RESEARCH ARTICLE

In Vitro and In Vivo Characterization of the Alkaloid Nuciferine

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Abstract

Rationale

The sacred lotus (Nelumbo nucifera) contains many phytochemicals and has a history of human use. To determine which compounds may be responsible for reported psychotropic effects, we used in silico predictions of the identified phytochemicals. Nuciferine, an alkaloid component of Nelumbo nucifera and Nymphaea caerulea, had a predicted molecular profile similar to antipsychotic compounds. Our study characterizes nuciferine using in vitro and in vivo pharmacological assays.

Methods

Nuciferine was first characterized in silico using the similarity ensemble approach, and was followed by further characterization and validation using the Psychoactive Drug Screening Program of the National Institute of Mental Health. Nuciferine was then tested in vivo in the head-twitch response, pre-pulse inhibition, hyperlocomotor activity, and drug discrimination paradigms.
Results

Nuciferine shares a receptor profile similar to aripiprazole-like antipsychotic drugs. Nuciferine was an antagonist at 5-HT$_{2A}$, 5-HT$_{2C}$, and 5-HT$_{2B}$, an inverse agonist at 5-HT$_{7}$, a partial agonist at D$_2$, D$_5$ and 5-HT$_{6}$, an agonist at 5-HT$_{1A}$ and D$_4$ receptors, and inhibited the dopamine transporter. In rodent models relevant to antipsychotic drug action, nuciferine blocked head-twitch responses and discriminative stimulus effects of a 5-HT$_{2A}$ agonist, substituted for clozapine discriminative stimulus, enhanced amphetamine induced locomotor activity, inhibited phencyclidine (PCP)-induced locomotor activity, and rescued PCP-induced disruption of prepulse inhibition without induction of catalepsy.

Conclusions

The molecular profile of nuciferine was similar but not identical to that shared with several approved antipsychotic drugs suggesting that nuciferine has atypical antipsychotic-like actions.

1. Introduction

The lotus plants, *Nelumbo nucifera* and *Nymphaea caerulea*, have been used by cultures, both past and present, for their medicinal properties.[1] In eastern medicine, one of the cited potential medical effects of the lotus is "calming emotional disturbance".[2] The alkaloid nuciferine (Fig 1) is thought to be responsible for the psychotropic effects of *Nelumbo nucifera* and *Nymphaea caerulea*, though its pharmacological properties are not entirely clear. Macko and colleagues [3] observed that nuciferine produces effects similar to those of the antipsychotic chlorpromazine in rodents. Bhattacharya and colleagues [4] observed that nuciferine produces antipsychotic-like

![Fig 1. The chemical structure of nuciferine.](https://doi.org/10.1371/journal.pone.0150602.g001)
behavior in rats, including inhibition of the conditioned avoidance response and amphetamine-induced behaviors. The behavioral effects previously observed in rodents, in addition to the cited potential medical effects of the lotus in eastern medicine, led the authors to hypothesize that nuciferine has a pharmacological profile similar to that of antipsychotic medications.

Most antipsychotics share common G protein-coupled receptor (GPCR) targets including the D2 dopamine and 5-HT2A serotonin receptors. [5–7] Beyond these shared targets, antipsychotic compounds exhibit diverse receptor affinity profiles. For example, the atypical antipsychotic clozapine binds with nanomolar affinities to nearly 50 targets (Fig 2). [8] This polypharmacological profile has been suggested as a path forward in therapeutic drug development.[6, 9] Patterns of pharmacological activity responsible for antipsychotic efficacy remain the subject of ongoing investigations. This polypharmacology approach presents a conundrum for drug discovery efforts as it is impossible to design a compound for a polypharmacological "target" (a pattern of molecular activity that engenders therapeutic efficacy) that has not yet been elucidated. We therefore used ethnobotanical records of *Nelumbo nucifera* and *Nymphaea caerulea* medicinal properties as suggestive evidence for potential therapeutic efficacy of a novel polypharmacological profile. Our *in silico* predictions of all phytochemicals identified in *Nelumbo nucifera* suggest that nuciferine (and its metabolites) cross the blood-brain barrier and have multiple protein targets. Furthermore, nuciferine has been shown to cross the blood brain barrier in rats.[10] These predictions and previously reported data suggest that nuciferine has a rich polypharmacology that is responsible for its psychotropic effects. We therefore investigated the *in vitro* and *in vivo* properties of nuciferine using cell-based pharmacology assays and animal behavioral models of antipsychotic drug action.

2. Materials and Methods

2.1 Drugs

Nuciferine was purchased from Sequoia Research Products (Pangbourne, United Kingdom) and Angene (Hong Kong, China) and was dissolved in DMSO at 10 mM for in vitro studies or in 0.9% saline with 1 drop 85% lactic acid per 50 ml for animal studies. D-amphetamine (AMPH), phencyclidine (PCP), and 2,5-dimethoxy-4-iodoamphetamine (DOI) were purchased from Sigma-Aldrich (St. Louis, MO, USA) and were dissolved in 0.9% saline. Clozapine (Sigma-Aldrich) was dissolved in 0.2% acetic acid—2% cyclodextran solution. For animal studies, all drugs were administered intraperitoneally unless otherwise noted.

2.2 Bioinformatics

The profile of phytochemicals in *Nelumbo nucifera* was obtained from Mukherjee et al.[11] The similarity ensemble approach (SEA) was utilized to predict molecular targets for each phytochemical, using Scitegic ECFP4 fingerprints on a target panel extracted from a "binding" subset of ChEMBL-12 [12] and standardized as previously described. [13, 14] Briefly, The SEA [13–15] uses the chemical similarity of a bait molecule, against that of a set of ligands annotated to a target, to predict whether the bait molecule will modulate that target. Briefly, SEA calculates the similarity of the bait molecule to every annotated ligand, typically using topological fingerprints such as ECFP4. Similarity is calculated as the Tanimoto coefficient (Tc), the number of feature (bits) in common between the bait molecule and any given ligand, divided by the total number of features (bits) in the two molecules; identical molecules will have Tc values of 1.0. The Tc values above a threshold value against all the ligands for the target are averaged, and that Tc is compared to that expected for a set of ligands of similar size that would be expected at random. An E-value is calculated by calculating a Z-score for the observed average Tc vs. the ligand set, then plotting this value against an extreme value distribution and using
Fig 2. Comparison of the polypharmacology of nuciferine with atypical and typical antipsychotics. The empirical affinity values of three antipsychotics are shown (clozapine, haloperidol, aripiprazole). The empirical nuciferine affinity profile from this study is shown in comparison. Values for clozapine, haloperidol, and aripiprazole compiled from data available on the PDSP website accessed 20150428. Only “PDSP verified” data was used for the figure. Data entries of “>10,000” were entered as 10,000 μM.

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the BLAST sequence comparison machinery. This E-value represents the likelihood of seeing the similarity one does, between the bait molecule and the known ligands for any given target, compared to what one would expect at random. This calculation is repeated against all of the > 2500 targets in ChEMBL [16] (https://www.ebi.ac.uk/chembl/). The blood-brain barrier penetrability of each compound was predicted using the online blood-brain barrier prediction (BBB) server [17] (http://www.cbligand.org/BBB/).

2.3 Psychoactive drug screening program affinity and functional profiling


For affinity determination, nuciferine was subjected to primary radioligand binding assays tested at a single 10 μM concentration to displace 50% of the radioligand at a given receptor target. If a more than 50% of the radioligand was displaced, nuciferine was selected for a secondary binding assay tested at 11 concentrations in triplicate in competition with the radioligand to generate an IC50 and Ki. Binding assays were performed in 96-well plates with 125 μL per well in appropriate binding buffer using radioligand at or near the Kd. Plates are incubated at room temperature in the dark for 90 min. Reactions are stopped by vacuum filtrations onto 0.3% polyethylenimine soaked 96-well filter mats using a 96-well Filtermate harvester, followed by at least three washes of cold wash buffer. Scintillation (MeltiLex) cocktail is melted onto dried filters and radioactivity is counted using a Wallac Trilux Microbeta (Perkin Elmer).

For receptor functional assays, Gs or Gi-coupled receptor activation was measured using a split-luciferase cAMP biosensor, GloSensor (Promega), and Gq-coupled receptor activation was measured as calcium flux using Fluo-4 Direct Dye (Invitrogen). HEKT cells (ATCC) transiently transfected or cells stably expressing the receptor were plated into 384-white (GloSensor) or black plates (Calcium flux) in DMEM containing 1% dialyzed FBS at least 6 hours to 24 hours before the assay. For GloSensor, the media was decanted and replaced with 20 μL drug buffer per well (HBSS, 20 mM HEPES, pH 7.4) containing GloSensor substrate. Cells were challenged with 10 μL of nuciferine or positive control (3X) to generate 16 point concentration curves and incubated for 15 minutes. For Gs-mediated cAMP accumulation, plates were read immediately. For Gi-mediated cAMP inhibition, 10 μL of isoproterenol (200 nM final concentration) was added to stimulate cAMP via endogenous β-adrenergic receptors and plates were read 15 minutes later. Luminescence was measured using Wallac TriLux Microbeta (Perkin Elmer) and luminescent counts per second (LCPS) was plotted. For Gq-mediated calcium flux, media was decanted and replaced with 20 μL drug buffer containing 2.5 mM probenecid and Fluo-4 dye and allowed to incubate for at least one hour at 37°C and 5% CO2 in a humidified incubator. Afterwards, 10 μL of nuciferine or positive control (3X) was added per well for 16 point concentrations and fluorescence was measured using FLIPR TETRA (Molecular Devices). Maximum-fold increase over basal fluorescence was plotted. Results were analyzed using non-linear regression to obtain EC50 using Graphpad Prism 5.0.

2.4 Dynamic light scattering to test for colloidal aggregation

Nuciferine was diluted into filtered water from a 10 mM stock in DMSO with 50 mM potassium phosphate, pH 7.0. Measurements were made at room temperature using a DynaPro MS/ X (Wyatt Technology) with a 55 mW laser at 826.6 nm. The laser power was 100%, and the detector angle was 90° with samples run in duplicate.
2.5 AmpC β-lactamase assay counterscreen to test for colloidal aggregation

AmpC β-lactamase inhibition was measured in 50 mM potassium phosphate (pH 7.0) at room temperature as described. [21] Nuciferine was diluted from a 10 mM stock in DMSO and incubated with 1 nM AmpC for 5 minutes before the reaction was initiated by adding 92 μM CENTA substrate (Tydock Pharma; Modena, Italy). The final reaction volume was 1 mL. Change in absorbance was monitored at 405 nm for 2 minutes using an HP 8453 UV-Vis spectrophotometer. The assay was performed in duplicate in methacrylate cuvettes.

2.6 Vesicular monoamine transporter (VMAT2) study

Cell culture: HEK293 cells (ATCC) lines stably expressing hDAT or hDAT and hVMAT2 constructs were cultured at 37°C and 5% CO₂ in DMEM with 10% FBS. All constructs were made in pcDNA3.1 (Life Technologies). hDAT and hVMAT2 expressing constructs contained a neomycin or zeocin resistance gene, respectively. Plasmids were transfected into HEK293 cells with Lipofectamine 2000. Stable cell lines were generated by repetitive rounds of limiting dilutions in selective media. Double stable cell lines were created by transfecting HEK-hDAT stable cells with the hVMAT2 construct and selecting for both plasmids with both neomycin and zeocin.

Whole cell ³H-dopamine (DA) uptake: Cells were plated into 48-well plates one day before uptake was performed. Cells were washed with 0.5 ml uptake buffer (4 mM Tris, 6.25 mM HEPES, 120 mM NaCl, 5 mM KCl, 1.2 mM CaCl₂, 1.2 mM MgSO₄, 5.6 mM D-glucose, 1.7 mM ascorbic acid, and 1 μM pargyline, pH 7.4). Cells were incubated with 225 μl uptake buffer with or without the indicated concentration of nuciferine for 15 minutes. After incubation, 25 μl uptake buffer containing ³H-DA and DA was added for a final concentration of 20 nM ³H-DA and 1 μM DA. Cells were incubated at 37°C for 20 minutes or for the time indicated. Nonspecific uptake was determined in the presence of 10 μM nomifensine. Uptake was terminated by aspirating uptake buffer and washing each well twice with 0.5 ml ice-cold uptake buffer. Cells were lysed in 0.1 N NaOH and transferred to vials containing 3 ml scintillation cocktail. Radioactivity was quantitated using a Beckman LS6500 counter. Data were analyzed in Graph Pad Prism 5.0.

Vesicular ³H-DA Uptake: Cells were plated in 10 cm dishes and grown to 100% confluency. Cells were washed with warm PBS without Ca²⁺ or Mg²⁺ and resuspended in uptake buffer (25 mM HEPES, 100 mM potassium tartrate, 100 μM EDTA, 50 μM EGTA, pH 7.4). Cells were homogenized with a glass/Teflon homogenizer 30 times on ice. The homogenate was centrifuged at 8000xg for 8 minutes at 4°C. Protein content of the resulting supernatant was determined by BCA assay. Uptake assays utilized 100 μg of protein in complete uptake buffer (uptake buffer with 1.7 mM ascorbate, 2 mM Mg²⁺-ATP salt, pH 7.4) and 20 μM tetrabenazine (TBZ) to define specific uptake. Samples were incubated in uptake buffer with or without the indicated concentration of nuciferine for 10 minutes at 30°C followed by addition of 1 μM dopamine with a 2% tracer of ³H-DA. Samples were incubated for 5 minutes at 30°C with gentle shaking. The assay was terminated by addition of 5 ml ice-cold assay buffer before filtration through 0.5% PEI-soaked Whatman GE/F filters (Brandel Inc., Gaithersburg, MD). Filters were then placed in vials containing 3 ml scintillation fluid and counted using a Beckman LS6500. Data were analyzed in Graph Pad Prism 5.0.

2.7 Animals

University of North Carolina at Chapel Hill: The head-twitch response, locomotor activity, and catalepsy studies were run with male C57BL/6J mice bred at UNC. University of Arkansas for
Medical Sciences: For DOI discrimination studies, adult male NIH Swiss mice weighing approximately 25 g were obtained from Harlan Laboratories (Indianapolis, IN, USA) and housed 3 mice per cage upon arrival. For PCP discrimination studies, adult male Sprague-Dawley rats weighing approximately 250 g were obtained from Harlan Laboratories (Indianapolis, IN, USA) and housed 3 rats per cage upon arrival. Virginia Commonwealth University: For clozapine substitution studies, adult male 129S2/SvHsd inbred mice were bred in-house. Duke University: adult male and female C57BL/6J mice, wild type (WT) mice, and dopamine transporter knockout (DAT-KO) mice were used in PPI experiments. The DAT mice were generated by heterozygous matings. DAT mice were group-housed in an environmentally-controlled room on a 14:10-h light/dark cycle (lights on 0800 h). The C57BL/6J, NIH Swiss, and 129S2/SvHsd mice and the Sprague-Dawley rats were housed in environmentally-controlled rooms on a 12:12-h light/dark cycle (lights on at 0700 hr). All experiments were conducted with approved protocols from the Institutional Animal Care and Use Committees of the university associated with each principal investigator: The University of North Carolina at Chapel Hill Institutional Animal Care and Use Committee, the Duke University Institutional Animal Care and Use Committee, the Virginia Commonwealth University Institutional Animal Care and Use Committee, and the University of Arkansas for Medical Sciences Animal Care and Use Committee. It should be noted that the range of species and breeds used in these studies was due to the “convenience and availability” nature of the collaboration, in which nuciferine was tested in the established experimental protocols of the collaborating laboratories dependent upon the availability of animals and openings in the experimental schedule.

2.8 Head-twitch responses

The head-twitch response procedure has been described elsewhere.[22–24] In brief, mice were injected with either nuciferine (1.0, 3.0, or 10.0 mg/kg, i.p.) or vehicle, n = 4 mice/condition. Fifteen minutes later, mice were injected with 1.0 mg/kg DOI (i.p.) and immediately placed in an observation chamber (new cage without bedding). Head-twitches (operationally defined as a rapid rotational jerk of the head that can be distinguished from species-appropriate grooming or scratching behaviors) were counted for 20 minutes in 5 minute bins. For the time-course study, mice were pretreated with 3.0 mg/kg nuciferine (i.p.) at 60, 45, 30, 15, or 0 minutes (co-injection) prior to the 1.0 mg/kg DOI (i.p.) injection, and head-twitches were counted as described above. In one experiment, mice (n = 4 per condition) were pretreated with an injection (s.c.) of 3.0 mg/kg nuciferine or vehicle 15 minutes prior to 1.0 mg/kg DOI injection (i.p.) and head-twitches were counted as described above. All experiments were performed by 3 observers, with 2 observers blinded to the experimental conditions which were evenly distributed. Power analyses were performed with the resulting data. The two highest doses of nuciferine tested (10.0 and 3.0 mg/kg), had 0.96 and 0.88 power to detect significance (α = 0.05). As these experiments were performed blinded and in distinct mice, further replication was not performed.

2.9 Open field activity

Locomotor activity was assessed in AccuScan activity monitors (41X41X30 cm; AccuScan Instruments, Columbus, OH) with photocells spaced at 1.52 cm as described. [22] In these monitors, the photocells create a grid of light beams, and breaks in the light beams (caused by the mouse) are recorded. The Accuscan software then calculates the total distance travelled (amongst other measurements) by analyzing the sequential order of beam breaks in the grid. Horizontal activity was measured as the total distance traveled in centimeters and was recorded in 5-minute bins. PCP-induced hyperlocomotor activity: mice (n = 16) previously acclimated
to activity chambers were placed into the chambers for 15 minutes. Mice were then injected (i.
p.) with either vehicle or 3 or 10 mg/kg nuciferine and returned to the chamber for 15 min.
Subsequently, mice were injected with 6.0 mg/kg PCP (i.p.) and returned to the chamber for 90
minutes. Induction of AMPH-induced hyperlocomotor activity followed an identical protocol
except locomotor activity was recorded for a total of 75 minutes and 3 mg/kg AMPH was used
(i.p.).

2.10 Catalepsy procedure
Mice (n = 3) were initially injected (i.p.) with vehicle (0.9% saline/0.2% lactic acid), 10.0 mg/kg
nuciferine, or 1.0 mg/kg haloperidol. Mice were placed upright on a 45° angled screen. The
time required for the animal to move all four paws was scored in seconds (maximum of 5 min)
and is reported as the latency to movement. An extended delay to move on the inclined screen
test is indicative of drug-induced catalepsy. Power analyses were performed with the resulting
data. The experiment had 100% power to detect a significant difference (α = 0.05) between
nuciferine and haloperidol at the 60 minute timepoint.

2.11 DOI drug discrimination
Adult male NIH Swiss mice (n = 6) were trained to respond under an FR5 reinforcement
schedule by presentation of evaporated milk in daily sessions using procedures similar to those
previously described. [25] Mice were trained in drug discrimination via injection of saline
(VEH) or 0.3 mg/kg R(-)-DOI presented in a pseudo-random order, with the constraint that
no animal could receive the same injection for more than 3 consecutive sessions (1 session /
day). Response assignments were counterbalanced across trials. Drugs were administered i.p.
and pre-treatment time was 10 minutes. During each training session the overall response rate,
overall distribution of responses on the drug-injection lever, and the distribution of responses
on this same lever prior to delivery of the first reinforcer were analyzed. When animals reliably
achieved a level of >85% correct responding prior to delivery of the first reinforcer over 3 con-
secutive sessions, a substitution test occurred the following day. During test sessions, a multiple
component cumulative dosing procedure was used, and no responses were reinforced. Each
component was terminated after the emission of five responses on either lever. Mice were then
removed from the chamber, administered the next cumulative dose, and returned to the cham-
ber. Ten minutes later, levers were re-extended into the experimental chamber. In this manner,
four doses of drug could be tested over ~40 min in a single session. The distribution of
responses between the two levers was expressed as a percentage of total responses emitted on
the drug-appropriate lever. Response rate was calculated for each session by dividing the total
number of responses emitted on both levers by the elapsed time prior to 5 responses on either
lever. Nuciferine was administered 15 minutes prior to the first injection of DOI.

2.12 PCP drug discrimination
Adult male Sprague-Dawley rats (n = 5) were trained to respond under an FR20 schedule and
were reinforced by presentation of food pellets in daily sessions using procedures similar to
those previously described. [26] Rats were trained in drug discrimination via a pre-session
injection of saline (VEH) or 3 mg/kg PCP chosen in a pseudo-random order (coin flip), with
the same constraints, criteria, and dosing procedures as described above except that during test
sessions, a given component of the cumulative dosing procedure was terminated after the emis-
ion of 20 responses on either lever. Drugs were administered i.p. and pretreatment time was
10 minutes. As described above for DOI, four doses of drug could be tested in a single ~40 min
test session. Nuciferine was administered 15 minutes prior to the first injection of PCP.
2.13 Clozapine drug discrimination study

Adult male B6129 hybrid mice (n = 12) were trained to respond under a FR10 schedule and were reinforced by presentation of sweetened milk as described previously. [27] The drug and vehicle lever positions were counterbalanced between groups to control for olfactory cues. [28] All injections were given subcutaneously with a pre-session injection time of 30 minutes. Training occurred on a double alternation injection schedule with two days of VEH followed by two days of CLZ and repeated (VEH, VEH, CLZ, CLZ, VEH, VEH etc.). In order for a mouse to pass a training day it had to meet three criteria: (1) complete the first fixed ratio (FR) on the condition-appropriate lever, (2) at least 80% of the total responses were made on the condition-appropriate lever, and (3) at least 10 responses per minute were made during the session. Drug testing was conducted approximately two times per week with at least two training days in between. To be eligible for testing, mice were required to pass both a drug and vehicle training day consecutively. During drug substitution tests, animals were injected subcutaneously with nuciferine (0.1, 0.3, 1.0, 3.0, 10.0 mg/kg) and placed in the operant chamber after 30 minutes. Responses on both levers were reinforced.

2.14 Prepulse inhibition (PPI)

PPI of the acoustic startle response was conducted as described elsewhere [29] using SR-LAB chambers (San Diego Instruments, San Diego, CA). To determine whether nuciferine could ameliorate or normalize PPI, two separate experiments were conducted. In the first, C57BL/6J mice were administered VEH, 5, or 10 mg/kg nuciferine (i.p.) and returned to their home-cages for 15 min. Subsequently mice were treated with either VEH or 6 mg/kg PCP (i.p.) and placed into the PPI apparatus for a 5 min habituation prior to the onset of testing. In the second study, WT and DAT-KO mice were given VEH or 2.5, 5, or 10 mg/kg nuciferine or 2 mg/kg clozapine (i.p.) and returned to their home-cages. Fifteen min later DAT mice were habituated to the PPI apparatus for 5 min and testing began. The startle trials consisted of a 40 msec burst of 120dB white-noise; prepulse trials consisted of 20 msec prepulse stimuli that were 4, 8, or 12 dB above the white-noise background (64dB) and were followed 100 msec later by the 120dB acoustic startle stimulus. Non-stimulus or null trials consisted of the 64dB white-noise background. PPI responses were calculated as a percentage score for each prepulse intensity, where \( \%\text{PPI} = \left[1 - \frac{\text{prepulse trials}}{\text{startle-only trials}}\right] \times 100 \).

2.15 Statistics

The data are presented as means and standard errors of the mean (SEM) and data from the locomotor activity, head-twitch, and PPI studies were analyzed by SPSS statistical software (IBM Corp., Armonk, NY). Locomotor data were assessed with repeated measures ANOVA (RMANOVA) for within subject effects of time and for between subjects effects of treatment; head-twitch data were similarly assessed with RMANOVA. ANOVA and independent measures t-test were used to also examine treatment differences in motor activity. For the PPI experiment, the responses to the null and startle stimuli were analyzed by two-way ANOVA, whereas the PPI data were subjected to RMANOVA where the within subjects effect was prepulse intensity and the between subjects effects were treatment and, in the case of the DAT mice, genotype. All post-hoc analyses were by Bonferroni corrected pair-wise comparisons. A \( p < 0.05 \) was considered significant. In the drug discrimination studies for each test session, mean (±SEM) percent responding on the drug-associated lever and the rate of responding (responses/sec) were calculated for each session component. Full substitution was operationally defined as >80% selection of the drug-associated lever, partial substitution was operationally defined as 40%-80% selection of the drug-associated lever, and no substitution was operationally defined as <40% selection of the drug-associated lever. Subjects failing to complete the
response requirement in a given component were excluded from the data analyses, but their data were included in response rate calculations. Discrimination and response rate data were not normally distributed, so the Kruskal-Wallis one-way ANOVA on ranks was used to analyze data across dose, comparing 3 treatment conditions: 1) training drug alone, 2) training drug + 1.0 mg/kg nuciferine, and 3) training drug + 3.0 mg/kg nuciferine. Significant ANOVAs were followed by pair-wise multiple procedures using the Dunn’s method (\( \alpha = 0.05 \)) to determine differences among means.

3. Results

3.1 Prediction, in vitro identification, and in vitro characterization of nuciferine

The \textit{in silico} assessment of phytochemicals in \textit{Nelumbo nucifera} (Fig 3) suggested that nuciferine and its metabolites are the most structurally similar to known compounds (Fig 3; color of circles). Additionally, nuciferine and its metabolites have high confidence protein-binding predictions (Fig 3; size of circles) and are predicted to cross the blood brain barrier (y axis). Finally, nuciferine is predicted to have a relatively large number of molecular targets (x axis).

Next we compared the polypharmacological profile of nuciferine with two atypical antipsychotic drugs (clozapine and aripiprazole) and the typical antipsychotic drug haloperidol (Fig 2). This convergence of predictions coupled with the potential utility of a compound with high polypharmacology suggested that nuciferine (Fig 1) would be the most pharmacologically interesting to investigate. Notable predictions from the SEA [13] are listed in Table 1, of which we were capable of testing 10 via the National Institute of Mental Health Psychoactive Drug Screening Program (PDSP).

The PDSP \textit{in vitro} affinity screening (Tables 1 and 2) revealed a total of 13 receptors with affinities less than 1 \( \mu \text{M} \), and 21 receptors with affinities less than 10 \( \mu \text{M} \). SEA successfully predicted 7 out of 13 G protein-coupled receptors that were determined by the PDSP to have \( K_i \) values of less than 1 \( \mu \text{M} \). Functional studies (Table 1) indicate that nuciferine shows appreciable potency as a D2 partial agonist (\( EC_{50} = 64 \text{nM} \)), as a 5-HT\(_7\) inverse agonist (\( EC_{50} = 150 \text{nM} \)), and as a 5-HT\(_{2C}\) antagonist (\( IC_{50} = 131 \text{nM} \)). [30] Nuciferine was a partial agonist at D2 receptors with an activity (\( E_{\text{max}} = 67\% \) of dopamine, Fig 4A) similar to aripiprazole (\( E_{\text{max}} = 50\% \) of dopamine).[19, 31] In line with its partial agonist activity, nuciferine inhibited dopamine-induced activation of G\(_i\) (Fig 4B) with a potency similar to clozapine (nuciferine \( K_B = 62 \text{nM} \); clozapine \( K_B = 20 \text{nM} \)) as determined via Schild regression analysis. [32] Also similar to clozapine, nuciferine exhibits 5-HT\(_{2A}\) antagonist activity (\( IC_{50} = 478 \text{nM} \)) and 5-HT\(_{2}\) inverse agonist activity (\( IC_{50} = 150 \text{nM} \)), [33] and may effectively antagonize 5-HT\(_{6}\) receptors as a low efficacy partial agonist (\( EC_{50} = 700 \text{nM} \), 17.3\% maximal effect). [2, 34] Finally, nuciferine exhibited micromolar potency as a 5-HT\(_{2B}\) antagonist (\( IC_{50} = 1 \text{\mu M} \)), a D\(_4\) agonist (\( EC_{50} = 2 \text{\mu M} \)), a D\(_{3}\) partial agonist (\( EC_{50} = 2.6 \text{\mu M}, 50\% \) maximal response), and a 5-HT\(_{1A}\) agonist (\( EC_{50} = 3.2 \text{\mu M} \)).

3.2 Assessment of aggregator properties

The predominantly low-potency antagonist activity of nuciferine suggested that nuciferine could be functioning as a colloidal aggregator, a well-known mechanism of promiscuous activity in early discovery, such as high-throughput screening. [35, 36] Recent studies have shown this mechanism can affect membrane-bound receptors such as GPCRs [37] in cell-based screening. At concentrations relevant to this study (\(< = 10 \text{\mu M} \)), dynamic light scattering determined that nuciferine did not scatter light above background in aqueous buffer, suggesting colloidal aggregates of nuciferine are not formed at this concentration. Consistent with this,
nuciferine did not inhibit AmpC β-lactamase at these concentrations; AmpC is a counter-screening enzyme widely used to test for colloidal aggregation [35, 36, 38] (data not shown). The lack of particle formation by light scattering and the lack of inhibition of the orthogonal counter-screening enzyme supports the idea that nuciferine is not acting promiscuously via colloidal aggregation in the assays described here.

3.3 Dopamine transport by DAT and VMAT2

SEA predicted that nuciferine may interact with the vesicular monoamine transporter-2 (VMAT2). To determine if nuciferine affects uptake at VMAT2, we conducted dopamine
uptake experiments in vesicles isolated from HEK cells overexpressing VMAT2 and found no effect of nuciferine on DA uptake (Fig 5A). We also tested whether nuciferine affected uptake in whole-cell uptake studies in HEK cells expressing DAT and VMAT2 or DAT alone as a control, as described elsewhere. [39] In whole cells expressing DAT and VMAT2, the total capacity of the cell to take up and store dopamine is measured. This has a DAT-mediated and a VMAT2-mediated component, as previously demonstrated.[39] Comparison of the effects of compounds in these two cell types, as well as in isolated vesicles, aids in separating out these two components of uptake in whole cells. Nuciferine pre-treatment had no effect in HEK-DAT/VMAT2 cells (Fig 5B). Counter to predictions, in the HEK-DAT control cells, nuciferine pre-treatment increased uptake in HEK-DAT cells by 60% over the vehicle control (Fig 5C; EC50 = 1.8 nM).

3.4 DOI-induced head-twitch response (HTR)
Mice pretreated with nuciferine (1.0, 3.0 and 10.0 mg/kg, i.p. or 3 mg/kg, s.c.; 15 min prior to DOI) showed a dose-dependent inhibition of the head-twitch response produced by 1.0 mg/kg DOI during the course of testing (Fig 6A). Bonferroni corrected pair-wise comparisons indicated that regardless of route of injection, nuciferine attenuated DOI-induced head-twitches at 10, 15 and 20 min (ps<0.05). When the different doses of nuciferine were examined, a RMANOVA revealed a significant effect of time [F[3,36] = 232.725, p<0.0001], and a significant time by treatment interaction [F[9,36] = 16.170, p<0.0001]. Bonferroni comparison results indicated that 3 and 10 mg/kg nuciferine decreased head-twitches at 5, 10, 15 and 20 min (ps<0.05) relative to DOI-treated mice. Differences in responses to the 3 and 10 mg/kg doses were not statistically significant (Fig 6A). By comparison, 1 mg/kg nuciferine exerted no statistical effect on the DOI induced head-twitches.

The time-course of 3 mg/kg nuciferine pre-treatment (i.p.) indicated that suppression of the DOI-induced head-twitches was most pronounced when the interval between the nuciferine

### Table 1. In silico and in vitro characterization of nuciferine.

<table>
<thead>
<tr>
<th>SEA Predictions (in order of confidence)</th>
<th>Radioligand used</th>
<th>In Vitro Affinity (nM)</th>
<th>Functional EC50 (nM)</th>
<th>Function Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>D1 (5.4e-32)</td>
<td></td>
<td>752 [3H]SCH23390</td>
<td></td>
<td></td>
</tr>
<tr>
<td>D2 (1.6e-25)</td>
<td></td>
<td>515 [3H]N-methyl Spiperone</td>
<td>65.07</td>
<td>Partial agonist</td>
</tr>
<tr>
<td>D3 (1.3e-12)</td>
<td></td>
<td>741 [3H]N-methyl Spiperone</td>
<td></td>
<td></td>
</tr>
<tr>
<td>D4 (5.0e-13)</td>
<td>(neg) [3H]SCH23390</td>
<td>2600</td>
<td></td>
<td>Partial agonist, ~50%</td>
</tr>
<tr>
<td>VMAT2 (2.1e-13)</td>
<td></td>
<td>NA</td>
<td></td>
<td></td>
</tr>
<tr>
<td>SK Channel (3.8e-11)</td>
<td></td>
<td>NA</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5-HT1A (1.2e-11)</td>
<td></td>
<td>77 [3H]WAY100635</td>
<td>3230</td>
<td>Agonist</td>
</tr>
<tr>
<td>5-HT2B (1.6e-7)</td>
<td></td>
<td>49.8 [3H]LSD</td>
<td>150</td>
<td>Inverse agonist</td>
</tr>
<tr>
<td>5-HT7 (4.1e-4)</td>
<td></td>
<td>312 [3H]Ketanserin</td>
<td>478</td>
<td>Antagonist</td>
</tr>
<tr>
<td>5-HT3A (4.1e-6)</td>
<td></td>
<td>41 [3H]LSD</td>
<td>1000</td>
<td>Antagonist</td>
</tr>
<tr>
<td>Unpredicted Hits</td>
<td></td>
<td>60.5 [3H]Mesulergine</td>
<td>131</td>
<td>Antagonist</td>
</tr>
<tr>
<td>5-HT6</td>
<td></td>
<td>268 [3H]LSD</td>
<td>700</td>
<td>Partial agonist, 17.3%</td>
</tr>
<tr>
<td>D4</td>
<td></td>
<td>1387 [3H]N-methyl Spiperone</td>
<td>2000</td>
<td>agonist</td>
</tr>
</tbody>
</table>

Receptor targets are listed with their respective SEA prediction value (if available), followed by their competition binding affinity value (if available), followed by their functional EC50 value (if available) and the corresponding function type.

Information about cell lines for all assays can be found at [http://pdsp.med.unc.edu/pdspw/binding.php](http://pdsp.med.unc.edu/pdspw/binding.php).

doi:10.1371/journal.pone.0150602.t001
and DOI injections was 15 min (Fig 6B). When the interval between nuciferine and DOI injections was varied, a RMANOVA found the main effect of time \([F[9,54] = 334.913, p < 0.001]\) and the time by pre-treatment interval \([F[15,54] = 5.838, p < 0.001]\) to be significant. Bonferroni analyses indicated that mice treated 45 or 60 min prior to DOI injection failed to show significant alterations in head-twitches compared to DOI-treated animals (Fig 6B). By contrast, mice treated 15 or 30 min prior to the DOI injection all had statistically significant \((p < 0.05)\) reductions in the numbers of head-twitches compared to mice given DOI at the 5, 10, 15, and 20-min sampling times. Moreover, nuciferine pretreatment at 15 min produced the most profound overall reductions in head-twitches relative to animals pretreated at 45 and 60 min and measured at 15 and 20 min sampling times, all of which were statistically significant \((p < 0.05)\). Collectively, these data demonstrate that nuciferine can antagonize DOI-induced head twitches and that these effects are time-dependent with the 15 min interval between nuciferine and DOI injection being the most effective.

### 3.5 Drug discrimination

Dose-dependent generalization for the DOI training dose was observed when cumulative doses were administered alone, with cumulative doses of 0.1 and 0.3 mg/kg producing full substitution (Fig 7A). In the presence of 1.0 mg/kg nuciferine, however, cumulative DOI doses only produced partial substitution, whereas in the presence of 3.0 mg/kg nuciferine, no cumulative dose of DOI substituted for the training dose, up to a dose that profoundly suppressed responding (Fig 7D). Although the overall ANOVA was significant \((p < 0.001)\), no within-dose

<table>
<thead>
<tr>
<th>PDSP Hits Without Functional Assay</th>
<th>In vitro Affinity (nM) Radioligand used</th>
</tr>
</thead>
<tbody>
<tr>
<td>5-HT1D</td>
<td>518 ([^{3}H]5-CT)</td>
</tr>
<tr>
<td>α1A</td>
<td>1386 ([^{3}H]Prazosin)</td>
</tr>
<tr>
<td>α1B</td>
<td>1995.3 ([^{3}H]Prazosin)</td>
</tr>
<tr>
<td>α1D</td>
<td>818 ([^{3}H]Prazosin)</td>
</tr>
<tr>
<td>α2A</td>
<td>1153.5 ([^{3}H]Rauwolscine)</td>
</tr>
<tr>
<td>α2B</td>
<td>686.8 ([^{3}H]Rauwolscine)</td>
</tr>
<tr>
<td>α2C</td>
<td>692.8 ([^{3}H]Rauwolscine)</td>
</tr>
<tr>
<td>β1-AR</td>
<td>7149 ([^{3}H]CGP12177)</td>
</tr>
<tr>
<td>β2-AR</td>
<td>1103 ([^{3}H]CGP12177)</td>
</tr>
<tr>
<td>DOR</td>
<td>10000 ([^{3}H]DADLE)</td>
</tr>
<tr>
<td>H2</td>
<td>1662 ([^{3}H]Cimetidine)</td>
</tr>
<tr>
<td>MOR</td>
<td>9549.00 ([^{3}H]DAMGO)</td>
</tr>
<tr>
<td>V1A</td>
<td>10000 ([^{3}H]Vasopressin)</td>
</tr>
<tr>
<td>5-HT5A</td>
<td>1113 ([^{3}H]LSD)</td>
</tr>
</tbody>
</table>

Receptor targets not predicted by SEA but identified by in vitro competition binding assays for which functional assays are not available. The following receptors assayed in the PDSP screen had no observed binding of nuciferine. Receptors are listed with their associated radioligand in (). Sets of receptors using the same radioligand are indicated in (): [5-HT1B, 5-HT1E] \([^{3}H]5-CT\), 5-HT3 \([^{3}H]GR65630\), β2-AR \([^{3}H]CGP12177\), BZP Rat Brain Site \([^{3}H]Flunitrazepam\), D5 \([^{3}H]SCH23390\), DAT \([^{3}H]WIN35428\), GabaA \([^{3}H]Muscimol\), H1 \([^{3}H]Pyrilamine\), H3 \([^{3}H]o-methylhistamine\), H4 \([^{3}H]Histamine\), KOR \([^{3}H]U69593\), NMDA PCP site \([^{3}H]MK801\), [M1, M2, M3, M4, M5] \([^{3}H]QNB or [^{3}H]NMS\), NET \([^{3}H]Nisoxetine\), Oxytocin \([^{3}H]Oxytocin\), SERT \([^{3}H]Citalopram\), Sigma 1 \([^{3}H]Pentazocine\), Sigma 2 \([^{3}H]DTG\), [V1B, V2] \([^{3}H]Vasopressin\).
pairwise comparisons reached statistical significance. Dose-dependent generalization for the PCP training dose was observed when cumulative doses were administered alone, with a cumulative dose of 3.0 mg/kg producing full substitution. In the presence of 1.0 or 3.0 mg/kg nuciferine, cumulative PCP doses produced similar substitution to PCP alone (Fig 7B). Although the overall ANOVA was significant (p<0.05), no within-dose pairwise comparisons reached statistical significance. In the clozapine-trained animals, a dose-dependent substitution for 1.25 mg/kg clozapine was seen at 10.0 mg/kg nuciferine (80.63% drug lever responding), with an ED₅₀ value of 5.42 mg/kg (95% CI 3.09–9.48 mg/kg) while the lower doses tested (0.1 mg/kg–3.0 mg/kg) failed to produce substitution for clozapine’s discriminative cue (Fig 7C). In addition to a

Fig 4. Nuciferine functional activity at the dopamine D₂ receptor as measured via D₂-mediated Gᵢo signaling in HEKT cells. (a) Concentration-response curve of nuciferine (red) compared to dopamine (black) and aripiprazole (blue). (b) Concentration-response curve of dopamine (DA) in the presence of 1 μM nuciferine (red) or clozapine (green) showing rightward-shift of DA indicative of competitive antagonist activity. The data represent concentration-response curves of normalized data with respect to dopamine performed in triplicate (mean +/- s.e.m.; n = 3).

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high percentage of responding on the clozapine-appropriate lever, 10.0 mg/kg nuciferine also produced significant rate suppression as compared to vehicle control points (p < 0.001) (Fig 7F).

3.6 Locomotor activity

Stimulation of locomotor activity by 6.0 mg/kg PCP was significantly blocked by 10.0 mg/kg, but not by 3.0 mg/kg, nuciferine (Fig 8A, left panel). RM ANOVA revealed a significant effect of time [F[17,765] = 30.553, p<0.001], and a significant time by treatment interaction [F[34,765] = 1.997, p<0.010]. Bonferroni corrections indicated that during the 0–30 min interval locomotion in all groups was similar. As expected, PCP stimulated locomotor activity from 35–90 min compared to baseline at 0–30 min (ps<0.001). Treatment with 3 mg/kg nuciferine prior to PCP failed to significantly alter the PCP-induced hyperlocomotion, whereas the 10 mg/kg nuciferine significantly depressed this locomotion compared to PCP-treated mice at 45–60 min (p<0.05). Nevertheless, the reductions in PCP-stimulated activity by 3 and 10 mg/kg nuciferine were not statistically different. When the results were presented as cumulative distance traveled between 45 and 60 min of testing, dose-dependent decreases in PCP-stimulated activity could be seen (Fig 8A, right panel). ANOVA found a significant effect of treatment [F[2,47] = 4.323, p<0.05] and Bonferroni tests confirmed that 10 mg/kg nuciferine significantly reduced locomotion compared to PCP (p<0.05). The 3 mg/kg dose decreased activity, but it was not significantly different from the PCP alone or the 10 mg/kg plus PCP group.

A second set of mice was used to assess the effects of 3 mg/kg nuciferine pre-treatment on amphetamine induced hyperlocomotion (Fig 8B, left panel). A RM ANOVA revealed a significant effect of time [F[13,390] = 177.243, p<0.001] and a significant time by treatment interaction [F[13,390] = 14.625, p<0.001]. Bonferroni corrected comparisons found that activities between the two group at 0–30 min were not statistically different. All animals showed a significant increase in activity following amphetamine treatment compared to their baseline activities (p<0.001). Those mice given nuciferine prior to amphetamine treatment showed significantly higher motor activity between 35–50 min compared to those given amphetamine alone (p<0.05); however, responses after this time were not significantly differentiated. When the distance travelled in the open field for the 40 min following amphetamine treatment was aggregated, mice given nuciferine followed by amphetamine had heightened activity compared to those animals given amphetamine alone (Fig 8B, right panel) [t[1,30] = 4.014, p<0.001].
Together, these findings indicate that while nuciferine is capable of attenuating the hyperlocomotion induced by PCP, amphetamine induced hyperlocomotion is exacerbated by the compound.

### 3.7 Pre-pulse inhibition

For the experiment with C57BL/6J mice a two-way ANOVA was applied with two levels of treatment: PCP (vehicle or VEH and PCP) and nuciferine (VEH and the two doses of nuciferine) constituted the 6 groups. A two-way ANOVA for null activity identified a main effect of PCP treatment \( [F_{(1,45)} = 31.70, p<0.001] \), but the nuciferine treatments and the PCP by nuciferine interaction was not significant. Although Bonferroni corrected pair-wise comparisons...
noted that null activity was higher in the PCP-treated groups than in the respective vehicle/nuciferine groups \((p<0.001)\), this activity was still less than 7% of the startle responses in the PCP-treated group (Fig 9A). A two-way ANOVA for startle responses also observed a significant main effect of nuciferine treatment \([F(2,45)=13.22, p<0.001]\); the PCP treatment effect and the PCP by nuciferine interaction was not significant. Bonferroni post-hoc tests reported that startle responses for the 5 and 10 mg/kg nuciferine-treated groups were lower than those of the vehicle-treated groups \((p<0.001; \text{Fig } 9B)\). RMANOVA for PPI revealed the within subjects main effects for prepulse intensity was significant \([F(2,90)=93.16, p<0.001]\); the prepulse-intensity by PCP treatment, prepulse-intensity by nuciferine treatment, and the prepulse-intensity by PCP by nuciferine treatment interactions were not significant (Fig 9C). Bonferroni tests demonstrated that the response to the 4dB prepulse was lower than that for the 8 and 12dB prepulse responses \((p<0.001)\) and that the 8dB response was lower than that for the 12dB response \((p<0.001)\). Regardless of prepulse intensity, the between subjects main effects

**Fig 7. Drug Discrimination studies of nuciferine.** (a,d) Nuciferine blocks the discriminative stimulus of DOI at doses that do not affect the rate of responding. (b,e) Nuciferine does not block the discriminative stimulus of PCP at any dose tested. (e,f) Nuciferine substitutes for clozapine (solid circles) but only at a dose that suppresses response rate (empty circles). Data are presented as % training-drug appropriate responding and response rate per second. doi:10.1371/journal.pone.0150602.g007
for the nuciferine $F_{2,45} = 3.43, p<0.05$ and PCP treatments $F_{1,45} = 33.85, p<0.001$, as well as the nuciferine by PCP treatment interaction, were significant $F_{2,45} = 3.31, p<0.05$.

Decomposition of the interaction observed that PPI in the vehicle-vehicle, 5 mg/kg nuciferine-vehicle, and 10 mg/kg nuciferine-vehicle groups were not differentiated from each other, but that PPI in the PCP-vehicle group was suppressed relative to these controls ($p<0.001$). Despite this fact, mice given nuciferine prior to PCP had higher PPI than the PCP-vehicle groups ($p<0.05$) and the 10 mg/kg nuciferine-PCP group was not statistically different from the respective control. Thus 10 mg/kg nuciferine rescued the PCP-disrupted PPI.

Analyses of responses from the DAT KO mice presented a different picture. A two-way ANOVA for null activity observed significant genotype $F_{1,102} = 12.410, p<0.005$ and treatment effects $F_{4,102} = 18.516, p<0.001$ and a significant genotype by treatment interaction $F_{4,102} = 3.956, p<0.01$. Bonferroni corrections demonstrated that null activity in DAT-KO mice was higher than that in WT mice with clozapine ($p<0.005$) and 2.5 mg/kg nuciferine.
Null activity in WT mice to clozapine and 5 mg/kg nuciferine were significantly enhanced relative to the vehicle control ($p<0.005$), whereas null activity in DAT-KO mice to clozapine and all doses of nuciferine were significantly augmented compared to vehicle group ($p<0.001$). A two-way ANOVA for startle responses revealed significant main effects of genotype [$F(1,102) = 9.227$, $p<0.005$] and treatment [$F(4,102) = 28.673$, $p<0.001$] (Fig 9E).

Regardless of genotype, clozapine-treated mice had higher startle responses than all other groups ($p<0.001$). For PPI a RMANOVA found significant within-subject effects of prepulse intensity [$F(2,204) = 60.090$, $p<0.001$] and a significant prepulse intensity by genotype interaction [$F(2,204) = 10.479$, $p<0.001$] (Fig 9E). Bonferroni tests demonstrated prepulse dependency in WT animals where inhibition at each prepulse intensity was significantly different from each other

Fig 9. PPI responses to nuciferine in mouse models of hypoglutamatergia and hyperdopaminergia. Nuciferine rescued PPI in the former, but not in the latter model. (a-c) Null activity (a), startle activity (b), and PPI (c) for C57BL/6J mice treated with vehicle (Veh), 5 or 10 mg/kg nuciferine (Nuc), and/or phencyclidine (PCP). (d-g) Null activity (d), startle activity (e), and PPI for wild-type (WT) (f) and dopamine transporter knockout (DAT-KO) mice (g) given Veh, 2 mg/kg clozapine (CLZ) or 2.5–10 mg/kg Nuc. N = 8–17 mice/group in the C57BL/6J experiment; *$p<0.05$, compared to Veh/Nuc groups; +$p<0.05$, compared to the Veh and PCP groups; †$p<0.05$, WT versus KO within dose; #$p<0.05$, dose effect within genotype; &$p<0.05$, overall drug effect regardless of genotype.

doi:10.1371/journal.pone.0150602.g009
By contrast, in DAT-KO animals responses to the 12 dB prepulse were higher than those to the 4 and 8 dB prepulses \((p < 0.005)\) and a significant genotype by treatment interaction \(F(4,102) = 3.288, p < 0.05\). Bonferroni comparisons found that PPI was significantly lower for DAT-KO than WT mice in the groups given vehicle or any dose of nuciferine \((p < 0.05)\). Clozapine significantly enhanced PPI in DAT-KO mice relative to the vehicle and nuciferine treatments \((p < 0.05)\). Together, these data show that PPI is intact in WT animals and it is not affected by either clozapine or nuciferine. By contrast, PPI is deficient in DAT-KO mice and it was restored by clozapine but not by nuciferine.

3.8 Catalepsy

Nuciferine (10.0 mg/kg) did not produce catalepsy at any of the time-points examined (Table 3).

<table>
<thead>
<tr>
<th>Post-injection timepoint (min)</th>
<th>Vehicle</th>
<th>Nuciferine10.0 mg/kg</th>
<th>Haloperidol1.0 mg/kg</th>
</tr>
</thead>
<tbody>
<tr>
<td>30</td>
<td>3 ± 2.08 s</td>
<td>2.3 ± 2.3 s</td>
<td>270 ± 30 s</td>
</tr>
<tr>
<td>60</td>
<td>2.67 ± 1.45 s</td>
<td>4.7 ± 2.6 s</td>
<td>300 ± 0 s</td>
</tr>
</tbody>
</table>

Nuciferine did not cause a latency to move in the inclined grid test compared to haloperidol at the doses and times tested.

doi:10.1371/journal.pone.0150602.t003

(p < 0.001) (Fig 9F). By contrast, in DAT-KO animals responses to the 12 dB prepulse were higher than those to the 4 and 8 dB prepulses \((p < 0.005)\), which were not significantly distinguished from each other (Fig 9G). The between subjects test discerned significant genotype \(F(1,102) = 25.246, p < 0.001\) and treatment effects \(F(4,102) = 4.238, p < 0.005\) and a significant genotype by treatment interaction \(F(4,102) = 3.288, p < 0.05\). Bonferroni comparisons found that PPI was significantly lower for DAT-KO than WT mice in the groups given vehicle or any dose of nuciferine \((p < 0.05)\). Clozapine significantly enhanced PPI in DAT-KO mice relative to the vehicle and nuciferine treatments \((p < 0.05)\). Together, these data show that PPI is intact in WT animals and it is not affected by either clozapine or nuciferine. By contrast, PPI is deficient in DAT-KO mice and it was restored by clozapine but not by nuciferine.

4. Discussion

Here we present an *in vitro* and *in vivo* characterization of nuciferine, an alkaloid found in *Nelumbo nucifera* and *Nymphaea caerulea* lotus plants. Our primary finding was that nuciferine has a pharmacological profile similar but not identical to some antipsychotic drugs (especially aripiprazole) and that nuciferine performed as an antipsychotic-like drug in some animal models predictive of antipsychotic drug-like actions. Here, we discuss these findings in the context of antipsychotic pharmacology and animal models of antipsychotic efficacy.

4.1 Molecular characterization

The *in silico* prediction and *in vitro* characterization of nuciferine revealed a molecular affinity profile containing receptors known to be modulated by established antipsychotic drugs. Within the set of 13 receptors with \(< 1 \mu M\) affinity, nuciferine showed greatest affinity at the serotonergic receptors and overall it exhibited a molecular profile with multiple entities implicated in clinical antipsychotic efficacy: 5-HT\(_7\) [29, 34, 40, 41], 5-HT\(_6\) [34], 5-HT\(_2A\) [5], 5-HT\(_1A\) [42], 5-HT\(_{1D}\) [43], D\(_2\) [44], D\(_{1A}\), D\(_{3}\), D\(_{4}\), D\(_{5}\) [45], \(\alpha_{2B}\), and \(\alpha_{2C}\) [46]. The preponderance of low affinity antagonism led us to test whether nuciferine forms colloidal aggregates. Parenthetically, colloidal aggregation is a phenomenon that affects molecular pharmacological screening efforts and has not been appreciated until relatively recently.[47] Compounds that form colloidal aggregates can present unique and irrelevant behavior *in vitro* [48], and we avoided these confounds by demonstrating that nuciferine does not form aggregates at concentrations up to 10 \(\mu M\). Of the non-GPCR targets that SEA predicted, SK channels have been shown to bind antipsychotic compounds, [49] while VMAT2 is involved in monoaminergic neurotransmission and has long been proposed as a target for antipsychotic drug activity.[50, 51] It is also interesting to note that nuciferine does not bind to any muscarinic receptors. Muscarinic antagonists are prescribed to prevent or treat extrapyramidal side effects of antipsychotics. [52–54] During the
course of our investigation, an independent group determined that nuciferine functions as a 5-HT2A antagonist [55], corroborating our findings at this receptor.

With respect to antipsychotic efficacy, the D2 receptor is a desired druggable target with all approved antipsychotic drugs having potent interactions with this target.[30] In addition to D2 receptor affinity, however, it is also known that partial agonists have antipsychotic efficacy, such as aripiprazole.[19, 31] Therefore, we examined the D2 functional properties of nuciferine in comparison to aripiprazole and found that nuciferine exhibited a similar degree of partial agonist activity compared to aripiprazole, suggesting that nuciferine could exhibit antipsychotic drug-like properties, albeit with lower potency. The ability to block or antagonize DA-stimulated D2 activity may be a better predictor of antipsychotic efficacy, especially in the case of D2 partial agonists such as aripiprazole with low intrinsic efficacy in vitro.[56] which may act as D2 antagonists in vivo. [57] To examine nuciferine’s antagonist activity, we chose to measure nuciferine’s ability to block DA-stimulated D2 activation using a Schild regression analysis.[32] Results indicated that nuciferine can antagonize DA-stimulated cAMP inhibition with a potency similar to clozapine. Clozapine, one of the most effective atypical antipsychotics, possesses lower affinity for the D2 receptor compared to typical antipsychotics, [5] and therefore it is conceivable that just a moderate degree of D2 antagonism is needed for antipsychotic efficacy, given that clozapine also possesses potent 5-HT2A antagonism, 5-HT1A partial agonist activity, and 5-HT7 inverse agonism, which nuciferine also possesses. In summary, nuciferine shares two properties of antipsychotic efficacy at the D2 receptor, namely partial agonist activity at the D2 receptor similar to aripiprazole, and antagonist activity with moderate affinity comparable to clozapine.

There were notable discrepancies within our study across paradigms and compared to previously published findings. For instance, the SEA predicted one entity (dopamine D5 receptor) that was not detected in the PDSP binding affinity assay but was detected in the functional assay. The micromolar efficacy of nuciferine at the D5 receptor as measured via functional assays is consistent with the low binding affinity, due to the fact that the antagonist radioligand [3H]SCH 23390 labels inactive states of the receptor. Additionally, the dopamine transporter (DAT) was neither predicted using the SEA nor was it detected in the PDSP screen, yet it was determined that nuciferine modulates the DAT in vitro using a functional assay, again suggesting that this compound may be modulating the DAT at a site other than that occupied by the radioligand [3H]WIN35428. Interestingly, our behavioral results with amphetamine in the open field also support the notion that nuciferine interacts with DAT. It was previously reported that nuciferine inhibits the hyperlocomotor effect of amphetamine [4], but in our study we observed an increase in hyperlocomotor activity following nuciferine pretreatment. Our methods differed substantially from those of Bhattacharya et al. [4] who used a considerably higher dose range (25–100 mg/kg) of nuciferine, compared to our lower dose range (1–10 mg/kg across the study with 3.0 mg/kg nuciferine in the amphetamine experiment). These discrepancies support the use of orthologous assays in screening efforts to fully elucidate a chemical’s pharmacology.

The discovery of DAT modulation occurred serendipitously during the verification of the SEA VMAT2 prediction due to the nature of the experimental system. In these experiments, nuciferine caused an increase in uptake in cells expressing DAT but showed no effect in cells expressing both DAT and VMAT2. Direct measurement of uptake in isolated vesicles failed to detect direct modulation of VMAT2 by nuciferine. The difference between the results of the DAT and DAT/VMAT2 cells suggests that nuciferine may be indirectly inhibiting vesicular uptake in the HEK-DAT/VMAT2 cells, counteracting the increased DAT-mediated uptake. This is likely an indirect modulation of VMAT2 because nuciferine did not directly affect VMAT2 uptake in the isolated vesicle assay. However, further experiments are necessary to determine the mechanism.
4.2 Behavioral characterization

Since the initial studies of Macko and colleagues, our ability to assess antipsychotic efficacy in animal models has improved. For example, deficits in sensorimotor gating observed in individuals diagnosed with schizophrenia are modeled in rodents using the pre-pulse inhibition (PPI) paradigm, in which the startle response to a stimulus is inhibited by a pulse of reduced magnitude preceding the acoustic startle stimulus. Deficits in PPI are observed in schizophrenia patients and can be rescued by antipsychotic medications.[58] This deficit in PPI can be reproduced in animal models via administration of phencyclidine (PCP), a compound that produces psychosis-like effects in humans, and the disrupted PPI can be rescued by antipsychotic compounds.[59, 60] Notably, the rescue of PCP-induced disruption of PPI has been a hallmark of preclinical antipsychotic drug discovery efforts as a predictive model of antipsychotic efficacy.[61]

We thus set out to determine whether the identified molecular profile would translate into antipsychotic drug-like actions as determined by several animal models commonly used to study antipsychotic pharmacology. As mentioned previously, atypical antipsychotics exhibit 5-HT2A antagonist activity, and the potency with which antagonists attenuate the head-twitch response is highly correlated with the antagonist's affinity for 5-HT2A receptors. [62, 63] The ability of nuciferine to block the hallucinogen DOI-induced head-twitch response is consistent with the in vitro measurements of 5-HT2A antagonism and suggests activity in vivo via 5-HT2A receptor blockade. Furthermore, the dissociative psychedelic PCP produces a psychomimetic state in humans and the inhibition of PCP-induced behavioral effects is used as an animal model for evaluating schizophrenia-like behaviors.[64, 65] Nuciferine blocked not only the PCP-induced hyperlocomotor activity in the open field, but it also rescued PCP-disrupted PPI without direct antagonism of the NMDA PCP binding site. Despite clozapine rescuing PPI in DAT-KO mice, nuciferine failed to normalize PPI in the DAT-KO genetic mouse model of hyperdopaminergia or in the pharmacological model using amphetamine (data not shown). At this time, the molecular basis of the distinctions between the differential responses in the hypoglutamateric and hyperdopaminergic models is obscure. To our knowledge, this is the first report of nuciferine’s efficacy in these animal models and our results indicate that this compound has greater efficacy in hypoglutamateric than in hyperdopaminergic animal models.

The drug discrimination paradigm utilizes the interoceptive properties of drugs as a means to study their pharmacology. We used this paradigm to assess nuciferine’s pharmacology due to its unique ability to measure a systems-level pharmacological effect (as would be caused by a compound with high polypharmacology) at a whole organism level of analysis and due to its use in drug discovery efforts in the past.[66, 67] Nuciferine’s antagonism of the DOI-induced discriminative stimulus is consistent with its capacity to also antagonize DOI-induced head twitches and this result further validates the 5-HT2A antagonism observed in vitro. Interestingly, nuciferine did not block the PCP-induced discriminative stimulus. It should be emphasized that other investigators have reported that antipsychotics do not antagonize the discriminative stimulus of PCP. [68] Finally, nuciferine fully substituted for clozapine at the highest dose tested (10.0 mg/kg), indicating that nuciferine produced an interoceptive state similar to that of clozapine. Previous studies in C57BL/6 mice have shown that 5-HT2A serotonergic and α1-adrenoceptor antagonism mediate clozapine’s discriminative stimulus.[69] It should be noted that the 10.0 mg/kg dose of nuciferine that substituted for clozapine also produced significant rate suppression compared to vehicle control rates; however, previous studies have shown that the doses of antipsychotics that produce full substitution for clozapine are often accompanied by significant rate suppression (in both rats [70, 71] and mice[72, 73]).
4.3 Conclusion

In conclusion, we have comprehensively elucidated a complex pharmacological profile for nuciferine, one of the main alkaloids present in *Nelumbo nucifera*. The molecular profile of nuciferine was similar but not identical to the profiles of several approved antipsychotic drugs suggesting that nuciferine has atypical antipsychotic-like actions.

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