliably label calcium signals in microscopy images, a promising approach consists in generating 3D+time synthetic datasets that realistically depict astrocyte calcium signals observed in real microscopy images. To be as realistic as possible, the simulation should be driven by a biophysical model that describes the calcium signals at the nanoscale, which requires close collaboration between image analysts, modelers (see section Modeling astrocyte calcium signals) and experimentalists (see section Acquisition of astrocyte calcium signals). For instance, a recent interdisciplinary collaboration [54] has resulted in the creation of a simulator to generate realistic sequences of 3D lattice LSFM images depicting calcium activity in the sponge-like network of astrocyte processes by integrating a simplified version of the kinetic model developed by Denizot et al., 2019 [55]. In addition to hopefully opening the door to the deployment of 3D+time image analysis tools to quantify astrocyte calcium activity, these simulators could also help modelers tuning their models and the parameters in a faster way than using computational simulations, which are often time and computationally expensive. A major challenge to develop such simulators is the complexity of evaluating the similarity between the generated synthetic images and real images. Implementing rigorous methods to evaluate synthetic astrocyte calcium images will thus be essential to ensure the success of this approach. Note that these synthetic data are essential to guide analysts in the development of their algorithms, but final qualitative validation on real images is still required.

Modeling astrocyte calcium signals

Models correspond to simplifications that describe relevant parameters of a system of interest (its elements, their states and their interactions), allowing for better quantification, visualization, and understanding of the system. The famous quotation from George Box, “All models are wrong but some are useful” [56], highlights that models are incomplete representations of the system as a whole, yet provide crucial insights into the system’s behavior and dynamics. Such insights would not be grasped by a model as complex as the system of interest itself.

Depending on the question and hypothesis that emerge from experimental data, modelers choose different approaches and toolkits (see Table 2). For example, models studying calcium activity in microdomains will need a higher spatial resolution than models of somatic signals. Further, the modeling approaches that are well-suited to study calcium microdomains, such as particle-based methods (see [57] for a review), are more accurate but extremely demanding in terms of computational power and simulation time. Simulating hundreds of seconds of calcium activity in a fine astrocyte process (e.g., 1 µm long, 200 nm in radius) can take days to compute, so that using those tools
to simulate signals in a whole cell or in a network of cells is currently unfeasible. Please note that the computation time to simulate e.g., 1 millisecond of calcium activity varies not only depending on the modeling technique used, but also on the computational resources available in each laboratory, on the volume and number of reactions modeled as well as the simulation time. To learn more about the different approaches that can be used to model reactions, their insights and limitations, please refer to dedicated reviews [57, 58]. The goal of this section is not to present an exhaustive list of astrocyte models (see [57, 59, 60]), to review existing models of calcium signaling (reviewed in [61–66]), or to present a detailed list of modeling tools to model calcium signals [67, 68]. Rather, we emphasize the key insights that can be gained from models of astrocyte calcium activity as well as the challenges that computational neuroscientists are currently facing.

Insights from modeling into biological processes

Mathematical and computational models are powerful tools that provide new insights in the mechanisms that regulate calcium activity in astrocytes and generate testable predictions. First, models can be used to conduct *in silico* experiments that are time-consuming or unfeasible experimentally. Models have for example been used to finely tune the spatial distribution of calcium channels (molecules that, when open, result in a calcium influx into the cytosol, forming a calcium signal) within the cell and explore its impact on astrocyte activity [55, 69]. Moreover, models can be used to generate realistic datasets that can be used to train tools developed to characterize the system’s behavior (see section Analysis of astrocyte calcium signals) [54]. Lastly, computational models are useful to go beyond correlational observations and to propose mechanistic principles that explain experimentally-observed data. For example, models have shown the effect of cellular morphology on the compartmentalization of calcium signals in dendritic spines [70–73] as well as in astrocyte processes [74]. For a recent review on the insights gained from computational approaches on astrocyte function as well as strategies to start incorporating astrocyte calcium signals in systems neuroscience to better understand how astrocytes contribute to brain computation, see [75]. Overall, modeling approaches can provide key insights to astrocyte physiology.

Main challenges associated with the development of models of astrocyte calcium activity

Computational neuroscientists are facing major challenges to build models of astrocyte calcium activity. First, a lot of data are currently missing or not shared publicly, so that most parameter values used in the astrocyte models that are currently available are derived from data obtained in other cell types. Those data include the concentration and sub-cellular distribution of endogenous buffers, the diffusion coefficient of diffusing molecules involved in calcium dynamics in astrocytes, the distribution of the major calcium channels and
Table 2 Brief summary of the main modeling approaches that are commonly used to model astrocyte calcium activity, their insights, limitations and examples. Biological processes are inherently noisy. When the system that is modeled contains a large number of molecules, this noise can be averaged. Such models are called deterministic and describe the variation of molecular concentrations over time. They are often used to describe calcium signals at the whole cell and at the network levels. When the system of interest contains a small number of molecules or ions, typically small subcellular compartments like astrocyte processes, this approximation is no longer valid and the stochastic nature of molecular reactions has to be taken into account in the model. Further, models can be spatial, i.e. take into account the position and potential diffusion of molecules in the cell, or well-mixed, i.e. at each time step, any molecule can virtually move anywhere in the cell. The location of the molecules and cell morphology is thus not taken into account in well-mixed models. * Calcium concentration in spatial stochastic simulations can be deducted from the number of molecules tracked and the system’s volume. ** Some spatial stochastic techniques track individual molecules (particle-based) while others track the number of molecules in small sub-compartments (voxel-based). See e.g. [76] for a review.

Note that the characteristics presented in this table are indicative as the usage and computational cost of a given model vary greatly depending on the precise method implemented and the number of molecules/reactions modeled (see [57, 58] for reviews).

| Name of the modeling approach | Spatial variation of concentration | Tracks individual molecules | Computational cost | Common use | Examples |
|-------------------------------|----------------------------------|-----------------------------|--------------------|------------|----------|
| Well-mixed, deterministic     | No                               | No                          | Very low           | Astrocyte network/w-hole cell | [77–79] |
| Well-mixed, stochastic        | No                               | Yes                         | Low                | Astrocyte network/w-hole cell | [80–82] |
| Spatial, deterministic        | Yes                              | Yes                         | Low-intermediate   | Whole cell/Signal propagation in major branches | [83–89] |
| Spatial, stochastic           | Yes                              | Yes *                        | Yes **             | High       | Spongiform domain [55, 90] |
Reinforcing interdisciplinary collaborations to unravel the astrocyte “Calcium Code” activity. Bridging those models together is critical to better understand the involvement of local calcium signals in higher-level brain functions such as cognition and learning. Building such multi-scale models is challenging but should provide unprecedented insights in the involvement of astrocyte calcium signals in the activity of neural circuits and overall in brain (dys-)function.

Is there such a thing as a generic astrocyte model?

Although astrocytes share common morphological and biochemical characteristics, they are remarkably heterogeneous. The diversity of astrocyte morphology has been described as early as 100 years ago by Cajal and morphology-based classifications of astrocytes have been proposed [91]. Astrocyte electrophysiological properties [92–94], gene [95–97] and protein expression levels [98] also vary drastically depending on the brain region under study. Those observations suggest that astrocytes are a heterogeneous cell population, questioning the specificities and roles of individual sub-types. For more details, see dedicated reviews on astrocyte heterogeneity [1, 99–101]. Whether the diverse functions of astrocytes in the brain rely on molecularly and morphologically distinct sub-populations of astrocytes is still poorly understood, yet crucial to uncover the functions of astrocytes in the healthy and diseased brain. A recent study identified sub-populations of astrocytes that selectively contributed to specific functions such as synaptogenesis and tumor invasion of glioma [102]. Incorporating this diversity in astrocyte models by building models of specific sub-populations of astrocytes rather than the currently available generic astrocyte models will be essential to provide insights into the functional implications of the molecular and morphological heterogeneity of astrocytes that have been reported recently.

Need for interdisciplinary collaborations to improve models of astrocyte calcium activity

Several strategies and perspectives could be developed to go beyond the aforementioned challenges to model astrocyte calcium activity. First, computational neuroscientists would highly benefit from the existence of open-source datasets, which could be used to build and test models. Such datasets are crucial for data-driven modeling practices, which rely on strong iterative collaborative work between experimentalists and computational neuroscientists. Moreover, several good practices and step-by-step modeling guides have been published to ensure the reproducibility of models [103–105]. Lastly, initiatives such as the Neuromatch Academy courses and conferences [106, 107] provide an unprecedented opportunity to build an accessible, democratic, inclusive, international and interdisciplinary community aiming at using computational approaches to improve our understanding of brain function.
Perspectives

Astrocytes are cells that display a highly complex activity that is essential to brain function. Characterizing the diverse signals displayed by active astrocytes and understanding their physiological roles, the “Calcium Code”, are the biggest challenges of the field and are crucial to understand the involvement of astrocytes in brain function. In this short review article, we highlighted the different insights that can be gained from each field that studies calcium signals in astrocytes and the major challenges that they are facing. Key challenges that prevent us from making sense of astrocyte calcium activity have arisen from our discussions during our interdisciplinary workshop, hosted by the 1st Virtual Conference of the European Society for Neurochemistry “Future perspectives for European neurochemistry – a young scientist’s conference”, in May 2021, entitled “Let’s join forces - Bridging the gap between experimental, computational and data sciences to disentangle astrocyte calcium activity”. Those challenges include:

• The development of analysis tools allowing accurate detection and characterization of individual calcium signals in astrocytes are lacking, notably in 3D+time.
• There is no consensus in key definitions and terminology, which further hinders efficient communication across fields (e.g., calcium microdomain/nanodomain, processes/leaflets, Calcium Code, gliapil/spongiform domain).
• A lot of data is missing to fully grasp the mechanisms regulating astrocyte calcium signals and their physiological roles. For example, local and regional variability of the expression levels of proteins involved in calcium signaling, both in health and disease, remain to be characterized. The morphology of perisynaptic astrocyte processes and their organelles, together with their dynamical remodeling, also remain to be uncovered in live tissue.
• Raw data are rarely shared in public repositories. Notably, labeled datasets are needed to evaluate image analysis tools and to train data-driven tools. Providing public access to such datasets has contributed to fast improvements in other fields, such as the development of tools detecting the onset of epileptic seizures [108].
• Interdisciplinary events and projects are rare, which constitutes a major barrier to our efforts to unravel the astrocyte “Calcium Code”. Indeed, scientists from different fields lack opportunities to discuss, share ideas and knowledge. The interactions between fields working on astrocyte calcium signals and opportunities for improvements are highlighted in Fig. 2. We believe that such joint efforts are essential to fully grasp the complex properties and functions of astrocytes.

Reinforcing interdisciplinary projects, bringing together experts from different fields, will be crucial to ensure our success in cracking the astrocyte “Calcium Code”. Such collaborative projects are still rare in the field, which might result from the high fragmentation of research projects and fields working on astrocyte physiology, often presenting their work in different, highly
specialized conferences and journals. We propose initiatives that will facilitate the emergence of new interdisciplinary projects:

- Agreement on shared definitions and terminology across fields.
- Sharing datasets, together with all the relevant information on the data acquisition, processing and modeling (if relevant) methods used. This might require the creation of an online platform to store and discuss data on astrocytes.
- Sharing user-friendly data analysis tools, including providing the code in open-access and the dataset(s) used to facilitate their dissemination to the whole community.
- Organization of recurrent meetings and events that bring together experts from various fields of expertise.

Because of the complexity of astrocyte morphology and signaling, interdisciplinary projects will be essential to not only crack the astrocyte “Calcium Code”, but also to successfully improve our understanding of astrocyte (patho-)physiology and to propose models of astrocyte function.
Data availability statement

Data sharing is not applicable to this article as no datasets were generated or analysed during this study.

Ethical statement

Ethics approval and consent to participate

Not Applicable.

Consent for publication

Not Applicable.

Availability of data and materials

Not Applicable.

Competing interests

The authors have no relevant financial or non-financial interests to disclose.

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Author contributions

Ana Covelo wrote the first draft of the data acquisition section, Anaïs Badoual wrote the first draft of the data analysis section and Audrey Denizot wrote the first draft of the introduction, modeling and perspectives sections. All authors commented on previous versions of the manuscript, read and approved the final manuscript.

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