Norepinephrine and E139 Interactions on Epileptiform Activity in the Rat Hippocampus in vitro

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Abstract
Objectives: We tested if E139, an anticonvulsant enaminone, interacts with norepinephrine (NE) to suppress population responses and chemically induced in vitro seizures in the rat hippocampus. Materials and Methods: Evoked field population spikes (PS) were recorded in the hippocampal CA1 area, and in vitro seizures were generated chemically using the zero Mg 2+ model. Results: Low concentrations of E139 (≤10 μM) reversibly inhibited PS amplitude while high concentrations (≥100 μM) enhanced them. For example, E139 (10 μM) depressed the PS amplitude by –23.9 ± 2.3%, while 1 mM caused an enhancement. NE also depressed the PS by –34.5 ± 6.0% and prevented E139 from subsequently depressing the PS amplitude. UK 14304, a selective α2-adrenoceptor agonist, also depressed the PS amplitude by –32.6 ± 9.4% and occluded E139 suppression. NE suppression of PS was blocked by phentolamine and yohimbine which also blocked the effect of E139. Prazosin, a selective α1-adrenoceptor antagonist, did not block NE (–24.8 ± 6.9%) or E139 (–29.7 ± 6.1%) effects. Zero Mg 2+ buffer transformed a single PS to multiple spikes (MS; 3–8 spikes) and also induced spontaneous bursts (SB; 5–20/min). NE suppressed the number of MS from 5.6 ± 0.3 to 3.8 ± 0.2. At its peak effect, E139 was able to further suppress the number of MS to 3.0 ± 0.3. Yohimbine did not change the number of MS but blocked the NE- and E139-induced suppression of MS. SB frequency was suppressed by NE (–60.8 ± 11.7%) which occluded E139 effects. Finally, SB were reversibly abolished by yohimbine (–94.5 ± 11.7%). Conclusion: E139 suppressed population responses and in vitro epileptiform activity by both adrenergic and non-adrenergic mechanisms.

Introduction

Structural analogues of some anilino-enaminones have demonstrated in vivo anticonvulsant actions [1, 2] which are now clearly supported by in vitro evidence at the cellular and network levels [3–5]. Multiple mechanisms of action may underlie these actions of anilino-enaminones to suppress seizures. Based on electrophysiological recordings and occlusion experiments, they were postulated to enhance extracellular γ-aminobutyric acid (GABA) [3]. Pharmacological evidence has also been published that shows that enaminones directly suppress postsynaptic sodium currents [4] and modulate adrenergic antiseizure actions [6]. In the latter case, E139, a prototypical anilino-enaminone, was reported to act as a pu-
tative agonist of α2-adrenoceptors through which it enhanced extracellular GABA levels. Norepinephrine (NE), the main endogenous ligand for these receptors in the central nervous system, is reported to have both convulsant and anticonvulsant properties [7–15]. These proconvulsant and anticonvulsant effects of NE are thought to depend on the receptor subtype that is activated [12, 16] and possibly the extracellular concentration of NE [17, 18]. The majority of reports appear to support an anticonvulsant role of NE. For example chemical depletion of NE by 6-hydroxydopamine is known to facilitate kindling [19, 20]; genetically engineered mice lacking dopamine β-hydroxylase, an enzyme responsible for converting dopamine to NE, have increased vulnerability to seizure stimuli [10], and knockout mice lacking the NE transporter are more vulnerable to seizures [14]. Pharmacological studies largely support this anticonvulsant role of NE [12, 21–23]. However, proconvulsant actions of NE have also been reported using various research approaches [16, 17, 24, 25].

By recording whole-cell currents from single cells, we recently reported that NE-induced depression of excitatory synaptic transmission occurred that induced by E139 [6] suggesting that E139 may employ mechanisms similar to NE to produce its reported in vivo anticonvulsant effects. In a recent study, we also showed that E139 suppresses electrically and chemically induced epileptiform activity recorded in rat hippocampal slices [5]. In this study, we investigated if, similar to the above interaction at the single-cell level, E139 employed adrenergic mechanisms to suppress population and network responses in the hippocampus that are more representative of seizures.

Materials and Methods

Animal Experiments

All animals used in this study were male Sprague-Dawley rats supplied by Kuwait University Animal Resource Centre. All experiments were done in accordance with guidelines on humane handling of experimental animals as established by the Canadian Council on Animal Care. The procedures were designed to minimize animal suffering, and the smallest number of animals necessary to produce the required results was used.

Slice Preparation

Extracellular electrophysiological experiments were performed in coronal hippocampal slices generated from rats (75–150 g) using previously published techniques and methods [26]. Briefly, each rat was deeply anaesthetized with halothane and killed by quick decapitation. The brain was quickly removed and placed in ice-cold (4°C) artificial cerebrospinal fluid (aCSF) bubbled continuously with 95% O2 and 5% CO2 (carbogen). Coronal slices (400 µm) of the forebrain containing the hippocampus were cut from a block of brain tissue in ice-cold aCSF using an Electron Microscopy Sciences (Hatfield, Pa., USA) OTS–4000 tissue slicer. Prior to recording, slices were incubated for 1 h in aCSF that was continuously bubbled with carbogen at room temperature. Most cortical and basal brain tissues were carefully trimmed off slices, and the remaining block containing the hippocampal formation was suspended on a nylon mesh in a 500-μl capacity recording chamber. The bath temperature was tightly maintained at 29–31°C to ensure that changes in responses were not due to variation in temperature [27]. Slices were perfused at a flow rate of 2–3 ml/min with aCSF that was bubbled with carbogen. An extracellular field recording glass electrode (3–15 MΩ) filled with 3 M NaCl was placed in the stratum pyramidale of area CA1, and a bipolar stimulating electrode was placed in the stratum radiatum near area CA1 to activate Schaffer collateral/commissural fibres.

Population Spikes and Chemically Induced in vitro Seizures

The composition of the aCSF used for dissection, storage and population spike (PS) recordings was (in mM): 120 NaCl, 3.3 KCl, 1.2 MgSO4, 1.3 CaCl2, 1.23 NaHPO4, 25 NaHCO3 and 10 d-glucose. First, a single PS was recorded by stimulating the appropriate afferents and moving the field recording electrode around until an optimal PS could be recorded. For recording of epileptiform-type multiple PS and spontaneously occurring epileptiform-type activity referred to as spontaneous bursts (SB), the zero Mg2+ model employed by Klapstein and Colmers [26] was used. We chose this model because, in our laboratory, it was more reliable than picrotoxin in generating in vitro seizure-like activity [5]. Furthermore, the synaptic and anticonvulsant actions of NE have been reported in some studies to be independent of the GABAergic system [18, 28]. In this model, the omission of Mg2+ from the buffer results in the removal of the voltage-dependent Mg2+ block of N-methyl-D-aspartate receptors [29]. This transformed a single PS following one afferent stimulation to multiple spikes [5, 30, 31]. Furthermore, SB that are thought to represent interictal events also accompany the multiple spikes (MS). Briefly, the perfusing solution was changed to zero Mg2+ aCSF (aCSF with the MgSO4 omitted), and then the slice was monitored over a period of 10–20 min for the development of epileptiform-type multiple PS following stimulation and SB (independent of stimulation). All control responses were monitored and shown to be stable for at least 30 min prior to the application of a drug.

Data Acquisition, Analysis and Statistics

All recordings were made using an Axopatch 1D amplifier and pClamp software (Clampex 8; Molecular Devices, Sunnyvale, Calif., USA) in current clamp mode at sampling rates of 50 kHz, filtered at 10 kHz, digitized and stored for off-line analysis. For PS, each stored trace was an average of 5 successively triggered responses elicited at 10-second intervals. The amplitude of the PS was measured from the peak of the positive going wave to the tip of the negative going wave. As well, the initial rising slope (from zero to just before the compound action potential) was also analysed in some experiments. For quantifying epileptiform activity, the number of spikes occurring following the stimulation was counted, and the number of SB occurring in 1 min, recorded in gap-free mode, was counted and used as the frequency. The duration of SB was measured and the number of spikes in each SB was
counted. Drugs were perfused for 5–30 min depending on the experiment, and these are specified in the results section. All data are expressed as mean ± standard error. Statistical significance of all measures was determined using Kruskal-Wallis multiple comparison followed by Student’s test (paired or unpaired where appropriate) and was considered significant at \( p \leq 0.05 \) using Sigmastat® (Systat Software Inc., San Jose, Calif., USA). PS amplitudes were normalized by taking the mean of 4–5 responses prior to drug application and dividing the rest of the values by this mean. These values were used for average plots and bar graphs. All graphical representations were done using Sigmaplot® (Systat Software Inc.), and Coreldraw® (Corel Corp., Ottawa, Canada) softwares.

Chemicals and Drugs

The enaminone E139 was synthesized in house, characterized [1, 32] and dissolved in dimethyl sulphoxide. Stock solutions of 10 mM were prepared, aliquoted and stored at −20°C and used within 3 weeks. All routine laboratory salts and tetrodotoxin (TTX), Norepinephrine and E139 Interactions Med Princ Pract 2008;17:365–372

\[ \text{E139, 100 M} \]

centrations below 10 nM of E139 were tested on the amplitude of the PS. At concentrations just above 100 nM with a peak depression of –23.9 ± 2.3% \((p < 0.05; n = 6)\) recorded at 10 ?M (fig. 1a–c). Analysis of the initial rising slope of these same responses revealed that E139 (10 μM) caused a depression of –26.4 ± 8.1% \((p < 0.05; n = 6; \text{fig. 1d})\). At 100 μM, the effects of E139 were mixed resulting in an average enhancement in the amplitude of the PS (fig. 1c). An even bigger enhancement was recorded at 1 mM (60.1 ± 35.6%; \( n = 3; \text{fig. 1c} \)). The effect of NE (100 μM) on the PS amplitude and on the initial rising slope were also tested. Similar to 10 μM E139, 100 μM NE depressed these by –34.5 ± 6.0 and –44.8 ± 7.6%, respectively \((p < 0.05; n = 8; \text{fig. 2a, b and fig. 1d})\). This effect of NE showed substantial recovery (>90%) following 10–20 min washout of NE. The NE-induced PS depression occluded E139 depression since in the presence of NE, 10 μM E139 no longer produced a depression (11.4 ± 10.0% \( p > 0.05 \) compared to the baseline amplitude in the presence of NE; \( n = 8; \text{fig. 2a, b} \)). UK 14304 (1 μM), a selective α2-adrenoceptor agonist, also depressed the PS amplitude (–32.6 ± 9.4%; \( p < 0.05; n = 6; \text{fig. 2b} \)) to a similar extent as 100 μM NE. It also occluded the ability of E139 to depress the PS amplitude (1.3 ± 0.5%; \( p > 0.05 \)) compared to the baseline amplitude in the presence of UK 14304; \( \text{fig. 2b} \)). To further characterize this E139/NE interaction, we performed antagonism experiments. Pretreatment of slices with prazosin (10 μM), an α1-adrenoceptor antagonist, did not prevent E139 (–24.8 ± 6.9%; \( p < 0.05; n = 5; \text{fig. 3a} \)) from depressing the PS amplitude, neither did it prevent NE from depressing the PS amplitude (–29.7 ± 6.1%; \( p < 0.05; n = 5; \text{fig. 3a} \)). By contrast, phentolamine (10 μM), a non-selective α-adrenoceptor antagonist, blocked both the NE-induced (5.7 ± 8.2%; \( p > 0.05; n = 5; \text{fig. 3b} \)) and E139-induced (5.1 ± 5.9%; \( p > 0.05; n = 6 \)) PS depression (fig. 3b).

Results

The results reported in this study were obtained from a total of 95 field recordings performed in the area CA1 of the rat hippocampus. Each slice was used only once. PS and epileptiform activities were all sensitive to TTX treatment applied at the end of selected experiments.

Effects of E139 and NE on PS Recorded in Area CA1 of the Hippocampus

The effects of various concentrations (1 nM to 1 mM) of E139 were tested on the amplitude of the PS. At concentrations below 10 μM, E139 consistently depressed the amplitude of the PS. The threshold of PS depression was just above 100 nM with a peak reduction of –23.9 ± 2.3% (\( p < 0.05; n = 6 \)) recorded at 10 μM (fig. 1a–c). Analysis of the initial rising slope of these same responses revealed that E139 (10 μM) caused a depression of –26.4 ± 8.1% (\( p < 0.05; n = 6; \text{fig. 1d} \)). At 100 μM, the effects of E139 were mixed resulting in an average enhancement in the amplitude of the PS (fig. 1c). An even bigger enhancement was recorded at 1 mM (60.1 ± 35.6%; \( n = 3; \text{fig. 1c} \)). The effect of NE (100 μM) on the PS amplitude and on the initial rising slope were also tested. Similar to 10 μM E139, 100 μM NE depressed these by –34.5 ± 6.0 and –44.8 ± 7.6%, respectively (\( p < 0.05; n = 8; \text{fig. 2a, b and fig. 1d} \)). This effect of NE showed substantial recovery (>90%) following 10–20 min washout of NE. The NE-induced PS depression occluded E139 depression since in the presence of NE, 10 μM E139 no longer produced a depression (11.4 ± 10.0% \( p > 0.05 \) compared to the baseline amplitude in the presence of NE; \( n = 8; \text{fig. 2a, b} \)). UK 14304 (1 μM), a selective α2-adrenoceptor agonist, also depressed the PS amplitude (–32.6 ± 9.4%; \( p < 0.05; n = 6; \text{fig. 2b} \)) to a similar extent as 100 μM NE. It also occluded the ability of E139 to depress the PS amplitude (1.3 ± 0.5%; \( p > 0.05 \)).

Effect of E139 and NE on Chemically Induced MS and SB

In this in vitro model of seizure that targets and enhances glutamate-mediated excitatory transmission, when Mg2+ was removed from the perfusing buffer, the voltage-dependent Mg2+ block of N-methyl-D-aspartate receptor was removed [29] leading to MS and SB. After recording in zero Mg2+ buffer for 20 min, a single PS was transformed into MS (5.6 ± 0.3; \( n = 6; p < 0.05; \text{fig. 4a} \)) in response to a single electrical stimulation of the afferents. Similar to our previous report on E139 [5], NE (100 μM) reversibly reduced the number of MS from 5.6 ± 0.3 to 3.8 ± 0.2 (–29.6 ± 5.9%; \( p < 0.05; \text{paired t test}; n = 6; \text{fig. 4a, b} \)). The amplitude of the first spike was also reduced by –28.1 ± 6.0% (\( p < 0.05; \text{paired t test}; n = 6 \)) in the presence of NE. When E139 (10 μM) was applied at the peak of the NE effect, E139 produced a further decrease in the number of MS to 3.0 ± 0.3 (–28.2 ± 1.8%; \( p < 0.05; \text{paired t test}; n = 6; \text{fig. 4a, b} \)) without affecting the amplitude of the first spike (–6.2 ± 11.7%; \( p > 0.05; n = 6; \text{fig. 4a} \)). Yohimbine (1 μM) which by itself did not affect the number of MS (4.2 ± 0.2 in control vs. 4.2 ± 0.4 in yohimbine; \( p > 0.05; n = 5; \text{fig. 4c} \)) blocked the effect of NE (3.2 ± 0.2; \( p > 0.05; \text{fig. 4c} \)).
Fig. 1. E139 depresses evoked field PS in the CA1 region of the hippocampus in a concentration-dependent manner. a Sample PS recorded in the cell body layer of the area CA1 of the hippocampus in control, presence of 10 μM E139 and following 10 min washout of E139. b An average time-effect plot showing the effect of E139 on the PS amplitude obtained from 6 field recordings. c Concentration-response bar graph showing that E139 has a biphasic concentration-dependent effect on PS whereby concentrations of 10 μM and below produce depression while 100 μM and above enhance PS. d Average bar graph showing that the effects of E139 (10 μM) and NE (100 μM) on PS amplitude (ampl.) were similar to their effects on synaptic response as measured by the slope of the initial rising phase of the PS. * p < 0.05. EPSP = Excitatory postsynaptic potential. In this and all other figures, the number above/below each bar represents the number of slices to which each concentration/treatment was applied.

Fig. 2. NE inhibits PS amplitude that occludes E139 effects. a An average time-effect plot of the effect of NE (100 μM) and that of E139 (10 μM) in the continued presence of NE recorded in 8 slices. b Bar graphs summarizing the effects of 10 μM E139 on PS amplitude (from fig. 1) and effects of NE alone and in combination with E139. Finally shown on the right of the graph are bars showing a similar inhibitory effect by UK 14304, a potent selective α1-adrenoceptor agonist, on PS amplitude and occlusion of the E139 effect. * p < 0.05 compared to control.
**Fig. 3.** NE- and E139-induced PS inhibitions are blocked by α-adrenoceptor antagonists. **a** Summary bar graphs showing that both NE (n = 5) and E139 (n = 5) are able to inhibit PS amplitude in the presence of prazosin (Praz), a selective α1-adrenoceptor antagonist. *p < 0.05. **b** By contrast to **a**, phentolamine (Phent; n = 5), a non-selective α-adrenoceptor antagonist, and yohimbine (Yoh; n = 6), a selective α1-adrenoceptor antagonist, both blocked the NE- and E139-induced PS amplitude inhibition. Also shown here is the effect of TTX (1 μM; n = 5) on PS.

**Fig. 4.** NE inhibits multiple PS frequency that does not occlude the effect of E139. **a** Sample traces of a PS (first) and multiple PS (next 3 traces) induced by zero Mg²⁺ buffer and the effect of NE (100 μM) alone and in the presence of E139 (10 μM) on the frequency of multiple PS. Note the depression of the first spike by NE without any further depression by E139 and the total elimination of the spikes by TTX. **b** Summary bar graphs (n = 6) showing the effect of zero Mg²⁺ on the number of spikes triggered by a single afferent stimulation and the effect of NE alone and subsequently NE + E139. *p < 0.05 compared to control, †p < 0.05 compared to zero Mg²⁺, ‡p < 0.05 compared to zero Mg²⁺ + NE. Note that the effect of E139 alone on PS numbers has been inserted for comparison (modified from Ananthakumar et al. [5]). **c** Summary bar graph (n = 5) showing that yohimbine (Yoh) blocked NE- and E139-induced depression of the frequency of MS.
Furthermore it also blocked the E139 effect on the number of MS when it was applied in the presence of NE (3.6 ± 0.3; p > 0.05; n = 5; fig. 4c). An application for 15–20 min of zero Mg²⁺ buffer induced SB that ranged in frequency from 5 to 20/min yielding a mean of 15.4 ± 3.3/min (p < 0.05; n = 5; fig. 5a–d). NE (100 μM) reduced this frequency to 6.8 ± 2.1/min, a depression of –60.8 ± 11.7% (p < 0.05; n = 5; fig. 5a, c). Application of E139 at the peak of this NE effect did not produce any further reduction in SB frequency (16.2 ± 12.7%; p > 0.05; n = 5; fig. 5a, c). SB had an average duration of 108.7 ± 9.3 ms and contained 9.9 ± 1.6 spikes (p < 0.05; n = 10). Further analyses of SB in the presence of E139 and NE revealed that they suppressed both the duration of SB (–57.1 ± 18.1 and –66.1 ± 11.3%, respectively) and the number of spikes in each SB (–74.0 ± 7.5 and –55.8 ± 7.5%, respectively; p < 0.05, n = 5 each) to a similar extent.

In another set of experiments to verify the pharmacology of the effects of NE and E139 on SB, bath application of yohimbine (1 μM) almost completely abolished the SB frequency, reducing it from 12.8 ± 1.7 to 0.7 ± 0.8/min, a suppression of –94.5 ± 11.7% (p < 0.05; n = 6; fig. 5b, d). This effect was about 85% reversible upon washing out of yohimbine (fig. 5b, d).

**Discussion**

In this study, we investigated a possible interaction between E139 and NE on population responses in a chemically induced in vitro seizure model. Our data on field-recorded PS whereby both NE and E139 suppressed the amplitude that mutually occluded each other are consistent with our previous finding in single-cell studies [6]. Analyses of the synaptic component (initial rising phase) of the PS indicate that suppression of this phase contributed the most to the entire PS suppression since for both

**Fig. 5.** NE inhibits SB frequency that occludes the effect of E139. a Sample traces showing SB induced by zero Mg²⁺ and the effect of NE alone and E139 in the presence of NE. All SB were eliminated by TTX (1 μM). b Sample traces showing SB induced by zero Mg²⁺ and the effect of Yoh (1 μM) on the SB. c Summary bar graphs (n = 5) showing the effect of zero Mg²⁺ on the number of SB and the effect of NE alone and, subsequently, NE + E139. Note that the effect of E139 alone on SB/min has been inserted for comparison (modified from Ananthalakshmi et al. [5]). d Summary bar graph (n = 6) showing that yohimbine (Yoh) reversibly suppresses zero-Mg²⁺-induced SB. *p < 0.05 compared to zero Mg²⁺.
NE and E139, the latter suppression was not significantly different from the suppression of the initial rising phase. This suggests that the compound action potential is tightly coupled to the synaptic response that drives the action potentials. Furthermore, the pharmacology of these actions (mimicry by UK 14304 and blockade by phentolamine and yohimbine) also point to the involvement of a similar mechanism in suppressing the population spike in the hippocampus, i.e. both E139 and NE appear to produce their suppressant effects through action on \( \alpha_2 \)-adrenoceptors which have been reported to mediate the anticonvulsant effects of NE \([12, 18, 33, 34]\). Consistent with previous reports on the anticonvulsant actions of NE \([16, 18, 35–37]\), it suppressed the MS frequency and SB. Interestingly, unlike their mutually occlusive actions on PS amplitude, the presence of NE did not prevent E139 from further suppressing the MS frequency. This effect of E139 in the presence of NE may be due to the fact that E139, in addition to employing adrenergic mechanisms to suppress neuronal activity, also has direct effects on sodium channels \([4]\). Thus, the non-adrenergic component of this action of E139 may be related to its effect on postsynaptic sodium channels.

Similar to its blockade of NE and E139 effects on PS, yohimbine also blocked the MS frequency suppression by NE and E139. Also, the suppression of SB by NE occluded that of E139 further implicating an adrenergic mechanism. Finally, contrary to expectation, yohimbine almost completely wiped out the SB suggesting that NE, possibly acting on a subtype of \( \alpha_2 \)-adrenoceptor \([16, 34, 38]\), may promote epileptogenesis, i.e. have proconvulsant effects. Pro- and anticonvulsant effects of NE acting on \( \alpha_2 \)-adrenoceptors have been reported to be mediated by distinct populations of the same subtype of \( \alpha_2 \)-adrenoceptors \((\alpha_2 A)\) located on presynaptic or postsynaptic sites \([16]\). Alternatively, the apparent anticonvulsant effect of yohimbine observed here may be due to its reported partial agonist action on some serotonin receptors \([39]\). Such a postulated serotoninergic action of yohimbine on epileptiform activity in the hippocampus needs to be critically examined in future studies.

**Conclusion**

The results of this study reveal that E139, a prototypical anilino-enaminone, suppresses chemically induced in vitro epileptiform activity via both adrenergic and non-adrenergic mechanisms. These findings support the hypothesis that E139 may employ multiple mechanisms to produce its anticonvulsant effects.

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**References**


