PKC epsilon Promotes Synaptogenesis through Membrane Accumulation of the Postsynaptic Density Protein PSD-95


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Running Title: PKCε induces synaptogenesis via PSD-95

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ABSTRACT

Protein kinase C epsilon (PKCε) promotes synaptic maturation and synaptogenesis via activation of synaptic growth factors such as BDNF, NGF, and IGF. However, many of the detailed mechanisms by which PKCε induces synaptogenesis are not fully understood. Accumulation of PSD-95 to the postsynaptic density (PSD) is known to lead to synaptic maturation and strengthening of excitatory synapses. Here we investigated the relationship between PKCε and PSD-95. We show that the PKCε activators DCPLA-ME and bryostatin 1 induce phosphorylation of PSD-95 at the serine-295 residue, increase the levels of PSD-95, and enhance its membrane localization. Elimination of the serine-295 residue in PSD-95 abolished PKCε-induced membrane accumulation. Knockdown of either PKCε or JNK1 prevented PKCε activator-mediated membrane accumulation of PSD-95. PKCε directly phosphorylated PSD-95 and JNK1 in vitro. Inhibiting PKCε, JNK, or CaMKII activity prevented the effects of PKCε activators on PSD-95 phosphorylation. Increase in membrane accumulation of PKCε and phosphorylated PSD-95 (p-PSD-95S295) coincided with increased number of synapses and increased amplitudes of excitatory post-synaptic potentials (EPSPs) in adult rat hippocampal slices. Knockdown of PKCε also reduced the synthesis of PSD-95 and the presynaptic protein synaptophysin by 30% and 44% respectively. Prolonged activation of PKCε increased synapse number by 2-fold, increased presynaptic vesicle density, and greatly increased PSD-95 clustering. These results indicate that PKCε promotes synaptogenesis by activating PSD-95 phosphorylation directly, through JNK1 and CaMKII, and also by inducing expression of PSD-95 and synaptophysin.

Protein kinase C epsilon (PKCε) is one of the novel PKC isotypes and is characterized as a calcium independent and phorbol ester/diacylglycerol–sensitive serine/threonine kinase. Among the novel PKCs, PKCε is the most abundant species in the central nervous system, mediating various neuronal functions (1-2). In neuroblastoma cells overexpression of PKCε, but not PKCα, βII or δ leads to neurite outgrowth through interaction of actin filaments and the C1 domain of PKCε (3-5). The actin binding site of PKCε is also implicated in exocytosis of neurotransmitters (6). PKCε is essential for many types of learning and memory (7-8) and neuroprotection (9-13). Neuronal contact with astrocytes also promotes global synaptogenesis through PKCε signaling (14). PKCε activation has been shown to promote the maturation of dendritic synapses during associative learning (9). PKCε activation also protects against neurodegeneration (10,15). Phosphorylation of long-tailed AMPARs GluA4 and GluA1 by PKC promotes their surface expression (16-17). PKCε activation induces protein synthesis required for long-term memory (12,18). PKCε activation is also required for HuD-mediated mRNA stabilization of neurotrophic factors (19) and ApoE mediated epigenetic
regulation of BDNF (20). PKC activation induces translocation of calcium/calmodulin-dependent kinase II (CaMKII) to synapses (21) where it participates in PSD-95-induced synaptic strengthening (22). PKC also promotes NMDA receptor trafficking by indirectly triggering CaMKII autophosphorylation and subsequent increased association with NMDARs (23).

Thus, a number of studies have suggested that PKC activators such as bryostatin and dicyclopropanated linoleic acid methyl ester (DCPLA-ME) may be useful therapeutic candidates for the treatment of Alzheimer's disease (AD) and other causes of synaptic loss such as ischemia, stroke, and Fragile X syndrome (5-6,14,24). Some of these benefits have been attributed to induction of neurotrophic factors such as BDNF or the activation of anti-apoptotic activity (10,13,20,25). However, the biochemical mechanisms by which PKCε induces synaptogenesis and mediates neuroprotection are still not fully understood.

At excitatory synapses, the postsynaptic density is characterized by an electron-dense thick matrix that contains key molecules involved in the regulation of glutamate receptor targeting and trafficking (26). PSD-95 is an abundant scaffold protein in excitatory synapses, where it functions to cluster proteins such as glutamate receptors on the postsynaptic membrane and couples them to downstream signaling molecules, thereby inducing the surface expression and synaptic insertion of glutamate receptors (27-29). In addition to its role in synaptic function, PSD-95 has also been proposed to affect synapse maturation and stabilization (30-32) and thus, synapse number. Phosphorylation of the serine-295 residue of PSD-95 enhances the synaptic accumulation of PSD-95 and its ability to recruit surface α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA) receptors and potentiate excitatory postsynaptic currents (33).

In the present study, we examined the role of PKCε signaling and PKCε activators in PSD-95 regulation and induction of synaptogenesis in cultured neurons and CA1 hippocampal slices. We report that PKCε activation induces membrane translocation and phosphorylation of PSD-95 at the serine-295 residue, coinciding with an increased number of synapses. Our data suggest that an important mechanism by which PKCε induces synaptogenesis is by increasing the phosphorylation of PSD-95 at the postsynaptic site and by regulating the expression of synaptophysin at the presynaptic site.

RESULTS
PKCε activation prevents degradation of primary human neurons- PKCε is present in high concentration in central neuronal tissues and has been implicated in broad spectrum neuronal functions. To determine the effect of PKCε activation on survival and maintenance, primary human neurons were treated for 40 days with two different PKC activators (bryostatin 1 and DCPLA-ME, which are relatively specific for PKCε) (13,34-36). Culture media and activators were changed every three days. Cells were imaged from three independent wells every five days and neurite positive cells were counted from 508μm² field images. Cells treated with either DCPLA-ME (100nM) or bryostatin 1 (0.27nM) showed an improved survival with increased neuritic branching (Fig. 1A). Untreated cells showed degeneration and 50% cell loss by 36 days, while the treated cells remained healthy for at least 40 days (Fig. 1B). The number of viable neurite- positive cells was also significantly higher at 40 days (F (2, 6)=705.4; ANOVA, P<0.0001) in the activator treated cells than untreated cells (bryostatin 1: 369.7 ± 12.2; DCPLA-ME: 334.7 ± 1.8; untreated 109.7 ± 6.4).

Prolonged PKCε activation prevents loss of synaptic proteins- We quantified the mRNA expression of PKCε, PSD-95 and synaptophysin at 40 days in untreated and PKCε activator-treated neurons. At 40 days the mRNA levels of PKCε (F(3,8) = 18.3; P=0.0006) and PSD-95 (F(3,8) = 44.6; P<0.0001) were significantly higher in the PKCε activator-treated cells than untreated cells (bryostatin 1: 369.7 ± 12.2; DCPLA-ME: 334.7 ± 1.8; untreated 109.7 ± 6.4).

Synaptophysin mRNA showed no significant change in between treated and untreated groups (Fig. 1E). We also quantified the protein expression of phosphorylated PSD-95 (p-PSD-95S295), PSD-95 and synaptophysin at 40 days in untreated and PKCε activator-treated neurons by immunoblot (Fig. 1F). Expression levels of PKCε (F(3,8) = 16.60; P<0.001), p-PSD-95S295 (F(3,8) = 66.83; P<0.0001), PSD-95 (F(3,8) = 21.22; P<0.001) and synaptophysin were significantly higher in the 40 day PKCε activator-treated cells compared to 40 day control cells (Fig.

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1G-J). Moreover, protein expression levels of PKCε, PSD-95 and synaptophysin showed a marked decrease in 40-day untreated cells compared to 1-day cells, even after correction for total protein, while PKCε activation prevented the time-dependent loss. This indicates an essential role of PKCε in maintenance of synapses and preserving normal levels of both PSD-95 and synaptophysin.

**Bryostatin 1 and DCPLA-ME specifically activate PKCε.** We then investigated whether this phenomenon is specific to PKCε or whether other PKC isoforms are involved. PKC translocation to the plasma membrane generally has been considered the hallmark of activation and frequently has been used as a surrogate measure of PKC isoform activation in cells (37). Expression levels of PKCα, PKCε and PKCδ in the soluble (cytosol) and particulate (membrane) were measured by immunoblot at 1hr, 4hr and 24hr after either bryostatin 1 (0.27 nM) or DCPLA-ME (100 nM) treatment (Fig. 2A, C). Both DCPLA-ME and bryostatin 1 increased membrane translocation of PKCε but not PKCα or PKCδ (Fig. 2B,D), confirming that both the compounds activate PKCε but not PKCα or PKCδ.

**PKCε activation induces membrane translocation of phosphorylated PSD-95 (serine 295).** Phosphorylation of PSD-95 on Serine-295 is known to promote localization of PSD-95 in the postsynaptic density (PSD), strengthening the excitatory synapse (33). To determine whether time-dependent PKCε activation has an effect on localization and expression of p-PSD-95S295, we measured the expression of p-PSD-95S295 in the soluble and particulate fractions of the primary human neurons at 1hr, 4hr and 24hr post PKC activator treatment (Fig. 2E, F). PKCε activation increased the level of p-PSD-95S295 in the particulate fraction of both bryostatin 1 (F (3,8)=4.9; ANOVA, P=0.03) and DCPLA-ME treated cells (F (3,8)=11.7; ANOVA, P=0.003) (Fig. 2F). The total PSD-95 expression in whole cell lysate from primary human neurons was unchanged among different groups (Fig. 2E). At 4hr p-PSD-95S295 levels were significantly higher in bryostatin 1 (156.4 ± 14.9 %; P=0.01) and DCPLA-ME (160.1 ± 9.5 %; P=0.003) treated neurons compared to untreated neurons. In adult rat hippocampal slices PKCε activation increased p-PSD-95S295 expression at 1hr and 4hr (bryostatin 1: F (3,8) =4.95; ANOVA P=0.031 and DCPLA-ME: F (3,8) =4.34; ANOVA P=0.043). (Fig. 2G, H). Negligible amounts of p-PSD-95S295 were detected in the soluble fraction. These results show that the increase in membrane localization of p-PSD-95S295 corresponded with the kinetics of PKCε activation at 1hr and 4hr.

**PKCε-mediated phosphorylation of PSD-95 at serine-295 is essential for its membrane association.** Purified recombinant human PSD-95 (r-PSD-95) was readily phosphorylated by activated recombinant PKCε (r-PKCε) in vitro, and the PKC inhibitor bisindolylmaleimide I (Go 6850) (BisI: 100nM) blocked the reaction (Fig. 3A). Both bryostatin 1 and DCPLA-ME increased the amount of p-PSD-95S295 in vitro compared to unactivated PKC alone (Bryostatin 1 + r-PKCε + r-PSD-95: 200.3 ± 5.06 %, P=0.004; DCPLA-ME+ r-PKCε + r-PSD-95: 194.6 ± 12.95 %, P=0.032; r-PKCε + r-PSD-95 control: 146.9 ± 7.06 %) (Fig. 3B). The PKC inhibitor BisI blocked the phosphorylation. These results show that PKCε can phosphorylate PSD-95 at serine 295 in vitro.

Next we tested if the serine-295 residue in PSD-95 is essential for its membrane translocation. We created two separate clones, one containing the wild type human PSD-95, and the other containing mutant-PSD-95 (S295K), in which the serine residue at 295 (AGT) was changed to lysine (AAA). Both the clones were transfected and expressed in HEK-293 cells and their expression was measured by immunoblot. Both transfected cell lines showed PSD-95 immunoreactivity against a PSD-95 antibody raised against the N-terminal region of PSD-95. The anti-p-PSD-95S295 antibody showed positive bands only with the wild type PSD-95 transfected cell lysate. Untransfected HEK-293 cells showed no PSD-95 expression (Fig. 3C). The wild PSD-95 and PSD-95S295K expressing HEK-293 cells were then treated with bryostatin 1 and DCPLA-ME for 4hr in presence or absence of PKCε translocation inhibitor (EAVSLKPT; 5µM) and fractionated into cytosol and membrane fractions. Only small amounts of p-PSD-95 were found in soluble fractions (Fig. 3D). As expected, bryostatin 1 and DCPLA-ME significantly increased membrane translocation of PKCε in both PSD-95 and PSD-95S295K expressing cells (in wild-type cells: bryostatin 1: +54.4 ± 4.9 %, P=0.0004 and DCPLA-ME = +19.1 ± 4.2 %, P=0.01; F (4,10)=28.8, ANOVA P<0.0001) (Fig.
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PKCε activators also increased translocation of wild-type PSD-95 (bryostatin 1: + 29.9 ± 2.3%, P=0.0002; DCPLA-ME: + 20.5 ± 2.5 %, P=0.001; F(4,10)= 35, ANOVA P<0.0001), but not mutated PSD-95S295K, which lacks the PKCε phosphorylation site (F (2,6)= 0.75, ANOVA P=0.51) (Fig. 3F). To further verify that phosphorylation at serine-295 is essential for membrane association, we immunoblotted the membrane fraction against anti-p-PSD-95S295 antibody. The level of p-PSD-95S295 in the membrane was increased in the PKCε activator treated cells (bryostatin 1: 151.7 ± 6.1 %, P=0.002 and DCPLA-ME: 161.1 ± 10.4 %, P=0.004 (100 ± 2.8 %); F(4,10)=27.4, ANOVA P<0.0001) (Fig. 3G). Membrane p-PSD-95S295 translocation was blocked by the PKC inhibitor. Together, these results indicate that PKCε activation phosphorylates the serine-295 residue of PSD-95 and this phosphorylation is necessary for membrane accumulation of PSD-95.

PKCε-mediated membrane localization of p-PSD-95S295 involves JNK I and CaM-KII: Previously it has been reported that accumulation of PSD-95 in the PSD is increased by synaptic activity and by a Rac1-JNK I signaling pathway (33). PKCε is involved in JNK activation in macrophages (38-39) and CaM-KII inhibitors inhibit PKC-mediated signaling in hippocampal neurons (40). Thus we investigated the involvement of PKCε, JNK and CaM-KII in PSD-95S295 translocation in primary human neurons. Cells were pretreated for 30 min with BisI (100 nM, PKC inhibitor), SP600125 (20μM, JNK inhibitor) or KN-93 (10μM, CaM-KII inhibitor) and then treated with PKCε activators for 4hr. Cells were fractioned into cytosolic and membrane fractions and the membrane fractions were analyzed for the expression of p-PSD-95S295. The inhibitors alone reduced the expression of membrane bound p-PSD-95S295 (F (4,10) = 23.04; ANOVA P<0.0001) (Fig. 4A,C). DCPLA-ME treatment increased membrane localization of p-PSD-95S295 (147.3 ± 2.8 %; F (5,12)=39.2; ANOVA P<0.0001). PSD-95 phosphorylation was prevented by blocking PKC activation using bisindolylmaleimide I (Fig. 4B,D), confirming the involvement of PKCε in localization of p-PSD-95S295 in the membrane. The JNK inhibitor SP600125 and the CaM-KII inhibitor K-93 also prevented PKCε-mediated phosphorylation and translocation of PSD-95 (Fig. 4B,D).

PKCε phosphorylated PSD-95 in vitro, incorporating 1.46 ± 0.05 moles of 32P-ATP per mole of PSD-95. Western blotting with p-PSD-95S295-specific antibody confirmed that this included the Ser-295 site (Fig. 3A,B). PKC and JNK inhibitors fully inhibited the PKCε-mediated PSD-95 phosphorylation, while a CaM-KII inhibitor partially prevented PSD-95 phosphorylation (Fig. 4E). PKCε also phosphorylated JNK1 in vitro, incorporating 1.02 ± 0.04 moles of 32P-ATP per mole of JNK1; BisI prevented JNK1 phosphorylation (Fig.4F). PKC is also reported to phosphorylate CaM-KII in vitro (41), we also found increase in phosphorylation of CaM-KII by PKCε (Fig. 4G). Since both JNK and CaM-KII inhibitors prevented PSD-95 phosphorylation by PKCε (Fig. 4E), we considered the possibility that the JNK inhibitor might not be specific. Therefore we performed a siRNA knockdown of PKCε and JNK in human neurons. PKCε or JNK knockdown caused a 50% reduction in their respective protein expression (Fig. 4H). DCPLA-ME failed to induce the membrane accumulation of p-PSD-95S295 in PKCε and JNK knockdown human neurons (F (5,12) =24.6; ANOVA P<0.0001(Fig. 4I,J). These results confirm that PKCε is required for membrane translocation of p-PSD-95S295 and that JNK and CaM-KII are intermediates in the pathway (Fig.4K).

PKCε activation induces synaptogenesis in adult hippocampal slices: Next we investigated if PKCε mediated phosphorylation of PSD-95 at serine-295 leads to synaptogenesis. Since 4hr PKCε activator treatment produced the highest p-PSD-95S295 level, we quantified the number of synapses from within 100 μm2 of 30–35 CA1 regions from untreated, bryostatin 1 and DCPLA-ME treated slices using electron microscopy (3 different slices in each group) (Fig. 5A). Both bryostatin 1 and DCPLA-ME increased the number of synapses at 4hr compared to only vehicle treated control (8.97 ± 0.63, P=0.002, n=35). 6.97 ± 0.50, P=0.04; n=30, and 5.77 ± 0.50; n=35 CA1 areas, respectively) (Fig. 5B). Presynaptic vesicle density was measured in a series of 3D stacked images from 6–10 presynaptic boutons from 3 different hippocampal slices. Bryostatin 1 treatment increased
presynaptic vesicle density at 4hr (93.23 ± 4.1, P<0.001, n=30 presynaptic boutons) in comparison to control (71.33 ± 4.45, n=22 presynaptic boutons).

Next we investigated the effect of bryostatin 1 on basal synaptic transmission of hippocampal CA1 pyramidal neurons to determine whether the new synapses are functional. Field potential recordings were measured from rat hippocampal slices. An input-output curve was calculated with stimulus intensity versus the slope of EPSPs elicited in response to increasing intensity of stimulation to the Schaffer collateral. The mean EPSP slope increased with stronger intensity of stimulus. Slices preincubated with bryostatin 1 for 1hr exhibited greater EPSP slope without any change in fiber volley amplitude. This was abolished with 30 min pretreatment with the PKC inhibitor bisindolylmaleimide I (Go 6850) (BiSt: 100nM) (Fig. 5C, D). Bryostatin 1 increased the area under the curve, which represents the overall basal synaptic transmission, and a PKC inhibitor prevented the increase (bryostatin1: 0.71±0.08, P=0.03; BiSt+bryostatin 1: 0.49 ± 0.07 and untreated (ethanol only): 0.51 ± 0.06) (Fig. 5E). Treatment of slices for 4hr with bryostatin (12 slices, 3 rats) dramatically increased the EPSP slope compared to the ethanol-treated slices (6 slices, 3 rats) (Fig. 5F,G). The smaller response in the 4hr untreated slices compared to 1hr untreated slices may be attributed to the vehicle (ethanol) added to the slices. EPSPs in hippocampal slices are reduced by a smaller percentage after ethanol treatment (42). Thus, the prolonged treatment of slices with ethanol for 4hr may have slightly reduced the EPSP slope in these groups. Bryostatin increased the area under the curve by nearly 2 fold (P<0.0001, Fig. 5H). These results suggest that bryostatin 1 treatment facilitates basal synaptic transmission in the Schaffer collateral commissural pathway of rat hippocampus and that the increase in EPSP slope is independent of the fiber volley.

Our results indicate that increased phosphorylation of PSD-95 by PKCε leads to an increase in synapse number with increased synaptic activity. Together these data demonstrate that the new synapses are functional.

**PKCε** knockdown reduces the expression of PSD-95 and synaptophysin - PKCε is known to perform important functions both in presynaptic (14) and postsynaptic sites. To investigate whether PKCε is essential for the expression of synaptic proteins, we measured the effect of PKCε knockdown (PKCε KD) and PKCε overexpression (PKCε OE) on the expression of postsynaptic PSD-95 and presynaptic synaptophysin. Knockdown of PKCε was achieved by transfecting the neurons with a mixture of siRNA containing a pool of three to five siRNAs. PKCε siRNA effectively reduced PKCε expression both at the mRNA and protein levels by 2- and 3.4-fold (Fig. 6A, I) after 72 hr. Scrambled siRNA did not cause any change in PKCε expression (Fig. 6F,G). PKCε overexpression in the neurons was obtained by transfecting pCMV6-ENTRY vector containing human PKCε cDNA. Transfected neurons showed a ~7.4 fold increase in PKCε mRNA level (Fig.6A) and 3.6 fold increase in PKCε protein level (Fig. 6H, I). Overexpressing PKCε by 7-fold increased the level of synaptophysin mRNA by 59.3 ± 1.3 % and also increased the level of PSD-95 by 71.6 ± 3.8 %. Knockdown of PKCε had opposite effects (Fig. 6B, C). PKCε overexpression or knockdown did not alter SNAP-25 and syntaxin-1 mRNA levels (Fig. 6D, E). Loss of PKCε expression reduced the protein levels of PSD-95 by 30% (0.71 ± 0.07; P=0.043 (Fig. 6K) and synaptophysin by 44% (0.56 ± 0.08; P=0.021) (Fig. 6J) compared to controls transfected with scrambled siRNA. PKCε OE produced a 50% increase in synaptophysin (1.51 ± 0.1 vs 1.0 ± 0.1 in control; P=0.015) (Fig. 6J) and a 30% increase in PSD-95 expression (1.31 ± 0.08 vs 1.0 ± 0.07 in control; P=0.024) compared to vector-only transfected cells (Fig. 6K).

**Knockdown of PKCε reduces synaptogenesis-** To further establish the role of PKCε in synaptogenesis and its underlying role in expression of PSD-95 and synaptophysin we used confocal microscopy to measure the effect of PKCε knockdown on the localization of PSD-95 and colocalization of PSD-95 and synaptophysin. Punctate colocalization (clusters of proximal pre- and post-synaptic markers on neurites) of PSD-95 and synaptophysin is widely accepted as an indicator of synapses (43-44). Primary human neurons were treated with bryostatin 1or DCPLAMA alone or after PKCε KD for 10 days. PSD-95 clusters and colocalized PSD-95 and synaptophysin (as recognized by staining grains along a 40μm length of neurite, n=10) were...
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counted in 4 independent experiments (Fig. 7A). In normal cells, PKCε activation by bryostatin 1 and DCPLA-ME significantly induced PSD-95 clustering in the neurites compared to untreated controls (P<0.05) (Fig. 7B). The number of synapses was also significantly higher in cells treated with bryostatin1 and DCPLA-ME than in untreated neurons at 10 days (Fig. 7C). The increase in number of synapses was independent of the neuron density. We found no change in neuron density (measured by NeuN staining) after 10 days of PKCε activator treatment (Supplementary Fig.1). In PKCε KD cells, immunofluorescence staining of human neurons showed a loss of synaptic networks, and bryostatin 1 and DCPLA-ME had no effect. PKCε KD prevented the effect of PKCε activators and, more importantly, reduced the basal level of PSD-95 clusters and synapses by 50%. (Fig. 7A-C)

We also quantified the expression levels of PKCε, p-PSD-95S295, PSD-95 and synaptophysin by immunoblot after 10 days of PKCε-siRNA transfection (Fig. 7D). PKCε KD cells expressed significantly lower amounts of PSD-95 (F (5, 12) = 19.24, ANOVA P<0.0001) (Fig. 7E, F), and synaptophysin (F (5, 12) = 12.79, ANOVA P=0.0002) (Fig. 7G). Bryostatin 1 and DCPLA-ME failed to induce PSD-95 and synaptophysin expression in PKCε KD neurons. Bryostatin 1, but not DCPLA-ME, produced a 40% decrease in PKCε protein staining (Fig. 7D). No loss in PKCε mRNA was found in bryostatin 1-treated neurons (data not shown). Downregulation of PKC following activation by bryostatin 1 is a well documented phenomenon (45-46).

We further confirmed the effect of long term PKCε activation on synaptogenesis using rat hippocampal brain slices. Slices were treated with bryostatin 1 and DCPLA-ME for 10 days. The serum-free culture medium was changed every 3 days with fresh addition of activators. Synapse number in each case was quantified using electron microscopy (Fig. 8A). Bryostatin 1 (7.97 ± 0.68; P=0.013, n=29 CA1 areas) and DCPLA-ME (8.71 ± 0.78; P=0.001, n=24 CA1 areas) treatment increased the number of synapses in hippocampal slices compared to vehicle-only treated slices (4.5 ± 0.45; n=24 CA1 area) (Fig. 8B). Presynaptic vesicle density was also significantly higher in the bryostatin 1 (59.6 ± 6.4, P<0.05, n=19 presynaptic boutons) and DCPLA-ME (60.4 ± 5.1, P=0.04, n=19 presynaptic boutons) treated slices than vehicle treated controls (48.4 ± 4.3, n=20 presynaptic boutons) (Fig. 8C, D). Together, these findings confirm that PKCε is essential for bryostatin 1 and DCPLA-ME-mediated increase in PSD-95 and synaptophysin expression leading to increased synaptogenesis at 10 days.

DISCUSSION

The outgrowth of neurites and formation of synapses depend on interactions among a number of regulatory proteins. These interactions are required for synaptic structure rearrangement, spinoogenesis, and synaptogenesis. PKCε is one of the key regulators of synaptogenesis (3,24) and PKCε activators promote the maturation of dendritic spines (9,47). PSD-95 is a scaffold protein which also plays an important role in formation of excitatory synapses (48-49).

Here we showed that PKCε activation induces translocation and phosphorylation of PSD-95 at serine-295 residue leading to PSD-95 accumulation at the postsynaptic density. Our findings showed that PKCε activation not only increased the survival of neurons but also preserved the neuronal structure. Untreated cells showed gradual degeneration over 25 days, suggesting that PKCε activation is beneficial for both maturation and survival of neurons, confirming a previous report by Hama et al.(14). We have shown that short term acute changes in PKCε activity induce structural and biochemical changes in post-synaptic density scaffolding protein PSD-95 as well as increased synaptic activity. Synaptic activity is important for neuronal survival. Synaptic activity induces expression of survival genes and suppresses pro-death genes (50). Therefore, the increased survival of neurons treated with PKCε activators may be due to the increased connectivity induced in the early stages; however, other factors such as elevated neurotrophins may also play a role. PKCε induces BDNF (10,19) and elevated expression and release of BDNF is associated with elevated synaptic activity, which contributes to neuroprotection (51-52).

PKCε activation and membrane translocation occur both presynaptically (14,53) and postsynaptically (8) where it phosphorylates important substrate proteins required for synaptic
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facilitation and synaptogenesis. We found that p-PSD-95S295 accumulation increased in the membrane of PKCε-activated neurons and followed the same time course as PKCε activation at 1hr and 4hr. The serine-295 residue was essential for the PKCε mediated membrane accumulation of PSD-95. In vitro, PKCε phosphorylated both PSD-95 and JNK1. The JNK1 inhibitor also prevented PKCε activation-mediated increase in p-PSD-95S295, confirming previous findings showing that serine-295 phosphorylation of PSD-95 is regulated by Rac1-JNK1 and PP1/PP2A signaling (33,54). PKCε is involved in JNK activation; PKD, a downstream effector of PKC, also regulates JNK (38-39,55). Knockdown of either PKCε or JNK inhibited the PKCε activator mediated p-PSD-95S295 accumulation in the membrane, thus confirming that PKCε and JNK act collectively in regulating PSD-95. Although it has been reported that synaptic localization of PSD-95 is regulated by JNK signaling and not by CaMKII (33,56), our data demonstrate a role of both JNK1 and CaMKII. This is possible as PKC activation induces a simultaneous translocation of calcium/calmodulin-dependent kinase II (CaMKII) to synapses (21) and CaMKII activation is needed for PSD-95-induced synaptic strengthening (22). CaMKII is a downstream target of PKCε in many pathways, including the events responsible for the induction of neuroplastic changes associated with hyperalgesic priming (57). In this study, we found that both JNK1 and CaMKII inhibitors prevented the PKCε mediated membrane association of p-PSD-95S295. These results suggest that JNK1 and CaMKII are downstream to PKCε in events responsible for phosphorylation and membrane accumulation of PSD-95.

We also demonstrated that PKCε activation increases the levels of PSD-95 and the number of synapses. In adult hippocampal slices, bryostatin 1 increased basal synaptic activity. Our results indicate an important link between PKCε activation and the membrane localization of PSD-95, specifically enriching the membrane with the p-PSD-95S295 form, which is known to strengthen the excitatory synapses (33). PSD-95 also regulates membrane insertion of AMPA receptor and dendritic spine morphology during synaptic plasticity (22,30-32).

Overexpression of PSD-95 converts silent synapses to functional synapses (58), while synaptophysin may be required for increased presynaptic vesicle density, thereby facilitating neurotransmitter release (59). We found that overexpressing PKCε in primary human neurons induces the mRNA and protein levels of PSD-95 and synaptophysin, while knockdown of PKCε reduces PSD-95 and synaptophysin mRNA and protein levels. Our results indicate that PKCε regulates the gene expression of PSD-95 and synaptophysin. PKCε may play a critical role synapse maintenance by regulating the synthesis of PSD-95 and synaptophysin (18). PKCε is known to drive the mitogenic response and DNA synthesis (60) via the Raf-MEK-ERK cascade and regulates transcription of essential genes through JNK/AP1, NF-κB and JAK/STAT cascades (61-62). PSD-95 is a critical transcriptional target of NF-κB, which is known to induce excitatory synapse formation and regulate dendritic spine formation and morphology in murine hippocampal neurons (63). Synaptophysin mRNA expression is induced by the BDNF-cFos pathway (64). NF-κB and synaptophysin have a common regulator in BDNF (65). PKCε upregulates BDNF expression (19-20,66).

In conclusion, PKCε has two specific roles in synaptogenesis: at the postsynaptic site it regulates PSD-95, either directly or through JNK1 and CaMKII, and at the presynaptic site it induces the expression of synaptophysin. Repeated treatment with PKCε activators induces synthesis of PKCε, PSD-95 and synaptophysin, resulting in an increased number of synapses. PKCε knockdown inhibits the synthesis of PSD-95 and synaptophysin leading to a reduced number of synapses. Besides the PKC-JNK1/CaMKII-PSD-95 pathway, PKCε can also induce synaptogenesis through the HuD-BDNF pathway. PKCε stabilizes HuD, which increases the stability and rate of translocation of target mRNAs. HuD increases as a result of PKCε activation after learning (67) and stabilizes the mRNA for BDNF, nerve growth factor (NGF), and neurotrophin-3 (NT-3) (19). PKCε activation induces the synthesis of BDNF (10,20,47) and BDNF induces transport of PSD-95 to the dendrites (68), which is required for maintenance of mature spines (69). Deficits of PKCε function could also contribute to the synapse loss in AD (15), while the therapeutic
PKCε induces synaptogenesis via PSD-95 elimination of such deficits may offer a strategy for the treatment of synaptic loss in AD and other synaptic disorders.

EXPERIMENTAL PROCEDURES

Materials—Bryostatin 1 was purchased from Biomol International (Farmingdale, NY, USA). DCPLA methyl ester (DCPLA-ME) was synthesized in our laboratory following the method described earlier (34,70) and shown to be specific for PKCε. Primary antibodies (rabbit polyclonal anti-PKCε (sc-214), rabbit polyclonal anti-PKCα (sc-208), rabbit polyclonal anti-PKCδ (sc-213), mouse monoclonal anti-synaptophysin (sc-17750) and mouse monoclonal anti-β-actin (sc-47778)) were obtained from Santa Cruz Biotechnology, Inc, (Santa Cruz, CA, USA). Rabbit polyclonal anti-synaptophysin (TA300431) and rabbit polyclonal anti-phospho-PSD-95 (serine-295) (TA303850) were obtained from Origene (Rockville, MD, USA) and rabbit polyclonal anti-PKD-95 (#3450) and rabbit polyclonal anti-JNK1/2 (#9258) were obtained from Cell Signaling Technology, Inc (Danvers, MA, USA). Chicken polyclonal anti-NeuN (ab134014) was obtained from Abcam (Cambridge, MA, USA). All secondary antibodies were purchased from Jackson ImmunoResearch Laboratories, Inc (Bar Harbor, ME, USA). The anti-chicken Cy5 conjugated antibody was purchased from Abcam (Cambridge, MA, USA). Bisindolylmaleimide I (Go 6850) and PKCε translocation inhibitor [EAVSLKPT] were obtained from Santa Cruz Biotechnology, Inc (Santa Cruz, CA, USA) and SP600125 and KN-93 were obtained from Cell Signaling Technology, Inc (Danvers, MA, USA).

Cell culture—Human primary neurons (Hippocampal neurons, Catalog No: 1540, ScienCell Research Laboratories, Carlsbad, CA, USA) were plated on poly-L-lysine coated plates and were maintained in neuronal medium (ScienCell) supplemented with the neuronal growth supplement (NGS, ScienCell). For maintenance of neurons half of the media was changed every 3 days. Fresh activators were added with every media change. Human HEK-293 cells were obtained from ATCC, Manassas, VA, USA. Cells were maintained in EMEM and 10% fetal bovine serum (FBS).

Organotypic slice culture—Organotypic hippocampal slices were prepared mainly according to the method described by Stoppini et al. (71) with slight modifications (72). Rats were sacrificed and immediately decapitated under sterile conditions. Brains were rapidly removed and placed into a chilled dissection medium (DM) composed of Hibernate A (BrainBits, Springfield, IL), 2% B27 supplement, 2 mM L-glutamine by GlutaMax and antibiotic–antimycotics (all from Invitrogen, Carlsbad, CA, USA). The hippocampi were dissected out in fresh chilled dissection medium. Isolated hippocampi were washed in new chilled DM and placed on a wet 3 mm paper on the Teflon stage of a manual tissue slice chopper (Vibratome Co., Saint Louis, MO, USA) for coronal sectioning at 300 μm. Each slice with intact pyramidal and granular layers was transferred to one membrane insert (Millipore, Bedford, MA, USA) in 12-well plates containing Neurobasal A, 20% horse serum, 2 mM l-glutamine, and antibiotics–antimycotics for four days. For long term maintenance slices were cultured in serum-free medium consisting of Neurobasal A with 2% B27, 2 mM l-glutamine, and antibiotic–antimycotics. Slices were incubated in a humidified 5% CO₂ atmosphere at 37 °C. The entire medium was replaced with fresh medium at day 1. After that, half the medium was removed and replaced with fresh medium twice a week.

Cell lysis and Western blot analysis—Cells were harvested in homogenizing buffer (HB) containing 10mM Tris-Cl (pH 7.4), 1 mM PMSF (phenylmethylsulfonylfluoride), 1 mM EGTA, 1 mM EDTA, 50 mM NaF and 20 µM leupeptin, and were lysed by sonication. The homogenate was centrifuged at 100,000 × g for 15 min at 4°C to obtain the cytosolic fraction (soluble) and membrane (particulate). The pellet was resuspended in the HB by sonication. For whole cell protein isolation from primary neurons the HB contained 1% Triton X-100. Protein concentration was measured using the Coomassie Plus (Bradford) Protein Assay kit (Pierce, Rockford, IL, USA). Following quantification, 20 μg of protein from each sample was subjected to SDS-PAGE analysis in a 4-20% gradient Tris-glycine polyacrylamide gel (Invitrogen, Carlsbad, CA, USA). The separated protein was then transferred to a nitrocellulose membrane. The membrane was blocked with BSA and incubated with primary
antibody overnight at 4°C. All the primary antibodies were used at 1:1000 dilution except rabbit polyclonal anti-p-PSD-95 \( (1:10000) \) and rabbit polyclonal anti-synaptophysin \( (1:10000) \). After incubation, it was washed 3× with TBS-T (Tris-buffered saline-Tween 20) and further incubated with alkaline phosphatase conjugated secondary antibody at 1:10000 dilution for 45 min. The membrane was finally washed 3× with TBS-T and developed using the 1-step NBT-BCIP substrate (Pierce, Rockford, IL, USA). The blot was imaged in an ImageQuant RT-ECL (GE Life Sciences, Piscataway, NJ) and densitometric quantification was performed using IMAL software. For quantifying expression of a protein, the densitometric value for the protein of interest was normalized against β-actin (loading control).

Electrophysiology—Rats (1 month old) were euthanized and hippocampus was isolated and sliced into 300-µm slices on a Leica VT1200S Vibratome. Slices were incubated in ACSF at room temperature for one hour until recording. The ACSF contains (mM): NaCl (124), KCl (3), MgSO4 (1.2), CaCl2 (2.1), Na2 PO4 (1.4), NaHCO2 (26), Dextrose (20) and is saturated with 95% O2 and 5% CO2 which maintains the pH at 7.4. Slices were treated with ethanol or bryostatin 1 for 1hr or 4hr. All recordings were made at room temperature. For synaptic stimulation and field excitatory synaptic potentials (EPSP) recordings pyramidal neurons in the CA1 field were identified with an Olympus BX50WI microscope. Field potential recordings were measured to determine synaptic function. A bipolar stimulating electrode (100-µm separation, FHC, Bowdoinham, ME) was placed in the hippocampal Schaffer collateral pathway to elicit EPSPs in CA1 stratum radiatum, EPSPs were recorded through patch pipettes (2-5 MΩ, 1.5 mm OD, 0.86 mm ID, P87 Brown-Flaming Puller, Sutter Instruments) filled with ACSF. All parameters including pulse duration, width, and frequency were computer controlled. Constant-current pulse intensity was controlled by a stimulus isolation unit. Basal synaptic transmission, represented by input-output responses, was determined by the slopes of stabilized EPSP to different stimulus intensities. The strength of EPSPs was assessed by measuring the slopes (initial 20-80%) of the EPSPs rising phase.

PKC Assay—To measure PKC activity, 100 ng recombinant PKCε (Sigma) was incubated for 15 min at 37°C in the presence of 100ng JNK1 or 100ng PSD-95 or 100 ng CaMKII, 4.89 mM CaCl2, 1.2 μg/µl phosphatidyl-L-serine, 0.18 μg/µl 1,2-diocanoyl-sn-glycerol, 10 mM MgCl2, 20 mM HEPES (pH 7.4), 0.8 mM EDTA, 4 mM EGTA, 4% glycerol, 8 μg/ml aprotinin, 8 μg/ml leupeptin, 2 mM benzamidine and 0.5 μCi of \( [\gamma-\text{32P}] \) ATP. \( [\text{32P}] \) Phosphoprotein formation was measured by adsorption onto phosphocellulose as described previously (70).

Knockdown and overexpression—Human PSD-95 was cloned into pCDNA3.1 plasmid (Genscript, Piscataway, NJ, USA). Mutant PSD-95 mutated at serine-295 residue was also cloned into pCDNA3.1 plasmid and was obtained from Genscript (Piscataway, NJ, USA). PKCε knockdown was done using PKCε-siRNA constructs purchased from Santa Cruz Biotechnology, Inc (Santa Cruz, CA, USA). JNK knockdown was done using SAPK/JNK-siRNA from Cell Signaling Technology, Inc (Danvers, MA, USA). Overexpression of PKCε was obtained by transfecting pCMV6-ENTRY vector containing human PKCε cDNA (Origene, Rockville, MD, USA). Transfection was done using Lipofectamine 3000 (Invitrogen, Carlsbad, CA, USA). Medium was changed after 6hr of lipofectamine treatment. Protein expression was measured after 72hr of transfection.

QRT-PCR—QRT-PCR was done following the method described earlier (13). Total RNA (500ng) was reverse-transcribed using oligo (dT) and Superscript III (Invitrogen, USA) at 50°C for 1 hr. The cDNA products were analyzed using a LightCycler 480 II (Roche) PCR machine and LightCycler 480 SYBR Green 1 master mix following the manufacturer’s protocol. Primers for PKCε (Forward Primer – AGCCTCGTTACGGTCTATGC, Reverse primer – GCAGTGACCTTCTGCATCCAGA), PSD-95 (Forward Primer – GCCACTTGACAGTGACCGA, Reverse primer – CGTCACTGTCTCGTAGCTCAGA), synaptophysin (Forward Primer – TCGGCTTTGTGAAGGTGCTGCA, Reverse primer – TCACTCTCGGTCTTGTTGGCAC), SNAP-25 (Forward Primer – CGTCTATGTGCTGCAACTGGTTG, Reverse primer – GTTCATGCTCTTTCTCGACACG),
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Syntaxin-1 (Forward Primer – TGGAGAACACGATCCGTAGCT, Reverse primer – CCTCTCACATACTCTACCGCG) and GAPDH (Forward Primer – GTCTCCTCTGACTTCAACAGCG, Reverse primer – ACCACCTGTGCTGTAGCCAA) were purchased from Origene (Rockville, MD, USA).

Electron microscopy—Electron microscopy of slices were done following methods described earlier (9). Hippocampi were sectioned with a vibratome at 100 μm. Hippocampi were fixed in 1% OsO4. Electron micrographs (100 μm² CA1 area at ×5,000) were made of Epon-embedded hippocampal sections with a JEOL 200CX electron microscope. These sections were 90 nm thick and had been previously stained with uranyl acetate and lead citrate. During quantification, electron micrographs were digitally zoomed up to ×20,000 magnification. Spines were defined as structures that formed synapses with axon boutons and did not contain mitochondria. Presynaptic vesicle density was measured from within the presynaptic axonal boutons that were seen to form synapses with dendritic spines of diameter larger than 600nm. Increased numbers of presynaptic vesicles in axon boutons were measured as an increase in the frequency of axon boutons with presynaptic vesicles that occupied more than 50% of the cross-section space not occupied by other organelles.

Immunofluorescence and Confocal microscopy—Cells were grown in four-chambered slides (Nunc, USA) at low density. For immunofluorescence staining the cells were washed with PBS (pH 7.4) and fixed with 4% paraformaldehyde for 4 min. Following fixation, cells were blocked and permeabilized with 5% horse serum and 0.3% Triton X-100 in 1× PBS for 30 min. Cells were washed 3 × with 1 × PBS and incubated with primary antibodies (rabbit polyclonal anti-PSD-95, mouse monoclonal anti-synaptophysin and chicken polyclonal anti-NeuN) for 1hr at 1:100 dilution. After the incubation slides were again washed 3 × in 1 × PBS and were incubated with the FITC anti-rabbit IgG, rhodamine anti-mouse IgG and Cy5 anti-chicken IgY for 1hr at 1:400 dilution. Cells were further washed and mounted in Pro Long Gold antifade mounting solution (Invitrogen, USA). Stained cells were viewed under the LSM 710 Meta confocal microscope (Zeiss) at 350 nm, 490 nm, 540 nm and 650nm excitation and 470 nm, 525 nm, 625 nm and 667nm emission for DAPI, FITC, rhodamine and Cy5 respectively. Six individual fields at 40× or 63× oil lens magnification were analyzed for the mean fluorescence intensity (MFI) in each channel. Punctate colocalization was done following methods described earlier (43-44).

Statistical analysis—All experiments were performed at least three times. Data are represented as mean ± SEM. All data were analyzed by one-way ANOVA and Newman-Keuls multiple comparison post test. Significantly different paired groups were further analyzed by two-tailed Student’s t-test using GraphPad Prism 6.1 software (La Jolla, CA, USA). P values < 0.05 were considered statistically significant.

CONFLICT OF INTEREST
The authors declare no competing financial interests.

AUTHORSHIP STATEMENT
AS, TJN and DLA designed the study and wrote the paper. AS performed and analyzed all the biochemical and immunofluorescence experiments, JH performed and analyzed all the electron microscopy data and DW performed and analyzed the electrophysiology experiments.

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PKCε induces synaptogenesis via PSD-95

Figure Legends:

Fig. 1: PKCε activation prevents degeneration of human primary neurons. Primary human neurons were treated with either DCPLA-ME (100 nM) or bryostatin 1 (0.27 nM) for 40 days. Fresh drug was added every third day with 50% media change. A. Image of 40 day old untreated (Control), bryostatin 1 and DCPLA-ME treated neurons. B. Number of neurite positive cells counted from three 20× fields (508 μm²) over time. DCPLA-ME and bryostatin 1 treatment stabilized cellular viability for at least 40 days. Viability of untreated cells declined after 20 days. C-E. PKCε, PSD-95 and synaptophysin mRNA levels in 40 day old neurons compared to 1 day neurons. F. Immunoblot analysis of PKCε, p-PSD-95S295, PSD-95 and synaptophysin in 40 day old neurons compared to 1 day neurons. G-J. Immunostaining of p-PSD-95S295, PSD-95 and synaptophysin calculated from immunoblots. Staining is significantly higher in DCPLA-ME and bryostatin 1 treated cells. Data are represented as mean ± SE of three independent experiments (Student’s t-test, *P<0.05and **P<0.005).

Fig. 2: PKCε activation induces membrane accumulation of p-PSD-95S295. Primary human neurons were treated with ethanol (C), bryostatin 1 (0.27nM) or DCPLA-ME (100nM) for 1hr, 4hr and 24hr. Neurons were separated into soluble (S) and membrane (P) fractions and immunoblotted against PKCε, PKCε and PKCδ. (A-C) and p-PSD-95S295. PSD-95 and β−actin (E). ‘M’ represent molecular weight marker. PKCε activation is reported as the percentage of total protein in the membrane. Bryostatin 1-treated neurons showed PKCε activation at 1hr (175.2 ± 9.5 %; P=0.002), 4hr (181.6 ± 10.2 %; P=0.0016) and 24hr (170.9 ± 7.4 %; P=0.001) compared to the untreated neurons (100.0 ± 3.1%; F (3,8)=22.5; ANOVA, P<0.005) (B) and DCPLA-ME treated neurons showed a significant increase in PKCε activation at 1hr (144.0 ± 6.8 %;P=0.004), 4hr (146.5 ± 6.8 %; P=0.003) and 24hr (133.6 ± 4.2 %; P=0.003) compared to the untreated neurons (F (3,8)=15.7; ANOVA, P=0.001) (D) induced PKCε activation at 1hr, 4hr and 24hr. PKCε activation significantly increased the p-PSD-95S295 expression in the membranes at 1hr and 4hr (F). Adult rat hippocampal organotypic slices were treated with ethanol (C), bryostatin 1 (0.27nM) or DCPLA-ME (100nM) for 1hr, 4hr and 24hr. Protein lysates were immunoblotted against p-PSD-95S295, PSD-95 and β−actin. ‘M’ represent molecular weight marker (G). PKCε activation significantly increased the p-PSD-95S295 expression in the membranes at 1hr and 4hr (H). Data are represented as mean ± SE of three independent experiments (Student’s t-test, *P<0.05and **P<0.005).

Fig. 3: PKCε mediated phosphorylation of PSD-95 at serine-295 is essential for its membrane accumulation. A. Immunoblot representing p-PSD-95S295 and PSD-95 expression following incubation of the different combinations of recombinant PKCε, PSD-95, PKCε activators and PKCε inhibitors mentioned above at 37°C for 10 min in vitro. (B.) Bryostatin 1 (0.27nM) and DCPLA-ME (100nM) induces the phosphorylation of PSD-95 at serine-295 position. C. Expression of p-PSD-95S295 and PSD-95 and β−actin in HEK-293 cells transfected with empty vector, wild type human PSD-95 and mutant PSD-95S295K. D. Expression of PSD-95, p-PSD-95S295, PKCε and β−actin in the cytosol and membrane fraction of wild-PSD-95 and mutant PSD-95S295K transfected HEK-293 cells treated with bryostatin 1 and DCPLA-ME for 4hr in presence or absence of EAVSLKPT (5μM). Percentage of total protein in the membrane; PKCε (E), PSD-95 (F) and p-PSD-95S295 (G). Data are represented as mean ± SE of three independent experiments (Student’s t-test, *P<0.05, **P<0.005and ***P<0.005).

Fig. 4: PKCε mediated membrane localization of p-PSD-95S295 involves JNK1 and CaMKII. Immunoblot showing p-PSD-95S295 expression in the membrane of neurons (A) treated with vehicle, Bisindolylmaleimide I (100nM, PKC inhibitor), SP600125 (20μM, JNK inhibitor) and K-93 (10μM, CaMKII inhibitor) for 4hr. B. p-PSD-95S295 expression in the membrane of neurons treated with DCPLA-ME in presence or absence of different inhibitors. C. The inhibitors alone reduced the expression of membrane bound p-PSD-95S295. D. PKC, JNK and CaMKII inhibitors prevented the effect of PKCε activation on p-PSD-95. Activated PKCε increases phosphorylation of PSD-95 (E), JNK1 (F) and CaMKII (G). H. Immunoblot showing the downregulation of PKCε and JNK in PKCε and JNK si-RNA
treated neurons respectively.  

**I.** Protein expression of p-PSD-95$^{S295}$ and β-actin in the membrane of control, PKCε KD and JNK KD neurons in presence or absence of DCPLA-ME.  

**J.** Either in PKCε or JNK knockdown human neurons, DCPLA-ME failed to induce the membrane accumulation of p-PSD-95$^{S295}$.  

**K.** Diagram representing PKCε mediated membrane translocation of p-PSD-95$^{S295}$ and involvement of JNK1 and CaMKII in the pathway. Data are represented as mean ± SE of three independent experiments (Student’s t-test, *P* < 0.05 and **P** < 0.005).

**Fig. 5:** PKCε activation induces synaptogenesis. Adult rat hippocampal organotypic slices were treated with ethanol (C), bryostatin 1 (0.27nM) or DCPLA-ME (100nM) for 1hr and 4hr. **A.** Electron micrographs of the stratum radiatum in the hippocampal CA1 area (100 μm$^2$ CA1 area at ×5,000) treated with bryostatin 1 and DCPLA-ME for 4hr. Thirty to thirty-five CA1 areas each from three different hippocampal slices were analyzed. Dendritic spines showing synapse are highlighted in yellow. **B.** PKCε activation increased the synapse number at 4hr (F (2, 96) =9.05; ANOVA P< 0.0005) in bryostatin 1 (8.97 ± 0.63, P< 0.005, n=34 CA1 area) and DCPLA-ME (6.97 ± 0.5, P< 0.05, n=30 CA1 area) treated slices compared to control (5.77 ± 0.50). Typical traces of EPSPs evoked at a stimulus intensity of 200μA from bryostatin 1 treated hippocampal slices after 1hr (C) and 4hr (F). The input–output response, reflecting basal synaptic transmission, increased after treatment with bryostatin1 (0.27 nM), after 1hr (D) and 4hr (G). Areas under the curves (AUCs) were calculated to compare the basal levels of synaptic transmission. Bryostatin 1 increased EPSP slope significantly at 1hr and 4hr (E, H). Data are represented as mean ± SE of three independent experiments (Student’s t-test, *P* <0.05 and **P** <0.005).

**Fig. 6:** PKCε is essential for the expression of PSD-95 and synaptophysin. Primary human neurons were transfected with empty vector /scrambled siRNA (C), PKCε siRNA (PKCε KD), or a PKCε over-expression vector (PKCε OE) following method described in the ‘Method’ section. Cells were analyzed 72hr after treatment.  

**A.** mRNA transcript levels of PKCε, B. PSD-95, C. synaptophysin, D. SNAP-25 and E. syntaxin-1 in PKCε KD and PKCε OE neurons. PKCε KD suppressed while PKCε OE induced PSD-95 and synaptophysin mRNA transcript.  

**F.** Agarose gel image showing no effect of scrambled siRNA on PKCε mRNA.  

**G.** Immunoblot showing protein expression of PKCε in untreated, PKCε siRNA and scrambled siRNA treated human neurons.  

**H.** Immunoblot staining of PKCε, PSD-95, synaptophysin and actin in control, PKCε KD and PKCε OE neurons. **I, J, K.** Graphical representation of protein expression levels of PKCε, PSD-95, synaptophysin in control, PKCε KD and PKCε OE neurons. Data are represented as mean ± SE of three independent experiments (Student’s t-test, *P* <0.05 and **P** <0.005).

**Fig. 7:** Loss of PKCε prevents synaptogenesis.  

**A.** Confocal images of untreated, DCPLA-ME (100 nM), bryostatin 1 (0.27 nM), PKCε siRNA + DCPLA-ME (100 nM) and PKCε siRNA + bryostatin 1 (0.27 nM) treated primary human neurons (10 days). Each condition is represented by five panels. Four square panels represents nucleus (blue), PSD-95 (green), synaptophysin (red) and merged image respectively. The rectangular panel represents magnified image of a 40μm neurite.  

**B.** Number of PSD 95 signal grains was measured along 40μm neurite length (10 individual neurites from 4 independent slides). Bryostatin 1 and DCPLA-ME significantly increased the PSD-95 clusters per 40μm neurite (F (2,9)=4.5; ANOVA P<0.05).  

**C.** Synapses were quantified by the number of colocalized PSD-95 and synaptophysin signals. PKCε activation increased synapse number (F (2,9)=6.1; ANOVA P<0.05) and PKCε KO prevented the synaptogenic effect of PKC activators.  

**D.** Immunoblot analysis of PKCε, p-PSD-95$^{S295}$, PSD-95 and synaptophysin. PKCε knockdown (KD) reduces PKCε expression by 50% after 10 days in human neurons. **E, F, G.** At 10 days PKCε activation increased the expression of PSD-95 and synaptophysin significantly, but in PKCε KD cells their expressions were lower even after treatment with activators. Data are represented as mean ± SE of at least three independent experiments (Student’s t-test, *P* <0.05 and **P** <0.005).
Fig. 8: PKCε activation induces synaptogenesis in hippocampal slices. A. Electron microscopy of the hippocampal CA1 area from adult organotypic brain slices treated with vehicle (Control), DCPLA-ME (100 nM) and bryostatin 1 (0.27 nM) for 10 days. Dendritic spines showing synapse are highlighted in yellow. B. PKCε activation by bryostatin 1 and DCPLA-ME increased synapse number in adult organotypic brain slices (F(2,6) = 11.9; ANOVA P<0.01). C. Electron micrograph showing increased presynaptic vesicle density in PKC activator treated slices. Grey level of presynaptic vesicle stack from six to seven presynaptic boutons was measured from three different hippocampal slices. “D” represents dendritic spine, red arrow marks synapse and yellow marks presynaptic vesicles. D. Bryostatin 1 and DCPLA-ME significantly induced the presynaptic vesicle density at 10 days. Data are represented as mean ± SE of at least three independent experiments (Student’s t-test, *P<0.05 and **P<0.005).
Figure 2
**Figure 4**

**A**

Graph showing protein bands at 100 kDa and 50 kDa, with corresponding protein labels: p-PSD-95\(^{5295}\), β-Actin, and Bisindolylmaleimide I (PKC) measured with treatments including SP600125 (JNK) and K-93 (CaMKII).

**B**

Similar to A, showing protein bands and treatments with DCPLA-ME and Bisindolylmaleimide I (PKC).

**C**

Bar graph comparing p-PSD-95\(^{5295}\) in membrane (percent of control) under various conditions: Control, PKC, JNK, CaMKII, and combinations thereof.

**D**

Similar to C, showing p-PSD-95\(^{5295}\) in membrane for different treatments including DCPLA-ME and combined inhibitors.

**E**

Bar graph illustrating PKCε phosphorylates PSD-95 with treatment results.

**F**

Bar graph showing JNK phosphorylation with treatment results.

**G**

Bar graph depicting CaMKII phosphorylation with treatment results.

**H**

Western blot images showing protein bands at 100 kDa, 75 kDa, 50 kDa, 37 kDa, and β-Actin under various conditions: PKCε KD, JNK KD, PKCε + PKCε, PKCε + JNK1, and CaMKII + CaMKII inh.

**I**

Western blot images showing p-PSD-95\(^{5295}\) in membrane with treatment results.

**J**

Bar graph comparing p-PSD-95\(^{5295}\) in membrane (percent of control) for different treatments including DCPLA-ME, PKCε KD, JNK KD, PKCε KD + DCPLA-ME, and JNK KD + DCPLA-ME.

**K**

Schematic diagram illustrating the regulation of PSD-95 by PKCε and CaMKII, with arrows indicating phosphorylation (+p) and membrane/cytosol localization. The diagram includes JNK1 with +p indication and PKCε and CaMKII with dashed lines.
Figure 5
Figure 6

A. PKCε mRNA
B. Synaptophysin mRNA
C. PSD95 mRNA
D. SNAP-25 mRNA

E. Syntaxin-1 mRNA

F. Gel electrophoresis images showing differential expression of PKCε and β-Actin with PKCε siRNA and scrambled siRNA.

G. Western blot images showing differential expression of PKCε and β-Actin with PKCε siRNA and scrambled siRNA.

I. PKCε expression

J. Synaptophysin expression

K. PSD95 expression

Figure 6
Figure 7

A 10 days untreated

B

C

D

E

F

G

Figure 7
PKC epsilon Promotes Synaptogenesis through Membrane Accumulation of the Postsynaptic Density Protein PSD-95
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