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Cell-type-specific visualisation and biochemical isolation of endogenous synaptic proteins in mice

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Abstract
In recent years, the remarkable molecular complexity of synapses has been revealed, with over 1,000 proteins identified in the synapse proteome. Although it is known that different receptors and other synaptic proteins are present in different types of neurons, the extent of synapse diversity across the brain is largely unknown. This is mainly due to the limitations of current techniques. Here, we report an efficient method for the purification of synaptic protein complexes, fusing a high-affinity tag to endogenous PSD95 in specific cell types. We also developed a strategy, which enables the visualisation of endogenous PSD95 with fluorescent-protein tag in Cre-recombinase-expressing cells. We demonstrate the feasibility of proteomic analysis of synaptic protein complexes and visualisation of these in specific cell types. We find that the composition of PSD95 complexes purified from specific cell types differs from those extracted from tissues with diverse cellular composition. The results suggest that there might be differential interactions in the PSD95 complexes in different brain regions. We have detected differentially interacting proteins by comparing data sets from the whole hippocampus and the CA3 subfield of the hippocampus. Therefore, these novel conditional PSD95 tagging lines will not only serve as powerful tools for precisely dissecting synapse diversity in specific brain regions and subsets of neuronal cells, but also provide an opportunity to better understand brain region- and cell-type-specific alterations associated with various psychiatric/neurological diseases. These newly developed conditional gene tagging methods can be applied to many different synaptic proteins and will facilitate research on the molecular complexity of synapses.

KEYWORDS
conditional tagging, fluorescent protein, mouse genetics, proteomics, PSD
1 | INTRODUCTION

Information processing in the mammalian brain depends on synapse proteins. Over the past two decades, proteomic studies have revealed remarkable synapse complexity and diversity. In these studies, biochemically purified protein complexes from the postsynaptic density were analysed and found to contain over 1,000 different protein components (Bayes et al., 2012, 2017, 2011; Collins et al., 2006; Distler et al., 2014; Emes et al., 2008; Grant, 2007; Peng et al., 2004; Roy et al., 2018; Trinidad, Specht, Thalhammer, Schoepfer, & Burlingame, 2006; Trinidad et al., 2008; Uezu et al., 2016). The functional diversification of synapses is often related to their molecular diversity. It has been reported that many synaptic proteins, including different subtypes of receptors, isoforms of scaffolding proteins and adhesion molecules, are differentially expressed in various brain regions (Bayes et al., 2012, 2011; Collins et al., 2006; Emes et al., 2008; Frank et al., 2016; Frank, Zhu, Komiyama, & Grant, 2017; Husi, Ward, Choudhary, Blackstock, & Grant, 2000; Komiyama et al., 2002; Lein et al., 2007; Micheva, Busse, Weiler, O’Rourke, & Smith, 2010; Roy et al., 2018).

Despite this recent progress, we still have insufficient knowledge on precisely how individual synapses differ in their molecular composition and how these different types of synapses are distributed with regard to neuronal cell type and brain region. This is partly due to experimental limitations in accurately tracking and dissecting specific anatomical regions and populations of neuronal cells. It is therefore necessary to establish an experimental system that allows precise and reliable identification and characterisation of synapse diversity in vivo.

In recent years, gene targeting approaches to engineer the mouse genome have proven to be a crucial tool for the detailed investigation of genes of interest in vivo. Combining gene targeting with the Cre/loxp-recombinase system makes it possible to achieve the precise spatiotemporal control of gene expression, introduction of point mutations or deletions of genomic sequence. Using gene targeting, we have previously reported the generation of a constitutive PSD95-TAP (tandem affinity purification) tag knock-in mouse line (Fernandez et al., 2009), in which PSD95 (also known as PSD-95 or DLG4) was in-frame fused with the small peptide TAP tag, for highly efficient synaptic protein purifications. PSD95 is an important postsynaptic scaffolding protein that plays key roles in synaptic development, protein complex assembly and plasticity. In the present study, we have further developed this methodology and now present the generation of two lines of conditional knock-in tagging mice: PSD95-c (mCherry/eGFP; monomeric Cherry, enhanced green fluorescent protein) and PSD95-cTAP (conditional TAP tag). In contrast to the global labelling of PSD95, here we applied a site-specific recombination strategy that allows Cre-mediated switching of fluorescent reporters that faithfully reflect endogenous expression. As a result, these genetically modified mice express red fluorescent PSD95-mCherry prior to Cre-mediated recombination and green fluorescent PSD95-EGFP after recombination, thereby allowing visualisation and distinction of PSD95 synaptic clusters in recombined and non-recombined cells. We also describe a conditional PSD95-cTAP tagging line, whereby the PSD95-TAP tag is expressed upon Cre-mediated recombination, thus allowing efficient isolation and purification of PSD95 protein complexes from recombined neurons.

Crossing these conditional tagging mice with different driver lines that express Cre recombinase under the control of specific types of neuronal promoters, we detected robust expression of PSD95-eGFP or PSD95-TAP in corresponding regions/neuronal populations. Intriguingly, we also detected two distinctive populations of PSD95-positive synaptic clusters in CA1 and CA3 pyramidal neurons. These populations differed not only in synaptic features such as synaptic density, size and punctate morphology, but also in molecular composition of PSD95 complexes as revealed by proteomic analysis.

2 | MATERIALS AND METHODS

The animal experiments described in this study were undertaken under the authority of a UK Home Office Project Licence (PPL 70/8140) under the terms, conditions and regulations of the UK Home Office Animals (Scientific Procedures) Act 1986.

2.1 | Targeting vector construction

A previously constructed intermediate targeting vector for the generation of PSD95-TAP tag mice (Fernandez et al., 2009) was used for tagging the C-terminus of endogenous PSD95 proteins. For generating the conditional fluorescent gene targeting vectors, a loxP-flanked cassette containing red fluorescent protein mCherry coding sequence (Clontech, 632523) and an FRT site-flanked neomycin resistance gene (PGK-neo) cassette were assembled into the backbone plasmid pNeoflox (Figure 1a). To generate the Cre-mediated conditional replacement of mCherry with eGFP, coding sequence for eGFP (Clontech, GenBank accession U55762) was further subcloned into the intermediate vector pNeofox-mCherry and immediately after the FRT-PGK-neo-FRT-loxP sequence (Figure 1a). Both mCherry and eGFP coding cassettes were in-frame-inserted just before the STOP codon of the murine PSD95 (Dlg4) gene using 11 amino acids encoded by nucleotide sequence for the loxP site as part of a flexible linker (Figure 1a). Approximately 4–6 kb upstream and downstream
regions of PSD95 last exon (with corresponding genomic sequence retrieved from BAC clones previously used in Fernandez et al., 2009) were cloned into the targeting vectors as 5’ and 3’ homology arms, respectively. All final targeting vectors contain a diphtheria toxin A (DT-A) fragment that allows for negative selection in embryonic stem cells.

The conditional PSD95-TAP targeting vectors were designed and constructed using a similar strategy to that mentioned above.
above, in which the last exon of PSD95 was engineered to in-frame fuse to a loxP site-flanked STOP codon, which is followed by a short G-S-G linker peptide coding sequence plus the TAP coding sequence, which includes a histidine affinity tag (HAT), TEV protease cleavage site and a triple FLAG tag. Therefore, in the presence of Cre recombinase, the strategically placed STOP codon is deleted, which drives the expression of the in-frame fusion protein PSD95-TAP (Figure 1b).

2.2 Embryonic stem (ES) cell gene targeting and generation of reporter mice

The targeting vectors were transfected into murine ES cells (E14 TG2a) via electroporation as previously described (Fernandez et al., 2009). G418 (300 μg/ml final concentration) was added to the ES cell culture 24 hr after plating for positive selection. Single G418-resistant colonies were picked 5–7 days after selection.

Correctly targeted ES cell colonies were identified by long-range PCR amplification (Expand Long Template PCR system, Roche, Cat No. 11681842001) and further injected into recipient blastocysts from C57BL/6J mice. Adult male chimaeras were selected to breed with C57BL/6J wild-type female mice (The Jackson Laboratory) to produce heterozygotes, which were mated with FLP deleter mice to remove the FRT-flanked neo cassette. Genomic DNA extracted from all F1 progeny ear clips was analysed by PCR to confirm the genotype (data not shown).

Two lines of transgenic mice with region-specific expression of Cre recombinase were used to establish the conditional reporter mouse colonies: (a) Grik4-Cre mice expressing Cre recombinase in the CA3 pyramidal neurons of the hippocampus (Nakazawa et al., 2002); and (b) Pvalb-Cre mice expressing Cre recombinase under the control of the endogenous parvalbumin (Pvalb) promoter (Hippenmeyer et al., 2005).

2.3 Histology, fluorescence microscopy imaging

Anaesthetised mice were transcardially perfused with 4% paraformaldehyde (PFA; Alfa Aesar, 30525-89-4) in 0.1 M phosphate buffer (pH 7.4). The brain was immediately post-fixed in the same fixative (4% PFA) for 3–4 hr at 4°C before transferring into 30% (w/v) sucrose in 0.1 M phosphate buffer; Sigma-Aldrich) at 4°C for at least 24 hr. Coronal or sagittal sections of 18 μm thickness were cryosectioned by a cryostat (Thermo Fisher).

For some tissue sections, immunohistochemical stainings were carried out. Mounted frozen sections were thawed at room temperature and rehydrated with PBS for 10 min. After incubating sections with blocking solution 5% bovine serum albumin (BSA) in Tris-buffered saline (pH 7.5) with 0.2% Triton X-100 for 2 hr at room temperature, sections were incubated with primary antibodies including anti-PSD95 (mouse monoclonal IgG1; NeuroMab, 1:250, 75-348), anti-mCherry (rabbit polyclonal; Abcam, 1:500, ab167453), anti-Synaptophysin (mouse monoclonal IgG1; Millipore, 1:250, MAB5258-20UG), anti-MAP2 (rabbit polyclonal; Abcam, 1:1,000, ab32454) or anti-parvalbumin (mouse monoclonal IgG1; Swant, 1:200, PV235) in a humidified chamber overnight at 4°C. After washing three times, sections were incubated with Alexa Fluor 488/546/633-conjugated secondary antibodies (Life Technologies) for 2 hr at room temperature before being mounted with mounting medium. Slides were imaged by a laser scanning confocal microscope (LSM510, Zeiss) using a 63x objective (NA 1.4) or a spinning disc confocal microscope (Andor Revolution XD system) with a 100x oil-immersion Plan-Apochromat lens (NA1.4). Z-series images were collected with an optical interval of 0.1 μm for 10–13 planes. Tile scans were stitched into larger mosaics. Images were adjusted using ImageJ (National Institutes of Health) or Photoshop software for visualisation (Adobe). For fluorescence image analysis, digital image stacks were analysed by ImageJ and in-house-designed algorithms written in the ImageJ macro language. Before processing, the original images were adjusted to subtract the background fluorescence intensity. All adjustments were applied equally to all images. Images were then subjected to the algorithms for punctate object segmentation and detection. Quantitative measurements of punctum size and number were performed using an ImageJ plugin 3D Object Counter (Bolte & Cordelieres, 2006), and statistical analyses were performed with GraphPad Prism 6.0 (GraphPad Software, Inc.).

2.4 Biochemical analysis

Cervical dislocation was used as a method of animal euthanasia. Whole forebrains or hippocampi dissected and snap-frozen in the liquid nitrogen and kept at −80°C until used. Whole forebrains from the constitutively TAP-tagged PSD95 line were dissected from heterozygous mice. The hippocampi were isolated from the conditional TAP-tagged PSD95 line crossed with Grik4-Cre mice. Single-step affinity purifications were carried out as described previously with minor modifications (Fernandez et al., 2009). In brief, brain tissue was homogenised in DOC buffer [50 mM Tris pH 9.0, 1% sodium deoxycholate, 50 mM NaF, 20 mM ZnCl2, 1 mM Na2VO4, 2 mM Pefabloc SC (Roche) and 1 tablet protease inhibitor cocktail/10 ml (Roche)] and clarified as described (Grant & Husi, 2001). Extracts were incubated with Dynabeads coupled with FLAG antibody for 2 hr at 4°C. The resin was washed with three cycles of 15 resin volumes of DOC buffer. Captured proteins were eluted with the same buffer containing FLAG peptide. Eluted fractions containing PSD95 were pooled, concentrated in a Vivaspin concentrator (Vivascience), reduced with DTT, alkylated with iodoacetamide and separated by one-dimensional 4%–12% SDS-PAGE.
(NUPAGE, Invitrogen). The gel was fixed and stained with colloidal Coomassie Blue, and gel lanes were cut into slices and each slice was destained and digested overnight with trypsin (Roche, trypsin modified, sequencing grade) as reported previously (Bayes et al., 2011).

Antibodies used for Western blot are FLAG Ab (anti-mouse, from Sigma diluted 1:1,000) and PSD95 Ab (anti-mouse, from Thermo, diluted 1:1,000).

2.5 Mass spectrometry

Extracted peptides (three fractions per sample) were analysed by nanoLC-MS/MS on an LTQ Orbitrap Velos (Thermo Fisher) hybrid mass spectrometer equipped with a nanospray source, coupled with an Ultimate 3000 Nano/Capillary LC System (Dionex). The system was controlled by Xcalibur 2.1 (Thermo Fisher) and DCMSLink 2.08 (Dionex). Peptides were desalted online using a micro-precolumn cartridge (C18 PepMap 100, LC Packings) and then separated using a 120 min RP gradient (4%-32% acetonitrile/0.1% formic acid) on an EASY-Spray column, 50 cm × 75 µm ID. PepMap C18, 2 µm particles, 100 Å pore size (Thermo Fisher). The LTQ Orbitrap Velos was operated with a cycle of one MS (in the Orbitrap) acquired at a resolution of 60,000 at m/z 400, with the ten most abundant multiple-charged (2+ and higher) ions in a given chromatographic window subjected to MS/MS fragmentation in the linear ion trap. FTMS target values of 3e6 and an ion trap MSn target value of 1e4 were used with the lock mass (445.120025) enabled. Maximum FTMS scan accumulation time of 500 ms and maximum ion trap MSn scan accumulation time of 60 ms were used. Dynamic exclusion was enabled with a repeat duration of 30 s with an exclusion list of 500 and exclusion duration of 60 s.

MS data were analysed using MaxQuant (Cox & Mann, 2008) version 1.2.7.4. Data were searched against mouse UniProt sequence databases (downloaded March 2012) using the following search parameters: trypsin with a maximum of two missed cleavages, 7 ppm for MS mass tolerance, 0.5 Da for MS/MS mass tolerance, with acetyl (protein N-term) and oxidation (M) set as variable modifications and carbamidoethyl (C) as a fixed modification. A protein false discovery rate (FDR) of 0.01 and a peptide FDR of 0.01 were used for identification level cut-offs. Label-free quantification was performed using MaxQuant LFQ intensities (Cox et al., 2014), and statistical analysis was performed using Perseus (Tyanova et al., 2016). Only proteins quantified in at least three replicate purifications were included in the statistical analysis. Protein label-free quantification (LFQ) intensities were log2-transformed, missing values were imputed, and t-testing was performed (using Perseus) with correction for multiple hypothesis testing using a permutation-based FDR of 0.05 to identify proteins quantitatively different between sets of purifications.

2.6 Gene set over-representation analysis

In order to investigate whether any GO biological processes or pathways (KEGG and Panther) were enriched in the CA3 versus the hippocampal PSD-95 protein complexes, we performed an over-representation analysis (ORA) using WebGestalt (http://www.webgestalt.org/). The frequency of terms associated with proteins in the CA3 set was compared to the constitutive hippocampal set and those that were statistically over-represented were determined with an FDR of 0.05 (Benjamini–Hochberg).

3 RESULTS

3.1 Generation and establishment of conditional PSD95 knock-in tagged mice

To precisely probe the expression of endogenous PSD95 protein, we applied a gene targeting-based knock-in approach in which different tags (either a TAP tag or different fluorescent protein tags with distinct excitation/emission spectrums) were in-frame-fused to murine PSD95 (Figure 1a,b). Previous findings have shown that the PSD95 gene has multiple isoforms with alternatively spliced N-terminals but that all share a common C-terminus (Bence, Arbuckle, Dickson, & Grant, 2005); therefore, these tags were inserted into the open reading frame immediately before the STOP codon (Figure 1a,b). To maintain the “in-frame” insertion of the reporter cassettes before and after Cre-mediated recombination, we designed two similar targeting vectors for the fluorescent or TAP tagging, in which the mCherry coding sequence or a STOP codon, together with the FRT-flanked selection marker cassette, was placed immediately after the first loxP site, which encodes 11 amino acids, plus a short glycine-rich linker sequence (G-G-G-S or G-S-G for the fluorescent or TAP tag, respectively; Figure 1a,b). Following excision of the mCherry or STOP codon sequences that reside between the two loxP sites, the resulting targeted allele will convert to the coding sequence for eGFP or TAP peptide, which again follows an “in-frame” loxP site together with the short glycine-rich linker sequence (Figure 1a and Materials and Methods).

Over 200 G418-resistant ES cell clones were screened, and correctly targeted clones were identified by long-range PCR using a 3’ external genomic reverse primer to validate the integrity of the targeted allele. Clones were selected for blastocyst injection to generate chimaeras for either the conditional fluorescent tagging or TAP tagging mouse lines. Germline transmitted mice from chimaeras were identified by PCR genotyping using specific primer sets. F1 heterozygous offspring were bred with FLPe deleter mice, expressing FLP recombinase, to remove the neo cassette flanked by two FRT sites (Figure 1a,b). Heterozygous progeny (without the PGK-neo
cassette) were used for further breedings with either germline-expressing CAG-Cre or cell-type-specific Cre driver lines.

To confirm that introducing fluorescent tags into the endogenous gene locus did not alter the expression levels of endogenous PSD95, we performed Western blot analysis using protein extracts of homogenates from heterozygous PSD95mCherry+/+ and littermate wild-type mice. Antibodies against murine PSD95 showed two bands from the heterozygous sample group, with the ~130 kDa and ~100 kDa bands being of similar intensity (Figure 1d).

In the absence of Cre, heterozygous PSD95c(mCherry/eGFP)/+ mice carry a PSD95-mCherry fusion allele, whereas the eGFP cassette was prevented from transcribing due to the downstream position of the STOP codon (Figure 1a). Therefore, brain sections from these mice should only be labelled with PSD95-mCherry. Fluorescence microscopy of fixed sagittal sections from adult PSD95c(mCherry/eGFP)/+ mice revealed that prominent mCherry expression can be observed in various brain regions (Figure 1c). Higher fluorescence intensity levels of PSD95-mCherry were detected in the forebrain, including cerebral cortex and hippocampus, whereas weaker fluorescence was detected in other regions including the midbrain, cerebellum and medulla oblongata. Confocal images revealed that most of the PSD95-mCherry puncta were situated in close opposition to the Synaptophysin-immunoreactive pre-synaptic terminals, thus confirming the synaptic localisation of PSD95-mCherry fusion protein (Figure 1c-i,ii,e). This pattern strongly resembles previous immunohistochemical findings using a specific PSD95 antibody (Fukaya & Watanabe, 2000).

Upon Cre recombination, the loxP site-flanked region (including mCherry coding sequence, PGK-neo cassette and the following STOP codon) was excised and the eGFP coding cassette brought immediately downstream of the PSD95 gene, creating a PSD95-eGFP fusion allele (Figure 1a). To test whether the targeted allele can be converted from PSD95-mCherry to PSD95-eGFP, we crossed a male heterozygous PSD95c(mCherry/eGFP)/+ mouse with female CAG-Cre transgenic mice (Sakai & Miyazaki, 1997). CAG-Cre is capable of ubiquitous deletion of loxP site-flanked gene sequence in the germline cells. As a result, brain sections from the progeny of these breedings should be labelled with the eGFP fluorescent reporter tag. As expected, breeding with the CAG-Cre deleter line resulted in heterozygous mice expressing prominent PSD95-eGFP across the brain (Figure 1c-iii,iv).

3.2 | Neuronal cell-type-specific expression of PSD95-positive synapses in the brain

To further examine whether the targeted allele is responsive to a tissue/cell-type-specific conversion, conditional heterozygous PSD95c(mCherry/eGFP)/+ mice were bred with Grik4-Cre or Pvalb-Cre transgenic mice, respectively. Grik4-Cre mice show Cre activity in nearly all pyramidal neurons of the hippocampal CA3 area (Akashi et al., 2009), whereas Pvalb-Cre mice express Cre recombinase in parvalbumin-positive neurons (Hippenmeyer et al., 2005). As described in Figure 2a, brain cryosections from double-transgenic PSD95c(mCherry/eGFP)/+; Grik4-Cre mice showed CA3 region-specific expression of PSD95-eGFP, in contrast to the uncombined PSD95-mCherry expression in other hippocampal regions. Interestingly, higher magnification views revealed distinctive PSD95+ synapse populations among different hippocampal subregions: in contrast to the densely packed PSD95-mCherry puncta in the CA1 region, the PSD95-eGFP puncta in the CA3 area (stratum lucidum) appeared to be larger and more sparsely distributed (Figure 2a). Quantification of PSD95 punctum density showed a significant difference between CA1 (~120/100 µm²) and CA3 (~40/100 µm²) subregions (Figure 2b). The punctum size of PSD95 in these regions also differs (0.015 µm³ in CA1 vs. 0.023 µm³ in CA3; Figure 2b). Taken together, these data suggest that distinctive kinds of PSD95+ synaptic puncta occur in different subregions of the hippocampus.

In addition, we also examined compound PSD95c(mCherry/eGFP)/+; Pvalb-Cre transgenic mice. In contrast to native condition, PSD95c(mCherry/eGFP)/+; Pvalb-Cre compound mice display higher GFP fluorescence level in the cerebellum (Figure 2a, panel iv compared to Figure 1c, panel i). Further examination of the hippocampal CA3 regions using IHC of parvalbumin antibody to highlight parvalbumin-positive cells has shown that discrete PSD95 puncta were observed distributed along the parvalbumin+ neuritip, suggesting a fraction of PSD95+ excitatory synapses identified in the hippocampal parvalbumin+ cells (Figure 2a, panels v and vi).

3.3 | Proteomic analysis of PSD95-cTAP in the hippocampus versus the CA3 subfield

Although we have observed distinctive features of PSD95+ synaptic puncta from CA1 and CA3 pyramidal cells of the hippocampus with imaging techniques, it remains unknown whether and how these PSD95 complexes differ at the molecular level.

To address this issue, we performed cell-type-specific tagging of PSD95 and characterised the composition of its protein complexes using high-affinity FLAG purification followed by analysis of complex components using mass spectrometry.

Mice carrying the conditional PSD95-TAP tag alleles (PSD95c(TAP)) were crossed with the Grik4-Cre line in which Cre recombinase was expressed in CA3 of the hippocampus. We checked the expression of tagged PSD95 protein in the lysates from each of the double-transgenic lines. No leaky
expression of the tag could be detected in the cortex (Cx), hippocampus (Hp) or striatum (Str) of the PSD95cTAP/+ mice (Figure 3a). We could detect FLAG-tagged PSD95 protein in the brain lysate from double-transgenic mice (PSD95cTAP; Grik4-Cre; Figure 3a,b).

In order to generate a robust list of PSD95 interactors in the hippocampus (previous data sets were generated from forebrain tissue; Fernandez et al., 2009), we performed four replicate purifications from mouse hippocampal tissue expressing the constitutively expressed TAP-tagged version of PSD95 (PSD95cTAP; Grik4-Cre; Figure 3a,b).
of PSD95 (Fernandez et al., 2009) and three replicate control purifications using wild-type mice. We then performed protein/peptide identification and label-free quantification using MaxQuant (Cox & Mann, 2008) on all preparations. To define a set of proteins that were enriched in constitutively expressed PSD95-TAP purifications compared with control purifications, \( t \)-testing was performed (using Perseus) on LFQ protein intensities with correction for multiple hypothesis testing using a permutation-based FDR of 0.05. Of the 403 proteins quantified in at least 3 replicates, 103 proteins were significantly enriched in PSD95-TAP purifications compared to controls (Figure 4a and Table S1), representing a data set of quantitatively enriched proteins associated with constitutively expressed TAP-tagged PSD95 in the hippocampus.

Next, we purified PSD95 complexes from hippocampi dissected from the PSD95\(^{cTAP}\); Grik4-Cre double-transgenic line (see Materials and Methods). Four replicate CA3 purifications were analysed by mass spectrometry, and we defined a set of proteins that were enriched in purifications from mice with CA3 specific expression of PSD95-TAP by comparison with data from control purifications from hippocampal tissue of wild-type mice. In total, of 234 proteins quantified in at least 3 replicates, 62 proteins were significantly enriched in purifications from Grik4-Cre mice representing a data set of quantitatively enriched proteins associated with TAP-tagged PSD95 expressed in the CA3 region of the hippocampus (Figure 4b and Table S2).

Comparison of the constitutive hippocampal and CA3-specific PSD95-cTAP data sets reveals an overlap of 53 proteins with 9 that are specific to the CA3 subfield (Figure 4c) and 50 that are specific to the constitutive hippocampal data set. The CA3 data set contains PSD95 complex components that are mostly (90%) independently identified as complex components in the hippocampus of the constitutively expressed PSD95-TAP mouse line. Importantly, this CA3-restricted set represents just over half (51%) of the components of constitutive hippocampal data set, indicating that this approach allows the identification of brain region-specific PSD95 complexes. These results demonstrate proof of concept that this tagging approach allows the purification and characterisation of synaptic protein complexes from a specific region of the hippocampus.

In order to investigate whether PSD-95-associated complexes in the CA3 subfield of the hippocampus might be functionally different to those obtained from the whole hippocampus, we performed a bioinformatic analysis to identify GO biological processes and pathways (KEGG and Panther) that were over-represented in the CA3 set compared to the constitutive hippocampal set (Figure S1). The CA3 PSD-95 set was enriched for biological processes associated with general synaptic functions (such as “synaptic signalling” and “chemical synaptic transmission”) and more specific pathways (such as ErbB, Rap1, ionotropic glutamate receptor and mAChR signalling pathways and “anterograde trans-synaptic signalling”).

**DISCUSSION**

The findings reported here provide a proof of principle that conditional PSD95 labelling can be used for both visualisation and biochemical analysis in the mouse brain. PSD95 is one of the most abundant synaptic scaffolding proteins. It is widely expressed in many different types of neuronal cells in the brain and forms large macromolecular complexes with neurotransmitter receptors, various ion channels, cell adhesion and cytoskeletal proteins and signalling molecules (Collins et al., 2006; Fernandez et al., 2009; Frank et al., 2016, 2017; Husi & Grant, 2001; Husi et al., 2000). These complexes play essential roles in synaptic function, and mutations in the genes encoding the components of these complexes are implicated in human psychiatric disorders (Bayes et al., 2011; Kaizuka & Takumi, 2018). However, it is still not clear how the molecular nature of these complexes differs between brain regions and neuronal cell types, or at different developmental stages. Therefore, it was important to develop an efficient approach...
by which a reporter transgene can be activated in a temporally and spatially regulated fashion in specific cell lineage or tissue.

Here, we present two conditional PSD95 reporter mouse lines generated with a gene targeting approach. These mouse lines not only allow simultaneous visualisation of distinctive
PSD95+ synaptic puncta in different populations of neurons, but also facilitate quantitative measurement and analysis of PSD95-containing synaptic protein complexes (Frank et al., 2016) with biochemical/proteomic methods.

A major concern with current technical efforts in molecular mapping/imaging of the synapse is the use of antibodies that can lack sufficient specificity. Another technical difficulty concerns the access of the antibody into the protein-dense PSD structure. In a study carried out by Fukaya and Watanabe (2000), processing of tissue sections with a carefully titrated pepsin treatment was required before specific labelling of PSD95 could be achieved, involved a substantial investment of time and effort. Furthermore, antibody cross-reactivity, especially when detecting closely related protein paralogues, also needs to be carefully considered.

Using fluorescent proteins genetically tagged to a target protein can overcome the above limitations. In contrast to conventional immunofluorescence staining or biochemical methods, which usually rely on antibodies, genetic tagging provides an easy method for direct visualisation or biochemical isolation/purification of the tagged protein.

It is also recognised that quantifying and comparing immunostaining signals between different brain sections is often difficult and unreliable, even when antibody dilution and staining conditions are kept consistent. In contrast, we found that fluorescent signals between different slices prepared from different individual PSD95-eGFP knock-in mice were remarkably consistent and reproducible. This has allowed us to survey and numerically quantify PSD95-eGFP puncta in terms of their number, size and intensity at the whole brain scale (Zhu et al., 2018). In addition, the genetic fluorescent labelling makes it possible to carry out live imaging in the intact animal, or acute brain slices, to study the molecular dynamics of PSD95 (Wegner, Mott, Grant, Steffens, & Willig, 2018).

Several lines of evidence have shown that a physiological expression level of PSD95 is crucial for maintaining the normal number, size and morphology of spine synapses. Overexpression of PSD95 modifies synaptic transmission and impairs synaptic plasticity (Ehrlich & Malinow, 2004; El-Husseini, Schnell, Chetkovich, Nicoll, & Bredt, 2000; Nikonenko et al., 2008). To precisely probe the expression of PSD95 and keep to a minimum the functional perturbation of the endogenous protein, we decided to apply a gene targeting-based approach. In contrast to other genetic labelling methods such as BAC transgenesis, which due to the saturation of targeting machinery or overexpression of labelled proteins may sometimes fail to report the expression and localisation of target protein properly, this gene targeting method enables tags to be integrated into a specific genomic locus. Gene targeting is also superior to transient transfection methods, which cause overexpression of the protein, and other genetic labelling techniques such as insertion of the LacZ reporter gene, which only allows a rather general evaluation of expression pattern at a cellular level (Porter, Komiyama, Vitalis, Kind, & Grant, 2005). In the reporter lines described in this study, the locus-specific in-frame insertion of the tag faithfully follows the endogenous expression of PSD95 and enables precise protein localisation at the subcellular level.

One potential limitation of constitutively GFP-tagged PSD95 knock-in is difficulties in probing the expression of tagged protein in a specific subset of cells. The conditional labelling of PSD95 in this study allows not only precise visualisation, but also faithful proteomic capture and analysis of PSD95 molecular complexes in a specific subset of excitatory synapses.

Combined with appropriate Cre driver mice, these reporter lines offer an opportunity to readily label endogenous PSD95 in selected subpopulations of neuronal cells/tissues. For example, with the compound conditional fluorescent reporter line, two distinctive populations of PSD95 synaptic puncta from CA1 and CA3 pyramidal neurons identified by spectrally different fluorescent proteins were detected simultaneously (Figure 2b). Therefore, different synaptic properties (synaptic density, size and morphology) can be readily characterised and compared at the single-synapse level. Although some difficulties remain in terms of direct comparison of fluorescence intensity between two different species of fluorescent protein, the other conditional PSD95-cTAP tagging line provides a key complementary approach.

In this study, we have also successfully isolated and purified PSD95 protein complexes from different brain regions using conditional affinity tagging methods. Our results suggest that there might be differential interactions in the PSD95 complexes in different brain regions. We have detected differentially interacting proteins by comparing data sets from the hippocampus and the CA3 subfield of the hippocampus. All the CA3 (PSD95-cTAP; Grik4 Cre) enriched proteins (Abr, Actn4, Bai3, CaCng8, Grik2, Hist1h2a1, Mbp, Nlg3 and Prdx5) have been reported to have important functions in brain that include roles in synapse formation, synaptic plasticity and psychiatric disorders (Duman, Tu, & Tolias, 2016; Ge et al., 2016; Kalinowska et al., 2015; Lanoue et al., 2013; Malty et al., 2017; Martinelli et al., 2016; Oh et al., 2010; Park et al., 2017; Sigoillot et al., 2015; Walikonis et al., 2001; Yamagata et al., 2017). This might indicate that different types of PSD95 complexes exist in different brain regions with distinct functions. Our initial bioinformatic analysis for gene enrichment suggested that there might be specific biological pathways, such as ErbB, Rap1, and mAChR signalling pathways and anterograde trans-synaptic signalling, mediated by the proteins found in the CA3 area. However, a full interrogation of these data sets awaits more comprehensive bioinformatic analysis in the future.
The functional importance of the proteins enriched in PSD95 complexes in this specific brain area can be further elucidated by using mouse lines with conditional/floxed allele for these genes crossed with same Cre driver line (e.g. Grik4 Cre).

We previously reported the expression pattern of PSD95 and SynGAP, a major interactor of PSD95, in different brain regions and at different developmental stages (Porter et al., 2005). That study revealed remarkable differences in the expression pattern of these two PSD proteins at the cellular level using lacZ reporter knock-in mouse lines. These results indicated that the composition of PSD95 macromolecular complexes may show a high degree of heterogeneity between different neuronal cells. This view is consistent with our current finding, which shows that the molecular composition of PSD95 complexes in hippocampal CA3 neurons differs from that in the rest of the forebrain. Also, our recent study combining genetics and proteomics methods demonstrated that there are many different types of PSD95 subcomplexes in the forebrain (Frank et al., 2016), and our new method could help to elucidate further details of these different complexes. Using standardised proteomic analysis methods, this conditional tagging mouse line will enable us to accurately quantify and characterise the various PSD95 supramolecular complexes in virtually any lineage of neuronal cells/tissues with the use of different Cre driver lines.

In the future, we can further genetically engineer current reporter mouse lines (i.e. via the CRISPR/Cas system) to incorporate an additional fluorescent reporter gene cassette designed to follow a multicistronic element in order to visualise target neuronal morphology. As a result, both PSD95 and neuronal processes can be observed simultaneously upon Cre-mediated recombination. Therefore, this type of reporter line will be a valuable tool in making useful connections between the data generated by ongoing research into the mouse connectome and molecular synaptome mapping.

Taken together, these novel conditional PSD95 reporter mouse lines will serve as a powerful tool for detailed dissection of synapse diversity in any selected subset population of neuronal cells/tissues via optical imaging or proteomic techniques. These lines can be also used in many other research fields, for example the generation of transgenic animals by crossing with various neurological/psychiatric disease mouse models. Our conditional tagging method can be applied to any protein complexes expressed in different tissues and cell types in the mouse to enable the study of the distinct composition and distribution of multiprotein complexes.

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CONFLICT OF INTEREST

The authors declare no conflict of interests.

AUTHOR CONTRIBUTIONS

NHK designed the study. NHK, JSC and SGNG supervised the research. FZ, JH, MOC and NHK performed the experiments. FZ, MOC and NHK analysed and interpreted the data. FZ, MOC and NHK wrote the manuscript.

DATA AVAILABILITY STATEMENT

Supporting data are available on request.

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**SUPPORTING INFORMATION**

Additional supporting information may be found online in the Supporting Information section.

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