Desvenlafaxine Succinate: A New Serotonin and Norepinephrine Reuptake Inhibitor

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ABSTRACT

The purpose of this study was to characterize a new chemical entity, desvenlafaxine succinate (DVS). DVS is a novel salt form of the isolated major active metabolite of venlafaxine. Competitive radioligand binding assays were performed using cells expressing either the human serotonin (5-HT) transporter (hSERT) or norepinephrine (NE) transporter (hNET) with Kᵢ values for DVS of 40.2 ± 1.6 and 558.4 ± 121.6 nM, respectively. DVS showed weak binding affinity (62% inhibition at 100 μM) at the human dopamine (DA) transporter. Inhibition of [³H]5-HT or [³H]NE uptake by DVS for the hSERT or hNET produced IC₅₀ values of 47.3 ± 19.4 and 531.3 ± 113.0 nM, respectively. DVS (10 μM), examined at a large number of nontransporter targets, showed no significant activity. DVS (30 mg/kg orally) rapidly penetrated the male rat brain and hypothalamus. DVS (30 mg/kg orally) significantly increased extracellular NE levels compared with baseline in the male rat hypothalamus but had no effect on DA levels using microdialysis. To mimic chronic selective serotonin reuptake inhibitor treatment and to block the inhibitory 5-HT₁₈ autoreceptors, a 5-HT₁₈ antagonist, N-[2-[4-(2-methoxyphenyl)-1-piperazinyl]ethyl]-N₂-pyridinylcyclohexanecarboxamide maleate salt (WAY-100635) (0.3 mg/kg s.c.), was administered with DVS (30 mg/kg orally). 5-HT increased 78% compared with baseline with no additional increase in NE or DA levels. In conclusion, DVS is a new 5-HT and NE reuptake inhibitor in vitro and in vivo that demonstrates good brain-to-plasma ratios, suggesting utility in a variety of central nervous system-related disorders.

Biogenic amines such as serotonin (5-HT), norepinephrine (NE), and dopamine (DA) are neurotransmitters found in areas of the central nervous system (CNS) known to be important for regulation of cognitive function, mood, thermoregulation, pain sensation, sexual function, and various aspects of endocrine function related to homeostasis. Their synthesis is highly regulated, and long-term disturbance of the regulatory pathways for these neurotransmitters may lead to disruptions in overall health and quality of life. As with any specific mechanism that controls a physiological process, regulation of these neurotransmitters is a target of many pharmacological agents. Agents that have been developed to modulate these key neurotransmitters are well characterized and have provided evidence that regulation of these neurotransmitters affects physiological outcomes specific to each neurotransmitter. There are multiple means of regulating these neurotransmitters. One means is by inhibiting their presynaptic reuptake. Monoamine reuptake inhibitors work by binding to their respective transporter proteins located presynaptically. This binding interaction results in an increase in elimination time of the neurotransmitter cycle; thus, the extracellular concentration of the neurotransmitters increases in the synaptic cleft. This increase in neurotransmitter in the cleft allows for increased downstream cellular signaling (Blakely and Bauman, 2000). Serotonin and norepinephrine reuptake inhibitors (SNRIs) work by blocking the presynaptic reuptake of 5-HT and NE, resulting in an increased sustained level of both of these neurotransmitters. The 5-HT and NE monoamine transporters are

ABBREVIATIONS: 5-HT, serotonin; NE, norepinephrine; DA, dopamine; CNS, central nervous system; DMSO, dimethyl sulfoxide; DVS, desvenlafaxine succinate; FBS, fetal bovine serum; hDAT, human dopamine transporter; WIN-35,428, (−)-2-β-carbomethoxy-3-β-(4-fluorophenyl)tropane 1.5-naphthalene disulfonate; hNET, human norepinephrine transporter; HPLC, high-performance liquid chromatography; hSERT, human serotonin transporter; PEI, polyethylenimine; MDCK, Madin-Darby canine kidney; SNRI, serotonin and norepinephrine reuptake inhibitor; SSRI, selective serotonin reuptake inhibitor; WAY-100635, N-[2-[4-(2-methoxyphenyl)-1-piperazinyl]ethyl]-N₂-pyridinylcyclohexanecarboxamide maleate.
members of the 12-transmembrane-spanning transporter family that moves monoamines across the cell membrane with high affinity in a sodium-dependent manner (Pacholczyk et al., 1991; Amara and Kuhar, 1993; Uhl and Johnson, 1994).

Currently, two drugs approved in the United States, venlafaxine and duloxetine, are selective inhibitors of neuronal 5-HT and NE reuptake (Muth et al., 1996; Owens et al., 1997; Bymaster et al., 2001). Neither has significant affinity for muscarinic, cholinergic, H₂-histaminergic, or α₁-adrenergic receptors in vitro, supporting the hypothesis that the various anticholinergic, sedative, and cardiovascular effects seen with activity at these receptors will not be present with their clinical use. In essence, the selectivity of these drugs for 5-HT and NE transporters will diminish the potential for adverse effects seen with less selective, centrally active treatments. Both drugs are classified as SNRIs and have been shown to have use in multiple therapeutic end points. Specifically, venlafaxine is indicated for the treatment of major depressive disorder, generalized anxiety disorder, social anxiety disorder, and panic disorder, whereas duloxetine is indicated for the treatment of major depressive disorder (Stahl et al., 2005) and for the treatment of diabetic peripheral neuropathic pain (Goldstein et al., 2005; Kirwin and Goren, 2005).

Desvenlafaxine succinate (DVS), the succinate salt of the isolated major active metabolite of venlafaxine, is pharmaco-logically distinct molecule currently in clinical development. This compound was evaluated in a series of competitive radioligand binding and transport inhibition bioassays at the three human monoamine transporters (5-HT, NE, and DA). In addition, DVS was also examined for potential affinity at a large number of other nontransporter targets to ascertain its selectivity for the 5-HT and NE monoamine transporters. Furthermore, DVS was evaluated in vivo to determine brain penetrability and effect on neurochemical levels in the brain of male rats. The results of these experiments indicate that DVS is a SNRI both in vitro and in vivo. DVS demonstrates good brain-to-plasma ratios, suggesting utility in CNS, peripheral nervous system, and peripheral-related disorders associated with changes in these neurotransmitters.

**Materials and Methods**

**Compounds.** Desvenlafaxine succinate was synthesized by the Discovery Medicinal Chemistry group of Wyeth Research. Desipramine (catalog number D-3900), fluoxetine (catalog number F-132), paroxetine (catalog number P-1372), and mazindol (catalog number M-2017) were purchased from Sigma-Aldrich (St. Louis, MO). Radioligands ([³H]NE, catalog number NET-048, 5–15 Ci/mmol; [³H]nisoxetine, catalog number NET-1084, 85.5 Ci/mmol; [³H]WIN-35,428, cat-alog number NET-1033, 85.6 Ci/mmol; and [³H]bromolytryptamine creatinine sulfate, catalog number NET-498, 25–35 Ci/mmol) and scintillation proximity assay beads (containing 0.5 mg; GE Healthcare, Little Chalfont, Buckinghamshire, UK) were purchased in 50 mm Tris-HCl, 5 and 5 mM NaCl, and 5 mM KCl, pH 7.4) and prepared to approximately 3 ± 1 μg of protein per 150 μl of aliquot. Binding reactions were run in 96-well OptiPlates (PerkinElmer Life and Analytical Sciences). Cell membranes (50 μl) were added to each well of the reaction plate followed by 40 μl of binding buffer, 10 μl of test compound or buffer, or 50 μl of [³H]citalopram (1 nM final concentration). [³H]citalopram was run at the approximate Kᵦ concentration, and 94 to 96% specific binding was achieved in this assay. Data from wells containing fluoxetine (10 μM) were used to define nonspecific hSERT membrane binding (minimum binding in the presence of a 5-HT reuptake inhibitor). All reaction wells received 50 μl of wheat germ agglutinin scintillation proximity assay beads (containing 0.5 mg; GE Healthcare, Little Chalfont, Buckinghamshire, UK) made in 50 mm Tris-HCl, 120 mM NaCl, and 5 mM KCl, pH 7.4. The reaction plates were then incubated for a minimum of 1 h to reach equilibrium. The plates were then counted on a TopCount scintillation counter (PerkinElmer Life and Analytical Sciences).

**NE Radioligand Membrane Binding Assay.** Fresh or frozen membranes were suspended in binding buffer (50 mM Tris-HCl, 300 mM NaCl, and 5 mM KCl, pH 7.4) and prepared to approximately 3 ± 1 μg of protein per 150 μl of aliquot. Binding reactions were run in polystyrene 96-well plates (general assay plate, catalog number 3359; lid, catalog number 3930; Corning Life Sciences, Acton, MA). Cell membranes (150 μl) were added to each well of a reaction plate followed by 10 μl of compound solution. A stock solution of desipramine was prepared in dimethyl sulfoxide (DMSO; 1 mM) and delivered to triplicate wells containing membranes for a final test concentration of 1 μM. Data from wells containing desipramine (1 μM) were used to define nonspecific hNET membrane binding (minimal binding in the presence of an NE reuptake inhibitor). Total radioligand bound is defined by the addition of 10 μl of binding buffer alone in the presence of [³H]nisoxetine. Stock solutions of DVS were prepared in DMSO/H₂O (1:1) at concentrations from 10 μM to 10 mM. On the day of the assay, DVS was diluted in assay buffer according to the test range (10–30,000 nM). Compounds and membranes were preincubated at 4°C for 15 min prior to initiating the binding reaction by the addition of radioligand. The binding reaction was initiated by the addition of 50 μl of [³H]nisoxetine diluted in binding buffer to yield a final assay concentration of 3 nM. The equilibrium binding affinity constant (Kᵦ) value estimated for [³H]nisoxetine was 4 nM using membranes containing hNET. The assay run under these conditions showed a 90 to 96% specific binding signal. The reaction was incubated on an orbital shaking platform (Belloco, Vineland, NJ) for 1 h at room temperature at a speed of 3 revolutions per min. The MultiScreen-PB opaque 96-well filtration plates containing glass fiber filters (glass fiber B, catalog number MAFBNOB; Millipore Corporation, Billerica, MA) were used to terminate the binding reactions and to separate bound from free radioligand. The plates were presoaked with 0.5% polyethyleneimine (PEI, catalog number P-3143;
Sigma-Aldrich) in water for a minimum of 2 h at room temperature to reduce nonspecific binding of [3H]nisoxetine during the harvest of the binding reactions. The PEI solution was aspirated using a vacuum manifold from the filter plates just before completion of the reaction incubation time. Aliquots of each reaction (180 μl of each 200-μl reaction well) were transferred from the reaction plates to the filter plates using a Zymark Rapid Plate-96 automated pipette station (Caliper Tech Corp., Hopkinton, MA). The binding reaction was terminated by vacuum filtration through the glass fiber filters. The filter plates were aspirated at 5 to 10 inches of Hg, and the wells were washed seven times with 200 μl of wash buffer (50 mM Tris-HCl, 0.9% NaCl, pH 7.4, 4°C) using a 12-channel aspiration/wash system. Plastic bottom supports were removed from the filter plates, and the plates were placed in plastic liners. A 50-μl aliquot of scintillation fluid was added to each well, and the top of each plate was sealed with adhesive film. The plates were vigorously shaken on an orbital shake table (Belloco) at 5 rpm for 10 to 15 min to ensure adequate equilibration of aqueous-to-solvent partitioning. The collection of raw counts per minute data were done using a Wallac Microbeta counter (PerkinElmer Life and Analytical Sciences).

**NE Whole-Cell Radioligand Binding Assay.** The hNET whole-cell radioligand binding assay protocol was developed based on the combined methods supplied with hNET membranes (catalog number RBHNETM; PerkinElmer Life and Analytical Sciences), and the hNET functional uptake assay. Twenty-four hours before assay, cells were plated in 96-well plates at 3000 to 5000 cells/well in growth medium and maintained in a cell incubator (37°C, 5% CO2). On day 2, growth medium was replaced with 75 μl of assay buffer (25 mM HEPES, 120 mM NaCl, 5 mM KCl, 2.5 mM CaCl2, 1.2 mM MgSO4, and 2 mg/ml ascorbic acid and 1 μM pargyline). Five-microliter aliquots of DVS in assay buffer were added directly to triplicate wells to yield final test concentrations of 10 to 30,000 nM. Data from wells containing desipramine (1 μM) were used to define nonspecific hNET binding (minimal hNET binding in the presence of an NE reuptake inhibitor). Total radioligand bound is defined by the addition of 5 μl of binding buffer alone in the presence of [3H]nisoxetine. The radioligand binding reaction was initiated by the addition of [3H]nisoxetine to 25 μl of assay buffer to each well for a final concentration of 3 nM. The Kd value estimated for [3H]nisoxetine was 10 nM using intact whole cells. This assay was modified under these conditions showed a 93 to 96% specific binding signal. The cells in assay buffer with test compound and radioactivity were incubated for 1.5 h at 37°C. The supernatant was decanted to terminate the reaction. The cells were washed twice with 200 μl of assay buffer (37°C, without pargyline and ascorbic acid) to remove unbound radioligand. The plates were inverted and left to dry for 2 min and then reinserted and air-dried for an additional 10 min. Scintillation cocktail (50 μl) was added to each well, and the plates were sealed with film tape. The plates were placed on an orbital shake table for 10 min to ensure adequate mixing. The collection of raw counts per minute data were done using a Wallac Microbeta counter.

**5-HT Uptake Assay.** The methods for 5-HT functional reuptake using the JAR cell line were modified using a previous report (Prasad et al., 1996). On day 1, cells were plated at 15,000 cells/well in 96-well plates containing growth medium (RPMI 1640 with 10% FBS) and maintained in a cell incubator (37°C, 5% CO2). On day 2, cells were stimulated with staurosporine (40 nM) to increase the growth medium to ambient oxygen concentration. Subsequently, the growth medium was replaced with 200 μl of assay buffer (25 mM HEPES, 120 mM NaCl, 5 mM KCl, 2.5 mM CaCl2, 1.2 mM MgSO4, and 2 mg/ml glucose, pH 7.4, at 37°C) containing 0.2 mg/ml ascorbic acid and 1 μM pargyline, and the cells were incubated for 5 min at 37°C. A stock solution of paroxetine was prepared in DMSO (10 mM) and delivered to triplicate wells containing cells for a final test concentration of 1 μM. The cells were incubated with the compound for 10 min (37°C). To initiate the reaction, [3H]hydroxytryptamine creatine sulfate diluted in assay buffer was delivered in 25-μl aliquots to each well for a final test concentration of 15 nM. The cells were incubated with the reaction mixture for 9 min at 37°C. Decanting the super-
natant from the plates terminated the reaction. The cells were washed twice with 200 μl of assay buffer (37°C) to remove free radioligand. The plates were inverted and left to dry for 2 min and then reinverted and air-dried for an additional 10 min. Subsequently, the plates were lysed in 25 μl of 0.25 N NaOH (4°C) and then placed on a shaker table and shaken vigorously for 5 min. After cell lysis, 75 μl of scintillation cocktail was added to the wells, and the plates were sealed with film tape and replaced on the shake table for a minimum of 10 min. The plates were counted in a Wallac Microbeta counter to collect the raw counts per minute data.

NE Uptake Assay. On day 1, cells were plated at 3000 cells/well in a 96-well plate in growth medium and maintained in a cell incubator (37°C, 5% CO2). On day 2, growth medium was replaced with 200 μl of medium containing 0.1% DMSO/H2O (1:1) (10 mM) and diluted in assay buffer according to the test concentration (minimal NE uptake) or various concentrations of DVS were added to initiate the NE uptake, [3H]NE diluted in assay buffer (120 nM final concentration) was delivered in 25-μl aliquots to each well, and the plates were incubated for 10 min at 37°C. Decanting the supernatant from the plates terminated the reaction. The plates containing cells were washed twice with 200 μl of assay buffer (37°C) to remove free radioligand. The plates were then inverted, left to dry for 2 min, and then reinverted and air-dried for an additional 2 min. The cells were lysed in 25 μl of 0.25 N NaOH (4°C), placed on an orbital shake table, and vigorously shaken for 5 min. After cell lysis, 75 μl of scintillation cocktail was added to each well, and the plates were sealed with film tape. The plates were returned to the shake table and vigorously shaken for a minimum of 10 min to ensure adequate partitioning of organic and aqueous solutions. The plates were counted in a Wallac Microbeta counter to collect the raw counts per minute data.

Animals. The brain penetration studies were done using male Sprague-Dawley rats purchased from Taconic Farms (Germantown, NY) and housed in a facility accredited by the Association for Assessment and Accreditation of Laboratory Animal Care International that was monitored, and the probe placement was verified by histology. Data from animals with incorrect probe placement were discarded. The probe placement was verified by histology. Data from animals with incorrect probe placement were discarded.

Determination of DVS Concentrations in Plasma, Brain, and Hypothalamus. DVS was prepared in DMSO/H2O (1:1) (10 mM) and diluted in assay buffer according to the test range (1–10,000 nM). Twenty-five microliters of assay buffer (maximal NE uptake) or various concentrations of DVS were added directly to triplicate wells containing cells in 200 μl of assay buffer. The cells were incubated with test compound for 10 min at 37°C. To initiate the NE uptake, [3H]NE diluted in assay buffer (120 nM final assay concentration) was delivered in 25-μl aliquots to each well, and the plates were incubated for 10 min at 37°C. Decanting the supernatant from the plate terminated the reaction. The plates containing cells were washed twice with 200 μl of assay buffer (37°C) to remove free radioligand. The plates were then inverted, left to dry for 2 min, and then reinverted and air-dried for an additional 2 min. The cells were lysed in 25 μl of 0.25 N NaOH (4°C), placed on an orbital shake table, and vigorously shaken for 5 min. After cell lysis, 75 μl of scintillation cocktail was added to each well, and the plates were sealed with film tape. The plates were returned to the shake table and vigorously shaken for a minimum of 10 min to ensure adequate partitioning of organic and aqueous solutions. The plates were counted in a Wallac Microbeta counter to collect the raw counts per minute data.

Evaluation of Neurochemical Samples. A 20-μl sample from the hypothalamus was analyzed for NE and 5-HT content by methods described previously (Beyer et al., 2002). In brief, five control samples (ficomole/100 μl sample) were taken prior to drug injection to demonstrate a steady baseline. All subsequent values were expressed as a percentage of the mean preinjection value (percentage of baseline). After baseline sampling was complete, animals were treated with WAY-100635 (0.3 mg/kg s.c.) or vehicle (water s.c.). DVS (30 mg/kg) or vehicle (0.2% Tween 80 and 0.5% methylcellulose dissolved in water) was orally administered 20 min later at the end of the sixth sample. A 3-h stabilization period was allowed after probe implantation, and dialysate sampling was conducted according to previously described methods (Beyer et al., 2002). In brief, control samples (fimole/20-μl sample) were taken prior to drug injection to demonstrate a steady baseline. All subsequent values were expressed as a percentage of the mean preinjection value (percentage of baseline). After baseline sampling was complete, animals were treated with WAY-100635 (0.3 mg/kg s.c.) or vehicle (water s.c.). DVS (30 mg/kg) or vehicle (0.2% Tween 80 and 0.5% methylcellulose dissolved in water) was orally administered 20 min later at the end of the sixth sample. A 3-h stabilization period was allowed after probe implantation, and dialysate sampling was conducted according to previously described methods (Beyer et al., 2002). In brief, control samples (fimole/20-μl sample) were taken prior to drug injection to demonstrate a steady baseline. All subsequent values were expressed as a percentage of the mean preinjection value (percentage of baseline). After baseline sampling was complete, animals were treated with WAY-100635 (0.3 mg/kg s.c.) or vehicle (water s.c.). DVS (30 mg/kg) or vehicle (0.2% Tween 80 and 0.5% methylcellulose dissolved in water) was orally administered 20 min later at the end of the sixth sample.
methanol, pH 4.8) was delivered by a PU1550 HPLC pump (Jasco Ltd, Essex, UK) at a flow rate of 0.5 ml/min. Neurochemical data were captured using the Atlas software package (Thermo Lab-systems, Beverly, MA).

Data Collection and Statistical Analysis. For all in vitro experiments, with the exception of hSERT binding, a data stream of counts per minute values collected from the Wallac Microbeta counter was downloaded to a Microsoft Excel statistical application program. Calculations of IC_{50} values were made using the transformed-logistic concentration-response program written by Wyeth Biometrics Department. The statistical program uses mean counts per minute values from wells representing maximal binding or uptake (assay buffer) and mean counts per minute values from wells representing minimal binding or uptake [1 uM desipramine (hNET), 10 uM mazindol (hDAT), or 10 uM fluoxetine (hSERT)]. Estimation of the IC_{50} values were completed on a log scale, and the line was fit between the maximum and minimum binding or uptake values. Pooling the raw data from multiple experiments and analyzing the pooled data as one experiment were done to generate the IC_{50} values. For hSERT binding, triplicate counts per minute values per concentration were collected from the PerkinElmer TopCount and analyzed by a Prism nonlinear regression analysis program (GraphPad Software Inc., San Diego, CA) for receptor binding. Calculation of the K_{i} values was based on the K_{D} values of the radioligand for each bioassay. The K_{i} value is a function of the concentration of the compound required to inhibit 50% of the radioligand (IC_{50} value) divided by the free radioligand concentration [L] divided by the K_{D} value plus one [K_{i} = IC_{50}/(1 + [L]/K_{D})] (Cheng and Prusoff, 1973). Experiments evaluating DVS were completed a minimum of three times in separate experiments for all assays described within this report.

Pharmacokinetic data were subject to noncompartmental analysis using WinNonLin version 4.1 software. Tissue concentration ratios were calculated as the concentration in a given brain tissue divided by the concentration in plasma at a given time point.

In the microdialysis experiments, the ficmole concentrations for the baseline samples were averaged, and this value was denoted as 100%. Subsequent sample values were expressed as a percentage change from this preinjection baseline value (percentage change from baseline). Neurochemical data, excluding preinjection values, were analyzed by a two-way analysis of variance with repeated measures (time). Post hoc analyses were made using the Bonferroni/Dunn adjustment for multiple comparisons. All statistical calculations were performed using the Statview software application (Abacus Concepts Inc., Berkeley, CA).

Results

Determination of Affinities of DVS for Human Monoamine Transporters

5-HT Membrane Radioligand Binding. Competitive [3H]citalopram (SSRI) scintillation proximity assays were performed using membranes prepared from human embryonic kidney 293 cells overexpressing hSERT to determine the affinity of DVS (Fig. 1A). Data from three separate experiments demonstrated that DVS is a potent inhibitor of [3H]citalopram binding at the hSERT with a K_{i} value of 40.2 ± 1.6 nM (Table 1). NE Whole-Cell and Membrane Radioligand Binding. In MDCK-hNET6 cells, DVS competitively inhibited binding of a known NE reuptake inhibitor, [3H]nisoxetine (Fig. 1B), with a K_{i} value of 558.4 ± 121.6 nM (Table 1). In further evaluation using membranes containing the hNET, DVS competed for the binding of [3H]nisoxetine with a K_{i} value of 3385.1 ± 349.3 nM. The diminished affinity of DVS in the membrane bioassay may reflect the artificial nature of using isolated membranes in contrast to intact cells. Minor technical variations in these two radioligand binding assays may account for this discrepancy.

DA Binding. To determine the activity of DVS for this third human monoamine transporter, competitive binding assays using [3H]WIN-35,428, a known selective DA reuptake inhibitor, were performed using purified membranes prepared from Chinese hamster ovary cells that overexpress hDAT. Data from three separate experiments demonstrated that DVS was a very weak inhibitor of [3H]WIN-35,428 binding at the hDAT, with only a 61.6% ± 1.7 inhibition at the highest concentration tested (100 uM; Table 1). Therefore, an
extrapolated logistic concentration curve was generated to determine a $K_i$ value (25 ± 5 μM) (Fig. 1C).

**Inhibitory Effects of DVS on Radioligand Uptake at hSERT and hNET.** DVS inhibited the uptake of $[^3]$H]5-HT at the hSERT (Table 1). The data generated and pooled from three separate experiments show a mean $IC_{50}$ value of 47.3 ± 19.4 nM (Fig. 2A). In addition, the inhibitory action of DVS on uptake of $[^3]$H]NE in MDCK-hNET6 cells was evaluated. Pooled data from nine separate experiments indicate that the $IC_{50}$ value of DVS for $[^3]$H]NE uptake was 531.3 ± 113.0 nM (Fig. 2B). Desipramine, a well characterized NE reuptake inhibitor, was tested in parallel in all NE transporter assays and showed a similar affinity in all three hNET assays with a $K_i/IC_{50}$ value of 3.4 ± 0.2 nM. DVS showed an approximate 10-fold separation between hSERT and hNET activity, with the reported activity for hSERT greater than hNET.

**Selectivity of DVS**

DVS was examined for potential affinity at a large number of other nontransporter targets to ascertain its selectivity for the 5-HT and NE monoamine transporters. DVS (10 μM) was tested for activity at 96 targets (i.e., receptors, transporters, enzymes, and channels) by NovaScreen (Hanover, Maryland) (Table 2). No significant activity was detected for any targets, with the exception of the 5-HT and NE monoamine transporters, demonstrating the selectivity of this compound for the two monoamine transporters.

**DVS Concentrations in Plasma, Brain, and Hypothalamus after Oral Administration.** Concentrations of DVS were determined in the hypothalamus and total brain for each time point over a 24-h period (Fig. 3). Total brain represents the remainder of brain tissue after dissecting the hypothalamus. The $C_{max}$ value in plasma of 940 ng/ml occurred at 30 min after DVS dosing. The terminal half-life of DVS in plasma was 2.1 h, and the compound was undetectable 24 h after the single oral dose. Extrapolated total exposure ($AUC_{0-24}$) in plasma was 1864 h · ng/ml. Similar to plasma, the $C_{max}$ value in the hypothalamus and 771 ng/g in total brain tissue was reached 30 min after dosing. The half-life of DVS was 2.2 h in hypothalamus and 2.1 h in total brain. Extrapolated total exposure ($AUC_{0-24}$) in hypothalamus was 3485 h · ng/g and 2923 h · ng/g in total brain. DVS was below the limits of detection in hypothalamus and total brain tissue in all rats at the 24-h time point. The brain-to-plasma ratio of DVS reached a maximum in hypothalamus (2.6) at 8 h and in total brain (2.2) at 2 h (data not shown). The ratio of hypothalamus AUC to plasma AUC was 1.8, and that of brain AUC to plasma AUC was 1.6.

**Effect of DVS on Hypothalamic Levels of Monoamines (5-HT, NE, and DA).** Acute oral administration of DVS (30 mg/kg) did not significantly [$F(2,19) = 0.74, P = 0.4898$] alter concentrations of 5-HT in the hypothalamus (Fig. 4A). However, pretreatment with WAY-100635 (5-HT1A antagonist, 0.3 mg/kg s.c.), which did not alter 5-HT levels on its own, resulted in a significant 78% increase in extracellular 5-HT levels [$F(1,9) = 36.09, P = 0.0001$] in the rat hypothalamus. In contrast to the observations with 5-HT levels, acute administration of DVS, when given alone, produced a significant [$F(2,16) = 6.11, P = 0.0107$] increase in NE con-

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**TABLE 1**

Characterization of desvenlafaxine at the human serotonin, norepinephrine, and dopamine transporters

<table>
<thead>
<tr>
<th>Radioligand Binding</th>
<th>Neurotransmitter Uptake</th>
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<tr>
<td></td>
<td>$K_i$ Values</td>
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<tr>
<td></td>
<td>hSERT Membrane</td>
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<tr>
<td>Desvenlafaxine</td>
<td>40.2 ± 1.6</td>
</tr>
<tr>
<td>Comparator</td>
<td>7.0 ± 1.2</td>
</tr>
<tr>
<td>Fluoxetine</td>
<td>Desipramine</td>
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$IC_{50}$, concentration inhibiting 50% of the radioligand uptake; hSERT, human serotonin transporter; hNET, human norepinephrine transporter; hDAT, human dopamine transporter.

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**Fig. 2.** Functional activity of desvenlafaxine demonstrates inhibition of radioligand uptake of serotonin (A) or norepinephrine (B). For each uptake transporter bioassay, a known comparator was used [fluoxetine (5-HT reuptake inhibitor) and desipramine (NE reuptake inhibitor)]. Each data point depicted represents the mean ± S.E.M. of three independent experiments performed in triplicate. The $IC_{50}$ values for desvenlafaxine are shown in Table 1.
Targets Examined

Desvenlafaxine Selectivity Profile: Lack of Significant Affinity at Numerous Targets Examined

<table>
<thead>
<tr>
<th>Neurotransmitter-related</th>
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<tr>
<td>Adenosine transporter</td>
<td>Adenosine A&lt;sub&gt;1&lt;/sub&gt; and A&lt;sub&gt;2&lt;/sub&gt;</td>
<td>Benzo diazepam periphery (h)</td>
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<tr>
<td>Cannabinoid CB1 and CB2 (h)</td>
<td>Dopamine transporter</td>
<td>Dopamine 1–5 (h)</td>
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<td>Glutamate (AMPA, Kiant e)</td>
<td>GABA&lt;sub&gt;A&lt;/sub&gt;, agonist</td>
<td>GABA&lt;sub&gt;A&lt;/sub&gt;, BDZ</td>
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<td>Purinergic, P2Y (h)</td>
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<tr>
<td>Brain/gut peptides</td>
<td>AT1 (h) and AT2</td>
<td>BK2</td>
</tr>
<tr>
<td>CCK A and B</td>
<td>ET A and B (h)</td>
<td>Galanin</td>
</tr>
<tr>
<td>Steroids</td>
<td>Glucocorticoid</td>
<td>Pregestosterone</td>
</tr>
<tr>
<td>Testosterone (cytosolic and nuclear)</td>
<td>Ion channels</td>
<td>Calcium channel L (Benz)</td>
</tr>
<tr>
<td>Calcium channel L (Dihyd)</td>
<td>Calcium channel N</td>
<td>K&lt;sup&gt;+&lt;/sup&gt; channel, ATP and CA2</td>
</tr>
<tr>
<td>Sodium, sites 1 and 2</td>
<td>GABA, chloride</td>
<td>Prostaglandins</td>
</tr>
<tr>
<td>Prostaglandins</td>
<td>Leukotriene B&lt;sub&gt;4&lt;/sub&gt;, D&lt;sub&gt;4&lt;/sub&gt;</td>
<td>Thromboxane A&lt;sub&gt;2&lt;/sub&gt; (h)</td>
</tr>
<tr>
<td>Second messengers</td>
<td>Nitric-oxide synthase</td>
<td>Adenylate cyclase</td>
</tr>
<tr>
<td>IP&lt;sub&gt;3&lt;/sub&gt;</td>
<td>Protein kinase C</td>
<td>Growth factors/hormones</td>
</tr>
<tr>
<td>Oxytocin Corticotropin-releasing factor</td>
<td>Platelet-activating factor</td>
<td>Thyrotropin-releasing hormone</td>
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<tr>
<td>Neurokinin 1–3 (h)</td>
<td>Neuropeptide 1–2 (h)</td>
<td>Neurotensin</td>
</tr>
<tr>
<td>Somatostatin</td>
<td>Vasopressin intestinal peptide (h)</td>
<td>Vasopressin</td>
</tr>
<tr>
<td>Other</td>
<td>Complement e5a (h)</td>
<td>Sigma 1 and 2</td>
</tr>
</tbody>
</table>

DVS selectively inhibits serotonin and norepinephrine uptake<sup>a</sup>

Desvenlafaxine (10 μM) was tested for activity at numerous drug targets by NovaScreen. No significant activity was detected, with the exception of the 5-HT and NE transporters.

TABLE 2

Desvenlafaxine Selectivity Profile: Lack of Significant Affinity at Numerous Targets Examined

![Graph showing tissue conc. over time](image)

**Fig. 3.** Orally administered desvenlafaxine (30 mg/kg) to intact male rats is found in plasma and penetrates the brain and hypothalamus over time. Each time point depicted in the figure represents the mean concentration ± S.E.M. of desvenlafaxine from three individual rats. A standard curve (5–2000 ng/ml) was generated in plasma, brain, and hypothalamus from vehicle-treated rats.

Concentrations (Fig. 4B). Post hoc analysis revealed significant differences in maximal NE levels, comparing doses of 10 (data not shown) and 30 mg/kg to vehicle (96% above baseline; P = 0.0221 and 118% above baseline; P = 0.0034, respectively), although the effects of the two doses of DVS were not significantly different from each other (P = 0.4470).

Pretreatment with WAY-100635 (0.3 mg/kg s.c.) did not significantly increase NE in this brain region when tested alone [F(2,15) = 1.986, P = 0.1768]. However, when combined with DVS, pretreatment with WAY-100635 did not alter the ability of DVS to significantly elevate extracellular NE levels [F(1,9) = 8.11, P = 0.0192]. The neurochemical effects of DVS were found to be specific for noradrenergic and serotonergic systems, because the acute administration of DVS (30 mg/kg p.o.) alone or in combination with WAY-100635 (0.3 mg/kg s.c.) did not significantly alter concentrations of DA in the hypothalamus [F(2,23) = 0.18, P = 0.8343; Fig. 4C].

Discussion

DVS is a novel succinate salt monohydrate of desvenlafaxine, the major active metabolite of venlafaxine. The results of these in vitro experiments show that DVS exhibits selective inhibitory activity of neurotransmitter uptake at the human 5-HT and NE monoamine transporters and increases extracellular levels of NE and 5-HT (in the presence of a 5-HT<sub>1A</sub> antagonist) when compared with baseline levels of monoamines in the hypothalamus of rats. Higher affinity was noted for the hSERT compared with the affinity for the hNET, whereas weak affinity for the hDAT was noted. DVS (10 μM) demonstrated no significant activity at numerous nontransporter targets. In addition, oral administration of DVS resulted in measurable concentrations in the brain. Taken together, these data support that this new chemical entity would be classified as a new SNRI.

The results demonstrate that DVS potently interacts with hSERT and, to a lesser degree, with hNET. The in vitro functional assay indicated that DVS is approximately 10-fold more potent at inhibiting 5-HT uptake than NE uptake. DVS competes for the binding of a known NE reuptake inhibitor [3H]nisoxetine in cells containing the hNET. These data support the hypothesis that DVS interacts with the hNET to inhibit NE uptake. In addition, DVS inhibits functional NE uptake in the same cells, supporting a functional conse-
The sequence of the interaction of DVS with the hNET. Competition of DVS for the binding of [3H]WIN-35,428, a known selective DA reuptake inhibitor, only showed inhibitory activity at high micromolar concentrations (\( \mu \text{M} \)). Based on these assays that were performed to evaluate the binding interaction of DVS, this weak binding interaction would suggest that no functional consequence on dopamine levels would occur at concentrations required to inhibit hNET and hSERT.

Studies comparing the receptor and transporter binding profile of several SSRIs and SNRIs have illustrated that these drugs and their metabolites each have unique functional activity ratios (Owens et al., 1997; Bymaster et al., 2001). However, in vitro assays may not accurately depict in vivo activity because of factors such as pharmacokinetics, blood/brain permeability, protein binding, and clearance of compound. In addition, the amino acid binding sites for specific drugs versus different radioligands can influence apparent in vitro affinities. Although DVS showed modest affinity for the hNET, its action at these sites is likely to be functionally important. Based on currently available data, it is unclear what levels of 5-HT and NE are required to restore “normal” function. Although the efficacy of other SNRIs has been shown for the treatment of specific disorders, the optimal activity ratio for the hSERT and hNET has not been determined for any specific disease indication.

It is postulated that, at clinically relevant doses, acute treatment with SSRIs and SNRIs do not acutely elevate extracellular 5-HT levels in terminal brain areas. The postulated reason for this lack of serotonergic effect is that increases in terminal neurotransmitter concentrations, resulting from blockade of transporter, are paralleled by increases in the somatodendritic cell body region of the dorsal raphe. Local increases in raphe 5-HT, via activating the inhibitory 5-HT<sub>1A</sub> autoreceptors in this region, will suppress serotonergic neuronal cell firing and result in little or no 5-HT release in terminal brain regions (see Discussion in Beyer et al., 2002). Reversing this inhibitory feedback loop with 5-HT<sub>1A</sub> receptor antagonists (e.g., WAY-100635) or chronic antidepressant treatment (i.e., 5-HT<sub>1A</sub> receptor desensitization) causes an immediate increase in extracellular 5-HT in the presence of an SSRI or SNRI, probably mimicking their long-term effects (Chaput et al., 1986; Jolas et al., 1994; Kreiss and Lucki, 1995; Le Poul et al., 1995, 2000; Beyer et al., 2002). This neurochemical hypothesis is corroborated by preclinical findings showing that acute combination treatment with a 5-HT<sub>1A</sub> antagonist and antidepressants produces similar increases in extracellular 5-HT to those seen following repeated antidepressant treatment alone (Kreiss and Lucki, 1995). Because it is speculated that combining 5-HT<sub>1A</sub> antagonism with acute DVS treatment would mimic the neurochemical effects of chronic DVS treatment, the results of the present studies suggest that chronic treatment with DVS is likely to result in elevations in 5-HT in this brain region over time. Similar to additional microdialysis studies conducted in other areas of the brain, including the frontal cortex (Wyeth Research, unpublished findings), DVS produced significant increases in extracellular NE concentrations in the hypothalamus. As expected and similar to reports with other transport blockers (e.g., SSRIs), the addition of WAY-100635 did not affect the ability of DVS to increase extracellular NE levels compared with baseline NE levels (Beyer et al., 2002).

The present study evaluated brain concentrations and brain-to-plasma concentration ratios of DVS over time after oral administration in male rats. DVS demonstrates rapid brain penetration with brain concentrations in excess of those noted in the plasma after a single oral dose of 30 mg/kg. This work supports the notion that DVS can exert direct effects on neuronal systems of the brain and that it can have utility in treating CNS-related dysfunctions. In addition, increasing 5-HT and NE levels in the local brain environment may result in increased levels of these key neurotransmitters in the circulation. These transmitters play an important role in the peripheral nervous system, as well as in the periphery (Teschemacher, 2005), and thus may have utility in treating disorders both centrally and peripherally that are associated with modified levels of 5-HT and NE, such as various pain.
modalities (Willis and Westlund, 1997; Sommer, 2004), inflammatory processes (Kaneko et al., 2005; Maleki et al., 2005), and urinary incontinence (Fraser and Chancellor, 2003).

Compounds with dual-acting 5-HT and NE monoamine transporter inhibition have been shown to have multiplicity utilities (Sayar et al., 2003; Schober and Ansani, 2003; Stearns et al., 2003; Arnold et al., 2004; Iyengar et al., 2004; Ladd et al., 2005; Mariappan et al., 2005). DVS, a new molecule in this class, is brain-penetrable and increases key neurotransmitters in the hypothalamus, a region of the brain involved in important physiological functions, including regulation of mood, temperature, and pain sensation. In addition, the hypothalamus is important in regulation of the sleep cycle, stress response, and sexual behavior. Based on these preclinical findings, DVS appears to be a selective and potent SNRI that warrants further evaluation in a variety of therapeutic end points associated with disrupted 5-HT and NE levels. The utility and precise functional activity at both the serotonergic and noradrenergic transporters of this new chemical entity in disorders, implicated to have decreased neurotransmitter levels and altered functional activity, needs to be explored clinically.

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