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Prenatal Cocaine Exposure Increases Synaptic Localization of a Neuronal RasGEF, GRASP-1 via Hyperphosphorylation of AMPAR Anchoring Protein, GRIP

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Abstract

Prenatal cocaine exposure causes sustained phosphorylation of the synaptic anchoring protein, glutamate receptor interacting protein (GRIP1/2), preventing synaptic targeting of the GluR2/3-containing alpha-aminooacid-3-hydroxy-5-methyl-4-isoxazolepropionic acid-type glutamate receptors (AMPARs; J. Neurosci. 29: 6308–6319, 2009). Because overexpression of GRIP-associated neuronal rasGEF protein (GRASP-1) specifically reduces the synaptic targeting of AMPARs, we hypothesized that prenatal cocaine exposure enhances GRASP-1 synaptic membrane localization leading to hyper-activation of ras family proteins and heightened actin polymerization. Our results show a markedly increased GRIP1-associated GRASP-1 content with approximately 40% reduction in its rasGEF activity in frontal cortices (FCX) of 21-day-old (P21) prenatal cocaine-exposed rats. This cocaine effect is the result of a persistent protein kinase C (PKC)- and downstream Src tyrosine kinase-mediated GRIP phosphorylation. The hyperactivated PKC also increased membrane-associated GRASP-1 and activated small G-proteins RhoA, cdc42/Rac1 and Rap1 as well as filamentous actin (F-actin) levels without an effect on the phosphorylation state of actin. Since increased F-actin facilitates protein transport, our results suggest that increased GRASP-1 synaptic localization in prenatal cocaine-exposed brains is an adaptive response to restoring the synaptic expression of AMPA-GluR2/3. Our earlier data demonstrated that persistent PKC-mediated GRIP phosphorylation reduces GluR2/3 synaptic targeting in prenatal cocaine-exposed brains, we now show that the increased GRIP-associated GRASP-1 may contribute to the reduction in GluR2/3 synaptic expression and AMPAR signaling defects.

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Introduction

Prenatal cocaine exposure results in long-lasting changes in synaptic plasticity that may be responsible for the cognitive deficits in humans and animal models [1], [2], [3]. Synaptic plasticity, such as long-term potentiation (LTP) and depression (LTD) are regulated in part by AMPARs [4], [5]. The notion that AMPARs are sensitive to cocaine is supported by our earlier findings that prenatal cocaine exposure reduces GluR2- and GluR3-AMPAR synaptic expression and attenuates AMPAR-mediated LTD [6]. We demonstrated that reduced GluR2-, GluR3-containing AMPARs on the synaptic membrane is the result of a deficient interaction between AMPARs and synaptic anchoring protein GRIP resulting from sustained PKC-mediated GRIP phosphorylation [6]. However, there are no data discerning how this cocaine-induced effect influences other GRIP-associated proteins such as GRASP-1, which is known to affect GluR2/3-GRIP interaction [7] and regulate AMPAR trafficking to the synaptic membrane.

GRASP-1 is a neuronal RasGEF (guanine nucleotide exchange factor) and a neuron- specific effector for monomeric guanine nucleotide-binding proteins (G proteins) such as Rab4 which have been implicated in the regulation of membrane protein trafficking [7], [8], AMPAR targeting, and JNK signaling [7], [9]. Through its RasGEF domain in the N terminal region, GRASP-1 associates with GRIP1 by binding to the seventh PDZ domain of GRIP located at the C terminal region that is distinct from the GluR2/3 binding site on GRIP [7]. This GRIP-GRASP-1 association enables the formation of the GRASP-1/GRIP/GluR2 complex which activates monomeric G proteins via GRASP-1’s RasGEF activity. In addition, the level of GRIP associated AMPAR(s) is reduced when GRASP-1 is overexpressed [7].

Monomeric G proteins are prominent regulators of the actin network that has profound influence on the trafficking of surface
proteins including GluR2/3-AMPARs. It is highly likely that hyperphosphorylation of GRIP in prenatal cocaine exposed brains noted in our earlier work [6] could affect the GRIP-GRASP-1 association, and thus the GRASP-1 cellular distribution and/or activity. The altered GRIP-GRASP-1 complex level in prenatal cocaine-exposed brain could in turn influence GluR2/3 trafficking via modulating the activity of monomeric G proteins.

In accordance with this hypothesis, monomeric G proteins such as Ras and Rap regulate AMPAR trafficking during LTP and LTD [10], and an increase in active Rap1 is associated with reduced synaptic targeting of GluR2 [11]. Monomeric G protein Rac1/cdc42, clusters AMPARs during spinogenesis [12] and regulates synaptic structure and function [13], whereas RhoA regulates actin polymerization in dendritic spines and modulates spine length and density [14], [15]. Altogether, these data suggest strongly that alteration of monomeric G protein levels and/or activity could influence AMPAR trafficking and consequently the synaptic function.

We have previously demonstrated that in prenatal cocaine-exposed brain PKC- and Src-mediated persistent phosphorylation of GRIP reduces GluR2/3 synaptic expression [6]. Here, we show that hyperphosphorylated GRIP resulting from persistently activated PKC also increases GRASP-1 association with GRIP, leading to higher activated RhoA, cdc42/Rac1, and Rap1, as well as F-actin levels in FCX of prenatal cocaine-exposed rats. Their relevance to AMPAR trafficking is discussed.

Results

Prenatal cocaine exposure markedly increases GRASP-1 localization and GRASP-1 - GRIP interaction in the synaptic membrane

The GRIP binding partner GRASP-1 can regulate AMPAR targeting by forming a complex with GluR2/3-AMPARs via its association with GRIP [7]. To test directly whether prenatal cocaine exposure changes the expression and/or cellular distribution of GRASP-1, the level of GRASP-1 was measured in total post-mitochondrial FCX lysate as well as in cytoplasmic and membrane fractions of FCX synaptosomes prepared from 21-day-old (P21) prenatal cocaine- and saline-exposed rats. Prenatal cocaine exposure did not alter the expression of GRASP-1 (Fig. 1a,b), GRASP-1 localizes predominantly in the cytosol of FCX synaptosomes of saline-exposed brains, corresponding to 82.9±1.7% of overall synaptosomal GRASP-1 expression. Prenatal cocaine exposure dramatically increased the membrane-associated GRASP-1 level to 63.7±2.6% of the overall synaptosomal GRASP-1 expression level (Fig. 1c,d). A higher level of membrane-localized GRASP-1 in prenatal cocaine-treated brain was also evidenced immunohistochemically by the increased GRASP-1 immunoreactive puncta on membrane (Fig. 1e).

Because GRASP-1 also associates with GRIP, we next examined whether an increase in GRASP-1 membrane localization is mediated by the heightened association between GRASP-1 and GRIP1. Using an in vitro association assay with individually purified GRASP-1 and GRIP1 from FCX of P21 prenatal saline- and cocaine-exposed rats, we demonstrate that prenatal cocaine exposure markedly increases GRASP-1 and GRIP1 association (Fig. 2a,b). Since GRASP-1 overexpression reduces AMPAR synaptic expression [7], the increase in membrane-localized GRASP-1 observed in prenatal cocaine-exposed brain suggests that the heightened GRIP-associated GRASP-1 may contribute to the reduced GluR2/3-AMPARs synaptic targeting observed here and previously [6].

PKC- and Src-mediated phosphorylation of GRIP1 increases GRASP-1 and GRIP1 interaction in prenatal cocaine-exposed brain

Because hyper-phosphorylation of GRIP1 mediates reduced GluR2/3 synaptic expression in prenatal cocaine-exposed brains [6], we examined whether hyper-phosphorylation of GRIP also promotes GRIP-GRASP-1 coupling. Confirming our previous observation, an increased GRIP1 phosphorylation on serine and tyrosine but not threonine residues was noted in FCX from prenatal cocaine-exposed rats (Fig. 2a). Whether hyperphosphorylation of GRIP1 increases the GRASP-1 – GRIP1 interaction was then assessed following treatment of GRIP1 with alkaline phosphatase: dephosphorylation sharply reduced GRIP1-associated GRASP-1 (Fig. 2a,b). Further, increases in GRIP1 phosphorylation on serine and tyrosine, respectively by PKC and Src, heightened GRIP1-coupled GRASP-1 levels (Fig. 2a,b). Together with our earlier work, these results support the conclusion that PKC and Src-mediated phosphorylation of GRIP1 increases GRASP-1 coupling but decreases GluR2/3 coupling to GRIP1.

Prenatal cocaine exposure reduces GRASP-1 RasGEF activity

Since GRASP-1 contains the catalytic domain of guanine nucleotide exchange factors (GEF) for the ras family of G proteins [7], we compared whether prenatal cocaine affects the RasGEF activity of GRASP-1. The kinetics and capacity of GRASP-1, immunoactivity-purified from FCX of prenatal cocaine- and saline-exposed, were analyzed using a time-course and the level of [3H]GDP was released from [3H]GDP pre-loaded immunopurified RhoA and Rap1. Our results show that although 76.7±3.3% and 81.3±5.2% of [3H]GDP was released from RhoA and Rap1, respectively, within 5 min following exposure to purified GRASP-1 obtained from prenatal saline-exposed brains, GRASP-1 from prenatal cocaine-exposed brains only reduced 62.0±1.5% and 69.7±3.2% of [3H]GDP from RhoA and Rap1, respectively in 5 min (Fig. 3). Further analysis revealed that prenatal cocaine exposure reduces GRASP-1 GEF activity on RhoA and Rap1, respectively, by approximately 40% from 0.87±0.07 to 0.69±0.08 to 0.46±0.03 and 0.42±0.03 nmol/min (Fig. 3).

Prenatal cocaine exposure increases activated RhoA, Rap1 and Rac1/cdc42 levels

To investigate the impact of increased levels of GRIP-associated GRASP-1 and reduced RasGEF activity in prenatal cocaine-exposed brains, we compared levels of activated RhoA, Rac1/cdc42 and Rap1, all of which modulate AMPAR trafficking through their effects on the actin-dependent cytoskeleton [10], [11], [12]. Using GST-conjugated Rhotekin-, RalGDS- and GST-Pak1-RBD beads to isolate GTP-bound (activated) RhoA, Rap1 and Rac1/cdc42 respectively, we found that prenatal cocaine exposure markedly increased activated RhoA, Rap1 and Rac1/cdc42 without a detectable difference in their expression levels (Fig. 4a,b).

To determine whether the increased activated RhoA, Rap1 and Rac1/cdc42 in the prenatal cocaine-exposed brain is also related to the hyper-activated PKC [6], organotypic FCX slice cultures were treated with combination of γ and ζ-PKC specific pseudosubstrate inhibitor peptides (GIP and ZIP) or control scrambled peptides (GIP SC and ZIP SC). The data summarized in Fig. 5a, b indicate that GIP and ZIP robustly reduce the active RhoA, Rap1 and Rac1/cdc42 levels to that of saline-treated animals. Together with the data showing that a blockade of PKC...
by a combination of GIP and ZIP normalizes GRASP-1 and GluR2 cellular distribution (Fig. 6), these data indicate that the increased active RhoA, Rap1 and Rac1/cdc42 in prenatal cocaine brains originated from a persistent activation of PKC. These data further support the hypothesis that prenatal cocaine exposure alters GluR2/3 trafficking by increasing the relative abundance of polymerized (filamentous) actin.

Prenatal cocaine exposure increases F-actin levels

Toda et al (2006) indicate that withdrawal from repeated cocaine in the adult rat increases actin cycling and actin polymerization in the nucleus accumbens, suggesting that actin cycling and polymerization may be altered by in utero cocaine exposure [16]. To test this hypothesis, we measured F-actin levels in synaptosomes prepared from FCX of prenatal cocaine- and
saline-exposed rats using rhodamine-conjugated phalloidin, a
toxin which binds tightly to F-actin. The data summarized in
Fig. 7a shows a 2.2-fold increase in F-actin level in FCX
synaptosomes from P21 prenatal cocaine-exposed rats. This
prenatal cocaine-induced effect was not caused by a heightened
actin phosphorylation since serine-, threonine- and tyrosine-
phosphorylated actin levels were comparable in both FCX and
prefrontal cortex between P21 prenatal saline- and cocaine-
exposed rats (Fig. 7b). Since treatment of synaptosomes with
alkaline phosphatase normalized F-actin levels, this result
indicates that phosphorylation of the upstream actin regulators
such as GRIP by PKC is pivotal in determining the F-actin level,
(Fig. 7a).

Discussion

Cocaine exposure in utero can modify synaptic plasticity at
excitatory synapses resulting in long-lasting deficits in brain
function and altered cognitive and psychological development.
Our previous findings show that prenatal cocaine exposure attenuates AMPAR-mediated LTD and reduces AMPARs in the synaptic membrane resulting from a disrupted GluR2/3–GRIP interaction [6]. The reduced GluR2/3–GRIP association in prenatal cocaine-exposed brains is the result of a sustained PKC-mediated phosphorylation of the AMPAR scaffolding protein GRIP [6]. Resonating with our finding of altered GRIP in prenatal cocaine-exposed brain, chronic cocaine exposure of adult brains also changes the expression or function of glutamate receptor scaffolding proteins such as PSD-95 and Homer [17]–[20]. Collectively, these data suggest that scaffolding proteins for the glutamatergic receptors such as AMPARs in the postsynaptic density are prominent targets of cocaine.

In addition to serving as a synaptic anchor for AMPARs, GRIP also interacts with other signaling molecules including GRASP-1 [7], liprin-α [21], ephrin B receptors [22], and matrix metalloproteinase 5 [23]. Although the precise mechanism through which GRIP-interacting signaling molecules contribute to the reduced GluR2/3 synaptic membrane localization in prenatal cocaine-exposed brain remains ambiguous, a previous demonstration that overexpression of GRASP-1 in cultured hippocampal neurons can reduce AMPAR synaptic targeting suggests that an overly active or abundant GRASP-1 may hinder GluR2/3-GRIP interaction [7]. This hypothesis is supported by our current data showing a markedly higher synaptic localization of GRASP-1 (and compensatory reduction in cytosolic GRASP-1 level) in FCX of prenatal cocaine-exposed brains.
cocaine-exposed rats. This prenatal cocaine-induced GRASP-1 synaptic localization is caused by a greater coupling of GRASP-1 to GRIP resulting from sustained PKC and Src-mediated phosphorylation of GRIP that we previously showed to reduce GluR2/3-GRIP association [6]. An increased GRASP-1 - GRIP association may present a physical hindrance preventing efficient GluR2/3 - GRIP1 binding, although GRASP-1 and GluR2/3 bind to different PDZ domains on GRIP. Alternatively, an increased GRASP-1 and GRIP interaction may alter GRIP conformation rendering the GluR2/3 binding sites on GRIP inaccessible.

Figure 6. Blockade of PKCγ and PKC/β with isozyme-specific pseudosubstrate inhibitors abolishes prenatal cocaine exposure induced increase and decrease in synaptic membrane associated GRASP-1 and GluR2, respectively. (a) Organotypic FCX slice cultures were serum-depleted and treated with combination of myristoylated PKCγ- and PKC/β-specific pseudosubstrate inhibitors (GIP and ZIP), 10 μM each or 10 μM control peptide, GIP/ZIP scrambled (GIP/ZIP SC) for 4 hr. The slices were homogenized to yield synaptosomes and the resultant synaptosomes were hypotonically lysed to yield synaptic membranes (M) and cytosol (Cy). The levels of GRASP-1 and GluR2 were then determined sequentially using Western blotting. The blots were stripped and re-probed sequentially to measure the exclusive cytosolic and membranous markers, caspase-3 and GRIP to illustrate equal loading. (b) Densitometric quantification of blots. n=4. Data are means ± s.e.m. of the optical intensity. *p<0.01 compared to respective protein in the respective GIP/ZIP SC-treated group. #p<0.01 compared to respective protein in the saline-treated group. doi:10.1371/journal.pone.0025019.g006

Figure 7. Prenatal cocaine exposure induces an increased F-actin level which is normalized by protein phosphatase treatment without alteration in phosphorylation state of actin. (a) Synaptosomes obtained from the frontal cortices (FCX) of P21, prenatal cocaine- or saline-treated rats were treated with alkaline phosphatases, alkaline phosphatases (1 mg/ml) plus phosphatase inhibitors or cytochalasin D (10 μM) in vitro. The reactions were terminated, synaptosomes solubilized, and actin-containing proteins extracted using biotin-conjugated anti-actin antibodies and placed into streptavidin-coated plates. The level of F-actin was determined by rhodamine-conjugated phalloidin and the fluorescence intensity of phalloidin was measured using Beckman multimode plate reader, DX880. n=6. Data are means ± s.e.m. of the fluorescence intensity. The statistical significance was evaluated by Newman-Keuls multiple comparisons that followed one-way ANOVA. *p<0.01 compared to native GRIP1 from saline-treated group. #p<0.01 compared to dephosphorylated GRIP in cocaine-treated group. +p<0.01 compared with native GRIP1 from respective group. (b) The phosphorylation state of actin was evaluated in synaptosomes derived from the FCX of P21, prenatal cocaine- or saline-treated rats. Total actin was purified by immunoprecipitation with anti-actin and the levels of phosphor-serine (pS), -threonine (pT) and -tyrosine (pY) in the anti-actin immunoprecipitate was determined by Western blotting using phosphoepitope-specific antibodies. n=4. Data are means ± s.e.m. of the ratios of pS-, pT- or pY-actin to total actin optical intensities. There was no discernible difference noted in any of the actin phosphoepitopes in FCX of the prenatal cocaine- comparing to saline-exposed rats. doi:10.1371/journal.pone.0025019.g007
In addition to a greater GRASP-1 synaptic localization, prenatal cocaine exposure slows enzymatic kinetics without altering GRASP-1 RasGEF capacity but dramatically increases the level of active monomeric G proteins such as RhoA, cdc42/ Rac1 and Rap1. These data suggest that the tremendous increase in membrane-associated GRASP-1 through interaction with GRIP overcomes slower enzymatic kinetics to promote monomeric G protein activation that may ultimately influence the AMPAR-regulated synaptic transmission. The GRASP-1 having rasGEF activity observed herein agrees with the earlier finding by Ye et al. (2000) but sharply contrasts to a recent report indicating that GRASP-1 lacks enzymatic activity [10]. While the precise reason for the discrepancy in whether GRASP-1 possesses GEF activity remains elusive, there are clear methodological and species differences in these three studies that may have contributed to the opposite findings.

In agreement with the notion that increased active monomeric G proteins contribute to AMPAR synaptic transmission regulation, Rap mediates NMDA receptor-dependent, activity-induced LTD by removing GluR2/3-containing AMPARs from the synaptic membrane [10] and suppresses synaptic transmission by reducing GluR2 surface expression [11]. Hence, increased active Rap1 observed here may contribute to the previously observed reduction in synaptic targeting of GluR2/3 containing AMPARs and LTD in prenatal cocaine-exposed brains [6]. In contrast to Rap1, activated cdc42/Rac1 induces clustering of AMPARs in dendritic spines [24]. Thus, an elevated active Rac1 level in prenatal cocaine-exposed brains may compensate for the reduced AMPAR synaptic transmission by enabling higher transportation and clustering of AMPARs. RhoA is localized in the postsynaptic density and is associated with excitatory glutamatergic receptors at the spine plasma membrane [25]. Since NMDAR and AMPAR activation dampens RhoA activity and destabilizes actin networks [25], the reduced glutamatergic NMDAR (unpublished findings) and AMPAR activity together with the elevated association of GRIP and GRASP-1 should afford a more stable actin cytoskeleton in the prenatal cocaine-exposed brain. This notion of a more stable actin network is supported by our demonstration that F-actin levels are higher in FCX synaptosomes of prenatal cocaine-exposed rats. The Rho family of small GTPases also plays a pivotal role in regulating spine architecture and synaptic plasticity [26], [27]. Hence, an abnormally upregulated RhoA activity in prenatal cocaine-exposed brains may adversely influence neuronal development leading to cognitive deficits [28], [29].

The increased F-actin level in FCX synaptosomes of prenatal cocaine-exposed rat may promote GluR2/3 trafficking since latrunculin A, an actin-depolymerizing agent, reduces AMPAR-containing spines in cultured hippocampal pyramidal neurons [30]. By contrast, Rac1 and RhoA activation reduces dendritic pruning in hippocampal pyramidal neurons [31]. Interestingly, increased activated Rac1 was shown to attenuate synaptic and cognitive functions such as learning and memory [32]. Nonetheless, these data indicate that excessive RhoA and Rac1 activation in prenatal cocaine-exposed brain may have facilitated GluR2/3 transport to membrane to compensate for GluR2/3-GRIP interaction blockade and altered dendritic morphology. The latter agrees with our findings that cocaine exposure in utero increases dendritic spine density in rats [33], [34], dendritic length in rabbits [35], as well as dendritic length, volume, and extension in mouse FCX [36]. Since three weeks of withdrawal from repeated cocaine exposure increased F-actin levels [16], [37], it is also possible that the increased F-actin levels in prenatal cocaine-exposed rats could be caused by extended cocaine abstinence. In contrast to prenatal cocaine affects on F-actin but not overall actin level shown here, a 24-hr exposure of the human fetal cortical cells derived at 20-week gestation to 100 μM cocaine results in down-regulation of the cytoskeleton-related genes [38]. Such discrepancy may be related to different experimental systems used, including rodent vs. human cortices, 24-hr constant exposure vs. in vivo administration, and cytoskeletal protein vs. gene levels.

Most importantly, we show here that the persistently increased membrane localization of the activated PKC in prenatal cocaine exposed brain reported previously [6] is the primary mechanism underlying excessive GRASP-1 association with GRIP that leads to the elevated monomeric G proteins. Our data showing that pseudosubstrate PKC inhibitors targeting PKCγ and PKC/βII normalize the monomeric G protein activity in prenatal cocaine exposed rats further demonstrates the critical role of membrane localized, activated PKC. Additional support can also be drawn from previous reports showing that PKC activation in cultured hippocampal neurons induces the formation of dendritic lamellae in a Rho/Rac-dependent manner [39], [40]. Given that aberrant PKC overactivation leads to abnormal dendritic spine density, morphology and function [41], [42], the abnormally hyperactivated PKC and monomeric G proteins may act in tandem to promote AMPAR synaptic transmission and dendritic abnormalities observed in prenatal cocaine-exposed brains [6], [33], [34]. Such drastic functional and structural defects most likely play an important role in mediating the eventual cognitive changes, including impaired reward processing in animal models [43]–[46], [34] and in humans [47]. Future experiments are needed to determine whether the observed increase in GRASP-1 membrane localization in brains from P21 prenatal cocaine-exposed rats is persistent or simply a transient modification of synaptic plasticity during early development.

Given that a sustained PKC activation, indicated by an overwhelming presence of synaptic membrane-associated multiple PKC isoforms, and a markedly reduced phorbol ester-induced PKC translocation were observed in adult rabbit brains exposed to cocaine in utero [48], it is highly likely that the elevated GRASP-1 membrane localization persists into adulthood. Previous studies conducted by us and others in rabbits also indicate that such synaptic plasticity changes last well into adulthood [49], [49], [45].

In summary, our results indicate that increased GRASP-1 membrane localization resulting from sustained PKC- and Src-mediated phosphorylation of GRIP plays a significant role in mediating AMPAR dysfunction and dendritic abnormalities in the prenatal cocaine-exposed brains observed previously [6], [33] (Fig. 8). AMPAR signaling is governed by their synaptic localization and association with scaffolding proteins. The scaffolding proteins in turn recruit proteins that regulate actin-dependent movement of subunits to and from the synaptic membrane. Therefore, alteration in the functional state of AMPAR scaffolding proteins can result in deficits in excitatory synaptic transmission [6]. Excessive PKC activation markedly impairs prefrontal cortex-mediated cognitive function and increases distractibility [50]. The above findings therefore suggest that preventing further PKC activation such as blocking PKC cytosol-to-membrane translocation may reduce the protracted PKC-mediated deficits and restore AMPAR-regulated neurotransmission in prenatal cocaine-exposed brains. In this regard, mood stabilizers such as valproate that block PKC translocation without interfering with the enzymatic activity [51] may help attenuate prenatal cocaine-induced synaptic plasticity and dendritic structural defects leading to AMPAR-related brain dysfunction.
Figure 8. Schematic illustration of the effect of prenatal cocaine on the scaffolding and signaling molecules involved in the AMPA-GluR2 mediated synaptic long-term depression (LTD). Based on the results of this and our earlier studies (Bakshi et al., 2009), prenatal cocaine exposure enhances membrane localization and activation of PKC (especially γ and ζ isoforms) leading to hyper-phosphorylation of GRIP. The heightened GRIP phosphorylation results in increased GRASP-1 and diminished GluR2 association with GRIP (synaptic targeting) and consequent reduction and elevation of the cytosolic GRASP-1 and GluR2, respectively. By virtue of its neuronal ras-GEF activity, the increased GRASP-1 presence in the membrane of prenatal cocaine-exposed brain elevates the level of activated monomeric G proteins, RhoA, Rap1 and cdc42/Rac1 and eventual increase in actin polymer (F-actin) content. The increased F-actin level should then promote the trafficking of GluR2 in an attempt to restore the synaptic GluR2 content and LTD. The size of blocks and thickness of arrows indicates the level of each protein in the prenatal saline- and cocaine-exposed brains. The striped or dotted patterns symbolize modification made to the proteins such as increased phosphorylation of GRIP1 and reduced neuronal ras-GEF activity in GRASP-1.

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Materials and Methods

Materials and Chemicals

Soybean trypsin inhibitor, phenylmethylsulfonyl fluoride [PMSF], 2-mercaptoethanol, NaF, Na2VO4, Digitonin, protein phosphatase inhibitor I & II cocktails, recombinant γPKC, alkaline phosphatase, phorbol 12-myristate, 13-acetate (PMA), anti-phosphoserine (P3430), anti-phosphothreonine (P3555) were purchased from Sigma (St. Louis, MO). Leupeptin and aprotinin were from Peptide International (Louisville, KY). Recombinant Src, celastrene and PP1 were from Cal-Biochem (La Jolla, CA). Antibodies against GRASP-1 (SC-15568 and SC-1569), phosphoactin (SC-508), caspase-3 (SC-7272), β-actin (SC-47778), RhoA (SC-32039, SC-418), Rap1 (SC-28197), cdc42/Rac1 (SC-217) and actin (SC-1616R) were purchased from Santa Cruz Biotechnology (Santa Cruz, CA). Seize-X immunoprecipitation kit, antigen elution buffer, EZ-link Biotinylation kit and West pico chemiluminescent reagents were purchased from Pierce-Eudogen (Rockford, IL). Bradford reagent, SDS-PAGE reagents, and pre-stained molecular weight markers were purchased from Bio-Rad (Hercules, CA). 10-KDa cut-off filters were obtained from Cole-Palmer (Vernon Hills, IL). The antibody against β-tubulin (MAB3408), GST-Rhoetkin, GST-RaiGDS and GST-Pak1 were from Upstate Biotechnology/Chemicon (Temecula, CA). Target Buffer was purchased from Dako (Carpentaria, CA). Avidin-peroxidase-labeled biotin complex (ABC) was from Vector Labs (Foster City, CA). 3-3-diaminobenzidine-4 HCl (DAB)/H2O2 was from Biomeda (Foster City, CA). Cell permeable [myristoylated] pseudosubstrate inhibitors for PKCγ [GIP] and PKC/Cζ [ZIP] were custom synthesize by Peptide2 Inc. (Chantilly, VA). Guanosine 5’-diphosphate, trisodium salt [8,5-3H] ([3H]GDP, 27 Ci/nmol, PerkinElmer, Boston, MA).

Animal treatment

Pathogen-free, 10-week-old male and female Sprague-Dawley rats weighing approximately 200–215 g (Taconic, Germantown, NY) were housed individually in a 12-hr light/dark cycle with free access to food and water. All animal procedures were in compliance with the National Institutes of Health Guide for Care Use of Laboratory Animals and were approved by the City College of New York Animal Care and Use Committee. The animal treatment was performed as described previously [6]. Briefly, pregnant dams were assigned to receive daily intraperitoneal (i.p) injections from GD 8–20 of either cocaine HCl, 30 mg/kg in 0.9% saline or saline, 2 ml/kg. The animals were injected daily between 9–10 AM. Following each injection, these pregnant rats were observed for 1 hr and behavioral abnormalities recorded. There were no discernible differences in litter size (between 7–13 pups) and body weight of the pups at 21-day of age (48.9±2.5 and 50.7±2.8 g for cocaine and saline, respectively; n= 40 each) and gender distribution (23 males/17 females and 19 males/21 females for cocaine and saline groups, respectively). Importantly, the dose of cocaine used in this study did not induce seizure or fatality.

The progenies were cross-fostered to a naıve mother until sacrificed at 21 days of age (P21). They were subjected to the minimum handling associated with routine animal husbandry. Importantly, we did not find gender differences in our previous studies conducted in rabbit and rats [52], [53], [6], [35], both sexes from separate litters were employed in these experiments. Pups were sacrificed by rapid decapitation, the brains removed immediately on ice, and coronal cuts at optic chiasm and +4 mm rostral to optic chiasm were made to dissect out the frontal cortex and prefrontal cortex.

Rat cortical slice organotypic cultures

Rat brain FCX from P21 prenatal cocaine- and saline-exposed pups were chopped coronally into 200 m slices using a McIlwain chopper (Brinkman Instruments) and suspended in 10 ml of ice-cold oxygenated K-R.

The rat brain slice organotypic culture was performed with a modified procedure [54]. Rat FCX slices were transferred to sterile, porous 0.4 μm Millicell-CM insert, 2 slices per insert per
well containing 2 ml medium: 50% MEM with Earl’s salts, 2 mM L-glutamine, 25% Earl’s balanced salt solution, 6.5 g/l D-glucose, 20% fetal bovine serum (FBS), 5% horse serum, 25 mM HEPES buffer, pH 7.2, and 50 mg/ml streptomycin and 50 mg/ml penicillin. Cultures were kept in an incubator for 2 days at 36°C in 5% CO2. On the day of experiment, medium was removed, the brain slices rinsed and incubated in 0.1% FBS-containing medium for 4 hr at 36°C in 5% CO2. Brain slices were then incubated with 10 μM of cell permeable (myristoylated) pseudosubstrate inhibitors for PKCγ [GIP] and PKC/MC [ZIP] or control peptides, GIP/ZIP scrambled [GIP/ZIP SC] in fresh 0.1% FBS-containing medium for 4 hr. The effect of GIP/ZIP on the active RhoA, cdc42/Rac1 and Rap1 levels was determined as described below in the Affinity precipitation of GTP RhoA, Rap1 and Rac1/Cdc42 section.

Preparation of synaptosomes and fractionation

Synaptosomes (P2 fraction) were prepared from frontal cortices as previously described with a few modifications [54], [55], [56], [6]. To further purify synaptosomal fractions, the synaptosome-enriched P2 fraction was washed twice in 5 ml of ice-cold Kreb’s-Ringer solution (25 mM HEPES, pH 7.4; 118 mM NaCl, 4.8 mM KCl, 25 mM NaHCO3, 1.5 mM CaCl2, 1.2 mM MgSO4, 1.2 mM KH2PO4, 10 mM glucose, 100 μM ascorbic acid, 50 μg/ml leupeptin, 10 μg/ml aprotinin, 2 μg/ml soybean trypsin inhibitor, 0.04 mM PMSF, 0.1 mM 2-mercaptoethanol, 10 mM NaF, 1 mM Na2VO4 and 0.5 μl/ml protein phosphatase inhibitor I & II cocktails). To obtain cytosolic and membranous fractions of the synaptosomes, the washed synaptosomes were sonicated for 10 sec on ice in 0.5 ml hypotonic homogenization solution (25 mM HEPES, pH 7.4; 120 mM NaCl, 4.8 mM KCl, 25 mM NaHCO3, 1.5 mM CaCl2, 1.2 mM MgSO4, 1.2 mM KH2PO4, 10 mM glucose, 100 μM ascorbic acid, 50 μg/ml leupeptin, 10 μg/ml aprotinin, 2 μg/ml soybean trypsin inhibitor, 0.04 mM PMSF and 0.1 mM 2-mercaptoethanol, 10 mM NaF, 1 mM Na2VO4 and 0.5 μl/ml protein phosphatase inhibitor I & II cocktails). Samples were centrifuged at 50,000 × g for 30 min. The resulting supernatant was reserved as the cytosolic fraction and the synaptic membrane pellet was resuspended in 0.5 ml of hypotonic solution. Protein concentrations of the synaptic membranes were determined using the Bradford method before solubilization by adding 6× SDS-PAGE sample preparation buffer and boiled for 5 min. Since GRASP-1 was shown to elicit endocytosis of AMPARs [8], we measured transferrin receptor level in our synaptosomal fraction to rule out endosomal contamination.

Immunoadfinity purification of native GRASP-1, GRIP1 and GRIP2

The immunoadfinity purification was performed as described in Bakshi et al (2009) [6] with some modifications. To isolate native GRASP-1 and GRIP1, frontal cortices of P21 prenatal cocaine- or saline-exposed rats were homogenized in hypo-tonic homogenization solution described above. The obtained homogenates were solubilized using 0.5% digitonin, 0.2% sodium cholate, 0.5% NP-40 and 0.2% SDS in the presence of cocktails of protease and protein phosphatase inhibitors for 20 min at 25°C followed by 60 min at 4°C with end-over-end constant shaking. Following centrifugation to remove insoluble debris, the obtained brain lysate was diluted 5-fold and GRIP1 and GRASP-1 were individually purified using immunoadfinity columns (Seize-X immunoadfinity purification kit, Pierce-Endogen) with covalently immobilized antibodies directed against GRASP-1 and GRIP1. GRASP-1 and GRIP1 were each eluted twice with 90 μl antigen elution buffer. The resulting eluates were neutralized immediately with 20 μl 1.5 M Tris, pH 8.8 and concentrated to 100 μl by passing through 10-KDa cut-off filter. Protein concentrations were determined using the Bradford method. The purity of each protein was validated by Western blotting. In each case, the purified protein yielded a single protein band with apparent molecular weight identical to that found using rat brain lysate.

Immunoprecipitation of native actin for measurement of phosphorylated actin

To measure the level of serine-, threonine- and tyrosine-phosphorylated actin, total actin was purified by immunoprecipitation with the method described in Bakshi et al (2009) [6]. FCX of P21 from prenatal cocaine- or saline-exposed rats were homogenized in hypo-tonic homogenization solution described above. The obtained brain lysate was diluted 5-fold and protein concentration was measured by the Bradford method. Total actin in the 200 μg FCX lysate was immunoprecipitated using covalently immobilized anti-actin conjugated protein A-agarose beads overnight at 4°C. The resultant anti-actin immunoprecipitate was centrifuged, washed 3 times with 1 ml phosphate-buffered saline, solubilized by boiling in 100 μl SDS-PAGE sample preparation buffer (62.5 mM Tris-HCl, pH 6.8; 10% glycerol, 2% SDS; 5% 2-mercaptoethanol, 0.1% bromophenol blue) and used for measurement of phosphorylated actin levels by Western blotting. The obtained blots were screened for phosphoactin first (anti-phosphoserine), stripped and re-probed twice sequentially with anti-phosphotyrosine and anti-phosphothreonine. The signals were detected using a chemiluminescent method and visualized by exposure to x-ray film.

In vitro determination of GRASP-1 – GRIP1 interaction and immunoprecipitation

To control the GRIP1 phosphorylation state, native GRIP1 proteins (10 μg) purified from frontal cortices of P21 prenatal saline- and cocaine-exposed rats were incubated with 100 μg/ml alkaline phosphatase in Tris, pH 8.0, 130 mM NaCl and protease inhibitors at 30°C for 20 min (total incubation volume 100 μl) as described in Bakshi et al (2009) [6]. The phosphatase activity was terminated by adding 10 mM NaF/1 mM Na3VO4 and specific PKC- and src-mediated phosphorylation was induced by incubation with 0.5 μg/ml recombinant γPKC, 20 μg phosphatidylycerine and 100 mM PMA or 10 μg/ml recombinant Src in the presence of 30 μM ATP in Kreb’s-Ringer at 30°C for 10 min (total incubation volume 125 μl). The actions of PKC and Src were terminated by addition of 1 μM caffeine and PP1, respectively. One-half of the GRIP1 solution (containing 5 μg) was immediately solubilized by adding 6× SDS-PAGE sample preparation buffer and boiled for 5 min for analysis of phosphoactin and -threonine and -tyrosine levels by Western blotting. To determine the influence of GRIP1 phosphorylation state on the interaction between GRIP1 and GRASP-1, purified brain GRASP-1 (5 μg) from gestational saline- and cocaine-exposed rats were individually added to 5 μg of GRIP1 with different phosphorylation states and incubated in 100 μg/ml brain phospholipids, 1% BSA-containing Kreb’s-Ringer at 30°C for 30 min with constant end-over-end shaking. The GRIP1-associated GRASP-1 was isolated along with GRIP1 by 20 μl immobilized anti-GRIP1 conjugated protein A-agarose beads and measured using Western blot with anti-GluR2. The obtained blots were screened for phosphoactin first (anti-phosphoserine), stripped and re-probed twice sequentially with anti-phosphotyrosine and anti-GRIP1. The signals were detected using a
chemiluminescent method and visualized by exposure to x-ray film.

Western blotting
To determine cellular distribution or the interaction between GRASP-1 and GRIP1, cytosolic and membranous fractions of frontal cortices or anti-GRIP1 immunoprecipitates were boiled for 5 minutes in 100 μl SDS-PAGE sample preparation buffer and then size fractionated on 7.5 or 10% SDS-PAGE based on the molecular mass of the protein. Proteins were electrophoretically transferred to nitrocellulose membrane and Western blotting was performed with antibodies for GRASP-1, phosphotyrosine, phosphoserine, phosphothreonine. The blots were stripped and re-probed with anti-GRIP1 or capase-3 to assess the level of sample loading.

To determine the expression level of GRASP-1 and GRIP1, protein extracts of the synaptosome-enriched P2 fractions (50 μg) were size fractionated on 7.5% SDS-PAGE and Western blotting was performed using specific antibodies. In some cases, the blots were stripped and re-probed with anti-β-actin, β-tubulin and GRIP1.

Immunoreactivity was detected by reacting with chemiluminescent reagents for exactly 5 min and visualized by immediately exposing to X-ray film for 10–30 sec. Specific protein bands were quantified by densitometric scanning (GS-800 calibrated densitometer, Bio-Rad Laboratories).

Immunohistochemistry
Immunohistochemical analyses were performed using antibodies directed against GRASP-1 were carried out on paraffin-embedded tissues as described previously [57]. Briefly, after removal of paraffin with xylene and rehydration through a graded series of decreasing concentrations of ethanol, protein antigenicity was enhanced by microwaving sections in Target Buffer for 2 min. Following 30-min incubation in 0.3% H2O2, sections were treated for 30 min in normal blocking serum and then incubated with primary antibodies at appropriate dilutions for 1 hr at room temperature. Following a thorough rinse in PBS, a secondary biotin-labeled antibody was applied for 30 min. Immunoreactions were treated with the avidin-peroxidase-labeled biotin complex (ABC) and visualized by treatment of sections with 3,3'-diaminobenzidine-4 HCl (DAB)/H2O2. Sections were lightly counterstained with hematoxylin, dehydrated through a graded series of increasing concentrations of ethanol, cleared in xylene and mounted in Permount. Controls consisted of comparable sections treated with non-immune serum, pre-absorbed antibody or omission of the primary antibody. Specimens were examined and photographed with a Nikon FXA microscope, and digital images were recorded using a Nikon DXM1200F digital camera and processed using Image Pro Plus (Phase 3 Imaging, Glen Mills, PA) imaging software.

GDP Dissociation Assay to assess GRASP-1 rasGEF activity
GRASP-1 rasGEF activity in the FCX of P21 in utero cocaine- and saline- exposed rats was measured using purified GRASP-1, by the method described by Ye et al. (2000) with some modifications [7]. To load RhoA or Rap1 with [3H]GDP, 0.5 μg immunopurified RhoA or Rap1 (linked to covalently immobilized agarose-protein A, PIERCE) was incubated with 10 μCi [3H]GDP in 50 μl nucleotide loading buffer (50 mM Tris with 10 mM EDTA, 5 mM MgCl2, 1 mM DTT, and 1 mg/ml BSA) at 37°C. Twenty minutes later, 60 μl of nucleotide loading stopping buffer (50 mM Tris-HCl [pH 7.4], with 5 mM MgCl2, 1 mM DTT and 1 mg/ml BSA) was added. The 110 μl sample was divided into two, and each sample was added to 500 μl dissociation reaction buffer (25 mM Tris-HCl [pH 7.4] with 2 mM MgCl2, 1 mM DTT, 1 mg/ml BSA, and 0.1 mM GDP) containing 5 μg of immunopurified GRASP-1 from frontal cortical synaptosomes of prenatally cocaine- and saline-treated rats, respectively. The two reactions were incubated at 25°C. A sample was then removed from each mixture every 30 sec up to 5 min and mixed with 200 μl ice-cold dissociation reaction stopping buffer (50 mM Tris-HCl [pH 7.4] with 10 mM MgCl2). Rap1 filtration was performed using GF/C filters under vacuum. The filter was washed twice with 5 ml ice-cold stopping solution, and the radioactivity was measured by liquid scintillation counting after air drying. The data were expressed as the percent of [3H]GDP bound at each time point comparing to input [3H]GDP level (at time 0). The specific activity of the RhoA and Rap1 was calculated using Prism. Data points are means and vertical bars are the s.e.m. of three independent experiments in each treatment group.

Affinity precipitation of GTP RhoA, Rap1 and Rac1/Cdc42
GTP-bound RhoA, Rap1 and Rac1/Cdc42 were affinity-purified from the synaptosome-enriched fraction of the FCX region from prenatal cocaine- and saline-treated rats. Precipitation of active RhoA was performed using the fusion protein GST-Rhotekin, which specifically recognizes the active GTP-bound form of RhoA. Similarly, GTP-Rap1 was precipitated using fusion protein GST-RalGDS and Rac1/Cdc42 from GST-Pak1, respectively. Synaptosomes were ruptured by sonication in 0.25 ml of immunoprecipitation buffer and solubilized using 0.5% digitonin, 0.2% sodium cholate and 0.5% NP-50 at 4°C for 1 hr. Following dilution with 0.75 ml of immunoprecipitation buffer and centrifugation, the GTP-bound RhoA, Rap1 and cdc42/Rac1 in the resultant synaptosomal lysates were purified by incubating at 4°C for 1 hour with Rhotekin-RBD beads (10 μg, Upstate Technologies), GST-RalGDS (10 μg, Upstate Technologies) and GST-Pak1 beads (10 μg, Upstate Technologies), respectively, with end-over-end rotation. Following centrifugation, the beads were washed twice with 1 ml of ice-cold 25 mM Tris-HCl, pH 7.4, 20 mM MgCl2 containing protease and protein phosphatase inhibitors. To determine the levels of GST-RhoA, GST-Rac1/Cdc42, and GST-Rap1 the proteins were size-fractionated on 12% SDS-PAGE and Western blot analysis was performed using antibodies specific to RhoA, Rac-1/Cdc42, and Rap1 (Santa Cruz). Signals were detected using a chemiluminescent method (Pierce) and visualized by exposing to x-ray film.

Determination of F-actin levels
To measure the level of F-actin, synaptosomes were prepared from the FCX of prenatal cocaine- and saline-treated P21 rats by the method described above. Synaptosomes (500 μg) were treated with vehicle, alkaline phosphatases (1 mg/ml) or alkaline phosphatases (1 mg/ml) plus phosphatase inhibitors (10 mM NaF, 1 mM Na3VO4) or cytochalasin D (10 μM) in vitro for 20 min at 30°C in 50 mM Tris HCl, pH 8.0, 100 mM NaCl, 1 mM MgCl2, 1 mM CaCl2, protease inhibitors (total incubation volume: 100 μl). Following termination of the reaction by diluting with 300 μl immunoprecipitation buffer, the synaptosomes were solubilized in 0.5% digitonin, 0.2% sodium cholate, and 0.5% NP-40. Actin-containing proteins were extracted by incubating with 2 μg of biotin-conjugated anti-actin antibodies for 1 hr at 4°C and the anti-actin-linked actin proteins were then immobilized by loading 50% of the reaction mixture into each well of the streptavidin-coated 96-well plate (Pierce) and incubating at 4°C for 1 hr. The solution was then removed and the plate was washed twice with 25 mM Tris HCl, pH 7.4, 100 mM NaCl (200 μl/
well). The level of F-actin was determined using rhodamine-conjugated phalloidin (0.5 μM/ml) (Molecular Probes/Invitrogen). After two washes with 100 μl 25 mM Tris, [pH 7.4] containing 100 mM NaCl, the fluorescence intensity of phalloidin was measured using Beckman multimode plate reader, DX890.

Data Analysis and statistical evaluation
Statistical differences between cocaine and saline groups were assessed using the two-tailed Student’s t test. Differences between

in vitro dose-response relations were analyzed by ANOVA followed by Newman-Keuls multiple comparisons.

Author Contributions
Conceived and designed the experiments: H-YW RN EF. Performed the experiments: KB MK. Analyzed the data: H-YW KB. Wrote the paper: H-YW KB EF.

References