NEUROSCIENCE

A switch in G protein coupling for type 1 corticotropin-releasing factor receptors promotes excitability in epileptic brains

Chakravarthi Narla,1,2 Tanner Scidmore,1,2 Jaymin Jeong,1,3 Michelle Everest,1 Peter Chidiac,2,4 Michael O. Poulter1,2,3*

Anxiety and stress increase the frequency of epileptic seizures. These behavioral states induce the secretion of corticotropin-releasing factor (CRF), a 40-amino acid neuropeptide neurotransmitter that coordinates many behavioral responses to stress in the central nervous system. In the piriform cortex, which is one of the most seizurogenic regions of the brain, CRF normally dampens excitability. By contrast, CRF increased the excitability of the piriform cortex in rats subjected to kindling, a model of temporal lobe epilepsy. In nonkindled rats, CRF activates its receptor, a G protein (heterotrimeric guanosine triphosphate–binding protein)–coupled receptor, and signals through a Gαq/11–mediated pathway. After seizure induction, CRF signaling occurred through a pathway involving Gαq. This change in signaling was associated with reduced abundance of regulator of G protein signaling protein type 2 (RGS2), which has been reported to inhibit Gαq-dependent signaling. RGS2 knockout mice responded to CRF in a similar manner as epileptic rats. These observations indicate that seizures produce changes in neuronal signaling that can increase seizure occurrence by converting a beneficial stress response into an epileptic trigger.

INTRODUCTION

Aversive emotional experiences increase both the frequency and the severity of seizures in humans and in animal models (1, 2). Epileptic patients are also more vulnerable to suicide, anxiety attacks, and depression than healthy individuals (3, 4). Negative emotions such as worry and fear resulting from stress and depression affect neuronal activity within many brain regions, including limbic structures, such as amygdala, hippocampus, and piriform cortex; these brain regions also support epileptiform activity. The epileptic brain could be more susceptible to otherwise normal stress responses or, alternatively, the brain state could change the nature of the stress response so that it now promotes seizures. Several hormones and neurotransmitters are increased by anxiety and stress conditions and might therefore be involved in stress-dependent seizurogenesis. Basal concentrations of stress hormones are increased in epileptic patients compared to healthy individuals. For instance, increased seizure frequency is observed with increased concentrations of cortisol in epilepsy patients (5). The polypeptide hormone corticotropin-releasing factor (CRF), which also acts as a neurotransmitter, might also serve to increase seizure frequency and has been linked with various psychological disorders (6). The presence of CRF immunoreactive cell bodies in the paraventricular nucleus (PVN) links CRF to the endocrine stress axis, and the dense distribution of CRF immunoreactivity in the dorsal raphe nucleus and locus coeruleus implicates CRF in the modulation of monoaminergic pathways. Individuals with major depressive disorder may have hyperactive hypothalamic-pituitary-adrenal axis functioning, including CRF hypersecretion and increased CRF mRNA in PVN (7).

In other portions of the central nervous system, CRF is released from axons arising from cells of the central amygdala and interneurons (8–10). It acts on type 1 CRF receptors (CRFR1) with high affinity and with low affinity on CRFR2, both of which are G protein [heterotrimeric guanosine triphosphate (GTP)–binding protein]–coupled receptors (GPCRs). We have shown that in the piriform cortex, CRF acting through CRFR1 dampens excitation of the piriform cortex circuitry (11), whereas it usually increases excitability in other brain regions (12). This activity is likely mediated through activation of Gαq/11 signaling path because protein kinase C (PKC) activation and antagonism either mimic or block the activity of CRF, respectively.

Because epileptic seizures are exacerbated by anxiety and stress, we decided to test the hypothesis that the effect of CRF could depend on the underlying condition of the brain or its disease state. We used the kindling model of epilepsy, which mimics many aspects of temporal lobe epilepsy in humans (12). In contrast to the normal brain state, we found that CRF augmented the principle cell excitation in kindled piriform cortex. This alteration in function appears to be due to a change in the signaling pathway activated by the CRFR1 receptor, which occurs through the down-regulation of a regulator of G protein signaling protein type 2 (RGS2).

RESULTS

Upon activation of the piriform cortex by stimulation of the lateral olfactory tract (LOT) in nonkindled rats, CRF reduced the excitation of layer II dorsal endopiriform nucleus (DEn) neurons and the inhibition of layer III inhibitory interneurons, resulting in reduced disinhibition of layer II neurons (Fig. 1, A to D), as indicated by voltage-sensitive dye imaging (VSDI) (movies S1 and S2). By contrast, in kindled rats, CRF increased layer II and DEn activity, whereas in layer III, it further decreased the neuronal activity, enhancing disinhibition (Fig. 1, E to H, and movies S3 and S4). Thus, after kindling, CRF enhanced circuit activation, an effect opposite to that in sham-operated control rats. We hypothesized that this switch in activity could be due to changes in the relative abundance of CRFR1 and CRFR2. Immunohistochemical analyses revealed that CRFR1 protein abundance was unaltered in all of the layers of the piriform cortex.
Fig. 1. Activation of CRFR1 by CRF dampened piriform cortex circuitry in normal rats but potentiated it in kindled rats. (A) Representative images showing activation of piriform cortex from nonkindled rats without (top) or with (bottom) CRF perfusion before recording. Each image in the bottom row was taken at the same time interval as that of its corresponding image in the top row. (B to D) Quantification of CRFR1 activation shows that the activity of layer II pyramidal cells and interneurons of DEn is increased, whereas that of layer III interneurons is decreased in nonkindled animals over the range of stimulation frequencies used to activate the circuit. **P < 0.01. n = 7 slices from six rats. (E) Representative images showing activation of layers of piriform cortical slices from kindled rats without (top) or with (bottom) CRF perfusion before recording. (F to H) Quantification of CRFR1 activation shows that the activity of layer II pyramidal cells and interneurons of DEn is decreased, whereas that of layer III interneurons is increased in nonkindled animals over the range of stimulation frequencies used in piriform cortical slices from kindled rats to activate the circuit. **P < 0.01. Kindled n = 13 slices from nine rats. Arrows in (A) and (E) indicate orientation of slice. D, dorsal; V, ventral; M, medial; L, lateral. Red, highest ΔF/F; orange, yellow, and green, medium ΔF/F; blue, low ΔF/F.

(fig. S1A) and that CRFR2 abundance was low and unchanged as well (fig. S1B). The distribution and abundance of the CRF peptide were also unchanged (fig. S1C). As in normal controls, CRF acted through the activation of CRFR1, because antalarmin, a CRFR1 antagonist, blocked the effects of CRF in kindled rats (Fig. 2). The median inhibitory concentration (IC50) for antalarmin increased to about 200 nM, which is in contrast to the 100 nM that has been previously reported by our group in nonkindled rats (11).

Because there was no change in CRFR or CRF peptide abundance and the effects were mediated through the same receptors, we next investigated the signaling pathways involved in CRF responses. We have previously reported that in nonkindled animals, CRFR1 signals by activating PKC through Gα11 (11). Gα11-activated phospholipase C (PLC) hydrolyzes phosphatidylinositol 4,5-bisphosphate (PIP2) to diacylglycerol (DG) and inositol 1,4,5-trisphosphate (IP3). DG, a second messenger, activates PKC and facilitates further signaling cascades. Here, we showed that CRFR1 signaled through the activation of a Gαs-mediated pathway. We found that forskolin, an activator of adenylyl cyclase that mimics the effect of activated Gαs, mimicked CRF responses, and this was antagonized by the protein kinase A (PKA) inhibitor H-89 (Fig. 3A). We also showed that H-89 blocked the effects of CRF (Fig. 3B). The application of the PKC inhibitor bisindolylmaleimide (BIS) did not block the effects of CRF (Fig. 3C), contrasting with our previous observations that application of BIS blocks the effects of CRF in kindled animals (11).

We first hypothesized that the change in signaling pathways after kindling could be due to changes in the relative abundance of Gαs and Gα11 mRNA and protein. For example, an increase in Gαs abundance or a decrease in Gα11 abundance could favor the coupling of CRFR1 to Gαs. However, Gαs mRNA expression decreased, and protein abundance was unchanged. The abundance of Gα11 mRNA and protein was also unchanged (fig. S2). These observations therefore do not provide an explanation as to why the signaling was altered. We next considered the possibility that another mRNA(s) coding for protein(s) involved in intracellular signaling was altered. To assay for this, we used commercially available quantitative polymerase chain reaction (qPCR) arrays that permit the quantitation of mRNAs encoding 84 differing proteins involved in second messenger sig-
Fig. 2. CRF signals through CRFR1 in kindled rat piriform cortex brain slices. (A) Representative images of the activation of the piriform cortex layers in slices at differing time points before (top) and after the application of CRF in the presence of the CRFR1 antagonist antalarmin (ANT). The responses are similar in the presence of CRF and CRFR1 antagonist. (B) Dose-inhibition curve for CRFR1 activity in layer II over the range of concentrations of antalarmin used. *n = 9 slices from seven rats. Red, highest ΔF/F; orange, yellow, and green, medium ΔF/F; blue, low ΔF/F.

Fig. 3. CRF signals through a Goαs-mediated signaling pathway. (A) Quantification of the effect of forskolin (an adenylyl cyclase activator) with or without subsequent administration of H-89 on CRFR1 activation in piriform cortical slices from kindled animals. *P < 0.05. *n = 11 slices from nine rats per group. (B) Quantification of the effect of the PKA inhibitor H-89 on CRF-mediated activation of CRFR1 in piriform cortical slices from kindled animals. *P < 0.05. *n = 9 slices from eight rats per group. (C) Quantification of the effect of the PKC antagonist BIS on CRF-mediated activation of CRFR1 in piriform cortical slices from kindled animals. *P < 0.05. *n = 9 slices from seven rats per group.

Fig. 4. RGS2 mRNA and protein abundance is decreased in kindled rats. (A) Δ cycle threshold (ΔCt) was compared between kindled (n = 11) and non-kindled (n = 9) samples. *P < 0.05. ΔCt is the Ct (reference gene) − Ct (gene of interest). Analysis shows that RGS2 mRNA transcripts are significantly decreased in kindled piriform cortex in comparison to control. No significant difference is observed in RGS17 mRNA transcripts between non-kindled and kindled animals. (B) Representative confocal images indicating reduced RGS2 abundance in layer II after kindling. The brightest pixels three times above background have been marked in white to indicate differences in expression (scale bar, 20 mm). (C) Relative percentage of pixels that were above this cutoff (n = 5 slices from five control rats; n = 4 slices from four kindled rats; *P < 0.05).
The epileptic state makes the brain more susceptible to an otherwise normal physiological response. The extent to which this alteration in brain state can activate different G protein cascades (18). Although CRFR1 was initially classified as a GPCR that activated adenylyl cyclase (18), CRFR activation can activate different G protein cascades (19–21). For example, the neuroprotective effects of CRF in the hippocampus are mediated by the activation of PKA and MAPK (mitogen-activated protein kinase) signaling pathways (22). Moreover, CRFR1 couples to G\textsubscript{q/11} in BALB/c mice hippocampal neurons, whereas it couples to G\textsubscript{a1} and G\textsubscript{a11} in C57BL/6N mice hippocampal neurons (19). In cultured hippocampal neurons, CRF regulation of N-methyl-D-aspartate currents is mediated by PKC, which correlates with increased PLC-\(\beta\)3 abundance (23). In prefrontal cortex pyramidal neurons, CRF activates CRFR through the activation of PKC to enhance stressor responses (24). As previously mentioned, we have shown that activated CRFR suppresses piriform cortex activity through a PKC-dependent mechanism (11). Thus, it seems clear that depending on the region and

**DISCUSSION**

In an animal model that resembles human temporal lobe epilepsy, we showed that the stress-related neurotransmitter CRF became excitatory and potentially more seizurogenic, reversing the polarity of its activity from inhibitory to excitatory in the piriform cortex. Our observations indicate that the epileptic state was accompanied by greater susceptibility to stressor-induced excitability in a brain region that supports seizures and seizurogenesis. These findings suggest that the underlying brain pathology may be an important determinant in the exacerbation of epilepsy in response to heightened anxiety. In effect, the neurochemical alterations support seizure onset. This observation is opposite to the one where the stressor responses are otherwise “normal” (cell signaling and polarity of the response is constant), but the epileptic state makes the brain more susceptible to an otherwise physiological response. The extent to which this alteration in brain state may account for other comorbidities or susceptibilities seen with those with epilepsy (for example, depression) is not known. In addition to epilepsy, it is also possible that similar alterations may occur in individuals who have suffered a traumatic brain injury and who can experience increased frequency of anxiety- and stress-induced seizures in brain regions that have not been directly injured. Our observations therefore provide an increased impetus to consider “bystander” outcomes that may exacerbate pathophysiological neurological outcomes.

The change in the polarity of the CRFR\textsubscript{1} signaling occurs through the switch in the signaling cascade by which CRF acts. Although CRFR\textsubscript{1} was initially classified as a GPCR that activated adenylyl cyclase (18), CRFR activation can activate different G protein cascades (19–21). For example, the interferon gamma inhibin antagonist H-89 blocked the effects of CRF, revealing that PKA activation was essential for signaling of CRFR\textsubscript{1} (Fig. 6, H to J). We also found that PKC antagonism using BIS in RGS KO mouse and PKA antagonism in wild-type mouse using H-89 did not affect CRFR\textsubscript{1} signaling (fig. S6).
pathological condition of the brain, CRFR₁ can activate distinct G proteins, resulting in activation of multiple signaling pathways. The present findings show that RGS2 protein may be an important mediator of this variation in CRFR₁ signaling.

GPCR signaling involves a series of complex events that lead to a conformational change of the receptor upon ligand binding and then activation of G protein by promoting exchange of GTP and GDP (guanosine diphosphate) associated with Gz. Many factors influence these events, including perceiving the extracellular signals through transducing them to heterotrimeric G proteins and to downstream effectors. RGS proteins interact with Gz subunits to attenuate GPCR-mediated signaling by accelerating Gz-GTPase (guanosine triphosphatase) activity (13) and/or by inhibiting G protein-effector interaction (25–27). RGS2 can inhibit Go₅₆-stimulated adenyl cyclase activity, even though it does not promote the GTPase activity of Go₅₆ (15, 28, 29). RGS2 may bind directly to Go₅₆ (29, 30) or to adenyl cyclase (16, 17, 31), and thus, RGS2 could produce its inhibitory effect on cAMP production by interfering with Go₅₆-adenyl cyclase coupling. Overall, these reports corroborate our findings that the presence of RGS2 in control rats decreased the production of cAMP and thus suppressed the CRFR₁-stimulated activity of Go₅₆. However, various RGS proteins (including RGS2) have GTPase activating protein activity for Go₁₁ and also function as potent inhibitors of Go₁₁ signaling compared to other types of RGS proteins (14, 15, 32–34). Thus, it is still not clear how RGS2 is working before and after kindling, namely, how changes in RGS2 abundance might selectively favor either Go₁₁- or Go₅₆-mediated effects of CRFR₁ activation. Our findings indicated that, in the absence of RGS2 (in the KO mouse) or when RGS2 abundance is low, as occurs in the kindled rat, CRFR₁ activation seemed to “prefer” to signal through Go₅₆. Although it is easy to rationalize how a decrease in RGS2 could increase Go₅₆-mediated signaling, a concomitant decrease in Go₁₁-mediated signaling would not be expected. This finding recalls the decreased G₁₂/₁₃ signaling effect that occurs upon a decrease in RGS5 in vascular smooth muscle (35), although in that case the G protein would not be considered a potential target of the RGS protein. Although RGS2 can inhibit CRFR₁-stimulated Go₁₁ signaling, the targeting of RGS proteins to particular pathways in vivo may be governed by additional factors such as scaffolding proteins (27), and thus, a decrease in RGS abundance would not necessarily result in increased activation of all of its potential G protein partners.

RGS2 is present in brain regions that are implicated in the pathophysiology of anxiety and depression, such as hippocampus, amygdala, cerebral cortex, hypothalamus, and raphé nucleus of rats and mice (15, 36–38). These regions have reciprocal connections with the piriform cortex and pathological changes in these regions may negatively affect the physiology of the piriform cortex and perhaps seizurogenesis. Hippocampal slices from RGS2 KO mice have increased excitation in CA1 pyramidal cells, and these mice are more anxious compared to their wild-type counterparts (39). Moreover, a mutation in the RGS2 gene causes anxiety in mice (40), and decreased RGS2 abundance is associated with depression-like behavior in mutant mice (41). Thus, the relationship between RGS2 abundance and behavior may depend on whether anxiety or depression is also present, which may limit the usefulness of drugs targeting RGS2.
Several studies have reported the clinical implications of RGS2 and complications associated with its polymorphisms. RGS2 has been proposed to be an important drug target in anxiety-related disorders, depression, and panic disorders in humans (42, 43). Several studies have reported the pathogenetic association of single-nucleotide polymorphisms (SNPs) especially rs4606 in the RGS2 gene and neuropsychiatric disorders. RGS2 gene polymorphism has been associated with the development of panic disorder related to agoraphobia in 173 subjects of German descent (38). Furthermore, abnormalities in serotonin receptor function due to RGS2 gene polymorphisms have been found in Japanese suicide victims (44). Examination of postmortem brains revealed significant differences in allele frequencies of two SNPs (SNP 2 and 3) in suicide victims compared to controls. Moreover, RGS2 immunoreactivity is greatest in the amygdala and BA9 region of the prefrontal cortex of suicide subjects compared to controls, suggesting that the increased RGS2 abundance alters GPCR intracellular signaling (44). In 2004 Florida hurricane victims, the C allele of rs4606 SNP of RGS2 has been correlated with increased chances of developing generalized anxiety disorder, posttraumatic stress disorder, and ideation to commit suicide (42, 45). Behavioral studies and functional magnetic resonance imaging have identified the association of the G allele of rs4606 SNP in anxiety-related phenotype and increased activation of the insular cortex, a part of limbic system, in human subjects (46). Genetic variation such as rs4606 SNP that leads to reduced RGS2 abundance is associated with antipsychotic-induced parkinsonism (47, 48). Together, polymorphisms in RGS2 in different brain regions are associated with various neurological disorders. More research is needed to understand the physiology and pathophysiology of RGS2 proteins and thus their clinical implications in human neurological disorders.

In summary, our observations provide a mechanism by which anxiety and stressors may exacerbate the occurrence of seizures. The epileptic state alters the function of a stress neurotransmitter that increases the excitability of a brain region that supports seizurogenesis, although it is uncertain how general this effect may be. Other brain pathophysiological states (such as stroke, repeated concussion, or traumatic brain injury) may also produce similar alterations in signaling in many GPCR responses. Thus, we will need to consider how mental disorders may be mitigated by the altered pathophysiological state of the brain.

MATERIALS AND METHODS

Procedures were performed in accordance with the guidelines of the Canadian Council on Animal Care and approved by the University of Western Ontario Council on Animal Care.

Kindling procedure

Kindling is a phenomenon in which the delivery of daily electrical impulses to the limbic region, such as the amygdala, leads to development of seizures and epilepsy. A detailed explanation of the surgical procedure for the electrode implantation and kindling procedure was described elsewhere (49).

Slice preparation and dye loading

Adult male Sprague-Dawley rats weighing 150 to 180 g or C57BL/6N mice weighing 20 to 25 g were used in our experiments. They were housed individually with free access to food and water under a continuous 12-hour light and dark cycle. Animals were anesthetized with a ketamine–medetomidine hydrochloride combination and then perfused through the heart with an ice-cold artificial cerebrospinal fluid (ACSF) in which sodium ions were replaced by choline ions (49). The ACSF used was composed of 110 mM choline chloride, 2.5 mM potassium chloride, 1.2 mM sodium dihydrogen phosphate, 25 mM sodium bicarbonate, 0.5 mM calcium chloride, 7 mM magnesium chloride, 2.4 mM sodium pyruvate, 1.3 mM ascorbate, and 20 mM dextrose (30). The brains were perfused to flush any blood out of the vessels and thereby prevent the iron in the blood from oxidizing and causing damage to neuronal cells. The brain was rapidly removed after perfusion, and the region containing anterior piriform cortex was carefully cut into a block to facilitate slicing by a Vibratome (slices were 400 μm thick). The slices were incubated at 37°C for 30 min and subsequently moved to a room temperature (22°C) bath for 45 min. The perfusion, slicing, and incubation procedures were carried out in choline-ACSF with a continuous supply of carbogen (95% O₂ and 5% CO₂ mixture). These slices were used for both VSDI and patch-clamp recording. For VSDI, slices were incubated in the voltage-sensitive dye Di4-ANEPPS (D-199, Invitrogen Molecular Probes Inc.) for 35 min. The stock solution of the dye was dissolved in ethanol (22 mg/ml). On the day of the experiment, the dye incubation was prepared by mixing 60 μl of dye stock with 500 μl of fetal bovine serum, 500 μl of ACSF, and 310 μl of 10% Cremophor EL solution. The concentration of dye in the final solution was 0.1 mg/ml. After incubation, slices were washed for 8 to 10 min with ACSF and transferred to a recording chamber. The temperature of the bath was maintained at 32°C during recordings and continuously supplied with carbogen-bubbled ACSF, which was composed of 110 mM NaCl, 2.5 mM KCl, 1.2 mM NaH₂PO₄, 25 mM NaHCO₃, 2.0 mM CaCl₂, 2.0 mM MgCl₂, and 20 mM dextrose. The pH and osmolarity of the solutions were adjusted to 7.3 to 7.4 and 297 to 305 mOsm, respectively.

A platinum-iridium electrode (MicroProbes Inc.) with a tip diameter of 200 to 300 μm was used to stimulate the LOT of the piriform cortex. The stimulation of each slice was in the range of 160 to 200 μA; each square pulse was 2.0 ms in length. The electrode was connected to a stimulator (S88X Dual Output Square Pulse Stimulator, Grass Technologies, Astro-Med Inc.), which controlled the pulse frequency and train duration (51).

Voltage-sensitive dye imaging

Each optical recording was about 10 s in length and consisted of two phases. The first contained a recording of the background activity for 2 s followed by the application of stimulus for 1 s with a train of frequencies ranging from 5 to 80 Hz. The acquisition rate was set at 5 ms per frame. The camera saturation was set at about 50% for each recording. Optical signals were recorded by a CMOS (complementary metal oxide semiconductor) camera (MiCam Ultima Brain Vision Inc.) mounted on top of an upright microscope (fixed-stage upright microscope, BX51WI, Olympus); sections were illuminated using the light from a 100-W halogen lamp source (HLX 64625, Mirolic Americas Scientific Corp.) passed through an excitation filter (λ = 530 ± 10 nm). A long-pass emission filter (λ > 590 nm) collected the fluorescent signals. A long working distance objective was used in the experiments [XLFuor 4X, NA (numerical aperture) 0.28, Olympus]. The movies were analyzed using the Brain Vision Analyzer software. A detailed explanation of the technique is described elsewhere (51).

qPCR analyses

Brain tissue from microdissected piriform cortex was placed in TRIzol (Invitrogen, catalog #75649) and homogenized. Extraction of RNA from the homogenate was performed following the manufacturer's protocol. To assess purity of extracted RNA, a NanoDrop 1000 spectrometer was used. The RNA (2 μg) samples with appropriate purity (≥260/280 levels) were reverse-transcribed using the Invitrogen SuperScript II with oligo(dT) primers. For the PCR, 2 μl (40 ng) of complementary DNA (cDNA) was placed in duplicate in flat cap PCR strips. A master mix containing SYBR Green Fast Mix for iQ, along with Milli-Q water and the primers for the protein of interest, was prepared according to the manufacturer's instructions (Quanta Biosciences). Twenty-three microliters of master mix was then added to each PCR tube to make the total volume 25 μl. The strips were then
placed in a Bio-Rad MyIQ thermocycler for subsequent PCR and fluorescence detection. Primers were designed and purchased (IDT) for the mRNAs encoding Go protein stimulatory (GNAS), G protein q polypeptide (GNAQ), RGS2, and RGS17. The primer sequences were as follows: GNAS, 5′-CTGTCCTGCAACASTAAAGAC-3′ (forward) and 5′-GCAGCTGTCTTCGATCTTTT-3′ (reverse); GNAQ, 5′-ATGGCAATAGACAGCGGCG-3′ (forward) and 5′-ATAAGGAAGGGTCAACCAC-3′ (reverse); RGS2, 5′-CGGGAGAAATGAGGCGGC-3′ (forward) and 5′-ATGGTGGCGGCTCCAGGAG-3′ (reverse); and RGS17, 5′-CTAAGATGGCGCACGTCGGG-3′ (forward) and 5′-CTGCGTGCTCCATTTTGT-3′ (reverse). Primer efficiency was quantified using a five-point 10× dilution series of rat brain cDNA. The PCR primer sets were evaluated for efficiency at 55°C annealing temperature. Cyclophilin A (PPIA) (Sigma-Aldrich, catalog #1475-017 and #1475-018) was the reference gene to normalize expression (forward, 5′-TGGGCTTCCCAGGAG-3′; reverse, 5′-GTCCTGCTCTTTTCGCCGC-3′). Furthermore, 84 genes (fig. S3) that encode proteins involved in the rat dopamine and serotonin pathways were evaluated using RT² Profiler rat PCR arrays (PARN 1582A-12, Qiagen). Equal amounts of pooled cDNA from nonkindled and kindled rat brains (five each) were used as template for their own respective plates. One plate was run for the nonkindled and kindled conditions; each well on the plate contained pooled genetic material of five individual rats. Each plate was run twice on separate occasions. The plate also contained five housekeeping genes, whose cycle thresholds were averaged and used for normalization of the data. Differences in the amount of mRNA present were evaluated through a comparison of the Δ cycle thresholds (ΔCt) between nonkindled and kindled brain samples. ΔCt is the Ct (reference gene) − Ct (gene of interest). Mean ΔCt between kindled and nonkindled rats was statistically analyzed using a Student’s t test. Data are expressed as means ± SE of the ΔCt values.

Phospho-site protein kinase screen
Samples of 250 mg of rat brain piriform cortex were washed three times in ice-cold phosphate-buffered saline (PBS) and then homogenized in lysis buffer as described in the Kinexus Bioinformatics protocol (www.kinexus.ca/). The tissue was sonicated four times for 10 s in an ice bath at 20% power with a Branson sonicator. The homogenate was then centrifuged at 20,000g for 30 min at 4°C in the Eppendorf Centrifuge 5810R. The supernatant was assayed for protein concentration using the Bradford assay (Bio-Rad). At a concentration of 1 mg/ml in SDS–polyacrylamide gel electrophoresis sample buffer, the samples were boiled for 4 min and then shipped on wet ice for the phospho-site protein kinase screen (catalog #KPPS 11.0) at Kinetworks Screening Services (Kinexus Bioinformatics Corp.). This assay consists of the analysis of 54 proteins that have a phosphorylation-dependent function in intracellular signaling. The complete list of the proteins assayed and the epitopes analyzed is provided in fig. S4, and further information can be found at www.kinexus.ca/. The assay quantitatively shows whether a protein has increased or decreased target phosphorylation.

Tissue preparation and fixation for immunohistochemistry
The following methodology has been described in detail elsewhere (32). The rats were anesthetized and perfused with 0.1 M PBS, followed by Lan’s fixative (4% paraformaldehyde and 20% picric acid in PBS). The brain was removed from the skull, stored in Lan’s fixative for 24 hours, and then placed in phosphate buffer solution with 30% sucrose for 48 hours at 4°C. The brains were then flash-frozen at −80°C and sectioned coronally (40 μm) using cryostat at −15°C. The sections were stored in cryoprotectant solution at −20°C.

Immunostaining
Sections were washed with PBS with 0.2% Triton X-100 three times for 5 min and blocked with 10% donkey serum and 10% goat serum in PBS with 0.025% Triton X-100 and 1% bovine serum albumin (BSA) for 1 hour. The primary antibodies were diluted in PBS with 1% BSA and 0.025% Triton X-100. The sections were labeled with a single primary antibody and incubated for 24 hours at 4°C. The primary antibodies used were anti-rabbit CRF (Abcam, ab11133, 1:250), anti-rabbit CRFR1 (Abcam, ab150561, 1:250), and anti-rabbit CRFR2 (Abcam, ab75168, 1:1000). The sections were then washed two times with PBS with 0.2% Triton X-100 for 5 min and incubated with a secondary antibody for 1 hour at room temperature. The secondary antibody used was Alexa Fluor 488 goat anti-rabbit immunoglobulin G (G IgG) (Molecular Probes, A-11034) diluted in PBS with 1% BSA and 0.025% Triton X-100 and was diluted 1:1000 for CRF and CRFR2 and 1:2000 for CRFR1. Sections were then washed three times with PBS with 0.2% Triton X-100 for 10 min and incubated in 1% Sudan Black B (Sigma-Aldrich, S2380) in 70% ethanol for 5 min. Sections were rinsed twice with 70% ethanol for 1 min followed by two 5-min washes with PBS. Sections were mounted on glass slides with glass coverslips using ProLong Gold Antifade Reagent with DAPI (4′,6-diamidino-2-phenylindole) mounting medium (Molecular Probes, P36935). Primary antibodies used were rabbit anti–G protein alpha S (Abcam, ab83735, 1:500) and rabbit anti-GNAQ (Abcam, ab75823, 1:250). The secondary antibody used was Alexa Fluor 488 goat anti-rabbit IgG (Molecular Probes, A-11034) diluted in PBS with 1% BSA and 0.02% Triton X-100 at a ratio of 1:1000. Primary antibody anti-RGS2 was used at 1:500 (Sigma-Aldrich, GW22245F) with secondary antibody Cy3 anti-chicken IgY at 1:500 (Cedarlane, 703-166-155).

Image acquisition and analysis
Confocal images were obtained using an Olympus IX51 inverted microscope using either a 60× oil immersion objective (NA, 1.4) or, for large field of viewing, a 20× objective (NA, 0.8) on a PerkinElmer UltraVIEW spinning disc confocal unit. The microscope was equipped with a Hamamatsu Orca-ER CCD (charge-coupled device) camera (1344 × 1024 pixels). All images were acquired using the Velocity software (version 4.2.1, Improvision). A stack of 10 images with 0.2-μm spacing in the z plane was taken. RGS2 images (Fig. 4) were acquired using a Zeiss LSM 510 Meta confocal microscope with a 40× (NA, 1.2) objective. For Fig. 4, the intensity of immunoreactivity is indicated by adding a channel (colored in white) that contains only pixels three times above background. For the semiquantitative analysis in Fig. 4C, we averaged the total amount of pixels in the field above this background value for comparison.

Statistical analysis
All statistical evaluations were done using the StatView software. Two-way analyses of variance (ANOVARs) were conducted to compare the neuronal responses after the application of BIS, PMA (phorbol 12-myristate 13-acetate), H-89, and forskolin in the piriform cortex layers in response to LOT stimulation. For CRF responses in the presence of BIS, PMA, forskolin, and H-89, a repeated-measures ANOVA was used in which responses of CRF in the absence of drugs served as the within-group measure. The pixel/voxel counts were protected least significance difference test to maintain P at 0.05.

SUPPLEMENTARY MATERIALS
www.sciencesignal.org/cgi/content/full/9/432/ra60/GDC1
Fig. S1. No changes in CRFR1, CRFR2, or CRF immunoreactivity in the layers of the piriform cortex.
Fig. S2. The expression of the mRNAs encoding Goα, Gαs, and Gαq/11 is unchanged after kindling.

www.SCIENTESIGNALING.org 14 June 2016 Vol 9 Issue 432 ra60 7
RESEARCH ARTICLE

Fig. S3. qPCR analysis of mRNAs encoding proteins involved in GPCR signaling. Fig. S4. Kindling-induced changes in the phosphorylation of signaling proteins. Fig. S5. CRFR1 antagonism did not block the effects of CRF. Fig. S6. PKC or PKA antagonism did not affect CRFR1 activation in wild-type and RGS2 KO mouse, respectively. Movie S1. Excitation and disinhibition in the layers of nonkindled rat piriform cortex imaged through VSDI. Movie S2. Change in the excitation and disinhibition in the layers of nonkindled rat piriform cortex after CRF application. Movie S3. Excitation and disinhibition in the layers of kindled rat piriform cortex imaged through VSDI. Movie S4. Change in the excitation and disinhibition in the layers of kindled rat piriform cortex after CRF application.

REFERENCES AND NOTES


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A switch in G protein coupling for type 1 corticotropin-releasing factor receptors promotes excitability in epileptic brains
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