Probabilistic Encoding of Stimulus Strength in Astrocyte Global Calcium Signals

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Astrocyte calcium signals can range in size from subcellular microdomains to waves that spread through the whole cell (and into connected cells). The differential roles of such local or global calcium signaling are under intense investigation, but the mechanisms by which local signals evolve into global signals in astrocytes are not well understood, nor are the computational rules by which physiological stimuli are transduced into a global signal. To investigate these questions, we transiently applied receptor agonists linked to calcium signaling to primary cultures of cerebellar astrocytes. Astrocytes repetitively tested with the same stimulus responded with global signals intermittently, indicating that each stimulus had a defined probability for triggering a response. The response probability varied between agonists, increased with agonist concentration, and could be positively and negatively modulated by crosstalk with other signaling pathways. To better understand the processes determining the evolution of a global signal, we recorded subcellular calcium “puffs” throughout the whole cell during stimulation. The key requirement for puffs to trigger a global calcium wave following receptor activation appeared to be the synchronous release of calcium from three or more sites, rather than an increasing calcium load accumulating in the cytosol due to increased puff size, amplitude, or frequency. These results suggest that the concentration of transient stimuli will be encoded into a probability of generating a global calcium response, determined by the likelihood of synchronous release from multiple subcellular sites.

Key words: astrocyte, calcium puffs, crosstalk, ATP, glutamate

Introduction

Astrocyte calcium signaling has emerged in the last few decades as a potential mechanism through which nonexcitable glial cells can contribute to information processing in the brain (De Pitta et al., 2012). Although the computational potential of calcium signaling has been much speculated upon, the basis through which information about neuronal (and vascular) network activity can be encoded into astrocyte calcium signals, and decoded through glial responses, remains uncertain.

Numerous neurotransmitters and hormones have been linked to calcium elevation in astrocytes (FiaccO and McCarty, 2006; Scemes and Giaume, 2006; Verkhratsky et al., 1998), and calcium has been shown to regulate many aspects of astrocyte function—from the more “passive” homeostatic roles (Kimelberg, 2010) to release of gliotransmitters that can directly modulate neuronal function (Araque et al., 2014). As such, the theory that astrocytes can detect synaptic activity and respond by modulating neuronal function at an individual and network level has gained currency (Haydon, 2001; Ransom et al., 2003). Despite this general consensus, many areas of dispute remain. The most striking current controversy is the equivocal effect of deletion of the type 2 inositol (1,4,5) trisphosphate receptor (InsP3R)—which abolishes somatic calcium responses in astrocytes—on the neurophysiological traits that had been hypothesized to depend on neuron–glial communication, such as synaptic plasticity and learning. Some investigators have reported that this intervention does not result in overt defects in synaptic transmission or plasticity (Aguilhon et al., 2010; FiaccO et al., 2009), while others have reported deficits (Navarrete et al., 2012; Takata et al., 2011).

This uncertainty has led to a refocusing of attention on the spatiotemporal nature of astrocyte calcium signals (Araque...
et al., 2014; Rusakov, 2015; Srinivasan et al., 2015). Localization of genetic calcium indicators to submembrane regions (Shigetomi et al., 2010) and ultrastructural analysis of astrocyte morphology (Patrushev et al., 2013) have reinforced the view that subcellular calcium signals can be restricted to defined microdomains of the astrocyte, where the receptors and signaling machinery linked to neuron–glial transmission are likely to be localized. As such, a hypothesis is emerging that such local signaling can mediate rapid feedback between the cell types, leading to synaptic modulation, without the need for the calcium increase to spread to a range that encompasses the somatic regions of the cell (Khakh and Sofroniew, 2015; Srinivasan et al., 2015). This fits with existing concepts of the hierarchy of calcium signals (Bootman et al., 1997), where subcellular puffs of calcium summate to trigger a positive feedback process of calcium-induced calcium release from internal stores that generates a cell-wide calcium wave—which we will term a “global” calcium signal.

Despite this proposal for the primacy of local calcium signals for mediating gliotransmission, a good deal of evidence exists for the generation of global (even network-wide) calcium waves in astrocytes under physiological conditions (Ding et al., 2013; Hirase et al., 2004; Nimmerjahn et al., 2009; Wang et al., 2006). These global signals must presumably have a role in physiology, and they must arise from underlying local signaling events. At present, however, the conditions under which global signals evolve from local signals and whether the global signals are able to encode information about the stimulus that triggered them are poorly understood.

Existing analyses of the encoding of stimulus strength into global calcium waves have focused predominantly on how the interval between waves varies with stimulus strength or type. By analogy to neuronal spike trains, these repetitive waves of calcium release from intracellular stores are often termed “calcium spikes” (Skupin et al., 2008). A leading hypothesis is that the frequency of calcium spiking is proportional to the strength of stimulus (Berridge et al., 2003), as exemplified by hepatocytes responding to increasing concentrations of a muscarinic receptor agonist (Woods et al., 1986). A more recent analysis explored this mechanism in detail, and demonstrated an exponential relationship between mean interspike interval and agonist concentration (Thurley et al., 2014). These authors concluded that fold changes in the average stochastic period of the interspike interval proved the most robust parameter for encoding changes in the stimulus strength into cellular calcium responses.

In general, however, astrocytes seem poorly equipped to encode stimulus strength into interspike interval, despite their capacity to readily generate oscillations in response to neurotransmitter stimuli (James et al., 2011). Evidence for frequency modulation is weak, and the capacity for information to be encoded in interspike interval (at least for spontaneous calcium signals) is small (Skupin et al., 2008). In some respects, this makes intuitive sense, as exploiting patterns in spike interval as an encoding mechanism requires prolonged stimulation. While this may occur for vasoactive compounds, the time scale for neurotransmitter and neuromodulator signaling is orders of magnitude too fast to allow for multiple calcium waves to be generated from a single stimulus event under normal conditions.

Previous work in our laboratory explored the possibility that the kinetics of the calcium response generated by different transmitters could encode information about the strength or type of stimulus, and reached the conclusion that discrimination on these grounds was not feasible due to the overlap in temporal profiles between different stimuli (James et al., 2011). Here, we explore an alternative hypothesis: that the strength of a transient stimulus could be encoded as a probability of generating a global calcium signal from underlying subcellular events. We show a robust concentration–response relationship between agonist concentration and probability of generating a global signal, that the probability varies between transmitters, and that neuromodulators that do not themselves generate calcium responses can act as gain modulators—both positively and negatively—for other transmitters. The origin of this probabilistic response is the requirement for the synchronous release of calcium from subcellular “puff” sites, leading to the initiation of a cell-wide wave.

As such, the pattern of global calcium signals in an astrocytic network will be a composite of the type of transmitter released, the concentration of transmitter, duration of transient, and coincidence detection between signaling pathways under the dynamic conditions encountered in vivo.

**Materials and Methods**

**Ethics Statement**

Experiments were performed in accordance with guidelines outlined in the code of practice for humane killing under Schedule 1 of the UK Home Office Animals (Scientific Procedures) Act 1986. Experiments were performed according to policies on the care and use of laboratory animals of British Home Office and European Community laws and were approved by The University of Nottingham Animal Welfare and Ethical Review Body.

**Preparation of Cerebellar Astrocyte Cultures**

Under sterile conditions, neonatal (P1–P2) rat pups were sacrificed by cervical dislocation and decapitation before whole brains were removed. A dissection microscope was used to remove the cerebellum before transferring to separate vessels containing ice cold dissection buffer composed of: Ca$^{2+}$- and Mg$^{2+}$-free Hank’s HEPES-buffered saline solution (Invitrogen, Paisley, UK), 22 mM glucose, 20 mM HEPES, 100 U mL$^{-1}$ penicillin, 100 µg mL$^{-1}$ streptomycin, 0.1 mM...
i-serine, 0.5 mM t-glutamine, and 1 mM sodium pyruvate (pH adjusted to 7.2 with NaOH). The meninges and blood vessels were removed using fine forceps before the dissected tissue was chopped finely and dissociated enzymatically in buffer supplemented with 15 μg mL⁻¹ papain (45 min, 37°C). Tissue was then mechanically dissociated via repeated trituration through fire-polished Pasteur pipettes in the presence of 20 mg mL⁻¹ DNase in medium containing DMEM (Invitrogen), 4500 mg L⁻¹ glucose, 4 mM t-glutamine, 110 mg t-pyruvate, 10% fetal bovine serum, 50 μg mL⁻¹ t-proline, 100 U mL⁻¹ penicillin, and 100 μg mL⁻¹ streptomycin. Cells were plated at a density of 5 × 10⁴ cells mL⁻¹ on poly-L-lysine-coated coverslips or 96-well plates. Medium was refreshed 24 h after plating, and every 3–4 days thereafter.

**Live Cell Ca²⁺ Imaging**

Methods were similar to James et al. (2011). Briefly, astrocytes were loaded with 1 μM of the Ca²⁺ indicator Fluo-4 AM (Invitrogen) for 30 min in buffer containing: 135 mM NaCl, 3 mM KCl, 10 mM HEPES, 15 mM t-glucose, 2 mM MgSO4, and 2 mM CaCl₂. Subsequently de-esterification of accumulated Fluo-4 AM occurred by leaving cells in buffer for a further 30 min before transferring to the stage of a BD Pathway 855 High Content cell analyser for 96 well plate experiments, a Brunell inverted microscope (10×, 0.25 NA) for WIN 55,212-2 treatment experiments or an Olympus IX70 inverted microscope (20×, 0.75 NA objective) for all other treatments. Cells were excited at 450/20 nm (exposure 100 ms, frame rate 1 Hz) and fluorescence emission detected at 535/50 nm with an ORCA-ER camera (Hamamatsu, Welwyn Garden City, UK).

For imaging Ca²⁺ puff events, astrocytes were loaded with 2 μM Oregon Green 488 BAPTA-1 AM (Invitrogen) for 30 min and then de-esterified in buffer for 30 min before transferring to the stage of a Nikon Eclipse Ti inverted microscope (60×, 1.4 NA oil immersion objective). Cells were excited at 470/10 nm with an exposure of 30 ms and continuously imaged for 60 s. Fluorescence emission of 550/10 nm was detected with a Zyla 4.2 sCMOS camera (Andor Technology, Belfast, UK) and recorded in Micromanager (Edelstein et al., 2010). Agonists/modulators were added by manual displacement of bath solution with excess buffer containing agonist/modulator at the indicated concentrations.

**Image Analysis**

For global Ca²⁺ imaging data, a time-series analyzer plugin to ImageJ (available at http://rsb.info.nih.gov/ij; Wayne Rasband, National Institutes of Health, Bethesda, MD) was used to manually define regions of interest (ROIs) centered on the cell nucleus. Mean fluorescence intensity within each ROI was quantified and expressed as the ratio of fluorescence at time t divided by mean intensity for 0–20 s before addition of agonist (Ft/F₀). Cells were classified as global responders if mean Ft/F₀ increased >1.045-fold (at least 3 SD above F₀). Area under the curve (AUC) is calculated as the integral of fluorescence intensity during the 3-min time window of agonist stimulation [(Ft/F₀) dt]. Cells were excluded from analysis if Ca²⁺ signals occurred before addition of agonist.

For Ca²⁺ puff imaging data, cells were first segmented manually in ImageJ to quantify whole cell mean fluorescence and a circular ROI centered within the nucleus to quantify mean nuclear fluorescence. A bespoke algorithm written using Matlab R2015a (The MathWorks, Natick, MA) was used to automate the detection of Ca²⁺ puff events within each cell. Raw data time series images of segmented cells were time-binned such that mean and standard deviation of each pixel were calculated every five frames. For each frame of the time-binned image sequence, the difference in pixel intensity to the previous frame was calculated. An individual pixel is defined as a candidate puff region pixel if its difference value to the previous frame is ≥1.5 × SD (SD taken from previous frame of time-binned SD values). Puff regions were subsequently identified as the regions where the number of spatially connected candidate pixels in a single frame was ≥16. A circular ROI (radius = 0.65 μm) is centered on the puff region and mean fluorescence intensity of the ROI expressed as the ratio of fluorescence at time t divided by mean intensity from the first 90 ms before addition of agonist/buffer (Ft/F₀). Subsequent filtering of puff regions discounts any regions that do not show a reduction in Ft/F₀ ≥50% puff amplitude within 750 ms of the detected puff time (to exclude long-term trends such as wave initiation). Each region matching these criteria is counted as an individual puff event. Puff area is calculated as pixel count within puff region × 0.108 (pixel to μm conversion factor). A cell is classified as having transitioned to a global wave if ΔFt/F₀ ≥0.045 relative to baseline and the wave is detectable in the nucleus.

**Statistical Analysis**

All statistical tests were performed with GraphPad Prism™ software (GraphPad Software, La Jolla, CA). The test applied, numbers of cells, and P values are reported in figure legends.

**Results**

**Number of Cells Responding**

Previous work has shown that the kinetic parameters governing global calcium signals generated in cultured cerebellar astrocytes, screened with high-throughput imaging methods, did not show any clear dependence on agonist concentration or agonist type (James et al., 2011). In contrast, we observed that the number of cells responding to a given agonist (ATP, glutamate, or histamine) appeared to show clear concentration dependence. To quantify this, we measured the percentage of cells responding to a stimulus in each well of a multiwall plate, and calculated the mean percentage of responding cells for all wells that were treated with the same stimulus (18 wells per agonist concentration).

The percentage of cells responding showed a concentration-dependent increase with agonist concentration for ATP, glutamate, and histamine (Fig. 1A,B). However, the maximal percentage of cells varied substantially between agonists: at the highest concentration tested (1 mM), ATP triggered Ca²⁺ signals in 85% of cells, whereas glutamate only triggered responses in 37% and histamine 46% of cells. The agonists therefore differ in the efficacy with which they can trigger a global Ca²⁺ elevation in astrocytes.
These features could be accounted for by two hypotheses. First, the concentration responsiveness may reflect heterogeneity in cell sensitivity. In this scenario, increasing agonist concentration would recruit more and more cells as the threshold for triggering a cell-wide response was breached (NB: the observation that the vast majority of cells exhibit a Ca\textsuperscript{2+} signal in response to high-affinity pharmacological agonists suggests that the existence of appreciable numbers of nonresponders is unlikely; James et al., 2011). The second possibility is that all cells are equally sensitive, and the percentage of responding cells reflects the probability that any given cell will respond. In this scenario, the concentration dependence will arise as the probability of any one cell responding increases with agonist concentration.

A method for discriminating between the foregoing possibilities would be to repetitively apply agonist to a population of cells at a submaximal concentration (leaving time for recovery between applications). If the percentage of cells responding reflects a continuum of sensitivities, then the same high-sensitivity subpopulation of cells should respond each time. Alternatively, if cells have equivalent sensitivities, but respond in a probabilistic manner, then there would be variability in the identity of responding cells between applications (i.e., any cell could respond, and chance determines which ones do).

**A Probabilistic Basis for Cellular Responses**

We applied ATP to a population of astrocytes at a submaximal concentration (0.1 μM) for 3 min, three times in succession, with a 5-min washout period between each addition (Fig. 2A). On each occasion, around 40% of cells responded with a global Ca\textsuperscript{2+} signal within the 3-min stimulation period (Fig. 2B). However, the specific cells responding varied from addition to addition. We observed all cases: cells that responded on 0, 1, 2, or 3 occasions (Fig. 2A,C). These results were consistent with the second hypothesis outlined in the last section—that each cell has a chance of responding, and the probability of response depends on ATP concentration.

There is a danger with repetitive application of agonists in this manner that later Ca\textsuperscript{2+} responses are altered by preceding ones; phenomena such as receptor desensitization (or internalization), store depletion, or feedback loops (such as autophosphorylation) may change the dynamics of the later Ca\textsuperscript{2+} signals. To test for this, we compared the AUC [integral of fold change, \((F/F_0)\times s\)] of the spikes evoked by 0.1 μM ATP for each stimulus. There was no detectible difference in this parameter over the three trials (Fig. 2B).

The results suggest that astrocytes respond to agonists in a probabilistic manner, rather than by varying in sensitivity. For a model based purely on chance, where all cells are equivalent, then the distribution of the total population into the four possible response classes (0, 1, 2, or 3 responses) can be predicted. If 40% of cells respond to 0.1 μM ATP, then the distribution of cells into the response classes (0, 1, 2, or 3 responses) would be 21.6, 43.2, 28.8, and 6.4%, respectively. The observed distribution diverged from this ideal case, with
fewer cells than predicted giving a single or two responses, and more cells than predicted giving three responses (Fig. 2C).

We next assessed the response of the astrocytes to glutamate and histamine, which generate a response in a lower percentage of cells, to investigate whether the same probabilistic trend was evident (Fig. 3A,B). For glutamate (5 μM), around 30% of cells responded for the first two additions and, as for ATP, the number of responses observed covered all possibilities: 0, 1, 2, or 3 events per cell. In contrast to ATP, however, the percentage of responding cells decreased for the third trial, and the AUC for glutamate responses progressively declined (Fig. 3A). These results suggest that although glutamate operates probabilistically, there is detectable desensitization over the course of the trials. Histamine responses were similar to ATP. Over the three trials, around 12% of cells responded, and the cells that responded varied from trial to trial, but with no apparent desensitization between trials (Fig. 3B).

Another potential complication in interpreting results is the capacity of astrocytes to sustain intercellular waves through gap-junctional coupling and paracrine signaling (Giaume and Venance, 1998). This could result in the transient application of agonists leading to a signal initiated in one cell propagating into neighboring cells, meaning response probability may be overestimated owing to indirect coupling (although under our culture conditions, cells are not confluent and rarely exhibit long-range calcium waves). To test for this directly, we applied 0.1 μM ATP to the cells for 3 min as usual, followed by addition of 100 μM of the gap junction blocker carbenoxolone (Rozental et al., 2001), and another 3-min transient application of ATP. The presence of carbenoxolone did not significantly alter the percentage of cells responding to ATP (51.3% before and 53.7% after; \( P = 0.623 \), Fisher’s exact test, \( n = 298 \)), or the mean AUC for fluorescence change in responding cells [240.4 ± 4.10 before and 243.3 ± 3.29 (\( F/F_0 \))s after treatment; \( P = 0.092 \) Mann–Whitney test, \( n = 160 \)], suggesting that gap-junctional coupling did not detectably affect responses under normal conditions. These results suggest that propagation of calcium responses between neighboring cells is unlikely to significantly distort interpretation of the percentage of responding astrocytes.

Overall, this analysis suggests that although there is heterogeneity in sensitivity among the cells to the different agonists, each cell displays a given probability of responding to a low amplitude, transient stimulus.

Gain Modulation of Response Probability

Many neuromodulators and neurotransmitters have been shown to trigger calcium signaling in astrocytes, directly or indirectly (Fiacco and McCarthy, 2006). There is also a wealth of research demonstrating crosstalk between the calcium signaling apparatus and other second messenger signaling networks (Bruce et al., 2003; Clementi and Meldolesi, 1997), perhaps best exemplified by the range of signals that converge on the InsP3 receptor (Foskett et al., 2007). This suggests that coincident activation of neurotransmitter receptors coupled to calcium release and modulatory receptors coupled to pathways that affect calcium signaling could alter the probability of generating a global calcium response.

Endocannabinoid signaling mediates neuromodulation in many neuronal classes and many brain regions (Howlett...
et al., 2004). In astrocytes, activation of cannabinoid receptor type 1 (CB1R) has been linked to calcium responses (Navarrete et al., 2014). To test for modulation of response probability, we stimulated cells with 0.1 mM ATP, with and without co-application of the CB1 agonist WIN 55,212-2 (WIN, 1 μM). When WIN was present (1-min preincubation), there was a significant increase in the percentage of cells responding to ATP with a global calcium signal (Fig. 4A). For those cells that responded, treatment with WIN had no effect on the total cytosolic calcium flux, as the AUC of the responses was unaffected (Fig. 4A). Washout of WIN and subsequent challenge with ATP alone showed that the potentiation of probability was reduced, but still statistically significant (Fig. 4A). This result suggests that WIN is able to act as a positive modulator of the probability of evoking a global calcium response in the astrocytes.

Nitric oxide (NO) signaling has been reported to affect multiple aspects of the calcium signaling network, including InsP₃ receptors (Cavallini et al., 1996; Komalavilas and Lincoln, 1996; Rooney et al., 1996), ryanodine receptors (Stoyanovsky et al., 1997; Takasago et al., 1991), store-operated and nonstore-operated calcium entry channels (Li et al., 2003; Moneer et al., 2003), and many other elements of the calcium-handling apparatus (Clementi and Meldolesi, 1997). The nature of this modulation appears to vary between cell types and experimental conditions, with NO being able to activate or inhibit several of these targets. In astrocytes, NO has been shown to accelerate the reloading of calcium stores (Li et al., 2003), increase the rate and range of calcium wave propagation through gap-junctionally coupling networks of cells (Willmott et al., 2000), and increase the frequency of spontaneous calcium oscillations in cultured astrocytes at physiological temperatures (Schipke et al., 2008). Direct activation of calcium responses by NO has also been reported (Bal-Price et al., 2002). We tested the effects of NO on ATP responses, under similar conditions to WIN.

Co-administration of NO (with 1-min preincubation) and 0.1 μM ATP led to a significant increase in the percentage of cells responding to the stimulus (Fig. 4B). NO also significantly increased the AUC of the calcium responses (Fig. 4B). In contrast to WIN, the enhancement due to NO fully reversed after washout, with the response to a third ATP stimulus returning to pretreatment levels (Fig. 4B).
To test whether these modulators alone are able to trigger calcium responses in astrocytes, we applied WIN and DETA/NO for 18 min (Fig. 4C). WIN had no detectible ability to trigger calcium responses under these conditions (Fig. 4C), but addition of the NO donor DETA/NO (100 μM) to the astrocytes triggered global calcium responses in

![FIGURE 4: Positive gain modulation of global responses. (A) Representative traces (upper panels) of cells repetitively stimulated with 0.1 μM ATP (filled bars), with 1 μM WIN 55,212-2 present for 1 min before and during the second stimulation (open bar). Aggregate data (lower panels) for % of responding cells and mean AUC (±SEM). ***P<0.0001 for % response (Fisher’s exact test), relative to the first ATP stimulus; n = 1,322. (B) Representative traces (upper panels) of cells repetitively stimulated with 0.1 μM ATP (filled bars), with 100 μM DETA/NO present for 1 min before and during the second stimulation (open bar). Aggregate data (lower panels) for % of responding cells and mean AUC (±SEM). ***P<0.0001 for % response (Fisher’s exact test) and AUC (Kruskal–Wallis test with Dunn’s correction for multiple comparisons), relative to the first ATP stimulus; n = 650. (C) Representative traces of astrocytes exposed to WIN 55,212-2 alone (upper trace) and DETA/NO alone or in the presence of 100 μM PPADS (lower traces) for 18 min (bars). Right panel: percentage of cells exhibiting global calcium responses in cells exposed to WIN alone (n = 464), DETA/NO alone (n = 439), or cells exposed to DETA/NO in the presence of PPADS (n = 373). ***P<0.0001 relative to DETA/NO alone (Fisher’s exact test).]
approximately 50% of cells (Fig. 4C). However, these responses had a long latency (mean = 318 ± 21 s), which could indicate an indirect effect on calcium signaling rather than direct calcium influx or release from stores due to NO acting as an agonist. We repeated the experiment in the presence of an antagonist for P2 receptors (PPADS, 100 μM), which substantially reduced the number of NO-evoked responses (Fig. 4C). This result indicates that the long-latency spikes triggered by NO depend on P2 receptor activation—suggesting that NO is acting indirectly by increasing the number of global calcium signals spontaneously occurring through release of ATP from the astrocytes. Accordingly, the positive modulation of calcium responses to transient ATP stimulation by WIN and DETA/NO is due to enhancement of P2 receptor-evoked responses, not direct release of calcium following CB1R or soluble guanylyl cyclase activation.

In addition to the positive gain modulation exhibited by NO and CB1 receptors, we explored whether negative modulation could also be observed. A number of TRP channels have been reported in astrocytes (Verkhratsky et al., 2014) and linked to basal elevation of cytoplasmic calcium concentration (Shigetomi et al., 2013). Gadolinium is a broad-spectrum blocker of TRP channels (Bouron et al., 2015), and so we tested the impact of this ion on the probability of generating a global response to ATP. Gd³⁺ applied 1 min before subsequent co-application with ATP caused a substantial reduction in the percentage of cells responding to ATP (Fig. 5A). Gd³⁺ also reduced the total calcium flux (as measured by AUC) in responding cells. This reduction in probability and magnitude of response was fully reversible after washout (Fig. 5A).

Finally, we also tested for the possibility of crosstalk with the second branch of the phospholipase C signaling

![Figure 5: Negative gain modulation of global responses. (A) Representative traces (upper panels) of cells repetitively stimulated with 0.1 μM ATP (filled bars), with 2 μM Gd³⁺ present for 1 min before and during the second stimulation (open bar). Aggregate data (lower panels) for % of responding cells and mean AUC (±SEM). ***P < 0.0001 for % response (Fisher’s exact test) and AUC (Kruskal–Wallis test with Dunn’s correction for multiple comparisons), relative to the first ATP stimulus; n = 641. (B) Representative traces (upper panels) of cells repetitively stimulated with 0.1 μM ATP (filled bars), with 10 nM PMA present for 1 min before and during the second stimulation (open bar). Aggregate data (lower panels) for % of responding cells and mean AUC (±SEM). ***P < 0.0001 for % response (Fisher’s exact test) relative to the first ATP stimulus. *P = 0.012 for AUC of second stimulus compared with first and *P = 0.024 for AUC of third stimulus compared with first stimulus (Kruskal–Wallis test with Dunn’s correction for multiple comparisons); n = 227.]
pathway by applying the direct activator of protein kinase C, phorbol 12-myristate 13-acetate (PMA, 10 nM). PMA alone had no noticeable effect on calcium concentration, but co-application with ATP after 1 min substantially reduced the percentage of cells responding and the AUC of the responding cells. In contrast to the reversible negative modulation exhibited by Gd$^{3+}$, PMA persistently decreased astrocyte responsiveness to ATP after washout (Fig. 5B).

The Origin of Probabilistic Responses
Global calcium responses originate from the summation of underlying subcellular events. To understand the factors that determine the probability of a global signal arising from calcium puffs, we need to understand how the number, frequency, size, and range of subcellular events vary in astrocytes, and so determine why the same submaximal treatment sometimes results in initiation of a global wave but sometimes fails (even within the same cell).

To explore this issue, we used high-speed epifluorescence imaging to measure fluorescence across a whole cell with sufficient spatiotemporal resolution to capture all subcellular events. Although this approach does not allow detailed spatial resolution of puffs, in the manner of optical sectioning methods (Sanderson et al., 2014), its advantage is that the entire cell can be imaged, and so the full complement of puffs can be detected. Our goal was to use this strategy to identify the determining factor in transition from local calcium puffs to a global calcium wave.

Previous studies in a variety of cell types have proposed different mechanisms for how changes in puff parameters lead to wave initiation after agonist addition (Bootman et al., 1997; Parker et al., 1996; Tovey et al., 2001); puff amplitude could increase, the spatial range of individual puff events could expand (enhancing coupling between clusters of InsP$_3$ receptors), or the frequency of puffs could increase. In principle, any of these effects could result in formation of a wave nucleation site, or they may act to simply increase cytosolic calcium concentration above a threshold level that causes a generalized increase in InsP$_3$R open probability (Marchant et al., 1999; Yamasaki-Mann et al., 2013).

High-speed imaging of astrocytes allowed subcellular calcium release events to be identified throughout the cell cytoplasm (Fig. 6A,B). Typically, cells exhibited multiple release sites, at which multiple release events could be observed, and readily distinguished from baseline fluctuations and the downward drift associated with photobleaching under continuous illumination (Fig. 6B). Puffs were resolvable at subcellular sites, despite the mean fluorescence change for the entire cell showing only minor baseline deflections (Fig. 6B). To quantify the parameters of these calcium puffs, we developed an algorithm for identifying fluorescence changes that matched the time course of typical release events (Fig. 6C; for details, see Materials and Methods section) and used it to analyze the number, spatial range, and amplitude of calcium puffs in the astrocytes.

Under resting conditions, in standard imaging buffer, astrocytes exhibited spontaneous calcium puffs that occurred with a mean frequency of 10.6 ± 1.1 min$^{-1}$ (median 7 min$^{-1}$), had mean amplitude of 1.07 ± 0.04-fold ΔF (median 1.06-fold ΔF), and mean area of 45.7 ± 1.6 μm$^2$ (median 31.39 μm$^2$) calculated from 65 cells. Round puff morphology was typically observed with diameters of 7.6 μm (mean) and 6.3 μm (median).

To determine how puffs were changed in response to agonist stimulation, we treated the cells first with a buffer exchange 10 s after initiating image acquisition and then imaged for a further 50 s to determine puff parameters under resting conditions. The same cells were then treated for a second time, with buffer containing 0.1 μM ATP (Fig. 7A–C). Cells were segregated into those that exhibited a global signal (ΔF/ΔF$_{0}$ ≥ 1.045 relative to baseline and increase detectable in the nucleus) and those that only exhibited subcellular responses (Fig. 7A).

The distributions of puff amplitudes in cells without a wave (−w) and cells that transitioned to a wave (+w) were closely similar (Fig. 7B). Puff amplitude was compared within
each cell during treatment with buffer and treatment with ATP (Fig. 7B). No significant changes in amplitude were observed after ATP treatment regardless of whether a global calcium response occurred.

The distribution of puff areas was also similar for cells that transitioned and cells that did not (Fig. 7C). As with amplitude, the area of puffs did not significantly change after treatment with ATP for either class of cell (Fig. 7C). Collectively, the results suggest that changes in the amplitude and size of puffs are not determining factors in whether cells transition from local to global signaling regimes.

We next tested the influence of puff frequency on the probability of transition. We measured the cumulative frequency of puffs in cells treated first with buffer exchange alone, followed by 0.1 μM ATP. For cells that underwent a transition to waves, we stopped counting puffs at the point that wave initiation occurred, as the widespread release of calcium during the rising phase of the wave could not be distinguished reliably from a torrential, widespread series of puff events (which is, in effect, the nature of the calcium-induced calcium release process that constitutes the wave; Bootman et al., 1997). A further rationale for this approach was that our goal was to determine the conditions that lead up to the point of transition, not the dynamics of cytoplasmic calcium after the tipping point had been breached.

The trajectory for cumulative puff events was similar for both classes of cells exposed to buffer initially (with the exception of a few outliers), suggesting that cells that transition in the presence of ATP were not especially “excitable” under resting conditions (Fig. 8A,B). After treatment with ATP, there was a small increase in mean puff number in cells that did not initiate waves (Fig. 8A), but this was not statistically significant (P = 0.059; Wilcoxon signed-rank test), suggesting that puff frequency was not reliability changed by addition of submaximal ATP concentrations.

Despite this apparently undetectable effect on puff frequency, 0.1 μM ATP caused the formation of a global wave in nearly half of the tested cells. However, even in those cells that exhibited global waves, the trajectory of puffs preceding the tipping point was not noticeably different from cells that did not transition (Fig. 8B). The total number of puffs between agonist addition and initiation of a wave in cells exhibiting a global response ranged from 1 to 12 (mode 6), a range of values frequently obtained in cells that did not give global responses (Fig. 8A,B). Accordingly, the absolute number of puffs over the time taken to reach wave initiation (and so, the baseline frequency of puffs) did not correlate with whether or not a global response occurred; it is not possible to predict from the trajectory of puffs whether or not a given cell is going to transition.

Finally, we examined the temporal structure of the puff events to investigate whether a particular pattern of puffs could be identified that occurred commonly in cells that respond globally, but rarely in cells which do not. Puff events were assigned to 300-ms bins (approximately half the duration of a single puff; see Fig. 6C), and aligned with the mean fluorescence intensity for the whole cell and for the nucleus (Fig. 9A–C). In cells exposed to two rounds of buffer exchange, the frequency of puffs did not appear to change before and after treatment, and the distribution of puffs appeared random over
time, with only one or two puffs per bin (Fig. 9A). For cells exposed to ATP which did not exhibit global signals, a similar pattern was observed. The total number of puffs occurring after treatment increased, but the typical number of puffs per bin again remained at one or two (Fig. 9B). In contrast, for cells that exhibited global signals, the moment of initiation was commonly preceded by a single 300-ms time bin in which three or more puffs occurred synchronously (Fig. 9C).

To test the predictive power of this synchronicity more directly, we found the maximum number of puffs/bin for nonresponding and responding cells (Fig. 10). Of the 18 cells with waves, 14 had three or more synchronous puffs, whereas only 6 of 27 nonwave cells had three or more synchronous puffs (Fig. 10).

To determine whether the spatial distribution of puff events influenced the probability of a global wave evolving, we determined the Euclidean distance between puff events (using the centroid of the region where fluorescence was suprathreshold) for all cells that exhibited three or more synchronous puffs. We plotted the minimum and maximum distance between puff sites at the moment of synchronous release for cells that did or did not transition to global responses.

The analysis showed that there was no statistically significant difference in the minimum distance between puff sites in responding and nonresponding cells, although the range of values was greater for responding cells (Fig. 10C). In contrast, responding cells showed a statistically significant increase in the maximum distance between puff sites compared with nonresponding cells (Fig. 10C).

These results suggest that synchrony in puff events (occurring in different regions of the cell)—rather than amplitude, range, or cumulative frequency—is the major determinant of the probability that an agonist will result in a global calcium response under submaximal stimulation conditions.

**Discussion**

Astrocyte calcium signaling is widely perceived to be a mechanism by which these electrically passive cells can make an active contribution to information processing within the brain (Araque et al., 2014; Nedergaard and Verkhratsky, 2012). Despite decades of research into the potential for modulation of neuronal development, synaptic transmission, and synaptic plasticity, there is still much about the computational properties of astrocyte calcium signaling that is poorly understood.

In particular, a recent controversy concerns the equivocal impact of deletion of type 2 InsP$_3$ receptors on the neurophysiological roles in which astrocytes have been implicated, despite overt loss of somatic calcium responses in astrocytes.
plasticity were apparently unaffected by loss of global calcium responses (Agulhon et al., 2010), but cholinergic-dependent long-term potentiation in both cortex (Takata et al., 2011) and hippocampus (Navarrete et al., 2012) requires InsP3R2-dependent calcium signals in astrocytes.

One explanation for apparently benign outcomes after deletion of InsP3R2 is that the spatial organization of calcium signals is crucial to the effectiveness of signal transduction, with InsP3R2-dependent somatic signals being distinct in function from near-membrane calcium signals or other subcellular/microdomain responses (Araque et al., 2014; Di Castro et al., 2011; Grosche et al., 1999; Shigetomi et al., 2010; Srinivasan et al., 2015). Despite this recognition of the importance of spatiotemporal segregation, widespread somatic calcium signals are readily observed in vivo (Hirase et al., 2004; Nimmerjahn et al., 2009; Wang et al., 2006). The roles of these signals are not well understood, however, nor are the potential computational rules that govern their emergence. Here, we have investigated the basis by which global signals evolve from subcellular events and the consequences of such all-or-nothing responses for computation in the face of local, low amplitude, or short-lived stimuli.

**The Origin of Global Calcium Responses in Astrocytes**

A hierarchy of calcium signaling has long been recognized in cell physiology (Berridge et al., 2000). Release of calcium from InsP3Rs can range in space from a unitary “blip” (opening of a single InsP3R channel) to puffs (release from clusters of channels) and waves that represent the coordinated coupling of adjacent clusters of InsP3Rs to result in a spread of release events at the wavefront that progresses through the cell. The mechanistic basis

![Diagram](https://example.com/diagram.png)
of the latter global calcium signals, and how they evolve from underlying elementary events, has been studied in many cell types (Bootman et al., 2001; Cheng et al., 1993; Falcke, 2003; Marchant et al., 1999; Ruckl et al., 2015; Tovey et al., 2001).

Increases in puff amplitude, puff frequency, and puff size have all been reported to follow agonist application, but the common consequence is that cytosolic calcium increases to a threshold that triggers the positive feedback mechanism of calcium-induced calcium release (Bootman et al., 1997).

This tendency results in a situation where immediately after stimulation, cells enter a stochastic phase of calcium signaling, where puff events summate, increasing the “calcium load” on the cytosol until the threshold for calcium-induced calcium release is reached and a wave initiates (Marchant and Parker, 2001; Thul and Falcke, 2004, 2006; Thul et al., 2009). This pushes the cell into a phase where a global signal progresses through the cell in an almost deterministic manner, and the InsP$_3$Rs are inactivated until local calcium concentrations fall once more. Such a bimodal signaling regime has been modeled and shown to be the basis of variation in inter-wave intervals in cells challenged with tonic stimuli (Skupin et al., 2008; Thurley et al., 2014).

Although calcium puffs have previously been reported in astrocytes (Koizumi et al., 2002), the factors determining transition from subcellular to global signals have not been determined. A detailed characterization of the parameters defining puffs in astrocytes could not be obtained with our imaging approach, because wide-field epifluorescent imaging allowed high-speed and high-resolution imaging of the whole cell but with less spatial resolution than optical sectioning methods. This compromise was needed to capture events throughout the whole cell, and means that resolving low amplitude or fast events would be constrained relative to methods such as TIRF or confocal microscopy (Ellefsen et al., 2014). Our goal, however, was to count all puffs and so gain a sense of whether a particular pattern or sequence of puffs gives a high probability of transition to a global wave. We found that astrocytes are in some respects atypical in their subcellular responses.

The analysis showed that the frequency of puffs in astrocytes was low compared with several cell lines (Tovey et al., 2001).
et al., 2001; especially given the more limited measurement volume due to optical sectioning used in these other studies), and that puff size and amplitude were unaffected by agonist application (Fig. 7). As a consequence, even after stimulation with ATP, the total number of puffs occurring remained sparsely distributed in time (Fig. 9). This consistency in the spatiotemporal properties of puffs has the consequence that, under our recording conditions, no obvious build up in bulk cytoplasmic calcium concentration was observed: there appears to be no cumulative increase in calcium “load” that leads inexorably to wave evolution. Instead, the best predictor for wave initiation seems to be the occurrence of the synchronous release of calcium from three or more puff sites (Fig. 10; with a 300-ms window for coincidence). This sudden concerted release appears to be the basis of the all-or-none trigger for a global signal.

Analysis of the spatial distribution of synchronous puffs indicated that an increase in the maximum distance between puff sites was correlated with a greater probability of wave initiation. This suggests that formation of a local hotspot of release from multiple neighboring sites is unlikely to be the trigger for initiation of a global wave; instead the implication is that spatially segregated puff sites causing calcium elevation over a wider area of the cytoplasm triggers the threshold crossing that leads to coordinated calcium-induced calcium release throughout the cell.

As a final note, in common with other reports (Wu et al., 2014), the spatial range of calcium signals varied considerably—indeed some subcellular events occupied a substantial fraction of the whole cell volumes, raising a question as to how a boundary between subcellular and global signals could be defined. In our case, the time course of waves was clearly distinguishable from that of puffs, and we further discriminated on the basis of whether the signal occupied a sufficient range so as to encompass the nucleus. Nevertheless, it should be acknowledged that while stereotypical puffs and waves were the most abundant signal classes (and could be easily distinguished), there is a continuum of intermediate signals that bridge these broad categories in space and time.

**Probabilistic Encoding of Stimulus Amplitude**

The principal consequence of an all-or-none signal based on the chance of several puffs occurring synchronously is that the signal transduction from agonist application to global response is probabilistic. At high concentrations of ATP, probability is high, but for other agonists, the maximum probability appears much lower, indicating that agonist efficacy manifests as the probability that a global signal will initiate (Fig. 1B).

Furthermore, submaximal concentrations, or transient or local applications, would also be predicted to provoke a response with lower probability. As a consequence, even a single cell presented with an identical stimulus on multiple occasions only has finite chance of responding to any individual challenge. As this is likely to be the character of a stimulus encountered in vivo following synaptic release of transmitters or modulators, it seems necessary to consider stimuli to be encoded not as stereotyped and predictable responses, but as a probabilistic signal. Such signals obviously recur in neuroscience, with synaptic transmission being the classic example.

When considering the potential for astrocyte calcium signaling to be involved in information processing, the input–output relationship between synaptic release and global calcium responses would manifest as a response probability. In contrast, the generation of localized calcium signals may be more reliable (high probability), again distinguishing the two modes of signal transduction on a computational as well as mechanistic basis. Ultimately, the determinant for whether a global signal evolves is the coordinated progression of a wave through the cytoplasm. If our results relate directly to the in vivo situation, local signaling in the vicinity of synapses (where receptors are clustered; Arizono et al., 2012) will be transmitted to the cell interior only if diffusion of InsP3 and Ca2+ reaches ER store receptors and triggers the synchronous release of calcium from multiple puff sites.

Determining the applicability of results from primary cultures to the in vivo state is always challenging. Nevertheless, if the astrocytic cytoplasm acts essentially as an “excitable medium” for propagating calcium waves, it seems reasonable to assume that the factors governing summation of subcellular events into cell-wide events could be common between the in vitro and in vivo states. That said, the morphology of the cell would also be a determining factor for wave propagation, and the difference between protoplasmic cultured cells and the more diffuse and ramified anatomy of astrocytes in vivo may well alter wave propagation trajectories. Interestingly, the analysis of puff spatial distribution suggests that puffs occurring synchronously at more distant sites increase the likelihood of transition to a global signal. In situ this may translate into global calcium signals becoming more probable if multiple local signaling events occur synchronously—for example, if there is coordinated release from multiple synapses that are all ensheathed by the same astrocyte. From a computational perspective, this offers the potential for spatial summation of activity in the synaptic network by the astrocyte.

**Gain Modulation of Global Signals**

A final consideration in understanding how probabilistic encoding can influence computation is that the response probability can be altered by stimuli that do not themselves cause calcium release. This modulation can be positive or negative, and transient or prolonged (Figs. 4 and 5).

There are numerous examples of crosstalk between neuromodulators, such as NO and endocannabinoids, and
calcium signaling. Modification of the pumps, channels, transporters, and buffers of the calcium signaling "toolkit" (Berridge et al., 2000) could alter many of the properties of calcium puffs, in this case, presumably altering the likelihood of synchronous events or the conditions under which puff to wave transitions occur.

Gain modulation of this type has been described in situ in hippocampal slices. Functional independence was demonstrated between Schaeffer collateral and alveus inputs to astrocytes, with selective engagement of glutamate and acetylcholine receptors, respectively. Co-stimulation of the inputs led to bidirectional, frequency-dependent modulation of astrocyte calcium responses, due to crosstalk between the signaling pathways (Perea and Araque, 2005).

Gain modulation has also recently been described in vivo. Astrocyte somatic calcium responses in the cortex depend on release of noradrenaline from locus coeruleus projections (Ding et al., 2013; Espallergues et al., 2007), despite ex vivo responses being mediated by glutamate released during local network activity. A possible explanation for this discrepancy comes from Paukert et al. (2014) who showed that astrocytes in the visual cortex did exhibit global responses to local transmission, but the responses were markedly enhanced by synergistic noradrenergic transmission.

These results suggest that the frequency of global responses that constitute the major part of somatic calcium signals detected by intravital imaging methods is highly sensitive to crosstalk between signaling pathways, suggesting that arousal increases the probability of global signaling. This phenomenon also suggests a note of caution for experimental studies, where application of agonists that provoke a calcium response in glia may not be reflective of a direct link between the agonist and calcium mobilization, but rather, a change in the probability of spontaneous puffs (or puffs generated by tonic signals) evolving into global responses.

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