Nanomolar Bifenthrin Alters Synchronous Ca$^{2+}$ Oscillations and Cortical Neuron Development Independent of Sodium Channel Activity

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ABSTRACT

Bifenthrin, a relatively stable type I pyrethroid that causes tremors and impairs motor activity in rodents, is broadly used. We investigated whether nanomolar bifenthrin alters synchronous Ca$^{2+}$ oscillations (SCOs) necessary for activity-dependent dendritic development. Primary mouse cortical neurons were cultured 8 or 9 days in vitro (DIV), loaded with the Ca$^{2+}$ indicator Fluo-4, and imaged using a Fluorescence Imaging Plate Reader Tetra. Acute exposure to bifenthrin rapidly increased the frequency of SCOs by 2.7-fold (EC$\text{_{50}}$ = 58 nM) and decreased SCO amplitude by 36%. Changes in SCO properties were independent of modifications in voltage-gated sodium channels since 100 nM bifenthrin had no effect on the whole-cell Na$^{+}$ current, nor did it influence neuronal resting membrane potential. The L-type Ca$^{2+}$ channel blocker nifedipine failed to ameliorate bifenthrin-triggered SCO activity. By contrast, the metabotropic glutamate receptor (mGluR)5 antagonist MPEP [2-methyl-6-(phenylethynyl)pyridine] normalized bifenthrin-triggered increase in SCO frequency without altering baseline SCO activity, indicating that bifenthrin amplifies mGluR5 signaling independent of Na$^{+}$ channel modification. Competitive [AP-5; (–)-2-amino-5-phosphonopentanoic acid] and noncompetitive (dizocilpine, or MK-801 [(5S,10R)-(–)-5-methyl-10,11-dihydro-5H-dibenzo[a,d]cyclohepten-5,10-imine maleate]) N-methyl-D-aspartate antagonists partially decreased both basal and bifenthrin-triggered SCO frequency increase. Bifenthrin-modified SCO rapidly enhanced the phosphorylation of Ca$^{2+}$-binding protein (CREB). Subacute (48 hours) exposure to bifenthrin commencing 2 DIV–enhanced neurite outgrowth and persistently increased SCO frequency and reduced SCO amplitude. Bifenthrin-stimulated neurite outgrowth and CREB phosphorylation were dependent on mGluR5 activity since MPEP normalized both responses. Collectively these data identify a new mechanism by which bifenthrin potently alters Ca$^{2+}$ dynamics and Ca$^{2+}$-dependent signaling in cortical neurons that have long term impacts on activity driven neuronal plasticity.

Introduction

Synthetic pyrethroid insecticides were introduced into widespread use for the control of insects and disease vectors more than three decades ago. Pyrethroids account for approximately one-fourth of the worldwide insecticide market, and the use of these compounds is increasing (Schleier and Peterson, 2011). Based on both their chemical structures and biologic responses to acute exposure, pyrethroids are classified into two major groups: type I and type II. Type I pyrethroids lack a cyano group at the a carbon of the 3-phenoxybenzyl alcohol moiety and produce hyperexcitation, tremors, and convulsions (T syndrome), whereas type II pyrethroids have a cyano group at the a carbon and produce hypersensitivity, choreoathetosis, salivation, and seizures (CS syndrome). A few pyrethroids producing both tremors and salivation were classified as type I/II (Soderlund, 2012; Casida and Durkin, 2013). The primary molecular target underlying insecticidal activity and presumably mammalian neurotoxicity of pyrethroids are voltage-gated sodium channels (VGSCs) (Soderlund, 2012; Casida and Durkin, 2013). Pyrethroids enhance sodium channel activity by shifting activation to more negative membrane potentials as well as by slowing channel inactivation. This action results in altered neuronal excitability characterized by in vitro and in vivo changes in neuronal firing rates (e.g., repetitive firing or depolarizing block of the neuron). In addition to their actions on VGSCs, pyrethroids interact with other types of ion channels in a variety of in vitro systems, including altering the function of voltage-gated calcium (Ca$^{2+}$)

ABBREVIATIONS: CREB, cAMP response element–binding protein; Cl, confidence interval; CNQX, 6-cyano-7-nitroquinazoline-2,3-dione; d-AP-5, d-[(–)-2-amino-5-phosphonopentanoic acid; DAPI, 4’,6-diamidino-2-phenylindole; DIV, days in vitro; mGluR, metabotropic glutamate receptor; MK-801, (5S,10R)-(–)-5-methyl-10,11-dihydro-5H-dibenzo[a,d]cyclohepten-5,10-imine maleate; MPEP, 2-methyl-6-(phenylethynyl)pyridine; PBS, phosphate-buffered saline; RyR1, ryanodine receptor type 1; SCO, synchronous Ca$^{2+}$ oscillation; VGSC, voltage-gated sodium channel.
channels and GABA<sub>α</sub> receptors (Clark and Symington, 2012; Soderlund, 2012; Casida and Durkin, 2013). It should be noted that all these actions invariably require exposure to pyrethroid a concentrations ≥ 1 μM.

Bifenthrin (Fig. 1) is one of the most potent commercialized pyrethroids to date, with LD<sub>50</sub> values of about 53 mg/kg in female rats and 70 mg/kg in male rats after oral gavage (Wolansky et al., 2006; Wolansky and Harrill, 2008). Bifenthrin, as low as 10 mg/kg, produces type I pyrethroid-like episodes of head and whole-body shakes, a strong and prolonged tremorogenic response, hyperthermia, vocalizations, prostration, and, at doses > 28 mg/kg, it eventually triggers convulsions and death (Holton et al., 1997; Wolansky et al., 2007). This dose range corresponds to a brain concentration of bifenthrin of about 550–650 ng/g of brain tissue (Scollon et al., 2011). The high-dose effect is consistent with pyrethroid-induced perturbation of voltage-dependent Na<sup>+</sup> and Ca<sup>2+</sup> fluxes at micromolar concentrations in primary cultured cortical neurons (Cao et al., 2011a,b). However, at a threshold dose of 1.28 mg/kg, which corresponds to a brain concentration of roughly 30 ng/g of brain tissue, bifenthrin decreases motor activity in rats (Wolansky et al., 2006; Scollon et al., 2011). These data suggest that bifenthrin can alter central nervous system physiology at doses much lower than those needed to induce tremors.

Extensive studies have been performed to elucidate the molecular mechanism and cellular effect of pyrethroids in cultured neurons. Previously we have demonstrated that at concentrations >1 μM, 9 of 11 commercially used pyrethroids enhance Na<sup>+</sup> and Ca<sup>2+</sup> fluxes with distinct potencies and efficacies (Cao et al., 2011a,b) by a mechanism that is consistent with their reported actions on VGSC expressed in Xenopus oocytes (Choi and Soderlund, 2006). For example, in mouse cortical neuronal cultures, bifenthrin induces modest Na<sup>+</sup> and Ca<sup>2+</sup> influx at concentrations above 3 μM, which can be eliminated by tetrodotoxin pretreatment, demonstrating that bifenthrin is a low-efficacy sodium channel agonist. Pyrethroids have also been demonstrated to act on Ca<sup>2+</sup>- or Cl<sup>-</sup> channels (Clark and Symington, 2012; Soderlund, 2012); however, no direct evidence has been presented demonstrating that bifenthrin can interact with these channels. Furthermore, the possible effect of pyrethroids on neuronal network electrical activity has been investigated. Both deltamethrin (0.1 μM) and permethrin (0.1 μM) are able to increase electric spike activity in the absence of GABAergic inhibition (Meyer et al., 2008); however, bifenthrin appears without effect, even at a concentration of 50 μM (McConnell et al., 2012).

Murine primary-cultured neurons mature to form synaptic networks and display spontaneous synchronized Ca<sup>2+</sup> oscillations (Cao et al., 2012a). These spontaneous Ca<sup>2+</sup> oscillations are dependent on the generation of action potentials and can be manifested by multiple Ca<sup>2+</sup> signaling pathways (Dravid and Murray, 2004). Synchronous Ca<sup>2+</sup> oscillations (SCOs) are important in mediating neuronal development and activity-dependent dendritic growth (Dolmetsch et al., 1998; Wayman et al., 2006). Genetic or environmental factors that interfere with activity-dependent dendritic growth influence the functioning of neuronal networks (Pessah et al., 2010; Stamou et al., 2013).

In the present study, we identified a novel mechanism by which bifenthrin, at nanomolar concentrations, alters the patterns of SCOs in primary mouse cortical neuronal cultures by amplifying glutamatergic glutamate receptor 5 (mGluR5) activity, which alters CREB signaling and patterns of neurite outgrowth in the absence of detectable perturbation of VGSCs. To date, these are the most potent neurotoxic actions attributed to bifenthrin and suggest that bifenthrin at low concentrations relevant to current exposures can persistently alter patterns of SCO and activity-dependent neuronal cell growth and network development.

**Materials and Methods**

Bifenthrin (purity: 99.1%, mix of isomers) was from Chemical Service, Inc. (West Chester, PA) (Fig. 1). Anti-p-CREB, anti-CREB, and anti-microtubule-associated protein 2 antibodies were from Cell Signaling Technology (Danvers, MA). Odyssey Blocking Buffer and IRDye-labeled secondary antibody were from LI-COR Biotechnology (Lincoln, NE). Fluo-4 and Alexa Fluor 488-conjugated goat anti-rabbit secondary antibody were from Life Technologies (Grand Island, NY). Tetrodotoxin, D-AP-5 ([R(-)-2-amino-5-phosphonopentanoic acid], MK-801 [(5S,10R(+)-5-methyl-10,11-dihydro-5H-dibenzo[d,c]cyclopepten-5,10-imine maleate), CNQX (6-cyano-7-nitroquinoxaline-2,3-dione), MPEP [2-methyl-6-(phenylethynyl)pyridine], and nifedipine (3,5-dimethyl 2,6-dimethyl-4-(2-nitrophenyl)-1,4-dihydropyridine-3,5-dicarboxylate) were from Tocris Bioscience (Minneapolis, MN).

**Cortical Neuronal Cultures.** The University of California Davis Animal Use and Care Committee approved this study, and animal experiments were conducted in accordance with the guidelines of the Animal Use and Care of the National Institutes of Health. Cortical neurons were dissociated from the cortex from C57Bl/6J mouse pups at postnatal day 0 to 1 as described previously (Cao et al., 2011b). The dissociated neurons were maintained in Neurobasal complete medium (Neurobasal medium supplemented with NS21 (Chen, et al., 2010), 0.5 mM l-glutamine, and HEPES) with 5% fetal bovine serum. The dissociated cortical cells were plated onto poly-l-lysine-coated (0.5 mg/ml; Sigma-Aldrich, St. Louis, MO) clear-bottom, black-walled, 96-well imaging plates (BD, Franklin Lakes, NJ) at densities of 1 x 10<sup>5</sup>/well for Ca<sup>2+</sup> imaging and 2000/well for immunochemistry, respectively. For the Western blot experiments, the dissociated cortical neurons were plated onto poly-l-lysine–coated six-well plates at a density of 3 x 10<sup>5</sup>/well. The medium was changed twice a week by replacing half the volume of culture medium in the well with serum-free Neurobasal complete medium. The neurons were maintained at 37°C with 5% CO<sub>2</sub> and 95% humidity.

**Measurement of Synchronous Intracellular Ca<sup>2+</sup> Oscillations.** Cortical neurons at 9 days in vitro (DIV) were used to investigate how acute exposure to bifenthrin alters synchronous Ca<sup>2+</sup> oscillations as described previously (Cao et al., 2012a). Briefly, the neurons were loaded with Fluoro-4 for 1 hour at 37°C in Locke’s buffer (in mM: 8.6 HEPES, 5.6 KCl, 154 NaCl, 5.6 glucose, 1.0 MgCl<sub>2</sub>, 2.5 CaCl<sub>2</sub>, and 0.0001 glycine, pH 7.4). After recording the baseline spontaneous Ca<sup>2+</sup> oscillations for 3 minutes, vehicle or bifenthrin was added to the well using a programmable 96-channel pipetting robotic system, and the intracellular Ca<sup>2+</sup> concentration ([Ca<sup>2+</sup>]<sub>i</sub>) was monitored for additional 50 minutes. The calculations for EC<sub>50</sub> for changes in oscillatory frequency and peak transient amplitude were from data acquired.

**Fig. 1.** Chemical structure of bifenthrin.
between 30 and 40 minutes after the additions were made (i.e., averages from a 10-minute period). To test the mechanisms influencing SCO activity, receptor-targeted inhibitors were added to designated wells after the Ca2+ response was induced by bifenthrin (0.1 μM).

To test the chronic influences, exposures of neurons to bifenthrin (0.01–1.0 μM) commenced 24 hours after plating, and on DIV 6, the patterns of SCO measured as described already.

**Membrane Potential Measurement.** A membrane-sensitive dye, FMPlblue, was used to investigate whether bifenthrin can depolarize the membrane potential in neurons that were 8 or 9 DIV. Briefly, the neurons were loaded with 180 μl of FMPlblue (1 vial diluted to 50 ml Lock’s; Molecular Devices, Sunnyvale, CA). After incubation with the dye for 30 minutes, the cells were excited at 510–545 nm, and the fluorescent signals emitted at 565–625 nm were recorded. After recording basal fluorescence for 1 minute, vehicle or bifenthrin (0.01–1.0 μM) was added to the wells, and the signals were recorded for an additional 40 minutes.

**Measurement of Neuronal Na+, K+, and Ca2+ Channel Currents.** Cultured cortical neurons were voltage-clamped by whole-cell patch-clamp using an EPC-10 amplifier and Pulse software (HEKA; Lambrecht/Pfalz, Germany). Cells were bathed in extracellular solution containing (in mM) 160 NaCl, 4.5 KCl, 1 MgCl2, 1 CaCl2, and 10 HEPES; pH was adjusted to 7.4 using NaOH (310 mOsm). Pipettes were pulled from 1.5-mm capillary tubing and filled with intracellular solution containing (in mM) either 145 KF, 2 MgCl2, 10 EGTA, and 10 HEPES; pH was adjusted to 7.2 with KOH (300 mOsm) or 145 CaF, 3 KCl, 2 NaCl, 1 MgCl2, 3 Na-ATP, 0.2 Na-GTP, 10 EGTA, and 10 HEPES (pH adjusted to 7.4 with KOH; 303 mOsm). Pipette-tip resistances were 2–4 MΩ. Series resistances of 3–10 MΩ were compensated 40–90%. All neurons were voltage-clamped to a holding potential of −80 mV. Data analysis, fitting, and plotting were performed with IGOR-Pro (WaveMetrics, Lake Oswego, OR) and Origin 9.0 (OriginLab, Northampton, MA). Na+ currents were elicited by 50-millisecond voltage steps from −80 to 0 mV applied every 10 seconds. Current decay and tail current were fitted with the following equation:

\[
I = A_1 \times \exp(-t/\tau_1) + A_2 \times \exp(-t/\tau_2) + C,
\]

where \(I\) is the current amplitude, \(t\) the time range of current chosen for fitting, \(\tau_1\) the fast time constant of inactivation, and \(\tau_2\) the slow time constant of inactivation. A1 and A2 are the percentages of current inactivating with the fast and slow time constants, respectively, and C is the non-inactivating current component.

K+ currents were recorded by 200-millisecond voltage steps from −80 to 40 mV applied every 10 seconds. L929 cells stably expressing mKv3.1 and mKV1.1 were previously described (Grissmer et al., 1994). Experiments were performed with the same external and internal solutions as used for the cortical neurons.

Bifenthrin’s influence on ryanodine receptor type 1 (RyR1) isolated from rabbit skeletal muscle was assessed in two ways: [3H]ryanodine binding analysis and measurements of ionic current from single internal solutions as used for the cortical neurons. Bifenthrin was previously shown to induce modest Ca2+ influx at concentrations >3 μM (Cao et al., 2011a,b). These studies focused on net Ca2+ influx associated with neuronal excitotoxicity but not possible alternations of patterns of SCO that are important for activity-dependent gene expression, neuronal cell growth, and plasticity. We therefore performed directed studies to investigate whether bifenthrin at submicromolar concentrations influences Ca2+ dynamics and their possible mechanisms. Primary murine cortical neurons grown in 96-well plates were loaded with Fluo-4 at 8 or 9 DIV, and cytoplasmic Ca2+ dynamics were monitored in real-time using FLIPR Tetra. Under these conditions, dissociated neuronal cultures form extensive networks that engage synchronized Ca2+ oscillations (Fig. 2A, top trace) essential for promoting dendritic complexity and activity-dependent plasticity (Chen et al., 2010; Cao et al., 2012a,b). The addition of bifenthrin (0.01–1 μM) produced a concentration-dependent increase in the frequency of SCO having an EC50 value of 57.7 nM (31.9–104 nM, 95% confidence interval (CI)) that attained 270% of baseline at 1 μM (Fig. 2, A, traces 2–8, and B). Coincident with the increasing frequency of

**Results**

**Acute Exposure to Nanomolar Bifenthrin Alters SCO Patterns in Cortical Neurons.** Bifenthrin was previously shown to induce modest Ca2+ influx at concentrations >3 μM (Cao et al., 2011a,b). These studies focused on net Ca2+ influx associated with neuronal excitotoxicity but not possible alternations of patterns of SCO that are important for activity-dependent gene expression, neuronal cell growth, and plasticity. We therefore performed directed studies to investigate whether bifenthrin at submicromolar concentrations influences Ca2+ dynamics and their possible mechanisms. Primary murine cortical neurons grown in 96-well plates were loaded with Fluo-4 at 8 or 9 DIV, and cytoplasmic Ca2+ dynamics were monitored in real-time using FLIPR Tetra. Under these conditions, dissociated neuronal cultures form extensive networks that engage synchronized Ca2+ oscillations (Fig. 2A, top trace) essential for promoting dendritic complexity and activity-dependent plasticity (Chen et al., 2010; Cao et al., 2012a,b). The addition of bifenthrin (0.01–1 μM) produced a concentration-dependent increase in the frequency of SCO having an EC50 value of 57.7 nM (31.9–104 nM, 95% confidence interval (CI)) that attained 270% of baseline at 1 μM (Fig. 2, A, traces 2–8, and B). Coincident with the increasing frequency of
SCO, bifenthrin partially decreased the mean amplitude of the Ca$^{2+}$ oscillations reaching 64% of control at 1 μM (Fig. 2, A and C). The IC$_{50}$ value for bifenthrin suppressing the mean amplitude of the Ca$^{2+}$ oscillations was 83.4 nM (49.8–141.0 nM, 95% CI) (Fig. 2C). To test whether the bifenthrin response on SCO was reversible, the cells were completely washed five times with Lock’s buffer after exposure to 0.1 μM bifenthrin for 30 minutes. The washing the cells itself had no detectable influence on the basal SCO (Supplemental Fig. 1, upper black traces); however, the bifenthrin-augmented pattern of SCO was not fully reversible after washout, suggesting persistent alteration in the pattern of SCO after acute exposure to a low concentration of bifenthrin (Supplemental Fig. 1, lower red traces).

**Nanomolar Bifenthrin Neither Alters Membrane Potential nor Voltage-Gated Currents.** The FMPblue dye was used to examine whether bifenthrin alters the resting membrane potential of cortical neurons. After recording basal FMPblue fluorescence for 1 minute, bifenthrin was added to the well and the signal was recorded for an additional 40 minutes. Bifenthrin, at a concentration that caused a nearly 2-fold increase in SCO frequency (0.1 μM), had no detectable influence on membrane potential (data not shown). As a positive control, application of KCl (10 mM) to the external bath showed that the FMPblue dye was capable of reporting even a modest (14 mV, calculated from Goldman–Hodgkin–Katz equation) shift in membrane depolarization (Fig. 3).

To explore more directly whether bifenthrin-induced changes in Ca$^{2+}$ oscillations might be due to direct effects on voltage-gated sodium channels, we tested the effect of bifenthrin on Na$^+$ currents in cultured cortical neurons using whole-cell patch-clamp. As shown in Fig. 4A, bifenthrin concentrations of 10 and 100 μM drastically slowed inactivation during depolarization, as evidenced by the broadening and incomplete inactivation of the Na$^+$ current elicited by a 50-millisecond depolarizing step from −80 to 0 mV, whereas concentrations of 0.1 or 1 μM had no effect. Bifenthrin further delayed deactivation as shown by the large Na$^+$ tail currents that became apparent after repolarization to −80 mV. This slow-tail current, which constitutes an additional component to the normal fast-tail current evoked on repolarization, was very sensitive to bifenthrin, and 0.1 μM already slightly changed the deactivation time constant $\tau_d$ of its biexponential decay (Fig. 4C).

However, since the physiologic relevance of the tail current is not clear, we fitted separate EC$_{50}$ values (Fig. 4B) for increasing

**Fig. 2.** Bifenthrin-induced Ca$^{2+}$ dysregulation in cortical neurons. (A) Representative traces showing how acute exposure to bifenthrin (0.01–1 μM) influences Ca$^{2+}$ fluctuations in cortical neurons at 8 or 9 DIV as a function of time. Bifenthrin produces a robust increase in the frequency of Ca$^{2+}$ oscillations accompanied by the slightly decreased Ca$^{2+}$ oscillations amplitude. (B) Concentration-response relationships for bifenthrin-stimulated increase in the frequency of Ca$^{2+}$ oscillation. The EC$_{50}$ value for bifenthrin-stimulated increase in the frequency of Ca$^{2+}$ oscillation frequency is 57.7 nM. (C) Concentration-response relationships for bifenthrin-stimulated decrease in the amplitude of Ca$^{2+}$ oscillation. The EC$_{50}$ value for bifenthrin-stimulated decrease in the amplitude of Ca$^{2+}$ oscillation frequency is 83.4 nM. These data were averaged from a time period of between 30 and 40 minutes after the addition of bifenthrin. Each data point represents mean ± S.E.M. (n > 20 well) from five independent cultures. VEH, vehicle.

**Fig. 3.** Bifenthrin has no effect on membrane potential in cortical neurons. Influence of bifenthrin (0.1 μM) or KCl (10 mM, positive control) on the rest membrane potential as a function of time. KCl produces a significant depolarization of the cortical neurons reflected by increased FMPblue fluorescence signals. However, bifenthrin has no response on the resting membrane potential. This experiment was repeated twice each with five replicates with similar results. VEH, vehicle.
the area under the current elicited by depolarization (16.8 ± 0.7 
\mu M) and the area under the tail current (4.4 ± 0.4 
\mu M), assuming the effect at 100 \mu M to be maximal. The more
meaningful of these EC_{50} values representing the bifenthrin-
induced delay in VGSC inactivation is 275-fold higher than the
EC_{50} for increasing SCO frequency. We further tested bifen-
thrin on cortical neuron K^\text{1} currents as well as heterologous
expressed Kv1.1 and Kv3.1, two of the major delayed-rectifier
type K^\text{1} channels expressed in cortical neurons (Grissmer et al.,
1994; Massengill et al., 1997; Goldberg et al., 2008), and found
that 1 \mu M of bifenthrin had absolutely no effect on K^\text{V} currents
(data not shown).

We tested whether bifenthrin-triggered SCO patterns could
be reversed by the addition of the L-type Ca^{2+} channel blocker
nifedipine (10 \mu M). Figure 5, A and B, shows that nifedipine
neither affected basal SCO activity nor the activity modified by
bifenthrin, discounting a possible contribution of L-type Ca^{2+}
channel modulation in eliciting these effects of bifenthrin.

Finally, bifenthrin was also tested for possible influences
toward ryanodine receptor activity as recently reported for
deltamethrin and selected type II pyrethroids (Morisseau
et al., 2009). Bifenthrin (≤1 \mu M) had no detectable influence
on the gating activity of RyR1 channels reconstituted in bilayer lipid membranes, nor did it alter the binding of
[^3]H\text{ryanodine to its high affinity sites (data not shown).}

**mGluR5 Contributes to Bifenthrin-Modified SCO Patterns.** Since SCO activity in cortical neurons was highly
dependent on extracellular Ca^{2+} (data not shown), we investi-
gated whether Glu receptors contributed to bifenthrin-triggered
alterations in SCOs. Although MPEP (1 \mu M), a selective an-
tagont of mGluR5, had no significant effect on basal SCO
patterns, it produced a significant 60% ± 8% amelioration of
bifenthrin-triggered SCO patterns (Fig. 5C). MPEP had no
effect on the bifenthrin-induced decrease in the amplitude of
SCSO (Fig. 5D). Both AP-5 (50 \mu M) and MK-801 (1 \mu M),
competitive and noncompetitive antagonists of N-methyl-D-
aspartate receptor, respectively, partially reversed bifenthrin-
triggered SCO frequency (Fig. 5, E and G). AP-5 reduced basal
SCO by 69% ± 12% and bifenthrin-triggered SCO
frequency by 78% ± 5% (Fig. 5E). Similarly, MK-801 reduced
basal SCO by 49% ± 14% and bifenthrin-triggered SCO
frequency by 47% ± 14% (Fig. 5G). MK-801 and AP-5 alone
also dramatically inhibited the amplitude of basal SCOS by
69% ± 4% and 71% ± 4% of basal SCOS, respectively (Fig. 5, F
and H). Both MK-801 and AP5 further inhibited bifenthrin-
induced decrease in the amplitude of SCOS (Fig. 5, F and H).

**Subacute Exposure to Bifenthrin Persistently Alters
SCO Patterns.** We next investigated whether more pro-
longed exposure to bifenthrin also persistently altered the
SCO patterns. Cortical neurons were exposed to the vehicle or
various concentrations of bifenthrin (0.01–0.30 \mu M) for 6 days
commencing 24 hours after plating. After washout of
bifenthrin, cells were loaded with Fluo-4 for 1 hours. After
five washes, the cell plate was transferred to the FLIPR Tetra

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**Fig. 4.** Low bifenthrin concentrations do not affect cortical neuron Na^+ currents, whereas high concentrations delay inactivation. (A) Representative
Na^+ current recording showing the effect of vehicle control (dimethylsulfoxide; DMSO) followed by increasing concentrations of bifenthrin (0.1, 1, 10, and
100 \mu M). Na^+ currents were elicited by 50-millisecond voltage steps from −80 to 0 mV followed by a 50-millisecond step back to −80 mV. Under these
recording conditions, higher bifenthrin concentrations delayed inactivation and induced a slow tail current visible after repolarization to −80 mV. (B)
Concentration-response curves for increasing the charge during depolarization (= delay of Na_\text{v} channel inactivation) or after repolarization (= increase of
tail current). Data were normalized to the maximal response elicited with the highest bifenthrin concentration. (C) Inactivation time constants for
delay of the current and the tail current. Experiments were repeated on three independent cells with similar results, and data are presented as mean ± S.D.
Statistical significance between different groups was calculated using Student’s t test. msec, millisecond.
imager and SCOs were recorded. Chronic exposure to bifen-
thrin produced a persistent increase in SCO frequency (Fig. 6,
A and B) and a reduction in SCO amplitude (Fig. 6, A and C)
compared with vehicle control, despite the extensive washout
protocol (Fig. 6). Bifenthrin increased the frequency of SCOs
with an EC\textsubscript{50} value of 52.8 nM (25.1–110.1 nM, 95% CI) and

**Fig. 5.** mGluR5-N-methyl-D-aspartate receptor (NMDAR)–mediated neurotransmission, but not L-type Ca\textsuperscript{2+} channels contributes to bifenthrin (BF)-triggered SCO patterns. After exposure to vehicle or bifenthrin for 30 minutes, cortical neurons were exposed to either an L-type Ca\textsuperscript{2+} channel blocker, nifedipine (1 \textmu M) (A and B), the specific mGluR5 antagonist MPEP (1 \textmu M) (C and D), MK-801 (1 \textmu M), a noncompetitive antagonist of NMDAR (E and F), or the competitive MNDAR antagonist AP-5 (50 \textmu M) (G and H) to assess further modification or reversal of SCO activity. The data shown were averaged from a 10-minute period immediately after the addition of vehicle or candidate inhibitors. Data represent mean ± S.E.M. (n > 12) from at least two independent cultures performed in at least triplicate. Statistical significance between different groups was calculated using an analysis of variance followed by a Dunnett’s multiple comparison test (**P < 0.01, bifenthrin versus vehicle; ###P < 0.01, inhibitor versus vehicle; $$$P < 0.01, inhibitor + bifenthrin versus bifenthrin). Veh, vehicle.
Western blot analysis demonstrated that exposure to bifenthrin (0.1 \mu M) enhanced p-CREB in a time-dependent manner (Fig. 7, A and B). Given that MPEP partially ameliorated bifenthrin-enhanced SCO activity, we tested whether MPEP could also diminish bifenthrin stimulated CREB phosphorylation. After preincubation with MPEP (1 \mu M) or vehicle (0.01% dimethylsulfoxide) for 15 minutes, the neurons were then treated with bifenthrin (0.1 \mu M) or vehicle (0.01% dimethylsulfoxide) for 3 hours. Western blot analysis showed that MPEP significantly inhibited bifenthrin-stimulated CREB phosphorylation without altering basal CREB phosphorylation (Fig. 7, C and D).

**Discussion**

In this study, we demonstrate that bifenthrin, at concentrations well below those that influence neuronal membrane potential and delay Na$^+$ channel inactivation (<1 \mu M), is capable of altering SCOs that are essential for normal activity–dependent growth of cortical neurons. It is well known that bifenthrin and related pyrethroid insecticides influence axonal voltage-gated Na$^+$ channels by delaying their inactivation and producing a significant tail current, and as such this mechanism is currently accepted as the primary mechanism for both insecticidal activity and mammalian neurotoxicity (Clark and Symington, 2012; Soderlund, 2012; Casida and Durkin, 2013). Currently, the US Environmental Protection Agency proposes that all pyrethroids share a common mechanism for promoting impaired motor activity and therefore present a cumulative human health risk (US EPA, 2011). However, neurotoxicological symptoms in mammalian models vary greatly among individual pyrethroids, and differences in their toxicokinetics are insufficient to predict divergent behavioral responses (Wolansky et al., 2006; Starr et al., 2012). The current results indicate that, at least for bifenthrin, alterations in the patterns of SCO are independent of voltage-gated Na$^+$ channel modulation and occur at concentrations approximately two log units below those required for altering neuronal membrane potential. In support of an alternative mechanism, bifenthrin does not alter the spontaneous electric activity of cortical neurons measured in multiple electrode arrays (data not shown), a finding consistent with an independent report using cortical neuronal cultures (McConnell et al., 2012). Thus, the newly found mechanism for bifenthrin may have implications for understanding differences in the potency and pattern of pyrethroid neurotoxicity. For example, bifenthrin differs little
from permethrin in structure and physicochemical properties, yet bifenthrin has a 10- to 100-fold greater oral toxicity, which cannot be accounted for by differential activity at voltage-gated Na⁺ channels (Wolansky and Harrill, 2008).

Since bifenthrin-altered SCO patterns are sufficient to modify CREB phosphorylation and the early growth trajectory of developing cortical neurons in culture, these results raise questions as to whether certain pyrethroids, such as

Fig. 7. Bifenthrin (BF)-enhanced p-CREB is dependent on the mGluR5 activity. (A) Representative Western blot for bifenthrin (0.1 μM)-stimulated CREB phosphorylation (p-CREB) in 8-9 DIV cortical neurons as a function of time post treatment. (B) Quantification of bifenthrin-induced CREB phosphorylation. Each data point represents mean ± S.E.M. (n = 4) from two independent cultures, each time performed in duplicate. (C) Representative Western blot for MPEP (1 μM) on bifenthrin (0.1 μM)-stimulated p-CREB in 8 or 9 DIV cortical neurons. (D) Quantification of MPEP response of MPEP on bifenthrin-induced CREB phosphorylation. Each data point represents mean ± S.E.M. (n = 3) from two independent cultures. Statistical significance between different groups was calculated using an analysis of variance and followed by a Dunnett’s multiple comparison test (*P < 0.05; **P < 0.01, bifenthrin versus vehicle; ##P < 0.01, bifenthrin versus MPEP + bifenthrin). VEH, vehicle.

Fig. 8. Bifenthrin (BF)-altered neurite outgrowth is dependent on mGluR5 activity in cortical neurons. (A) Representative neurons stained with microtubule-associated protein 2 (MAP2) and Hoechst 33342 after exposure to vehicle [(0.01% dimethylsulfoxide (DMSO)] or increasing concentrations of bifenthrin (0.01–1.0 μM) for 48 hours (neurons were fixed and stained on 3 DIV). (B) Quantification of the total neurite length/cell after vehicle and bifenthrin exposure. Bifenthrin produces a biphasic response in the neurite outgrowth, which peaked at 0.1 μM. (C) Representative neurons stained with MAP2 and Hoechst 33342 after exposure to vehicle (0.02% DMSO), MPEP (1 μM), bifenthrin (0.1 μM) or MPEP + bifenthrin for 48 hours. MPEP was added to the culture medium 15 minutes before bifenthrin addition. (D) Quantification of the total neurite length per cell after MPEP, bifenthrin, or MPEP + bifenthrin exposure. The experiments were repeated on two separate culture days with similar results. Statistical significance between different groups was calculated using an analysis of variance and followed by a Dunnett’s multiple comparison test (*P < 0.05; **P < 0.01, bifenthrin versus vehicle; #P < 0.01, bifenthrin + MEPE versus bifenthrin).
bifenthrin, engage other primary molecular targets at concentrations much lower than those needed to modify voltage-gated Na channels. Mechanisms that alter the integrity of SCO may not only contribute to higher acute toxicity but may also affect more chronic neurodevelopmental outcomes related to long-term changes in Ca^{2+}-dependent processes, including activity-dependent dendritic growth regulated by CREB-dependent signaling pathways (Wayman et al., 2006). The observation that bifenthrin persistently alters SCO patterns after prolonged (i.e., 6 days) exposure indicates that this pyrethroid could alter Ca^{2+}-regulated pathways over the critical neurodevelopmental time frame.

Although we were unable to delineate the direct molecular targets for bifenthrin that elicited altered patterns of SCO, our data suggest that bifenthrin augments mGluR5 signaling since an mGluR5 receptor antagonist, MPEP, produced a maximal suppression in the bifenthrin-augmented SCOs frequency without affecting the basal spontaneous Ca^{2+} oscillations frequency. This finding is consistent with previous reports that have demonstrated Glu receptor signaling pathways regulate SCO behavior (Dravid and Murray, 2004; Koga et al., 2010). We also show that both competitive and noncompetitive N-methyl-D-aspartate receptor antagonists (AP5 and MK-801) suppress both basal and bifenthrin-augmented SCO activity. However, increased mGluR5 activity cannot fully explain how bifenthrin alters SCO activity since metabotropic glutamate receptor antagonists could not recover the SCO baseline amplitude.

Bifenthrin-augmented Ca^{2+} signaling is sufficient to rapidly increase the phosphorylation of CREB and neurite extension. Consistent with the inhibition of SCO activity of MPEP, bifenthrin response on CREB phosphorylation and neurite outgrowth is also dependent on amplification of the mGluR5 activity since both outcomes are mitigated by MPEP. Activity-dependent calcium signaling has been shown to regulate dendritic growth and branching (Konur and Ghosh, 2005), which is dependent on the phosphorylation of CREB (Wayman et al., 2012). Small to moderate amplification of Ca^{2+} signaling stimulates dendritic growth, whereas excessive Ca^{2+} influx may stall dendritic growth (Gomez and Spitzer, 2000; Hui et al., 2007). The concentration-dependent profile of bifenthrin-triggered neurite outgrowth is therefore consistent with previously reported mechanisms responsible for regulating activity-dependent dendritic growth (Gomez and Spitzer, 2000; Hui et al., 2007).

Numerous pyrethroids have also been demonstrated to interact with GABA\textsubscript{A} receptors and Ca^{2+} channels (Clark and Symington, 2012; Soderlund, 2012). However, the effect of bifenthrin on SCO was unlikely from an interaction with GABA\textsubscript{A} receptors. Bicuculline, a competitive GABA\textsubscript{A} antagonist, was reported to increase the spike rate using multiple electrode arrays recording in cortical neurons (McConnell et al., 2012). In addition, we found that both bicuculline and picrotoxin increased the synchronicity of neuronal firing in cortical neuronal cultures (data not shown). However, bifenthrin fails to alter the spike rate and firing synchronicity. Bifenthrin-triggered alterations in SCO are also unlikely to have been induced by direct interaction with Ca^{2+} channels since nifedipine has no effect on bifenthrin-augmented SCO, and we found no detectable influences of ryanodine receptor RyR1, as previously reported for deltamethrin (Morisseau et al., 2009). Collectively, the current data are consistent with a previous report in which no direct interaction of bifenthrin with GABA\textsubscript{A} receptor or Ca^{2+} channels was observed (Burr and Ray, 2004).

In the 1999 National Health and Nutrition Examination Survey, more than 70% of a nationally representative sample of persons aged 6 years or older had detectable levels of pyrethroid metabolites in their urine (Burr et al., 2010). Although pyrethroids are generally thought to be rapidly metabolized, and therefore are unlikely to bioaccumulate in breast milk, several recent studies have found that pyrethroid levels vary greatly in human breast milk, ranging from 1<sub>–</sub>2000 ng g<sup>–1</sup> depending on geographic location and insecticide use patterns (Sereda et al., 2009; Weldon et al., 2011; Corellas et al., 2012). For example, a recent analysis of pyrethroids in human breast milk indicated that bifenthrin is most abundant in Brazilian samples, whereas \textalpha\textsubscript{-}cyhalothrin and permethrin are most abundant in Colombian and Spanish samples, respectively (Corellas et al., 2012). In this study, the mean bifenthrin levels ranged from 1.44 to 2.8 ng g<sup>–1</sup> (lipid weight) and maximum levels approached 7.48 ng g<sup>–1</sup>. The bifenthrin concentration even in these highly exposed individuals would be below the minimum effective concentration (10 nM) that alters neurite outgrowth (Fig. 8). However, considering the very low solubility of bifenthrin in aqueous media (logP 6.4) (Schleier and Peterson, 2011), the actual concentration of bifenthrin in the culture medium at the time measurements of neurite length and SCO properties after “chronic” exposures would be expected to be lower than nominal concentrations at the start of exposures.

The present results also raise the important question of whether exposure to bifenthrin could produce additive effects with other pyrethroids (Starr et al., 2012). It is also possible that the effects of bifenthrin may be additive with exposure to persistent chemicals such as nonoxyin-like polychlorinated biphenyls (Pessah et al., 2010) or genetic mutations known to produce abnormal SCO activity and dendritic and synaptic architecture and network connectivity (Stamou et al., 2013). For example, fragile X mental retardation, a neurodevelopmental disorder caused by loss of fragile X mental retardation protein, is the most widespread single-gene cause of developmental disorders, including autism (FMR; Hagerman and Hagerman, 2013). In FMR1 gene knockout mouse model, enhanced mGluR5 signaling has been demonstrated and contributes to long-term synaptic depression during synaptic transmission. Such disturbances in mGluR5 signaling are thought to elicit cognitive as well as syndromic features of fragile X syndrome (Huber et al., 2002; Bear et al., 2004; Dolen and Bear, 2008), and FMR1 premutation (Chen et al., 2010; Cao et al., 2012b; Liu et al., 2012). Our in vitro observation that nanomolar bifenthrin alters neuronal Ca^{2+} dynamics and mGluR5 signaling activity with consequential changes in neurite outgrowth provides evidence for a new paradigm in understanding how certain pyrethroids might influence health outcomes that involve abnormal neuronal network development associated with developmental disorders.

**Authorship Contributions**

**Performed data analysis:** Cao, Cui, Nguyen, Jenkins, Wulff, Pessah.

**Wrote or contributed to the writing of the manuscript:** Cao, Cui, Nguyen, Wulff, Pessah.
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Nanomolar Bifenthrin Alters Synchronous Ca^{2+} Oscillations And Cortical Neuron Development

Independent of Sodium Channel Activity

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Supplementary Figure 1 | (A), Representative traces for spontaneous and bifenthrin-augmented Ca^{2+} oscillations before and after washout. (B), Quantification of bifenthrin-augmented Ca^{2+} oscillations frequency before and after complete wash for 5 times. Complete washout out of bifenthrin produces a partial recovery on the frequency of Ca^{2+} oscillations suggesting a persistent response on the spontaneous Ca^{2+} oscillations of bifenthrin. Each data point represents Mean±SEM (n=10) from two experiments each performed in quintuplicates.