cGMP-dependent protein kinase Iα associates with the antidepressant-sensitive serotonin transporter and dictates rapid modulation of serotonin uptake

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Abstract

Background: The Na+/Cl−-dependent serotonin (5-hydroxytryptamine, 5-HT) transporter (SERT) is a critical element in neuronal 5-HT signaling, being responsible for the efficient elimination of 5-HT after release. SERTs are not only targets for exogenous addictive and therapeutic agents but also can be modulated by endogenous, receptor-linked signaling pathways. We have shown that neuronal A3 adenosine receptor activation leads to enhanced presynaptic 5-HT transport in vitro and an increased rate of SERT-mediated 5-HT clearance in vivo. SERT stimulation by A3 adenosine receptors derives from an elevation of cGMP and subsequent activation of both cGMP-dependent protein kinase (PKG) and p38 mitogen-activated protein kinase. PKG activators such as 8-Br-cGMP are known to lead to transporter phosphorylation, though how this modification supports SERT regulation is unclear.

Results: In this report, we explore the kinase isoform specificity underlying the rapid stimulation of SERT activity by PKG activators. Using immortalized, rat serotonergic raphe neurons (RN46A) previously shown to support 8-Br-cGMP stimulation of SERT surface trafficking, we document expression of PKGI, and to a lower extent, PKGII. Quantitative analysis of staining profiles using permeabilized or nonpermeabilized conditions reveals that SERT colocalizes with PKGI in both intracellular and cell surface domains of RN46A cell bodies, and exhibits a more restricted, intracellular pattern of colocalization in neuritic processes. In the same cells, SERT demonstrates a lack of colocalization with PKGII in either intracellular or surface membranes. In keeping with the
ability of the membrane permeant kinase inhibitor DT-2 to block 8-Br-cGMP stimulation of SERT, we found that DT-2 treatment eliminated cGMP-dependent kinase activity in PKGI-immunoreactive extracts resolved by liquid chromatography. Similarly, treatment of SERT-transfected HeLa cells with small interfering RNAs targeting endogenous PKGI eliminated 8-Br-cGMP-induced regulation of SERT activity. Co-immunoprecipitation studies show that, in transporter/kinase co-transfected cells, PKG\(\alpha\) specifically associates with hSERT.

**Conclusion:** Our findings provide evidence of a physical and compartmentalized association between SERT and PKG\(\alpha\) that supports rapid, 8-Br-cGMP-induced regulation of SERT. We discuss a model wherein SERT-associated PKG\(\alpha\) supports sequentially the mobilization of intracellular transporter-containing vesicles, leading to enhanced surface expression, and the production of catalytic-modulatory SERT phosphorylation, leading to a maximal enhancement of 5-HT clearance capacity.

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**Background**

Signaling by the neurotransmitter serotonin (5-hydroxytryptamine, 5-HT) plays a critical, modulatory role in brain pathways supporting mood, appetite, sexual behavior, and reward. The inactivation of synaptic 5-HT is achieved largely through the actions of presynaptic serotonin transporters (SERT, SLC6A4). SERTs also participate in presynaptic 5-HT recycling to sustain neuronal 5-HT stores: this is most evident in studies with SERT KO mice that display a 60–80% reduction in brain 5-HT level [1]. These mice also display profound changes in 5-HT receptor coupling and a loss of psychostimulant (MDMA, "ecstasy") sensitivity [1-5], underscoring the importance of SERT for 5-HT signaling and drug response.

Increasing evidence indicates that SERT-mediated 5-HT clearance is controlled by multiple regulatory pathways that dictate both SERT plasma membrane expression and catalytic activity (reviewed in [6]). By analogy with the structure of the recently crystallized, SLC6 homolog LeuT\(\alpha\) [7], SERTs are modeled to possess twelve transmembrane domains [8] with intracellular amino and carboxy termini. These latter domains possess multiple canonical Ser/Thr phosphorylation sites and are known to interact directly with other proteins, including Hic-5, neuronal nitric oxide synthase (nNOS), as well as heteromeric integrins that contain a β3 subunit [9-11]. Further specification and localization of the key molecules involved in SERT regulation, clarification of whether they form stable or transient associations with SERT, and evaluation of how loss of regulation contributes to SERT dysfunction may offer important insights into 5-HT-linked brain disorders such as depression, autism, and obsessive-compulsive disorder (OCD) [12-15]. In this regard, we and others have shown that multiple human SERT coding variants display compromised regulation through protein kinase C (PKC)-, cGMP-dependent protein kinase (PKG)-, and p38 mitogen-activated protein kinase (p38 MAPK)-linked pathways [16-19]. Among these coding variants are mutants found to associate with autism and OCD [20,21].

A detailed analysis of the signaling pathways that regulate SERT may thus make important contributions to our understanding of multiple neuropsychiatric disorders and offer new strategies for intervention.

Our previous work has elucidated the actions of multiple G-protein coupled receptors that can regulate SERT negatively [e.g. α2 adrenergic receptors [22]] or positively (e.g. A3 adenosine receptors [A3ARs, [23,24]). A3ARs enhance SERT trafficking and 5-HT clearance capacity as a consequence of elevations in intracellular Ca++ and cGMP and PKG activation [23,24]. We have shown that PKG-dependent regulation of SERT by A3ARs is present both in cultured RBL-2H3 mast cells [23] and in serotonergic neuronal terminals monitored ex vivo or in vivo [24]. Consistent with these findings, SERTs are phosphorylated by treatment of cells or neurons with the general PKG activators 8-pCPT-cGMP and 8-Br-cGMP [17,25]. Two genes encoding homologous PKG proteins have been identified, PKGI and PKGII (also identified as cGK1 and 2, [26]), both of which can be stimulated by membrane-permeant cGMP analogs. PKGI is a soluble, cytosolic protein that is abundant in smooth muscle, platelets, and brain [27,28]. Mice lacking PKGI have profound smooth muscle deficits that lead to eventual death [29]. These deficits can be rescued in constitutive PKGI KO mice by smooth muscle-specific rescue using either of the two PKGI amino-terminal splice variants PKGI\(\alpha\) or PKGI\(\beta\) [30]. PKGII is a myristoylated, membrane-associated protein that is most abundant in intestine, brain, and kidney [31,32]. PKGII knockout mice exhibit dwarfism and defects in intestinal secretion [33]. Which of the two PKG genes participate in SERT regulation, or whether both contribute, is presently unknown though critical for further mechanistic studies that pursue whole animal genetic or pharmacological manipulations.

Neuroanatomical studies are lacking that document the precise localization in brain of PKG isoforms in relation to SERT expression, likely due to the insensitivity of available...
 PKGI stimulation, regulation that arises from elevated trans-
 siRNAs completely eliminate 8-Br-cGMP-triggered SERT 
ular trafficking. Consistent with this idea, PKGI-targeted 
dominant role of PKGI in regulating of transporter vesic-
 tion is most evident when labeling of the transporter is 
immunofluorescence approaches. SERT/PKGI colocaliza-
strate expression of DT-2 sensitive, cGMP-dependent 
kinase activity that comigrates with PKGI immunoreactive 
protein in RN46A cells, where we also document colocal-
ization of SERT with PKGI, but not PKGII, by quantitative 
immunofluorescence studies. SERT/PKGI colocalization 
is most evident when labeling of the transporter is 
achieved in permeabilized cells, consistent with a pre-
localization studies, RN46A cells were probed with a SERT 
secondary antibodies alone and in the case of PKGI, when 
the presence of PKGI labeling, but positive 
for SERT, were clearly evident at the tips of processes (Fig. 
1C, right panel, open arrowhead). With P conditions, 
PKGI/SERT colabeling was more extensive than under NP 
conditions, filling both the cell soma as well as processes 
(Fig. 1D, right panel). In keeping with a lack of detection 
of PKGII in the RN46A cells, low signal for PKGII was evi-
dent by immunofluorescence (Fig. 1E, middle panel) and 
was detected as a nuclear pattern surrounded by diffuse 
fluorescence labeling. SERT/PKGII colocalization evalua-
tion in double labeling preparations (Fig. 1E, right panel) 
was quite distinct from that observed with PKGI, and 
showed no evidence for overlap in patterns of expression.

To assess PKG and SERT colocalization using quantitative 
approaches, we calculated the Intensity Colocalization 
Quotient (ICQ) under either NP or P conditions for both 
cell bodies and processes. As described by Li and col-
leagues [35], ICQ values represent a measure of the extent 
of correlation of intensity values in space for 2 separate 
fluorophores, where values range between -0.5 and 0.5 for 
negatively and positively correlated signals, respectively. 
As shown in Fig. 2, PKGI/SERT ICQ values fall signifi-
cantly above 0 under some conditions, whereas PKGII/
SERT analyses yield no ICQ values different from 0 under 
any condition, suggesting a random relationship of stain-
ing. For PKGII/SERT under NP conditions, significant colo-
calization is evident only in the cell body area. Under P 
conditions, significant colocalization of PKGII/SERT is evi-
dent for both the cell soma and processes, in keeping with 
the qualitative evaluations noted above. These findings 
provide quantitative evidence of PKGII/SERT colocaliza-
tion (our PKGII antibody cannot distinguish PKGIα and β) 
and reveal that PKGI/SERT complexes are largely intracel-
ular in distal processes.

**Results**

**Expression and colocalization of PKGI with SERT in 
RN46A cells**

To initiate studies of PKG gene products engaged in neu-nal SERT regulation, we explored the expression of PKG 
isoforms in RN46A cells, a serotonergic cell line that we 
have previously found to display 8-Br-cGMP-stimulated 
SERT activity. Initially, we performed qualitative RT-PCR 
analysis on total RN46A RNA using oligonucleotide prim-
ers specific for PKGIα, PKGIβ and PKGII. Compared to rat 
midbrain RNA, where we readily identified expression of all 
three gene products, RN46A RNA only yielded amplifi-
cation of PKGIα and PKGIβ (Fig. 1A). Next, we pursued 
immunofluorescence studies of RN66A cells, where we 
confirmed readily detectable expression of PKGI and SERT 
(Fig. 1B–D) and low expression of PKGII (Fig. 1E). Staining 
with PKGI or PKGII primary antibodies was elimi-
nated with preabsorption against their respective antigens 
(Fig. 1B, middle panel; also see Additional File 1). Staining 
was also absent in cells treated with fluorescent second-
ary antibodies alone and in the case of PKGI, when 
cells were not permeabilized (data not shown). For SERT 
localization studies, RN46A cells were probed with a SERT 
 ectodomain antibody under two conditions. One set of 
assays involved probing for SERT prior to permeabiliza-
tion (NP, Fig. 1), followed by permeabilization of cells to 
detect cytoplasmic PKG. A second evaluation involved use 
of both SERT and PKG antibodies under permeabilized 
(P, Fig. 1) conditions. P cells displayed a broader cellular 
patern of SERT staining than nonpermeabilized cells 
(compare Fig. 1B, left and right panels), consistent with 
detection of a pool of surface SERTs in NP cells. Dual-
labeling experiments under these conditions revealed evi-
dence of significant colocalization of PKGI with SERT 
under NP conditions, localized primarily to RN46A cell 
membranes in vivo can be antagonized by the peptide 
DT-2, a membrane-permeant inhibitor proposed to block 
selectively PKGI [24,34]. In the present work, we demon-
strate expression of DT-2 sensitive, cGMP-dependent 
kinase activity that comigrates with PKGI immunoreactive 
protein in RN46A cells, where we also document colocal-
immunofluorescence approaches. SERT/PKGI colocalization 
is most evident when labeling of the transporter is 
achieved in permeabilized cells, consistent with a pre-
odominant role of PKGI in regulating of transporter vesic-
ular trafficking. Consistent with this idea, PKGI-targeted 
siRNAs completely eliminate 8-Br-cGMP-triggered SERT 
stimulation, regulation that arises from elevated trans-
surface expression surface. Co-immunoprecipitation stud-
ies reveal that SERT forms a stable complex with PKGI, 
with a greater efficiency of association achieved with 
PKGIα over PKGIβ. We discuss the impact of our findings 
with respect to a model for how intracellular SERT/PKGI 
associations support both trafficking-dependent and -independent modes of SERT regulation.

**DT-2 sensitive PKGI activity in RN46A cells**

DT-2 is a membrane permeant peptide that was developed as 
a PKGI inhibitor [34] and has been previously demonstrated 
to eliminate SERT stimulation following A3AR stimulation 
or exogenous 8-Br-cGMP treatments [24,36]. Prior studies of 
kine specificity in vitro using purified kinases, indicate a Ki 
of 12.5 ± 3.0 nM for purified PKGI (either PKGIα or PKGIβ) 
versus 75.5 ± 18 μM for PKA [34]. We extended these find-
ings to PKGII and PKCα, yielding Ki values of 9.10 ± 1.3 μM 
and 26.6 ± 4.0 nM, respectively. Although DT-2 can inhibit 
PKCα at concentrations comparable to PKGI inhibition,
Colocalization of SERT and PKG1 in RN46A cells. A) Total RNA from RN46A cells or rat midbrain was isolated and subjected to RT-PCR as described in Methods. Figure shows agarose gel analysis of cDNA products, revealing expression of PKGI but very little PKGII. B-E) Double immunocytochemical labeling of SERT, PKGI, and PKGII. RN46A cells were fixed, stained, and imaged by confocal microscopy as described in Methods. SERT staining was achieved with an ectodomain antibody (Advanced Targeting AB-N09). NP, nonpermeabilized. P, permeabilized by 0.2% NP40. NP/B, nonpermeabilized with antibody preabsorption by peptide (Advanced Targeting PR-03). DIC image is inset in B, middle, to demonstrate presence of cells. Inset image in D, right, shows staining in process. Closed arrows indicate perinuclear staining while open arrows indicate process staining. Scale bars represent 10 μm. Data are representative of at least three independent experiments.
PKC activation leads to SERT internalization rather than a movement to the surface and an enhancement of 5-HT uptake. Regardless, our findings add support to the contention that the DT-2 sensitivity of cGMP-dependent SERT regulation is evidence of PKGI versus PKGII involvement. To explore this issue further in the RN46A model, we fractionated RN46A cell lysates via DEAE-Sephadex chromatography, testing fractions for levels of PKGI-immunoreactive protein as well as levels of cGMP-dependent phosphorylation of the heptapeptide substrate (RRKSRAE). In Fig. 3A, we display a representative analysis, plotting both PKGI protein level and kinase activity from fractions where significant cGMP-dependent kinase activity was detected. Activity in our kinase assays does not necessarily derive from PKG activity as other kinases, notably PKA, can utilize cGMP for activation [37]. Thus, in Fig. 3B, we analyze the sensitivity of cGMP-stimulated kinase activity to DT-2. The fractions bearing the most significant level of PKGI protein (fractions 13–17; 93.9% of total PKGI immunoreactivity) exhibited a significantly greater degree of DT-2 sensitivity as compared to fractions where PKGI immunoreactivity was low or absent (fractions 18–22; 6.1%). Indeed, in fractions 13–17, all of the cGMP-stimulated kinase activity (Fig. 3B, dotted line) was eliminated whereas we detected DT-2 insensitive kinase activity in fractions 18–22. Together, these studies demonstrate the presence of DT-2-sensitive PKGI in RN46A cells and support the likelihood that DT-2 blockade of cGMP-dependent SERT stimulation in intact cells arises from antagonism of PKGI.

**PKGI is required for 8-Br-cGMP-mediated increases in SERT activity in transfected cells**

We have shown that SERT activity can be stimulated by 8-Br-cGMP in multiple heterologous expression systems, including transfected HeLa cells [17,20,36]. These observations allowed us to implement transient transfection of PKGI-directed siRNAs to assess whether evidence, independent of DT-2, could be gathered that PKGI is essential for 8-Br-cGMP triggered SERT stimulation. Mock or siRNA-treated HeLa cells (60 hrs) were subjected to either vehicle or 8-Br-cGMP treatment (100 μM, 10 min) followed by [3H]5-HT transport assays (20 nM, 10 min), as described in Methods. As shown in Fig. 4, PKGI siRNA treatments had no significant effect on basal SERT activity in either mock or siRNA transfections. Additionally, mock-transfected cells demonstrated the previously documented stimulation of SERT activity following 8-Br-cGMP treatment [17]. In contrast, cells treated with PKGI siRNA exhibited no 8-Br-cGMP-induced SERT stimulation.

**hSERT forms a stable association with PKGIα**

Although our studies above implicate PKGI in SERT regulation, they do not address whether PKGI/SERT interactions are transient or stable, nor do they identify which isoform of PKGI is most likely involved in SERT regulation since our PKGI antibodies are not isoform specific. As SERT protein levels were too low in RN46A cells to permit co-immunoprecipitation of SERT/PKGI complexes, we performed transient co-transfections of HEK-293T cells using His-tagged hSERT and HA-tagged PKGI isoforms. HA-PKGI complexes were isolated via anti-HA beads followed by immunoblotting of total and co-immunoprecipitated extracts using an antibody against SERT. Anti-HA blots of total cell lysates demonstrated equivalent expression of HA-PKGIα and HA-PKGIβ (Fig. 5A, arrow). Anti-SERT blots revealed equivalent expression of hSERT (Fig. 5B, arrow). We did note a broader, higher Mr smear of SERT-immunoreactive species in lanes cotransfected with either PKG isoform. The origin of this altered SERT migration is as yet unclear, but could reflect a capacity of either isoform to trigger SERT phosphorylation or lead indirectly to the stabilization of multimeric, detergent-resistant SERT complexes. However, only anti-HA bead eluates of PKGIα revealed the presence of hSERT (Fig. 5C). These studies provide evidence that stable complexes of hSERT with PKGI are formed in transporter/kinase transfected cells and uncover an unexpected specificity for the PKGIα isoform in this model.

**Discussion**

Increasingly, we appreciate that biogenic amine transporters typified by SERT are under complex regulatory control mediated by associated proteins and multiple signaling pathways [9-11,17,18,22,23,25,36,38-50]. Understanding the molecules and contingencies involved in SERT regulation is likely to enhance our abilities to manipulate SERT for therapeutic ends, as well as expand our understanding as to how SERT dysregulation contributes to mental illness [16,17,20,21]. Miller and Hoffman first identified a cGMP-linked pathway promoting rapid enhancement of SERT activity following adenosine recep-
tor activation [44]. Thus, 8-Br-cGMP treatments recapitulate the actions of nonselective adenosine receptor agonists on RBL-2H3 cells (e.g. 5'-N-ethylcarboxamideadenosine, NECA) and these authors hypothesized that cGMP-stimulated protein kinases might direct phosphorylation of SERT or a SERT-associated protein to trigger increased SERT activity. We now understand that A3ARs and/or 8-Br-cGMP treatments can increase SERT activity in multiple contexts, including the rat raphe cell line RN46A [36], transfected CHO and HeLa cells [17,18], rat midbrain and cortical synaptosomes in vitro and the mouse hippocampus in vivo [47]. In all cases, pharmacological inhibition of PKG blocks stimulation of SERT activity. Furthermore, biochemical and surface expression studies reveal that PKG activators lead to both SERT phosphorylation [17,18,47] and enhanced SERT insertion into the plasma membrane [17,19,23], providing an opportunity for steady-state increases in 5-HT uptake capacity. SERT trafficking regulation leads to p38 MAPK activation that we have shown shifts the transporter to a higher affinity state for 5-HT recognition [36]. Progress has also been achieved in identifying a number of SERT-interacting proteins [9-11,43,49] whose associations can impact both SERT trafficking and transport rates. In this report, we provide evidence that PKGIIα is responsible for rapid, 8-Br-cGMP-dependent, DT-2-sensitive SERT stimulation. Moreover, we demonstrate that mammalian cells can organize a stable association of PKGIIα with SERT, an interaction likely to support the enhanced surface trafficking of the transporter evident after treatment of cells with PKG activators.

In RN46A cells, we established using qualitative and quantitative approaches that endogenous PKGII and SERT colocalize. Although PKGII could not be detected by RT-PCR, low levels of PKGII immunoreactivity were evident via confocal-assisted immunofluorescence studies. However, no evidence for SERT/PKGII colocalization was evident. Interestingly, SERT/PKGII colocalization was nonuniform, being most prominent in permeabilized preparations and absent in the distal segments of nonpermeabilized cells. One possible explanation for these findings is that SERT recruits PKGII to vesicles as part of a coordinated mechanism to achieve regulated plasma membrane trafficking and that, in distal processes, SERT moves to the plasma membrane after dissociation of PKGII. Extensive studies to date in our lab have failed to document consistent interactions between PKGII and SERT (Steiner and Blakely, unpublished findings), indicating a role for an intermediate in formation of SERT/PKGII complexes. This intermediate could result in a SERT (or PKG) post-translational modification or represent a SERT-associated protein lacking under the conditions of in vitro incubations of purified proteins. Additional studies are clearly needed to elaborate the manner by which transporter/kinase associations are formed. Future studies should also explore dynamic features of the SERT/PKGII association, such as whether these interactions change upon vesicle membrane fusion and localization of SERT to plasma membrane subdomains.

PKG activation has been found to alter actin dynamics and vesicular trafficking [51,52] as well as promote phosphorylation of a single Thr (Thr276) on SERT whose

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**Figure 3**

**DT-2 sensitive, PKGII activity in RN46A cells.** A) DEAE chromatography and phosphorylation assays of RN46A cell extracts were performed as described in Methods. Chromatogram displays cGMP-dependent (background-subtracted) kinase activity and quantitative protein measurement. cGMP-independent kinase activity was measured as 27.3% of total kinase activity. Representative immunoblot of fractions with detectable kinase activity is shown below chromatogram. B) Evaluation of DT-2 sensitivity in fractions containing detectable (13–17) or nondetectable (18–22) PKGII protein by immunoblot. Both fraction bins 13–17 and 18–22 displayed DT-2-sensitive kinase activity (***, p < 0.001; **, p < 0.01), with fractions overlaying PKGII protein being significantly more sensitive to DT-2 (****, p < 0.001). Dashed line represents cGMP-independent kinase activity equivalent to 27.3%. Data shown are from a single experiment, are representative of three independent experiments, and are presented as means ± SEM. Statistical significance was calculated via two-way ANOVA, followed by Bonferroni’s post-test.
Gly56Ala is a catalytically activated variant [17,19] that
also displays diminished sensitivity to the PP2A specific
inhibitor fostriecin [19]. Thus, it is possible that the SERT/
PKGIα associations documented here are mutually exclu-
sive with, or are modulated by SERT/PP2A interactions
and that these two enzymes govern phosphorylation sites
linked to SERT trafficking, as well as catalytic modulation.
Interestingly, p38 MAPK is known to phosphorylate and
activate PP2A [55], providing a path by which PKGI acti-
vation could lead to both p38 MAPK-dependent trans-
porter catalytic activation and efficient
dephosphorylation of trafficking regulatory SERT sites. A
mirror image of this idea related to SERT inactivation and
internalization has been proposed involving PKC and
PP2A by our lab [40,43] and the work of Ramamoorthy's
group [41]. Future studies that elucidate the sites targeted
by PP2A and PKC should greatly clarify these ideas. Since
OCD and autism-associated SERT variants appear to
impact PKG, p38 MAPK and PP2A-dependent transporter
regulation [18], this effort may also have significant
impact on our understanding of the origins of 5-HT linked
brain disorders.

We have recently proposed by analogy to the classical pic-
ture of insulin-dependent glucose transporter regulation
[56] that two modes of SERT trafficking to the plasma
membrane exist: one, a finite capacity, regulated pathway
overseen by PKG, p38 MAPK, PP2A and PKC, and a sec-
ond, constitutive pathway that is relatively insensitive to
these proteins but which can be more greatly populated
by heterologous overexpression. RN46A cells endog-
ously express a low level of SERT and display substan-
tial elevations (≥200%) in SERT activity following 8-Br-
cGMP treatment or p38 MAPK stimulation [36]. High-
level heterologous expression of SERT in transfected cells,
in our hands, leads to SERT proteins that demonstrate
greatly reduced, and in some cases, no cGMP-dependent
regulation. Due to this characteristic, we routinely con-
strain SERT expression in transfected cells where SERT reg-
ulation is being assessed, including the HeLa model we
have used in this study. Our goal is to approach the low
level of SERT expression observed in RN46A and RBL-2H3
cells. Unfortunately, such low expression levels make bio-
chemical studies, such as SERT/PKGI co-immunoprecipi-
tation experiments, extremely challenging, if not
impossible. Although we have utilized the higher express-
ing HEK-293T model for our co-immunoprecipitation
studies, our goal here was not to parallel functional stud-
ies, but rather to determine whether a portion of PKGI can
assemble in a SERT complex and whether PKGI isoform
specificity is evident. Additionally, we expect the pool of
PKGI that associates with SERT to be small, and likely dif-
ferentially localized, compared to the total PKGI pool.
This possibility likely explains why we do not observe ICQ
values for SERT and PKGI approaching 0.5, the theoretical
maximum, even in permeabilized cells. Our blots of

![Figure 4](http://www.molecularbrain.com/content/2/1/26)

**Figure 4**

**Treatment of transfected HeLa cells with PKGI siRNA prevents 8-Br-cGMP-induced SERT stimulation.** Cells were transfected with hSERT and siRNA directed against human PKGI and assayed for 8-Br-cGMP stimulation of SERT activity as described in Methods. Upon 8-Br-cGMP stimulation, cells expressing hSERT (closed bar) demonstrate elevated 5-HT transport activity (147 ± 3.56%) absent from cells treated with PKGI siRNA (dotted bar, 96.4 ± 6.96%). Cells treated with siRNA in the absence of 8-Br-cGMP (checkered bar) show no change in 5-HT uptake (94.5 ± 1.88%) from control (open bar, 100 ± 0.010%). Each assay was repeated three times. Values are means ± SEM. ***, p < 0.001 versus control, calculated via one-way ANOVA followed by Tukey's post-test.
siRNA-treated HeLa extracts demonstrate only a modest reduction in total PKGI level [see Additional file 2] despite full elimination of 8-Br-cGMP triggered SERT stimulation. We also observed a full elimination of p38 MAPK-dependent regulation of SERT after only a modest reduction in p38α MAPK level by p38α MAPK-targeted siRNA treatment [36]. Since siRNAs are cotransfected with hSERT to evaluate transporter regulation, we thought it important to determine whether the full inhibition of regulation could be due to full suppression of kinase expression, though limited to the small percentage of cells transfected. Indeed, when we determined the percentage of cells transfected with siRNA using toxic siRNAs [see Additional file 2], we found a good match to the percent reduction in PKGI protein (~15% in both cases).

**Conclusion**

We provide evidence that native SERT and PKGI are coexpressed in rat serotonergic RN46A cells where the two proteins colocalize, particularly when intracellular transporter pools are visualized in situ using permeabilized cells. In these cells, we corroborate the utility of the membrane-permeant peptide inhibitor DT-2 to eliminate cGMP-stimulated kinase activity, sensitivity that co-migrates with immunoreactive PKGI following ion exchange chromatography. Using PKGI siRNA, we establish a critical role for endogenous PKGI in 8-Br-cGMP triggered SERT regulation. We demonstrate that stable SERT/ PKGIα complexes can be isolated from kinase/transfected or co-transfected cells, revealing an unexpected isoform specificity that will propel future studies to evaluate how the alternatively spliced amino-terminal sequences that distinguish between PKGIα and β support PKGIα associations. Our studies thus illuminate a key facet of the molecular network that underlies rapid regulation of SERT via cell signaling pathways and compel further investigation of the mechanisms by which PKGIα/SERT associations are modulated by antidepressants, drugs of abuse, and disease-associated SERT mutations.

**Methods**

**Reagents**

Most chemicals were purchased from Sigma-Aldrich (St. Louis, MO). Culture medium (Dulbecco’s modified Eagle’s medium and Ham’s F-12) was prepared by the Vanderbilt Media Core using Invitrogen (Invitrogen Corporation, Carlsbad, CA) reagents. Trypsin-EDTA, glutamine, and penicillin/streptomycin were also purchased from Invitrogen. Fetal bovine serum was obtained from Atlanta Biologicals (Norcross, GA). Anti-HA High Affinity Matrix beads and rat anti-HA monoclonal antibody were purchased from Roche (Basel, Switzerland). H8 hydrochloride was obtained from Cayman Chemical (Ann Arbor, MI). 5-hydroxy [3H]tryptamine trifluoroacetate ([3H]5-HT, 76 Ci/mmol) was purchased from Amersham Biosciences Inc. (Piscataway, NJ). Rabbit polyclonal anti-PKGI antibodies were a generous gift from Dr. Robert Feil (University of Tübingen, Tübingen, Germany) or were purchased from Assay Design, Inc. (Ann Arbor, MI). Rabbit polyclonal antibody directed against PKGII was kindly provided by Dr. Hugo De Jonge (Erasmus University, The Netherlands). Mouse SERT monoclonal antibody was purchased from Roche (Basel, Switzerland).

**Cell culture and transfection**

RN46A cells (a generous gift from Dr. Scott Whittemore, University of Louisville, Louisville, KY) were maintained in DMEM/F-12 (1:1 in volume) containing 250 mg/L G418, 10% FBS, 2 m L-glutamine, 0.1 U/mL penicillin-0.11 g/mL streptomycin. HEK-293 T cells and HeLa cells were cultured in DMEM, 10% FBS, 2 m L-glutamine, 0.1 U/mL penicillin-0.11 g/mL streptomycin. For co-

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**Figure 5**

hSERT forms a stable association with PKGIα. hSERT and HA-PKGIα or HA-PKGIβ were coexpressed in HEK-293 T cells and coimmunoprecipitations were performed as described in Methods. Total cell extracts were blotted for A) HA to reveal PKG expression or B) SERT. C) Co-immunoprecipitations of SERT and PKG using anti-HA for immunoprecipitations and anti-SERT for blotting. Experiment shown was replicated in four independent experiments with similar results.
expression studies, HEK-293 T cells were plated at a density of 100,000 cells/mL in individual wells of 6-well plates one day before transfection. Transfections were performed with TransIt™-LT1 (Mirus Bio Corporation, Madison, WI) and 500 ng of each cDNA, in accordance with manufacturer’s instructions. For the siRNA studies, HeLa cells were plated at a density of 10,000 cells/mL in individual wells of a 24-well plate. Transfections were performed with Lipofectamine™ 2000 (Invitrogen), 25 ng pcDNA3-hSERT, and 100 nM siRNAs. For the siRNA studies, pcDNA3-hSERT and ON-TARGETplus SMARTpool human PRKG1 siRNAs (Thermo Fisher Scientific, Pittsburgh, PA) were used for transfections. To evaluate siRNA transfection efficiency, Allstars Hs Cell Death Control siRNAs (Qiagen, Valencia, CA) were transfected as described for PKG siRNAs as described by the manufacturer and dying/dead cells counted as a proportion of the total number of cells transfected by phase microsopy.

**Plasmid constructs**

The pcDNA3-His-hSERT construct was generated from previously described pcDNA3-hSERT [39] by adding a [His]_{10}-Gly epitope sequence after the ATG start site using the QuikChange site-directed mutagenesis kit (Stratagene, La Jolla, CA). pcDNA3-HA-PKGα and pcDNA3-HA-PKGβ were kindly provided by Dr. Bo Cen (Columbia University, New York, NY).

**RT-PCR Studies**

Total RNA from adult male, Sprague-Dawley rat midbrain and from RN46As was isolated using 5 Prime PerfectPure RNA kits (Fisher Scientific, Pittsburgh, PA) as described in the manufacturer’s protocol. 1 μg of RNA was subjected to reverse transcription followed by PCR (RT-PCR) using the TITANIUM™ One-Step RT-PCR Kit (Clontech Laboratories, Inc., Mountain View, CA) and PKGα (forward-5’ GGTACATAACAATCCACAGTCTC 3’, reverse-5’ CTCCTCCTCTAATTAG 3’), PKGI(forward-5’ GGTACATACAATCCACAGTCTC 3’, reverse-5’ AATCTGCACACTGAAACTCTCG 3’) and PKGII (forward-5’ GTATGTTCCACGCACGTCT 3’, reverse-5’ CGATGTAATCGCCCTGTTGATA 3’) specific oligonucleotide primers (Invitrogen Corporation, Carlsbad, CA). Reaction products were resolved by 2% agarose gel electrophoresis and sequenced using dideoxy terminator sequencing (Vanderbilt University Medical Center DNA Sequencing Core).

**Immunostaining**

The immunofluorescence protocol was adapted from previous work [57]. Briefly, 1–2 days before fixation, RN46A cells were seeded onto circular poly-D-lysine-coated coverslips in 24-well plates. Cells were fixed with 4% paraformaldehyde for 30 min., followed by either permeabilization with 0.2% NP40 or application of primary antibodies overnight at 4°C without detergent (nonpermeabilized conditions). Treatment with secondary antibodies conjugated to fluorophores Cy2 or Cy5 for 1 hr was followed by mounting of coverslips to charged slides. Indirect immunofluorescence images were captured using an LSM510 confocal microscope (Vanderbilt University Medical Center Cell Imaging Shared Resource supported by National Institutes of Health grants CA68485, DK20593, DK58404, HD15052, DK59637 and EY08126). Images were quantified using the Integration Colocalization Analysis section of WCIF Image J [http://www.uhnresearch.ca/facilities/wcif/imagej/]. Intensity Correlation Quotient (ICQ) values are presented and were taken as evidence for dependent staining or colocalization when the ICQ value falls between 0 and + 0.5. Independent staining or lack of colocalization was indicated when ICQ values were not different from 0 or were between 0 and -0.5. Data were analyzed by one-sample t-test comparing means versus a value of 0.

**Kinase inhibition assays using DT-2**

Kinetic assays (K<sub>i</sub>, IC-50) were performed as previously described [34,58] with purified recombinant kinases and multiple concentrations of DT-2 (YGRKKRRQRRPP- LRKKKKKH). Recombinant PKG Iα, PKG Iβ, PKA Iα (catalytic subunit) and PKGII were expressed from insect cell cultures using the Bacmid baculovirus expression system (Gibco/Invitrogen) and purified to apparent homogeneity as described recently [34]. Recombinant PKCα (human) was obtained from Calbiochem. All enzymes were assayed using saturating concentrations of the peptide TQAKRKSLAMA as substrate. PKG isoforms were activated with 1 μM cGMP, PKA was stimulated with 1 μM cAMP, and PKC was activated using 50 μg/μL phospholipids and 100 ng/mL 12-O-Tetradecanoylphorbol-13-acetate.

**Fractionation and activity assays**

Approximately 500 million RN46A cells were homogenized in ice-cold 10 mM potassium phosphate pH 6.8, 1 mM EDTA, and 25 mM β-mercaptoethanol (KPEM) and centrifuged at 10,000 × g for 20 min. at 4°C. The supernatant was applied to a pre-equilibrated 0.9 × 8 cm DEAE-Sephael column and the column was washed with KPEM with 20 mM NaCl. A linear gradient (20 mM to 280 mM NaCl) was next run on the column and 30-drop fractions were collected. As previously described [59], kinase assays to determine PKG activity in each fraction were conducted in the presence and absence of 10 μM cGMP (a PKG activator) and 136 μg/mL heptapeptide substrate (RKRRSAE) as the PKG substrate. The cAMP-dependent protein kinase (PKA) inhibitor PKI-tide (5–24) was also included in the assay in order to block PKA activity. Activity was expressed as picomoles of [32P] incorporated per min per mL of sample.
Co-immunoprecipitations and immunoblotting
Two days post-transfection with his-hSERT and HA-PKGI plasmids, HEK-293 T cells were detergent-extracted with 1% TRITON-X-100 and prepared for immunoprecipitation, as previously described [43]. Briefly, protein concentrations of lysates were determined with a bicinchoninic acid assay kit (Pierce, Rockford, IL), using bovine serum albumin as a standard. 50–100 µg total protein was immunoprecipitated with anti-HA High Affinity beads (Roche) for 1 hr and eluted with Laemmlli sample buffer. Samples were subjected to SDS polyacrylamide gel electrophoresis and electroblotted to polyvinylidene difluoride membrane (Amersham), followed by incubation with primary antibodies. Blots were thoroughly washed with PBS containing 0.5% Tween-20 and developed with enhanced chemiluminescence (Amersham) with multiple exposures captured to ensure nonsaturation conditions.

Transport assays
60 hrs after transfection with pcDNA3-hSERT and siRNAs, HeLa cells were subjected to 8-Br-cGMP treatment and 5-HT transport assays as previously described [17]. Briefly, transfected cells were treated with 100 µM 8-Br-cGMP for 10 min., then incubated with 20 nM [3H]5-HT for 10 min. After thorough washing with Krebs-Ringers-Hepes, pH 7.4, cells were solubilized in MicroScint 20 (PerkinElmer, Waltham, MA) and [3H]5-HT accumulation was quantified using a TopCount plate scintillation counter (Perkin Elmer). Specific 5-HT uptake was determined by subtracting the amount of [3H]5-HT taken up by non-transfected cells.

Statistical analyses
All data were derived from experiments replicated a minimum of three times. Values are expressed as mean ± SEM. All Statistical analyses were completed in GraphPad Prism (GraphPad, San Diego, CA). In Fig. 2, one-sample Student’s t tests were performed by comparing mean values to 0, and a value of p < 0.05 was taken as statistically significant. Analysis by one-way or two-way ANOVA was used followed by post-hoc testing (Bonferroni or Tukey) for Fig. 3 and 4, and a value of p < 0.05 was accepted as statistically significant. In Additional file 2, Analysis via a Repeated Measures One-Way ANOVA was utilized followed by Bonferroni post-tests, and a value of p < 0.05 was accepted as statistically significant.

Competing interests
The authors declare that they have no competing interests.

Authors' contributions
JAS designed and carried out experiments and data analysis resulting in Figures 1, 2, 3, 4, 5, and drafted the manuscript. AMDC provided direction in experimental design, data analysis and interpretation, and paper layout/editing. JW generated DNA constructs and provided assistance with immunohistochemical studies. HJGM provided training and guidance with experiments resulting in Figure 2. HCP completed experiments resulting in Figure 4. CKN and WRD performed kinase specificity assays noted in Results. JDC and SHF oversaw and participated in experiments comprising Figure 3 and assisted in data interpretation. RDB designed and coordinated the full study and assisted in manuscript construction. All authors read and approved the final manuscript.

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Additional material

Additional file 1
PKGI antibody specifically labels RN46A cells. The data provided demonstrate the specificity of the PKG antibody. Additional Figure 1. PKGI antibody specifically labels RN46A cells. RN46A cells were fixed, permeabilized, stained, and imaged by confocal microscopy as described in Methods. Left panel) RN46A cells stained with anti-PKGI (Assay Designs, KAP-PK002E). Right panel) Antibody preabsorption with bovine lung PKGIa blocks the majority of PKGI staining, indicating antibody specificity. DIC image is inset in right panel to demonstrate presence of cells. Scale bars represent 10 µm. Data are representative of at least three independent experiments.
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Additional file 2
siRNA knockdown of PKGI due to low transfection efficiency. The data provided support the results shown in Figure 4 by demonstrating analysis of the siRNA-mediated protein knockdown and corresponding transfection efficiency. Additional Figure 2. siRNA knockdown of PKGI due to low transfection efficiency. HeLa cells were plated identically to those used for Figure 4, and assayed for PKGI protein and cell viability as described in Methods. A) Knockdown of PKGI protein by siRNAs. Transfected and siRNA-treated HeLa cells were lysed and analyzed via immunoblot for PKGI and β-actin immunoreactivity (IR). Analysis via a Repeated Measures One-Way ANOVA (control, mock, PKGI siRNA) of raw values for PKGI abundance normalized for β-actin signal in the same sample indicates a significant overall effect of PKGI siRNA (p < 0.05), with Bonferroni post-tests indicating no significant difference between mock and control but a significant (* = p < 0.05) difference between PKGI siRNA and control. Representative blot image is shown below quantitation. Percent changes are plotted for ease of evaluation. B) Measurement of transfection efficiency. The AllStars Hs cell death control siRNA (cell death siRNA; QIAGEN Inc., Valencia, CA) was used to quantitate siRNA transfection efficiency in the HeLa cells 48 hrs. post-transfection. Data are representative of at least three independent experiments, and results are presented as means ± SEM.
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