Amphetamine-like Neurochemical and Cardiovascular Effects of α-Ethylphenethylamine Analogs Found in Dietary Supplements

Charles W. Schindler, Eric B. Thorndike, John S. Partilla, Kenner C. Rice, and Michael H. Baumann

Designer Drug Research Unit (C.W.S., J.S.P., M.H.B.) and Preclinical Pharmacology Section (C.W.S., E.B.T.), National Institute on Drug Abuse Intramural Research Program, Baltimore, Maryland and Drug Design and Synthesis Section, National Institute on Drug Abuse and National Institute of Alcohol Abuse and Alcoholism Intramural Research Programs, Rockville, Maryland (K.C.R.)

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ABSTRACT

Dietary supplements often contain additives not listed on the label, including α-ethyl homologs of amphetamine such as Nα-diethylphenethylamine (DEPEA). Here, we examined the neurochemical and cardiovascular effects of α-ethylphenethylamine (AEPEA), N-methyl-α-ethylphenethylamine (MEPEA), and DEPEA as compared with the effects of amphetamine. All drugs were tested in vitro using uptake inhibition and release assays for monoamine transporters. As expected, amphetamine acted as a potent and efficacious releasing agent at dopamine transporters (DAT) and norepinephrine transporters (NET) in vitro. AEPEA and MEPEA were also releasers at catecholamine transporters, with greater potency at NET than DAT. DEPEA displayed fully efficacious release at NET but weak partial release at DAT (i.e., 40% of maximal effect). In freely moving, conscious male rats fitted with biotelemetry transmitters for physiologic monitoring, amphetamine (0.1–3.0 mg/kg, s.c.) produced robust dose-related increases in blood pressure (BP), heart rate (HR), and motor activity. AEPEA (1–10 mg/kg, s.c.) produced significant increases in BP but not HR or activity, whereas DEPEA and MEPEA (1–10 mg/kg, s.c.) increased BP, HR, and activity. In general, the phenethylamine analogs were approximately 10-fold less potent than amphetamine. Our results show that α-ethylphenethylamine analogs are biologically active. Although less potent than amphetamine, they produce cardiovascular effects that could pose risks to humans.

SIGNIFICANCE STATEMENT

The α-ethyl homologs of amphetamine have significant cardiovascular, behavioral, and neurochemical effects in rats. Given that these compounds are often not listed on the ingredient labels of dietary supplements, these compounds could pose a risk to humans using these products.

Introduction

Nutritional supplements often contain ingredients that are not listed on the product labels, including analogs of phenethylamine (PEA) that display structural similarity to amphetamine (Eichner, 2014; Pawar and Grundel, 2017). We recently showed that the supplement additive β-methylphenethylamine (BMPEA), a positional isomer of amphetamine (i.e., α-methylphenethylamine), increases blood pressure (BP) in rats and may therefore produce adverse effects in humans (Schindler et al., 2019). Another PEA analog previously found in dietary supplements is Nα-diethylphenethylamine (DEPEA; see Fig. 1 for chemical structure). DEPEA has been detected in powdered material confiscated for drug trafficking (Lee et al., 2013) and in supplement products destined for human consumption (ElSohly and Gul, 2014; Cohen et al., 2014; Wahlstrom et al., 2014; ElSohly et al., 2015).

Urine toxicology testing has also confirmed the presence of DEPEA in users of some dietary supplements (Wójtowicz et al., 2015). For example, in urine samples obtained through routine toxicological testing, DEPEA was detected in samples from individuals who used a dietary supplement suspected of containing added DEPEA (Uralets et al., 2014). In addition to DEPEA, α-ethylphenethylamine (AEPEA) was also found in some of the same samples, potentially the result of N-dealkylation of DEPEA via hepatic metabolism. Both DEPEA and AEPEA have been found in urine samples collected by the World Anti-Doping Agency, demonstrating exposure to these substances among athletes (World Anti-Doping Agency, 2012, 2013). In a survey of German athletes who were asked about their supplement use, some respondents specifically reported seeking products containing DEPEA (Dreher et al., 2018). Since DEPEA and related analogs are not listed as ingredients on the labels of supplement products, the FDA considers them adulterants and requires companies...
saying the products to remove them from the market (Pawar and Grundel, 2017). However, PEA analogs may still be present in those supplements not tested by the FDA.

Despite the apparent widespread use of DEPEA and similar compounds, very little is known about their biologic effects. Oberlender and Nichols (1991) reported that AEPEA partially generalizes to the discriminative stimulus effects of amphetamine. In that study, rats were trained to respond on one lever when injected with amphetamine and another lever when injected with saline. When the trained rats received a non-contingent injection of AEPEA, they responded on the amphetamine-associated lever, suggesting that AEPEA and amphetamine might share common effects. Santillo (2014) reported that AEPEA inhibits human monoamine-oxidase type A in vitro in a competitive and reversible manner similar to amphetamine, whereas DEPEA has much weaker effects in this regard. Liu and Santillo (2016) reported that DEPEA inhibits activity of the hepatic cytochrome P450 enzyme CYP2D6, which could alter the effects of other drugs taken in combination with DEPEA. Because of the paucity of information about the pharmacology of α-ethyl PEA analogs, we sought to study the effects of AEPEA and its amine-substituted analogs, N-methyl-α-ethylphenethylamine (MEPEA) and DEPEA, as compared with the effects of amphetamine (see Fig. 1 for chemical structures). Based on their structural similarities to amphetamine, we expected that these PEA analogs would have similar effects to amphetamine, although at potentially different potencies.

**Materials and Methods**

**Drugs and Reagents.** AEPEA, MEPEA, and DEPEA were synthesized using standard organic chemical reactions and techniques as follows: AEPEA was prepared by reductive amination of 1-phenyl-2-butanone (TCI America Research Chemicals, Portland, OR) using the method of González-Sab et al. (2002). The distilled base was then converted to the HCl salt in acetonitrile-ether. MEPEA was prepared in two steps by first N-formylation of AEPEA with ethyl formate at 135°C for 18 hours in a pressure bottle. The resulting N-formyl derivative was distilled and then reduced to MEPEA with Vitride in refluxing Toluene. The distilled base was then converted to the oxalate salt in acetonitrile. DEPEA was synthesized in two steps by N-acetylation of AEPEA with acetic anhydride in a chloroform-saturated sodium bicarbonate two-phase system, followed by reduction of the resulting N-acetyl derivative with lithium aluminum hydride in tetrahydrofuran. The distilled base was converted to the HCl salt in acetone-ether. Each compound was fully characterized with appropriate high-resolution mass spectral and 400-MHz NMR analyses and with appropriate combustion analyses for carbon, hydrogen, and nitrogen. Each compound was chromatographically homogenous by thin-layer chromatography. Chemical purity for each compound was estimated to be greater than 98%. The corresponding α-methyl comparator compound (S)-amphetamine sulfate (amphetamine) was obtained from the pharmacy at the National Institute on Drug Abuse (NIDA), Intramural Research Program (IRP), in Baltimore, MD. [3H]Methyl-4-phenylpyridinium ([3H]MPP⁺; 80 Ci/mmol) was purchased from American Radiolabeled Chemicals (St. Louis, MO), and [3H]-neurotransmitters (30–50 Ci/mmol) were purchased from Perkin Elmer (Shelton, CT). All other chemicals and reagents were acquired from Sigma-Aldrich (St Louis, MO) unless otherwise noted. Drugs for the in vivo experiments were dissolved in sterile saline, and doses are expressed as the salts.

**Animals.** Male Sprague-Dawley rats were used for all experiments, as described in detail below. All procedures were approved by the Animal Care and Use Committee of the NIDA IRP and were carried out in accordance with the Guide for the Care and Use of Laboratory Animals. Animals were housed in facilities fully accredited by the Association for the Assessment and Accreditation of Laboratory Care.

**In Vitro Transporter Assays.** In total, 28 male Sprague-Dawley rats (Charles River, Kingston, NY) weighing 250–300 g were used for the synaptosome assays. Rats were group-housed with free access to food and water under a 12-hour light/dark cycle with lights on at 0700 hours. Rats were euthanized by CO₂ narcosis, and synaptosomes were prepared from brains using standard procedures (Rothman et al., 2002). Transporter uptake and release assays were performed as described previously (Solis et al., 2017). In brief, synaptosomes were prepared from caudate tissue for dopamine transporter (DAT) assays and from whole brain minus caudate and cerebellum for norepinephrine transporter (NET) and serotonin (5-HT) transporter (SERT) assays.

For uptake inhibition assays, 5 nM [3H]dopamine, [3H]norepinephrine, or [3H]5-HT was used for DAT, NET, or SERT assays, respectively. To optimize uptake for a single transporter, unlabeled blockers were included to prevent the uptake of [3H]-transmitter by competing transporters. Uptake inhibition was initiated by incubating synaptosomes with various doses of test compound and [3H]-transmitter in Krebs-phosphate buffer. Uptake assays were terminated by rapid vacuum filtration, and retained radioactivity was quantified with liquid scintillation counting (Baumann et al., 2013).

For release assays, 9 nM [3H]MPP⁺ was used as the radiolabeled substrate for DAT and NET, whereas 5 nM [3H]5-HT was used for SERT. All buffers used in the release assay contained 1 μM reserpine to block vesicular uptake of substrates. The selectivity of release...
assays was optimized for a single transporter by including unlabeled blockers to prevent the uptake of $[^{3}H]$MPP$^+$ or $[^{3}H]$5-HT by competing transporters. Synaptosomes were preloaded with radiolabeled substrate in Krebs-phosphate buffer for 1 hour to reach steady state. Release assays were initiated by incubating preloaded synaptosomes with various concentrations of the test drug. Release was terminated by vacuum filtration, and retained radioactivity was quantified by liquid scintillation counting. For substrate reversal experiments, the effects of AEPEA and MEPEA on transporter-mediated release were examined as described above in the presence or absence of 1 nM 1-[2-[bis(4-fluorophenyl)methoxyethyl]-4-(3-phenylpropyl)piperazine (GBR12909) for DAT assays or 8 nM desipramine for NET assays. Effects of test drugs on release were expressed as a percentage of maximal release, with maximal release (i.e., 100% $E_{\text{max}}$) defined as the release produced by tyramine at doses that evoke the efflux of all “releasable” tritium by synaptosomes (10 μM tyramine for DAT and NET assay conditions, and 100 μM tyramine for SERT assay conditions).

Effects of test drugs on uptake inhibition and release were analyzed by nonlinear regression using GraphPad Prism 8 (GraphPad Scientific, San Diego, CA). Dose-response values for the uptake inhibition and release were fit to the equation $Y(x) = Y_{\text{max}} + (Y_{\text{max}} - Y_{\text{min}})(1 + 10^{\log IC_{50} - \log x}) \times n$, where $x$ is the concentration of the compound tested, $Y(x)$ is the response measured, $Y_{\text{max}}$ is the maximal response, $Y_{\text{min}}$ is either IC$_{50}$ (the concentration that yields half-maximal uptake inhibition response) or EC$_{50}$ (the concentration that yields half-maximal release), and $n$ is the Hill slope parameter. We employed the “EC$_{50}$ shift test” in Prism to statistically evaluate the possibility of significant differences in potency across the drugs. Briefly, the EC$_{50}$ shift compares dose-response curves for a series of drugs and can be used to test the null hypothesis that all drugs are equipotent (i.e., the EC$_{50}$ ratio equals 1). When significant differences in drug potency are observed, the null hypothesis is rejected, and supporting F scores and $P$ values are given.

Receptorome Screening. AEPEA, MEPEA, DEPEA, and amphetamine were submitted to the psychoactive drug screening program program of the National Institute on Mental Health and evaluated for binding affinity at a variety of human G protein–coupled receptors (GPCRs) according to established protocols (Besnard et al., 2012; https://pdbp.unc.edu/pdbpweb/content/UNC-CH%20Protocol%20Book.pdf). Compounds were first screened at a fixed concentration of 10 μM to assess inhibition of receptor binding. In those instances in which binding was inhibited by more than 50% at 10 μM, full dose-effect functions were obtained, and $K_i$ values were calculated by nonlinear regression using the Cheng-Prusoff equation.

In Vivo Biotelemetry. Five adult male Sprague-Dawley rats (Charles River, Kingston, NY) were used as subjects for the telemetry experiments. Rats were purchased by Data Sciences International (St. Paul, MN) and received surgically implanted HD-S10 biotelemetry transmitters. For the surgery, the rats were anesthetized with isoflurane and the abdominal cavity was opened. The descending aorta was isolated, and the catheter from the transmitter was inserted into the aorta and glued in place. The abdominal muscles and skin were then sutured to close the incision. Rats were treated with subcutaneous meloxicam after surgery. After recovery at Data Sciences International, the rats were shipped to the NIDA IRP in Paul, MN) and received surgically implanted HD-S10 biotelemetry transmitters. For the surgery, the rats were anesthetized with isoflurane and the abdominal cavity was opened. 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As expected, amphetamine was a fully efficacious substrate-type releasing agent with high potency at DAT (EC50 = 5 nM) and NET (EC50 = 8 nM). AEPEA and MEPEA were much less potent than amphetamine as releasers at DAT but were approximately equipotent with each other at this transporter. It is noteworthy that DEPEA was a weak partial releaser at DAT, achieving only 40% of the maximal release response (i.e., partial agonist effect). EC50 shift analysis showed that amphetamine was significantly more potent as a releaser at DAT when compared with PEA compounds (F3,89 = 374, P < 0.0001), whereas DEPEA was significantly less potent than AEPEA and MEPEA in this regard (F2,57 = 158, P < 0.0001). All of the PEA compounds were efficacious substrate-type releasers at NET. EC50 shift analysis demonstrated that amphetamine was a more potent releaser at NET when compared with PEA compounds (F3,89 = 96.2, P < 0.0001), and DEPEA was less potent than AEPEA and MEPEA (F2,66 = 29.6, P < 0.0001). In general, AEPEA and MEPEA displayed greater potency as releasers at NET (EC50 = 80 and 58 nM) when compared with DAT (EC50 = 273 and 179 nM).

To explore the mechanism of 3H-neurotransmitter efflux produced by the efficacious releasers AEPEA and MEPEA, we carried out “substrate reversal” studies. These experiments involved testing the ability of selective uptake blockers to antagonize the releasing actions of the drugs. Figure 3 illustrates the effects of the selective DAT uptake inhibitor GBR12909 (1 nM) or the selective NET uptake inhibitor desipramine (8 nM) on releasing activity produced by AEPEA and MEPEA. The results show that GBR12909 produced a parallel rightward shift in the DAT release curves for AEPEA and MEPEA, suggesting that both drugs interact at the orthosteric site on DAT to elicit their releasing actions. An EC50 shift analysis confirmed that GBR12909 significantly reduced the potency of AEPEA (F1,43 = 1875, P < 0.0001) and MEPEA (F1,43 = 504, P < 0.0001) at DAT. For the NET substrate reversal experiments, desipramine shifted the NET release curves for AEPEA and MEPEA to the right, demonstrating that these drugs interact at NET sites. The EC50 shift results confirmed that desipramine significantly reduced the potency of AEPEA (F1,43 = 64, P < 0.0001) and MEPEA (F1,43 = 109, P < 0.0001) at NET.

Receptorome Screening. Table 2 presents the results for AEPEA, MEPEA, and DEPEA in the human GPCR screening in comparison with amphetamine. In general, the PEA analogs had little activity at GPCRs when tested at a 10 μM concentration (see receptors listed in footnote of Table 2).
Specific exceptions included the 5-HT$_{1A}$ receptor, in which MEPEA and DEPEA had low micromolar affinities (1966–1588 nM) and the $\alpha_2$ receptor subtypes, in which all three PEA analogs showed affinities in the range of their transporter releasing potency (411–2320 nM). DEPEA showed activity at DAT and NET (108–124 nM), whereas AEPEA and MEPEA did not. AEPEA and DEPEA had low micromolar affinities to inhibit binding to the $\sigma_2$ site, whereas AEPEA and MEPEA had low to mid micromolar affinities to inhibit binding at the histamine H1 site. Amphetamine showed activity at the $\alpha$ receptor subtypes and also at the NET but, interestingly, not at the DAT. Although this latter finding seems counterintuitive, previous studies demonstrate that amphetamine displays weak ability to displace high-affinity phenyltropane analogs at monoamine transporters (Rothman et al., 1999; Eshleman et al., 1999, 2017).

**Biotelemetry in Rats.** Rats quickly adapted to the experimental injection procedure, and results for the control conditions remained relatively stable throughout testing. Figure 4 depicts representative time course effects of saline or AEPEA injection on BP, HR, activity, and body temperature. In saline-treated rats, BP rapidly decreased and stabilized over the 3-hour session, whereas AEPEA produced time-related ($F_{17,338} = 14.9, P < 0.0001$) and dose-related ($F_{4,20} = 18.0, P < 0.0001$) elevations in BP that were maintained throughout the session. The hypertensive effect of AEPEA was significantly greater than saline at all doses (Dunnett’s $P < 0.05$). The HR response to AEPEA displayed significant effects of time ($F_{17,338} = 7.3, P < 0.0001$) but not dose ($F_{4,20} = 2.3, P = 0.098$), even though the higher doses of AEPEA caused modest tachycardia toward the end of the session. AEPEA produced small increases in motor activity at the beginning of the session, which were significant with respect to time ($F_{17,338} = 17.1, P < 0.0001$) but not dose ($F_{4,20} = 1.7, P = 0.1905$). Core body temperature was not significantly altered by AEPEA administration ($F_{4,20} = 1.7, P = 0.1922$), despite a hypothermic response at the highest dose administered.

<table>
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<td><strong>G Protein–coupled receptorome screening for amphetamine and PEA analogs</strong></td>
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<td>Data represent $K_i$ (nanomolar) values obtained from nonlinear regression using the Cheng-Prusoff equation when inhibition was $&gt;50%$. The results for amphetamine (AMPH) were reported previously (Schindler et al., 2019).</td>
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<tr>
<th></th>
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$^*$K$_i$ values for the following receptors were $>10,000$ nM for all four drugs tested: serotonin receptors 5-HT$_{1B}$, 5-HT$_{1D}$, 5-HT$_{2A}$, 5-HT$_{2C}$, 5-HT$_{3}$, 5-HT$_{5A}$, 5-HT$_{6}$, 5-HT$_{7}$, and SERT; norpinephrine receptors $\alpha_1A$, $\alpha_1B$, $\alpha_1D$, $\beta_1$, $\beta_2$, and $\beta_3$; dopamine D$_{1}$, D$_{2}$, D$_{3}$, D$_{4}$, and D$_{5}$; histamine receptors H$_{2}$, H$_{3}$, and H$_{4}$; opioid receptors $\mu$, $\kappa$, and $\delta$; muscarinic receptors M$_{1}$, M$_{2}$, M$_{3}$, M$_{4}$, and M$_{5}$; $\sigma_1$ receptor; benzodiazepine and peripheral benzodiazepine receptors.
The time course data from Fig. 4 demonstrated that BP effects of AEPEA lasted for most of the 3-hour session, and the effects of MEPEA and DEPEA followed similar time courses (data not shown). Given this information, we examined dose-response relationships for the averaged responses over the 3-hour sessions. Figure 5 shows the effects of amphetamine and the PEA analogs on BP, HR, locomotor activity, and body temperature averaged over the full 3-hour session. As expected, amphetamine produced dose-dependent increases in BP ($F_{4,24} = 27.7, P < 0.0001$) and HR ($F_{4,24} = 9.4, P = 0.0002$), with the highest three doses (0.3–3.0 mg/kg) producing increases in BP and HR significantly above saline control.

Amphetamine also produced increases in locomotor activity ($F_{4,24} = 5.4, P = 0.0041$), although those effects were not dose-dependent. Activity increased as dose increased up to 1 mg/kg, which was significantly above saline levels. However, at the dose of 3 mg/kg, activity decreased when compared with the effect of 1 mg/kg. This apparent suppression of activity most likely reflects an increase in stereotypy that is not measured by the telemetry devices. At the doses tested, amphetamine did not significantly affect body temperature.

Similar to amphetamine, all three PEA analogs increased BP in a dose-dependent manner (AEPEA $F_{4,24} = 28.4, P < 0.0001$; MEPEA $F_{3,19} = 17.2, P < 0.0001$; DEPEA $F_{3,19} = 10.6$).
P = 0.0004), with higher doses of each drug producing increases in BP significantly above saline control. The maximal effects of each drug on BP were similar. When only the doses inducing maximal effects were compared by one-way ANOVA, the overall effect was significant (F3,16 = 3.5, P = 0.04); however, comparisons between the drugs (Tukey) failed to show any significant differences between any of the PEA analogs and amphetamine. MEPEA and DEPEA also produced dose-dependent increases in HR (MEPEA F3,19 = 6.6, P < 0.01; DEPEA F3,19 = 8.1, P = 0.0017), with the effects of each drug at the highest dose tested being significantly above saline control. The effects of AEPEA (F4,24 = 8.6, P = .0003) were more complicated, with HR increasing after treatment doses up to 3 mg/kg but then returning toward saline levels at 10 mg/kg. For both their effects on BP and HR, the PEA analogs appeared to be approximately 10 times less potent than amphetamine. Like with BP, the maximal effects of each drug were similar (F3,16 = 2.1, P = 0.14).

Because all of the test drugs produced dose-dependent increases in BP, ED50 values were calculated based on the maximal effect of amphetamine compared with saline. The calculated ED50 for amphetamine was 0.31 mg/kg (95% CI = 0.17–0.48 mg/kg). The ED50 values for the PEA analogs were from 8- to 14-fold greater than that of amphetamine, and the 95% CIs for the PEA analogs did not overlap with those of amphetamine (AEPEA ED50 = 2.54 mg/kg, 95% CI = 1.89–3.53 mg/kg; DEPEA ED50 = 3.74 mg/kg, CI = 2.33–6.71 mg/kg; MEPEA ED50 = 4.47 mg/kg, CI = 2.80–9.02 mg/kg). Moreover, an EC50 shift analysis confirmed that the potency of amphetamine to induce increases in BP was significantly greater than the potencies of the PEA compounds (F3,56 = 13.3, P < 0.0001), although the effects of AEPEA, MEPEA, and DEPEA on BP did not differ from each other. Taken together, these results support the conclusion that the PEA analogs are generally 10 times less potent than amphetamine at increasing BP.

Similar to amphetamine, MEPEA and DEPEA produced significant increases in locomotor activity (MEPEA F3,19 = 7.4, P = 0.0025; DEPEA F3,19 = 10.9, P = 0.0004), with the increases at 10 mg/kg being significantly above saline controls. In contrast to amphetamine, there was no downturn in activity at the highest doses of MEPEA or DEPEA, although doses above 10 mg/kg were not tested. For their effects on locomotor activity, MEPEA and DEPEA were at least 10 times less potent than amphetamine. There was a trend for AEPEA to also produce small dose-related increases in activity (F4,24 = 3.1, P = 0.0376); however, follow-up tests failed to reveal any significant changes from saline control at any dose tested. Unlike with BP and HR, the maximal effects for the drugs on locomotor activity were different for the dose range tested. When the maximal effects were compared (F3,16 = 4.1, P = 0.02), the effects of AEPEA were significantly different from amphetamine. None of the PEA analogs produced significant effects on body temperature.

Discussion

PEA analogs continue to be found in dietary supplements, despite being banned by the FDA (Cohen et al., 2014; Elsohly et al., 2015; Zhao et al., 2018). Here, we studied compounds that possess an ethyl group at the α-position carbon, in contrast to amphetamine, which has a methyl group at this position. Our results show that compounds found in nutritional supplements can have effects that mimic those produced by the abused psychostimulant amphetamine. Like amphetamine, all three of the structurally related PEA analogs produced statistically significant elevations in BP, and MEPEA and DEPEA also increased HR. MEPEA and DEPEA produced increases in locomotor activity similar to amphetamine. In general, the PEA analogs were about 10 times less potent than amphetamine. If supplement products are taken orally according to the manufacturer’s recommendations, the amount of DEPEA ingested in these nutritional supplements would be 35–45 mg per serving (Cohen et al., 2014; Elsohly et al., 2015). In humans, the threshold oral dose of amphetamine that can be discriminated is 5 mg (Chait et al., 1985), whereas amphetamine increases BP and HR at oral doses as low as 7.5–10 mg (Brauer and de Wit, 1996; Kelly et al., 2006; Harvanko et al., 2016). If the relative potency estimates for PEA analogs observed here for subcutaneous administration in rats hold for oral consumption in humans, a single serving of the supplement containing one of the compounds may be sufficient to produce similar subjective effects to amphetamine, and as little as two servings may produce cardiovascular effects. Thus, it is possible for an individual to use these supplements in amounts that could produce amphetamine-like effects, including mood stimulation and cardiovascular complications.

Although the PEA analogs produced similar effects on BP with comparable potencies, there were substantial differences in their other in vivo effects. DEPEA appeared to produce larger and more potent effects on HR. MEPEA may produce larger HR effects at higher doses, but the effect of AEPEA appeared to peak at 3 mg/kg. Unlike DEPEA and MEPEA, AEPEA had minimal effects on locomotor activity at the dose range tested. These differences in effects on activity may translate into differences for other behavioral effects, such as drug self-administration, although additional research would be needed to confirm this. None of the drugs, including amphetamine, had significant effects on body temperature at the doses tested.

As expected, in the release assays, amphetamine was a potent efficacious releaser at both DAT and NET in rat brain synaptosomes (Rothman and Baumann, 2003). AEPEA and MEPEA were also releasers, but they were less potent than amphetamine and displayed selectivity for the NET over the DAT. The releasing abilities of AEPEA and MEPEA were antagonized by inhibitors of DAT and NET, implicating monoamine transporters in their effects. However, any definitive conclusions about the molecular mechanism of action for PEA compounds will require the replication of our findings using transfected cells expressing pure populations of DAT and NET. The releasing abilities of AEPEA and MEPEA were antagonized by inhibitors of DAT and NET, implicating monoamine transporters in their effects. However, any definitive conclusions about the molecular mechanism of action for PEA compounds will require the replication of our findings using transfected cells expressing pure populations of DAT and NET. Despite the similarity of effects produced by AEPEA and MEPEA in the in vitro assays, MEPEA significantly increased locomotor activity in vivo, whereas AEPEA did not. DEPEA displayed fully efficacious release at NET but only weak partial releasing activity at DAT (i.e., 40% of Emax). The precise molecular underpinnings of DEPEA’s partial releasing activity at DAT is not known, but we speculate that uptake inhibition at DAT is the predominant effect of DEPEA at this transporter. Like the other PEA analogs, DEPEA was slightly more potent at NET than DAT. In general, DEPEA was less potent than AEPEA or MEPEA at both DAT and NET in vitro but was at least equipotent to both MEPEA and AEPEA on BP and HR in vivo and clearly more potent than AEPEA on...
locomotor activity. This pattern of results suggests that in vivo effects of these PEA analogs are influenced by factors other than uptake inhibition or release properties at DAT and NET, such as pharmacokinetic factors or actions at other binding sites.

The α-ethyl–substituted PEA analogs failed to interact (i.e., \( K_i > 10 \, \mu M \)) with most of the GPCRs studied (footnote of Table 2). Further, for the few receptors they bind to, the compounds do not show potency relationships that mirrored any of the in vivo effects observed. For example, at the 5-HT1A receptor, amphetamine was equipotent to MEPEA and DEPEA, but amphetamine was much more potent than both of these compounds in all measures of in vivo activity. At the α2B receptor, MEPEA appears to have twice the potency of DEPEA; however, MEPEA displayed equivalent or less potency than DEPEA for the in vivo tests. Therefore, it does not appear as though activity at these other receptors is responsible for the observed in vivo effects. Given the fact that amphetamine showed clearly greater potency than the PEA analogs on the DAT and NET release assays and that the same potency relationship held for the in vivo tests, actions at DAT and NET seem most likely to mediate the in vivo effects, although we cannot rule out that pharmacokinetic effects may be responsible for some of the observed differences.

In a previous study (Schindler et al., 2019), we examined the pharmacological effects of a PEA analog in which the α-methyl group of amphetamine was moved to the β-position to form BMPEA. This change reduced the potency of BMPEA to increase BP about 30-fold when compared with amphetamine, and at the doses tested, BMPEA did not increase HR or activity. BMPEA maintained substrate activity at DAT and NET but was more potent at NET than DAT. By comparison, in the experiments reported here, replacing the α-methyl group of amphetamine and methamphetamine with an α-ethyl group to yield AEPEA and MEPEA led to a reduction in potency of only about 10-fold for the in vivo effects and also maintained releaser efficacy. Importantly, the α-ethyl compounds seem to show a preference for the NET over the DAT, which is similar to the profile of BMPEA. Replacing the amine methyl group of MEPEA with an ethyl group to produce DEPEA markedly reduced the efficacy of release at DAT, and this observation agrees with findings reported for amphetamine analogs with extended N-alkyl chain length (see Solis et al., 2017). Overall, replacing the α-methyl group of amphetamine with an ethyl group maintains the amphetamine-like effects of PEA analogs more effectively than moving the methyl group to the β-position for BMPEA. Thus, α-ethyl–substituted analogs may also have abuse potential similar to amphetamine.

Although not directly studied here, it is likely that locomotor-activating effects of α-ethyl compounds are related to their effects at the DAT. Activity at DAT is known to be primarily responsible for the locomotor-activating effects of a variety of stimulants, including cocaine, amphetamine, and related analogs (Uhl et al., 2002; Rothman and Baumann, 2003; Zolkowska et al., 2009). Activity at DAT is also known to be primarily responsible for maintaining drug self-administration (Wise and Bozarth, 1985; Uhl et al., 2002; Howell and Kimmel, 2008), and our present findings suggest that at least some of the α-ethyl–substituted analogs will be self-administered and have abuse potential in humans. Likewise, previous work with other psychomotor stimulants has shown that central or peripheral noradrenergic systems are primarily responsible for their cardiovascular effects. For example, increases in BP produced by BMPEA were blocked by the α-adrenergic antagonist prazosin (Schindler et al., 2019). β-Adrenergic antagonists can block the HR-increasing effects of psychomotor stimulants, including some amphetamines (Schindler et al., 1992b, 2014), cathinones (Varner et al., 2013; Alsufyani and Docherty, 2015; Schindler et al., 2016), and cocaine (Branch and Knuepfer, 1992; Schindler et al., 1992a). With respect to cocaine, although pretreatment with β antagonists can antagonize tachycardia, some studies show that such antagonists can exacerbate the hypertensive effects (Branch and Knuepfer, 1992; Schindler et al., 1992a).

Limitations of the current study include the measurement of only BP and HR as indices of cardiac function. Although increases in both BP and HR would be expected to increase the workload on the heart and lead to potential adverse effects, the assessment of other relevant endpoints, such as cardiac output, cardiac electrical activity, and contractility (Mladenka et al., 2018), would produce a more complete picture of the effects of α-ethyl–substituted PEA analogs on cardiac function. Nevertheless, the effects shown here do raise concern for the presence of these substances in dietary supplements, particularly if they are not listed on the ingredients label. In addition, the in vivo effects reported here were observed after passive administration of the compounds. In humans, these substances would be self-administered, which could influence the effects of the compounds on cardiac function, as well as other measures (Hemby et al., 1997; Graziella De Montis et al., 1998).

In summary, similar to amphetamine, AEPEA and MEPEA function as efficacious neurotransmitter releasers at DAT and NET, but with greater potency at NET relative to DAT. By contrast, DEPEA displays efficacious release at NET but low-efficacy partial release at DAT. Despite these minor differences from amphetamine in neurochemical mechanism, all three α-ethyl PEA analogs increased BP similar to amphetamine, although at reduced potency. MEPEA and DEPEA also increased HR. As a result, these compounds could produce toxic effects if taken in large enough amounts. Like amphetamine, MEPEA and DEPEA increased locomotor activity, suggesting these agents could have central effects, including abuse potential in humans.

Authorship Contributions
Participated in research design: Schindler, Baumann.
Conducted experiments: Schindler, Thorndike, Partilla, Baumann.
Contributed new reagents or analytic tools: Rice.
Performed data analysis: Schindler, Thorndike, Baumann.
Wrote or contributed to writing of the manuscript: Schindler, Partilla, Rice, Baumann.

References

Address correspondence to: Charles W. Schindler, Designer Drug Research Unit, IRP, NIDA, NIH, DHHS, 251 Bayview Blvd., Baltimore, MD 21224.
E-mail: cshind@helix.nih.gov