Research Article

Modulation of Neuronal Activity on Intercalated Neurons of Amygdala Might Underlie Anxiolytic Activity of a Standardized Extract of *Centella asiatica* ECa233

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Received 3 December 2017; Revised 7 March 2018; Accepted 25 March 2018; Published 24 April 2018

Academic Editor: Hyunsu Bae

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GABAergic intercalated neurons of amygdala (ITCs) have recently been shown to be important in the suppression of fear-like behavior. Effects of ECa233 (a standardized extract of *Centella asiatica*), previously demonstrated anxiolytic activity, were then investigated on ITCs. Cluster of GABAergic neurons expressing fluorescence of GFP was identified in GAD67-GFP knock-in mice. We found that neurons of medial paracapsular ITC were GABAergic neurons exhibiting certain intrinsic electrophysiological properties similar to those demonstrated by ITC neurons at the same location in C57BL/6J mice. Therefore, we conducted experiments in both C57BL/6J mice and GAD67-GFP knock-in mice. Excitatory postsynaptic currents (EPSCs) were evoked by stimulation of the external capsule during the whole cell patch-clamp recordings from ITC neurons in brain slices. ECa233 was found to increase the EPSC peak amplitude in the ITC neurons by about 120%. The EPSCs in ITC neurons were completely abolished by the application of an AMPA receptor antagonist. Morphological assessment of the ITC neurons with biocytin demonstrated that most axons of the recorded neurons innervated the central nucleus of the amygdala (CeA). Therefore, it is highly likely that anxiolytic activity of ECa233 was mediated by increasing activation, via AMPA receptors, of excitatory synaptic input to the GABAergic ITC leading to depression of CeA neurons.

1. Introduction

Emotional symptoms especially fear/anxiety-like behavior are considered to be regulated through the amygdala region. The stimuli from the environment reach the basolateral amygdala (BLA) via thalamic and cortical afferents and transmit to the central nucleus of amygdala (CeA) where a proper anxiogenic response occurs [1]. Between BLA and CeA, the relay local processing happens at the intercalated cells (ITCs) level. The ITCs of the amygdala consist of densely packed GABAergic neurons surrounding BLA providing inhibitory control of neurons in BLA and CeA [2]. This group of neurons receives information from both intra- and extra-amygdaloid nuclei. The external capsule is the major connecting fiber to the ITCs. However, the exact fiber in external capsule that inputs to the ITCs is not clearly identified [3, 4]. Nevertheless, the projections from the medial prefrontal cortex to ITCs are thought to be a critical component of the forebrain circuitry for fear extinction [3]. Furthermore, ablation of ITCs has led to extinction retrieval deficits indicating the requirement of ITCs for the expression of fear extinction [5]. Thus, previous data suggest that ITCs play a critical role in fear extinction; a very important translational process since extinction-based exposure therapies is the mainstay for the treatment of anxiety disorder [6]. Therefore, it is our interest to explore how ITCs would respond to ECa233, a standardized extract of *Centella asiatica* previously shown to possess anxiolytic activity by our group [7].
**Centella asiatica** (CA), one of psychoactive medicinal plants, has been used in Ayurvedic medicine for centuries to relieve the symptom of anxiety and to contribute a deep state of relaxation and/or mental calmness throughout meditation practices. A recent clinical study showed the beneficial effects of CA on generalized anxiety disorder [8]. As to the mechanism of its action in brain function, CA is considered to enhance the GABAergic neurotransmission at the presynaptic level by facilitating the glutamic acid decarboxylase activity [9]. However, the possible function of CA on the glutamatergic synaptic transmission to GABAergic neurons remained unclear. A standardized extract of *Centella asiatica* or ECa233 was prepared to avoid the large variation in composition of herbal extracts and to ensure consistency in physical and chemical constituents. ECa233 contained triterpenoid glycosides not less than 80% and the ratio between madecassoside and asiaticoside should be 1.5 ± 0.5. Previous studies demonstrated that ECa233 exerted the neuroprotective effects on learning and memory deficit induced by 2-vessel occlusion (2VO) in mice at the doses of 10 and 30 mg/kg [10]. It protected the neuronal cell death in hippocampus. Moreover, the anxiolytic activity of ECa233 on stress-induced anxiety in chronic immobilization models was observed [7]. Interestingly, the effect of ECa233 on ERK1/2 signaling pathways was reported to promote the neurite outgrowth of neuronal cells [11].

**2. Materials and Methods**

**2.1. Animals.** The protocol of animal housing and treatment used in this study was approved by the Ethics Committee of the Faculty of Pharmaceutical Sciences, Chulalongkorn University, Thailand, and the Institutional Animal Care and Use Committee of the National Institutes of Natural Sciences, Japan. The GAD67-GFP knock-in and wild type C57BL/6j mice aged between 14 and 21 days were used for slice preparation. The procedures for the generation and genotyping of the GAD67-GFP knock-in mice were described by Kaneko et al., 2008 [12]. Mice homozygous for the GAD67-GFP allele were mated with C57BL/6j wild type mice to obtain the heterozygous mice. Wild type C57BL/6j mice were obtained from the National Laboratory Animal Center, Mahidol University.

**2.2. Electrophysiological Studies.** Male GAD67-GFP knock-in or wild type C57BL/6j mice were decapitated under deep isoflurane anesthesia. Their brains were quickly removed and submerged for 2-3 min in ice cold modified Ringer's solution containing (in mM): 234 sucrose, 2.5 KCl, 1.25 NaH2PO4, 10 MgSO4, 0.5 CaCl2, 26 NaHCO3, and 11 glucose and bubbled with 95% O2/5% CO2, pH 7.4. Coronal slices (~300 μm thickness) of the amygdala at approximately 1.64 mm caudal to bregma were cut with Microsystems (DTK-2000; Dosaka EM, Kyoto, Japan) and incubated with standard Ringer solution at room temperature for 1 h before recording. The standard Ringer solution contained (in mM) 125 NaCl, 2.5 KCl, 2 CaCl2, 1 MgCl2, 26 NaHCO3, 1.25 NaH2PO4, and 25 glucose and bubbled with 95% O2/5% CO2, pH 7.4. After incubation, slices to be used for recording were placed individually in a recording chamber on an upright microscope (DM LFS; Leica, Wetzlar, Germany) and continuously superfused with standard Ringer solution at a rate of 3–5 ml/min using a peristaltic pump (Minipuls 3, Gilson, Villiers, France). The tested substance, ECa233, was bath applied at the final concentration in standard Ringer solution.

The procedures of whole cell patch-clamp recordings are described previously [13]. Whole cell patch-clamp recordings were conducted on the neurons in ITC under visual control of patch pipettes using a patch-clamp amplifier (EPC-7; List, Darmstadt, Germany). GFP-positive neurons were selected using epifluorescence microscopy and then visualized with Nomarski optics with ×43 water-immersion objective lens. Patch pipettes were pulled from borosilicate glass capillaries (GC150T-10, 1.5 mm OD, Harvard Apparatus, Kent, UK) using the horizontal electrode puller (P-97; Sutter Instrument, CA, USA). The pipettes were filled with a glutaconate-based intracellular solution containing (in mM): 140 K-glutamate, 20 KCl, 0.2 EGTA, 2 MgCl2, 2 Na2ATP, 0.5 Na2GTP, 10 HEPES, and 0.1 spermine (pH 7.3) for the study of firing pattern, passive, and active properties of neurons. The Cs-based intracellular solution was used to study the AMPA and NMDA receptor-mediated excitatory postsynaptic currents, effect of ECa233 on eEPSCs consisting of (in mM): 120 Cs-glutamate, 10 CsCl, 2 MgCl2, 4 Na2ATP, 10 EGTA, 10 HEPES, and 0.1 spermine (pH 7.3). To stain the recorded neurons, biocytin (4 mg/ml; Sigma, St. Louis, MO) was dissolved in the intracellular solution. As the liquid junction potential between the standard Ringer’s solution and the gluconate-based intrapipette solution was estimated at ~10 mV, the actual membrane potential was corrected by this value. The osmolarity of the intrapipette solution was 280–290 mOsm/L. The resistance of the electrodes is 5–8 MΩ in the bath solutions. All recordings were performed at 32–34°C. Data were acquired using the pClamp system (pClamp 8.1; Molecular Devices, CA, USA).

**2.3. Comparative Electrophysiological Properties of ITCs between GAD67-GFP Knock-In and Wild Type C57BL/6j Mice.** Initially electrophysiological recordings were made on GABAergic neurons of ITC, easily identified by GFP fluorescence, in GAD67-GFP knock-in mice. Depolarizing current pulses were applied with 400 ms duration and step amplitudes up to 200 pA (40 pA in each step). The resting membrane potential (RMP), input resistant (Rin), threshold, and action potential amplitude were recorded. The same procedure was applied on neurons at the same location in wild type C57BL/6j mice and similar electrophysiological characteristics were obtained indicating GABAergic properties of the recorded neuron which were further used for analysis of ECa233 activity on ITC of amygdala.

**2.4. Effect of ECa233 on EPSCs of ITCs.** Electrical stimulation of external capsule was applied as a cathodal square wave pulse of 200 μs duration with an intensity of up to 150 μA using a concentric bipolar electrode (Clark Electromedical Instruments, Pangbourne, UK) to activate the input fibers to ITCs (Figures 1 and 2). ECa233 was bath applied to examine the EPSCs on ITCs. Data was analyzed with Clampfit.
pH 7.4, for 2-3 days at 4°C, fixed with 4% paraformaldehyde in 0.12M phosphate buffer, detached from the cells after recording. The slices were fixed throughout the recorded neuron (estimated to take 30 min), in the patch-clamp recordings. In order to distribute biocytin in the intrapipette solution and diffused into the neuron during recording neurons, 4 mg/ml biocytin was dissolved in the intracellular solution and applied to the patch electrode. To visualize biocytin we used avidin-biotin peroxidase complex solution (1%) (Vector Laboratories, Burlingame, CA) containing 0.3% Triton X-100 and 1% nickel ammonium sulfate, and 0.0003% H₂O₂ for 30 min. After washes with PBS, the slices were incubated in methanol containing 0.6% H₂O₂ for 30 min. The patch pipettes were carefully detached from the cells after recording. The slices were fixed with 4% paraformaldehyde in 0.12 M phosphate buffer, pH 7.4, for 2-3 days at 4°C, rinsed in 0.05 M PBS, pH 7.4, and incubated in methanol containing 0.6% H₂O₂ for 30 min. After washes with PBS, the slices were incubated in avidin-biotin peroxidase complex solution (1%) (Vector Laboratories, Burlingame, CA) containing 0.3% Triton X-100 for 3 h. After washes with PBS and 0.05 M Tris-buffered saline (TBS), pH 7.6, they were incubated in a TBS solution containing 0.01% dianinobenzidinetetrahydrochloride (DAB), 1% nickel ammonium sulfate, and 0.0003% H₂O₂ for 30 min. All procedures for visualization of biocytin were performed at room temperature. The slices were mounted on gelatin-coated slides, counterstained with cresyl violet, dehydrated, and then coverslipped. The morphological properties of stained cells were drawn using a camera lucida attached to a light microscope (Nikon, Japan).

### Table 1: Comparison of electrophysiological properties between ITC in GAD67-GFP knock-in and ITC in C57BL/6J mice.

| Parameter                        | ITC in GAD67-GFP | n  | ITC in C57BL/6J | n  | P value (t-test) |
|----------------------------------|------------------|----|----------------|----|-----------------|
| Resting membrane potential (mV)  | −71.16 ± 4.10    | 18 | −69.16 ± 4.66   | 18 | P = 0.195       |
| Input resistant (MΩ)             | 529.00 ± 59.18   | 18 | 534.33 ± 67.18  | 18 | P = 0.822       |
| Threshold (mV)                   | −41.41 ± 3.02    | 18 | −40.70 ± 3.44   | 18 | P = 0.565       |
| Action potential amplitude (mV)  | 77.2339 ± 5.29   | 18 | 76.47 ± 3.76    | 18 | P = 0.586       |

2.5. Chemicals and Test Compounds. The preparation and the phytochemical profile of ECa233 were described by Wanasuntronwong et al., 2012 [7]. All drugs and chemicals in the electrophysiological study were purchased from Sigma (St. Louis, MO). We used 10 μM bicuculline methobromide (Bic), 10 μM 6-cyano-7-nitroquinoxaline-2,3-dione (CNQX), and 50 μM D-2-amino-5-phosphonovaleic acid (AP-5) for suppression of GABAₐ receptors, AMPA/kainite receptors, and NMDA receptors, respectively.

2.6. Statistical Analyses. Results are represented as mean ± SEM from eight to ten animals per group. Statistical differences of the mean were analyzed using one-way analysis of variance (ANOVA) followed by Dunnett's post hoc test for a significance level when the "P" value was less than 0.05. Simple comparison between two groups was performed using Student's t-test. All statistical analyses were performed using PASW Statistics 18 (SPSS Inc, Chicago, USA).

3. Results

3.1. The Morphological and Electrophysiological Properties of ITCs in GAD67-GFP Knock-In and Wild Type C57BL/6J Mice. Firstly, we use GAD67-GFP knock-in mice to identify the GABAergic neurons in the ITCs. We found that ITC was a well-established structure with clear boundaries and almost all were GABAergic neurons. Comparative structure between GAD67-GFP knock-in and wild type C57BL/6J mice was shown in Figure 2. The boundary of this area was also visible in wild type C57BL/6J mice. Electrical properties of ITC neurons in GAD67-GFP knock-in and wild type C57BL/6J mice were investigated. The firing pattern and morphological properties of neuron in ITC between GAD67-GFP knock-in and wild type C57BL/6J mice showed no differences (Figure 3). Three types of firing pattern were demonstrated: regular spiking, late spiking, and fast spiking. The morphological properties were typically bipolar with the I-V plots of input resistance (Figures 3(c1) and 3(f1)). The passive and active membrane properties of neurons in both mice showed no difference (Table 1). The P value of RMP, Rin, threshold, and action potential amplitude was 0.195, 0.832, 0.565, and 0.586, respectively, by Student's pair t-test.

3.2. AMPA/NMDA Receptors Mediated eEPSCs Recorded in the ITC. In this study, the stimulating electrode was placed on the external capsule and the recording electrode was patched on neurons in the ITCs cluster (Figures 1 and 2). In order to examine the voltage dependence of the evoked responses.
**Figure 2:** The comparison of amygdala area morphology between GAD 67-GFP knock-in and wild type C57BL/6J mice. The whole cell patch-clamp recording was made in medial paracapsular ITC (Imp) neurons. Stimulating electrode was placed in external capsule. LA, lateral amygdala; BLA, basolateral amygdala; Imp, medial paracapsular ITC; IN, main ITC nucleus; CEL, lateral subdivision of the central amygdala; CEm, medial subdivision of the central amygdala; AStr, amygdalostrial transition area; and AL, nucleus of the ansa lenticularis.

**Figure 3:** Intrinsic electrophysiological properties of ITCs. (a) \((n = 25)\), (b) \((n = 5)\), and (c) \((n = 20)\) are the properties of ITCs in GAD67-GFP knock-in mice. (d) \((n = 24)\), (e) \((n = 3)\), and (f) \((n = 18)\) are the properties of ITCs in C57BL/6J mice. (c1) and (f1) are the light microscopy image of a representative biocytin filled in ITC neurons (typical bipolar structure) with their input resistance with \(I-V\) plot (scale bar = 50 \(\mu m\)).
EPSCs (eEPSCs), recording was performed in the presence of bicuculline (10 μM) to block GABA_A receptor activity and the Cs-based intracellular patch solution was used to reduce outward potassium current and block GABA_B receptors. The ionotropic glutamate receptor-mediated component at these synapses had a linear current-voltage (I-V) relationship and reversal potential of +1.2 mV (Figures 4(a) and 4(b)). NMDA-EPSCs were isolated by blocking AMPA/kainate receptor with CNQX (50 μM) in the presence of bicuculline. The NMDA receptor-mediated component at these synapses had a nonlinear I-V relationship, with a region of negative slope between −40 and +10 mV and reversal potential of 7.8 mV (Figures 4(a) and 4(c)). Additionally, the application of 10 μM APV (a selective NMDA receptor antagonist) in the presence of bicuculline and CNQX totally abolished the response (Figures 4(a) and 4(d)).

3.3. **ECa233 Increased the Amplitude of eEPSCs in ITC.** To study the effect of ECa233 on the eEPSCs, we continuously recorded the eEPSCs in ITC neurons every 2 min for 1 h. We found that the eEPSCs gradually increased 10 min after initiation of bath application of ECa233 (Figure 5). The maximal response was noted approximately at 15 min. Upper traces showed the examples of the eEPSCs recorded at 0, 10, 15, 20, 30, 40, and 60 min after bath application of 1 μg/ml ECa233 compared with the application of PBS. All of the solutions contained bicuculline 10 μM.
3.4. Concentration-Response Curve of ECa233. To characterize the excitatory effects of ECa233 on ITC neurons, the concentration-response curve of ECa233 was obtained by testing the effects of different concentrations of ECa233 (Figure 6). ECa233 (0, 0.03, 0.1, 0.3, 1.0, and 3.0 μg/ml) generated a concentration-dependent enhancement of EPSCs in ITC neurons. The estimated maximal increase of eEPSCs was 120%, and the estimated value of EC50 was 0.25 μg/ml. Effects of ECa233 were completely abolished after being washed by PBS for 30 min. Example tracing of eEPSCs before and after a bath application of ECa233 (1 μg/ml) for 15 min was shown in the inset of Figure 6. All of the solutions contained 10 μM bicuculline.

3.5. Effect of ECa233 in the Presence of Blockers of Fast Synaptic Transmission. We examined the ECa233-induced increase of eEPSC amplitudes in ITC neurons in the presence of various blockers of synaptic transmission. Bicuculline, a GABA<sub>A</sub> receptor antagonist, was used to exclude the interference of IPSCs. In the presence of bicuculline, 1 μg/ml ECa233 enhanced the eEPSCs in ITCs to 151.19 ± 17.11% suggesting that the enhancement was mediated via fast synaptic glutamatergic transmission. To determine whether the excitatory effect of ECa233 on ITC neurons was mediated by AMPA or NMDA receptors, eEPSCs were recorded in the presence of bicuculline + CNQX or bicuculline + APV. It was found that bicuculline + APV failed to block the effect of ECa233 (160.69 ± 15.23%) on eEPSCs, whereas bicuculline + CNQX completely abolished this effect (102.81 ± 17.16%), suggesting involvement of AMPA/kainate receptors in the facilitation of eEPSCs in ITC neurons by ECa233 (Figure 7).
3.6. Postsynaptic Targets of ITC Neurons. We examined further the postsynaptic targets of biocytin filled ITC axon terminal. The result showed that ITC neurons had medium ovoid cell bodies, largely bipolar dendritic trees, and axons that send collaterals into several areas such as LA, BLA, ITC, main nucleus of ITC (IN), and amygdalostriatal transition area (Astr) and mostly in CeA (Figure 8).

4. Discussion

In the present study, bath application of ECa233 significantly increased evoked EPSCs of ITCs up to 120%. Further experiments conducted in the presence of a series of antagonist(s) suggested that modulation of AMPA receptors was responsible for the increased evoked EPSPs observed and ultimately leading to anxiolytic activity in animal models. To our knowledge, this is the first report to demonstrate a modulation of ITCs of amygdala as possible mechanism underlying anxiolytic activity of a natural product, the standardized extract of *Centella asiatica* ECa233 in animal models. Furthermore, our findings could possibly suggest the prominent role played by ITCs in fear extinction as a potential pharmacological treatment of anxiety as well.

Due to the lack of supply of GAD67-GFP knock-in mice in Thailand and our findings that the morphological and electrical properties including active and passive membrane properties of specific cluster of neurons in GAD67-GFP knock-in and C57BL/6J mice were not different [14, 15], we conducted experiments using C57BL/6J mice to study the effect of ECa233, a natural extract of *Centella asiatica*, on ITCs neurons.

Stimulation of ITCs was made by a concentric bipolar electrode placing on external capsule encompassing a number of fibers from various nuclei including mPFC-ITCs pathway responsible for anxiolytic-like behavior seen in elevated plus maze and open field test [7]. Recently, there were numerous studies of fear extinction-based therapies for the treatment of anxiety disorder focusing on the mPFC-ITCs pathway. The density of mPFC fibers innervated the ITCs is a subject of controversy. Several laboratories indicated that the mPFC provides a direct innervation to ITCs [3, 16]. However, Strobel and their coworker showed the di-synaptic of mPFC-ITCs via the BLA [4]. Therefore, at the moment, anxiolytic activity of ECa233 acting on ITCs may or may not involve mPFC-ITCs pathway. Synaptic inputs of ITC neurons coming from the external capsule are glutamatergic synapses similar to the result of main intercalated cells [14], suggesting that long term depression and potentiation can occur in the ITCs. It is well known that CeA promotes the anxiogenic-like behaviors. For example, the injection of muscimol was found to impair the avoidance of rats to the open arms of the elevated plus maze [17]. So, the strongest feed forward inhibition of ITCs to CeA should present the anxiolytic activity. ECa233 increased the eEPSCs in ITCs via the AMPA receptors as mentioned in this experiment. The finding that brain slices had to be perfused with ECa233 in 5–10 min before the effects were detected suggested the possibility that ECa233 might increase the phosphorylation of AMPA receptor. Previous study of our group demonstrated the upregulation of ERK1/2 and PI3K/Akt signaling pathways after ECa233 exposure [11]. Increased PI3K/AKT signaling was associated with increased expression of AMPA receptors in the surface membrane [18]. In addition, superfusion of ECa233 for 10 min in brain slice preparation increased the long-term potentiation and the BDNF protein expression in hippocampus (unpublished observation). Unfortunately, in the present study, no attempt is made to probe for a presynaptic or postsynaptic site of action for ECa233. This topic should be further investigated in the future.

In line with other studies, the collateral axon of ITCs in the present study mainly innervates CeA [2, 15] suggesting that facilitation of ITCs activity by ECa233 could therefore lead to an increased GABAergic inhibition of CeA. GABA is known as major inhibitory neurotransmitter of the brain and dysfunction of GABAergic neurotransmission causes neurological disorder. Thus, activities of ECa233 on other neuronal circuitry using GABA are plausible and await proof of evidence.

Taken all together, it is highly likely that ECa233 modulated the AMPA receptor and consequently activated ITC neuron to release GABA neurotransmitter in CeA, which
then suppressed the CeA neuronal activity. Thus, it is possible that ECa233 acutely reduced the anxiety behavioral responses in animal models by increasing the GABA release in CeA via modulation of ITC neurons. Similar results were previously reported on neuregulin-1, the neurotrophic factor that could u-regulate AMPA receptor in GABAergic neurons and hence explained its benefit in the treatment of mental disorder [19].

5. Conclusions

The present study demonstrated the mechanism of anxiolytic activity of ECa233, a standardized extract of *Centella asiatica*, on ITCs of the amygdala. ECa233 facilitates the AMPA receptor-mediated excitatory synaptic inputs in ITCs. Since most of the ITC axons innervated CeA. We suggest that increasing the excitatory responses of ITCs by ECa233 would promote the GABA release in the CeA suppressing the output signals from the amygdala.

Disclosure

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Conflicts of Interest

The authors declare that they have no conflicts of interest.

Acknowledgments

This research was supported by the 90th Anniversary (Ratchadapiseksomphot Endowment Fund), Postgraduate Scholarship for Research Aboard (G-RSAB) of Chulalongkorn University Fund, and NIPS Internship 2010.

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