

# Anxiolytic-like effect of natural product 2-hydroxy-3,4,6-trimethoxyacetophenone isolated from *Croton anisodontus* in adult zebrafish via serotonergic neuromodulation involvement of the 5-HT system

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## Research Article

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# Abstract

Benzodiazepines are highly effective in combating anxiety; however, they have considerable adverse effects, so it is important to discover new safe anxiolytic agents. This study was designed to investigate the anxiolytic and anticonvulsant effect of natural product 2-hydroxy-3,4,6-trimethoxyacetophenone (HTMCX) and its possible mechanisms of action in adult zebrafish. The open field and light / dark tests (n = 6 animals/group) were used to assess anxiety and pentylenetetrazole (PTZ) as a seizure inducer. The 96-hour acute toxicity of HTMCX was also investigated. HTMCX (1, 3, and 10 mg / kg; *v.o.*) was not toxic and affected locomotor activity. The highest doses (3 and 10 mg / kg; *v.o.*) produced signs of anxiolytic action in the light / dark test, and this effect was abolished by the pizotifen (antagonist 5HTR1 and 5HTR2A / 2C), having the potential to form a complex in the same region of the site indicating that the anxiolytic effect via the serotonergic mechanism. However, the anxiolytic effect of HTMCX has not been abolished by flumazenil (antagonist GABA<sub>A</sub>), cyproheptadine (antagonist 5HTR2A), and granisetron (antagonist 5HTR3A / 3B). Therefore, HTMCX demonstrated an anxiolytic effect, suggesting that the 5HTR1 and 5HTR / 2C receptors may be involved in the pharmacological performance of this acetophenone in the central nervous system.

## 1. Introduction

Neurological diseases affect millions of people worldwide (Goni et al. 2021). According to the World Health Organization (WHO), approximately 264 million people suffer from some type of anxiety disorder, and Brazil contains one of the highest rates of this pathology (Depression and other common mental disorders 2017). In the United States and Europe, anxiety has considerable social and economic costs (Ham et al. 2020). In this circumstance, it is necessary to develop research for the discovery of new drugs that have anxiolytic activity without toxicity and withdrawal effect (Sedláčková et al. 2011).

Although benzodiazepines have high effectiveness in combating anxiety, they have considerable adverse effects, which is why selective serotonin reuptake inhibitors (SSRIs) are currently the first-line drugs of choice for the treatment of anxiety disorders due to a combination of efficacy and safety. However, its effects may take time to occur. The pathological state of anxiety is associated with changes in different neurotransmission pathways, such as serotonergic, noradrenergic, GABAergic, dopaminergic, and nitrenergic pathways (Martin et al. 2009; Spolidorio et al. 2007). Animal models are generally used to investigate the anxiolytic action of new compounds that act on these neuromodulation pathways.

The zebrafish (*Danio rerio*) has received attention as an animal model for pharmacological studies of anxiety because it has conserved neurotransmitters, and in addition, its genome has more than 80% of ortholog genes related to human diseases (Khan et al. 2017). Serotonergic neurotransmission in zebrafish has been shown to be similar to a mammal in terms of its physiology and pharmacology (Connors et al. 2014; Maximino et al. 2010; Panula et al. 2010), and studies have shown that the activation of serotonergic receptors through the use of selective agonists alters anxiety-like behaviors in zebrafish (Nowicki et al. 2014).

Modern pharmacological research indicates that natural products derived from plants have an anxiolytic effect through neuromodulation GABAergic or serotonergic (Lei et al. 2015). Based on the reported data, this study aimed to evaluate the anxiolytic and anticonvulsant effect of 2-hydroxy-3,4,6-trimethoxyacetophenone isolated from *C. anisodontus* and the possible mechanisms of action in adult zebrafish.

## 2. Material And Methods

### 2.1 Drugs and reagents

Granisetron hydrochloride (Corepharma / Inglaterra-Mx), pizotifene maleate (Central Manipulation Pharmacy / Brasil-SP), fluoxetine (Eli Lilly / EUA-IN), Cyproheptadine (Evidencia Soluções Farmacêuticas / Brasil-CE), Diazepam and Pentilenotetrazol (Sigma-Aldrich / USA-MO).

### 2.2. Plant material, extraction, isolation, and NMR analysis

In this work, the 2-hydroxy-3,4,6-trimethoxyacetophenone natural product was extracted, isolated, purified, and characterized of the *C. anisodontus* (Fig. 1) (Santiago et al. 2018).

### 2.3 Animals

The fish (from 90 to 120 days;  $0.4 \pm 0.1$  g,  $3.5 \pm 0.5$  cm), wild, male and female, were purchased at a local store (Fortaleza, CE) and kept for a week before the experiments in a glass aquarium (30 x 15 x 20 cm) of 10 L ( $n = 3 / L$ ), at a temperature of  $25 \pm 2$  °C, in light-dark cycles for 24 h and with chlorinated water (ProtecPlus®) and air pump with submerged filters, at 25 °C and pH 7.0, under a 14:10 h circadian cycle (light/dark). The animals received food (Alcon Gold Spirulina Flakes®) ad libitum 24 hours before the experiments.

### 2.4 Zebrafish

Adult fish were randomly selected from males and females and transferred to a wet sponge; then, they were treated orally with samples of HTMCX, drugs, or controls. After treatment, the fish were transferred (individually) to beakers (250 mL) containing 150 mL of water and left to recover before testing. After the tests, the fish were sacrificed by immersion in cold water ( $2-4$  °C) for 10 min until the end of the opercular movement (Matthews and Varga 2012). All procedures were approved by the Animal Use Ethics Committee of the State University of Ceará (CEUA-UECE; # 3344801/2017).

### 2.5 Toxicity to adult Zebrafish (ZFa)

Acute toxicity was performed against the ZFa for lethal concentration ( $LC_{50}$ ) for 96 hours in accordance with the guidelines of the Organization for Economic Cooperation and Development (OECD) (Test Guideline No. 203 - Fish, Acute Toxicity Testing 2019). After the OFT (see the section below), the fish ( $n = 6 / group$ ) were transferred to tanks, separated by treatment groups, and observed for 96 h. Dead fish in each group were counted, and the  $LC_{50}$  was determined.

## 2.6 Assessment of locomotor capacity

The open-field test (OFT) was carried out to identify changes in the zebrafish's motor coordination, either by sedation and/or muscle relaxation (Ahmad and Richardson 2013). The animals ( $n = 6$  / group) received 20  $\mu\text{L}$  of HTMCX orally at a dose of 1 mg / kg, 3 mg / kg or 10 mg / kg. The negative control group in each experiment was treated with the vehicle (3% DMSO; 20  $\mu\text{L}$ ), and the positive control group received Diazepam (10 mg / kg; 20  $\mu\text{L}$ ; *v.o.*). An untreated group (Naive) was also included. After 1 h, the fish were transferred to Petri dishes (10 cm  $\times$  15 cm; with quadrants at the bottom of the plate) containing water from the tank. Locomotor activity was assessed based on the number of times that each individual crossed the lines drawn in the Petri dishes in 5 min (Gonçalves et al. 2020).

## 2.7 Anxiolytic evaluation

An animal's anxiety behavior can be observed through the light / dark test (LDT). Similar to rodents, the zebrafish naturally avoids bright areas (Gonçalves et al. 2020). The experiment was carried out in a glass aquarium (30 cm  $\times$  15 cm  $\times$  20 cm) divided into a light area and a dark area. The water tank was filled to 3 cm with tap water without chlorine and without drugs, which simulated a new shallow environment different from the conventional housing aquarium and capable of inducing anxiety behavior. For zebrafish ( $n = 6$  / group) 20  $\mu\text{L}$  of HTMCX were administered orally at doses of 1 mg / kg, 3 mg / kg or 10 mg / kg. The negative and positive control groups consisted of 3% DMSO and 10 mg/kg Diazepam solution, respectively. An untreated group (Naive) was also included. After 1 h, zebrafish were placed individually in the clear zone, and the anxiolytic effect was measured based on the time spent in the clear zone within 5 minutes of observation (Gebauer et al. 2011).

## 2.8 Evaluation of the mechanism of action

The receptor (s) involved in the HTMCX anxiolytic type effect was identified by pretreatment with flumazenil (GABAA antagonist) and cyproheptadine serotonergic antagonists (5-HTR2A antagonist), pizotifen (antagonist of 5-HTR1 and 5-HTR2A / 2C), and granisetron (5-HTR3A / 3B antagonist) before LDT (Benneh et al. 2017). Zebrafish ( $n = 6$  / group) were pretreated with flumazenil (4 mg / kg; 20  $\mu\text{L}$ ; *i.p.*), cyproheptadine (32 mg / kg; 20  $\mu\text{L}$ ; *v.o.*), pizotifen (32 mg / kg; 20  $\mu\text{L}$ ; *v.o.*), or granisetron (20 mg / kg; 20  $\mu\text{L}$ ; *v.o.*). After 15 min, the highest effective dose of HTMCX (10 mg/kg; 20  $\mu\text{L}$ ; *v.o.*) found in the pilot test was administered. The 3% DMSO (vehicle; 20  $\mu\text{L}$ ; *v.o.*) was used as a negative control. Diazepam (Dzp; 10 mg / kg, 20  $\mu\text{L}$ ; *v.o.*) and fluoxetine (Flx; 0.05 mg / kg; *i.p.*) were used as GABA<sub>A</sub> and 5-HT agonists, respectively. After 1 hour of the treatments, the animals were submitted to the light / dark test described in the previous section.

## 2.9 Pentylentetrazole-induced seizure (PTZ)

PTZ-induced seizure reversal was investigated (Siebel et al. 2015). The animals ( $n = 6$  / group) were treated with HTMCX (10 mg / kg; 20  $\mu\text{L}$ ; *v.o.*), Diazepam (10 mg / kg; 20 $\mu\text{L}$ ; *v.o.*), vehicle (3% DMSO; 20 $\mu\text{L}$ ; *v.o.*). An untreated group ( $n = 6$  / group) was included (Naive). After 1 h, the animals were exposed to PTZ (7.5 mM), and the behavior similar to seizure in three stages was evaluated: stage I - dramatically

increased swimming activity; stage II - swirling swimming behavior; stage III - clonus-like seizures, followed by loss of posture when the animal falls to one side and remains immobile for 1–3 s. At the end of the evaluation of the three stages of the test, the animals were euthanized on the ice.

## 2.10. Docking procedure

The structure of the human 5-HT<sub>1B</sub> receptor (PDB 4IAQ) was obtained from the Protein Data Bank (<https://www.rcsb.org/>), identified as “Crystal structure of the chimeric protein of 5-HT<sub>1B</sub>-BRIL in complex with dihydroergotamine (PSI Community Target)”, deposited with a resolution of 2.80 Å, being determined by X-Ray diffraction, classified as signaling protein and electron transport, Homo sapiens organisms, *Escherichia coli* and *Spodoptera frugiperla* expression system (Wang et al. 2013). Molecular docking simulations were performed using the Auto Dock Vina code (version 1.1.2) (Trott and Olson 2010). The grid box was defined by centralizing the entire protein, with parameters of 70Åx100Åx76Å and dimensions (x, y, z) = (-21,133, -0,855, 17,314), with 50 independent simulations with 20 poses each. The Discovery Studio Visualizer (Biovia 2017) and UCSF Chimera (Pettersen et al. 2004) codes were used to analyzing the results.

### 2.11 Statistical analysis

The results were expressed as mean ± standard deviation from the mean for *in vivo* tests (n = 6 / group). After confirming the normality and homogeneity distribution of the data, differences between the groups were subjected to analysis of variance (one-way ANOVA), followed by the Tukey test, using the GraphPad Prism v software. 7.0. The level of statistical significance was considered to be 5% (p < 0.05).

## 3. Results

### 3.1 Acute toxicity (96 h)

The natural product HTMCX was non-toxic to adult zebrafish up to 96 h of analysis (LC<sub>50</sub> > 10 mg/kg), as there was no death and did not cause any apparent anatomical changes in the animals during this period.

### 3.2 Open field test (OFT)

HTMCX [**\*\*** p < 0.01, **\*\*\*** p < 0.001 (1; 3 or 10 mg / kg)] and Diazepam [**\*\*\*** p < 0.001 (10 mg / kg)] significantly decreased the locomotor activity of the adult zebrafish compared to control groups (Naive and vehicle) (Fig. 2A).

### 3.3 Anxiolytic evaluation (LDT)

HTMCX (3 or 10 mg / kg) caused (\*\*\*\*  $p < 0.0001$  vs Naive or vehicle) anxiolytic effect on ZFa in LDT (Fig. 2B). This effect was significantly similar to the effect of Diazepam (Dzp; 10 mg / kg; *v.o.*), positive control. Only the lowest dose of HTMCX (1 mg/kg) did not have an anxiolytic effect.

### 3.4 Involvement of the GABAergic system (GABA)

Flumazenil did not reduce the anxiolytic effect of acetophenone HTMCX (10 mg / kg, *v.o.*), and reduced (# #  $p < 0.01$  vs. Dzp) the anxiolytic effect of Diazepam (Dzp; 10 mg / kg; *v.o.*) (Fig. 2C).

### 3.5 Involvement of the serotonergic system (5-HT)

#### 3.5.1 System involvement 5-HTR<sub>2A</sub>

Cyproheptadine did not reverse the anxiolytic effect of HTMCX (10 mg / kg, *v.o.*), however, it reduced (# # #  $p < 0.0001$  vs. Flx) the anxiolytic effect of fluoxetine (Flx; 0.05 mg / kg; *i.p.*) (Fig. 2D).

#### 3.5.2 Involvement of 5-HTR<sub>1</sub> and 5-HTR<sub>2A/2C</sub> systems

Pizotifen reduced (# # # #  $p < 0.0001$  vs. HTMCX or Flx) the anxiolytic effect of HTMCX (10 mg / kg, *v.o.*) and fluoxetine (Flx; 0.05 mg / kg; *i.p.*), indicating that HTMCX has an anxiolytic effect through neuromodulation of the 5-HTR<sub>1</sub> and 5-HTR<sub>2C</sub> channels (Fig. 2E).

#### 3.5.3 Involvement of 5-HTR<sub>3A/3B</sub> systems

Granisetron did not reverse the anxiolytic effect of (10 mg / kg, *v.o.*), however, it reduced (# # # #  $p < 0.0001$  vs. Flx) the anxiolytic effect of fluoxetine (Flx; 0.05 mg / kg; *i.p.*) (Fig. 2F).

### 3.6 Pentylentetrazole-induced seizures (PTZ)

HTMCX (10 mg / kg; *v.o.*) did not reverse the convulsive behavior induced by PTZ, unlike [(\*\*  $p < 0.01$ , Stage I; \*\*\*  $p < 0.001$ , stage II; \*\*  $p < 0.01$ , stage III)] of Dzp (10 mg / kg; *v.o.*) that delayed the onset of the three stages of the seizure in the aZF compared to the control groups (Naive and vehicle) [(\*\*\*  $p < 0.001$ , Stage I and stage II; \*\*  $p < 0.01$ , stage III)] (Fig. 3).

### 3.7 Docking of the anxiolytic effect

The best conformation simulation between HTMCX and the receptor coupled to the human G 5-HT<sub>1B</sub> protein presented a RMSD (Root Mean Square Deviation) value in the order of 1,452 and an affinity energy value in the order of -6.0 kcal mol<sup>-1</sup>. The trimethoxyacetophenone-5-HT<sub>1B</sub> complex formed showed four interactions, two hydrophobic with residues ILE130A, VAL201A, and two hydrogen bonds of strong intensity with residues THR134A and ASP129A (Table 1).

## 4. Discussion

The OFT was initially performed to evaluate the effect of 2-hydroxy-3,4,6-trimethoxyacetophenone (HTMCX) on the locomotion of adult zebrafish. This test allows an evaluation of the stimulating or depressing activity of compounds and can also indicate more specific behaviors, such as anxiety. Thus, it was observed that HTMCX caused locomotor impairment (Fig. 2A) a result similar to those obtained with anxiolytic drugs that caused a sedative effect and decreased locomotor activity in animals (Gupta et al. 2014). In addition, HTMCX was non-toxic during the 96 h of analysis.

Zebrafish are anxious under natural conditions, preferring dark environments. Moreover, new environments are potentially risky and trigger anxiety in them. The preference of these animals for the clear region of the aquarium is characterized by the action of anxiolytic substances in the central nervous system (CNS) (Maximino et al. 2011). LDT was performed to confirm the possible effect of the HTMCX anxiolytic type observed in OFT. The higher doses of acetophenone HTMCX and Diazepam significantly increased the time of the animals in the clear region of the aquarium (Fig. 2B), allowing the assumption that this natural product has an acute anxiolytic effect, confirming the results observed in the OFT.

LDT and OFT showed that HTMCX and Diazepam have similar effects on zebrafish. Benzodiazepines have a depressant/sedative effect on the CNS of zebrafish and mammals through positive allosteric neuromodulation of the GABA<sub>A</sub> receptor, which causes hypnotic effects, locomotor deficiency, and anxiolytic effects (Griffin et al. 2013). Flumazenil is a GABAergic receptor antagonist well known for antagonizing the sedative / hypnotic effects caused by benzodiazepine overdoses. It reverses these effects by binding to the  $\alpha$  [1–3, 5]  $\beta\gamma$  subunits of these receptors (Penninga et al. 2016). Pretreatment with flumazenil did not alter the effects of HTMCX on zebrafish (Fig. 2C). However, flumazenil sharply reduced the anxiolytic effect in fish treated with Diazepam. However, the anxiolytic activity of HTMCX is independent of the GABAergic system.

Pentylenetetrazole is also an antagonist of GABAergic receptors; however, it is used in zebrafish to induce epileptic-like effects and to study seizures (Shaikh et al. 2013). The anticonvulsant effect of HTMCX and Dzp on PTZ-induced seizures in adult zebrafish was evaluated. However, HTMCX did not increase the latency time for the onset of seizures (Fig. 3). Unlike Dzp, which increased the latency time for the onset of crises in the three stages. These results corroborate data obtained from the mechanism of action of

the GABA<sub>A</sub> receptor with flumazenil (Fig. 2C), demonstrating that there is no action of HTMCX in the GABAergic system.

Serotonergic neuromodulation is also involved in the mechanisms of anxiety mediation, and the role of serotonin (5-HT) in this disorder is widely investigated (Maximino et al. 2014). Anxiety behavior in LDT is positively associated with extracellular levels of 5-HT in the zebrafish brain (Gonçalves et al. 2020). Thus, high levels of 5-HT can cause effects similar to those of anxiety, while low levels generally cause anxiolytic behaviors (Nowicki et al. 2014).

Considering the hypothesis that serotonergic neurotransmission mediates anxiolytic effects, the participation of this system in the anxiolytic action of HTMCX after pretreatment with cyproheptadine antagonists (a 5-HTR<sub>2A</sub> antagonist), pizotifen (a 5-HTR<sub>1</sub> and 5-HTR<sub>2A/2C</sub> antagonist), and granisetron (a 5-HTR<sub>3</sub> antagonist) was evaluated. Fluoxetine was used as a positive control. Unlike granisetron (5-HTR<sub>3A/3B</sub>; Fig. 2E) and cyproheptadine (5-HT<sub>2A</sub>; Fig. 2D), pretreatment with pizotifen significantly inhibited the acute anxiolytic effect of HTMCX, suggesting that the mechanism of action of this acetophenone involves the 5-HTR<sub>1</sub> and/or 5HT<sub>2C</sub> serotonergic receptors (Fig. 2D). However, it does not involve the 5HT<sub>2A</sub> receptor because cyproheptadine did not reverse the anxiolytic effect of HTMCX on LDT. In addition, it was observed that all the mentioned antagonists reversed the effects of fluoxetine (Fig. 2C, D, and F).

5HT<sub>1A</sub> receptor agonist substances exhibit antidepressant and/or anxiolytic effects. 5-HTR<sub>1</sub> is an inhibitory receptor coupled to protein G, and its activation reduces the rate of firing of serotonergic neurons, promotes the synthesis, renewal, and release of 5-HT in various areas of the CNS (Gonçalves et al. 2020). Studies have shown that zebrafish treated with 5HT<sub>1A</sub> receptor agonists showed anxiolytic behaviors (Benneh et al. 2017; Gonçalves et al. 2020; Maximino et al. 2014). Therefore, HTMCX is possibly a 5-HTR<sub>1A</sub> agonist, as it prevented the animals' anxiolytic effect (Fig. 2E).

The serotonin 2C receptor (5-HTR<sub>2C</sub>) is a molecular target of drugs developed for the treatment of behavioral conditions, such as eating and mood disorders, anxiety, and motor behavior (Lee et al. 2010). The genetic similarities of mammalian 5-HTR<sub>2C</sub> with that of zebrafish indicate the use of this model to study the role of the 5-HT<sub>2C</sub> receptor in the behavior, development, and discovery of drugs that act in this channel. Reports indicate that 5-HTR<sub>2C</sub> activation induces panic and/or anxiety while blocking it causes anxiolytic effect (Chagraoui et al. 2016). Thus, HTMCX may have competed with pizotifen and was replaced by it in 5-HTR<sub>2C</sub>, being indicative of the action of this acetophenone in the anxiolytic effect under this route.

For the study of the anxiolytic effect *in silico*, via serotonergic neuromodulation of HTMCX, the structure of the receptor coupled to the human G 5-HT<sub>1B</sub> protein was selected due to its orthostatic site composed of residues conserved in practically the entire family of the receptor 5-HT (Morgan et al. 2013). The dihydroergotamine agonist (2GM) is complexed in the protein at the orthostatic site, composed of

residues ILE130, CYS133, THR134, VAL201, ALA216, PHE331, ASP352, and THR355. After the molecular docking simulations, it was possible to observe that the HTMCX coupled in the same region of the orthostatic site (Fig. 4), showing four interactions that vary between 2.51Å and 3.89Å in distance, with three interactions with the residues of the dihydroergotamine site ILE130A, VAL201A, THR134A (two hydrophobic interactions and a hydrogen bond of strong intensity) and a hydrogen bond with the ASP129A residue (Table 1).

## 5. Conclusion

The results of this study show that acetophenone HTMCX was non-toxic and demonstrated an anxiolytic effect in adult zebrafish. Acetophenone HTMCX exhibited an anxiolytic effect through the 5-HT<sub>1</sub> and 5-HT<sub>2C</sub> systems, having the potential to form a complex in the same region of the orthosteric site, indicating that the anxiolytic effect via the serotonergic mechanism. Our results confirm the pharmacological relevance of HTMCX as an anxiety inhibitor, enabling new therapeutic approaches to be investigated.

## Declarations

**Authors' contributions** AWS performed the experiments, analyzed the data, prepared figures and tables, authored and wrote the manuscript. MKAF and JESAM conceived and designed the experiments. ELR, FRSM and ALSM analyzed the data and reviewed drafts of the manuscript. MMM, and ESM performed the docking molecular study and reviewed the paper. HSS and AMRT contributed reagents/materials/analysis tools, supervised the research and reviewed the manuscript. All authors read and approved the final version of the manuscript. The authors declare that all data were generated in-house and that no paper mill was used.

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**Conflicts of interest/ Competing interests** The authors declare no conflicts of interest. The authors also declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

**Availability of data and material** All data generated or analyzed during this study are included in the Supplementary Material.

**Code availability** The authors thank Centro Nacional de Processamento de Alto Desempenho (CENAPAD) of the Federal University of Ceará (UFC) for the use of the AutoDock Vina software. The Discovery Studio Visualizer and UCSF Chimera free software were used to analyzing the results of the research.

### **Compliance with ethical standards**

**Ethics approval** All procedures performed in studies involving zebrafish-based animal were approved by the Animal Use Ethics Committee of the State University of Ceará (CEUA-UECE; # 3344801/2017). All methods were performed in accordance with the relevant guidelines and regulations.

**Consent to participate** All authors gave their consent to participate in this study.

**Consent for publication** All authors gave their consent for the publication of this manuscript.

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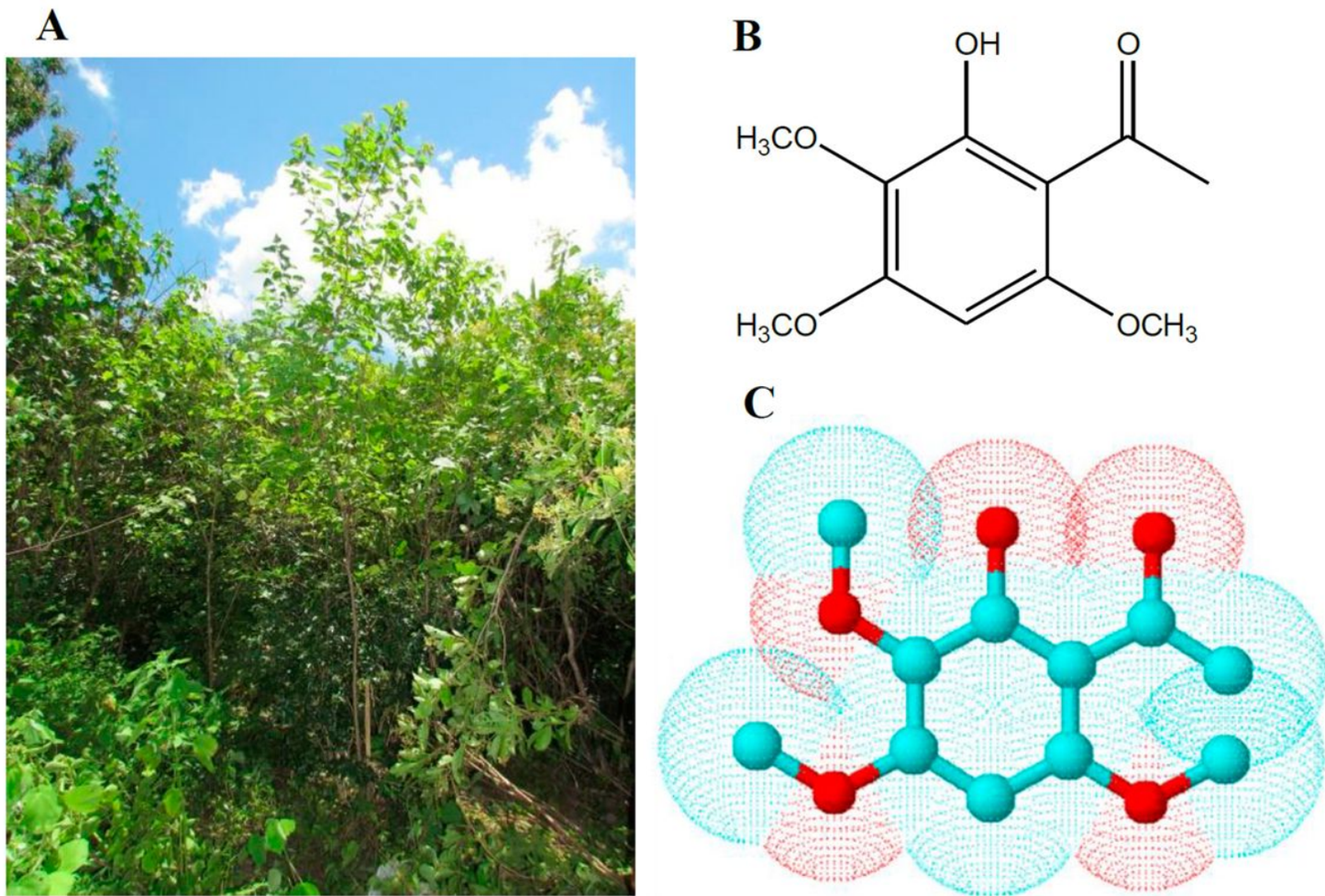
## Tables

**Table 1.** Interactions between the 5-HT1B receptor and the HTMCX ligand.

Ligand	Receptor	Interaction	Distance (Å)
HTMCX	ILE130A*	Hydrophobic	3.89 Å
	VAL201A*	Hydrophobic	3.86 Å
	ASP129A	H-Bond	2.77 Å
	THR134A*	H-Bond	2.51 Å

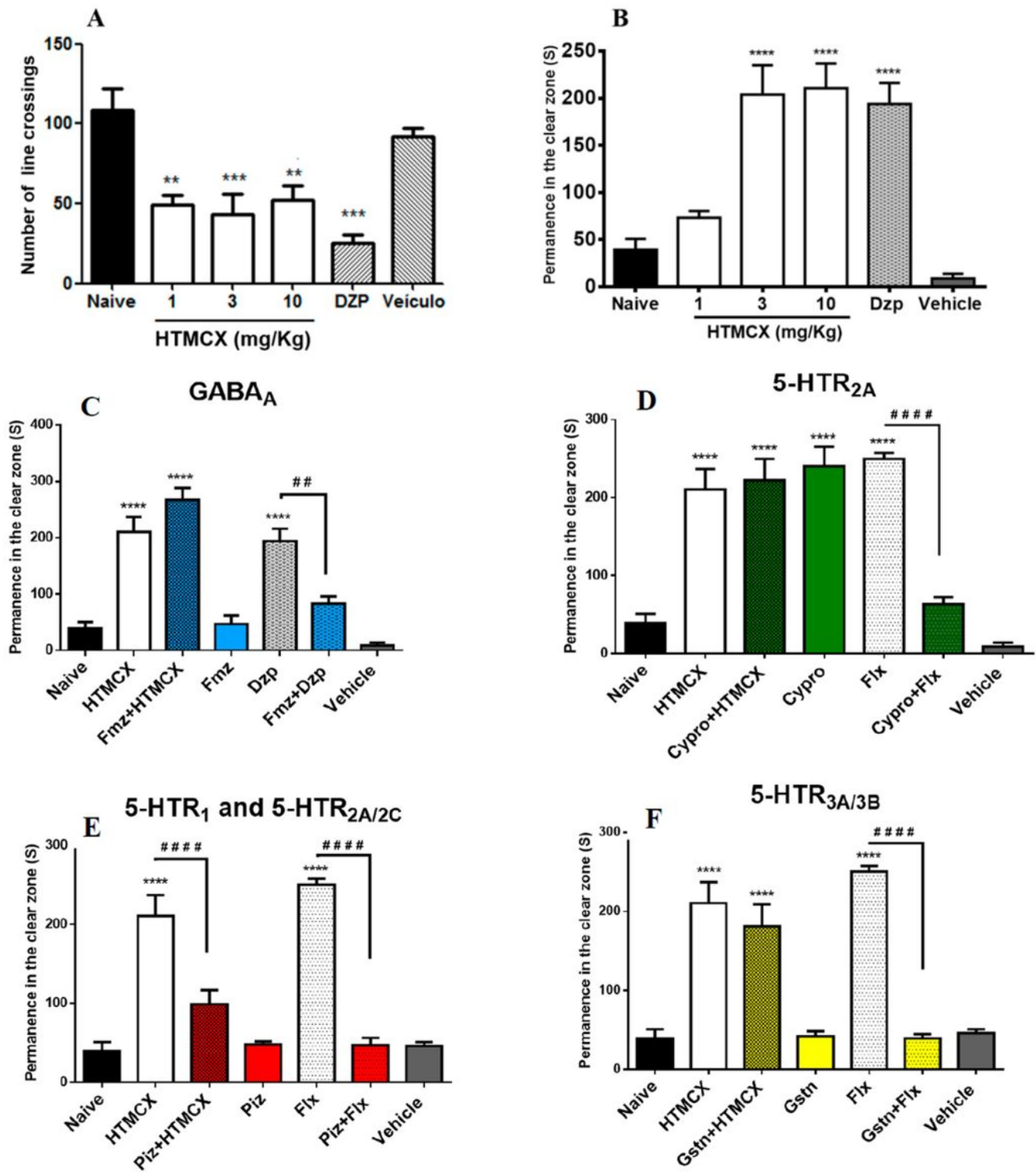
\* 2GM orthostatic site residue.

## Figures



**Figure 1**

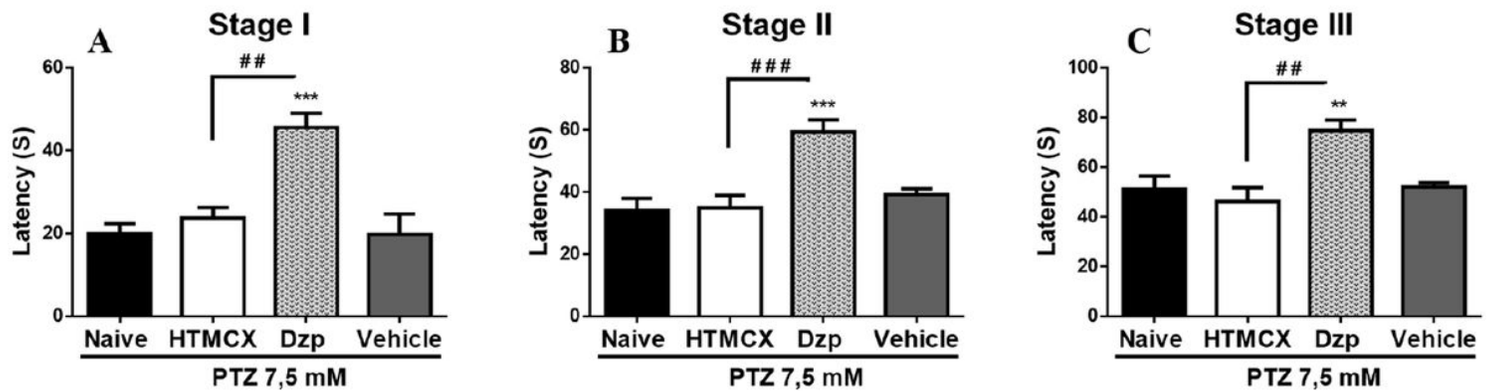
Croton anisodontus (A), structural representation (B) and three-dimensional structure (C) of the 2-hydroxy-3,4,6-trimethoxyacetophenone (HTMCX)



**Figure 2**

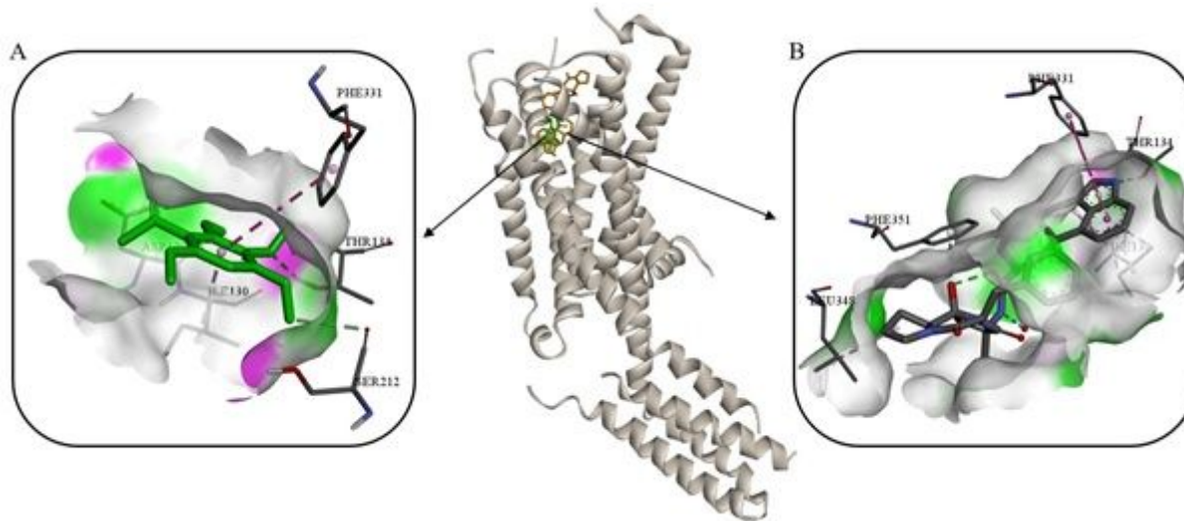
Effect under the locomotor behavior of zebrafish (*Danio rerio*) adult in the Open Field Test (A) and anxiolytic-like effect in the Light & Dark Test (B) of the HTMCX (20  $\mu$ L; v.o.). Anxiolytic-like effect of HTMCX after pretreatment with flumazenil (C), cyproheptadine (D), pizotifen (E), and granisetron (F) in adult zebrafish in the light/dark test (0–5 min). Naive-untreated animals. Dzp - Diazepam (10 mg/kg; 20  $\mu$ L; v.o.). Vehicle (DMSO 3%) (20  $\mu$ L; v.o.). The results are expressed as mean values  $\pm$  S E M. (n =

6/group). ANOVA followed by the Tukey test (\*\*p < 0.01, \*\*\*p < 0.001, \*\*\*\*p < 0.0001 vs. Naive or Vehicle; ##p < 0.01, ####p < 0.0001 vs. HTMCX or Flx or Dzp). Dzp – Diazepam, Cipro – cyproheptadine; Gsxn – granisetron; Piz – pizotifene; Flx – fluoxetine.



**Figure 3**

Effect of HTMCX on pentylentetrazole-induced seizure in adult zebrafish, Stage I (A), Stage II (B), Stage III (C). Dzp – Diazepam (10 mg/kg; 20  $\mu$ L, v.o.); Vehicle – 3% DMSO (20  $\mu$ L; v.o.). The values represent the mean  $\pm$  standard error of the mean (E.P.M.) for 6 animals/group. ANOVA followed by Tukey (\*\* p < 0.01, \*\*\* p < 0.001 vs. Naive or Vehicle; # # p < 0,01; # # # p < 0.001 vs. Dzp).



**Figure 4**

5-HT1B receptor interaction complex with acetophenone (A) and the co-crystallized inhibitor Dihydroergotamine (B).

## Supplementary Files

This is a list of supplementary files associated with this preprint. Click to download.

- [TABLE1Interaction5HT1BreceptorwiththeHTMCXligand.rar](#)

- [Excel file with the data generated in Figures 13.xlsx](#)
- [FIGURE4MolecularDocking.rar](#)
- [Prism file of the statistical data.pzfx](#)