

“Kiss-and-Run” Glutamate Secretion in Cultured and Freshly Isolated Rat Hippocampal Astrocytes

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Under physiological conditions, astrocytes not only passively support and nourish adjacent neurons, but also actively modulate neuronal transmission by releasing “glial transmitters,” such as glutamate, ATP, and D-serine. Unlike the case for neurons, the mechanisms by which glia release transmitters are essentially unknown. Here, by using electrochemical amperometry and frequency-modulated single-vesicle imaging, we discovered that hippocampal astrocytes exhibit two modes of exocytosis of glutamate in response to various stimuli. After physiological stimulation, a glial vesicle releases a quantal content that is only 10% of that induced by nonphysiological, mechanical stimulation. The small release event arises from a brief (~2 ms) opening of the fusion pore. We conclude that, after physiological stimulation, astrocytes release glutamate via a vesicular “kiss-and-run” mechanism.

Key words: kiss-and-run; fusion pore; amperometry; single-vesicle FM imaging; glutamate; freshly isolated astrocyte

Introduction

Glial cells, including astrocytes, oligodendrocytes, and Schwann cells, have generally been considered passive components in the nervous system. Recently emerging evidence indicates that astrocytes, by releasing signaling molecules that include glutamate (Glu) and ATP, may also play active roles in various neural functions including neurogenesis, synaptogenesis, and synaptic modulation and plasticity (Carmignoto, 2000; Bezzi and Volterra, 2001; Haydon, 2001; Ullian et al., 2001; Fields and Stevens-Graham, 2002). One way in which astrocytes respond to neuronal activity is via activation of their membrane receptors, elevation of $[Ca^{2+}]_i$, and then release of glutamate to modulate synaptic plasticity. The mechanisms by which astrocytes release glutamate has been intensively studied (Carmignoto, 2000; Haydon, 2001). Recently, by using total internal reflection fluorescence (TIRF) microscopy, Bezzi et al. (2004) showed that vesicular release does occur in cultured astrocytes. Their study revealed that the vesicular glutamate release is Ca^{2+} and soluble N-ethylmaleimide-sensitive attachment protein (SNAP) receptor (SNARE) dependent. However, because of the limitations of TIRF, four major questions remain concerning the vesicular release of glial transmitters. (1) What are the detailed kinetics of the quantal transmitter release during fusion between vesicle and

plasma membrane? (2) Does the vesicle release all (“full fusion”) or only part of its content (“kiss-and-run”) during a release event? (3) What type of vesicle (310 vs 30 nm in diameter) is responsible for the Ca^{2+} -triggered release? (4) Does the Ca^{2+} -dependent vesicle release exist not only in cultured but also in freshly isolated astrocytes? Clearly, these critical questions about glial transmitter release cannot be answered using TIRF.

Several techniques allow the recording of single-vesicle activity in real time. Micro-carbon fiber electrodes (CFEs) are used to record the release of easily oxidizable monoamines (catecholamines and serotonin) from adrenal chromaffin cells, mast cells, and neurons (Wightman et al., 1991; Chow et al., 1992; Zhou and Misler, 1995a). At a holding potential of 500–800 mV, this electrochemical assay, termed amperometry, permits high temporal (in milliseconds) and spatial (in micrometers) resolution recording of the kinetics of vesicular release (Zhou et al., 1996). Application of amperometry can be extended to cells without native monoamines by preloading their vesicles with serotonin or dopamine (Zhou and Misler, 1996; Kim et al., 2000; Zhang and Zhou, 2002). Single-vesicle release can also be detected by imaging vesicles preloaded with fluorescent styryl dyes (Angleson et al., 1999; Aravanis et al., 2003). After stimulation, the preloaded dyes are discharged from single vesicles, and this can be recorded by confocal microscopy. This optical method has very high spatial resolution ($<1 \mu\text{m}$).

In the present work, we examine the kinetics of stimulus-induced vesicular release in cultured as well as freshly isolated astrocytes. We demonstrate that astrocytes release glutamate via two secretion modes from the large vesicles (310 nm in diameter) in response to different types of stimulation. Surprisingly, in response to physiological stimulation, astrocytes release their vesicular contents via a kiss-and-run mechanism.

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Materials and Methods

Cell culture and immunocytochemistry. Purified hippocampal astrocytes were obtained from postnatal day 1 (P1) Sprague Dawley rats (Shanghai SLAC Laboratory Animal Co., Shanghai, China), maintained in low-density primary culture, and used after 7–20 d in culture (Yang et al., 2003; Zhang et al., 2003). Freshly isolated astrocytes were obtained from rat (P10–P15) hippocampal slices (Zhou and Kimelberg, 2000, 2001) and were used for FM dye destaining experiments within 1 h. For immunostaining, cultures were stained with anti-GFAP antibody (Chemicon, Temecula, CA) and fluorescence-conjugated secondary antibody anti-rabbit-IgG-FITC (Jackson ImmunoResearch, West Grove, PA), as described previously (Yang et al., 2003). The specificity of the GFAP antibody was confirmed with mixed cultures of astrocytes and neurons (Yang et al., 2003). In Figure 4B, AM1-43 (Biotium, Hayward, CA), a fixable fluorescent dye, was preloaded into the astrocytes. After washout of AM1-43 and culture medium, cells were fixed and subjected to permeabilization with 0.1% Triton X-100 for 45 min at 4°C. Then, the rabbit anti-glutamate antibody (Sigma, St. Louis, MO) was added to the fixed astrocytes. Cyanine 5-conjugated anti-rabbit-IgG (Chemicon) was used to visualize the glutamate content of vesicles. Excess antibody was removed and cells were imaged using a Zeiss (Oberkochen, Germany) LSM 510 confocal microscope.

The use and care of animals in this study complied with the guidelines of the Animal Advisory Committee at the Shanghai Institutes for Biological Sciences.

Dopamine loading and electrophysiology. Purified astrocytes were loaded in a bath solution containing 70 mM dopamine (DA) for 45 min at 37°C (Zhang and Zhou, 2002). Ascorbate (1.2%) was added to protect against DA oxidation. After DA loading, the astrocytes were washed five times with standard external solution. The loading of DA had little effect on the Ca^{2+} signals in response to glutamate stimulation (Glu-stim; data not shown). Control solutions and drugs were puffed locally onto the cell during recording via a multichannel microperfusion system (Zhang and Zhou, 2002). The cell was stimulated by either physiological chemical reagents (e.g., glutamate) or mechanically. The mechanical stimulation (Mstim) was applied by poking the astrocyte for 1–2 s with a micro-patch-pipette (Araque et al., 1999). After Mstim, the cell membrane appeared to seal immediately, leaving no leak, because the elevated $[Ca^{2+}]_i$ returned to resting level in the presence of thapsigargin (data not shown). All data were collected and analyzed by a computer using IGOR software (WaveMetrics, Lake Oswego, OR). Mean \pm SEM was calculated, and the paired *t* test was applied to determine significance (***p* < 0.01).

Amperometry. Highly sensitive, low-noise, carbon fiber electrodes (ProCFE; Dagan, Minneapolis, MN) were used for electrochemical monitoring of quantal DA release from astrocytes. The amperometric current (I_{amp}) was measured at 780 mV using a patch-clamp amplifier (Zhang and Zhou, 2002) (PC-2C; INBIO, Wuhan, China). Amperometric signals were low-pass filtered at 0.3 kHz and digitized at 1 kHz. To increase the contact area between the flat glial cell and the CFE, the 5- μ m-diameter sensor tip was abraded to form a 45° angle using a beveller (EG44; Narishige, Tokyo, Japan). For the tetanus toxin (TeNT) experiment shown in Figure 5, astrocytes from the same preparation were divided into two groups (with and without TeNT), and both were incubated at 37°C for 24 h; the number of release events within 1 min were counted for statistics.

FM and $[Ca^{2+}]_i$ imaging. Astrocytes were treated with 100 μ M glutamate and 10 μ M FM 1-43 dye or 50 μ M FM 2-10 for 5 min, and then washed 10 times. FM was excited at 800 nm by a Zeiss LSM 510 two-photon upright microscope, and destaining was sampled at 1 Hz. Fura-2 was excited at 760 nm laser, which activates Ca^{2+} -sensitive fura-2 via two-photon effect (equivalent to single photon of 380 nm = 760/2 nm, with F380 as emission signal). For combined FM and Ca^{2+} recording, astrocytes were preloaded with 2 μ M fura-2 AM for 45 min before FM loading. Two-photon laser stimulation (800 nm, 0.5 mW, 25 \times 25 pixels) was used to mimic Mstim in the FM imaging experiments. The laser-induced photobleach at the stimulation site was not counted in Figure 3. In some experiments, the nucleus served as a photon-section marker to adjust the focal plane before and after laser stimulation. For intracellular

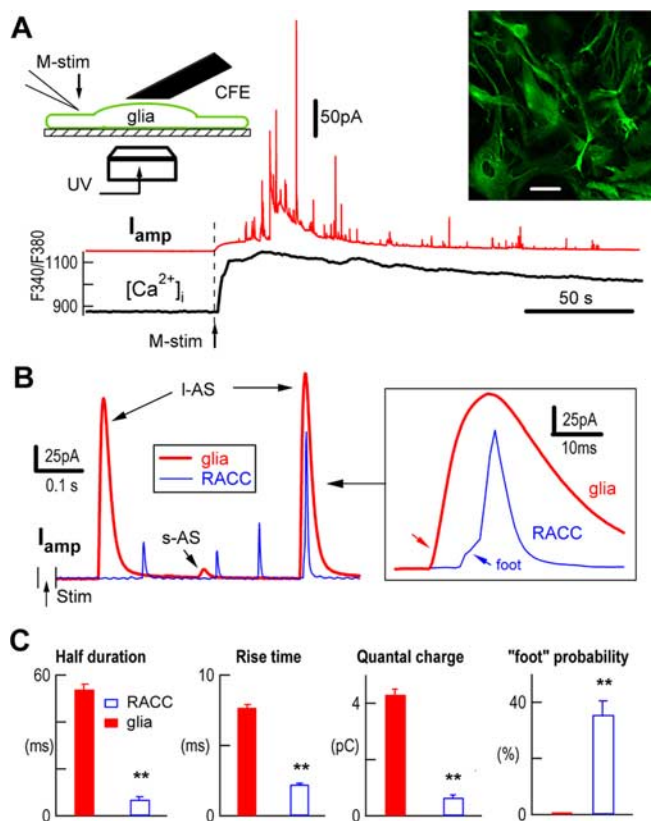


Figure 1. Stimulus-induced quantal secretion in astrocytes preloaded with dopamine. **A**, Mstim (M-stim)-induced $[Ca^{2+}]_i$ changes and amperometric spikes (I_{amp}) in astrocytes. GFAP immunostaining of astrocytes (right). Scale bar, 20 μ m. The diagram shows the experimental protocol (typical of CFE recording from >100 cells). **B**, Superimposed I_{amp} traces of glial and RACC ASs. Two classes of glial ASs are marked as s-AS and l-AS. Inset, Glial l-AS and RACC AS at expanded time scale. **C**, Statistical comparisons of half-width, rise time, quantal charge, and the probability of occurrence of a foot in the largest 10% of quantal ASs between astrocytes and RACCs. Error bars represent SEM.

Ca^{2+} imaging, cells were loaded with 2 μ M fura-2 AM and measured as described previously (Zhang and Zhou, 2002).

FM photoconversion and electron microscopy. FM photoconversion was performed as described previously (Harata et al., 2001). After FM staining, astrocytes cultured on coverslips were fixed with 2% glutaraldehyde in 100 mM sodium phosphate buffer (PB) for 20 min and washed with 100 mM glycine (in PB) for 1 h. The coverslips were washed further in 100 mM ammonium chloride (in distilled water) for 5 min. After brief rinsing in PB, coverslips were incubated in DAB (1 mg/ml in PB) for 10–20 min. Fluorescence excitation light was then continuously applied for 10–20 min in DAB solution. For electron microscopy (EM), photoconverted astrocytes were osmium postfixed (2% OsO_4 in 0.1 M PBS, pH 7.3, 4°C) for 1 h and dehydrated through an ascending series of acetone solutions. Dehydration was continued after staining and then the tissue was embedded in Epon. Thin serial sections (60–80 nm) were cut parallel to the coverslip surface and mounted on Formvar-coated slot grids. Grids were poststained with 2% aqueous uranyl acetate and Sato's lead citrate and examined on a JEM-1230 transmission electron microscope (Jeol, Tokyo, Japan).

Results

Quantal secretion in astrocytes

Hippocampal astrocytes were cultured from P1 Sprague Dawley rats (Zhang et al., 2003). As confirmed by immunostaining (Fig. 1A) and electrophysiology, astrocytes and neurons were distinguished by their morphologies. To record stimulus-induced quantal exocytosis by amperometry, astrocytes were preloaded

with DA (Zhou and Mislisler, 1996; Kim et al., 2000; Zhang and Zhou, 2002). To increase the contact area between the cell and the CFE, the tip-polished sensor surface faced the flat membrane (Fig. 1A, diagram). Mstim is a well established method to selectively stimulate astrocytes in mixed glia–neuron cultures (Araque et al., 1999). As shown in Figure 1A, Mstim induced a rapid and sustained $[Ca^{2+}]_i$ increase followed by amperometric spikes (ASs) in the glia. The slow component of the amperometric current was induced by the remote release of large amounts of dopamine, which diffused to the CFE (Zhou and Mislisler, 1995b).

To compare the stimulus-induced glial ASs with classic ASs from chromaffin cells, CFE recordings from an astrocyte (2 s Mstim) and a rat adrenal chromaffin cell (RACC) (0.2 s depolarization) were superimposed (Fig. 1B), giving similar profiles of latency distributions of AS events, except that the glial AS latency, defined as the time from the stimulus onset to an AS event, was 4.3 times longer (data not shown). Since the ASs of RACCs reflect established quantal events of single-vesicle release (Wightman et al., 1991; Chow et al., 1992; Alvarez de Toledo et al., 1993; Zhou and Mislisler, 1995a), the glial ASs should also represent quantal vesicular exocytosis. Among all ASs, vesicles immediately under the sensor surface give fast and large events, whereas vesicles distant from the sensor give slow and small events. The largest (and fastest) 10% of ASs are those induced immediately under the sensor surface (Zhou and Mislisler, 1995a; Zhou et al., 1996). By analyzing the largest 10% of ASs, the half-width, rise time, total charge, and the probability of observing a “foot” were 54 ± 3 versus 6.9 ± 0.7 ms, 7.7 ± 0.9 versus 2.2 ± 0.1 ms, 4.3 ± 0.2 versus 0.6 ± 0.1 pC, and 0.4 versus $36 \pm 6\%$ for astrocytes and RACCs, respectively [the largest 10% of ASs from nine astrocytes (460 ASs) and from seven RACCs (650 ASs)] (Fig. 1C). The kinetics of glial ASs, however, had two distinct features. First, the rise time, quantal size, and half-width of glial ASs were much greater (Fig. 1C). For example, the half-width (54 ms) was 7.8 times that in RACCs. Second, compared with neurons (Bruno and Jahn, 1995), endocrine cells (Wightman et al., 1991; Chow et al., 1992), and mast cells (Alvarez de Toledo et al., 1993), glial ASs had very few events (2 of 564) in which a foot (transmitter pre-release from flickering fusion pores) preceded an individual major spike (Chow et al., 1992; Zhou et al., 1996), indicating that the astrocyte fusion pores are more rigid after opening (Fig. 1B,C). This distinct feature in glia is, however, fully consistent with the following reports: (1) instead of the fusion protein SNAP-25, astrocytes express SNAP-23 (Hepp et al., 1999); and (2) compared with SNAP-25, SNAP-23-mediated Ca^{2+} -dependent ASs have reduced “feet” (Sorensen et al., 2003).

As indicated in Figure 1B, Mstim induced two classes of ASs from astrocytes: large (l-ASs) and small (s-ASs). Compared with the two l-ASs in Figure 1B, the s-AS has a 10-fold smaller current integral (or quantal size, corresponding to the number of molecules released from the vesicle), which may represent another class of vesicle release (see below).

Partial vesicle release induced by physiological stimuli in cultured astrocytes

Glutamate is an endogenous transmitter that activates metabotropic glutamate receptors to release Ca^{2+} from intracellular stores (data not shown). Local application of 0.5 mM glutamate (Schneggenburger et al., 1993; Zhou and Kimelberg, 2001) induced a burst of s-ASs in DA-loaded astrocytes (Fig. 2A). The s-ASs induced by glutamate (Fig. 2C) were 10-fold smaller than the l-ASs induced by Mstim (Fig. 2E). We then examined signal transmission between two astrocytes by stimulating glia 1 and

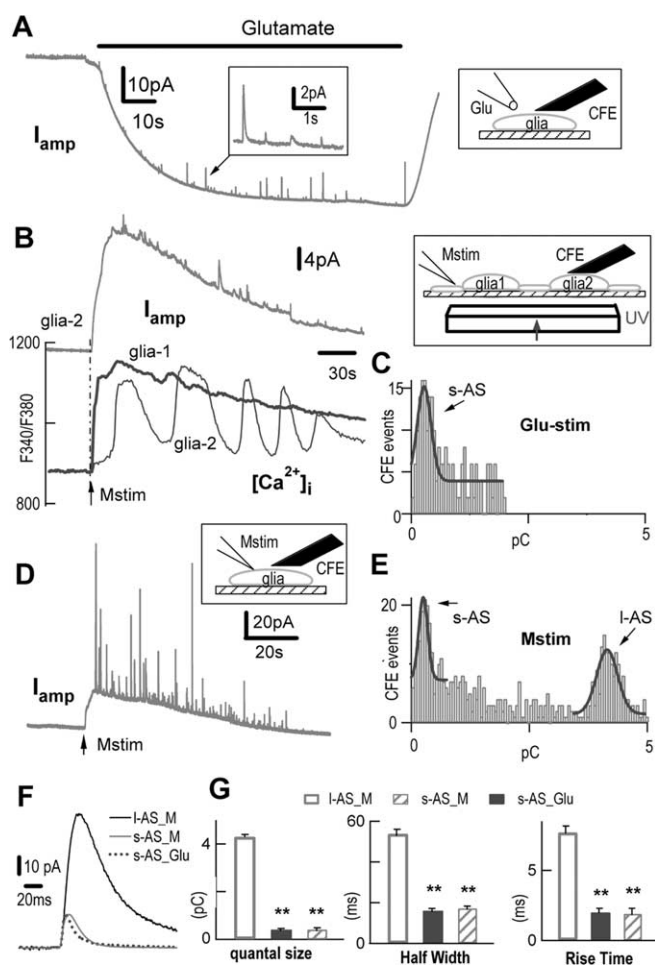


Figure 2. Evidence for kiss-and-run release from amperometry in astrocytes under physiological stimulation. **A**, Glutamate (0.5 mM)-induced amperometric spikes in an astrocyte ($n = 11$). The slowly descending background current was an artifact caused by puffing glutamate (Zhou and Mislisler, 1995a). Inset, An AS burst. **B**, Stimulating glia 1 induced a $[Ca^{2+}]_i$ increase in glia 1 (bottom trace, thick line) and caused $[Ca^{2+}]_i$ oscillations (bottom trace, thin line) and ASs (top trace) in glia 2 ($n = 8$). **C**, Quantal size distribution of Glu-stim-induced ASs and their Gaussian fit (214 ASs from 11 cells). **D**, Mstim-induced ASs. **E**, Quantal size distribution of Mstim-induced ASs and their Gaussian fit (726 ASs from 18 cells). The two peaks represent the two AS classes, s-AS and l-AS (see also Fig. 2A). **F**, Comparison of s-AS and l-AS induced by Mstim (s-AS_M and l-AS_M), and s-AS induced by Glu-stim (s-AS_Glu) at an expanded time scale. **G**, Statistics. Corresponding to Mstim-induced l-ASs, Mstim-induced s-ASs, and Glu-stim-induced s-ASs, the quantal sizes were 4.3 ± 0.1 , 0.4 ± 0.1 , and 0.4 ± 0.1 pC, the half-widths were 54 ± 3 , 17 ± 1 , and 16 ± 1 ms, and the rise times were 7.7 ± 0.5 , 1.9 ± 0.4 , and 2.0 ± 0.3 ms, respectively. Diagrams in **A**, **B**, and **D** illustrate the setup for each experiment. Error bars represent SEM.

observing the responses in adjacent glia 2 (Fig. 2B, diagram). Mstim induced a sustained $[Ca^{2+}]_i$ elevation in glia 1, followed by ASs and Ca^{2+} waves in glia 2 (Fig. 2B). This result in cultured astrocytes is consistent with the calcium wave seen in astrocytes in brain slices (Parri et al., 2001). Figure 2, C and E, shows the quantal size distributions of Glu-stim- and Mstim-induced ASs, respectively. The histogram of Glu-stim-induced ASs has a single peak at 0.4 pC. However, the histogram of Mstim-induced ASs gives two distinct peaks at 0.4 and 4.3 pC, corresponding to the s-ASs and l-ASs shown in Figure 1B. Interestingly, the ASs induced by Glu-stim and the s-ASs induced by Mstim are virtually identical in both quantal size and kinetics (Fig. 2F,G), indicating that they belong to the same class of vesicle release. The fact that two classes of ASs were induced by Mstim, but not Glu-stim, suggests that l-ASs need the sustained high $[Ca^{2+}]_i$ that is in-

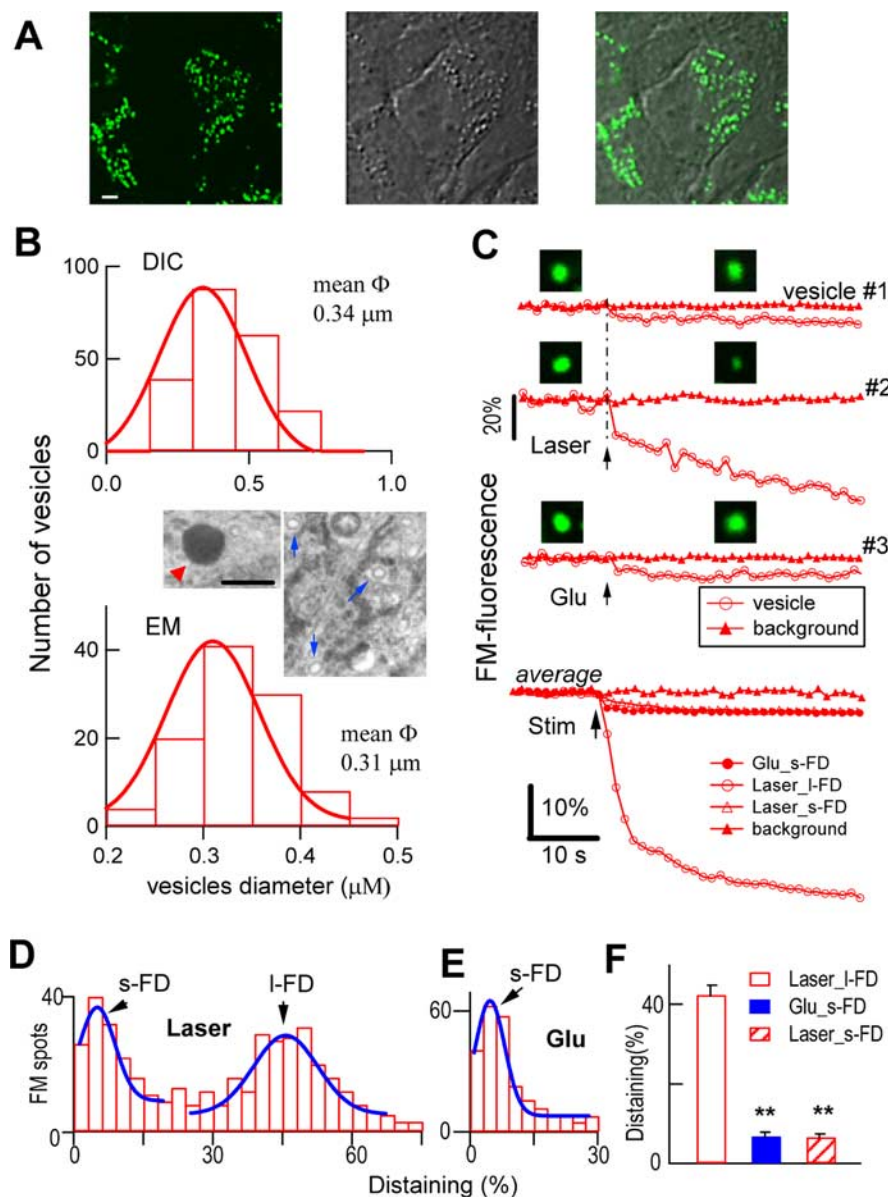


Figure 3. Evidence for kiss-and-run release from FM imaging in cultured astrocytes under physiological stimulation. **A**, Confocal images. Left, FM fluorescence; middle, DIC; right, merged. Scale bar, $2 \mu\text{m}$. **B**, Diameter distribution and Gaussian fit curves of FM-loaded vesicles. Spot diameters under DIC (top) and EM (bottom). Insets, Glial vesicles stained by FM and photoconverted for EMs from an astrocyte. Scale bar, 300 nm . The red arrow indicates a photoconverted vesicle (left); the blue arrows indicate small vesicles (right). **C**, Time course of FM destaining from three representative single vesicles induced by laser (vesicle #1 for s-FD, #2 for l-FD), or $100 \mu\text{M}$ glutamate (vesicle #3, s-FD). The vesicle images before and after the FM discharge are placed near the imaging time on the traces. Fluorescence signals of FM discharge from single vesicles and nearby background are marked as circles and triangles, respectively. The bottom panel shows average traces of 393 vesicles from 14 cells with laser stimulation and 243 vesicles from 11 cells with glutamate stimulation. **D**, FM destaining distribution of vesicles induced by laser and their Gaussian fits (393 vesicles from 14 cells). The two peaks represent the two FD classes, s-FD and l-FD. **E**, FM destaining distribution of Glu-stim-induced FD and their Gaussian fit (243 vesicles from 11 cells). **F**, Histogram of percentage of FM destaining in vesicles similar to those in **C** at 25 s after the onset of stimulation. Error bars represent SEM.

duced by the nonphysiological Mstim. In contrast, in addition to quantal size, the rise time and half-width of s-ASs were also three to four times smaller than those of l-ASs (Fig. 2G). The three recordings of representative s-ASs and l-ASs show that the two s-ASs have similar kinetics; however, the decay of the l-AS is very different from the decay of the s-ASs. Note that the rising phase of the l-AS and the s-AS are nearly identical, implying that the DA release was from two vesicles with similar kinetics of fusion to the plasma membrane (Fig. 2F).

The two AS classes induced by Mstim may come from either two kinds of vesicles with different content sizes or from a single type of vesicle releasing its contents under two modes, full fusion and kiss-and-run, as for neuronal vesicles (Zhou et al., 1996; Aravanis et al., 2003; Staal et al., 2004). To determine whether kiss-and-run release is responsible for the s-ASs, single vesicles were visualized by two-photon imaging in astrocytes preloaded with the fluorescent dye FM 1-43 (Angleton et al., 1999; Aravanis et al., 2003). Because cultured astrocytes have a flat morphology, individual vesicles were clearly visible in both fluorescence and differential interference contrast (DIC) transmission images (Fig. 3A). Surprisingly, most DIC-visible vesicles were FM loaded, indicating that most of the vesicles experienced endocytosis/FM loading during the preloading process. The distribution of vesicle diameters gave a single peak as measured by DIC and EM (photoconverted FM-loaded vesicle) images, indicating that FM dye was loaded into only one type of vesicle. The average vesicle diameters measured from DIC and EM were 340 ± 9 and $310 \pm 22 \text{ nm}$, respectively (Fig. 3B,C). The diameter distributions of FM 1-43 spots had similar single peak as well, except that the peak was at 810 nm (data not shown). According to calibration of fluorescence beads, the apparent larger (2.4-fold) size of FM spots was attributable to the fluorescence diffraction effect (data not shown). This confirmed that each FM-loaded fluorescent spot represented a single vesicle. To minimize focus errors of FM imaging caused by mechanical disturbance, Mstim was replaced by applying a brief laser beam (800 nm , 25 ms , 0.5 mW) of $0.25 \times 0.25 \mu\text{m}^2$ to the astrocyte. Similar to nonphysiological Mstim, the laser also induced a sustained $[\text{Ca}^{2+}]_i$ increase. In contrast, glutamate (0.1 mM) application was used as the Glu-stim. The time courses of release from two representative vesicles, which showed laser-induced small and large FM fluorescence destaining (s-FD and l-FD, respectively), and from a typical glutamate-induced FM destaining, are illustrated in Figure 3C. The distribution of laser-induced

single-vesicle destaining contained two distinct peaks at 7% (s-FD) and 43% (l-FD) (Fig. 3D). In contrast, the distribution of glutamate-induced vesicle destaining had only a single peak at 7% (Fig. 3E), which matched the laser-induced s-FD. These FM imaging data are entirely consistent with the amperometric results, with s-FD and l-FD corresponding to s-AS and l-AS. These data strongly suggest that FM-loaded vesicles are the same ones loaded by dopamine shown in Figures 1–3.

We also examined the effects of two subsequent stimulations by glutamate and laser on the FM 2-10 destaining in the same vesicle (Fig. 4A, left). FM 2-10 has a faster destaining rate than that of FM 1-43 ($\tau_{\text{off}} = 0.6$ vs 2.5 s) (Ryan et al., 1996; Klingauf et al., 1998). Statistically, a single vesicle destained $5.4 \pm 1.2\%$ in response to the first stimulation (glutamate for 10 s), and the same vesicle destained $45 \pm 3\%$ in response to the second stimulation (Mstim-like laser) (Fig. 4A, right). This experiment provides direct and compelling evidence that (1) a single vesicle releases only 5% of its total FM content after physiological stimulation; and (2) the same vesicle can release nine times as much after very strong stimulation. Note that the onset of FM destaining took 2 s, which was considerably longer than the half duration (60 ms) of a l-AS detected by CFE, probably because of the slow dissociation constant of the FM dye (0.6 s for FM 2-10) (Klingauf et al., 1998).

Glutamate is the glial transmitter

Subsequently, we showed that glutamate-containing vesicles are the same vesicles as those loaded with FM 1-43, illustrated in Figures 3 and 4. We double-labeled the vesicles using a fixable FM 1-43 analog, AM1-43 (Renger et al., 2001), and anti-glutamate antibody (Carder and Hendry, 1994). As shown in Figure 4B, 87% of AM1-43-labeled vesicles were glutamate positive. On average, the anti-glutamate antibody and AM1-43 colocalization was $66 \pm 7\%$ ($n = 8$). Thus, the releasable vesicles underlying the different kinds of stimulation were glutamate-releasing vesicles.

Glial quantal release is Ca^{2+} and SNARE dependent

Combined CFE and $[\text{Ca}^{2+}]_i$ measurements were applied in single astrocytes to determine the Ca^{2+} dependence during a Mstim-induced sustained $[\text{Ca}^{2+}]_i$ elevation and ASs. The frequency of ASs and $[\text{Ca}^{2+}]_i$ were reduced by 77 ± 7 and $63 \pm 8\%$ when external Ca^{2+} was changed from 2 mM to 0 ($n = 7$) (Fig. 5A). The same Ca^{2+} dependence was observed with chemical stimulation, when the Ca^{2+} ionophore ionomycin (10 μM) was puffed onto the astrocytes (Fig. 5B). Finally, combined measurements of FM and fura-2 confirmed that laser-induced FM destaining from glial vesicles (revealed by FM) was Ca^{2+} dependent (revealed by F380) (Fig. 5C). In 2 mM $[\text{Ca}^{2+}]_o$, the stimulation reduced F380 and FM by $56 \pm 4\%$ (108 images from eight cells) and $42 \pm 3\%$ (308 vesicle images from 12 cells), respectively. In contrast, the laser caused little change in F380 and FM in 0 $[\text{Ca}^{2+}]_o$. These data demonstrate that stimulus-induced quantal secretion depends on $[\text{Ca}^{2+}]_i$ in astrocytes.

Fusion proteins are required for Ca^{2+} -dependent exocytosis in neurons and other cells (Jahn and Sudhof, 1999). To test whether the fusion protein complex SNARE is involved in quantal secretion in astrocytes, TeNT, a specific toxin against SNARE (Xu et al., 1998), was introduced into astrocytes and RACCs (positive control). In astrocytes, TeNT significantly inhibited Mstim-induced ASs by $44 \pm 6\%$ (71 cells for control, 40 cells for TeNT; $p < 0.01$) (Fig. 5D) (Pasti et al., 2001). In RACCs, $79 \pm 3\%$ of secretion was inhibited by TeNT (79 cells for control, 58 cells for

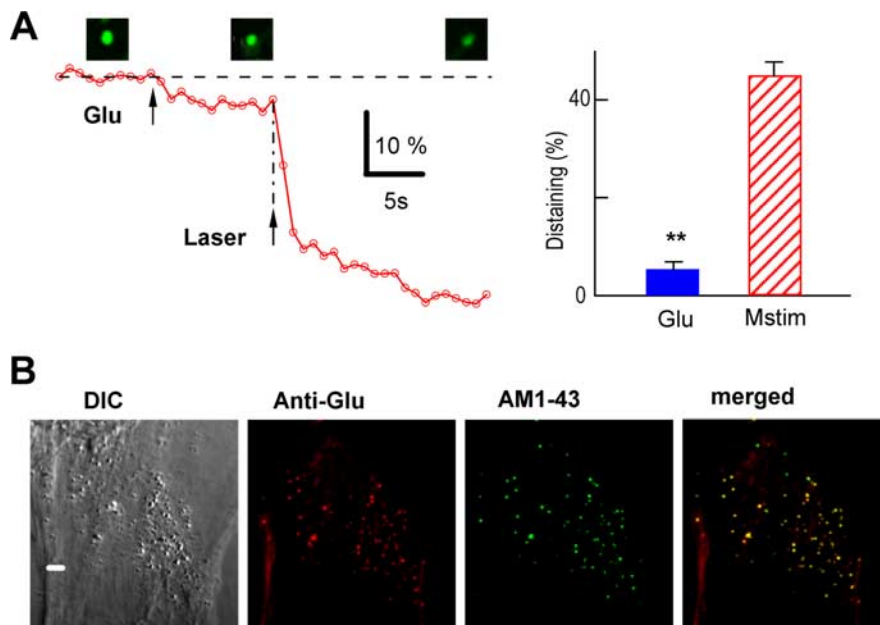


Figure 4. Kiss-and-run glutamate release from single vesicles. **A**, Averaged FM destaining recordings from single vesicles induced by two subsequent stimulations with glutamate (0.1 mM for 10 s) and laser (32 vesicles from 4 cells). Histogram of the glutamate- and laser-induced FM discharges (right, 10 s after the onset of stimulation). Error bars represent SEM. **B**, In these images, the colocalization of anti-glutamate antibody and AM1-43-labeled vesicles was 87%. The average colocalization of anti-glutamate antibody and AM1-43 was $66 \pm 7\%$, ranging from 45 to 92% ($n = 8$). Scale bar, 2 μm .

TeNT; $p < 0.01$) (Xu et al., 1998). The different efficiencies of tetanus toxin in suppressing secretion in chromaffin cells and astrocytes may come from the different combinations of fusion proteins in the two kinds of cell (Sorensen et al., 2003; Zhang et al., 2004). These data indicate that SNARE is involved in quantal secretion from astrocytes.

Partial vesicle release in freshly isolated astrocytes

Thus far, the results are consistent with hypothesis that release of glutamate from cultured astrocytes was by the kiss-and-run mode. However, long-term culture (2–3 weeks) may cause the artificial expression of some proteins, such as Ca^{2+} -permeable AMPA receptors (Janssens and Lesage, 2001). To determine whether the glutamate quantal release is an artifact of culture, we examined the laser-stimulated FM destaining in freshly isolated astrocytes (FIAs) (Zhou and Kimelberg, 2000, 2001). FIAs were identified by a soma with an oval, elongated, or triangular shape and with multiple, long, bushy processes extending from the cell body (Fig. 6A, right). After the standard FM destaining experiments, we also retrospectively stained with GFAP to confirm the identity of each cell as an astrocyte (Fig. 6A, middle). Under DIC imaging, vesicles in the processes of FIA could be clearly identified and had similar diameters to those in cultured astrocytes (Fig. 6B). The diameter distributions of FM 1-43 spots had a similar single peak as well, except that the peak was at 780 nm (data not shown). (The apparent larger size of FM spots is attributable to the fluorescence dispersion effect.) By applying the laser stimulation protocol of Figure 3 to FIAs, the FM-loaded vesicles showed an FM destaining distribution with two populations ($6.2 \pm 1.5\%$ for s-FD and $45.5 \pm 6.3\%$ for l-FD) (Fig. 6C–E), and the ratio of s-FD to l-FD ($\sim 1:7.5$) was also similar to that in cultured astrocytes (Fig. 3). Thus, the partial vesicular release also occurs in FIAs.

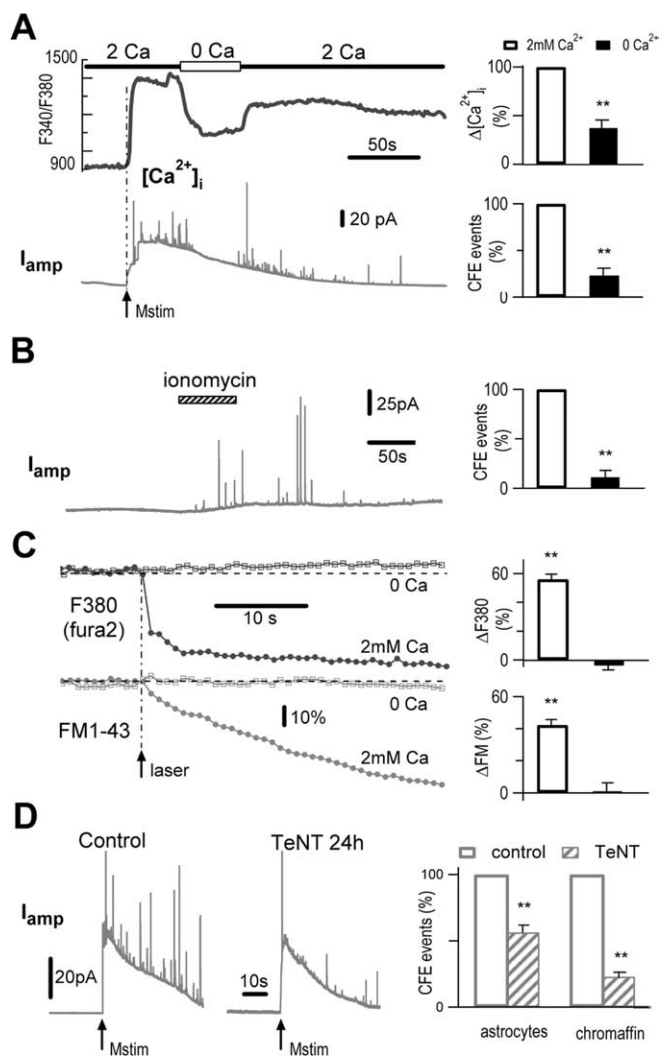


Figure 5. Stimulus-induced ASs are dependent on Ca^{2+} and SNARE. **A**, Ca^{2+} dependence of amperometric spikes. Left, Mstim-induced $[Ca^{2+}]_i$ and ASs (I_{amp}) at 2 mM versus 0 $[Ca^{2+}]_o$. Right, Statistics. **B**, Ionomycin-induced ASs in 2 mM $[Ca^{2+}]_o$ (left). ASs were reduced by $89 \pm 8\%$ in 0 $[Ca^{2+}]_o$ (right; $p < 0.01$; $n = 5$). **C**, Laser-induced destaining of FM 1-43 was Ca^{2+} dependent. Left, Ca^{2+} -sensitive F380 and FM 1-43 at 2 mM (filled symbols) versus 0 $[Ca^{2+}]_o$ (open symbols). Right, Statistics. **D**, ASs were inhibited by tetanus toxin (control vs TeNT). Right, Statistics. All experiments were from three batches of astrocytes and three batches of RACCs. Error bars represent SEM.

Discussion

Kiss-and-run as the release mechanism in astrocytes

The most important finding in the present work is that a glial vesicle releases only part ($\sim 10\%$) of its total vesicle content during a release event triggered by physiological stimulation. The partial release is termed as kiss-and-run. This conclusion is supported by the following experimental evidence: (1) In quantal content distribution, Mstim induced two populations of amperometric spikes, the s-ASs and l-ASs, with quantal release contents of 1:10 (Fig. 2). (2) Glu-stim-induced ASs comprised only one population, which was equivalent to the Mstim-induced s-AS (Fig. 2). (3) There were two populations of Mstim-induced FM destaining from single vesicles, s-FD and l-FD, with quantal release contents of 1:9 (Fig. 3). (4) Glu-stim-induced FDs comprised only one population, which was equivalent to the Mstim-induced s-FDs (Fig. 3). (5) Single-vesicle FM imaging revealed that, although initial Glu-stim induced a s-FD, a subsequent

Mstim-like laser pulse induced a l-FD from the same vesicle (Fig. 4A). (6) The distribution of vesicle sizes (both by DIC and EM imaging) gave only a single vesicle population (Fig. 3B), excluding the possibility that s-AS and l-AS were from small and large vesicles, respectively. (7) The distributions of Mstim- and Glu-stim-induced release of quantal contents were virtually same for ASs and FDs, indicating that AS and FD were from the same kind of vesicle (Figs. 2, 3). (8) Glutamate was colocalized with FM dyes, indicating that the FM release and glutamate release were from the same kind of vesicle (Fig. 4B). (9) Two populations of Mstim-induced FM destaining from single vesicles were also found in freshly isolated astrocytes (Fig. 6). Furthermore, although the rising phases of the l-AS and the s-AS were nearly identical, the s-AS decayed twice as fast as the l-AS (Fig. 2F). Following the kiss-and-run hypothesis, we propose that the rapid decay in s-ASs was caused by the fusion pore closing before release of the major content. In other words, the rise time of the s-AS was equal to the open time of the fusion pore (2.0 ± 0.3 ms), whereas the slower decay of the l-AS was attributable to vesicle release during a full (or at least more complete) fusion event.

The advantage of kiss-and-run release is that a single vesicle can release transmitter multiple times after multiple stimulations, providing an efficient means of exocytosis with a limited number of release vesicles. This mechanism has been found in endocrine cells (Zhou et al., 1996) and neurons (Aravanis et al., 2003; Staal et al., 2004). However, because the glial vesicle size (310 nm) is much larger than that in RACCs (125 nm; data not shown) and synapses (50–100 nm), the fractional release during each fusion is much smaller in astrocytes ($< 5\%$ of total FM dye) (Fig. 4) than that in chromaffin cells [$> 80\%$ of total transmitter (Zhou et al., 1996)] and synapses [30% of total transmitter for DA neurons (Staal et al., 2004); 47% of total FM dye for hippocampal neurons (Aravanis et al., 2003)], permitting more rounds of release in a short time before refilling of the glial vesicles. Assuming that the vesicle releases 10% of its dopamine contents during each round (Fig. 2), after 10 rounds, 35% of the original dopamine content still remains. According to diffusion theory (Pusch and Neher, 1988), the release rate is dependent on molecular weight (release rate, $k \times M^{-1/3}$, where M is the molecular weight of the released substance and k is a constant). Given the similar molecular weights of dopamine (170 Da) and glutamate (147 Da), if both molecules are free to diffuse from the vesicles, the 10% dopamine release per exocytotic event should also apply to glutamate in astrocytes.

Our findings suggest that two distinct exocytotic modes correspond to physiological stimulation and nonphysiological stimulation. Kiss-and-run is the exocytotic mode for physiological stimulation, whereas strong stimulation induced by Mstim or laser causes the exocytotic mode of permanent vesicle fusion, during which most/all vesicle contents are released. The dependence of vesicle fusion mode on stimulus/ Ca^{2+} has been reported previously by others and by us in chromaffin cells (Zhou et al., 1996; Ales et al., 1999). In addition, different isoforms of synaptotagmin have been reported as underlying different modes of vesicle fusion. Synaptotagmin I is for full fusion, and synaptotagmin IV is for kiss-and-run (Wang et al., 2003). Recently, synaptotagmin IV was found to be expressed exclusively in astrocytes for glutamate release (Zhang et al., 2004). Future work should address which fusion protein is responsible for kiss-and-run fusion in astrocytes.

Single flicker fusion pore

In chromaffin cells and DA neurons, in which kiss-and-run is characterized by transmitter release during rapid fusion-pore flickering, the fusion pore switches on and off multiple times within a second (Zhou et al., 1996; Ales et al., 1999; Staal et al., 2004). Surprisingly, in astrocytes, although only a small portion of the total content was released after physiological stimulation, there was little detectable fusion pore flickering (very few foot events). As revealed by direct measurement with FM imaging, Glu-stim triggered only a partial release of the vesicle content (Fig. 4A). Interestingly, few feet or repeated fusion pore flickers occurred during each release event in astrocytes. Thus, the kiss-and-run in astrocytes is distinct from that in neuronal cells: it is a “single kiss” or “single flicker.” The molecular mechanism of “single-kiss-and-run” in astrocytes is unclear, but it may be related to distinct fusion proteins including SNAP-23, which differs from the SNAP-25 in neurons (Parpura et al., 1995; Hepp et al., 1999; Sorensen et al., 2003).

What type of vesicle is responsible for the glutamate release?

Comparing the properties of quantal vesicle release in astrocytes and chromaffin cells, the two cell types differ in their kinetics and fusion protein subunits. Consistent with the large size of glial vesicles (310 nm in diameter), the quantal event in glia is much larger and wider than that in neurons and endocrine cells but similar to that in mast cells. This result differed from those of Bezzi et al. (2004), who identified the vesicular glutamate transporter (VGLUT)-positive vesicle with small size (~30 nm in diameter) in brain slices. However, it is not clear whether the exocytotic vesicles that were evoked were these small vesicles. It seems, rather, that their exocytotic vesicles were large (~700 nm in diameter for their GFP fluorescent spots). This spot size is far larger than the 30 nm vesicles seen in the slice, but similar to the FM fluorescent spots in our cultured and freshly isolated astrocytes. In contrast, our single-vesicle FM imaging showed that all Ca^{2+} -triggered exocytosis was from large (310 nm) vesicles. Thus, we propose that large vesicles (310 nm diameter), but not small vesicles (30 nm), are responsible for the Ca^{2+} -dependent vesicle release in cultured and freshly isolated astrocytes.

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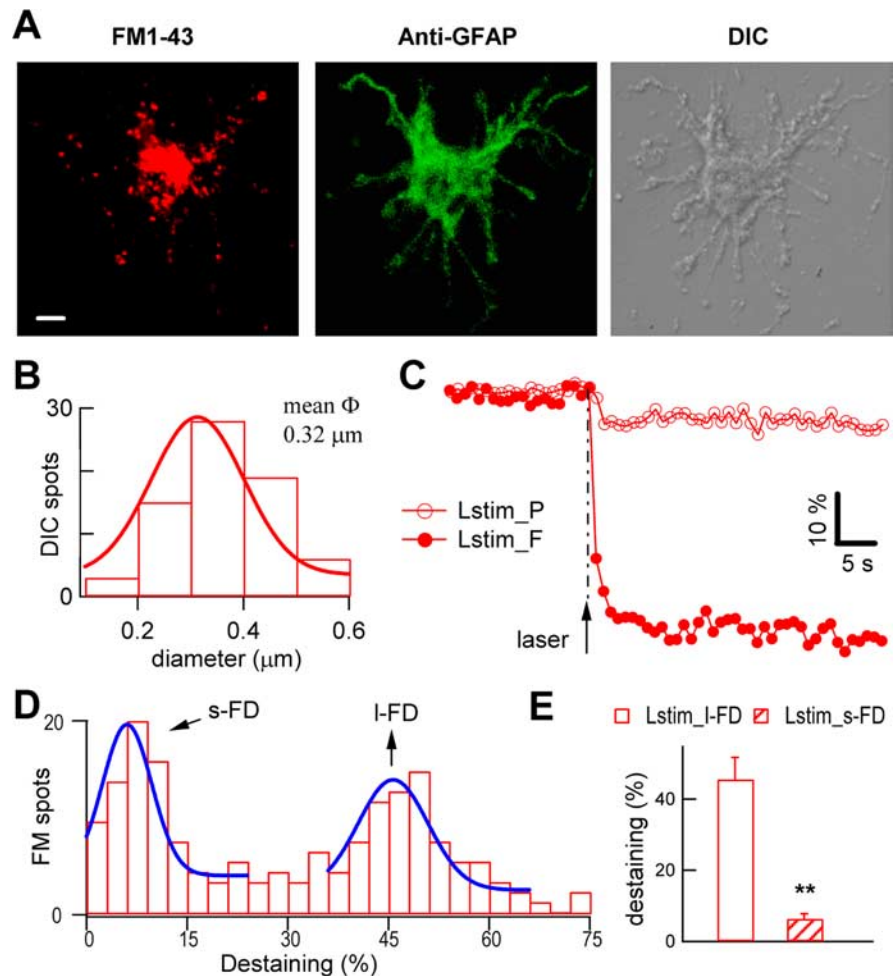


Figure 6. FM imaging of kiss-and-run vesicle release in freshly isolated astrocytes. **A**, A typical FIA for single-vesicle FM imaging. Left, The FIA loaded with FM; middle, the FIA examined by GFAP immunostaining after FM destaining; right, DIC image. Scale bar, 5 μ m. Without the laser stimulation, the GFAP immunoreactivity was distributed in distinct fibers (data not shown). **B**, Diameter distribution and Gaussian fit curve of DIC-resolved vesicles. **C**, Average FM traces of 86 large (I-FD) and 83 small (s-FD) vesicles from 11 cells with laser stimulation. Lstim_P and Lstim_F indicate laser-induced partial and full release, respectively. **D**, FM destaining distribution of vesicles induced by laser and their Gaussian fits (214 vesicles from 11 cells). The two peaks represent the two FD classes, s-FD and I-FD. **E**, The percentage destaining of s-FD and I-FD were 6 ± 2 and $46 \pm 6\%$, respectively. Error bars represent SEM.

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