Gray Matter NG2 Cells Display Multiple Ca\textsuperscript{2+}-Signaling Pathways and Highly Motile Processes

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Abstract

NG2 cells, the fourth type of glia in the mammalian CNS, receive synaptic input from neurons. The function of this innervation is unknown yet. Postsynaptic changes in intracellular Ca\textsuperscript{2+}-concentration ([Ca\textsuperscript{2+}]\textsubscript{i}) might be a possible consequence. We employed transgenic mice with fluorescently labeled NG2 cells to address this issue. To identify Ca\textsuperscript{2+}-signaling pathways we combined patch-clamp recordings, Ca\textsuperscript{2+}-imaging, mRNA-transcript analysis and focal pressure-application of various substances to identified NG2-cells in acute hippocampal slices. We show that activation of voltage-gated Ca\textsuperscript{2+}-channels, Ca\textsuperscript{2+}-permeable AMPA-receptors, and group I metabotropic glutamate-receptors provoke [Ca\textsuperscript{2+}]\textsubscript{i}-elevations in NG2 cells. The Ca\textsuperscript{2+}-influx is amplified by Ca\textsuperscript{2+}-induced Ca\textsuperscript{2+}-release. Minimal electrical stimulation of presynaptic neurons caused postsynaptic currents but no somatic [Ca\textsuperscript{2+}]\textsubscript{i} elevations, suggesting that [Ca\textsuperscript{2+}]\textsubscript{i} elevations in NG2 cells might be restricted to their processes. Local Ca\textsuperscript{2+}-signaling might provoke transmitter release or changes in cell motility. To identify structural prerequisites for such a scenario, we used electron microscopy, immunostaining, mRNA-transcript analysis, and time lapse imaging. We found that NG2 cells form symmetric and asymmetric synapses with presynaptic neurons and show immunoreactivity for vesicular glutamate transporter 1. The processes are actin-based, contain ezrin but not glial filaments, microtubules or endoplasmic reticulum. Furthermore, we demonstrate that NG2 cell processes in situ are highly motile. Our findings demonstrate that gray matter NG2 cells are endowed with the cellular machinery for two-way communication with neighboring cells.

Introduction

In addition to astrocytes, oligodendrocytes, and microglia, NG2 cells are now recognized as a fourth glial cell type in the CNS [1,2]. NG2 cells display long narrow processes and lack gap junction coupling. Fate mapping analysis has demonstrated that in white matter the majority of NG2 cells are oligodendrocyte precursors (OPCs). In contrast, gray matter NG2 glia only rarely give rise to oligodendrocytes or astrocytes but keep their phenotype throughout postnatal life [3], but see also [4,5].

NG2 cells are unique among glial cells in receiving synaptic input [reviewed by [2,6]], but the physiological impact of this innervation is unknown. Specifically, it remains unclear whether pre-synaptic transmitter release generates Ca\textsuperscript{2+}-elevations in post-synaptic NG2 cells, which might evoke cellular motility or release of neuroactive substances. This ignorance is quite astonishing in view of the increasing knowledge of glia-mediated modulation of CNS signaling, such as astrocyte-neuron interactions which gave rise to the tripartite synapse concept [7–9]. Moreover, it is known for more than a decade that 'complex' glial cells [10], which display properties similar to NG2 cells, express Ca\textsuperscript{2+}-permeable AMPA receptors [11–13] and voltage-gated Ca\textsuperscript{2+}-channels (Ca\textsubscript{s},Ca\textsubscript{v}) [14]. In cultured presumed glial progenitor cells, Ca\textsubscript{s} are activated by the depolarizing action of GABA [15]. However, despite these previous reports the presence of Ca\textsubscript{s} in NG2 glia is still disputed. Instead, a role for the Na\textsuperscript{+}-Ca\textsuperscript{2+} exchanger (NCX) in NG2 cell Ca\textsuperscript{2+}-signaling has recently been proposed [16,17].

There are different terms in the literature describing NG2-like cells in acute preparations of wild type or different transgenic mouse lines: complex glial cells (e.g. [10]); GluR cells (e.g. [18]), OPCs (e.g. [19]), synantocytes [20], and polydendrocytes (e.g. [21]). It is currently unknown to which degree these cellular populations overlap [6]. In the present study, we employed transgenic mice with fluorescence labeling of NG2 and GluR cells to study their process structure and Ca\textsuperscript{2+}-signaling mechanisms. Morphological, molecular and functional analyses revealed that NG2 cells (i) generate transient elevations of the intracellular Ca\textsuperscript{2+}-concentration ([Ca\textsuperscript{2+}]\textsubscript{i}) upon different types of stimulation and (ii) display \textit{in situ} highly motile actin-based processes.
Results

Cell identification and basic electrophysiological properties

Cell identification in the hippocampus was based on EYFP or EGFP fluorescence, morphology, and physiological criteria as reported previously [18,22,23]. Cells used for Ca²⁺-imaging (n = 636; 691 of them genotyped) were EYFP positive, had an input resistance of 193±157 MΩ, a resting membrane potential of −77±6 mV, and a membrane capacity of 33±6 pF (K⁺-based pipette solution). All cells tested (n = 23) received glutamatergic and/or GABAergic synaptic input (not shown). EYFP positive cells from homozygous (n = 351) and heterozygous (n = 340) mice did not differ with respect to the above membrane parameters, expression of Cav channel transcripts, and Ca²⁺-responsiveness upon somatic depolarization or high frequency stimulation of presynaptic fibers (see below for details). Therefore, data were pooled.

Ultrastructure of neuron-NG2 cell synapses in the hippocampus

Applying correlated light and electron microscopy, we investigated synapses onto NG2 cells in the CA1 region. The typical current pattern and light microscopic morphology of the filled cells analyzed ultrastructurally (n = 3) are shown in Figs. 1A, B. Axon terminals form synapses with processes of all three NG2 cells (Fig. 1D, E). This confirms earlier findings demonstrating synapses on processes of NG2 cells in the hippocampus [6,23–25]. However, only 3, 6, and 8 synapses, respectively, were found on the three cells analyzed, (Table 1), although all serial sections from a given biocytin filled NG2 cell were examined over its full process extent. The total

Figure 1. Neuron-NG2 cell synapses in mouse hippocampus. (A) Whole-cell current pattern (de- and hyperpolarization between −160 and +20 mV; 10 mV increments, holding potential −70 mV). (B) The morphology of the cell in (A) is still visible after biocytin-filling, signal conversion to DAB, and araldite-embedding for EM. Note the oval soma (asterisk) and varicose, branched processes. (C–F) Ultrastructural details of the same cell. Typical features of neuron-neuron synapses, viz. pre-synaptic vesicles, synaptic cleft (arrows) and post-synaptic density are also displayed by neuron-NG2 cell synapses, which are identified by dark DAB reaction product. Enlargements from consecutive sections of the boxed areas in (D, E, F) are shown. The synapses in (C, D₁, D₂, E₁) are asymmetric, whereas that in (F₁) is symmetric. Several pre-synaptic vesicles are docked (arrowheads in D₁, D₂, E₁). Note that the diameter of post-synaptic NG2 cell processes can be very small (approx. 200 nm in C and E) or >1 μm (D). Scale bars, 5 μm (B), 200 nm (all others).

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number of synapses on the three cells was estimated to be 30 (as
described above; Table 1). These synapses were very similar in
structure to neuron-neuron synapses, displaying pre-synaptic
vesicles, post-synaptic density and cleft material (Figs. 1C, D1, E1).
In several axon terminals, docked vesicles were observed at the pre-
synaptic membrane (Figs. 1D1, E1). In some cases, the DAB
reaction product was faint enough to reveal distinct post-synaptic
detail, which was indistinguishable from neuron-neuron synapses.
Thus, several neuron-NG2 cell synapses could be unequivocally
classified as either asymmetric (7/17) or symmetric (1/17) (see
Table 1, Figs. 1D–F). All synapses were on the processes of NG2
cells, none on the soma. The post-synaptic NG2 cell process was
described in some detail, which was indistinguishable from classical
synapses between neurons.

The physiological properties of these neuron-NG2 cell-synapses are characterized in some detail [6]. So far, however, it is largely
unclear whether neuronal innervation initiates Ca2+-signaling in
post-synaptic NG2 cells. Therefore, we tested for potential pathways provoking [Ca2++]i, elevation in NG2 cells, which might
be activated by the synaptic input.

NG2 cells express functional voltage-gated Ca2+-channels

Previous work has demonstrated that complex glial cells in wild
type mice express different types of Ca2+ channels, as described above [14], although later on in its
presence in NG2 cells has been disputed [16,17]. To investigate this issue in NG2/EYFP positive cells, putative Ca2+ currents were
isolated using Na+- and K+-free bath and pipette solutions. In addition, solutions were supplemented with Na+, K+, channel blockers, and [Ca2++]i in the bath was increased to 5 mM (see Materials and Methods and [14]). To remove steady-state
inactivation from putative Ca2+ channels, conditioning pre-pulses to −110 mV and −10 mV were applied for 1.5 s, respectively.
Afterwards, current families were subtracted at corresponding
membrane potentials. This procedure isolated transient membrane
currents in NG2 cells (peak amplitudes 100±30 pA at −20 mV, n = 14) (Fig. 2B). Plotting the I/V relationship of the evoked currents revealed a threshold potential of −60 mV, while peak inward currents occurred at about −20 mV (Fig. 2B). The L-type
channel blocker Verapamil (100 μM) reduced the maximum
inward currents from 167±35 pA to 65±33 pA (n = 9, Fig. 2C) and significantly shifted the half-maximum voltage of the steady
state inactivation curve (from −86.5±7.2 mV to −64.3±4.5 mV, n = 4, paired T-test, Fig. 2C). Coaplication of the T-type channel blocker Mibefradil (50 μM) further diminished Ca2+ currents in 4/5
cells tested (to 25±10 pA). These properties resemble Ca2+ currents in complex glial cells of the hippocampal CA1 region [14].

To identify the subtype(s) of Ca2+ expressed by NG2/EYFP
positive cells, transcript analysis was performed employing single
cell RT-PCR (Tab. S1). We found predominant expression of mRNA encoding the L-type channel isoforms Ca2+ 1.2 and Ca2+ 1.3
(Fig. 2D) and the T-type channels Ca2+ 3.1 and Ca2+ 3.2. Transcripts for P/Q and N-type channels, Ca2+ 2.1 and Ca2+ 2.2, were less abundant, while mRNAs for Ca2+ 2.4, Ca2+ 2.3 and Ca2+ 3.3 were never detected (Fig. 2D). Interestingly, the majority of NG2 cells tested (n = 39/46) expressed mRNA for the glial marker S100β. This is in line with our previous data showing that some of the NG2/EYFP positive cells express S100β while the astrocytic
marker GFAP was consistently lacking [Karam et al., 2008].

To further confirm the presence of functional Ca2+ in NG2 cells
of the hippocampus, Ca2+-imaging was combined with patch-
clamp recording in the whole-cell mode. Train stimulation via the
patch-pipette (15 consecutive depolarizing voltage steps (100 ms)
from −100 mV to +20 mV, see lower traces in Fig. 3B and 3C) produced reversible elevations of [Ca2++]i in NG2/EYFP cells
(Fig. 3A). It is important to note that in the same cell, several
[Ca2++]i elevations could be elicited up to 30 min after establishing
the whole-cell configuration (Fig. 3B). Next, we tested the sensitivity of the [Ca2++]i elevations to Ni2+. At high concentrations Ni2+ is known to non-specifically block Ca2+ channels [26,27]. Indeed, application of 200 μM Ni2+ abolished the [Ca2++]i elevations in the NG2/EYFP cells tested (n = 4) (Fig. 3B).

At these high concentrations, Ni2+ might also inhibit the NCX
[28]. To exclude that the observed block of [Ca2++]i elevations by Ni2+ was due to its action on NCX rather than Ca2+-, we tested the
sensitivity of evoked [Ca2++]i elevations to the NCX inhibitor SN-6. SN-6 has no effect on Ca2+- while blocking NCX operating in the
Ca2+-influx mode [29]. The amplitudes (103±34 pA vs. 86±22 pA, n = 5) and decay time-constants (39.4±6.8 ms vs. 39.2±3.6 ms, monoexponential fit, n = 4) of depolarization-
induced Ca2+ currents (at −10 mV) were not affected by SN-6 (10 μM; paired Student’s T-test, p>0.05; not shown; but see Fig. 2B). Together, these data demonstrate functional expression
of Ca2+ by NG2 cells in the hippocampus, corroborating previous
findings in complex glial cells of wild type mice [14].

We further analyzed the kinetics and amplitudes of depolarization-
induced [Ca2++]i elevations by Ca2+-imaging. Calibrated Ca2+-imaging measurements with Fura-2 revealed a free basal [Ca2++]i of
60 nM. Train stimulation led to an increase in [Ca2++]i (by 49±60 nM
(n = 8). The [Ca2++]i elevation immediately ceased after the last pulse (Fig. 3C). In contrast, [Ca2++]i elevations by a single pulse considerably outlasted the pulse duration. Maximal [Ca2++]i was observed about 1.2 s after stimulus offset. During this time Δ[Ca2++]i, almost doubled (from 4.3±1.6 nM to 8.1±1.6 nM; n = 6; Fig. 3C).

To improve time resolution of Ca2+-imaging we also performed
LSM based x-t line scans. Therefore, individual NG2/EYFP cells
were loaded with 400 μM Fluo-4 via the patch-pipette (Fig. 4A).

Table 1. Synopsis of ultrastructural analysis of neuron-NG2 cell synapses.

| cell | number of synapses observed | synaptic contacts | estimated total number of synapses |
|------|-----------------------------|------------------|----------------------------------|
|      |                            | asymm | symm | unclear | indication of perforation |                 |
| 1    | 3                           | 1     | 2    |         |                     | 5                |
| 2    | 6                           | 6     | 3    |         |                     | 11               |
| 3    | 8                           | 6     | 1    | 1       |                     | 14               |
| total| 17                          | 7     | 1    | 9       |                     | 30               |

For estimation of total synapse numbers (rounded), observed numbers were multiplied by 1.75 (see text).

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This approach confirmed the long-lasting $[\text{Ca}^{2+}]_i$ elevation and its slow kinetics as observed with the calibrated Fura-2 method. During single pulses, $\Delta F/F_0$ increased by 0.12±0.15 ($n=70$). Peak $\Delta F/F_0$ (0.20±0.20), however, only occurred 1.15 s after stimulus offset, and significantly exceeded the values registered at the end of the voltage step (paired Student’s T-test, $p<0.001$). Thus, kinetics and amount of $[\text{Ca}^{2+}]_i$ elevation were almost the same using either imaging technique (cf. Fig. 4B, C with Fig. 3C2, C1, respectively).

Obviously, there was a ceiling effect because the $[\text{Ca}^{2+}]_i$ elevations during train stimulation were much smaller than the calculated superposition of the responses to 15 single pulses (Fig. 4C). Saturation in $[\text{Ca}^{2+}]_i$ elevation and the prolonged kinetics of this signal cannot simply be ascribed to $\text{Ca}^{2+}$ influx through $\text{Cav}$s. First, saturation is unlikely to occur under these conditions because the limited $\text{Ca}^{2+}$-influx during the short stimulus trains can be expected to leave the driving force for $\text{Ca}^{2+}$ largely unchanged.

**Figure 2. Hippocampal NG2 cells express functional Cav s.** (A) Typical whole-cell current pattern of an EYFP positive NG2 cell (voltage steps between $-160$ and $+20$ mV with 10 mV increment, holding potential $-80$ mV). This cell had an input resistance of 221 MΩ, a membrane capacitance of 23 pF, and a resting potential of $-78$ mV. (B) Cav currents. (B1) Depicted Cav currents were separated by conditioning pre-pulses (1.5 s) to $-110$ and $-10$ mV (voltage-step duration 150 ms, upper schematic) while recording in Na$^+$ and K$^+$ free solution containing 1 μM TTX and 10 μM SN-6. Dotted line represents zero current level. (B2) Current voltage relationship of 5 pooled cells (upper curve, normalized to peak) and one exemplary cell (lower curve, corresponds to B1). (C) Basic pharmacological properties. (C1) Steady state inactivation curve before (open circles) and after (filled circles) wash in of verapamil (100 μM) (C2) Cav$^{2+}$ currents elicited at voltage steps to $-10$ mV after hyperpolarizing prepulses ($-110$ mV, 1.5 s, artifacts canceled for clarity). Verapamil (100 μM) reduced the initial peak current from 253 pA to 138 pA. Additional application of Mibefradil (50 μM) diminished the current to 28 pA. Upper traces represent baseline currents at $-80$ mV. (D) Single cell RT-PCR identified mRNA coding for different Cav subtypes. (D1) Representative agarose gel of mRNA-transcripts for Cav 1.2, 1.3, 1.4, and S100β. (D2) Relative abundance of Cav expression in NG2 cells. Cell numbers in parentheses. doi:10.1371/journal.pone.0017575.g002
Second, the \([\text{Ca}^{2+}]\) elevation outlasted channel open time more than tenfold but the binding kinetics of the \([\text{Ca}^{2+}]\) -indicator dyes used are in the range of microseconds [30]. Therefore, this can not account for the phenomenon.

Recently, it was suggested that in NG2 cells \([\text{Ca}^{2+}]\) elevation evoked by depolarization is mainly due to NCX operating in the \([\text{Ca}^{2+}]\)-influx mode in a tetrodotoxin (TTX) sensitive manner [17]. In our hands, TTX (1 \(\mu\)M, \(n = 13\)) neither affected the amplitudes nor the kinetics of depolarization-induced \([\text{Ca}^{2+}]\) elevations in NG2/EYFP cells (\(n = 13\), Fig. 4D). This goes in line with our finding, that Cav channels were not influenced by the specific NCX reverse mode blocker, SN-6 (Fig. 2B).

\(\text{Ca}^{2+}\)-influx through \(\text{Ca}_{\text{s}}\) evokes \(\text{Ca}^{2+}\)-induced \(\text{Ca}^{2+}\)-release in NG2 cells

\(\text{Ca}^{2+}\)-influx through the plasma membrane may evoke further increase in \([\text{Ca}^{2+}]\), by triggering \(\text{Ca}^{2+}\)-release from intracellular stores [31], which might account for the observed saturation and prolonged kinetics of \([\text{Ca}^{2+}]\), elevations. To investigate whether \(\text{Ca}^{2+}\)-induced \(\text{Ca}^{2+}\)-release (CICR) is operative in NG2 cells we performed recordings in nominal \(\text{Ca}^{2+}\)-free bath solution supplemented with 2 mM EDTA. Under these conditions \(\text{Ca}^{2+}\)-traces represent the average of 4 cells. \(\text{Ca}^{2+}\)-traces evoked by train stimulation (bottom). Note that the increase in \([\text{Ca}^{2+}]\), stopped immediately at the end of the stimulation (gray). \(\Delta[\text{Ca}^{2+}]\), amounted to 49 nM. \(\text{Ca}^{2+}\)-traces represent the average of 8 cells. (C) In contrast, single step depolarization (100 ms) typically elicited prolonged \([\text{Ca}^{2+}]\), elevation, outlasting depolarization (gray). Peak \([\text{Ca}^{2+}]\), was reached 1.3 s after stimulus onset and amounted to \(\Delta[\text{Ca}^{2+}]\), \(= 8.1\) nM. At the end of the depolarization the \([\text{Ca}^{2+}]\), elevation reached only 50% of the maximum (\(\Delta[\text{Ca}^{2+}]\), \(= 4.3\) nM). \(\text{Ca}^{2+}\)-traces represent the average of 6 cells.

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**Figure 3. Fura-2 based calibrated \(\text{Ca}^{2+}\)-imaging.** (A) Depolarization of NG2 cells reproducibly generated \([\text{Ca}^{2+}]\), elevations, recorded as \(-F_{380}/F_{362}\) fluorescence ratio. (A1) Repeated train stimulations (15 depolarizations from \(-100\) to \(+20\) mV, 100 ms each; indicated by the gray box) were applied to three exemplary NG2 cells. (A2) Run down of the \([\text{Ca}^{2+}]\), elevation over time as revealed with successive stimulation. Amplitudes shown in (A1) were normalized to the first response, which was recorded 9 min after establishing the whole cell configuration. (B) 200 \(\mu\)M Ni\(^{2+}\) abolished the train depolarization-induced \([\text{Ca}^{2+}]\), elevations. The upper traces illustrate the \([\text{Ca}^{2+}]\), elevation in NG2 cells, before (circles) and after (triangles) application of Ni\(^{2+}\). The lower panel shows the simultaneously recorded current responses. \(\text{Ca}^{2+}\)-traces represent the average of 4 cells. (C) \(\text{Ca}^{2+}\)-imaging. (C1) \([\text{Ca}^{2+}]\), elevations (upper traces) evoked by train stimulation (bottom). Note that the increase in \([\text{Ca}^{2+}]\), stopped immediately at the end of the stimulation (gray). \(\Delta[\text{Ca}^{2+}]\), amounted to 49 nM. \(\text{Ca}^{2+}\)-traces represent the average of 8 cells. (C2) In contrast, single step depolarization (100 ms) typically elicited prolonged \([\text{Ca}^{2+}]\), elevation, outlasting depolarization (gray). Peak \([\text{Ca}^{2+}]\), was reached 1.3 s after stimulus onset and amounted to \(\Delta[\text{Ca}^{2+}]\), \(= 8.1\) nM. At the end of the depolarization the \([\text{Ca}^{2+}]\), elevation reached only 50% of the maximum (\(\Delta[\text{Ca}^{2+}]\), \(= 4.3\) nM). \(\text{Ca}^{2+}\)-traces represent the average of 6 cells.
AMPA and GABA_A receptor-mediated depolarization evokes $[\text{Ca}^{2+}]_i$ elevation

Due to a relatively high $[\text{Cl}^-]_i$ in NG2 cells, activation of GABA_A receptors has a depolarizing effect [6]. We tested if application of AMPA or GABA_A receptor agonists induce elevations in $[\text{Ca}^{2+}]_i$, in NG2/EYFP cells. TTX (1 $\mu$M) was added to the bath solution to reduce indirect effects. In the current-clamp mode, the AMPA/kainate receptor agonist kainate (500 $\mu$M, $n = 4$) as well as the GABA_A receptor agonist muscimol (250 $\mu$M, $n = 4$) induced $[\text{Ca}^{2+}]_i$ elevations (Fig. 6A). In the voltage-clamp mode, only kainate (100 $\mu$M, $n = 4$) evoked increases in $[\text{Ca}^{2+}]_i$ (Fig. 6B), due to activation of $\text{Ca}^{2+}$-permeable AMPA receptors [11,12,16,24,33]. Muscimol (10 $\mu$M), although evoking larger inward currents than kainate, failed to affect $[\text{Ca}^{2+}]_i$ (n = 4) (Fig. 6B). These data demonstrate that AMPA/kainate receptor activation may produce direct ($\text{Ca}^{2+}$)-influx through the receptor pore or possibly through metabotropic effects [34] and indirect (depolarization-induced opening of $\text{Ca}_{\text{L}}$ followed by $\text{Ca}^{2+}$-influx) $[\text{Ca}^{2+}]_i$ elevations. In contrast, GABA_A receptor-induced $\text{Ca}^{2+}$-influx in NG2 cells is indirect, i.e. due to membrane depolarization and $\text{Ca}^{2+}$-activation.

NG2 cells express functional group I metabotropic glutamate receptors

Next, we tested whether NG2 cells express metabotropic glutamate receptors (mGluRs). The group I mGluR-specific agonist 3,5-DHPG was locally applied, while membrane currents and $[\text{Ca}^{2+}]_i$ were monitored by simultaneous patch-clamp recording in the whole cell mode and line scan imaging. All cells tested responded to 3,5-DHPG with $[\text{Ca}^{2+}]_i$ elevations (100 $\mu$M, $n = 7$; $\Delta F/F_0 = 1.14 \pm 0.79$, n = 6, 10 $\mu$M). This was never accompanied by current responses (Fig. 7A). The delay between substance arrival and the onset of $[\text{Ca}^{2+}]_i$ rises (see [Material and Methods] for details) varied among cells ($3.4 \pm 3.3$ s, $n = 7$, range between 0.6 and 9.4 s), but not between multiple 3,5-DHPG applications to the same individual cell.

Pre-application of the unspecific group I mGluR antagonist LY341495 [35] (10 $\mu$M, $n = 7$), immediately followed by co-application of 3,5-DHPG and LY341495 (11 s, 10 $\mu$M both), reversibly blocked the $[\text{Ca}^{2+}]_i$ elevations (n = 2, Fig. 7A). Although indicating the involvement of mGluRs, these responses might have been produced indirectly, i.e. via mGluR activation of neighboring cells that innervate the NG2 cell. Another
constraint of these experiments was the significant run down of the [Ca\(^{2+}\)] for 40±16% of the initial amplitudes, two applications, n = 4), probably due to wash-out of cytosolic constituents during whole cell recording. To circumvent these limitations we added TTX (1 μM) to block action potentials and inhibited P2Y receptors (with 100 μM PPADS, 100 μM suramin), mACh receptors (with 5 μM ipratropium), 5-HT2 receptors (with 10 μM methysergide), NMDA receptors (with 10 μM prazosin), and GABA\(_{\mathrm{A}}\) receptors (with 2 μM CGP55845). In addition, local loading of groups of NG2/EYFP cells with Fluor-4 AM was employed using focal pressure application (Fig. 7B, B). Under these conditions, almost all NG2 cells tested (96%) showed robust [Ca\(^{2+}\)]i elevations upon application of 3,5-DHPG (11 s; 10 μM; ΔF/F\(_0\) = 0.56±0.36, n = 108). Further analysis using mGluR group I subtype-specific antagonists [reviewed by [36,37,38]] indicated a non-uniform distribution of mGlnR1 and mGlnR5 in NG2/EYFP cells. The mGluR1 antagonist 3-MATIDA (30 μM) abolished the [Ca\(^{2+}\)]i elevations in 20% of the cells (n = 2/10; Fig. 7C, for both antagonists: 23 s pre-application followed by 11 s co-application with 10 μM 3,5-DHPG). In the remaining cells, 3-MATIDA (n = 8) and MPEP (n = 6) exerted partial inhibition of 3,5-DHPG-induced responses (to 47±17%) that did not differ significantly between the antagonists. Co-application of both antagonists abolished [Ca\(^{2+}\)]i elevations in 88% of the NG2 cells tested (n = 13/17) (Fig. 7C). The [Ca\(^{2+}\)]i transients recovered after wash out of the antagonists to 78±23% (n = 47) of the initial value. We noted that all cells were sensitive to at least one of the two antagonists.

Pre-synaptic fiber tract stimulation evokes [Ca\(^{2+}\)]i elevations in the soma of NG2 cells

Next, we investigated whether pre-synaptic stimulation of GABAergic interneurons or axons of glutamatergic CA3 neurons provokes [Ca\(^{2+}\)]i elevations in NG2 cells. Minimal stimulation induced post-synaptic currents in NG2/EYFP cells matching those observed in weakly fluorescent hGFAP/EGFP cells (previously termed GluR cells) [23] or wild type hippocampus (termed OPCs, not shown) [24,25]. Tetanic stimulation (100 Hz, 10 s) caused robust depolarization (ΔV = 15±5 mV, n = 11) (Fig. 8A, bottom) while producing only small elevations of somatic [Ca\(^{2+}\)]i, (ΔF/F\(_0\) = 0.039±0.030, n = 11). To simulate more physiological conditions, single pulses (20 μs) were applied. With this protocol, a failure rate of about 60% was observed. Excluding failures, the post-synaptic depolarization now amounted to 1.5±0.6 mV (resting membrane potential = -71±6 mV, n = 12). These depolarizations were never accompanied by somatic [Ca\(^{2+}\)]i elevations (n = 12) (Fig. 8B). Obviously, the sparse innervation of hippocampal NG2 cells is insufficient to provoke [Ca\(^{2+}\)]i elevations at the cell soma under these conditions.

NG2 cells express vesicular glutamate transporters

The observation of stimulus-induced [Ca\(^{2+}\)]i elevations prompted us to search for potential downstream signaling mechanisms in NG2 cells. Astrocytes express vesicular glutamate transporters (vGLUTs) in their distal processes, and were reported to communicate with neurons by Ca\(^{2+}\)-dependent release of vesicular glutamate [39–41]. To investigate whether vGLUTs may also be expressed by NG2 cells, transcript analyses were performed. vGLUT1 and vGLUT2, but not vGLUT3 could be detected by post-recording single cell RT-PCR from NG2 cells of hGFAP/EGFP mice (p9–15). Gene transcripts for vGLUT1 and vGLUT2, but not vGLUT3 could be detected by post-recording single cell RT-PCR from NG2 cells of hGFAP/EGFP mice (p9–15). Gene transcripts for vGLUT1 and vGLUT2, but not vGLUT3 could be detected by post-recording single cell RT-PCR from NG2 cells of hGFAP/EGFP mice (p9–15). Gene transcripts for vGLUT1 and vGLUT2, but not vGLUT3 could be detected by post-recording single cell RT-PCR from NG2 cells of hGFAP/EGFP mice (p9–15). Gene transcripts for vGLUT1 and vGLUT2, but not vGLUT3 could be detected by post-recording single cell RT-PCR from NG2 cells of hGFAP/EGFP mice (p9–15). Gene transcripts for vGLUT1 and vGLUT2, but not vGLUT3 could be detected by post-recording single cell RT-PCR from NG2 cells of hGFAP/EGFP mice (p9–15). Gene transcripts for vGLUT1 and vGLUT2, but not vGLUT3 could be detected by post-recording single cell RT-PCR from NG2 cells of hGFAP/EGFP mice (p9–15).
recording and biocytin filling. Staining was observed for vGLUT1 (2/3 cells) and vGLUT2 (2/2 cells). Larger vGLUT1 positive puncta, putative vesicle groups, were found in the fine NG2 cell processes (Fig. 9). The inclusion of vGLUT-immunoreactivity (vGLUT-IR) within NG2 cell profiles was verified at high magnification by 3D inspection (Fig. 9A), and by increasing the opacity of surface-rendered, 3D-reconstructed NG2 cells (Fig. 9B, Video S1). Based on the rigorous thresholding, we assume that in our analysis the amount of vGLUT-IR in NG2 cells is underestimated. vGLUT1 or vGLUT2 positive puncta did not display a preference for the varicosities of NG2 cell processes but occurred all over the process tree, also at any proximo-distal distance. The immunohistochemical and RT-PCR data indicate heterogeneity among NG2 cells with regard to expression of vGLUTs.

NG2 cell processes are motile and display actin and ezrin, but not tubulin

Recent reports suggested a link between [Ca\(^{2+}\)]\(_i\) elevation and migration of NG2 cells in vitro [17]. To investigate the possibility of process motility in situ, we performed time-lapse recordings in acute hippocampal slices. We detected process motility in 5 out of 11 dye-labeled NG2/EYFP cells (Fig 10A). At least three types of process motility were observed; including elongation (Fig. 10B) and retraction (Fig. 10C) of processes (see also Videos S2, S3). Additionally, we observed that strongly dye-labeled varicosities, which are characteristic of NG2 cells, move along the processes (Fig 10D). The varicosities traveled up to 2.9 \(\mu\)m within 6 min (Fig. 10D). Some varicosities showed bi-directional motility. Thus, NG2 cell processes and their varicosities exhibit motility on a minute time range.

Next, we investigated cytoskeletal constituents potentially relevant to motility of NG2 cells. Therefore, cells were freshly isolated from tg[hGFAP/EGFP] mice and selected according to their characteristic morphology and specific immunolabeling (GFP positive, GFAP negative) [18]. Antibodies against \(\alpha\)-tubulin, \(\beta\)-actin, ezrin (a microvillus-associated, actin-binding protein [42]), or protein disulfide isomerase (PDI) were combined with both, anti-GFP and anti-GFAP staining. Noteworthy, \(\alpha\)-tubulin (6/6) was not present in the processes but restricted to the soma and in a few cases to the proximal portion of processes (Fig. 10E). At the same time, the processes of nearby astrocytes were positive for \(\alpha\)-tubulin (Fig. S1). \(\beta\)-actin (10/10) and ezrin (10/10) were distributed all over the cell including the fine NG2 cell processes (Fig. 10F,G). PDI-IR (10/10) was restricted to the soma and never detected in the NG2 cell processes (Fig. 10H).
**Figure 7. NG2 cells express mGluRs.** (A1) The group I mGluR agonist 3,5-DHPG (100 μM, 11 s) produced [Ca^{2+}]_{i} increases (upper trace) but not membrane currents (bottom). (A2) Pre-application of LY341495 (10 μM, 11 s) followed by co-application of 3,5-DHPG (10 μM, 11 s) completely blocked the 3,5-DHPG-induced [Ca^{2+}]_{i} elevation. Three min later, 3,5-DHPG (10 μM, 11 s) again provoked a [Ca^{2+}]_{i} elevation in the same cell. (B1) Region of interest with five NG2/EYFP cells (bar, 20 μm) which was selected for focal application of Fluo-4 AM (B2). Arrows mark NG2 cells from which [Ca^{2+}]_{i} elevations were recorded (also seen in C1). Note Fluo-4 labeled, EYFP-negative cells located in the lower left corner. (C1–C3) [Ca^{2+}]_{i} elevations upon 3,5-DHPG in the presence (middle) and absence (left, control; right, wash) of subtype-specific mGluR antagonists. Antagonists were pre-applied for 23 s, followed by 11 s co-application with 3,5-DHPG (10 μM, gray boxes). Applications were separated by 3 min. (C1) The mGluR1 specific antagonist 3-MATIDA (50 μM) mostly exerted partial block of Ca^{2+}-responses. (C2) In most cells, 3,5-DHPG-mediated [Ca^{2+}]_{i} elevations were abolished by the mGluR5 specific antagonist, MPEP (20 μM). (C3) In a few cells, co-application of both antagonists failed to inhibit 3,5-DHPG-induced [Ca^{2+}]_{i}.
Discussion

NG2 cells display several mechanisms of intracellular Ca\(^{2+}\)-elevation

Our data demonstrate the capability of gray matter NG2 cells to increase [Ca\(^{2+}\)]. via several independent pathways: G-protein coupled receptors, as well as ligand- and voltage-gated ion-channels. While the presence of mGluRs in NG2 cells represents a new finding, expression of Ca\(_{\text{v}}\)s is under discussion. Recently, it was reported that NG2 cells in the hippocampus lack Ca\(_{\text{v}}\)s [16,17]. In contrast, earlier work on complex glial cells in the hippocampus was reported that NG2 cells in the hippocampus lack Cavs [16,17]. The discrepancy with the former data may be due to different recording conditions. Ca\(^{2+}\)-currents through NG2 cell Ca\(_{\text{v}}\)s may be different recording conditions. Ca\(^{2+}\)-currents in NG2 cells are small in amplitude, compared with the dominating K\(^+\) currents. Its reliable separation requires use of Na\(^+\) - and K\(^+\)-free solutions, elevated [Ca\(^{2+}\)] in the bath solution and application of conditioning pre-pulses.

The small amplitudes and high activation threshold of the Ca\(^{2+}\)-currents through NG2 cell Ca\(_{\text{v}}\)s raise the question of its physiological relevance. To tackle this question, we employed Ca\(^{2+}\)-imaging. Using aGFP, depolarization evoked reversible [Ca\(^{2+}\)], elevations in NG2 cells. This was due to influx of Ca\(^{2+}\) through Ca\(_{\text{v}}\)s, but not to the activation of NCXs, as recently suggested [17]. A possible explanation for this conflicting finding might be that in the latter study, KB-R 7943 was used as an inhibitor of NCX, which blocks Ca\(_{\text{v}}\)s with almost the same affinity [45]. Similarly, Ni\(^{2+}\) does not only block Ca\(_{\text{v}}\)s but also NCXs [20]. SN-6, on the other hand antagonizes with high affinity only the Ca\(^{2+}\)-influx mode of NCXs, preferentially of NCX1, while not interfering with Ca\(_{\text{v}}\)s at the concentration used here [29]. Because (i) SN-6 did not affect the electrophysiologically recorded Ca\(^{2+}\)-currents (Fig. 2B) and (ii) TTX did not diminish the voltage-step induced [Ca\(^{2+}\)], elevations (Fig. 4D) we believe that in NG2 cells Ca\(^{2+}\)-influx through NCXs plays only a minor role, if any. The functional characterization of the NG2 cell Ca\(_{\text{v}}\) subtypes is a challenging task for future studies. The transcript data reported here together with the pharmacological findings by Akopian [14] might provide first clues.

[Ca\(^{2+}\)], elevation through Ca\(_{\text{v}}\) activation was almost doubled due to CICR. Notably, this led also to a significant prolongation of the [Ca\(^{2+}\)], elevations. Thus, CICR represents a powerful mechanism to amplify small inward currents through Ca\(_{\text{v}}\)s in NG2 cells. The observed saturation effect (Fig. 4C) suggests the involvement of Ca\(^{2+}\) binding sites with low affinity acting as intracellular Ca\(^{2+}\) sensors, analogously to myocardial cells [e.g. [46]]. This may regulate the gain of CICR depending on ambient [Ca\(^{2+}\)] levels. Currently, we do not know whether Ca\(^{2+}\) amplification exists in NG2 cell processes. The absence of PDI-IR from processes (Fig. 10H) precludes CICR in these structures, and potential amplification mechanisms would have to be independent of endoplasmic reticulum.

In agreement with previous findings [6] our data suggest the presence of Ca\(^{2+}\)-permeable AMPA/kainate and GABA\(_{\text{A}}\) receptors in NG2/EYFP cells. Activation of the latter receptors depolarizes NG2 cells, which might trigger the activation of Ca\(_{\text{v}}\)s. Such indirect GABA receptor-mediated [Ca\(^{2+}\)], elevations have been observed in cultured OPCs [15]. Depolarizations induced by AMPA/kainate receptor activation might have similar effects, although we can not exclude a contribution of metabotropic kainate receptors to the [Ca\(^{2+}\)], elevations [34]. It will be a challenge to determine whether in the fine processes, receptor activation produces depolarization sufficient for Ca\(_{\text{v}}\) activation in NG2 cells under physiological conditions.

We further report that NG2 cells in the hippocampus express functional group I mGluRs. Pharmacological analysis indicated...
preferential expression of mGluR5, while only a minority of the 3,5-DHPG-induced [Ca\textsuperscript{2+}]\textsubscript{i} elevations were sensitive to an mGluR1 antagonist. Whether these receptors are activated upon pre-synaptic release of glutamate needs to be demonstrated. In the present study, focusing on post-synaptic NG2 cell depolarization, fiber tract stimulation-induced [Ca\textsuperscript{2+}]\textsubscript{i} elevations have only been monitored in the soma during whole cell recording. It is very likely that dialysis of the cytosol led to an attenuation of the [Ca\textsuperscript{2+}]\textsubscript{i} elevations. NG2 cell processes are highly motile, actin-based surface extensions

Our live microscopic data demonstrate, for the first time, motility of NG2 cell processes in \textit{vivo}. We investigated the presence of cytoskeletal proteins in NG2 cell processes to test for prerequisites of process motility. The cytoskeleton of NG2 cell processes is found to be actin-based, since GFAP-positive glial (intermediate) filaments or microtubules were not observed by immunolabeling and electron microscopy. This appears astonishing in respect of their length (30–50 μm) and small diameter (0.2–1 μm) in between the varicose expansions. Of the many actin-binding proteins ezrin was chosen as a further marker, because its (de)phosphorylation-based mode of membrane-to-cytoskeleton linking enables rapid shape changes [47]. Ezrin, and its close relatives, radixin and moesin (the ERM protein family), are typically involved in establishing highly motile and very narrow structures in the CNS, such as neuronal growth cone filopodia [47,48] or peripheral astrocyte processes [49,50]. Also, ERM proteins are required for maintaining stereocilia integrity in cochlear and vestibular hair cells [51]. Altogether, the set of features displayed by NG2 cell processes classifies them as actin-based stereocilia and surface extensions. They constitute a rare example of an actin-based surface extension that is directly involved in synaptic signaling.

Possible impact of the synaptic input onto NG2 cells

Recent findings suggest a role of neuron-NG2 cell synapses in migration. Thus, in the corpus callosum adult-born migrating NG2 cells receive glutamatergic synaptic input from demyelinated axons [52], and GABA-mediated [Ca\textsuperscript{2+}]\textsubscript{i} elevation is essential for migration of subventricular zone NG2 cells to and within white matter \textit{in vitro} [17]. Ca\textsubscript{s}s might be important in this context as they have been reported to govern migration in newborn neurons, e.g. in the postnatal olfactory bulb [53]. However, the reported data relate to lesioned white matter, where neuron-glia synapses are transient [52]. In contrast, gray matter NG2 cell synapses are lesion independent and functional under physiological conditions. An alternative function of synaptic input on NG2 cells in gray
Figure 10. Properties of NG2 cell processes. (A–D) Two-photon time-lapse recordings. (A) Overview of an Alexa-594-labeled NG2/EYFP cell (maximum projection, 100 µm x 100 µm x 15 µm, 60 equidistant planes, scale 10 µm). (B–D) Pairs of maximum projections (16 µm x 14 µm x 5 µm, 20 planes, scale 2 µm) taken at time points $t_0$ (left) and $t_0 + \Delta t$ (right). Arrows mark processes that were elongated (B, $\Delta t = 185$ s) or retracted (C, $\Delta t = 370$ s). Additionally we observed varicosities traveling along the process (D, $\Delta t = 370$ s, start and end point marked by arrows). See also Videos S2, S3. (E–H) NG2 cell processes do not contain $\alpha$-tubulin and PDI. Cortical tissue from an hGFAP/EGFP mouse (p13) was freshly dissociated and quadruple-stained with a nuclear marker (bisbenzimidine, blue) and antibodies against GFAP (also blue channel), GFP (green) and one of the proteins of interest (red): $\alpha$-tubulin (E), $\beta$-actin (F), ezrin (G) or the ER marker PDI (H). The cells analyzed were GFAP negative, GFP positive. Note nearby GFP negative cells (overviews, left in F–H, E). Areas boxed in the overviews (F–H) are enlarged for colocalization analysis. $\beta$-actin and ezrin were localized in the NG2 cell processes. Note the fine dimensions of these varicose processes visualized in the GFP channel (green). The PDI signal is present both in a
Ca2+-Signaling and Process Motility in NG2 Cells

non-identified, nearby cell (H, overview) and in the soma of the NG2 cell, but not in its processes (H, red, merge). The same is observed for α-tubulin (E red, merge). Note that α-tubulin/microtubules are well-preserved in the processes of nearby non-identified cells (E, merge) and of GFAP positive astrocytes (cf. Fig. S1). Scale bar 5 μm.

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Materials and Methods

Maintenance and handling of animals used in this study was according to local government regulations. Experiments have been approved by the State Office of North Rhine-Westphalia, Department of Nature, Environment and Consumerism (LANUV NRW, approval number 9.93.2.10.31.07.130). All measures were taken to minimize the number of animals used.

Slice preparation

Transgenic mice with human GFAP promoter-controlled expression of EGFP (tgGFAP/EGFP) mice [55] or knockout mice in which the chromophore EYFP has been inserted after the start ATG of the endogenous NG2 gene [22] aged postnatal day (p) 7–15 were anaesthetized, decapitated, and the brains were removed. Coronal hippocampal slices (200 μm thick) were cut in ice-cold oxygenated solution consisting of (in mM): 87 NaCl, 2.5 KCl, 1.25 NaH2PO4, 7 MgCl2, 0.5 CaCl2, 25 NaHCO3, 25 glucose, 75 sucrose (347 mOsm). Slices were stored for 30 min in carbogen (95% O2 and 5% CO2) to a pH of 7.4 (room temperature). The Ca2+-signaling pathways reported here. However, because the processes are devoid of endoplasmic reticulum, these [Ca2+]i elevations are unlikely to be amplified by CICR and might occur locally confined. Local [Ca2+]i elevations might play a role in regulation of process motility. In addition, restricted Ca2+-signaling might be interesting in the light of the demonstrated vGLUT expression. In neurons, vGLUT expression is sufficient for defining a glutamatergic phenotype [54]. In astrocytes vGLUTs mediate vesicular transmitter release, at least in the cell culture [39–41]. The scattered vGLUT organelles within NG2 cell processes might serve a similar function. The intriguing perspective that NG2 cells might signal to neighboring cells in a Ca2+-dependent manner remains to be addressed in future studies.

Electrophysiological recordings

Slices were transferred to a recording chamber and constantly perfused with aCSF at room temperature. Whole-cell recordings were obtained using an EPC7 or EPC8 amplifier (HEKA Elektronik, Lambrecht, Germany). The holding potential in the voltage clamp mode was −80 mV if not stated otherwise. Signals were digitized with an ITC 16 or LIH 1600 (HEKA). Patch-pipettes, fabricated from borosilicate capillaries (Hilgenberg, Malsfeld, Germany), had resistances of 4–7 MΩ when filled with a solution consisting of (in mM): 130 KCl, 2 MgCl2, 3 Na2-ATP, 5 1,2-bis(o-aminophenoxy)ethane-N,N,N',N'-tetraacetic acid (BAPTA), 10 2(-4(-2-hydroxyethyl)-1-piperazinyl)-ethanesulfonic acid (HEPES) (pH 7.25).

For separation of Ca2+-currents, Na+- and K+-free bath and pipette solutions were used as described by Akopian et al. [14]. HEPES-based bath solution contained (in mM): 130 tetraethyl-ammonium chloride (TEA), 10 HEPES, 5 CaCl2, 4-aminopyrrolidin-4-AP, 10 glucose, supplemented with 1 μM TTX. HEPES-buffered solutions were continuously bubbled with O2. The pipette solution contained (in mM): 120 N-methyl-D-glucamine chloride (NMDG), 20 TEA, 0.5 CaCl2, 5 ethylene glycol bis-(β-aminoethylether) N,N'-tetraacetate (EGTA), 2 MgCl2, 3 Na2-ATP, 10 HEPES (pH 7.2). Liquid junction potentials have been corrected for.

Recordings were monitored with TIDA software (HEKA). Series and membrane resistance were checked in constant intervals with self-customized macros using Igor Pro 6 software (WaveMetrics Inc., Lake Oswego, USA). Visual control was achieved by a microscope equipped with an infrared DIC system (Leica DM6000, Leica, Mannheim, Germany) and an IR objective (HCX APO L 20x/1.0 W; Leica). Infrared and epifluorescence images were captured with a digital CCD camera (DFC350FX R2; Leica).

Membrane currents were compensated offline for stimulus artifacts using Igor Pro 6 software according to the following procedure: Ten traces evoked by voltage steps from −80 to −70 mV were averaged and fitted monoexponentially. Compensated current traces were obtained by multiplying the fitted curve with the respective factors and subsequent subtraction from the original current traces at different membrane potentials.

Evoked post-synaptic currents in NG2 cells were compensated for stimulus artifacts by subtracting averaged failure traces.

Substances were pressure-applied focally using a multichannel Octalflow superfusion system (ALA Scientific Instruments, Farmingdale, USA). The 20–80% rise time of agonist concentration amounted to ~100 ms. Short test pulses of GABA were used to assess the delay between valve opening and arrival of the substance at the recorded cell, which ranged between 0.4 and 0.8 s. All agonist responses were corrected for this delay. In some cases, substances were applied by changing the bath solution. All statistical data are given as mean ± SD.

Two-photon time-lapse imaging

Individual NG2/EYFP-positive cells were filled for 2 min with Alexa-594 (Invitrogen, Karlsruhe, Germany) via the patch-pipette [56]. Dye was allowed to diffuse for >30 min before imaging. Subsequent two-photon imaging was performed on a confocal laser scanning microscope (LSM/SP5, Leica) equipped with a mode-locked infrared laser (MaiTai BB, Newport/Spectra Physics, Irvine, USA). The dye was excited at 610 nm and emitted light was detected with built-in non-descan detectors below 680 nm. These experiments were performed at 35°C to increase process motility. The bicarbonate concentration of aCSF was reduced to 20 mM to achieve correct pH values. Image stacks of up to 60 optical sections were acquired for 20 to 60 min (z-step distance 250 nm, aCSF). We assured by inspection of all optical planes that the observed cellular motility was not caused by drift of slices, recording chamber, or microscope.

Ca2+-imaging

NG2/EYFP cells in the stratum radiatum of the CA1 area were used for Ca2+- imaging. To determine absolute [Ca2+]i, and achieve a high time resolution of Ca2+-transients two different methods were applied.
(i) Changes in $[Ca^{2+}]_i$, were monitored by a CCD camera (SensiCam; TILL photonics, Martinsried, Germany) mounted on a wide-field epifluorescence system (Polychrome II, TILL photonics). It was attached to an upright microscope (Axioskop FS2; Zeiss, Oberkochen, Germany) equipped with a 60x LUMPlan HI/IR objective (Olympus Optical Co., Hamburg, Germany). Fluorescence excitation was achieved by a monochromator. Individual cells in acute hippocampal slices were loaded via the patch-pipette with Fura-2 (200 μM; Invitrogen). Dye filling lasted ≥3 min before Ca^{2+}-imaging was started. If not stated otherwise, Fura-2 was excited at 380 or 340 nm for 40 ms and emission was detected at an acquisition rate of 25 Hz during, and 3 Hz after depolarization. Single frames were recorded at the isosbestic point (362 nm) before and after each sequence. This allowed offline calculation of pseudo-ratio images to correct for bleaching. The latter was assumed to be proportional to exposure time. A linear function was calculated from the first and the last 362 nm frame of each of the 380 or 340 nm sequences. This function was used to determine the 362 nm values for each recorded frame. Pseudo-ratios F_{380}/F_{340} were calculated from the measured F_{380} or F_{340} and the extrapolated F_{362} values for each time point. F_{380}/F_{362} pseudo-ratios were inversely plotted so that $[Ca^{2+}]_i$, elevations are always indicated by upward deflections.

Absolute $[Ca^{2+}]_i$, was estimated through calibration according to Grynkiewicz et al. [57]:

$$[Ca^{2+}]_i = K_{diff} \cdot \left( \frac{R_{min} - R}{R_{max} - R} \right)$$

with $R = F_{340}/F_{380}$

$$F_{340} = SF_{340} \cdot c_f + SB_{340} \cdot c_b; \quad F_{380} = SF_{380} \cdot c_f + SB_{380} \cdot c_b$$

($c_f$: concentration of Ca$^{2+}$ bound Fura-2)

$$R_{min} = SF_{340}/SB_{340}; \quad R_{max} = SB_{340}/SB_{380}$$

$$K_{diff} = SF_{380}/SB_{380} \cdot K_d; \quad K_d = c_f \cdot [Ca^{2+}]_i \cdot c_b$$

$R_{min}$ and $R_{max}$ were determined with 10 mM BAPTA or 10 mM CaCl2 in the pipette solution, respectively. $K_d$ was determined with a pipette solution buffered to 11 mM free Ca$^{2+}$ and amounted to 51 nM. $R(t)$ curves were calculated from two successive recordings at 380 nm and 340 nm. F_{340(0)} and F_{340(t)} were corrected for bleaching using the pseudo-ratio method described above. Calibration was performed using self-customized IGOR 6 functions.

(ii) Alternatively, an LSM (Leica) was used for Ca$^{2+}$-imaging, allowing for higher time resolution. Individual NG2/EYFP positive cells were loaded with Fluo-4 (400 μM, Invitrogen) via the patch-pipette. Subsequent line-scans, taken at the soma, were recorded with an acquisition rate of 488 nm. Emission was detected between 500 and 650 nm. Signals were sampled at 1–0.4 kHz. Changes in $[Ca^{2+}]_i$, measured as change in fluorescence intensity ($\Delta F$), were offline related to the baseline fluorescence ($F_0$) according to $\Delta F/F_0 = (F - F_0)/F_0$. Time-correlated signals from individual cells were averaged to improve signal-to-noise ratio. For local loading of groups of EYFP positive cells, Fluo-4 AM (10 μM, Invitrogen) with 0.01% Pluronic F127 was focally pressure-applied for 5 min employing an Octaflow System (ALA Scientific Instruments). x-y-t scans of 2.2 μm thick single optical planes were recorded. $\Delta F/F_0$ was determined in separate regions of interest (ROIs) placed in each NG2 cell soma in the field of view. Data analysis was performed with LAS Live Data Mode (Leica) and IgorPro 6 software. 3-MATIDA, (S)-3,5-DHPG, CGP 55845, GABA, ipratropium, kainic acid, methysgalride, MPEP, muscimol, PPADS, prazosin, SN-54, suramin, and thapsigargin were from Tocris (Bristol, UK).

Fiber tract stimulation

Stimulation was performed with monopolar glass pipettes filled with ACSF. Pipette resistance ranged between 0.5 and 2 MΩ. Biphasic constant voltage-pulses of 100–200 μs were applied with a stimulus generator (STG 2004, Multi-Channel-Systems, Reutlingen, Germany). High-frequency stimulation was accomplished using Mc Stimulus 2 software (Multi-Channel-Systems). Time correlation was achieved by synchronizing TTL pulses generated by the recording software (TIDA 5.22, HEKA).

Single cell RT-PCR

After electrophysiological characterization in situ, the cytoplasm of individual cells was harvested under microscopic control as reported previously [18]. Reverse transcription (RT) was started after addition of RT-buffer, 10 mM DTT (final concentration; Invitrogen), 4×250 μM dNTPs (Applied Biosystems, Darmstadt, Germany), 50 μM random hexamer primer (Roche, Mannheim, Germany), 20 U RNase inhibitor (Promega, Madison, USA), and 100 U SuperscriptIII reverse transcriptase (Invitrogen). Final volume was ~10 μl. A multiplex two-round PCR with single-cell cytosol was performed with primers for the Ca_1, Ca_2, and Ca_3 families or vesicular glutamate transporters (vGLUT) 1/2 and vGLUT3, respectively (Table S1). Primers were located in conserved regions to amplify all members of the respective family. The first PCR was performed after adding PCR buffer, MgCl2 (2.5 mM), and primers (200 nM each) to the reverse transcription product (final volume 50 μl). Taq polymerase (3.5 U; Invitrogen) was added after denaturation, 45 cycles were performed (denaturation at 94°C, 25 s; annealing at 54°C, first five cycles: 2 min, remaining cycles: 45 s; extension at 72°C, 25 s; final elongation at 72°C, 7 min). An aliquot (2 μl) of the PCR product was used as a template for the second PCR (35 cycles; annealing at 54°C, first five cycles: 2 min, remaining cycles: 45 s) using nested, subunit-specific primers (Table S1). The conditions were the same as described for the first PCR-round, but dNTPs (4×50 μM) and Platinum Taq polymerase (2.5 U; Invitrogen) were added. Products were identified by gel electrophoresis using a molecular weight marker (Phi X174 HincII digest; Eurogentec, Seraing, Belgium).

Primer specificity was tested with total RNA from freshly isolated mouse brain (p20). For optimization, a two-round RT-PCR was performed with 2 ng of total RNA and primers as described above. Subsequent gel analysis did not detect unspecific products. The primers for different targets were located on different exons to prevent amplification of genomic DNA. Omission of the RT-enzyme and substitution of template by bath solution served as negative controls for reverse transcription and PCR amplification and confirmed the specificity of the reaction.

Electron microscopy

Acute hippocampal slices were prepared from juvenile (p9–12) hGFAP-EGFP mice. Weakly fluorescent cells with a typical electrophysiological current-pattern (previously termed GluR cells; [18]) were filled with biocytin (0.5%) via the patch-pipette during whole-cell recording. Slices were then fixed for 2 h in a solution containing paraformaldehyde (PFA) and glutaraldehyde (2% each)
in 0.1 M phosphate buffer, PB). Fixation delay after decapitation ranged from 45–120 min. Slices containing a biocytin-filled cell were rinsed, cryoprotected in sucrose solution (30% in PB), snap-frozen in liquid nitrogen and thawed [50]. Cells were visualized for correlating light and electron microscopy by overnight incubation in a combination of avidin-biotin complex (1:100, Vector, Burlingame, USA; [59]) and streptavidin-CY3 (1:1,000, Vector). After rinsing, the biocytin-filled cells were overlapped in PB and documented by recording image z-stacks under a fluorescence microscope. Subsequently, the peroxidase was developed by diaminobenzidine (DAB) and 0.07% H2O2 for ultrastructural staining. Sections were osmicated (1% OsO4), block stained (1% uranyl acetate in 70% ethanol), dehydrated and flat embedded in Araldite. Ultrathin sections were contrasted with lead citrate and uranyl acetate. To analyze overall synaptic contacts on NG2 cells at the ultrastructural level, these flat embedded cells were completely sectioned. Inspecting all ultrathin sections from a given cell, the complete process tree was scanned for synapses on DAB-containing profiles. Most synapses found in one section could also be documented in subsequent sections. To estimate the total number of synapses, the observed number of synapses was documented (Table 1) and then multiplied by 1.75 (1+0.5+0.25). An estimated factor of 0.5 was introduced to account for the missed, nearly tangentially sectioned synapses above and below a DAB-labeled profile. This corresponds to missing unrecognized synaptic profiles which are obliquely sectioned between 30 and 0 degrees (tangential). Further, we amply estimated to have missed unrecognized synaptic profiles which are obliquely sectioned between 30 and 0 degrees (tangential). Further, we amply estimated to have missed, nearly tangentially sectioned synapses above and below a DAB-labeled profile. This corresponds to missing unrecognized synaptic profiles which are obliquely sectioned between 30 and 0 degrees (tangential). Further, we amply estimated to have overlooked ¼ of the NG2 cell profiles, because most synapse-bearing profiles were below 0.3 μm (comp. Figs. 1 C, E), which was corrected for by a factor 0.25.

Dissociation of NG2 cells

Unequivocal determination of antigen presence in the NG2 cell processes is hampered by light microscopic resolution because they are frequently only 200–500 nm thick. We either studied freshly dissociated NG2 cells by conventional immunofluorescence or NG2 cells in brain slices using deconvolution microscopy with higher resolution.

The isolation method applied adapts previous cell-isolation protocols [60–63] to permit dissociation of glial cells within 2–3 h with morphological preservation of their thin processes. Briefly, hGFAP/EGFP mice at p13–15 were anaesthetized using isoflurane and decapitated. Cortical vibratome sections were incubated for 10 min at 37°C in papain solution (20 units/ml papain, 1 mM L-cysteine, 0.5 mM ethylenediaminetetraacetate (EDTA) in Ca2+/Mg2+-containing EBSS, Worthington Biochemical Corporation, Lakewood, USA). Subsequently, sections were disaggregated using pipettes, centrifuged, and resuspended in inhibitor solution (1 mg/ml ovomucoid, 1 mg/ml BSA, 0.0005% DNase I in Ca2+/Mg2+-containing EBSS, Worthington Biochemical Corporation). Finally, the cells were centrifuged onto silane-coated slides and immediately fixed with 4% PFA.

Immunofluorescence and microscopy

Dissociated cells on slides were quadruple-stained; incubation was with a mixture of the three primary or secondary antibodies according to standard procedures. The primary antibodies were chicken anti-GFAP (1:300, Chemicon/Millipore, Billerica, USA), sheep anti-GFP (1:4,000, Serotec, Düsseldorf, Germany), and a label for the protein of interest, viz. mouse anti-α-tubulin (1:500, Sigma, Deisenhofen, Germany), mouse anti-β-tubulin (1:500, Sigma), mouse anti-β-actin (1:500, Sigma), or rabbit anti-PDI (1:200; Stressgen, Assay Designs, Ann Arbor, USA). For cell identification and nucleus localization, AMCA-coupled donkey anti-chicken (1:100) and dylight488-coupled donkey anti-sheep (1:100) were combined with bisbenzimidine (1:200,000). For visualization of the antigen of interest, cells were incubated with CY3 coupled to donkey anti-rabbit (1:250) or anti-mouse (1:250). NG2 cell identification was based on morphology (small soma, multiple, very thin processes directly emanating from the soma), presence of staining with anti-GFP but absence of staining with anti-GFAP [18]. GFAP-positive nearby astrocytes served as a positive control. These specimens were documented using a fluorescence microscope (Axioskop, Zeiss), controlled by Metaview software (Molecular Devices, Sunnyvale, USA) and equipped with 100×1.3, and 40×0.75 (Plan-Neofluar) lenses, and a 4 MP b/w camera (Spot Insight, KAI+2110; Diagnostic Instruments, Sterling Heights, USA).

To detect putative glutamate vesicles in NG2 cells in situ, vGLUT1 or vGLUT2 immunofluorescence was combined with fluorescence detection of biocytin-filled NG2 cells. Cells were identified and filled as above, and fixed in 4% PFA (in PB, 2 h). After freeze-thawing, the sections were incubated with streptavidin-CY3 (1:1,000, overnight). Subsequent immunostaining was carried out by incubating sequentially with normal goat serum (10% in PB including 0.2% Triton X100, 30 min), rabbit anti-vGLUT2 (1:2,000 including 0.2% TritonX100, overnight, Synaptic Systems, Gottingen, Germany), and goat anti-rabbit-Alexa 647 (1:100, Invitrogen). For visualization of vGLUT1, only rabbit anti-vGLUT1 directly coupled to Oyster-645 (1:200, Synaptic Systems) was applied overnight.

Detection of vGLUT-IR in NG2 cells is challenging because it is abundant and dense in brain, and NG2 cell processes are frequently thinner than 0.5 μm, as observed in the electron microscope (cf. Fig. 1C). We carried out subresolution microscopy on an appropriate microscopy setup (Zeiss 200M; Orca AG camera, Hamamatsu, Hamamatsu City, Shizuoka, Japan; Openlab software, Improvision, Coventry, UK; 40×1.3, 63×1.4, 100×1.45 oil immersion lenses, Zeiss). We applied on-chip magnification (100–160x), imaging the cells at 50–100 nm steps in two fluorescence channels (filter sets [I] ex 475/20, bp 495, em 513/17 and [II] 632/22, 660, 700/75). The resulting image stacks underwent iterative deconvolution (Openslabs) based on calculated point spread function that has previously been applied and validated for antigen colocalization in single vesicles [40,64]. Image analysis and 3D reconstruction (Openslabs) included intensity thresholding in both channels. In particular, intensity thresholding in the vGLUT channel was rigorous and led to disappearance of most smaller vGLUT-positive puncta, with many false negatives to avoid false positives. Thresholding in the GFP channel frequently resulted in discontinuous glial cell processes. Post hoc exclusion of all vGLUT-IR outside the cell facilitated visualization. All instances of vGLUT-IR within in the glial cells were checked for full inclusion in 3D cardbox view (see Fig. 9). No vGLUT-IR was detected in controls without primary antibody. Further processing of electron or light microscopic images was done with Photoshop (Adobe Systems), and comprised only linear operations for optimizing brightness and contrast, but no selective processing of image detail.

Supporting Information

Figure S1 Microtubules are well-preserved in the processes of freshly dissociated, identified astrocytes. Labeling for both, cell nuclei (bisbenzimidine) and glial filaments (GFAP, Alexa 360) is revealed in the blue channel. An astrocyte (center) and two unidentified cells (right) are displayed. Microtubules (α-tubulin, red) are obvious in the astrocyte processes.
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demonstrating that the dissociation method does not interfere with microtubule integrity even in the processes.

Figure S2 Exemplary agarose gels of mRNA-transcripts for Ca₃ channel family and S100β.

Table S1 Primers used for single-cell RT-PCR.

Video S1 Demonstration of full inclusion of vGLUT1 positive objects in NG2 cell processes (3D reconstruction). The cell is the one shown in Fig. 9. NG2 cells from hippocampus (CA1) were identified by electrophysiology, biocytin-filled, fixed and visualized by streptavidin CY3 (red channel). The green channel displays immunocytochemical detection of vGLUT1. For clarity, all vGLUT staining outside the cells has been removed. After deconvolution of 75 nm optical sections, the movies demonstrate full inclusion of the vGLUT1 objects in the small processes (<0.5 μm, often 0.2 μm). Unit of the 3D grid scale: 5.5 μm.

Video S2 Elongation of an NG2 cell process. (cf. Fig. 10B). Two-photon time-lapse video was obtained from Alexa-594 dye-loaded NG2/EYFP cell processes located in an acute brain slice. Optical stacks of 20 planes were recorded every 34 s. Maximum z-projections are shown with 1 frame per second (volume 16 × 14 × 5 μm, total time 330 s, aCSF, 35 C).

Video S3 Retraction of an NG2 cell process and movement of intracellular varicosities. (cf. Fig. 10C, D). Similar recording parameters as in Video S2 were used.

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Author Contributions

Conceived and designed the experiments: CH AD CS RJ. Performed the experiments: CH AW JH JP KK RJ. Analyzed the data: CH AD AW JH JP KK JT GS MF CS RJ. Contributed reagents/materials/analysis tools: JT RJ. Wrote the paper: AD GS RJ.

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