Chrysin, but not flavone backbone, decreases anxiety-like behavior in animal screens

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- 31 Conflicts of interest: None declared
- 32

33 **Running title:** Anxiolytic-like effects of chrysin and flavone

34 Abstract

Chrysin (5,7-dihydroxyflavone), a nutraceutical flavonoid present in diverse plants, has 35 a backbone structure shared with the flavone backbone, with additional hydroxyl groups 36 that confers its antioxidant properties and effects at the GABA_A receptor complex. 37 However, whether these effects are due to the hydroxyl groups is unknown. Here we 38 39 report the effects of chrysin or the flavone backbone (1 mg/kg) in rats subjected to the 40 elevated plus-maze and the locomotor activity test, as well as in the zebrafish evaluated in light/dark model. Chrysin, but not flavone, increased entries and time in the open 41 arms of the elevated plus-maze, as well as time on white compartment of the light/dark 42 model in zebrafish. These effects were comparable to diazepam, and were devoid of 43 44 motor effects in both tests, as well as in the locomotor activity test. On the other hand, 45 flavone decreased risk assessment in the light/dark test but increased rearing in the locomotor activity test in rats, suggesting effects threat information gathering; important 46 47 species differences suggest new avenues of research. It is suggested that the specific effects of chrysin in relation to flavone include more of a mechanism of action in which 48 49 in addition to its action at the GABA_A/benzodiazepine receptor complex also could be involved its free radical scavenging abilities, which require specific research. Preprint: 50 https://doi.org/10.1101/575514; Data 51 and scripts: https://github.com/lanec-52 unifesspa/chrysin

53 Keywords: Chrysin; Flavone; Anxiety; Rat; Zebrafish

54 Introduction

55 Flavonoids are polyphenolic compounds present in diverse plants (Ghosh & 56 Scheepens, 2009; Krafczyk et al., 2009) and have been considered as nutraceuticals 57 due to they are part of the human diet (Marventano et al., 2017) and produce diverse pharmaceuticals actions. Also, they are able to cross the blood-brain barrier and modify 58 59 the brain function (Johnston et al., 2015). Particularly, the flavonoid chrysin (5,7-60 dihydroxyflavone) has been studied by its antioxidant properties (Siddiqui et al., 2018), but currently also due to its neuropharmacological actions produced on specific brain 61 62 structures implicated in neuropsychiatric disorders (Filho et al., 2016; Bortolotto et al., 63 2018). Chrysin is a natural flavonoid present in honey, propolis and diverse plant extracts (German-Ponciano et al., 2018), and is one of the most studied polyphenolic 64 nutraceuticals (Chadha et al., 2017). Chrysin is used as a nutraceutical as a 65 66 "testosterone boosting agent" (a claim that is probably very exaggerated; Gambelunghe 67 et al., 2003), but there is some preclinical evidence that this molecule ameliorates indices of hepatic and renal functioning in diabetic animals (Ramanathan & 68 Thekkumalai, 2014; Yamamoto, 2014). There is also preclinical evidence that this 69 70 flavonoid exerts anxiolytic-like effects (Wolfman et al., 1994; Salqueiro et al., 1997; Zanoli et al., 2000; Rodríguez-Landa et al., 2019). 71

72 Chrysin has a backbone structure that consists in two aromatic rings (A and C), and a 73 phenyl B ring, which is attached to the second position of ring C and shares the flavone backbone, with an additional hydroxyl group in the 5 and 7 positions of the A ring (Mani 74 75 et al., 2017; Figure 1). The potential of chrysin to act as free radical scavenger has been 76 attributed to the presence of these hydroxyl groups (Sathiavelu et al., 2009; Souza et 77 al., 2015), and it has been suggested that this functional group represents the main 78 action site of this flavonoid to produce a great variety of potential therapeutic effects to 79 ameliorate diverse physiological and psychiatric disorders (Sheela & Augusti, 1995). 80 However, no studies have directly compared the effects of similar doses of chrysin and the flavone backbone on the potential therapeutic effects reported at preclinical 81 82 research.

83

84

--- Insert Figure 1 about here ----

85 Chrysin (either isolated from Passiflora Coerulea, Passiflora incarnata, Matricaria Chamomilla or as a synthetic drug) exerts anxiolytic-like effects through action on the 86 GABA_A receptors without the associated side effects (i.e. sedation, amnesia, 87 88 myorelaxation) that have been attributed to other GABA_A modulators (benzodiazepines, ethanol, etc.); these effects have been evaluated in behavioral models using rodents 89 90 (Wolfman et al., 1994; Salgueiro et al., 1997; Zanoli et al., 2000; Rodríguez-Landa et al., 2019). Although it has been suggested that presence of the hydroxyl groups in the 91 92 chrysin structure is the responsible for this pharmacological action, this effect has not 93 been compared with the effects produced by the flavone backbone in preclinical models 94 of anxiety-like behavior.

In addition to mammalian model organisms used to study the anxiolytic-like effects of 95 diverse substances, non-mammalian models have been developed to explore the 96 97 potential therapeutic effects of drugs in a short time (Norton & Bally-Cuif, 2010). The 98 use of zebrafish in preclinical research in the psychopharmacology field has increased recently. Biobehavioral assays of anxiety and screening tests for anxiolytic-like effects, 99 100 such as the rodent light/dark test (LDT), have been adapted to zebrafish, considering 101 that behavioral phenotypes are similar between rodents and zebrafish (Maximino et al., 102 2010a). Behavioral screens using zebrafish have evaluated the effects of clinically 103 effective anxiolytic drugs, including benzodiazepines (Bencan et al., 2009; Gebauer et 104 al., 2011; Maximino et al., 2011; Schaefer et al., 2015) and ligands of both the central 105 and peripheral BZD sites (Lima-Maximino et al., 2018), suggesting good 106 pharmacological isomorphism and the ability of such screens to identify potential anxiolytic drugs that can progress in the drug development pipeline. In addition to 107 108 increasing the toolbox of behavioral screens for psychoactive drugs, zebrafish screens 109 and biobehavioral assays also increase our understanding of the "core" mechanisms of behavioral functions, such as fear and anxiety, that are altered in human patients and 110 111 shared among fish, rodents, and humans (Gerlai, 2014).

112 The aim of the present study was to compare the effects of the nutraceutical flavonoid chrysin and the flavone backbone (the basic structure of the flavone without the 113 hydroxyl radical groups), using the LDT in zebrafish, as well as the elevated plus-maze 114 115 (EPM) and locomotor activity test in rats. This could contribute to identify if anxiolyticlike effects produced by chrysin are due to the presence of hydroxyl radicals in its basic 116 structure, which could impact in the specific design of novel molecules destined to 117 pharmacotherapy of anxiety. This manuscript is a complete report of all the studies 118 performed to test the effect of chrysin, flavone, or diazepam on anxiety-like behavior in 119 120 rats and zebrafish. We report all data exclusions (if any), all manipulations, and all 121 measures in the study.

122 Materials & methods

123 Animals

124 Rats. 32 adult male Wistar rats (2.5 months old), weighing 200-250 g, were used. Animals were housed in Plexiglas cages (4-5 rats per cage) in the Laboratorio de 125 126 Neurofarmacología (Universidad Veracruzana, Xalapa, Mexico) under a 12 h/12 h light/dark cycle (lights on at 7:00 AM) at 25°C ± 1°C with ad libitum access to food and 127 water. The experimental procedures were performed in accordance with national 128 (Especificaciones Técnicas para la Producción, Cuidado y Uso de Animales de 129 Laboratorio NOM-062-ZOO-1999) and international (National Research Council (US) 130 Committee for the Update of the Guide for the Care and Use of Laboratory Animals, 131 2011) ethical recommendations. All efforts were made to minimize animal discomfort 132 133 during the study.

Zebrafish. 48 adult unsexed zebrafish of the longfin phenotype were used in the present experiments. Animals were acquired in a local aquarium shop (at Belém, Brazil) and kept in collective tanks at the Laboratório de Neurofarmacologia e Biofísica (UEPA, Marabá, Brazil) for at least 2 weeks before experiments started. Conditions in the maintenance tank were kept stable, as described by Lawrence (2007) and recorded in protocols.io (Maximino et al., 2019; [https://dx.doi.org/10.17504/protocols.io.rupd6vn]). Recommendations in the Brazilian legislation (Diretriz brasileira para o cuidado e a

- utilização de animais para fins científicos e didáticos DBCA. Anexo I. Peixes mantidos
 em instalações de instituições de ensino ou pesquisa científica, 2017) were followed to
- 143 ensure ethical principles in animal care and throughout experiments.
- 144

145 **Drugs and treatments**

Chrysin (CAS #480-40-0; Sigma-Aldrich, C80105) and flavone (CAS #525-82-6; Sigma-146 Aldrich, F2003) were dissolved in 5% dimethyl sulfoxide (DMSO) and injected *i.p.* For all 147 148 drugs, the dose was 1 mg/kg, a dose of chrysin shown to be anxiolytic in elevated plusmaze (Wolfman et al., 1994). Thirty-two rats were assigned to four independent groups 149 (n = 8/group). The vehicle group received 5% DMSO solution (Golden Bell Reactivos, 150 México City, México) in which chrysin and diazepam were prepared; a second group 151 (CHR) received 1 mg/kg chrysin; a third group (FLAV) received 1 mg/kg flavone; and a 152 153 fourth group (DZP) received 2 mg/kg diazepam (CAS #439-14-5; Tocris, 2805; German-Ponciano et al., 2020) as a pharmacological control of anxiolytic activity. Sixty minutes 154 after the injection, the rats were evaluated in the elevated plus maze and then in the 155 locomotor activity test. The time elapsed between injection and the behavioral test has 156 previously demonstrated anxiolytic-like effects of both chrysin and diazepam at the 157 158 doses that were uesed in present study (Taiwo et al., 2012; Rodríguez-Landa et al., 2019; Cueto-Escobedo et al., 2020; German-Ponciano et al., 2020). 159

Zebrafish were randomly drawn from the holding tank immediately before injection and 160 161 assigned to four independent groups (n = 12/group). Groups were identical to those used for rats, including vehicle (5% DMSO). The injection volume was 1 µL/0.1 g of 162 163 body weight, and animals were injected intraperitoneally (Kinkel et al., 2010). The order with which groups were tested was randomized via generation of random numbers 164 using the randomization tool in http://www.randomization.com/. Experimenters were 165 blinded to treatment by using coded vials for drugs. The data analyst was blinded to 166 phenotype by using coding to reflect treatments in the resulting datasets; after analysis, 167 data was unblinded. 168

169 Elevated plus-maze

170 The rats were separately evaluated in the elevated plus maze, and then in the locomotor activity test. On the testing day, the rats were brought to the experimental 171 room at 11:00 AM and left for 1 h to acclimate to the novel surroundings. The behavioral 172 tests began at 12:00 AM. The apparatus consisted of two opposite open and closed 173 arms set in a "+" configuration, and it was situated in an illuminated room at 40 lux. The 174 open arms were 50 cm length x 10 cm width, and the closed arms were 50 cm length x 175 176 10 cm width, with 40 cm-high walls. The entire maze was elevated 50 cm above the floor (Walf & Frye, 2007). To evaluate the effects of the treatments, the rats were 177 individually placed in the center of the maze, facing an open arm. The following 178 variables were scored during 5 min: 179

- 180 1. Number of entries into the open arms (N);
- 181 2. Number of entries into the closed arms (N);
- 182 3. Total number of entries (open arms + closed arms), N;
- 4. *Percentage of open arm entries* (Proportion of entries made in the open arms in relation to the total number of entries)
- 185 5. *Time spent on the open arms* (s);
- 186 6. *Time spent in head-dipping* (s);
- 187 7. Number of head-dipping events (N);
- 188 8. Time spent in stretched-attend posture (SAP; s);
- 189 9. Number of SAP events (N).
- 190
- After the elevated plus maze test, the rats underwent the locomotor activity test.Approximately 2 min elapsed between tests.
- 193

194 Locomotor activity test

195 Each rat was individually placed in the locomotor activity cage (44 cm length \times 33 cm 196 width \times 20 cm height). The floor of the cage was delineated into twelve 11 cm \times 11 cm 197 squares to evaluate spontaneous locomotor activity, grooming, and rearing for 5 min. At

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the beginning of the test, the rat was gently placed in one of the corners of the cage.The following variables were scored:

- Number of squares crossed (crossings; a crossing was considered when the rat passed from one square to another with its hind legs);
- 202
 2. *Time spent rearing* (rearing was considered when the rat emitted a vertical position relative to the cage floor);
- *Time spent grooming* (including paw licking, nose/face grooming, head washing, body grooming/scratching, leg licking, and tail/genital grooming).

Digital video cameras (Sony DCR-SR42, 40 optical zoom, Carl Zeiss lens) were 206 207 installed above each apparatus (elevated plus maze and locomotor activity test) to 208 record activity. Two independent observers measured the behavioral variables using ex profeso software to record the number and time (in seconds) of each evaluated 209 behavioral variable until >95% agreement was reached among the observers. After 210 each test session, the apparatus was carefully cleaned with a 10% ethanol solution to 211 212 remove the scent of the previous rat, which can influence spontaneous behavior of the subsequent rat. 213

214

215 Light/dark test

The light/dark preference (scototaxis) assay was performed as described elsewhere 216 (Maximino, 2018 [https://10.17504/protocols.io.srfed3n]; Maximino et al., 2010b). Briefly, 217 30 min after injection animals were transferred individually to the central compartment of 218 219 a black/white tank (15 cm height X 10 cm width X 45 cm length) for a 3-min acclimation period, after which, the doors which delimit this compartment were removed and the 220 221 animal was allowed to freely explore the apparatus for 15 min. While the whole experimental tank was illuminated from above by a homogeneous light source, due to 222 the reflectivity of the apparatus walls and floor average illumination (measured just 223 above the water line) above the black compartment was 225 ± 64.2 (mean \pm S.D.) lux, 224 while in the white compartment it was 307 ± 96.7 lux. The following variables were 225 recorded: 226

- 1. *Time spent on the white compartment:* the time spent in the white half of the tank(s);
- 229 2. *Transitions to white:* the number of entries in the white compartment made by the 230 animal throughout the session;
- 3. *Entry duration:* the average duration of an entry (time on white / transitions);
- 4. *Erratic swimming:* defined as the number of zig-zag, fast, and unpredictable
 swimming behavior of short duration;
- 5.*Freezing:* the duration of freezing events (*s*), defined as complete cessation of movements with the exception of eye and operculum movements;
- 6. *Thigmotaxis:* the duration of thigmotaxic events (*s*), defined as swimming in a distance of 2 cm or less from the white compartment's walls;
- 7.Frequency of risk assessment: defined as a fast (<1 s) entry in the white
 compartment followed by re-entry in the black compartment, or as a partial entry in
 the white compartment (i.e., the pectoral fin does not cross the midline);
- 241 242

A digital video camera (Samsung ES68, Carl Zeiss lens) was installed above the apparatus to record the behavioral activity of the zebrafish. Two independent observers, blinded to treatment, manually measured the behavioral variables using X-Plo-Rat 2005 (<u>https://github.com/lanec-unifesspa/x-plo-rat</u>). Inter-observer reliability was at least > 0.95.

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- 249

250 Statistical analysis

251 Data were analyzed using Asymptotic General Independence Tests, a permutationbased analogue of one-way ANOVA, followed by pairwise permutation tests, analogous 252 to pairwise t-tests with *p*-values adjusted for the false discovery rate. Analyses were 253 made in R (R Core Team, 2019), using the package 'coin' (Hothorn et al., 2006). 254 Normality was not assumed, and thus no specific test for normality was performed; 255 however, this type of analysis is resistant to deviations from the assumptions of the 256 257 traditional ordinary-least-squares ANOVA, and is robust to outliers, thus being 258 insensitive to distributional assumptions (such as normality) (Mangiafico, 2015). This test is based on the permutation of a statistic maxT, the maximum of several competing univariate statistics. Outliers were detected using an *a priori* rule based on median absolute deviation (MAD) of time on white (the main endpoint of the LDT) or time in the open arms (the main endpoint of the EPM), with values above or below 3 MADs being removed (Leys et al., 2013). Using this procedure, one outlier was removed from the flavone-treated group in zebrafish.

265

266 **Open science practices**

267 Hypotheses and methods were not formally pre-registered. Data and analysis scripts 268 can be found at a GitHub repository (<u>https://github.com/lanec-unifesspa/chrysin</u>).

269

270 Results

271 Elevated plus-maze

In the elevated plus-maze (Figure 2A), animals made more entries in the open arms 272 (Figure 2B; maxT = 2.4407, p = 0.05) after the treatment with chrysin (p = 0.02) and 273 diazepam (p = 0.02) in comparison to the control group. No effect was found for entries 274 in the closed arm (Figure 2C; maxT = 1.0295, p = 0.63) or total entries (Figure 2D; 275 maxT = 1.5331, p = 0.36). A significant effect was found for percentage of entries in the 276 277 open arms (Figure 2E; maxT = 2.7848, p = 0.02), but differences disappeared after adjusting *p*-values in *post-hoc* contrasts. A significant effect was found for time spent in 278 the open arms (Figure 2F; maxT = 2.6129, p = 0.03), this variable was increased after 279 treatment with chrysin (p = 0.02) and diazepam (p = 0.01). No effects were found for 280 281 time spent head-dipping (Figure 2G; maxT = 1.3025, p = 0.51) or number of headdipping events (Figure 2H; maxT = 0.853, p = 0.8). A significant treatment effect was 282 found for time spent in SAP (Figure 2I; maxT = 2.5326, p = 0.04), but meaningful 283 differences disappeared after adjusting p-values in post-hoc contrasts. Finally, a 284

285 significant effect was found for SAP frequency (Figure 2J; maxT = 3.2229, p = 0.01), with flavone (p = 0.02) and diazepam (p = 0.01) reducing SAP. 286 287 --- Insert Figure 2 about here ---288 Locomotor activity test 289 In the locomotor activity test (Figure 3A), none of the treatments modified the number of squares crossed (Figure 3B; maxT = 1.6296, p = 0.31) and grooming behavior (Figure 290 3C; maxT = 1.5029, p = 0.38); but significant differences were founded in rearing 291 (Figure 3D; maxT = 4.0090, p = 0.01). Post-hoc test revealed that flavone backbone 292 293 increased this variable (p = 0.01) respect to the control group.

295

296 Light/dark test

In the light/dark test (Figure 4A), chrysin (p = 0.02) and diazepam (p = 0.02), but not the flavone backbone (p = 0.14), significantly increased (maxT = 3.7331, p = 0.01) the time spent into the white compartment (Figure 4B) compared to the control group. No significant changes were detected in the duration of entries (maxT = 2.0981, p = 0.12) or transitions (maxT = 0.6707, p = 0.89) to the white compartment (Figures 4C and 4D, respectively), associated with the treatments.

On the other hand, the flavone backbone (p = 0.02), but neither chrysin (p = 0.06) nor diazepam (p = 0.06), significantly (maxT= 2.6404, p = 0.03) reduced risk assessment (Figure 4E) in relation to the control group. Diazepam (p = 0.031), but not chrysin (p =0.60) or flavone (p = 0.50), increased erratic swimming (Figure 4F; maxT = 3.9468, p =0.01). No significant changes were observed in thigmotaxis (Figure 4G; maxT = 1.4946, p = 0.39), or freezing duration (Figure 4H; maxT = 2.3935, p = 0.06) between treatments. 310

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--- Insert Figure 4 about here ---

312 **Discussion**

The present study demonstrated that 1) the anxiolytic-like effect of the flavonoid chrysin 313 314 in the rat EPM and LDT, without locomotor effects, seems to be associated with hydroxyl groups substituted in 5 and 7 positions of the A ring in the flavone backbone; 315 and 2) the flavone backbone is devoid of anxiolytic-like effects in the rat EPM in rats, but 316 317 decreases risk assessment in the LDT in zebrafish and rearing in the locomotor activity test. These results contribute to the knowledge that positions of hydroxyl in the flavone 318 319 backbone are important structural characteristic to produces its pharmacological action 320 on the central nervous system, and suggest that a comparative approach is best suited to detect conserved drug effects across species that are more likely to be shared with 321 322 humans.

323 In the present experiments, after chrysin treatment the number of entries into and time spent into the open arms were increased, without locomotor effects either in the EPM or 324 in the locomotor activity test, an effect that is consistent with anxiolysis and is 325 comparable to DZP. The effects of chrysin in the EPM have been shown before. 326 327 Wolfman et al. (1994) showed that chrysin (1 mg/kg), isolated from Passiflora coerulea, 328 exerts an anxiolytic-like effect in the EPM. We have previously shown that chrysin (2 329 and 4 mg/kg) blocked the anxiogenic-like effects of long-term ovariectomy in female 330 rats, which were attributed to the activation of GABAA receptors, as they were blocked by pretreatment with picrotoxin (Rodríguez-Landa et al., 2019). 331

Similar effects were observed in the LDT, a biobehavioral assay that is based in the preference of zebrafish for dark environments (Serra et al., 1999) and has been validated pharmacologically as a model to screening diverse anxiolytic drugs (Maximino et al., 2011; Magno et al., 2015). In this sense, the flavonoid chrysin increased the time spent in the white compartment in comparison to the control group. This variable is the main endpoint of the LDT (Maximino et al., 2011; Lima-Maximino et al., 2018), also clinically effective anxiolytic drugs that target different sites of the GABA_A receptor, like
diazepam and other benzodiazepines, can increase it (Gebauer et al., 2011; Maximino
et al., 2011; Magno et al., 2015), which has been considered as an anxiolytic-like effect
at preclinical-research.

However, the flavone backbone did not modify the time spent into the open arms in the 342 343 EPM neither the time spent in the compartment in the LD, in relation to the control 344 groups. The basic flavone moiety (backbone) has a hydrogen bond donor site, lipophilic pockets, and an electron rich site (Marder & Paladini, 2002), which are thought to be 345 important for the binding of both flavonoids and 1,4-benzodiazepines to the central 346 benzodiazepine (BZD) site at GABA_A receptors (Marder & Paladini, 2002). Since the 347 348 chrysin hydroxyl groups are located in one region of the flavone basic ring, this could be important for the steric interactions associated with binding, and therefore chrysin could 349 350 bind to BZD sites with a higher affinity than flavone (Marder & Paladini, 2002; Huen et 351 al., 2003). This could explain the wider anxiolytic-like effect of chrysin, but not flavone backbone, in the present study. 352

353 In addition to the effects of chrysin and flavone on the BZD site, another potential mechanism is a free radical scavenging effect: chrysin, like most flavonoids, exerts 354 powerful antioxidant effects, because it possesses highly reactive hydroxyl groups in its 355 356 basic structure, which confers a potent free radical scavenging activity (Panche et al., 2016). It has been reported that the generation of free radicals promote the inactivation 357 358 of the enzyme glutamate decarboxylase (responsible for the synthesis of GABA), leading to a decrease of GABA concentrations and facilitating excitatory 359 neurotransmission (Singh et al., 2014), and thus the scavenging potential of chyrsin 360 could enhance the GABAergic neurotransmission (Harvey, 2015). On another hand, it 361 362 has been reported that chrysin increases testosterone levels (Ciftc et al., 2011), which could be associated to its aromatase inhibitor activity (Campbell & Kurzer, 1993). In this 363 sense, Fernández-Guasti et al. (2005) demonstrated that testosterone administration 364 exerts an anxiolytic-like effect in male Wistar rats due to the GABA_A-benzodiazepine 365 receptor modulation by testosterone. This suggest that the flavonoid chrysin can exert 366 more than one effect in the GABA_A/benzodiazepine site complex. 367

368 In the other hand, chrysin did not alter risk assessment behavior in the present experiments, a variable that has been shown to be sensitive to non-benzodiazepine 369 treatments as well as to benzodiazepines (Maximino et al., 2014; Lima-Maximino et al., 370 371 2018). In a multivariate approach, risk assessment has been shown to co-vary with other variables in the LDT, forming a separate cluster from thigmotaxis and time on 372 white (Maximino et al., 2014); thus, risk assessment is sensitive to anxiolytic treatments, 373 but represents a different dimension or factor in the multivariate structure of zebrafish 374 behavior in the LDT. 375

376 Surprisingly, the flavone backbone decreased risk assessment in the LDT and SAP in the rat EPM, and increased rearing in the locomotor activity test. These outcomes are 377 378 paradoxical; effects on rearing are difficult to interpret, but it has been suggested (Lever et al., 2006) that rearing is associated with environmental novelty, with a possible 379 380 cognitive function in information-gathering for escape behavior. However, risk 381 assessment in the LDT and SAP in the EPM can be interpreted as having an information-gathering function as well, suggesting different mechanisms for rearing, on 382 the one hand, and risk assessment behavior in anxiety tests, on the other. Taken 383 384 together, these results suggest that, although not able to reduce the avoidance of potentially adverse environments such as the open arm of the EPM or the light 385 compartment of the LDT tank, the flavone backbone is able to reduce risk assessment 386 387 behavior. Risk assessment has been suggested as a cognitive component of anxietylike behavior (Blanchard et al., 2011), and is more sensitive than spatio-temporal 388 389 measures (time spent on open arms) in the EPM to detect effects of serotonergic anxiolytics (Griebel et al., 1997). 390

Finally, the present study compared the effect of a single dose of chrysin (1 mg/kg) *versus* the flavone backbone. We have previously reported other doses (2 and 4 mg/kg) of this flavonoid with anxiolytic-like effect in the EPM (Rodríguez-Landa et al., 2019). However, that study used female Wistar rats with long-term ovariectomy, while in this work male Wistar rats were used. In other work, we demonstrated that, in male Wistar rats, only 1 mg/kg, but no 0.5 neither 2 mg/kg, of chrysin exerted an anxiolytic-like effect in the EPM in male Wistar rats (German-Ponciano et al., 2020). Thus, we used a single dose of chrysin in agreement with the ethical recommendations of the 3R's principles in
order to reduce the number of animals used in this study (Russell and Burch, 1959).
Nevertheless, future work will be required in order to compare the possible doseresponse curves of chrysin *versus* flavone backbone.

402 Overall the results of the present study show that, in zebrafish and rats, the anxiolytic-403 like effect of the flavonoid chrysin might be due to more than one effect in the 404 GABA_A/benzodiazepine receptor complex including its antioxidant effect, which could be 405 related with the presence of the hydroxyl groups in its basic structure. However, 406 additional work is required to determine the molecular mechanism through this 407 nutraceutical compound and other flavonoids exert their potential therapeutic to 408 ameliorate anxiety symptoms at preclinical research.

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635 **Figure captions**

Figure 1 – Chemical structures of flavone and chrysin.

Figure 2 – Effects of chrysin, flavone, and diazepam on the elevated plus-maze. A. Scheme of the apparatus. B. Entries on the open arms. C. Entries on the closed arms. D. Entries on all arms. E. Percent entries on the open arms. F. Time spent on the open arms. G. Time spent in headdipping. H. Number of head-dipping events. I. Time spent in stretched-attend posture (SAP). J. Frequency of SAP events. For variables with the same letter, the difference is not statistically significant. Likewise, for variables with a different letter, the difference is statistically significant ($p \le 0.05$).

Figure 3 – Effects of chrysin, flavone, and diazepam on the locomotor activity test. A. Scheme of the apparatus and tested molecules. B. Squares crossed. C. Time spent in grooming. D. Time spent in rearing. For variables with the same letter, the difference is not statistically significant. Likewise, for variables with a different letter, the difference is statistically significant ($p \le 0.05$).

Figure 4 – Effects of chrysin, flavone, and diazepam on the light/dark test. A. Scheme of the apparatus and tested molecules. B. Time spent on the white compartment. C. Duration of entries on the white compartment. D. Transitions to white compartment. E. Frequency of risk assessment events. F. Frequency of erratic swimming events. G. Time spent in thigmotaxis. H. Duration of freezing events. For variables with the same letter, the difference is not statistically significant. Likewise, for variables with a different letter, the difference is statistically significant ($p \le 0.05$).

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for

"Chrysin, but not flavone backbone, decreases anxiety-like behavior in animal screens"

León Jesús German-Ponciano: Conceptualization, Methodology, Investigation, Formal analysis, Writing – Original draft, Writing - review & editing; Bruna Patricia Dutra Costa: Investigation, Writing – Original draft; Leonardo Miranda Feitosa: Investigation, Data curation; Kimberly dos Santos Campos: Investigation, Data curation; Suianny Nayara da Silva Chaves: Investigation, Data curation; Jonathan Cueto-Escobedo: Conceptualization, Methodology, Investigation, Writing – Original draft; Monica Lima-Maximino: Conceptualization, Validation; Juan Francisco Rodríguez-Landa: Conceptualization, Methodology, Writing – Original draft; Caio Maximino: Conceptualization, Methodology, Validation, Data Curation, Writing – Original draft, Supervision, Project administration

Highlights

- Chrysin produced anxiolytic-like effects in zebrafish and rats
- The flavone backbone decreased risk assessment behavior
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