Systematic Quantification of Synapses in Primary Neuronal Culture

HIGHLIGHTS
- Synapse antibody library screen using segmentation-independent image analysis
- Colocalization of pre- and postsynaptic markers partially reveals mature synapses
- Transsynaptic proximity ligation assay enhances sensitivity
- Guidelines for synapse-oriented in vitro screening assays

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Systematic Quantification of Synapses in Primary Neuronal Culture

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SUMMARY

Most neurological disorders display impaired synaptic connectivity. Hence, modulation of synapse formation may have therapeutic relevance. However, the high density and small size of synapses complicate their quantification. To improve synapse-oriented screens, we analyzed the labeling performance of synapse-targeting antibodies on neuronal cell cultures using segmentation-independent image analysis based on sliding window correlation. When assessing pairwise colocalization, a common readout for mature synapses, overlap was incomplete and confounded by spurious signals. To circumvent this, we implemented a proximity ligation-based approach that only leads to a signal when two markers are sufficiently close. We applied this approach to different marker combinations and demonstrate its utility for detecting synapse density changes in healthy and compromised cultures. Thus, segmentation-independent analysis and exploitation of resident protein proximity increases the sensitivity of synapse quantifications in neuronal cultures and represents a valuable extension to the analytical toolset for in vitro synapse screens.

INTRODUCTION

Synapses are the prime mediators of neuronal communication, and their plasticity defines learning and memory. They are vulnerable cell structures in many neuropathological conditions. In Alzheimer disease, for instance, soluble oligomers of amyloid beta (Aβ) and hyperphosphorylated tau localize to synaptic terminals, causing their number to decline even in the early stages of the disease (Fein et al., 2008; Pickett et al., 2016; Scheff et al., 2006). Synapse loss is recapitulated in mouse models that overexpress human amyloid precursor protein (Koffie et al., 2009; Mucke et al., 2000) and in wild-type mice that have received an intracerebroventricular injection of Aβ oligomers (Jun and Selkoe, 2015). As synapse loss is considered to adversely affect cognition, reversal or prevention thereof may represent a therapeutic strategy for neurodegenerative diseases (Jackson et al., 2019).

Discovery of novel compounds that modulate synapse density most often starts with in vitro screens. In its simplest form, screening is done on primary neuronal cultures, stained for a specific synapse marker (Evans et al., 2008; Harrill et al., 2011; Nguyen et al., 2012). Among the applications, this approach has been used to document the synaptotoxic effect of Aβ1-42, and its prevention by an oligomerization inhibitor (Evans et al., 2008). However, using a single marker to identify synapses is complicated by the presence of non-synaptic (e.g., vesicular) or degenerate synaptic structures. To focus more specifically on mature synapses, the colocalization between a pre- and postsynaptic marker has been introduced as readout (Flavell et al., 2006; Nieland et al., 2014; Paradis et al., 2007; Schatzle et al., 2012; Sharma et al., 2013; Verschuuren et al., 2019). Several gene silencing screens have made use of this approach to identify regulators of excitatory and inhibitory synapses (Nieland et al., 2014; Sharma et al., 2013). Yet, this approach is also not free from caveats, because the use of antibodies limits the sensitivity of the assay to their labeling performance. Although elegant genetic strategies for synapse labeling have been conceived (Lee et al., 2016), from a screening perspective, immunostaining is preferred for its ease of use, universal applicability, and lack of overexpression artifacts. However, validation of antibody specificity relies on knocking down target genes (Paradis et al., 2007; Sharma et al., 2013), which can be an expensive and unreliable approach suffering from complications such as off-target effects, incomplete knockdown, or alterations of cellular phenotype (e.g., cytotoxic). Hence, a more systematic validation of antibody performance that does not
require experimental interventions would be welcome. To satisfy this need, we introduced a segmentation-independent microscopic image analysis, which we applied to a set of commercially available antibodies raised against pre- and postsynaptic proteins on primary cortical and hippocampal cultures. We also quantified the degree of colocalization between pre- and postsynaptic markers and found significant differences in synapse count between different marker combinations. Finally, to improve the sensitivity and reproducibility of synapse density quantification, we implemented a transsynaptic proximity-ligation assay (PLA). Considering the unique property of PLA to only detect protein interactions at distances below 40 nm (Fredriksson et al., 2002), we identified antibodies that label bona fide partners at the pre- and postsynaptic sides obviating the need for overexpression of myc-tagged variants as was recently reported (Dore et al., 2020). We show that transsynaptic PLA has the specificity and sensitivity necessary to provide reliable synapse counts for in vitro experiments aimed at synaptic modulation.

RESULTS
Not all Synapse Markers can Be Used for Synapse Quantification
Synapses harbor a unique proteome for which a wealth of dedicated antibodies has been developed. Yet, despite their ample use, the specificity and applicability of many of these antibodies remains questionable. Therefore, we screened a panel of synapse-targeting antibodies for their labeling performance on hippocampal and cortical cultures at 14 days in vitro (DIV), a time point at which synaptic connections are well-established (Grabrucker et al., 2009; Ichikawa et al., 1993; Lesuisse and Martin, 2002; Verschuuren et al., 2019). Targets included proteins of presynaptic (vesicular, active zone) and postsynaptic (receptor, scaffold) compartments, as well as transsynaptic adhesion proteins (Table S1, Figures 1A and 1B). As expected, immunoblotting on bulk protein extracts had very limited predictive value for immunofluorescent (IF) staining performance because of sample denaturation (leading to [un-]masking of epitopes) and the presence of alternative isoforms (Figure S1). Upon IF, however, most antibodies yielded a punctate staining that predominantly decorated microtubule-associated protein 2 (MAP2)-positive neurites, suggesting specificity. Yet, when quantifying the absolute spot density using intensity threshold-based segmentation, several observations pointed to an imperfect labeling (Figure 1B). For instance, antibodies targeting the same antigen yielded significantly different spot densities (e.g., Neurexin1, Neuroligin1, GluN1). For other antibodies (e.g., PSD95a, Shank) there was high variability between replicates (Figure 1B for cortical and Figure S2 for hippocampal cultures). Also, the sum of excitatory (vGLUT positive) and inhibitory (vGAT positive) spots differed substantially from the total number of spots stained by a pan-presynaptic marker (Synapsin, Synaptophysin1a, Synaptophysin1b, Bassoon). Thus, not all antibodies show equal labeling performance, and the presence of spurious signals may represent a serious confounding factor for synapse density quantification.

To rule out bias arising from the synapse segmentation process, we adopted a method for segmentation-independent analysis of staining performance, which automatically determines the auto-correlation function (ACF) of individual images as a function of a lateral shift (van Steensel et al., 1996) (see Transparent Methods; Figures 1C and S3). The amplitude of the ACF was used as proxy for the local signal-to-background ratio (SBR), the full width at half max (FWHM) reported on the average spot size, whereas the spread (2 x standard deviation at dx = 50 px) around the average ACF informed on the variability between replicates. Based on the dimensions and quality of representative markers, we determined a window in which antibodies were considered to yield an optimal staining result (ACF amplitude >0.75, FWHM <11 [= expected spot size], and spread <0.08). All synaptic vesicle and active zone markers met these criteria (Figure 1D for cortical and Figure S2B for hippocampal cultures; see Figure S4 for the individual ACF plots). Except for Neuroligin1c, GluA1-4, Homer, PSD95a, and PSD95b, antibodies that target adhesion molecules, neurotransmitter receptors, or scaffold proteins yielded more blunted (amplitude <0.75; FWHM >11) and variable (spread >0.08) ACFs, indicative of lower labeling performance (Figure 1D). Antibodies that target GluA2 or SAP102 yielded exceptionally high FWHM values because they also labeled nuclei. For some inhibitory markers (Gephyrin and Neuroligin2), we observed considerable differences in the ACF of hippocampal and cortical cultures, which could point to variations in antigen abundance and inhibitory contacts between the two cell culture types (Figures 1D and S2B). We conclude from these experiments that a segmentation-independent analysis of staining performance facilitates the selection of antibodies for synapse quantification.
Genetic and Spine Labeling Strategies Reveal Partial Synaptic Localization

Although ACF analysis narrowed the selection of suitable antibodies that yielded a crisp punctate staining, it did not reveal whether the obtained signals solely represent synapses. This was exemplified by a non-

Figure 1. Not All Synapse Markers can be Used for Synapse Quantification

(A–D) (A) Markers of different functional classes considered in this work (adapted from Servier medical art); (B) Quantification of the spot density after immunostaining yields variable results even between antibodies that target the same marker (mean + SD, n = 6 wells with 15 fields/well, *p < 0.05, one-way ANOVA post-hoc Sidak’s multiple comparisons test). (C) The auto-correlation function (ACF) allows segmentation-independent evaluation of staining performance, by calculating the Pearson’s correlation coefficient between an image and its duplicate as a function of a lateral shift (sliding window). The ACF amplitude and spread report on the SBR and the variability across images, respectively, the full width at half maximum (FWHM) correlates with spot size. A crisp synapse spot image has a high amplitude (>0.75) with narrow spread (<0.08) and small FWHM (<11 μm), whereas a noisy image has a small amplitude with a large standard deviation and large FWHM. (D) Scatterplot of ACF parameters for the different synapse antibodies tested on primary cortical cultures at 14 DIV (n = 90 images originating from 6 wells), along with representative images for a selected subset (scale bar, 10 μm). Inset shows which combinations are retained (in color) after gating for optimal parameter settings. See Also Figures S1–S4.

Genetic and Spine Labeling Strategies Reveal Partial Synaptic Localization

Although ACF analysis narrowed the selection of suitable antibodies that yielded a crisp punctate staining, it did not reveal whether the obtained signals solely represent synapses. This was exemplified by a non-
specific PSD95 antibody that yielded randomly dispersed spots of a similar size than synapse spots (Figure S3E and S3F). Hence, we sought for a strategy that could provide more certainty and serve as benchmark for determining synaptic localization. To this end, we introduced a cross-correlation function (CCF), which performs the same sliding window Pearson correlation analysis as defined for ACF, but now between two different fluorescence channels instead of within a single channel. When applied to two synapse markers, the CCF amplitude reports on the degree of colocalization and the FWHM on the combined size of the markers, as evidenced by simulations (Figure 2A). Of note, a CCF between a synapse marker and a neurite marker already allows discriminating randomly dispersed non-specific spots from more specific synaptic spots (Figure S3E and S3F).

We applied the CCF analysis to PSD95-mVenus (ENABLED) transgenic mouse cultures (Fortin et al., 2014), in which endogenous excitatory postsynaptic protein PSD95 is labeled with minimal overexpression artifacts. Having noticed that the PSD95-mVenus signal increased with culture age (Figure S3), we analyzed the colocalization of PSD95-mVenus with immunostained synaptic markers at both 14 and 28 DIV (Figures 2B and S6–S8). We also measured the percentage of PSD95-mVenus spots with an overlapping synapse marker spot using a segmentation-based approach (defined as overlap coefficient [OC]). To determine the maximum level of colocalization attainable with both CCF and segmentation-based approaches, an anti-GFP antibody (which cross-reacts with mVenus) was used as a positive control. Representative images are from 28 DIV cortical cultures (scale bar, 10 μm). (C) Exemplary dendrite stretches from 14 DIV cortical cultures after co-labeling with postsynaptic synapse markers and DiI (scale bar, 10 μm). The bottom panel shows a binarized version of synapse spots within the MAP2 boundary. Quantification of the association with spines was done in both cortical and hippocampal cultures. Data are represented as mean ±SD; at least 200 spines were counted per condition.

See Also Figures S5–S8.

In a second approach to validate synaptic localization, we specifically focused on the excitatory postsynaptic compartment. This part of the synapse has a unique structure that is referred to as a dendritic spine (Segal, 2017). Spines can be readily visualized, e.g., by using lipophilic dyes such as Dil (Papa et al., 1995; Verstraelen et al., 2014). When combining Dil with postsynaptic antibody staining, we found considerable differences in colocalization between synapse markers in cultures of 14 DIV (Figure 2C). At the lower limit of the dynamic range, GABA-R puncta were rarely located on spines (6% ± 4% for cortical and 12% ± 9% for hippocampal cultures), which corresponds with findings on cortical tissue at the ultrastructural level (Kubota et al., 2016; Kwon et al., 2018). To gauge the upper limit of the dynamic range, we considered the genetic PSD95-mVenus signal and found that nearly 80% of the spots were located on spines and that 80% of all spines contained PSD95-mVenus signal. AMPA-R clusters, as labeled with GluA1-4 antibody, were partially located on dendritic spines, yet the majority of spots were found on the shaft. For both PSD95 antibodies, of which we theoretically expected spine labeling similar to PSD95-mVenus (~80%), colocalization was less than half (30%–40%) of the anticipated result. This implies that the antibodies did not label their target antigens sufficiently, as also suggested by the PSD95-mVenus colocalization measurements (Figure 3B). With ~50% of spines being labeled and ~60% of all spots residing in spines, the scaffold protein
Homer approached the upper limit of the dynamic range, proving (again) more selective for the excitatory compartment than both PSD95 antibodies.

Taken together, there is a substantial fraction of the IF signal that does not overlap with bona fide synapses, and not all synapses are efficiently labeled by antibodies. Of those antibodies tested, Homer showed the highest colocalization with the PSD95-mVenus signal and the strongest enrichment in dendritic spines, indicating that this antibody is best suited to stain the majority of the postsynaptic compartments.

**Synapse Marker Colocalization Improves Selectivity for Mature Synapses**

The previous experiments revealed that synaptic antibodies show variable labeling performance and that non-synaptic spots are present, which both complicate robust synapse quantification. Yet, using expansion microscopy, we confirmed that pre- and postsynaptic markers are often found juxtaposed in cultures at 14 DIV (Figures 3A and S9). The diffraction limit inherent to conventional fluorescence microscopy precludes signals closer than 200 nm from being resolved. With 20 nm, the synaptic cleft is well below this limit. Hence, a simple approach to detect mature synapses—i.e., synapses that dispose of a clear pre- and postsynaptic compartment—with more certainty consists in quantifying the apparent overlap between pre- and postsynaptic markers (Nieland et al., 2014; Paradis et al., 2007; Sharma et al., 2013;...
Proximity of Markers Allows Selective Labeling of Mature Synapses

As the colocalization of two markers might be confounded by the presence of spurious signals (e.g., extrasynaptic staining), we sought for an approach that would only give rise to positive signals in case pre- and postsynaptic markers are sufficiently close. We reasoned that the PLA (Leuchowius et al., 2010) would represent a plausible method to achieve this, because it only leads to a reaction when two markers (and their respective antibodies) are closer than 40 nm, a distance range that bridges the synaptic cleft (Savtchenko and Rusakov, 2007) (Figure 4A). We first validated the method using a positive control—antibodies for two nuclear envelope markers that are known to interact directly (lamin A and lamin B1) (Kapinos et al., 2010)—and a negative control—antibodies targeting lamin A and the mitochondrial marker Translocase Of Outer Mitochondrial Membrane 20 (TOMM20), which are spatially separated by much more than 40 nm. The positive control yielded nuclear-localized PLA spots, whereas the negative control produced virtually no signal, as was the case for a negative technical control in which only one of the primary antibodies was used (Figure S13A). Next, we tested whether targeting pairs of proteins present in excitatory and inhibitory synapses resulted in a transsynaptic PLA reaction. To maximize the efficiency of the PLA reaction, we selected within our panel of antibodies those that would bind markers that have been described to directly interact. From our panel, Neurexin1b was the most suitable pan-synaptic marker antibody because it interacts at the molecular level with the extracellular epitopes of four postsynaptic proteins with available antibodies, namely, GluA1-4, GABA-R, Neuroligin1c, and Neuroligin2 (Dore et al., 2020; Mondin et al., 2011; Sudhof, 2017; Zhang et al., 2010). Indeed, all four combinations yielded the characteristic punctate PLA pattern at the expected location: inhibitory synapse markers (GABA-R and Neuroligin2) preferentially located near neuronal somas, whereas excitatory PLA signals (GluA1-4 and Neuroligin1c) were found evenly distributed along the neurites (Figure 4A). The negative control—targeting vGLUT(excitatory) and Neuroligin2 (inhibitory)—yielded low PLA signal, whereas the positive control—vGLUT and Neuroligin1c (both excitatory synapse markers)—produced a strong PLA signal. We also noticed that the amount of transsynaptic PLA signal with the combination VGAT-Neuroligin1c (a discordant excitatory-inhibitory combination) was higher than that of the combination VGAT-Neuroligin2, which should detect reaction on inhibitory synapses only (Figure S13B). ACF plots from transsynaptic PLA displayed a large amplitude (0.82–0.93) and small FWHM (7.5–10.5 px), which would set the quality of the PLA staining as one of the best performing compared with single staining antibodies (Figure S13C) and outperforming all combinations for double staining.

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Transsynaptic PLA Signals Accurately Reflect the Number of Mature Synapses but Are Spatially Displaced from the Synapse

We next investigated the subcellular location of transsynaptic PLA signal for the PLA pairs Neurexin1b/Neuroligin1c and Neurexin1b/Neuroligin2. To this end, we used PSD95-mVenus (ENABLED) neuronal cultures at 28 DIV and counterstained for the pan-synaptic marker Synapsin (Figure 4B). Transsynaptic PLA density measurements were significantly higher for Neurexin1b/Neuroligin1c than for Neurexin1b/Neuroligin2.
Neurexin1b/Neuroligin2 (0.18 ± 0.02 versus 0.12 ± 0.01 spots/μm²; two-way ANOVA, post-hoc Sidak’s test p value = 0.036), in line with the expected predominance of excitatory synapses in cortical cultures. Importantly, the densities of transsynaptic PLA spots, and PLA/Synapsin and PLA/mVenus pairs, were within the ranges obtained previously by double IF colocalization (0.05–0.2 spots/μm²) (Figure 4B). As PSD95-mVenus resides in excitatory synapses, we focused on spot densities and overlap of the transsynaptic PLA for the excitatory combination Neurexin1b/Neuroligin1c. The OC of transsynaptic PLA signals with PSD95-mVenus and Synapsin was 21.5% ± 2% and 21.3% ± 1%, respectively, indicating that less than half of the PLA signals colocalized with one of these markers. This was lower than the OC between mVenus and Synapsin (35.7% ± 4%) (Figure S14A). However, when putting the images under scrutiny, we noticed that many of the PLA spots were juxtaposed to the other markers (Figure 4B). To corroborate this, we performed a nearest neighbor correlation analysis between the distributions of detected objects of each marker pair (PLA/mVenus, PLA/Synapsin, mVenus/Synapsin). In brief, we calculated the probability that the nearest neighbor of a given spot is of a different marker type versus the probability that it could be either of both markers (random distribution), taking into account the spatial constraints of the neurite mask. Thus, if both markers were distributed in a completely independent manner, the resulting nearest neighbor ratio value (nncorr) would be 1, whereas a completely correlated set would have a ratio value close to 0 (Figure S14B) (Baddeley and Turner, 2005). We found that the nncorr measures for transsynaptic PLA combinations were below 0.2, suggesting strong dependence with the other markers, as also confirmed by positive (Homer-Bassoon) and negative (vGAT-PSD95a) controls (Figure S14B).

To obtain a more resolved view on the actual distances between the PLA signals and the other markers, we employed super-resolution radial fluctuations (SRRF) microscopy (Figure 4C). The SRRF algorithm calculates the likelihood of the true position of a fluorophore by measuring radial symmetries informed by successive acquisitions of the same field. Once image segmentation and object detection for each channel on SRRF images was performed, we expanded the individual spots by 10 pixels (~230 nm) to account for the increased resolution and searched for overlapping objects within that distance. This way, we found that 95% of the transsynaptic PLA signals were closer than 230 nm to Synapsin spots (Figures 4C and S15A). The average number of Synapsin spots associated with every transsynaptic PLA spot was 0.95 ± 0.1, which suggests that PLA spots are in a 1:1 ratio with synapses. When quantifying the spot density in SRRF images, we found that the density of PSD95-mVenus and Synapsin spots was 0.5 ± 0.1 and 1.1 ± 0.2 spots/μm², respectively, whereas that of transsynaptic PLA (or PLA in proximity with other marker combinations) was ≤0.25 spots/μm², similar to IF measurements (Figure S15B). Taking advantage of the increased resolution in SRRF images, we measured the average distance between the nearest neighbors from pairs of all marker types. This confirmed that PLA spots were always found in closer proximity to mVenus (0.49 μm) or Synapsin (0.54 μm) spots than to other PLA spots (Figure 4C). Despite our restriction to optical sections of 0.4 μm, triplets of transsynaptic PLA, mVenus, and Synapsin were frequently encountered at a density of 0.13 spots/μm² and accounted for 46% ± 12% of total transsynaptic PLA particles (Figure S15). Thus, the density of transsynaptic PLA spots aligns well with that of the best-performing transsynaptic marker combinations, although they occupy a spatial location that is slightly offset from the actual synapses.

Transsynaptic PLA Increases Sensitivity for Detecting Synapse Density Changes

Having previously shown that synapse density scales with culture time (Verschuuren et al., 2019), we decided to further validate the transsynaptic PLA on a set of cultures of increasing maturity (7, 14, and 21 DIV). We thereby used the validated Neurexin1b–Neuroligin1c combination (as direct interactors), as well as the Bassoon-Homer combination (as best antibody pair), and directly compared PLA with the colocalization approach. As expected, we found an increase in synapse density for the Neurexin1b–Neuroligin1c and also for the Bassoon-Homer combination, suggesting that it is likely that most pre-postsynaptic partners are still in sufficiently close proximity for the PLA to work (Figure 5A). Moreover, observed synapse densities were consistently higher (2 up to ~8-fold) for PLA than colocalization, suggesting increased sensitivity for the former. To further investigate the differential sensitivity, we tested whether transsynaptic PLA could also pick up changes in synapse density as evoked by pathological conditions. We previously found that overexpression of hTau-P301L reduces neuronal connectivity from 14 DIV onwards (Verschuuren et al., 2019). Therefore, we measured synaptic density at 21 DIV after applying increasing multiplicities of infection (MOI 200, 300, 400, and 500) of the hTau-P301L construct. For PLA with both antibody combinations, a concentration-dependent decrease of the synapse density was detected with the highest concentration (MOI 500) becoming statistically significant (two-way ANOVA p < 0.001 and p < 0.05 in Sidak’s post-hoc test with MOI 0 within staining type). This dose-dependent decrease was not detected by the colocalization approach. In addition, colocalization of Neurexin1b and Neuroligin1c yielded highly variable

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Thus, these results suggest that transsynaptic PLA allows detecting changes in synapse density in a more sensitive and reproducible manner than a colocalization approach.

**DISCUSSION**

A wealth of studies has used IF to quantify synapse density changes in primary neuronal cultures (Evans et al., 2008; Flavell et al., 2006; Hagmeyer et al., 2015; Harrill et al., 2011; Jin and Selkoe, 2015; Nguyen et al., 2012; Nieland et al., 2014; Paradis et al., 2007; Schatzle et al., 2012; Sharma et al., 2013; Verschuuren et al., 2019; Verstraelen et al., 2014). Yet, very few have truly documented the labeling performance of the utilized antibodies in detail. Labeling performance encompasses different features such as specificity and sensitivity. A specific antibody binds its antigen with minimal cross-reactivity, whereas a sensitive one will bind a high fraction of the available antigens. Experimental assessment of both parameters is not trivial as every experimental intervention to identify putative errors (such as a knockout or overexpression approach) is bound to influence cell physiology. Furthermore, an antibody may have a high score for both parameters, but may not be compatible with synapse screens, when it labels an antigen that is not sufficiently enriched at the structure of interest. That is why we have chosen to evaluate labeling performance as a whole from an application perspective. Although extensively used (Signore and Reeder, 2012), we found that western blotting is not useful for routine antibody validation because neither the presence of additional bands (alternative splice forms) nor the absence of a specific band (epitope masking) can truly predict non-specificity. Therefore, we first introduced a segmentation-independent method to assess the staining performance of individual antibodies in IF images. At the magnification and resolution used, the method does not inform on size, shape, or orientation of individual synapse spots. Instead, it capitalizes on the high density and heterogeneous orientation of the labeled structures in the field of view, to give a reliable indication of the SBR and average spot size at the population level. These analyses revealed a large variability between antibodies, which partially depended on the functional class to which these markers belong, i.e., vesicle markers, markers associated with the active zone of the presynapse, adhesion proteins, neurotransmitter receptors, and scaffold markers. Synaptic vesicle markers yielded consistently better results in ACF analysis than neurotransmitter receptors or adhesion proteins, likely because the local protein abundance is higher for the former. We further found that the absolute spot density (as quantified by

**Figure 5. Transsynaptic PLA Increases Sensitivity for Detecting Synapse Density Changes**

(A and B) (A) Representative images and quantification of synapse density in cortical neuronal cultures at 7, 14, and 21 DIV, after transsynaptic PLA or colocalization with Neurexin1b/Neuroligin1c and Bassoon/Homer combinations (scale bar, 10 μm). (B) Representative images and quantification of synapse density at 21 DIV after infection with a hTau-P301L AAV vector (scale bar, 10 μm). Infection with increasing multiplicities of infection (MOI) reveals a concentration-dependent decrease that is detected by the PLA but not the colocalization approach (*p < 0.05 in post-hoc Sidak’s test with MOI 0 within staining type). Neurexin1b-Neuroligin1c colocalization is considered inaccurate given the suboptimal IF labeling performance (Figure 1D) and consequent high spot count variability; (A–B) mean ± SD; n = 4 wells for Bassoon-Homer PLA, n = 8 wells for Neurexin1b-Neuroligin1c PLA, n = 6 wells for colocalization—always 15 fields/well).
segmentation-dependent analyses) was highly variable even for antibodies that target the same marker. This may be caused by non-specific labeling, low signal-to-noise ratio complicating thresholding, and the presence of spurious signals such as markers being trafficked in the neurites or extrasynaptic neurotransmitter receptors. These caveats do not necessarily preclude relative quantification in a setting where experimental conditions are compared with an internal control (Evans et al., 2008; Harrill et al., 2011; Nguyen et al., 2012), but they may significantly affect sensitivity and reproducibility, and also render the assay susceptible to confounding factors such as density differences (e.g., due to neurotoxicity). By quantifying the association with the postsynaptic compartment, we discovered that Homer outperformed PSD95 antibodies, in terms of its overlap with the genetic marker PSD95-mVenus and its presence in dendritic spines. Surprisingly, and in contrast with other reports, we could not recapitulate equally good results with PSD95 antibodies (Lagache et al., 2018). This may be due to the different species (rat versus mouse primary hippocampal cultures) or cultivation protocol (Lagache et al. used astrocyte-conditioned medium). Another explanation may be that Homer, as a polyclonal antibody, is more resistant to epitope masking than the PSD95 monoclonals. As exponent, the GluA1–4 antibody showed remarkably low correlation with the postsynaptic compartment, suggesting that many AMPA-R clusters reside outside of the synapse, potentially in reserve pools awaiting synaptic potentiation (Kneussel and Hausrat, 2016).

We further found that the selectivity for mature synapses could be enhanced by considering the apparent colocalization between a pre- and postsynaptic marker, while maintaining high-content screening compatibility. The positive results from ACF analyses for Bassoon and Homer antibodies were further substantiated by their high colocalization metrics (CCF amplitude >0.45, OC > 30%). This makes the latter combination preferred for global assessment of structural connectivity.

Although colocalization helps focusing on the relevant fraction of immunolabeled samples, it is subject to bias and suffers from a limited dynamic range. Therefore, we explored whether PLA could serve as a more sensitive alternative. PLA had been used before to confirm synaptic interactions in the pre- or post-synaptic compartment (Almandoz-Gil et al., 2018; Eagleson et al., 2013; Lundgren et al., 2013). Very recently, the utility of PLA for monitoring transsynaptic interactions was demonstrated as well (Dore et al., 2020). In this work, co-cultures of neurons were used in which one population was transfected with myc-tagged Neurexin and another with hemagglutinin-tagged Neuroligin. While physiological interventions were used to illustrate the potential of the PLA approach in reporting changes in synapse plasticity, the overexpression of synaptic markers may perturb the delicate balance in the synapse. Furthermore, transfection may not be equally efficient for all cellular models, and mature synapses will only be detected between differentially transfected partner cells. Therefore, it has limited applicability for high-throughput screens that aim at interrogating synaptic modulators across neuronal networks. That is why we sought an approach applicable to endogenous markers, using validated antibodies. Our results demonstrate that transsynaptic PLA has advantages over classical pre-post marker colocalization. First, it allows for simpler signal detection, as also proven by ACF analysis. Importantly, we found that on average each transsynaptic PLA spot had ~1 Synapsin spot, which supports the notion that transsynaptic PLA is a reliable proxy for synapse estimates. Second, using one single marker for synaptic connections liberates one channel for additional staining. Finally, our results of synapse quantification in cultures through stages of maturation and pathological perturbations strongly endorse the potential of transsynaptic PLA being more sensitive and thus capable of detecting subtle changes in synapse number. This is especially relevant for dense cultures, in which crowding may complicate signal detection. PLA reported a much more consistent and higher synapse density for different marker combinations than colocalization did. Remarkably, despite lesser IF performance (e.g., AFC metrics), the PLA combination for Neurexin1b-Neuroligin1 performed equally well as Homer-Bassoon. Neurexin1b was even shown to be homogeneously expressed at the dendritic surface (Dore et al., 2020) reducing its SBR below a standard required for spot detection. This suggests that PLA is able to discriminate the synaptic pool by virtue of its proximity dependence.

Notwithstanding its advantages, it should be mentioned that transsynaptic PLA displayed some unexpected characteristics and potential drawbacks. For one, there is the offset we encountered between PLA signals and antibody labeling. This may be explained by the effect of the amplification protocol or the characteristics of the antibody-bound DNA probes that produce a reaction adjacent to the synapse. However, it should be kept in mind, when aiming at performing spatial proteomics. Another potential issue is the detection of PLA signal in controls that were initially considered to be negative (such as VGAT and Neuroligin1c). Yet, it is becoming clearer that the histological definition of inhibitory versus excitatory
synapses is not clear-cut. For example, in cortical culture, electron microscopy of dendritic spines, once thought to be exclusive of excitatory synapses, also contains inhibitory contacts (Kubota et al., 2016). In addition, Neuroligin1 and Neuroligin2 have been shown to colocalize with both vGAT and vGlut1, with only enrichment of each isoform on excitatory and inhibitory synapses, respectively, but by no means an exclusive colocalization (Chih et al., 2006).

In summary, we have introduced a workflow to validate antibodies that target synapse markers, which can be extended to additional culture types (primary cultures originating from other brain regions, induced pluripotent stem cell [iPSC] derived, etc.). In doing so, we found combinations of pre- and postsynaptic markers that report on synapse density in cortical and hippocampal cultures, of which Bassoon and Homer offered the largest dynamic range. Building further on the concept of marker proximity, we demonstrated that transsynaptic PLA using endogenous markers can offer an even more sensitive readout than synapse detection by marker colocalization. Our work therefore contributes to the refinement of synapse-oriented in vitro screening assays.

Limitations of the Study

1) We have screened a small library of antibodies for their labeling performance in primary cortical and hippocampal cultures and revealed variability between markers, culture types, and maturity stages. We have not shown the exact origin of this variability (antigen turnover, location heterogeneity, etc.) as the very gist of our work was to identify the best possible strategy for labeling mature synapses with an eye on high-throughput screening.

2) We have used the same staining protocol for all antibodies. Optimizing blocking conditions might increase the staining performance of certain antibodies.

3) Although the ACF/CCF method is much less prone to bias than spot segmentation based on a user-defined threshold, it only provides information at the population level. It does not report on size, orientation, or shape of individual synapses and the readout depends on the image resolution.

4) We have introduced an approach for gauging labeling performance, which may serve for validation of additional markers, well beyond the scope of this work. However, we also emphasize the need for multiple validation and benchmarking methods. In doing so, we also reveal limitations of current gold standards such as transgenic reporters or spine labeling.

5) We show that transsynaptic PLA is a sensitive and screening-format-compatible method. Although it liberates a channel for multiplexing, the increased cost and faster degradation of the PLA signal may currently limit its applicability. We also observed a physical displacement of signal, which we attribute to the amplification protocol but do not prove.

6) We have considered primary cortical and hippocampal cultures from mouse. Alternate models such as rat cultures or iPSC-derived human cultures may require re-assessment.

Resource Availability

Lead Contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Winnok H. De Vos (winnok.devos@uantwerpen.be)

Materials Availability

This study did not generate new unique reagents.

Data and Code Availability

The datasets generated during this study are available from the corresponding author on request. The image analysis script for spot segmentation in Acapella (PerkinElmer) is available on Github (https://github.com/VerschuurenM/NeuronalConnectivity), and the Fiji scripts used in this study are available from the corresponding author on request.

METHODS

All methods can be found in the accompanying Transparent Methods supplemental file.
SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2020.101542.

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AUTHOR CONTRIBUTIONS

Conceptualization, P.V., G.G.-D.B., and W.H.D.V.; Methodology, P.V., G.G.-D.B., and W.H.D.V.; Investigation, P.V., G.G.-D.B., and B.A.; Software, M.V., G.G.-D.B., and W.H.D.V.; Visualization, P.V., G.G.-D.B., and W.H.D.V.; Writing – Original Draft, P.V., G.G.-D.B., and W.H.D.V.; Writing – Review & Editing, P.V., G.G.-D.B., and W.H.D.V.; Supervision and Funding Acquisition, W.H.D.V., P.L., J.-P.T., and R.N.

DECLARATION OF INTERESTS

Authors R.N. and P.L. are employees of Janssen Pharmaceutica NV, yet all authors declare no competing interests.

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Supplemental Information

Systematic Quantification of Synapses
in Primary Neuronal Culture

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Figure S1. Antibody validation by western blot. (Related to Fig. 1)
Antibodies were tested on lysates of 14 DIV primary hippocampal (left lane) and cortical (right lane) cultures. The molecular weight marker in the visible spectrum is shown on the left. An arrowhead indicates the height of the predicted molecular weight of each marker and is shown in green or red when the predicted band is present or absent, respectively. Additional or missing bands may be the result of alternative isoforms or (un-)masked epitopes due to sample denaturation, which makes the predictive value of immunoblotting for IF performance very weak.
Figure S2. Synapse marker evaluation in 14 DIV primary hippocampal cultures. (Related to Fig. 1)
(A) Quantification of the spot density by manual thresholding reveals variable results (Mean + SD, n=6 wells with 15 fields/well, *p<0.05, one-way ANOVA post-hoc Sidak's multiple comparisons test); (B) Synapse antibody evaluation in 14 DIV primary hippocampal cultures. A scatter plot of ACF-derived parameters (and inset with gating on optimal parameter conditions) along with representative images are shown (n=90 images originating from 6 wells, scale bar 10 µm).
Figure S3. Segmentation-independent analysis of primary neuronal networks. (Related to Fig. 1) (A-C) ACF/CCF readout depends on the image resolution. (A) ACF and CCF plots for all channel combinations of DAPI (nuclei) / MAP2 (neurites) / Bassoon (Pre) / Homer (Post)-stained primary cortical cultures, acquired with different objective lenses using the same sliding window of 50 pixels. For all markers, the FWHM (in µm) increases
with lower numerical aperture due to the broadening of the point spread function. The more pronounced increase in FWHM for the nuclei channel is most likely due to the chromocenters becoming less resolved and contrasted at lower resolution, causing them to now longer be picked up as individual entities. The amplitude remains largely unchanged, except for the nuclei channel which is due to the sliding window not capturing the whole size of the structure at higher magnifications (n=90 images originating from 6 wells); (B) Scatter plot of the amplitude and FWHM (±std) derived from the ACF/CCF plots shown in panel A; (C) Crops of 30x30 µm from representative images acquired with different objective lenses (scale bar 5 µm); (D) The direction of the lateral shift does not affect the ACF/CCF readout as shown by the non-significant differences in amplitude and FWHM for horizontal vs. vertical ACF / CCF of three marker combinations with different overlap (n=90 images originating from 6 wells); (E) ACF and CCF plots of primary cortical cultures stained for MAP2, Synapsin and one of the above-mentioned postsynaptic markers. In contrast with Homer and PSD95a antibodies, one antibody (PSD95 abcam ab2723) yields non-specific puncta that are uniformly distributed across the image and have a size comparable to that of synaptic spots. This results in a seemingly ‘good’ ACF readout (green asterisk), yet CCFs indicate that the colocalization with neurites and presynaptic spots is lower than expected (red asterisks), allowing identification of this non-specific antibody (n=90 images originating from 6 wells); (F) Microscopic images confirm the lack of colocalization of the non-specific PSD95 antibody with neurites and the presynaptic marker (scale bar 20 µm).
Figure S4. Labeling specificity of synapse marker antibodies. (Related to Fig. 1)
IF results of all tested antibodies on 14 DIV primary cortical and hippocampal cultures are shown, subdivided into five different functional classes (see Fig. 1a). A representative image of MAP2 (red) with the synapse marker (cyan - contrast optimized) is shown along with the auto-correlation function (ACF) of the image dataset, which served for extracting amplitude, spread and FWHM in Fig. 1 (n=90 images originating from 6 wells, scale bar 20 µm). The average ACF of the functional class is superimposed in red.
Figure S5. PSD95-mVenus signal evolves between 14 and 28 DIV. (Related to Fig. 2)
Representative images of 14 and 28 DIV cortical cultures. The contrast-matched images reveal a marked increase in the overall intensity of the PSD95-mVenus signal between 14 and 28 DIV (scale bar 50 µm). The insets, shown with optimized contrast settings per DIV, show a larger difference between back- and foreground for 28 as compared to 14 DIV (scale bar 10 µm), leading to an ACF with higher amplitude (n=90 images originating from 6 wells). Note that the ACF relies on the Pearson’s Correlation Coefficient (PCC) which is independent of the average intensity.
Figure S6. Colocalization of synapse marker antibodies with PSD95-mVenus in primary hippocampal cultures. (Related to Fig. 2)

The colocalization is shown as a scatter plot of the CCF-derived parameters amplitude (reporting on colocalization) and FWHM (reporting on combined size), and colored by the overlap coefficient (OC), defined as the percentage of PSD95-mVenus spots that have an overlapping antibody spot (n=90 images originating from 6 wells). Images are from 28 DIV hippocampal cultures (scale bar 10 µm).
Figure S7. Auto- (A) and cross-correlation (B) functions of PSD95-mVenus and synapse markers in cortical cultures (n=90 images originating from 6 wells). (Related to Fig. 2)
Figure S8. Auto- (A, B) and cross-correlation (C) functions of PSD95-mVenus and synapse markers in hippocampal cultures (n=90 images originating from 6 wells). (Related to Fig. 2)
Figure S9. Measurement of the expansion factor in expansion microscopy. (Related to Fig. 3)
HeLa cells were stained for α-tubulin using standard ICC protocols and expansion microscopy was performed using the protocol described by Chozinski et al. (2016). Before digestion of the gel, the samples were imaged a first time. Large sample fields were tile-scanned with a 10x objective to have a topographical overview of the sample. Next, a number of cells were randomly selected within the sample and imaged using a 63x 1.4 NA objective (see ‘pre-ex’ image as an example; scale bar 25 µm). After the expansion, the same cells were relocated based on the topographical information at low magnification, and cells were reimaged using 63x 1.3 WI objective (see ‘post-ex’ image for the same example cell; scale bar 100 µm). The green-magenta overlay shows the registration of both images (‘Similarity’ method in ImageJ; translation + rotation + isotropic scaling, no warping/deformation). To quantify the expansion factor, clearly distinguishable local subcellular hallmark features were manually selected on both the pre- and post-expansion images of the same cells, such as microtubule ends, microtubule crossings or bright signal accumulations, and the distances between those points were measured. The data presented in the scatter plot is from 7 cells originating from two separately expanded samples. The expansion factor for this dataset, as given by the slope of the linear regression line, was 3.98.
Figure S10. Double labeling of synapse markers in 14 DIV hippocampal cultures. (Related to Fig. 3)
(A) Quantification of spot and synapse density in 14 DIV cortical cultures shows the limited fraction of colocalized signals (Mean + SD, n=6 wells with 15 fields/well); (B) Representative images (scale bar 10 µm) and a scatter plot of CCF parameters reporting on the colocalization of marker pairs in 14 DIV hippocampal neurons (n=90 images originating from 6 wells). The color code indicates the overlap coefficient, defined as the percentage of spots that reside in synapses. Pairs of inhibitory with excitatory synapse markers were considered as negative controls, while two pan-presynaptic markers and primary antibodies raised in the same species were used as positive controls.
Figure S11. Cross-correlation functions of double stainings in cortical and hippocampal cultures. (Related to Fig. 3) CCFs showing the colocalization (amplitude) and combined length (FWHM) of synapse marker combinations in 14 DIV cortical (A) and hippocampal (B) cultures (n=90 images originating from 6 wells).

Color code (antibody species): Mouse, Rabbit, Guinea Pig
Figure S12. Spot quantification of double stainings with low and high colocalization. (Related to Fig. 3)
(A) The combination of an inhibitory presynaptic (vGAT) with excitatory postsynaptic marker (PSD95a) yields a compact CCF and low OC after segmentation. (B) In contrast, a positive control in which two rabbit primary antibodies are used, yields a CCF that approximates 1 at $\Delta x = 0$ (nearly perfect colocalization), while the OC is still only 56%, displaying the better sensitivity of the segmentation-independent CCF approach. (A-B) n=90 images originating from 6 wells, scale bar 5 µm.
Figure S13. PLA controls and characteristics. (Related to Fig. 4)
(A) Positive and negative technical and biological controls show the specificity of PLA in neurons, when comparing the amount of signal and the location of known interacting nuclear proteins (Lamin A / Lamin B1) with proximal but out-of-PLA-reach markers (Lamin A / TOMM20) as well as the technical control of the omission of a singular primary antibody (scale bar 100 µm); (B) Synaptic PLA with permutations of inhibitory and excitatory markers show that congruent combinations produce increased numbers of PLA spots (mean + SD, n=4 wells with 15 images/well, scale bar 100 µm); (C) ACF plots for PLA signal from specified markers (n=60 images originating from 4 wells); and scatterplot location of synaptic PLA marker combinations as compared with single-staining ACF properties (right, cfr. also Fig. 1).
Figure S14. Analyses of PLA triple staining (trans-synaptic PLA-Neurexin1b/Neuroligin1c, mVenus-PSD95 and synapsin) of cortical 28 DIV cultures. (Related to Fig. 4)

(A) Quantification graph of overlap percentage for the specified pairs of spots derived from the segmentation of confocal images. Values are the percentage of each pair against the total number by marker (e.g., PLA – mVenus is the percentage of PLA spots that have an overlapping mVenus signal); (B) Nearest neighbor correlation schematic and quantification. \( \text{nncorr} \) measures the relationship between two types of spots by calculating the proportion of spots that have a nearest neighbor of the same type, and normalizing by what would be expected from an independent (random) distribution; thus values close to 1 suggest independence between the sets of spots, whereas a value closer to 0 is expected if spots are correlated. PLA triple staining pairs (mVenus-PLA, PLA-synapsin and mVenus-synapsin), present lower \( \text{nncorr} \) values implying that spots are correlated. This was also validated by comparing positive (Homer/Basoon) and negative (VGAT/PSD95a) control IF combinations. Note that in the latter case a different microscopy setup was used, leading to allover higher values (n=4 wells for PLA, n=6 wells for IF 15 field images per well).
Figure S15. Quantification of spots from SRRF images with triple staining (trans-synaptic PLA, Synapsin and mVenus-PSD95). (Related to Fig. 4)

(A) Quantification of proximity for the specified spots. Spot objects were identified in SRRF images of a single confocal plane of cortical 28 DIV cultures and were then expanded by 10 pixels (~230nm) to identify juxtaposed spots. Values represent the percentage of spots with juxtaposition of another marker spot versus the total number of spots (e.g., PLA – mVenus is the percentage of PLA spots that have an mVenus spot within 230 nm); (B) Spot density for each staining, pairwise combinations and triplets, derived from SRRF images on PLA triple stained cortical 28 DIV cultures (n=4-7 images stacks per PLA combination, 200 frames per image were captured for SRRF)
**Table S1.** Primary and Secondary antibodies employed in this study (antibodies used in PLA are marked with an asterisk) (Related to transparent methods)

**PRIMARY ANTIBODIES**

| Antigen       | Pre/Post | Type      | Localization      | Epitope            | Species/Clonality          | Clone      | Company     | Catalog       | Lot         | IF conc. | WB conc. |
|---------------|----------|-----------|-------------------|--------------------|---------------------------|------------|-------------|---------------|-------------|----------|----------|
| Bassoon       | Pre      | Pan       | Active zone       | intracellular      | Mouse Monoclonal          | L124/59    | Neuromab    | 73-491        | 463-3DH-50  | 2 µg/ml  | 3 µg/ml  |
| Neurexin1a    | Pre      | Pan       | Adhesion          | extracellular      | Mouse Monoclonal          | N170A/1    | Neuromab    | 75-216        | 447-1JH-73A | 20 µg/ml | 1 µg/ml  |
| Neurexin1b*   | Pre      | Pan       | Adhesion          | Intracellular      | Rabbit Polyclonal         | NA         | Abcam       | Ab222806      | GR3192323-13| 40 µg/ml | 1 µg/ml  |
| Synapsin1/2   | Pre      | Pan       | Vesicular         | intracellular      | Guinea Pig Polyclonal     | NA         | SySy        | 106004       | 106004/1-21 | 2 µg/ml  | 0.5 µg/ml|
| Synaptophysin1a| Pre     | Pan       | Vesicular         | Intracellular      | Guinea Pig Polyclonal     | NA         | SySy        | 101004       | 101004/2-26 | 2 µg/ml  | 0.25 µg/ml|
| Synaptophysin1b| Pre     | Pan       | Vesicular         | Intracellular      | Rabbit Polyclonal         | NA         | Abcam       | ab14692       | GR200179-2  | 2 µg/ml  | 0.65 µg/ml|
| vGAT          | Pre      | Inhibitory| Vesicular         | intracellular      | Rabbit Polyclonal         | NA         | SySy        | 131003       | 131003/1-45 | 2 µg/ml  | 0.25 µg/ml|
| vGlut1        | Pre      | Excitatory| Vesicular         | intracellular      | Rabbit Polyclonal         | NA         | SySy        | 135303       | 135303/2-63 | 2 µg/ml  | 0.25 µg/ml|
| AMPA-R (GluA1)   | Post    | Excitatory| Receptor          | extracellular      | Mouse Monoclonal          | N355/1     | Neuromab    | 75-327        | 455-5JD-89  | 2 µg/ml  | 1 µg/ml  |
| AMPA-R (GluA1-4)* | Post    | Excitatory| Receptor          | extracellular      | Mouse Monoclonal          | 248B7      | SySy        | 182411       | 182411/10   | 2 µg/ml  | 1 µg/ml  |
| AMPA-R (GluA2)   | Post    | Excitatory| Receptor          | intracellular      | Mouse Monoclonal          | L21/32     | Neuromab    | 75-002        | 455-6JD-81C | 2 µg/ml  | 1 µg/ml  |
| GABA-R (β2/3)*  | Post     | Inhibitory| Receptor          | extracellular      | Mouse Monoclonal          | 62-3G1     | Neuromab    | 75-363        | 455-2JD-15  | 2 µg/ml  | 1 µg/ml  |
| Gephyrin       | Post     | Inhibitory| Scaffold          | intracellular      | Mouse Monoclonal          | L106/93    | Neuromab    | 75-444        | 455-8JD-47  | 2 µg/ml  | 1 µg/ml  |
| Homer1         | Post     | Excitatory| Scaffold          | intracellular      | Rabbit Polyclonal         | NA         | SySy        | 160003       | 160003/31   | 2 µg/ml  | 1 µg/ml  |
| Neuroligin1a   | Post     | Excitatory| Adhesion          | intracellular      | Mouse Monoclonal          | N97A/31    | Neuromab    | 75-160        | 443-2KS-20C | 20 µg/ml | 1 µg/ml  |
| Neuroligin1b   | Post     | Excitatory| Adhesion          | extracellular      | Rabbit Polyclonal         | NA         | Almone      | ANR-035       | ANR035AN0125| 8 µg/ml  | 1 µg/ml  |
| Neuroligin1c*  | Post     | Excitatory| Adhesion          | intracellular      | Mouse Monoclonal          | A-4        | Santa Cruz  | Sc-365110    | B0618      | 4 µg/ml  | 1 µg/ml  |
| Neuroligin2*   | Post     | Inhibitory| Adhesion          | intracellular      | Mouse Monoclonal          | L107/39    | Neuromab    | 75-451        | 455-8JD-93  | 20 µg/ml | 1 µg/ml  |
| NMDA-R (GluN1a) | Post  | Excitatory| Receptor          | extracellular      | Mouse Monoclonal          | N308/48    | Neuromab    | 75-272        | 455-7JD-90B | 2 µg/ml  | 1 µg/ml  |
| NMDA-R (GluN1b) | Post   | Excitatory| Receptor          | extracellular      | Mouse Monoclonal          | M68        | SySy        | 114011       | 114011/1-24 | 2 µg/ml  | 0.25 µg/ml|
| NMDA-R (GluN2A) | Post   | Excitatory| Receptor          | extracellular      | Mouse Monoclonal          | N327/95    | Neuromab    | 75-288        | 455-7JD-90B | 2 µg/ml  | 1 µg/ml  |
| NMDA-R (GluN2B) | Post   | Excitatory| Receptor          | extracellular      | Mouse Monoclonal          | N59/20     | Neuromab    | 73-097        | 463-1DH-64  | 2 µg/ml  | 2 µg/ml  |
| PSD93         | Post     | Excitatory| Scaffold          | intracellular      | Mouse Monoclonal          | N18/30     | Neuromab    | 75-057        | 455-3JD-72  | 2 µg/ml  | 1 µg/ml  |
| PSD95a        | Post     | Excitatory| Scaffold          | intracellular      | Mouse Monoclonal          | 7E3-1B8    | ThermoFisher| MA1-046       | SA242215   | 2 µg/ml  | 0.5 µg/ml|
| PSD95b        | Post     | Excitatory| Scaffold          | intracellular      | Mouse Monoclonal          | K28/43     | Neuromab    | 75-028        | 455-7JD-22F | 2 µg/ml  | 1 µg/ml  |
| SAP102        | Post     | Excitatory| Scaffold          | intracellular      | Mouse Monoclonal          | N19/2      | Neuromab    | 75-058        | 449-1AK-85B | 2 µg/ml  | 1 µg/ml  |
| Shank 1/2/3   | Post     | Excitatory| Scaffold          | intracellular      | Mouse Monoclonal          | N23B/49    | Neuromab    | 75-089        | 443-1KS-78B | 2 µg/ml  | 1 µg/ml  |
| MAP2          | NA       | NA        | Dendrite          | intracellular      | Chicken Polyclonal        | NA         | SySy        | 188006       | 188006/1-3  | 2 µg/ml  | NA       |
| GAPDH         | NA       | NA        | Cytoplasm         | intracellular      | Mouse Monoclonal          | GT239      | GeneTex     | GTX627408    | 41323      | NA       | 1 µg/ml  |
| GFP/mVenus    | NA       | NA        | NA                | intracellular      | Alpaca Monoclonal         | +Atto594   | Chromogen   | Gba594        | 81212001AT3 | NA       | 1 µg/ml  |
| Type                  | Conjugate       | Company                  | Catalog       | Lot         | Concentration |
|----------------------|-----------------|--------------------------|---------------|-------------|---------------|
| Donkey-anti-Guinea Pig | HRP             | Jackson Immunoresearch   | 706-035-148   | 128221      | 0.1 µg/ml     |
| Donkey-anti-Mouse    | HRP             | Jackson Immunoresearch   | 715-035-151   | 125909      | 0.5 µg/ml     |
| Donkey-anti-Rabbit   | HRP             | Jackson Immunoresearch   | 711-035-152   | 104907      | 0.2 µg/ml     |
| Donkey-anti-Chicken  | AlexaFluor647   | Jackson Immunoresearch   | 703-605-155   | 134612      | 1 µg/ml       |
| Goat-anti-Mouse      | AlexaFluorPlus488 | ThermoFisher           | A32723        | SG251135    | 2 µg/ml       |
| Goat-anti-Rabbit (Fab) | FITC           | Jackson Immunoresearch   | 111-097-003   | 104632      | 0.5 µg/ml     |
| Donkey-anti-Guinea Pig | Cy3              | Jackson Immunoresearch   | 706-165-148   | 127715      | 1 µg/ml       |
| Donkey-anti-Mouse    | Cy3             | Jackson Immunoresearch   | 715-165-151   | 125797      | 1 µg/ml       |
| Goat-anti-Rabbit (Fab) | Cy3              | Jackson Immunoresearch   | 111-167-003   | 78942       | 0.5 µg/ml     |
Transparent Methods

Primary neuronal cell culture

The preparation of primary neuronal cultures was carried out in accordance with the recommendations of the ethical committee for animal experimentation of the University of Antwerp (approved ethical file 2015-54). Hippocampi and cortex were dissected from wild type C57Bl6 and PSD95-CreNABLED (Fortin et al., 2014) (purchased from the Jackson Laboratories, ref 029242) E18 mouse embryos in HEPES (7 mM)-buffered Hanks Balanced Salt Solution (HBSS-HEPES), followed by trypsin digestion (0.05%; 10 min; 37°C) and mechanical dissociation. After centrifugation (5 min at 200g), the cell pellet was resuspended in Minimal Essential Medium supplemented with 10% heat-inactivated normal horse serum and 30 mM glucose. Cells were plated in Poly-D-Lysin-coated 96-well plates (Greiner µClear) at 30,000 cells/cm² for immunocytochemistry, or in 6-well plates at 60,000 cells/cm² for western blot and kept in a humidified CO₂ incubator (37°C; 5% CO₂). After 4 hours, the medium was replaced with 150 µl B27-supplemented Neurobasal medium (NB-B27), containing Sodium Pyruvate (1 mM), Glutamax (2 mM), glucose (30 mM) and Penicillin-Streptomycin (0.5%). To suppress proliferation of non-neuronal cells, 0.5 µM arabinosyletosine was added in 25 µl NB-B27 at the third and tenth day in vitro (DIV). AAV6-mediated transduction of hTau-P301L was done at 3 DIV. Cell culture supplies were purchased from ThermoFisher (Waltham, MA, USA).

Western blotting

14 DIV cortical and hippocampal cultures were lysed using ice-cold RIPA buffer supplemented with phosphatase and protease inhibitor (HALT cocktail, ThermoFisher 78445) and 5 mM EDTA. The lysate was centrifuged (10,000g, 20 min, 4 °C) and the protein concentration of the supernatant was determined using a BCA assay (ThermoFisher 23225). Samples were denatured (70% sample, 25% LDS, 5% DTT) for 10 min at 70 °C before being loaded on a 4-12% Bis-Tris gel (ThermoFisher NP0322BOX) at 10 µg/lane. A stained ruler was included in the third and eight well (ThermoFisher 26616). The gel tank was filled with NuPage MOPS SDS running buffer and NuPage anti-oxidant, and was cooled during electrophoresis (200V, ± 1h). Proteins were transferred to a PVDF membrane using NuPage transfer buffer (30V, 1h). To check the transfer, blots were reversibly stained with 0.1% Ponceau S solution in 5% acetic acid. Blots were subsequently blocked with 5% ECL blocking solution in Tris-buffered Saline with 0.1% Tween (TBS-T). Using the rulers as landmarks, blots were cut into five pieces so that two lanes (hippocampal and cortical) with or without the ruler could be stained in one reaction. Primary antibody was applied overnight at 4 °C on a roller, followed by a TBS-T wash (3 x 5 min). Horseradish Peroxidase-coupled secondary antibodies were incubated for 2h at room temperature (RT), followed by a final TBS-T wash. All antibodies were diluted in blocking buffer and are listed in Table S1. After reconstructing the cut blots into their original positions, bioluminescent detection was performed using Immobilon Western HRP substrate (Merck Millipore WBKLS0500, 30 sec) and a Chemidoc Touch imager (Bio-Rad, Temse, Belgium). After completion, the blots were re-stained for GAPDH as loading control. The global contrast of the individual (cut) blots was adjusted with Fiji image analysis freeware (Schindelin et al., 2012). The positions of the molecular weight markers were written on the blots after overlaying the bioluminescent image with its marker (visible spectrum).

Immunofluorescence staining (IF)

Paraformaldehyde-fixed cultures (2%, 20 min, RT) were permeabilized with 1% Triton X-100 in blocking buffer (0.1% bovine serum albumin and 10% normal horse serum in PBS) for 10 min, followed by an overnight incubation with the primary antibodies (Table S1) at 4 °C in blocking buffer. After washing with PBS, secondary antibodies (Table S1) were added for 2h. Finally, 4’,6-diamidino-2-phenylindole (DAPI) was applied to the cultures for 10 min at a concentration of 2.5 µg/ml, followed by a PBS wash. Primary antibodies targeting extracellular epitopes were incubated prior to permeabilization to reduce non-specific intracellular background staining. This was followed by a PBS wash and permeabilization with antibodies targeting intracellular epitopes. All presynaptic markers were designated to the 561 nm excitation channel, while postsynaptic markers were labeled for the 488 nm channel. Secondary antibodies were kept identical where possible, depending on the species of the primary antibody.

Expansion microscopy

The protocol for expansion of the samples was adapted from (Chozinski et al., 2016). In brief, immunostained samples were crosslinked for 10 min in 0.25% glutaraldehyde in PBS. Gelation was done in a mixture of 2M NaCl, 2.5% (w/w) acrylamide, 0.15% (w/w) N,N’-methylenebisacrylamide, 8.625% (w/w) sodium acrylate in PBS with polymerization initiated with TEMED and APS. Polymerized gels were incubated for 30 min at 37°C in a digestion buffer containing 8 U/ml proteinase K. Cover glasses were removed from the digested gels, which were placed in high volumes (> 30 mL) of distilled water that were exchanged at least five times until full expansion of the gels. Finally, the gels were...
trimmed, nuclei counterstained with Hoechst 33342 (1/5000 in water for 30 min at room temperature), positioned in 50 mm diameter glass bottom dishes (WillCo Wells GWS-5040) and immobilized using 2% agarose. The lateral expansion factor of samples processed identically to those shown throughout the manuscript was calculated to be 3.98, based on distance measurements between of the same cellular hallmark structures imaged pre- and post-expansion (Fig. S9). This is in line with the expansion factor described in the original protocols (Chozinski et al., 2016; Chen et al., 2015). In an approximation, this expansion factor was applied to all other experiments.

**Dendritic spine labeling**

To label spines in a sparsely distributed subset of neurons, 14 DIV cultures were incubated with the lipophilic dye CM-Dil (ThermoFisher C7000, 5 µg/ml, 20 min), followed by a PBS wash. The next day, after the dye had spread throughout the plasma membrane, the cultures were gently permeabilized using a glycerol gradient (50% glycerol for 20 min, 80% for 20 min and 100% for 50 min) followed by an extensive PBS wash. This permeabilization protocol preserved Dil fluorescence while allowing antibody penetration. A similar, but longer protocol was used for IF, in which the primary antibody was applied overnight at RT and the secondary for 4 hours.

**Synaptic Proximity Ligation Assay (synaptic PLA)**

Primary cultures fixed and permeabilized as described for IF were used to perform PLA using a commercial kit (Duolink® PLA starter kit, Sigma-Aldrich, Overijsse, Belgium) according to the manufacturer’s instructions. Briefly, cells were incubated with a commercial blocking buffer, and then incubated overnight at 4°C, with primary antibodies diluted in the commercial solution at concentrations described in Table S1. After PLA probe incubation, ligation and amplification steps, samples were immunostained for MAP2 and Synapsin for triple staining experiments, counterstained with Duolink® Nuclear Stain and mounted with anti-fade buffer (Sigma-Aldrich). Images were taken no more than two days after the PLA procedure to ascertain signal preservation.

**Microscopy**

 Cultures were imaged with a spinning disk confocal high-content imager (Opera Phenix, PerkinElmer) equipped with a robotic arm for plate loading. A 40X water immersion objective (numerical aperture (NA) 1.1) was used. At 488 nm excitation, the optical resolution of the system is 0.271 μm (and corresponding pixel size 0.149 μm), which is considerably larger than the distance of the synaptic cleft (15–25 nm), yet sufficiently small to allow signals of corresponding pre- and postsynaptic markers to partially overlap. Per well, 15 fields were acquired in four channels (405nm, 488nm, 561nm, and 640nm excitation) at four axial positions separated by 1 μm spacing. Different fluorescence channels were separated using standard excitation/emission filters and dichroic mirrors. Owing to the large dynamic range of the Opera Phenix system, acquisition settings could be kept identical for all experiments. For assessing localization of postsynaptic markers in dendritic spines and triple stainings (trans-synaptic PLA/mVenus/Synapsin), a spinning disk confocal research microscope (UltraVIEW VoX, PerkinElmer) was used, equipped with a 60X water objective (NA 1.2) resulting in an optical resolution of ~200 nm and corresponding pixel size of 120 nm. Finally, for super-resolution radial fluctuations (SRRF) microscopy (Culley et al., 2018), the same setup was used to acquire stacks of 200 images of the same confocal plane, using the Perfect Focus System to prevent focus drift.

**Image analysis**

To assess the degree of colocalization between two synaptic markers, we adopted the van Steensel approach (van Steensel et al., 1996), which we implemented via a home-written script for Fiji image processing freeware. This method calculates the Pearson’s correlation coefficient (PCC) between two images as a function of a lateral shift (Δx), resulting in a correlation function (CF). When the image is compared to its (shifted) duplicate, the result is an auto-correlation function (ACF). In case of complete colocalization, the correlation function (CF) has a maximum (PCC = 1) at Δx = 0 (no shift); in case of exclusion, it reaches a minimum at this position. The shape of the CF not only informs on the presence or absence of colocalization, when colocalization is present, it also provides information on the size of the overlapping spots (the width of the CF measured as full width at half maximum (FWHM)) and the dynamic range – and thus quality – of the staining (measured as the amplitude of the CF). Exactly because of this latter property, the approach was also used to assess the specificity of single synapse marker staining, the assumption being that the ACF of an image decreases with Δx much stronger for a specific staining than it does for less specific ones. The maximal pixel shift used for both analyses was 50 pixels in both directions, which is well beyond the typical spot diameter of ~10 pixels. The variation across images, or the spread, was expressed by a band of 1 standard deviation above and below the average cross-correlation function (CCF) and measured at dx = 50 px for plotting. The auto- or cross-correlation function (ACF/CCF) reports on labeling performance without differentiating between specificity,
sensitivity or local antigen abundance, as its amplitude will increase when background signal arising from non-specific binding is low, more target epitopes are labeled and when epitopes are highly enriched in synapses. Given that no preprocessing or segmentation is required, the CF method is not sensitive to user bias, yet it is influenced by the magnification and image resolution (Fig. S3A-C) which is why all acquisitions were done with one and the same objective lens (40X NA 1.1). Given the high density and heterogeneous distribution of the synaptic spots, the orientation of the sliding window does not significantly influence the readout (Fig. S3D).

Spot and synapse density quantification was carried out in Acapella® software (PerkinElmer). Nuclei (DAPI) and neurites (MAP2) were segmented based on a user-defined fixed threshold. Non-neuronal nuclei were removed based on their larger size, lower circularity and limited overlap with the neurite mask. Next, both the neurite and neuronal nuclei mask were dilated and subtracted from each other to obtain a search region for synapse marker spots. Spots were enhanced using a Difference of Gaussian filter with a ratio of 1.6. To set thresholds in an unbiased manner, a Fiji script was written that allowed user-friendly interaction with the images while being blinded for the synapse marker at hand. Spot counts were normalized to the neurite area to obtain the spot density. For double stainings and PSD-mVenus experiments, the density of colocalizing spots in two fluorescence channels was calculated as those spots for which at least one-pixel overlaps. We previously determined that measurements were not biased by chromatic aberration using fluorescent beads (the lateral and axial shifts between channels were below one pixel and manually introduced one-pixel shifts did not affect the colocalization measurement significantly) (Verschuuren et al., 2019).

For colocalization of antibodies with PSD95-mVenus, the percentage of PSD95-mVenus spots with an overlapping synapse marker spot was measured. For double immunostainings, the percentage of all spots that reside in synapses was calculated as the density of colocalizing spots divided by the density of all (colocalizing and non-colocalizing) spots multiplied by 100.

For double labeling with dendritic spines, the puncta within a CM-Dil-positive stretch were counted manually on anonymized images and assigned to either the dendritic shaft or spine. The percentage of spots residing in dendritic spines, and the percentage of spines containing spots were calculated. To quantify trans-synaptic PLA spot density on samples counterstained with MAP2, the same Acapella® script was used as described above. For nearest neighbor analyses, center coordinates were extracted for all spots (using a similar CellProfiler pipeline as described for Acapella®), and the pairwise distance distribution between spot types (PLA, mVenus, Synapsin) was analyzed in R using the package spatstat by applying the nearest neighbor correlation (nncorr) method after correcting for the search region (neurite mask) (Baddeley and Turner, 2005). For SRRF, the NanoJ plugin for Fiji was used (Culley et al., 2018). Briefly, after drift correction with NanoCore, a temporal radial auto-correlation of order 2 was applied, with gradient smoothing, intensity weighting and gradient weighting (point spread function FWHM = 3.17) for spot channels, and the previous settings with renormalization for the MAP2/DAPI channel. To account for the increase in resolution, subsequent spot overlap was determined between different channels of the SRRF images after expanding detected objects with 10 pixels (~240 nm). The center locations of the detected spots were used to calculate the average nearest neighbor distance between marker combinations with the nnist and marktable functions of the spatstat R package.

Data visualization and statistics

For displaying purposes, the contrast of each fluorescence channel was optimized and this was done for the entire image. Graphs were constructed in Graphpad Prism and statistics performed in SAS JMP. The number of replicates is indicated in the figure legends. For segmentation-independent Steensel analyses, 90 images originating from six wells were considered per condition (marker or marker combination and cell type), and for spot segmentation the same number of images were considered but averaged per well (n=6 wells with 15 images/well). For dendritic spine analysis, at least 200 spines were counted per condition. For trans-synaptic PLA experiments 4 or 8 wells with 15 fields per well were averaged per condition. For SRRF of trans-synaptic PLA staining, between of four and seven images were used per combination. For statistical comparison of spot densities, a one-way ANOVA was performed followed by post-hoc testing for predefined comparisons (antibodies that had the same target; seven comparisons) with Sidak’s correction. For triple stainings, a two-way ANOVA was performed with marker and PLA type as fixed factors, post-hoc tests for comparisons within conditions are specified in the results. For hTau-P301L overexpression a two-way ANOVA was performed with staining type and MOI as fixed factors, followed by post-hoc testing for predefined comparisons (to MOI 0 within each staining type) with Sidak’s correction.
Supplemental References

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