

Rapid modulation of long-term depression and spinogenesis via synaptic estrogen receptors in hippocampal principal neurons

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Abstract

Rapid modulation of hippocampal synaptic plasticity by estrogen has long been a hot topic, but analysis of molecular mechanisms via synaptic estrogen receptors has been seriously difficult. Here, two types of independent synaptic plasticity, long-term depression (LTD) and spinogenesis, were investigated, in response to 17 β -estradiol and agonists of estrogen receptors using hippocampal slices from adult male rats. Multi-electrode investigations demonstrated that estradiol rapidly enhanced LTD not only in CA1 but also in CA3 and dentate gyrus. Dendritic spine morphology analysis

demonstrated that the density of thin type spines was selectively increased in CA1 pyramidal neurons within 2 h after application of 1 nM estradiol. This enhancement of spinogenesis was completely suppressed by mitogen-activated protein (MAP) kinase inhibitor. Only the estrogen receptor (ER) α agonist, (propyl-pyrazole-trinyl)tris-phenol (PPT), induced the same enhancing effect as estradiol on both LTD and spinogenesis in the CA1. The ER β agonist, (4-hydroxyphenyl)-propionitrile (DPN), suppressed LTD and did not affect spinogenesis. Because the mode of synaptic modulations by estradiol was mostly the same as that by the

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Abbreviations used: ACSF, artificial cerebrospinal fluid; BSA, bovine serum albumin; CNQX, cyano-nitroquinoxaline; DES, diethylstilbestrol; DG, dentate gyrus; DPN, (4-hydroxyphenyl)-propionitrile; EPSP, excitatory postsynaptic potential; ER, estrogen receptor; KO., Knockout; LTD, long-term depression; LTP, long-term potentiation; PBS, phosphate-buffered saline; PP2, amino-chlorophenyl-butylpyrazolo-pyrimidine; PPT, (propyl-pyrazole-trinyl)tris-phenol; PRE, presynaptic membrane-rich fraction; PSD, postsynaptic density; SDS, sodium dodecyl sulfate.

ERalpha agonist, a search was made for synaptic ERalpha using purified RC-19 antibody qualified using ERalpha knockout (KO) mice. Localization of ERalpha in spines of principal glutamatergic neurons was demonstrated using immunogold electron microscopy and immunohistochemistry.

ERalpha was also located in nuclei, cytoplasm and presynapses.

Keywords: hippocampus, estrogen, neurosteroid, synaptic plasticity, spine, estrogen receptor.

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Estradiol exerts a rapid (e.g. 1 h) influence on the synaptic plasticity of rat hippocampal glutamatergic neurons in slices, as has been demonstrated by a number of electrophysiological investigations in rats and mice concerning long-term potentiation (LTP) in CA1 (Foy *et al.* 1999; Bi *et al.* 2000), long-term depression (LTD) in CA1 (Vouimba *et al.* 2000) or kainate current in CA1 (Gu and Moss 1996; Gu *et al.* 1999). To explain this modulation, the identification of molecular type and localization of estrogen receptors should be clarified. The synaptic estrogen receptors may also be required from the recent observation of endogenous synaptic synthesis of estradiol in adult male hippocampal neurons (Kawato *et al.* 2002; Shibuya *et al.* 2003; Hojo *et al.* 2004). On the other hand, extensive studies have been performed to investigate the role of estrogens in slowly (1–4 days) modulating hippocampal plasticity, because the hippocampus is known to be a target for the actions of gonadal estrogens reaching the brain via blood circulation. For example, the density of dendritic spines in the CA1 pyramidal neurons is modulated *in vivo* by supplement of estrogens in ovariectomized animals (Gould *et al.* 1990; Woolley *et al.* 1990; Woolley and McEwen 1992; Leranath *et al.* 2000, 2002) and of androgens in castrated animals (Leranath *et al.* 2003), resulting in an increase in/recovery of the number of spines. *In vitro* investigations have also shown that spine density is increased following several days of treatment of cultured hippocampal slices with estradiol (Murphy and Segal 1996; Pozzo-Miller *et al.* 1999).

The site of estrogen action in the hippocampus has been a matter of debate for more than a decade. So far, two distinct types of estrogen receptor have been identified in the mammalian brain: estrogen receptor alpha (ER α) (Simerly *et al.* 1990; McEwen 2002) and ER β (Shughrue *et al.* 1997; Mitra *et al.* 2003). The rapid effect of estrogen may be achieved by either ER α or ER β , possibly localized at the membrane in analogy with cultured cells of peripheral origin (McEwen and Alves 1999; Razandi *et al.* 1999). The subcellular and cellular localization of estrogen receptors is still not fully elucidated even for ER α , particularly in adult rat hippocampus. Many studies have reported, in female rats, that ER α -immunoreactivity has been found in the nuclei of scattered inhibitory gamma-aminobutyric acid (GABA)ergic interneurons by light or electron microscopy using AS409 antiserum against ER α (Weiland *et al.* 1997; Orikasa *et al.* 2000; Milner *et al.* 2001). It is therefore assumed that interneurons are the targets of estrogen action. However, it is also possible that estrogen may exert its effects directly on

principal neurons, based on growing evidence such as an NMDA receptor-dependent mechanism of estradiol regulation on dendritic spine density (Woolley and McEwen 1994), and increase of glutamate binding to NMDA receptors by estradiol (Woolley *et al.* 1997).

The present study was designed, using acute hippocampal slices from adult male rats, (i) to examine mechanisms of rapid modulation of two representative types of synaptic plasticity (LTD and spine morphology) by estradiol and agonists for ER α and ER β , and (ii) to determine the synaptic localization of ER α , which may predominantly catalyze these modulations. When considering the role of estrogen in memory processing, its effect on LTD is essential. LTD is not simply a ‘forgetting’ mechanism; it may be a positive mechanism used to ‘correct’ wrong memories formed by initial LTP processes which store not only correct information but also wrong information (Migaud *et al.* 1998). Regulation of spinogenesis is another important role of estrogen in memory processes via producing new spines for creating new neuronal contacts. The use of acute hippocampal slices has a particular importance with regard to the direct effect of estradiol on glutamatergic neurons, because results from *in vivo* investigations may reflect not only direct, but also indirect effects of estradiol via cholinergic or serotonergic neurons projecting to the hippocampus (Leranath *et al.* 2000, 2003). To identify the localization of ER α at the synapses, purified antibody RC-19 was used to avoid non-specific staining. The reactivity of RC-19 was examined using ER α knockout (KO) mice and mass spectroscopy.

Materials and methods

In the current work, estradiol refers to 17 β -estradiol, unless otherwise stated.

Animals

Twelve-week-old adult male Wistar rats were purchased from Saitama Experimental Animal Supply (Saitama, Japan) and Harlan Sprague-Dawley (Indianapolis, IN, USA). Male rats are particularly used in order to elucidate an essential function of endogenously synthesized estradiol in the male hippocampus (Hojo *et al.* 2004). ER α knockout mice [ER α ($-/-$)] were obtained by inbreeding ER α ($-/+$) mice. The development of ER α KO mice was accomplished by deleting the whole exon 2 of the mouse ER α gene (Dupont *et al.* 2000). It should be noted that nomenclature of ER α exon changed recently, and the current exon 1 and exon 2 (Kos *et al.* 2002; Pendaries *et al.* 2002) correspond to the previous exon 2 and exon 3,

respectively (Dupont *et al.* 2000). All experiments using animals in this study were conducted according to institutional guidelines.

Chemicals

N-methyl-D-aspartate (NMDA), cyano-nitroquinoxaline-dione (CNQX), Lucifer Yellow, 17 α - and 17 β -estradiol, ICI 182,780, (propyl-pyrazole-trinyl)tris-phenol (PPT), (4-hydroxyphenyl)-propionitrile (DPN), amino-chlorophenyl-butylpyrazolo-pyrimidine (PP2), diethylstilbestrol (DES) and PD98059 were purchased from Sigma (St. Louis, MO, USA). MC-20 antisera were purchased from Santa Cruz Biotechnology (Santa Cruz, CA, USA) in the years of 2001 and 2002. Anti-synapsin I antibody was from Chemicon (Temecula, CA, USA).

Measurements of the LTD with custom multi-electrode probes

Adult male rats aged 12 weeks were deeply anesthetized with ethyl ether and decapitated. The brains were removed and placed in artificial cerebrospinal fluid (ACSF) at 4°C. The hippocampus was then dissected and 300 μ m thick transverse dorsal slices to the long axis, from the middle third of the hippocampus, were prepared with a vibratome (Dosaka, Japan). ACSF consisted of (mM): 124 NaCl, 5.0 KCl, 1.25 NaH₂PO₄, 2.0 MgSO₄, 2.0 CaCl₂, 22 NaHCO₃ and 10 glucose, and was equilibrated with 95% O₂/5% CO₂.

MED64 multi-electrode apparatus (Panasonic) was used for the electrophysiological measurements (Oka *et al.* 1999; Shimono *et al.* 2000). After incubation for at least 1 h with ACSF at 25°C for recovery, slices were positioned on a novel custom multi-electrode probe (see Fig. S1 in Supplementary material) in which 64 planar microelectrodes were particularly arranged to cover densely the important regions containing essential synaptic contacts of pyramidal and granule neurons, i.e. the stratum radiatum in CA1, the stratum radiatum in CA3 and the molecular layer in dentate gyrus (DG). By using the current custom microelectrode arrangement, correct stimulation of both recurrent collateral fibers in the stratum radiatum of CA3 (recording at CA3 pyramidal neurons) and the medial perforant pathways in the molecular layer of DG (recording at DG granule cells) was performed with simultaneous stimulation of Schaffer collaterals in the stratum radiatum of CA1 (recording at CA1 pyramidal neurons) (Fig. S1). With a conventional checkered uniform arrangement of 64 microelectrodes (Oka *et al.* 1999; Shimono *et al.* 2000), simultaneous selection of the recurrent collateral fibers and the medial perforant pathways was not easy, because the density of electrodes over such particular regions was not sufficiently high. Bipolar constant current pulses (approximately 60 μ A, 0.1 ms) served as a test stimulus at three selected electrodes as shown in Fig. S1. The responses of excitatory postsynaptic potential (EPSP) were measured with three selected electrodes at which EPSP were filtered through a 1 Hz–10 kHz bandpass filter and recorded at a 20 kHz sampling rate. Estradiol (0.1–10 nM) was perfused (at 30°C with perfusion rate of 2 mL/min) for 30 min prior to the induction of LTD, which was induced by a transient perfusion of 30 μ M NMDA for 3 min (Lee *et al.* 1998). The significance of the estradiol or drug effect was analyzed at 60 min via statistical analysis using ANOVAS (* p < 0.05; ** p < 0.01).

Imaging and analysis of dendritic spine morphology

A schematic diagram of the spine analysis is presented in Fig. S2 in the Supplementary material.

Current injection of Lucifer Yellow

Hippocampal acute slices from adult male rats aged 12 weeks were prepared with a vibratome in a manner identical to that described in the LTD analysis. Hippocampal slices were transferred into an incubating chamber containing ACSF and held at 25°C for 2 h for recovery. Slices were then incubated with 0.1–10 nM estradiol or drugs such as PPT or PD98059. Next, the slices were pre-fixed with 4% paraformaldehyde in phosphate-buffered saline (PBS) at 4°C for 2–4 h. Neurons within slices were visualized by an injection of Lucifer Yellow (Molecular Probes, Eugene, OR, USA) under a Nikon E600FN microscope (Japan) equipped with a C2400–79H infrared camera (Hamamatsu Photonics, Japan) and with a 40 \times water immersion lens (Nikon). Dye was injected with a glass electrode whose tip was filled with 5% Lucifer Yellow for 15 min, using Axopatch 200B (Axon Instruments, USA). Approximately five neurons within a 100–200 μ m depth from the surface of a slice were injected (Duan *et al.* 2002). After labeling, slices were fixed again with 4% paraformaldehyde overnight.

Confocal laser microscopy and morphological analysis

The imaging was performed from sequential z-series scans with a MRC-1024 confocal microscope (Bio-Rad, Hercules, CA, USA) at high zoom (1.5–3.0) with a 60 \times oil immersion lens, NA 1.4 (Nikon). For Lucifer Yellow, the excitation and emission wavelengths were 488 nm and 515 nm, respectively. For analysis of spines, a three-dimensional image was reconstructed from approximately 40 sequential z-series sections of every 0.5 μ m with a 60 \times oil immersion lens, NA 1.4 (Nikon, Japan). The applied zoom factor (1.5–3.0) yielded 9.1–18 pixels per 1 μ m. The z-axis resolution was approximately 0.34 μ m. The confocal lateral resolution was approximately 0.18 μ m. Our resolution limits were regarded as sufficient to allow the determination of the density of thorns or spines. Confocal images were then deconvoluted using AUTODEBLUR software (AutoQuant, USA). In each slice, two to three neurons with more than 100 spines were analyzed, and at least 50 spines were counted on each frame. In total, n = 5 neurons, approximately 20 dendritic segments and n = 800–2000 total spines were analyzed for each drug treatment, except that n = 11 neurons were analyzed for control. Density and morphology of dendritic spines were analyzed by tracing neurons with NeuroLucida (MicroBrightField, USA). The single apical dendrite was analyzed separately. The spine density was calculated from the number of spines along secondary dendrites having a total length of 30–100 μ m. These dendrites were present within the stratum radiatum, between 100 and 200 μ m from the soma. All protrusions from the dendrites were treated as 'spines', although with confocal microscopy, it was not possible to determine whether they formed synapses, or whether some of them were filopodia protrusions which did not form synapses (Sorra and Harris 2000). While counting the spines in the reconstructed images, the position and verification of spines was aided by rotation of three-dimensional reconstructions and by observation of the images in consecutive single planes. Spine shapes were classified into four categories as follows (Sorra and Harris 2000). (1) A mushroom spine that has a large distinguishable head and neck. The head diameter (D) is greater than 0.6 μ m, and the total spine length (L) is less than twice the head diameter, i.e. $D > 0.6 \mu\text{m}$ and $L < 2D$. (2) A thin spine that has a small distinguishable head and an elongated spine neck. $D < 0.6 \mu\text{m}$,

$L > 2D$. (3) A filopodium that is a long filamentous protrusion with no distinguishable head. L is much longer than the neck diameter (d). (4) A stubby spine that does not have a distinguishable head, and has an L -value similar to d . The significance of the drug effect was examined using the Tukey–Kramer posthoc multiple comparisons test when one way ANOVA tests yielded $p < 0.05$.

RC-19 antibody preparation and purification

MC-20 is a widely used antiserum against the C-terminal of ER α , but it has serious problems in non-specific staining of ER α in the hippocampus of ERalphaKO mice. Therefore, purification of ER α -specific antibody fractions from antiserum was necessary. As purification of ER α -specific antibody fractions requires a large amount of antiserum, we prepared home-made purified antibody RC-19. The most C-terminal 19 amino acids (C-HSLQTYIIP-PEAEGFPNTI) of rat ER α conjugated with bovine serum albumin (BSA) via *m*-maleimidobenzoyl-*N*-hydroxysuccinimide ester (Sigma) were used as an antigen. Antibody was elicited in male white rabbits. A 40 ml aliquot of the collected sera from the rabbits was passed through a BSA immobilized Sepharose 4B column (1 \times 5.5 cm) containing 5 mg BSA to remove the antibody against BSA. Subsequently, the specific antibody against the peptide was adsorbed on the peptide-BSA immobilized Sepharose 4B column (1 \times 2 cm). The column was washed with 30 mL of 50 mM potassium phosphate buffer (pH 7.2) containing 1% (w/v) sodium cholate and 500 mM NaCl. RC-19 antibody was eluted with 100 mM glycine (pH 4.0) containing 0.15 M NaCl. The pH of the eluted antibody was adjusted to pH 7.4 by immediate mixing with 1 M Tris-HCl (pH 8.0) to prevent denaturation of antibody. The elution was performed within 30 min. Purified RC-19 was stored at -80°C , with 0.5% BSA, in many separate tubes with a small volume. Once a tube of fresh antibody had been used after thawing we never re-froze it for further use, but discarded it. Repetitive thawing and freezing of antibody should be avoided, otherwise the good immunoreactivity of RC-19 in hippocampal tissues is lost.

Preparation of synaptic, cytoplasmic and nuclear fractions

The same hippocampal slices were used in order to compare the results obtained with electrophysiology and spine morphology investigations. For western blot of hypothalamus, cerebral cortex and cerebellum, whole tissues of these brain regions (instead of slices) were used. Fractionation was performed by a combination of centrifugations at 4°C for the hippocampus, hypothalamus, cerebral cortex and cerebellum (Cohen *et al.* 1977). Detailed descriptions of centrifugation procedures to obtain the nuclear fraction, the PSD fraction, the presynaptic membrane-enriched fraction and the cytoplasmic fraction are described in *Supplementary Material*.

Western immunoblot analysis

Purified fractions prepared by centrifugation were suspended in 125 mM Tris-HCl buffer (pH 6.8) containing 5 mM 2-mercaptoethanol, 10% sucrose, 6% sodium dodecylsulfate (SDS) and 0.002% bromophenol blue. The fractions were subjected to electrophoresis using a 10% polyacrylamide gel. After transfer to polyvinylidene fluoride membranes (Immobilon-P; Millipore Co., Bedford, MA, USA), the blots were probed with purified antibody RC-19 (diluted to 1/2000), MC-20 antiserum (1/3000) or AS409 antiserum (1/2000, a gift from Prof. Shinji Hayashi) for 12–18 h at 4°C , and incubated

with horseradish peroxidase (HRP)-conjugated goat anti-rabbit IgG (Cell Signaling). The protein bands were detected with enhanced chemiluminescence (ECL) plus western blotting detection reagents (Amersham, Piscataway, NJ, USA). To obtain high quality images of chemiluminescence from protein bands using ECL plus, we used LAS3000 Image Analyzer (Fuji Film) with a 16-bit wide dynamic range.

Immunohistochemical staining of hippocampal slices

Immunohistochemical staining was performed essentially as described elsewhere (Kimoto *et al.* 2001; Kawato *et al.* 2002). Briefly, hippocampal slices were prepared from a rat or mouse deeply anesthetized with pentobarbital and perfused transcardially with PBS (0.1 M phosphate buffer and 0.14 M NaCl, pH 7.3), followed by fixative solution of 4% paraformaldehyde. The hippocampi were post-fixed, cryoprotected and frozen-sliced coronally at 20 μm thickness with a cryostat (Leica CM1510, Germany). Brains from several animals were used and from each brain, a single representative coronal section including the dorsal hippocampus was selected.

Staining for ER α was performed using the avidin-biotin peroxidase complex (ABC) technique. RC-19 was pre-treated with 5% BSA for 5 h to pre-absorb non-specific contaminated IgGs against BSA. After application of RC-19 IgG (1/1000) or MC-20 antiserum (1/500), the slices were incubated for 24 h at 4°C , in the presence of 0.5% Triton X-100 and 3% skim milk, with gentle shaking. Triton X-100 treatment was necessary to facilitate penetration of IgGs into cells in slices. Biotinylated anti-rabbit IgG (1/1000) in PBS was then applied, followed by a 30 min incubation with streptavidin-horseradish peroxidase complex (Vector Laboratories, Burlingame, CA, USA). Immunoreactive products were detected by immersing the slices in a detection solution (0.1 M Tris-HCl, pH 7.2, containing 0.05% diaminobenzidine, 0.1% H_2O_2 and 0.3% ammonium nickel sulfate). After dehydration and embedding in Entellan Neu (Merck), the immunoreactive cells in the slices were examined under a microscope, and digital images with a 2272×1704 pixel resolution were taken by a digital camera (COOLPIX4500, Nikon). For pre-absorption of RC-19 with antigen, excess amount of antigen was pre-incubated with RC-19 for 15 h at 4°C .

Post-embedding immunogold method for electron microscopy

Rat hippocampal slices were prepared in essentially the same manner as that used for the immunohistochemical staining, except that slicing was performed at 4°C using a vibratome instead of frozen slicing. Brains from five animals were used and from each brain, a single representative coronal section including the dorsal hippocampus was processed for ultrathin sectioning. Freeze substitution and low-temperature embedding of the specimens was performed as described previously (Adams *et al.* 2002). Briefly, slices were plunged into liquid propane. The samples were immersed in uranyl acetate in anhydrous methanol (-90°C), then infiltrated with Lowicryl HM20 resin (Electron Microscopy Sciences, USA); polymerization was performed with ultraviolet light. Ultrathin sections were cut using a Reichert-Jung ultramicrotome. For immunolabeling, sections were incubated with primary antibody for RC-19 (1/1000) or MC-20 (1/2000) in the above diluent overnight, then incubated with secondary gold-tagged (10 nm) Fab

fragment in Tris-buffered saline (TBS). Sections were counterstained with 1% uranyl acetate, and viewed on a JEOL 1200EX electron microscope (Japan). Images were captured using the CCD camera (Advanced Microscopy Techniques, USA). Controls omitting the primary antibody were performed and no immunogold labeling was observed.

Mass spectrometric analysis of RC-19 reacted ER α

To perform immunoprecipitation with RC-19, a hippocampal cytoplasmic fraction (1 mg/mL) was suspended in 5 mM HEPES buffer. A 20 μ L volume of protein-G-Sepharose beads (Sigma) covalently coupled with 10 μ g RC-19 by dimethyl pimelimidate (Pierce, Rockford, IL, USA) was incubated with 15 mg protein overnight at 4°C. The beads were eluted with sample buffer of SDS–polyacrylamide gel electrophoresis (PAGE) for 5 min at 100°C, and reduced with dithiothreitol (DTT). The eluted immunoprecipitant proteins were suspended in 125 mM Tris-HCl buffer (pH 6.8) containing 5 mM 2-mercaptoethanol, SDS and bromophenol blue, and subjected to SDS–PAGE. After protein separation, the gel was stained overnight with 0.1% silver using the silver stain MS kit (Wako, Japan). A silver-stained band of approximately 67 kDa was excised from the gel, de-stained with the de-staining solution of the silver stain kit, and the proteins digested in-gel with trypsin (Izumi *et al.* 2003). Digested peptides were extracted from gel pieces with 50 μ L 50% acetonitrile and 0.1% trifluoroacetic acid. A C18 microcolumn (Millipore) was used for purification of peptides. A matrix-assisted laser desorption/ionization (MALDI) time-of-flight (TOF) mass spectrometer (Bruker Daltonics, USA) was used to detect ER α protein directly. Samples were prepared by mixing 0.5 μ L peptide solution with 0.5 μ L matrix solution on a MALDI plate. All masses are reported as monoisotopic mass values. Peptides were identified using the Mascot search program (<http://www.matrixscience.com>) to perform theoretical trypsin digests, and to search for potential unmodified tryptic peptides (with up to one missed cleavages) or suspected modified species. Further details are described in Supplementary material.

RT-PCR and Southern hybridization

Total RNAs were isolated from rat brain tissues using a total RNA purification kit (Nippongene, Japan) (Furukawa *et al.* 1998). The purified RNAs were quantified on the basis of absorbance at 260 nm, and then treated with RNase-free DNase I to eliminate the possibility of genomic DNA contamination. RNA was reverse-transcribed, using a T-primed first-strand kit (Pharmacia, Sweden). Specific primer pairs for subregions of ER α exons (Exon 1'-2-3-4', Exon 3'-4-5', Exon 4'-5-6', Exon 5'-6-7' and Exon 6'-7-8'), and their distribution along exons, are illustrated in Fig. S6 in Supplementary material. Specific primer sequences of ER β (Exon 7'-8') and the PCR protocol for both molecules are described in Supplementary material. Exon 3'-4-5' primer pairs of ER α were used for semi-quantitative analysis performed for comparison of several brain regions and developmental analysis (Ivanova and Beyer 2000). To achieve specific detection of mRNAs, we chose primer pairs which had Gibb's free energy around -8 kcal/mol (forward five bases) and -9 kcal/mol (backward five bases) upon recombination with cDNA. The amplified RT-PCR products were directly cloned into TA-cloning vector (Promega, Madison, WI, USA), sequenced, and used as DNA probes of the following Southern hybridization. The

RT-PCR products were separated on 1.8% agarose gels, stained with ethidium bromide and then transferred to nylon membranes. Southern hybridization was performed with 32 P-labeled cDNA probes for ER α . The hybridization signals were measured using a BAS-1000 Image analyzer (Fuji film, Japan). Four independent analyses produced identical results. In addition to ethidium bromide staining, cloning of cDNA and Southern hybridization were used to ensure the detection of target cDNA.

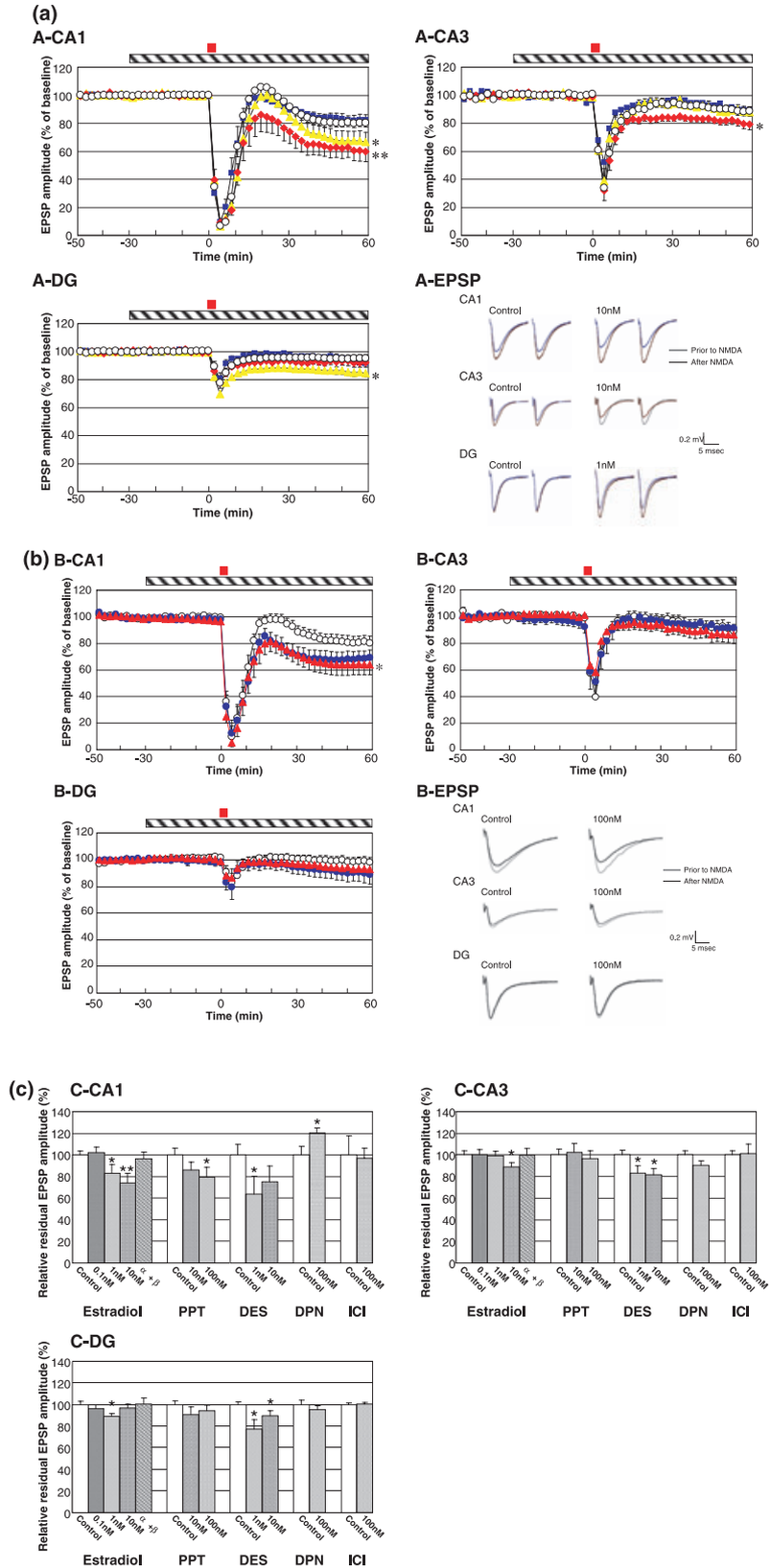
Results

Rapid effect of estradiol on long-term depression (LTD)

We carried out electrophysiological investigations to analyze the effect of 17 β -estradiol on LTD. LTD was simultaneously investigated in CA1, CA3 and DG of the same adult male hippocampal slices. Recordings were performed using novel 64 multi-electrodes particularly arranged to stimulate the Schaffer collaterals in the stratum radiatum of CA1, the recurrent collateral fibers in the stratum radiatum of CA3 and the medial perforant pathways in the molecular layer of DG (Fig. S1). LTD was induced pharmacologically by the transient application (3 min) of NMDA. Upon application of 30 μ M NMDA, the maximal amplitude of EPSP was transiently decreased to a minimal value, and then recovered to reach an approximate plateau level ($< 100\%$) within 30–50 min after NMDA application, indicating LTD establishment (Fig. 1a). This LTD was induced by the activation of phosphatase due to a moderate Ca $^{2+}$ influx through NMDA receptors (Lee *et al.* 1998). The plateau EPSP amplitude at 60 min was 80.4% (CA1), 88.8% (CA3) and 95.1% (DG). A 30 min pre-perfusion of 10 nM estradiol, for example, significantly enhanced LTD, resulting in the EPSP amplitude at 60 min of 59.7% (CA1), 79.1% (CA3) and 92.2% (DG) (Fig. 1a). The dose dependency of the enhancement effect by 17 β -estradiol was heterogeneous in different regions. LTD enhancement was least significant at 0.1 nM, moderately significant at 1 nM and most significant at 10 nM for 17 β -estradiol in both CA1 and CA3. In DG, however, LTD enhancement was most significant at 1 nM, and a smaller enhancement was observed at 0.1 nM and 10 nM for estradiol. Investigations using specific estrogen agonists indicated that the contribution of ER α (but not ER β) was essential to these estradiol effects. PPT (ER α agonist) at 100 nM exhibited a significant LTD enhancement in CA1 (Figs 1b and c), while DPN (ER β agonist) did induce a suppression of LTD in CA1 (Fig. 1c), suggesting a contribution of ER β opposite to that of ER α in the observed estradiol effect on LTD.

The co-perfusion of 10 nM 17 α -estradiol (used as an antagonist) with 1 nM 17 β -estradiol suppressed the 17 β -estradiol effect on LTD. The co-perfusion of 1 μ M ICI with 10 nM 17 β -estradiol, however, did not block the 17 β -estradiol-induced enhancement of LTD. DES at 1–10 nM enhanced LTD, and DES was more effective at

Fig. 1 (a) Rapid modulation by 17 β -estradiol of LTD in the CA1 (A-CA1), CA3 (A-CA3) and DG (A-DG) of the same hippocampal slice. Estradiol concentration was 0 nM (open circle, $n = 17$) and 0.1 nM (blue filled square, $n = 8$), 1 nM (yellow filled triangle, $n = 11$) and 10 nM (red filled diamond, $n = 10$), respectively; $n =$ number of independent experiments. Vertical axis indicates maximal amplitude of EPSP. Here, 100% refers to the EPSP value at $t = -40$ min prior to NMDA stimulation, irrespective of the test condition. LTD was induced by 30 μ M NMDA perfusion at time $t = 0-3$ min (filled bar above the graph). Hatched bar above the graph indicates period of time during which estradiol was administered. The significance of the estradiol effect was confirmed at 60 min via statistical analysis using ANOVA ($*p < 0.05$; $**p < 0.01$) as indicated in the figure. Illustrated data points and error bars represent the mean \pm SEM from n independent slices. A-EPSP = EPSP traces in the presence of 10 nM estradiol, showing sample recordings taken prior to ($t = -40$ min) and after ($t = 60$ min) NMDA stimulation. (b) Rapid modulation by PPT of LTD in the CA1 (B-CA1), CA3 (B-CA3) and DG (B-DG) of the same hippocampal slice. PPT concentration was 0 nM (open circle, $n = 10$), 10 nM (blue filled square, $n = 8$) and 100 nM (red filled triangle, $n = 12$), respectively. B-EPSP = EPSP traces in the presence of 100 nM PPT, showing sample recordings taken prior to and after NMDA stimulation. (c) Comparison of modulation effect of LTD by 17 β -estradiol and agonists in the CA1, CA3 and DG of hippocampal slices. Vertical axis is relative EPSP amplitude at $t = 60$ min, where EPSP amplitude at $t = 60$ min of the control slice without drug application is taken as 100%. From left to right, 17 β -estradiol (Estradiol), PPT, DES, DPN and ICI at indicated concentrations. In Estradiol, $\alpha + \beta$ indicates that 10 nM 17 α -estradiol was perfused with 1 nM 17 β -estradiol. The significance of the estradiol effect was confirmed at 60 min via statistical analysis using ANOVA ($*p < 0.05$; $**p < 0.01$) as indicated in the figure.



1 nM than 10 nM. The GPR30 agonist, 100 nM ICI, did not affect the NMDA-induced LTD. In Fig. 1(c), a quantitative comparison was shown by normalizing the EPSP value of the control LTD curve at 60 min to 100%. This normalization was performed for all pairs of control and drug-treated slices because the control LTD responses varied slightly between different drug treatment pairs, depending on differences in the animals. Here, we investigated LTD in particular because LTD has not been extensively analyzed compared with long-term potentiation. The comparison between three subregions indicated that estradiol modulation of LTD is more effective in CA1 than in CA3 and DG. The enhancing effect of LTD by estradiol suggests that estradiol could facilitate erasing processes of memory. Because the baseline EPSP was stable during the 120 min estradiol perfusion (in the case of absence of NMDA application), newly-generated spines (see below) during the 120 min incubation with estradiol probably did not form active synapses that contribute EPSP and LTD signals.

Rapid effect of estradiol on spinogenesis

Next, we analyzed the ability of 17β -estradiol to modulate spine density and morphology. To do this, single spine imaging was performed for Lucifer Yellow-injected neurons in hippocampal slices from 12-week-old male rats. These investigations were independent of synaptic transmission investigations such as LTD and LTP, which probe the characteristics of pre-formed synapses. The single spine imaging not only probes pre-existing spines but also, newly-formed spines upon estradiol treatment. The spinogenesis investigations also analyzed both spine synapses (spines forming synapses) and free spines (spines without forming synapses), whereas in the LTD measurements we obtained information about a particular population of spines that form synapses (spine-synapse). As we found the greatest modulation effect of estradiol in CA1 in the LTD experiments, we focus here on CA1 neurons for analysis of spinogenesis. It should be noted that the analysis of spinogenesis in CA3 has been reported elsewhere (Tsurugizawa *et al.* 2005).

We analyzed secondary branches of the apical dendrites located 150–250 μm distant from the pyramidal cell body around the middle of the stratum radiatum of CA1 region. Following a 0.5–2 h treatment with 17β -estradiol, treated dendrites have significantly more spines than control dendrites (i.e. with no estradiol). Time dependency was examined by treating slices for 0.5, 1 and 2 h with 1 nM estradiol. The enhancing effect on the total spine density was approximately proportional to the incubation time, showing 1.05 (0.5 h), 1.13 (1 h) and 1.31 spines/ μm (2 h). Dose dependency was also examined after a 2 h incubation, and showed that the enhancing effect was most significant at 1 nM estradiol (1.31 spines/ μm) compared with 0.1 nM (1.03 spines/ μm) and 10 nM (1.25 spines/ μm) estradiol. Because a

2 h treatment with 1 nM estradiol was most effective for spinogenesis, this incubation time was used in the following investigations unless specified. The different contributions of ER α , ER β and GPR30 in the effect of estrogen can be investigated using the ER α -specific agonist, PPT, the ER β -specific agonist, DPN (Harrington *et al.* 2003) and the GPR30-specific agonist, ICI (Thomas *et al.* 2005).

A 2 h treatment increased the total spine density to 1.31 spines/ μm (1 nM estradiol) compared with 0.85 spines/ μm (control, 0 nM estradiol) (Figs 2a and b). PPT (Harrington *et al.* 2003) at 100 nM induced a significant enhancement of spine density to 1.20 spines/ μm (Fig. 2b). However, DPN (Harrington *et al.* 2003) at 100 nM increased the spine density only slightly (0.95 spines/ μm). Diethylstilbestrol (DES, agonist of ER α and ER β) significantly increased spine density. A stronger effect was observed for 1 nM DES (1.29 spines/ μm) than 10 nM DES (0.99 spines/ μm). Blocking of ER α by 1 μM ICI 182,780 and of NMDA receptors by MK-801 completely suppressed the enhancing effect of estradiol on spine density. Blocking phosphorylation of mitogen-activated protein (MAP) kinase by 50 μM PD98059 completely prevented estradiol-induced spinogenesis. Blocking Src kinase by PP2 did not significantly suppress the estradiol effect. Blocking of AMPA receptors by CNQX at 20 μM only partially prevented the estradiol-enhancing effect by decreasing spine density to 1.07 spines/ μm . Washout of estradiol for an additional 2 h after the 2 h estradiol treatment abolished the estradiol-induced enhancement of spinogenesis, implying that these actions are reversible. The specific ligand of GPR30, 100 nM ICI and tamoxifen (Thomas *et al.* 2005), did not significantly increase spine density (Fig. 2b).

The morphological changes in spines induced by 2 h drug treatments were quantitatively assessed (Figs 2d,e). In the control slices (0 nM estradiol), the relative population of spines was approximately 24% for mushroom spines, 62% for thin spines, 1% for filopodia and 13% for stubby spines (Fig. 2c). Upon estradiol treatment, the densities of thin spines and filopodia were increased significantly, from 0.57 spines/ μm and 0.01 spines/ μm to 0.97 spines/ μm and 0.11 spines/ μm , respectively, while the densities of mushroom spines (from 0.16 to 0.12 spines/ μm) and stubby spines (from 0.10 to 0.11 spines/ μm) were not significantly altered within experimental error (Fig. 2d). As a result, the average spine diameter became significantly smaller following estradiol application (Fig. 2e). PPT induced a change nearly identical to that obtained with estradiol. Blocking by PD98059, ICI and MK-801 completely suppressed the enhancing effect of estradiol on the density of thin spines (Fig. 2d).

We estimated the population of spine-synapses that are spines forming synapses. In control slices, approximately $47.3 \pm 9.8\%$ of total spines, $61.8 \pm 12.7\%$ of mushroom spines, $45.2 \pm 9.2\%$ of thin spines, 0% of stubby spines and

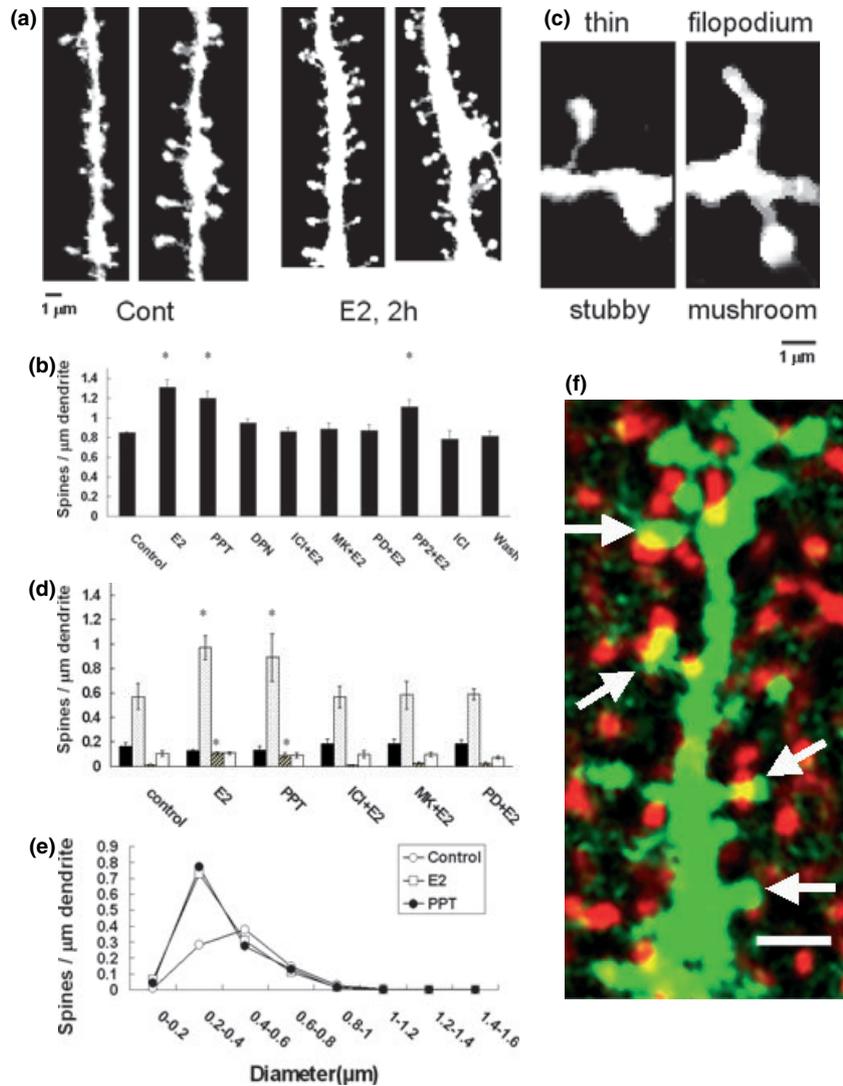


Fig. 2 Changes in the density and morphology of spines upon treatment with 17 β -estradiol (E2) and drugs in hippocampal slices. Spines were analyzed along the secondary dendrites of pyramidal neurons in the stratum radiatum of CA1. (a) Maximal intensity projections onto XY plane from z-series confocal micrographs, showing spines along the secondary dendrites of hippocampal CA1 pyramidal neurons. Control spines without drug treatments (Cont) and spines after E2 treatment (E2). Bar = 1 μm . (b) Effect of drug treatments on the total spine density in CA1 neurons. Vertical axis is the average number of spines/1 μm . A 2 h treatment in ACSF without drugs (Control), or with 1 nM E2 (E2), 100 nM PPT (PPT), 100 nM DPN (DPN), 1 nM E2 and 1 μM ICI 182,780 (ICI + E2), 1 nM E2 and 50 μM MK-801 (MK + E2), 1 nM E2 and 50 μM PD98059 (PD + E2), 1 nM E2 and 25 μM PP2 (PP2 + E2), 100 nM ICI (ICI) or 1 nM E2, followed by 2 h washout with ACSF (Wash). (c) Four different spine subtypes are shown: mushroom type, thin type, filopodium type and stubby type. Bar = 1 μm . (d) Density of four subtypes of spines. A 2 h treatment in ACSF without drugs (Control group), or with 1 nM E2 (E2 group), 1 nM E2 and 1 μM ICI (ICI + E2 group), or 1 nM E2 and 50 μM MK-801 (MK + E2 group). In each group, from left to right: mushroom (black column)

column), filopodium (hatched column) and stubby (open column). (e) Histogram of spine head diameters after a 2 h treatment in ACSF without drugs (Control, open circle), or with 1 nM E2 (open square) and 100 nM PPT (filled circle). (f) Distribution of spine-synapses in slices. Synapse formation is indicated as yellow regions that are superimposed regions of spines visualized with Lucifer Yellow (green) on axon terminals stained with anti-synapsin I IgG (red). Spine-synapses are indicated by arrows. Other yellow regions may be shaft-synapses, which are synaptic contacts ending directly on the dendrite. Lucifer Yellow was injected into a single neuron in a slice (400 μm thick), then the slice was further frozen-sliced (30 μm thick) and immunostained with anti-synapsin I IgG (1/500). Distribution of secondary antibody conjugated with Cy3 was analyzed by confocal microscopy. Scale bar = 2 μm . In (b), (d) and (e), results are reported as mean \pm SEM from $n = 11$ (control) or $n = 5$ (drug-treated) different neurons, and for each neuron 150–300 spines were analyzed. The significance of the drug effects was confirmed via statistical analysis using ANOVA ($*p < 0.05$) between control and drug-treated samples as indicated in the figure. In (b), (d) and (e), vertical axis is the average number of spines/1 μm of dendrite.

0% of filopodia were observed to form synapses, as judged from the superimposed region of immunostaining with synapsin I (presynaptic marker) on Lucifer Yellow-injected spines ($n = 5$ neurons, approximately 100 spines) (Fig. 2f). The rest of the spines were free spines without partners of axon terminals.

Western blot analysis of ER α in synaptic and nuclear fractions

Because the 67 kDa ER α protein is present in extremely low levels in the brain relative to the ovary, non-specific IgGs contained in the antisera used in histochemical investigations may result in serious non-specific binding, producing misleading results. Accordingly, the specific reactivity of ER α antibodies should be carefully tested via western immunoblot analysis. We therefore used a novel affinity column-purified RC-19 antibody raised against the 19 C-terminal residues of rat ER α . RC-19 was used as a type of purified MC-20. We also performed careful purification of synaptic membranes, cytoplasm and nuclei using a combination of density gradient centrifugations for the hippocampus and hypothalamus from adult 12-week-old male rats. Using RC-19 (1/2000 dilution), only a single protein band was observed, with a molecular mass of approximately 67 kDa, in the PSD and cytoplasmic fractions as well as the nuclear fraction (Fig. 3a). As a positive control, the 67 kDa band was stained for rat ER α protein expressed in COS-7 cells and ovary from a 4-week-old female rat. The excellent quality of RC-19 was demonstrated by the absence of immunoreactivity in the vicinity of 67 kDa for the hippocampus and hypothalamus from 8-week-old ER α knockout (ER α KO) mice, used as a negative control (Fig. 3c). Extended analysis indicated that ER α was certainly localized in the purified PSD fraction, which was characterized by PSD-95 (Fig. 3b). It should be noted that MAP kinase and Src kinase were both observed in the PSD fraction. ER α was observed in the presynaptic membrane-rich fraction (PRE, low density synaptic membrane fraction, characterized by synaptophysin) and in the cytoplasmic fraction (CYT). The integrated total amount of ER α was greater in the CYT fraction than in other PSD and PRE fractions. It should be noted that the densest band of ER α that appeared in the PSD fraction (Fig. 3b) was simply due to the application of a threefold excess amount of protein in PSD compared with CYT and PRE. The 67 kDa band was not observed when RC-19 antibody was pre-adsorbed with ER α antigen peptides (30 μ g/mL).

Western blot with other antisera

For comparison, the reactivity was examined for other antisera widely used in earlier studies. Unfortunately, these antisera failed to react primarily with the 67 kDa ER α protein in the hippocampus. When MC-20 (antiserum against the most C-terminal 20 amino acids of mouse ER α) or

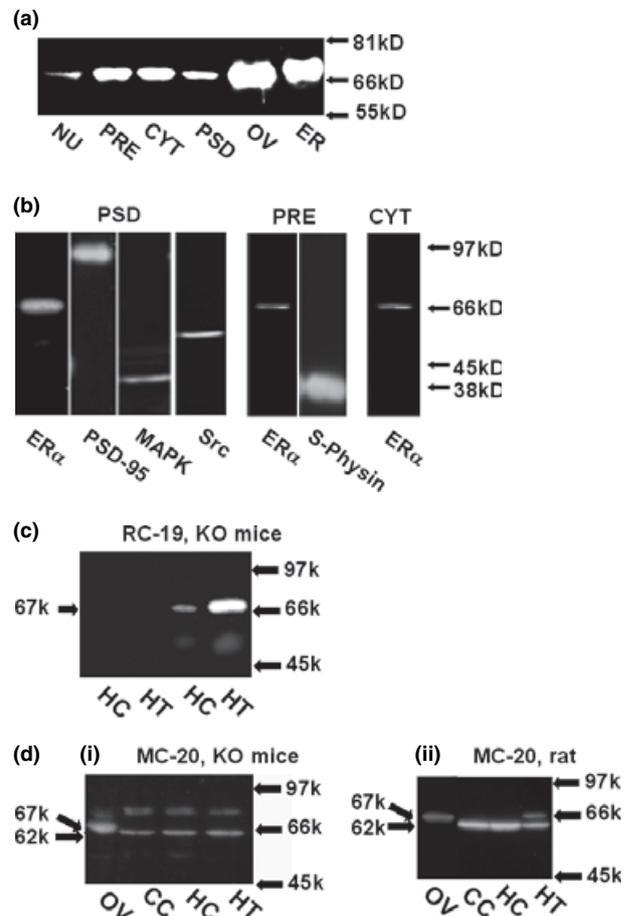


Fig. 3 Western immunoblot of ER α in subcellular fractions of male rat hippocampus. (a) Blot with RC-19 antibody of the subcellular fractions from adult rat hippocampus. From left to right: the nuclear fraction (NU), presynaptic membrane-rich fraction (PRE), cytoplasmic fraction (CYT), postsynaptic density fraction (PSD), ovary (OV) and ER α expressed in COS-7 cells (ER). The amount of protein applied to the gels was 20 μ g for each hippocampal fraction, and 1 μ g for ER α -expressed COS-7 cells and ovary. Ovary was taken from a 4-week-old rat. ER α genetically expressed in COS-7 was used as a positive control. (b) Western blot of PSD, PRE and CYT of the hippocampus. From left to middle, blot of PSD with RC-19 IgG (ER α), PSD-95 IgG (PSD-95), MAP kinase IgG (MAPK) and Src IgG (Src). From middle to right, blot of PRE with RC-19 (ER α) and synaptophysin IgG (S-Phyisin). At rightmost lane, blot of CYT with RC-19 (ER α). It should be noted that the amount of protein applied was 20 μ g for each lane, but 60 μ g for the PSD fraction. The densest staining of ER α was therefore shown in PSD, although the staining of ER α in PSD was weaker than that in the CYT and PRE fractions. (c) Staining with RC-19 of tissue homogenates from ER α KO mice and wild mice. From left to right, hippocampus (HC) and hypothalamus (HT) from ER α KO mice, HC and HT from wild mice. (d) (i) Staining with MC-20 antiserum of tissue homogenates from adult ER α KO mice brain. From left to right, ovary (OV, 4-week-old rat), cerebral cortex (CC), HC and HT. (dii) Staining with MC-20 antiserum of tissue homogenates from adult male rat brain. From left to right, OV from rat, CC, HC and HT. The amount of protein applied to the gels was 20 μ g for each lane in panels (c) and (d).

AS409 (polyclonal antiserum against rat ER α expressed in *Escherichia coli*) (Weiland *et al.* 1997) was used, the molecular weights of the main immunoreactive bands were different from 67 kDa. Genetically-expressed ER α or ovary was used as a control to differentiate 67 kDa from 62 kDa. MC-20 (1/3000) yielded a very weak 67 kDa band and a major, approximately 62 kDa band (Fig. 3d) (Toran-Allerand *et al.* 2002), whereas AS409 (1: 2000) gave a weak 67 kDa band and a major, approximately 90 kDa band (Fig. S3 in Supplementary material). Note that MC-20 also demonstrated a significant reactivity with a 62 kDa band in the hippocampus of ER α KO mice (Fig. 3d).

In the ER α KO mice (exon 2 deleted ER α - Δ 2 KO) used in this study, no ER α RNA containing transcripts of any of the exons located downstream of exon 2 was detected by RT-PCR, implying that this 62 kDa protein could not be a splice variant of ER α (Dupont *et al.* 2000; Pendaries *et al.* 2002). Our ER α KO mice are essentially different from ER α -

Neo KO mice (constructed by Neomycin insertion into exon 1) that have expressed an ER α splice variant, i.e. N-terminal modified ER α (61 kDa) (Couse *et al.* 1995; Kos *et al.* 2002).

Western blot examination for RC-19, MC-20 and AS409 has shown that the use of antibody purified from antiserum is essential to obtain real staining of ER α in the hippocampus.

Immunohistochemistry of ER α in hippocampal principal neurons

Immunohistochemical staining of ER α proteins via column-purified RC-19 antibodies (1/1000) was performed to determine the cell-specific localization of ER α in the adult male rat hippocampus. An intense immunoreactivity was observed in extranuclear, as well as nuclear regions of principal neurons in the CA1–CA3 and the dentate gyrus (DG) regions (Fig. 4a). Pre-adsorption of the antibody with an excess of ER α antigen (30 μ g/mL) resulted in the

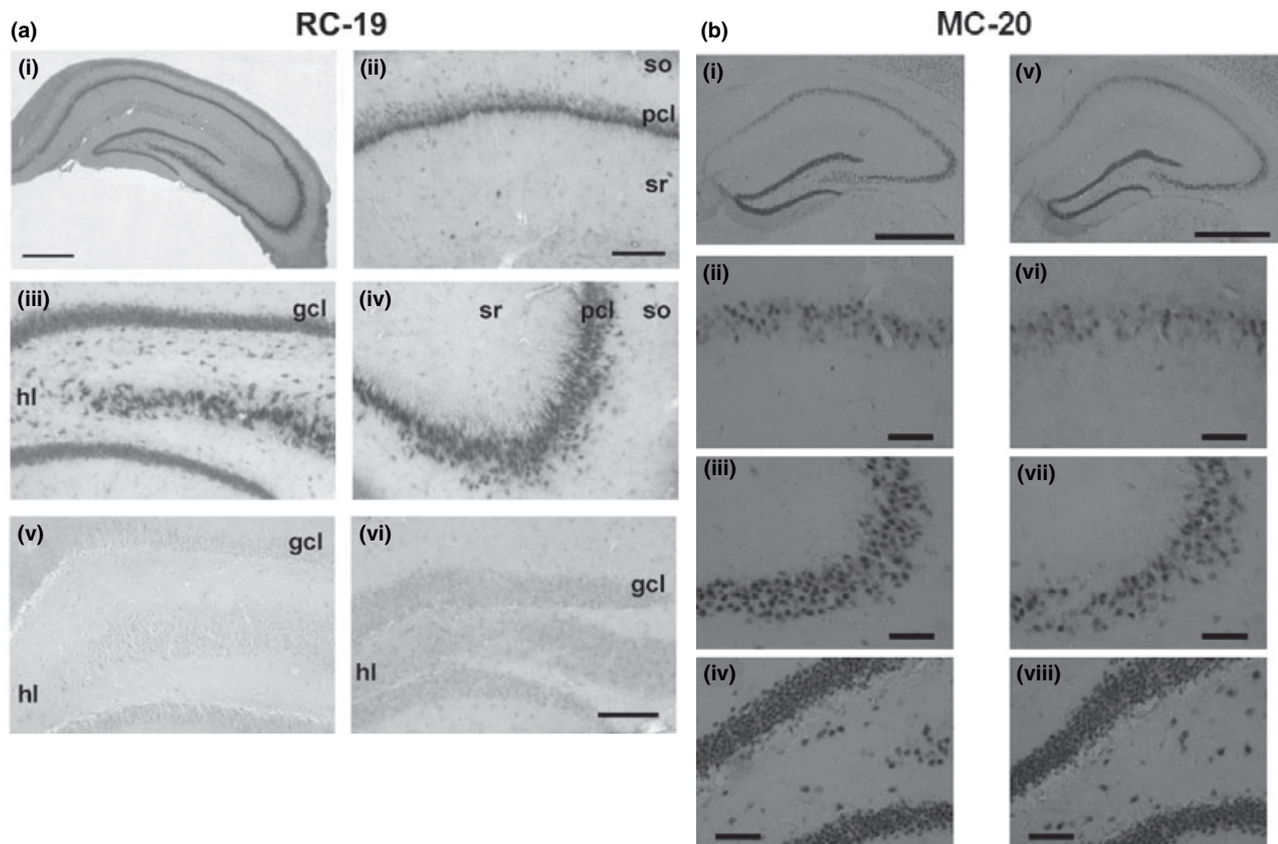


Fig. 4 (a) Immunohistochemical staining of ER α with RC-19 antibody in the hippocampal slices of adult male rat (i–v) and adult ER α KO mouse (vi). (i) Coronal section of the whole hippocampal formation; (ii) CA1; (iii) DG; (iv) CA3; (v) DG incubated with RC-19 antibody pre-adsorbed with antigen of C-terminal 19 amino acids of ER α ; (vi) the DG of ER α KO mouse. Abbreviations: so, stratum oriens; pcl, pyramidal cell layer; sr, stratum radiatum; gcl, granule cell layer; hl, hilus.

Scale bar = 500 μ m in (i) and 200 μ m in (ii)–(vi). Several independent experiments were performed. (b) Immunohistochemical staining with MC-20 antiserum in the hippocampal slices of adult ER α KO (i–iv) and wild male mice (v–viii): (i), (v) coronal section of the whole hippocampal formation; (ii), (vi) CA1 region; (iii), (vii) CA3 region; (iv), (viii) dentate gyrus. Scale bar = 50 μ m for (i) and (v); 10 μ m for (ii–iv) and (vi–viii). Three different batches of MC-20 were used.

complete disappearance of immunoreactivity in the hippocampus. No positive immunoreactivity was obtained using RC-19 in the hippocampus from ER α KO mouse (Fig. 4a). MC-20 (1/500) without purification, on the other hand, showed a significant reactivity for principal neurons in the hippocampal slices from ER α KO mouse (Fig. 4b) as well as from adult male rat (Fig. S4 in Supplementary material). Here, three independent batches of MC-20 showed nearly identical results. Note that RC-19 immunoreactivity was not observed in glial-like cells from immunohistochemical and immunogold investigations, excluding a significant contribution of glial cells.

Ultrastructural analysis for synaptic and nuclear localization of ER α

To explain the site of rapid modulation of spinogenesis and LTD of glutamatergic neurons by the activation of ER α , clarification of subcellular localization (particularly synaptic localization) of ER α in glutamatergic neurons is essential. The extranuclear and nuclear localization of ER α was clarified via ultrastructural investigations using RC-19 (1/1000). An immunoelectron microscopic analysis using post-embedded immunogold was performed to determine the localization of ER α immunoreactivity in the hippocampal neurons of adult male rats. ER α was localized not only in the nuclei but also, in both the axon terminals and dendritic spines of principal neurons (Fig. 5, Fig. S5 in Supplementary material). Gold particles were clustered in the postsynaptic and presynaptic compartments, as well as in the nuclei, dendrites and cytoplasm. At postsynapses, gold particles were distributed within the cytoplasm of the spine head. In some cases, gold particles were affiliated within the postsynaptic density. Significant labeling along dendrites was also found (Fig. 5), although many fewer gold particles were observed in axons. In glial-like cells, gold particles were not clearly recognized. Multiple labeling (three or more) of immunogold in the pre- and postsynaptic compartments was confirmed to ensure the specific labeling. A search for immunogold-labeled ER α proteins was performed for at least 30 synapses for each CA1, CA3 and DG region from more than 100 independent images. Almost no mitochondrial labeling was observed, indicating very low background non-specific labeling as RC-19 did not recognize the ER α band in the mitochondria by western blot. The topological distributions of gold particles within the neurons of the CA1, CA3 and DG were essentially identical. Pre-adsorption of the antibody with ER α antigen (30 μ g/mL) resulted in the disappearance of immunoreactivity. MC-20 immunoreactivity was strong in synapses and MC-20 (1/2000) often stained synaptic mitochondria (Fig. S5). These results raised some doubts about the staining specificity of MC-20. It should be noted that to date, ultrastructural investigations have suggested the presence of ER α mainly in GABAergic interneurons and cholinergic neurons using AS409 antiserum

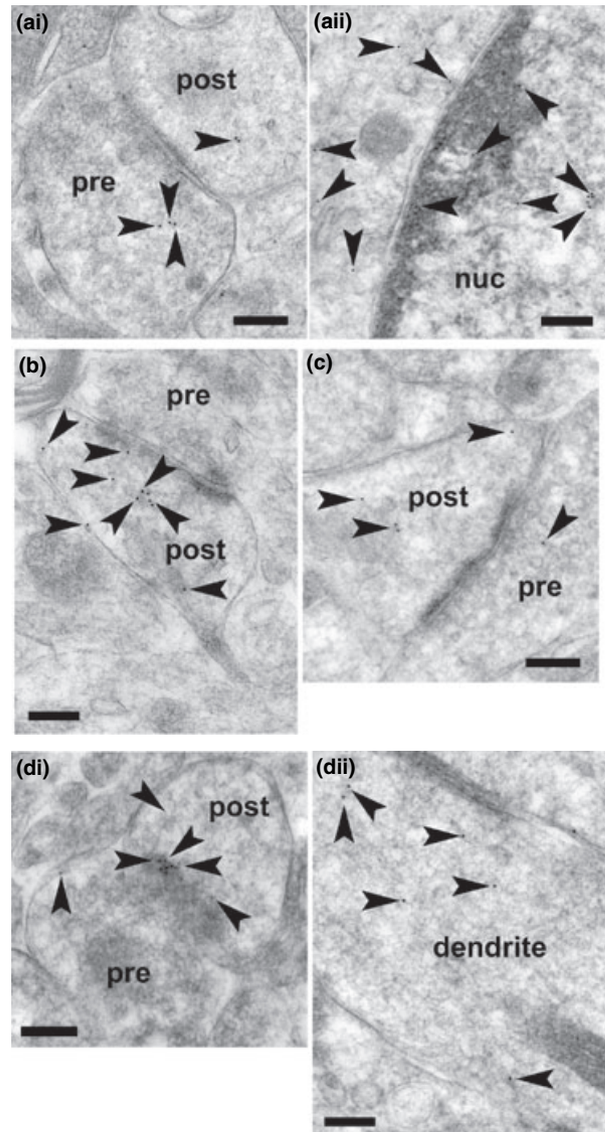


Fig. 5 Immunoelectron microscopic analysis, using RC-19 antibody, of the distribution of ER α within the axospinous synapses and nuclei of the hippocampal principal neurons in the stratum radiatum of CA1 (a), stratum radiatum of CA3 (b) and hilus of DG (c) regions. Representative images are shown from approximately 150 photographs from 32 independent slices from five animals. Gold particles (arrowheads) were localized in the pre- and postsynaptic regions (ai, b and c). In dendritic spines, gold particles were found within the spine head and in some cases, associated with PSD regions (di, axospinous synapses of the hippocampal principal neurons in the stratum radiatum of CA1). Gold particles were also localized in the nuclei (aii). In dendrites, gold particles were often found in the cytoplasmic space (dii, DG region). A 1 : 1000 dilution of RC-19 was used to prevent non-specific labeling. Abbreviations: pre, presynaptic region; post, postsynaptic region; nuc, nucleus. Scale bar = 200 nm.

(Milner *et al.* 2001; Adams *et al.* 2002; Towart *et al.* 2003), with one report showing ER α in pyramidal neurons (Adams *et al.* 2002).

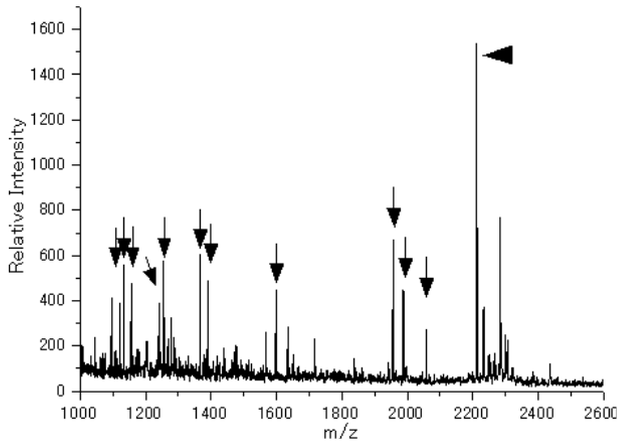


Fig. 6 MALDI-TOF mass spectrum of tryptic peptides from the 67 kDa band obtained after the separation of RC-19 reactive precipitants of the cytoplasmic fractions of hippocampus using SDS-PAGE. Eleven peptides appeared in the spectrum and were attributed to tryptic peptides from ER α as presented in Table 1. The masses matched with tryptic peptides of ER α are indicated by arrows. A large peak at m/z 2211 (arrow head) is the tryptic fragment of trypsin used for validation.

Mass spectrometric identification of RC-19-reacted ER α

To prove that the RC-19-reactive protein in the above experiments was ER α , mass spectrometric analysis was performed using the cytoplasmic fraction of the hippocampus. RC-19-reactive protein was immunoprecipitated, and an approximately 67 kDa protein was separated by SDS-PAGE. The protein was treated by in-gel protease (trypsin) digestion, and digested peptides were analyzed by MALDI-TOF mass spectrometry. Peptides were identified using the MASCOT search program (Matrix Science). As indicated in Fig. 6, more than 20 tryptic peptides were detected in mass spectrum, and 11 of the major peaks were attributed to the tryptic peptides from ER α . The identified peptides are listed in Table 1. These results strongly suggest that the RC-19-reacted protein is real ER α in the hippocampus.

Analysis of ER α and ER β mRNA

The possibility of the presence of a splice variant of ER α mRNA, corresponding to the 62 kDa protein detected by MC-20, was excluded in wild-type adult male rat hippocampus using RT-PCR combined with western blot analysis. The potential presence of isoforms of ER α is indicated by the observed reaction of ER α antisera (e.g. MC-20, AS409) with bands of approximately 62 and 90 kDa. ER α is transcribed from eight exons, denoted 1–8. The start and stop codons, corresponding to the ER α N- and C-termini, are included in exons 1 and 8, respectively. Several sets of forward and reverse primer pairs (Fig. S6) were used to examine the transcripts from several distinct subregions of mRNA. No PCR products lacking any of exons 2–7 were observed (Fig. 7). It should be noted that the inclusion of exon 8 was not explicitly examined using RT-PCR because the most

Table 1 Mass spectrometric identification of tryptic peptides from the 67 kDa protein, separated by SDS-PAGE of the RC-19 reacted-precipitants of the cytoplasmic fraction. The observed mass and the corresponding tryptic peptides of ER α are listed with the deviation from the theoretical mass (see Fig. 6)

Observed (m/z)	Deviation	Peptide sequence
1097.41	-0.03	NEMGTSGDMR (283–292)
1122.50	0.01	GMEHLYNMK (526–534)
1132.62	0.0	DHIHRVLDK (478–486)
1240.66	-0.01	ELVHMINWAK (358–367)
1268.73	0.03	INDTLIHLMAK (487–497)
1366.69	0.04	GNMIMESAKETR (177–188)
1389.94	0.05	LAQLLLSHIR (509–520)
1598.80	-0.01	KCYEVGMMGGIRK (249–262)
1956.10	0.67	SIILLNSGVYFLSSTLK (455–472)
1987.21	0.0	LAQLLLSHIRHMSNK (509–525)
2057.91	-0.12	KCYEVGMMKGGIRKDRR (249–265) 2 oxidation (Met)

C-terminal subregion must be included for western blot-reacted protein with C-terminal antibody RC-19. Although we did not investigate exon 1 deleted form of mRNA, 46 kDa protein should be expressed as reported in uterine tissue (Pendaries *et al.* 2002) in case of the expression of exon 1 deleted splice variant. However, scarcely any 46 kDa protein was observed in the hippocampus and hypothalamus, as judged from western blot with RC-19 (Fig. 3b). The combination of mRNA and western blot analyses therefore demonstrated that splice variants with molecular weights of approximately 62 kDa were not present in the wild-type rat hippocampus. It should be noted that no splice variants of ER α mRNA were expressed in our ER α KO mice (Dupont *et al.* 2000; Pendaries *et al.* 2002).

The expression levels of ER α transcripts in the hippocampus and other brain regions were compared by semi-quantitative RT-PCR using primer pairs designed for the region of exon 3'–4'–5'. The relative levels of ER α mRNA expression observed in various brain regions were approximately 1/20 (hippocampus), 1/2 (hypothalamus), 1/15 (cerebral cortex) and 1/80 (cerebellum) compared with levels in the 12-week-old ovary (Fig. S7A in Supplementary material). The relative levels of ER α mRNA transcripts in the hippocampus varied with age, from 100% (post-natal 10-day-old) to approximately 48% (4-week-old) and approximately 56% (adult) for the male rat. In addition, the relative level of ER α mRNA expression was not significantly different between male and female rat hippocampus from investigations of approximately 10 animals. These results were consistent with western blot investigations with RC-19, which also showed almost the same levels of ER α protein in the adult hippocampal homogenates of male and female rats.

The expression levels of ER β transcripts in the hippocampus and other brain regions were also compared using

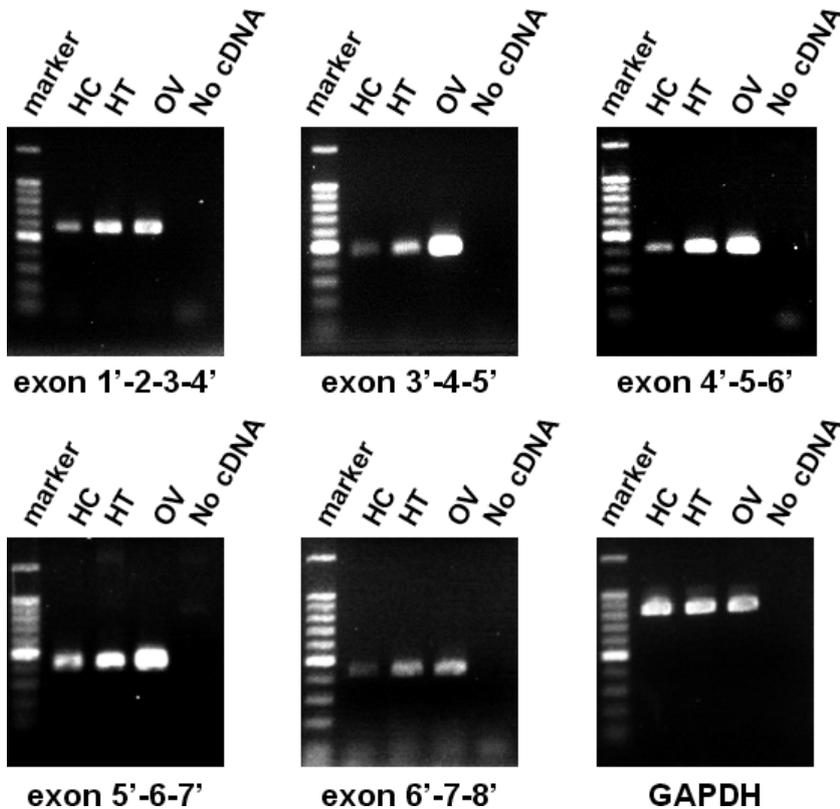


Fig. 7 Investigations regarding the absence of splice variants of ER α mRNA. RT-PCR of mRNA subregions of ER α in the adult rat hippocampus (HC), hypothalamus (HT) and 4-week-old rat ovary (OV). Five subregions were analyzed. Subregion 1'-4' contained full exons 2 and 3, in addition to partial sequences of nearest exons 1 and 4, as described in the primer pair design (see Supporting information). Subregions 3'-4-5', 4'-5-6', 5'-6-7' and 6'-7-8' contained full exon 4, exon 5, exon 6 and exon 7, respectively, in addition to partial sequences of the nearest exons. RT-PCR products (each from 50 ng of template) were visualized with ethidium bromide. Left-most lane shows a marker (100 bp ladder); right-most lane is water, used as a negative control.

primer pairs designed for the region of exons 7 and 8. The relative levels of ER β mRNA expression were approximately 1/80 (hippocampus), 1/20 (hypothalamus), 1/50 (cerebral cortex) and 1/20 (cerebellum) compared with levels in the 12-week-old ovary (Fig. S7B).

The relative expression level of ER α mRNA was approximately 10-fold that of ER β mRNA in the adult hippocampus, as judged from the PCR cycles needed for ER α (34 cycles) and ER β (36 cycles) (Figs. S7A and S7B). This level of ER α 10-fold higher than ER β is in good agreement with the observation that the mode of modulation by estradiol of LTD and spinogenesis is mostly the same as that by the ER α agonist, PPT, and very different from that by the ER β agonist, DPN.

Discussion

The current study demonstrates, in pyramidal and granule neurons from the acute slices of adult male hippocampus, the postsynaptic localization of ER α , and its modulatory function in rapid synaptic transmission (LTD) and spinogenesis of glutamatergic neurons. These two types of synaptic plasticity reflect, in principle, different processes triggered by estradiol.

Estradiol modulation of LTD occurred only in pre-existent synapses because spines newly generated by estradiol treatments did not form new synapses within 2 h, as judged

by no increase in the baseline magnitude of the EPSP signal during 2 h (data not shown). However, estradiol generated new thin spines that served as bases for new spine-synapse formation after, for example, more than 24 h (Pozzo-Miller *et al.* 1999). We used isolated hippocampal slices because we focused on the effect of estradiol on the glutamatergic circuits of the hippocampus. In this sense, the current results have a meaning different from many previous *in vivo* studies that investigated the effect of gonadal estradiol on the hippocampus via serotonergic or cholinergic afferents from the limbic system (Woolley *et al.* 1997; Rudick and Woolley 2000, 2003; Leranath *et al.* 2003; MacLusky *et al.* 2005). Both LTD and spinogenesis are enhanced by either 17 β -estradiol or the ER α agonist, PPT. In contrast, application of the ER β agonist, DPN, resulted in a suppressing effect (LTD) or hardly any enhancing effect (spinogenesis), implying a less significant role for ER β in estradiol modulation of synaptic plasticity (Szymczak *et al.* 2006). Two different types of synaptic plasticity may therefore be modulated dominantly by ER α in glutamatergic excitatory neurons.

Single spine imaging was used to examine all the dendritic spines. Spine morphological investigations detected not only spine-synapses that form functional synapses but also, dendritic protrusions that do not form synapses. Double-staining investigations with Lucifer Yellow and synapsin I demonstrated that approximately 53% of spines were free spines without synapse formation. Electrophysiological

investigations of synaptic transmission were therefore used to examine, in particular, spine-synapses that form functional synapses. Because the estradiol-induced modulation of spine morphology and LTD appeared so rapidly, in the time range of 1–2 h, the synaptic ER α observed at the postsynaptic density (PSD) and postsynaptic compartment (Figs 3a and b, Fig. 5) probably plays an essential role in driving rapid processes.

Rapid effect of estradiol on synaptic transmission via synaptic ER α

Novel multi-electrode probes covering the CA1, CA3 and DG enabled simultaneous measurements of LTD in both pyramidal and granule neurons. Results demonstrated for the first time that both estradiol and the ER α agonists, PPT and DES, induced rapid modulation of synaptic transmission (Fig. 1). Previous electrophysiological studies have shown the rapid effect of estradiol only on the CA1 region of hippocampal slices (Foy *et al.* 1999), although other hippocampal subregions, CA3 and DG, have not been investigated in comparison with CA1.

Not only LTP but also LTD are necessary for complete memory processing (Migaud *et al.* 1998). In the current study, LTD was chosen to probe the modulation effects of estradiol because estradiol was much more effective on LTD than LTP. When the enhancement of LTP by estradiol was observed in CA1, an immediate increase of approximately 20% was induced upon the onset of estradiol perfusion in the baseline of the EPSP slope. A further approximate 130% increase was induced by high-frequency tetanic stimulation of Schaffer collaterals (Foy *et al.* 1999; Bi *et al.* 2000). However, without this approximate 20% baseline increase in EPSP slope (before the tetanic stimulation), the enhancement by estradiol was not apparent with regard to the pure tetanic stimulation-induced LTP (Ito *et al.* 1999; Kawato 2004; Mukai *et al.* 2006). In other words, the magnitude of tetanic stimulation-induced LTP was nearly the same between the presence and absence of estradiol. On the other hand, in the current LTD enhancement by estradiol, an immediate increase in EPSP by the onset of estradiol perfusion was not induced. The current results are in agreement with previous investigations of LTD. For example, estradiol treatment for 2 days *in vivo* facilitated the induction of LTD in ovariectomized rats (Desmond *et al.* 2000). *In vitro* perfusion of 0.1 nM estradiol (very low concentration) was not effective in modulation of LTD in young adult rat hippocampal slices (Vouimba *et al.* 2000), although 0.1 nM estradiol suppressed LTD in aged rats; this suggests that the response to estradiol may be age-dependent. At estradiol concentrations above 1 nM, we obtained enhancement of LTD (Fig. 1).

Important investigations have been performed by Moss and co-workers using knockdown mice (Gu and Moss 1996; Gu *et al.* 1999). They reported no essential

contribution of ER α to estradiol-induced rapid enhancement of the kainate currents of CA1 neurons. This conclusion was based on the very small difference observed in the estradiol effect on kainate currents between wild-type and ER α -Neo KO mice constructed by Neomycin insertion into exon 1 (the previously named exon 2) (Couse *et al.* 1995). It should be noted that in Neomycin insertion ER α -Neo KO mice (practically knockdown mice), N-terminal-modified ER α (61 kDa) is expressed (Couse *et al.* 1995; Kos *et al.* 2002; Pendaries *et al.* 2002). Because the N-terminal-modified ER α has been shown to remain active on estradiol binding and carries genomic processes (Kos *et al.* 2002; Pendaries *et al.* 2002), the participation of ER α in the electrophysiological properties of CA1 cannot be excluded from their investigations.

Woolley and co-workers (Woolley *et al.* 1997; Rudick and Woolley 2000, 2003) reported a series of investigations on the slow disinhibitory action of estradiol on the CA1 pyramidal neurons via interneurons by analyzing dendritic spines, electrophysiological properties and c-Fos responses 24–48 h after *in vivo* estradiol treatment of ovariectomized adult female rats. Increase of dendritic spine density by 22% after estradiol s.c. injection correlates well with an increase in glutamate binding to NMDA receptors and a change in electrophysiological properties (Woolley *et al.* 1997). They also found rapid (2 h) and slow (24 h) expression of c-Fos in CA1 pyramidal neurons induced by estradiol (Rudick and Woolley 2000, 2003). Their explanation is based on the observation that the immunoreactivity of AS409 and MC-20 is located primarily in interneurons (Hart *et al.* 2001; Milner *et al.* 2001).

Rapid and delayed effect of estradiol on spine density and morphology

Spine morphology was demonstrated for the first time to be significantly affected by short (0.5–2 h) estradiol treatments in adult male hippocampus (Fig. 2). The density of thin spines was dramatically increased, although the density of mushroom and stubby spines was not significantly altered. Such a selective modulation of thin spines is certainly a novel observation, and is essential for consideration of the neurogenesis and synaptic plasticity controlled by estrogens. The rapid effects of estradiol may be mediated by a signal transduction pathway via postsynaptic ER α (at the PSD), including MAP kinase but not Src kinase, as further judged from inhibitor analysis (Fig. 2b). Because both ER α and MAP kinase are present in the PSD fraction, an efficient coupling between these two proteins can occur at post-synapses (Fig. 3). The abolition of estradiol-induced spinogenesis, by blocking calcium-permeable NMDA receptors by MK-801 (Fig. 2b), was also observed in investigations of a slow effect (48 h) by *in vivo* estradiol treatment, suggesting the particular correlation of an estradiol signaling pathway with a weak spontaneous Ca²⁺ influx into spines via NMDA

receptors, even in the presence of Mg^{2+} blocking in ACSF (Woolley and McEwen 1994).

It should be noted that spine density is not always increased by the estradiol treatment. In CA3 pyramidal neurons, our recent observation demonstrated that the total density of thorns of thorny excrescences (spine-like post-synaptic structures in CA3), having contacts with mossy fiber terminals originated from granule cells, decreased dramatically to approximately 70% upon a 2 h application of 1 nM estradiol (Tsurugizawa *et al.* 2005). Therefore, the estradiol-induced spinogenesis is highly region-specific and heterogeneous.

There have been several *in vivo* studies on the effect of gonadal steroid supplement on the density of spines of pyramidal cells in the CA1 region. Estrogen supplement in ovariectomized female rats (Gould *et al.* 1990; Woolley *et al.* 1990; Woolley and McEwen 1992; Leranath *et al.* 2000, 2002) and androgens in castrated male rats (Leranath *et al.* 2003) increased the number of CA1 spines, resulting in recovery of spines to the level of the wild type. Morphological changes in spines have also been observed in ovariectomized female mice (Li *et al.* 2004). It should be noted that the mechanisms of estradiol action may be different between *in vivo* and *in vitro* experiments, because indirect effects of estradiol may also occur *in vivo* via cholinergic or serotonergic neurons, projecting to the hippocampus, in addition to the direct influence of estradiol on glutamatergic neurons. Leranath *et al.* (2000) demonstrated that s.c. injection of estradiol modulates the cholinergic or serotonergic neurons that project to the hippocampus from subcortical region, resulting in CA1 spinogenesis of ovariectomized female rats. In addition, ovariectomy, necessary to avoid the contribution of plasma estrogens, may have changed the condition of the cholinergic, serotonergic or glutamatergic neurons so that they were short of estrogens and therefore, estradiol supplement had a significant impact. For more discussion see Supplementary material.

Until the current investigations, acute slices of adult hippocampus had not been extensively investigated with regard to the rapid effect of estradiol on spinogenesis and spine morphology. We selected acute hippocampal slices, as the most intact *in vitro* adult neuron networks, for investigations of the rapid effect of estradiol free from the indirect effect of estradiol on glutamatergic neurons via serotonergic/cholinergic neurons. *In vitro* investigations have been carried out, using mainly neonatal hippocampal slice cultures, focusing on slow (1–4 days) effects of exogenous and endogenous estradiol (Pozzo-Miller *et al.* 1999; Bi *et al.* 2000; Rune *et al.* 2002; Prange-Kiel *et al.* 2006).

On the other hand, there have been a number of *in vivo* studies on the effect of gonadal estradiol supplement on the recovery of the density (not morphology) of spines of pyramidal cells in the CA1 region of ovariectomized female rats. In ovariectomized female rats, circulating estrogen is

undetectable and the spine density decreases significantly. The significant effect of gonadal estradiol on the recovery of the density of spines has been demonstrated to be dependent on subcortical efferents, because the surgical transection of fimbria/fornix, which contains subcortical efferents to the hippocampus, has prevented estradiol-induced spinogenesis. A direct comparison between *in vivo* studies and the current *in vitro* slice study is, in principle, not possible, because *in vivo* studies demonstrate the effect of gonadal estradiol on serotonergic or cholinergic afferents projecting to glutamatergic and GABAergic neurons in the hippocampus in addition to the effect of estradiol on hippocampal intrinsic neurons (Woolley *et al.* 1997; Rudick and Woolley 2000; 2003; Leranath *et al.* 2003; MacLusky *et al.* 2005).

Rapid action of hippocampal-derived estradiol

Hippocampus-derived estradiol should play a key role in the rapid action via synaptic $ER\alpha$ as an intracrine modulator of synaptic plasticity. In the current *in vitro* slice study, the exogenous application of estradiol (1–10 nM) was carried out to elevate rapidly the hippocampal estradiol level from the resting level. Exogenous application of estradiol was used because, until now, the method of rapid elevation of endogenous estradiol concentration via activation of its synthesis pathway had not been established. The concentration of endogenous estradiol is approximately 0.6 nM in the hippocampus in the resting state of neural activity (Hojo *et al.* 2004). In accordance with this, we observed a significant effect on LTD and spinogenesis only at >1 nM estradiol (Figs 1 and 2). For measurement in a more intrinsic condition, methods for the rapid activation of endogenous synthesis of estradiol should be developed, such as NMDA-induced estradiol synthesis (Kawato *et al.* 2002; Hojo *et al.* 2004), or 1–10 Hz low frequency electric stimulation.

The decrease in endogenous estradiol level also effectively modulates synaptic plasticity. The endogenous estradiol level can be decreased, by inhibiting P450arom activity with letrozole for more than 24 h, using hippocampal slice cultures and cell cultures. The suppression of endogenous estradiol synthesis decreased the spine density in CA1 after 4 days of letrozole treatment (Kretz *et al.* 2004). However, with the letrozole treatment, only the slow estradiol depletion effect can be measured (Prange-Kiel *et al.* 2003; Hojo *et al.* 2004). It is virtually impossible to measure the rapid effect on LTD or spinogenesis in the presence of letrozole because a significant decrease in endogenous estradiol by letrozole cannot be achieved within 1–2 h (time range of LTD and rapid spinogenesis experiments).

Previous uncertainty regarding the distribution of $ER\alpha$ immunoreactivity

The current study demonstrates the critical importance of using purified antibody RC-19 for the target protein 67 kDa

ER α in the hippocampus, where its expression level is extremely low (less than 1/100 relative to levels in the endocrine organs). The specificity of RC-19 was validated by careful western blot analysis, including ER α KO mice, in which no protein reacted with RC-19 in the hippocampus (Figs 3c and 4). Mass spectrometric analysis clearly demonstrated that RC-19 reacted with real ER α (Fig. 6). As judged from the presence of only a single ER α mRNA species, and the observation of the single 67 kDa protein band, the synaptic ER α is concluded to be identical to the classical nuclear ER α observed in the nuclei of hippocampal neurons. The mechanisms of stable ER α binding to the PSD layer (Fig. 3b) should be investigated.

Staining with RC-19, in turn, indicates that antiserum MC-20, after purification, could be an appropriate antibody for ER α in the hippocampus. MC-20, used without purification, significantly reacted with unknown proteins in the hippocampus of ER α KO mice. MC-20 antiserum reacted with the 62 kDa protein in the hippocampus, hypothalamus and cerebral cortex not only from wild-type rats but also from ER α KO mice (Fig. 3d). However, in the ovary, MC-20 did not react with the 62 kDa protein and only reacted with the 67 kDa ER α (Fig. 3dii), suggesting that the 62 kDa protein may be selectively expressed in the brain. These results are consistent with those of Toran-Allerand and co-workers (Toran-Allerand *et al.* 2002), who reported the reaction of MC-20 with an approximately 62 kDa protein in the neocortex of both wild-type and ER α KO mice (Toran-Allerand *et al.* 2002). The 62 kDa band stained by MC-20 in the rat hippocampus cannot be a splice variant of ER α , having the same C-terminal sequence as 67 kDa ER α because MC-20 also stained the 62 kDa band in ER α KO mice (exon 2 deleted type) (Fig. 3d) in which no splice variant of ER α mRNA was expressed (Dupont *et al.* 2000; Pendaries *et al.* 2002). Our RT-PCR analysis, which indicated that the only possible splice variant was 46 kDa and not 62 kDa (see Fig. 7), also supports this conclusion. The assertion that there is a relatively small difference between the 62 kDa and 67 kDa MC-20-reacted bands is supported by the coincidence with the 67 kDa band of genetically-expressed ER α , and the 62 kDa of the hypothalamus in which these two bands always appear in parallel (Fig. 3dii).

The present study indicates that the synaptic plasticity of principal neurons is modulated by ER α expressed in the entire principal neurons, and that the hypothesis of interneuron-mediated estradiol modulation is probably not a main mechanism of the synaptic plasticity of principal neurons. The current ER α staining is a goal of previous attempts of ER α staining that demonstrated overall or partial staining of nuclei and cytoplasm of principal neurons of hippocampal slices using *in situ* hybridization, MC-20 or C1355 antiserum (Cardona-Gomez *et al.* 2000; Solum and Handa 2001; Rune *et al.* 2002).

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Supplementary Material

The following material is available online.

Appendix S1

Figure S1 The arrangement of stimulation electrodes (black circle) and recording electrodes (white circle) on a custom multielectrode probe (dotted line represent pyramidal and granule cell layers).

Figure S2 Schematic illustration for diagram of the spine analysis.

Figure S3 (A) Blot with RC-19 antibody of the nuclear and synaptosomal fractions from adult rat brain. From left to right: homogenates of ovary (OV), nuclear fractions of cerebral cortex (nCC), hippocampus (nHC), hypothalamus (nHT), cerebellum (nCL), synaptosomal fractions of cerebral cortex (sCC), hippocampus (sHC), hypothalamus (sHT), cerebellum (sCL), and ER α expressed in COS-7 cells (ER). (B) Staining with AS409 antiserum of tissue homogenates from adult rat. From left to right, OV from 4 week rat, HC, and HT. Representative blots are shown, but several independent experiments are performed for each blot. Molecular weights (97, 66, 45, 38 kDa) indicated in the figure are taken from molecular weight standard proteins.

Figure S4 Immunohistochemical staining with MC-20 antiserum in the hippocampal slices of adult male rat (A-D). (A) The coronal section of the whole hippocampal formation. (B) CA1 region. (C) dentate gyrus. (D) CA3 region. so, stratum oriens; pcl, pyramidal cell layer; sr, stratum radiatum; gcl, granule cell layer; hl, hilus. Scale bar, 500 μ m for (A), and 200 μ m for (B)-(D).

Figure S5 Immunoelectron microscopic analysis, using RC-19 antibody, of the distribution of ER α within synapse of stratum radiatum of CA3 neuron (A), nucleus of CA3 neuron (B), and nucleus of DG (C) regions. A 1:1000 dilution of RC-19 was used to prevent nonspecific labeling. pre, presynaptic region; post, postsynaptic region. Scale bar, 200 nm. (D) Immunoelectron microscopic analysis, using MC-20 antiserum, of the distribution of antigens of MC-20 in the presynaptic regions, in the stratum radiatum of the hippocampal CA3 region. Heavy staining was observed and several gold particles (arrowheads) were associated with the mitochondria (mito). Scale bar, 200 nm.

Figure S6 The distribution of the following subregions of ER α mRNA amplified with specific primer pairs: Exon1'-2-3-4', Exon3'-4-5', Exon4'-5-6', Exon5'-6-7', and Exon6'-7-8'.

Figure S7 (A) RT-PCR analysis of mRNAs for ER α expressed in the brain subregions. RT-PCR products for mRNAs (50 ng each) of ER α were electrophoresed on 1.5% agarose gel, and visualized with ethidium bromide. (Top) From left to right, size marker (Marker), ovary diluted at 1/50 (1/50 OV), ovary diluted at 1/500 (1/500 OV), cerebral cortex (CC), hippocampus (HC), hypothalamus (HT), cerebellum (CL), no cDNAs (No cDNA) as a negative control. (Middle) Southern hybridizations. (Bottom) The ethidium bromide

staining of GAPDH. The PCR cycles are 34 cycles. (B) RT-PCR analysis of mRNAs for ER β expressed in the brain subregions. RT-PCR products for mRNAs (50 ng each) of ER β were electrophoresed on 1.5% agarose gel, and visualized with ethidium bromide. (Top) From left to right, size marker (Marker), cerebral cortex (CC), hippocampus (HC), thalamus (TL), hypothalamus (HT), cerebellum (CL), no cDNAs (No cDNA) as a negative control, and ovary (OV). (Middle) Southern hybridizations. (Bottom) The ethidium bromide staining of GAPDH. The PCR cycles are 36.

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