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OPEN Neurobehavioral effects of fungicides in zebrafish: a systematic review and meta-analysis

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Pesticides are widely used in global agriculture to achieve high productivity levels. Among them, fungicides are specifically designed to inhibit fungal growth in crops and seeds. However, their application often results in environmental contamination, as these chemicals can persistently be detected in surface waters. This poses a potential threat to non-target organisms, including humans, that inhabit the affected ecosystems. In toxicologic research, the zebrafish (Danio rerio) is the most commonly used fish species to assess the potential effects of fungicide exposure, and numerous and sometimes conflicting findings have been reported. To address this, we conducted a systematic review and meta-analysis focusing on the neurobehavioral effects of fungicides in zebrafish. Our search encompassed three databases (PubMed, Scopus, and Web of Science), and the screening process followed predefined inclusion/exclusion criteria. We extracted qualitative and quantitative data, as well as assessed reporting quality, from 60 included studies. Meta-analyses were performed for the outcomes of distance traveled in larvae and adults and spontaneous movements in embryos. The results revealed a significant overall effect of fungicide exposure on distance, with a lower distance traveled in the exposed versus control group. No significant effect was observed for spontaneous movements. The overall heterogeneity was high for distance and moderate for spontaneous movements. The poor reporting practices in the field hindered a critical evaluation of the studies. Nevertheless, a sensitivity analysis did not identify any studies skewing the meta-analyses. This review underscores the necessity for better-designed and reported experiments in this field.

Chemical pesticides are synthetical active ingredients used to control pests that may threaten the productivity of crops¹. To yield high productivity levels, modern agriculture employs large amounts of pesticides². In 2020, the global consumption of these products reached almost 3 million tonnes³. The substantial quantity and the method by which they are applied results in environmental contamination of the soil, surface waters, and food⁴⁻⁶. Data shows that less than 0.1% of the pesticide hits the intended target species, leaving the remaining residual impacting the environment and public health⁷. Its presence in superficial waters generates risk to the non-target organisms by decreasing biodiversity and the population of primary food chain producers and reducing the prey for the aquatic organisms⁸⁻¹⁰. Moreover, the dissemination of pesticides in the environment represents a risk to humans, whereas their presence in the water supply leads to potential consumption¹¹⁻¹⁴.

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According to the target organism, these substances can be classified as herbicides, insecticides, rodenticides, and fungicides¹⁵, being the fungicides one of the most used chemicals¹⁶. Their application aims to kill and/or inhibit fungal growth in agriculture, both in seeds and crops¹⁷.

Due to the need to understand the effects of exposure to these products, the scientific literature presents several studies with animals in this area¹⁸. The model organism zebrafish (*Danio rerio*, Hamilton 1822) is widely used in toxicology, mostly because of its high fecundity, fast development, transparency of the embryo, and high homology of organs and genetics concerning humans¹⁹⁻²¹. In addition, the zebrafish is an aquatic animal that dwells in potentially contaminated ecosystems, representing the eventual consequences of exposure to other cohabitant species²². It has been reported that exposure to fungicides in zebrafish causes behavioral, neurochemical, developmental, metabolic, hormonal, hepatotoxic, cardiotoxic, enzymatic, morphological, and molecular alterations²³⁻²⁸.

From 2012 to 2019, more than 100 articles were published investigating the effects of fungicides in zebrafish, which represents the second most investigated type of pesticide in this organism²⁹. However, there is a high methodological heterogeneity between the studies. The interventions, developmental stages, and outcomes addressed are extremely variable between studies. Regarding the intervention, plenty of compounds used as fungicides exhibit distinct mechanisms of action³⁰ and can be administered over a wide range of durations through multiple routes of administration. As for the developmental stage, in vivo exposure can be performed in embryos, larvae, or adults; the outcomes are distinctly selected according to the research question and the capabilities of the research group (neurotoxicity, hepatotoxicity, cardiotoxicity, among others)³¹.

Many studies were published on the toxic effects of fungicides on neurobehavioral parameters in zebrafish^{22,32}. However, no secondary studies systematically synthesize these results to obtain an understanding supported by published evidence to optimize the planning of new research. An accurate description of these preclinical data and a meta-analysis can help avoid redundant studies and the consequent use of animals. Furthermore, considering the reproducibility issues raised for the zebrafish research field^{33,34}, it is essential to identify possible sources of bias and conflicting results, including assessing the quality of available publications. This systematic review and meta-analysis of literature aimed at synthesizing the data from neurobehavioral effects of fungicide exposure in zebrafish, also analyzing reporting quality and publication bias.

Methods

Before screening studies and data extraction, a protocol guiding this review was registered in Open Science Framework, and preregistration is available at https://osf.io/f2d38³⁵. The reporting of this study follows the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines³⁶.

Search strategy

The studies were identified through a search in the literature using three different databases: PubMed, Scopus, and Web of Science. The search strategies were designed to adapt to each database characteristics. The terms were combined for the intervention (fungicide exposure) and the population of interest (zebrafish), aiming to conduct a comprehensive search, including all the available articles that fulfilled the inclusion criteria. The complete query for each database can be found at https://osf.io/5ae9q³⁵. The strategy did not apply any search filter, language restriction, or limit of year. The search was performed on the 1st of December, 2021, and the articles were imported to Rayyan software³⁷ to identify and remove the duplicates.

Study screening

Initially, the retrieved studies from the three databases were analyzed to filter and exclude duplicates (performed by CGR). The remaining articles were pre-selected based on their title and abstract. If a reason to exclude the record was not found, at this stage, it was carried forward to the full-text screening stage. In both stages (title/ abstract and full-text), two independent reviewers (CGR and LMB, RC or SZB) examined each study. Disagreements between the decisions of the reviewers were resolved by a third reviewer (QKZ, AP, or APH).

Experimental studies evaluating the effects of exposure to fungicides in zebrafish on the following parameters were included: motor function, sensory function, learning and memory, social behavior, sexual behavior, eating behavior, anxiety-like or fear-related behaviors, behaviors related to the reward system, and behaviors related to circadian rhythms. The parameters were included only if they were linked to the central nervous system. The identity of the compound as a fungicide was consulted in the Pesticide Properties Database³⁸.

In the first phase (screening of title/abstract), papers were excluded according to the following criteria:

- 1. Type of study design: reviews, comments, abstracts published in conference proceedings, corrections, editorials;
- 2. Type of population: in vitro investigations or studies with species other than zebrafish;
- 3. Type of intervention: biological and commercial formulations or mixtures of fungicides, non-interventional studies;

In the next phase (full-text screening), the following criteria were added, and the articles were excluded based on the above items plus:

- 4. Comparison: when there is no proper control group (same organism, same procedure, except for fungicide exposure);
- 5. Outcome measures: if there is no assessment of any previously cited neurobehavioral outcome.

More information about this section is available at https://osf.io/wmsvg.

Data extraction

Two independent investigators (CGR and LMB, RC or SZB) performed the data extraction, and a discussion between the two reviewers resolved disagreements. The information and values of interest were directly extracted from the text and tables. When not possible, WebPlotDigitizer software (v4.5, Rohatgi, A., Pacifica, CA, USA, https://automeris.io/WebPlotDigitizer) was used to determine the values from the graphs manually. The following data were extracted: (1) study identification: study title, digital object identifier (DOI), first author, last author, year of publication, and last author affiliation; (2) model animal specifications: strain, sex, the developmental stage during exposure, age during exposure, the developmental stage during the test, age during the test; (3) fungicide exposure characteristics: fungicide, administration route and type (i.e., static, semi-static or flow through), frequency of renewal, frequency of exposure, duration of exposure, dose/concentration and the interval between exposure and test; (4) test properties: test nomenclature, category of measured variable (e.g., anxiety, locomotor, social) and the measured variable.

Regarding the authors of the studies, co-authorship networks were elaborated using VOSviewer software version 1.6.18 (https://www.vosviewer.com)^{39,40}.

Data were collected for each variable according to the outcomes of interest, including the mean and the number of animals (n) for both the control and exposed groups. The standard deviation (SD) or standard error of the mean (SEM) was extracted for the reported mean value. If the SEM was reported, the SD was calculated by multiplying SEM by the square root of the sample size (SD = SEM * \sqrt{n}).

In instances where the sample size was reported as a range, the lowest value was used. Whenever information was unclear or missing, attempts were made to contact the corresponding author of the study via email, with two separate attempts made at least two weeks apart.

Reporting quality

To assess the reporting quality of included studies, two independent reviewers (CGR and LMB, RC, or SZB) evaluated each paper based on⁴¹, which proposes criteria for transparent reporting. The observed topics were: (1) mention of any randomization process; (2) sample size estimation; (3) mention of inclusion/exclusion criteria; (4) mention of any process to ensure blinding during the experiments. A score of "yes" or "no" was given for each topic, meaning that it was or was not reported, respectively. The outcome measurements performed by any automated software were considered blinded. Reporting quality plots were created using robvis⁴².

A complete guide for assessing the reporting quality associated with each item in this review is available at https://osf.io/uy5v3.

Meta-analysis

To perform a meta-analysis, at least 5 studies with the same outcome were required a priori³⁵. Whenever two or more experimental groups shared the same control, the sample size of the control group was divided by the number of comparisons and then rounded down. Further information about the basic aspects of our method can be found at⁴³.

Effects sizes were determined with the standardized mean difference (SMD) using Hedge's G method⁴⁴. SMD was used because studies examined a common outcome while employing different measurement approaches, which makes it necessary to standardize the findings in a uniform scale to allow combination across studies. Briefly, SMD expresses the size of an intervention effect relative to the observed variability^{45,46}. Analyses were conducted using R Project for Statistical Computing with packages meta⁴⁷ (https://cran.r-project.org/package= meta) and ggplot2⁴⁸ following Hedge's random effects model, given the anticipated heterogeneity between studies. Values for SMD were reported with 95% confidence intervals. Heterogeneity between studies was estimated using I²⁴⁹, τ^2 , and Cochran's Q⁵⁰ tests. Heterogeneity variance (τ^2) was estimated using the restricted maximum likelihood estimator^{51,52}. The confidence intervals around pooled effects were corrected using Knapp-Hartung adjustments⁵³. Values of 25%, 50%, and 75% were considered as representing low, moderate, and high heterogeneity, respectively, for I², and a p-value ≤ 0.1 was considered significant for Cochran's Q. Prediction intervals were estimated and represent the range of effects expected for future studies⁴⁵. Furthermore, a subgroup meta-analysis was performed to evaluate if the developmental stage of the animals was a potential source of heterogeneity. Studies were grouped into two categories: larval and adult. Subgroup analysis was only performed when there were at least five unique studies for each subgroup. A p-value ≤ 0.1 was considered significant for subgroup differences⁵⁴.

We conducted an exploratory meta-analysis to investigate an association between the effect and the fungicide class by categorizing them based on their chemical structure. Even without reaching the minimum of 5 studies, we ran a meta-analysis with 4 articles investigating fungicides of the triazole and anilide groups.

A mixed-effects meta-regression analysis was conducted to explore the relationship between the effect sizes and fungicide concentration as a moderator variable. The random effects structure accounted for potential heterogeneity across studies⁵⁵. Meta-regressions excluding studies based on the sensitivity analysis were also performed.

Publication bias was investigated by generating funnel plots and performing Duval and Tweedie's trim and fill analysis⁵⁶ and Egger's regression test⁵⁷. Analyses were only conducted when at least five studies were available within a given outcome for funnel plots and at least ten studies for the regression test. A p-value < 0.1 was considered significant for the regression test.

Sensitivity analysis

A sensitivity analysis was conducted to assess if any experimental or methodological difference between studies was biasing the main effect found in the meta-analysis. Analyses were performed following the "leave-one-out

jackknife method"⁵⁸. A minimum of three comparisons were required for each outcome to conduct a sensitivity analysis. Furthermore, we conducted complementary meta-analyzes excluding studies that, when omitted in the leave-one-out, had observations that changed the overall effect direction. We also ran meta-analyses excluding studies containing experiments with atypically high SMD, as seen in the forest plots⁵⁹.

Results

Search results

The search in the three databases retrieved a total of 2139 results. After removing duplicates, 1140 articles were screened for eligibility by analyzing the titles and abstracts. As a result of the first screening phase, 369 studies remained to be assessed based on their full-text. At this phase, 3 were not retrieved, and 60 fulfilled the criteria and were included in the review (Fig. 1). The main overall reasons for the exclusions were outcome (n = 234), population (n = 260), and intervention (n = 350). Concerning the quantitative synthesis, 8 studies were excluded because the minimum number of studies to perform a meta-analysis was not reached for the reported outcomes and 10 because of missing information. There were 18 experiments measuring distance using luminous transitions (dark/light) in larvae that were not included due to the variations between the protocols⁶⁰, which makes the comparison infeasible. This resulted in 24 studies included in the quantitative synthesis. Detailed reasons for excluding studies from the meta-analysis are available at https://osf.io/qpcew.

Study characteristics

A qualitative description of the studies is provided in Table 1. The identification of the studies was attributed according to the table available at https://osf.io/85d2p. A total of 43 different fungicides were addressed in the articles included in this review. Studies with the fungicides difenoconazole (n = 5, 8.3%), boscalid (n = 4, 6.6%), and pyraclostrobin (n = 4, 6.6%) were the most frequent.

All the studies used immersion as the exposure method, whereas exposure durations ranged from 11 min to 217 days. The most recurrent duration of exposure among the publications was 24 h (n = 21, 35%), followed by 96 h (n = 12, 20%). It is important to emphasize that 24 h is usually employed to verify the outcome of spontaneous movements, while 96 h is recommended by the Organization for Economic Co-operation and



Figure 1. Flowchart diagram of the collection of studies and selection process.

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Study ID	Fungicide	Concentration (mg/L)	Duration of exposure	Developmental stage during exposure/outcome assessment	Main findings
Domingues et al. ⁸²	Prochloraz	0.3, 0.6, 1.2, 2.4, 4.8, 7.2, 9.6	96 h	Embryo/Larvae & Adult/Adult	Locomotor behavior ↑ Spontaneous movements ↓ Distance traveled ↑ Distance in the dark ↑ Distance in the light ↑ Velocity ↑ Acceleration ↑ Absolut turn angle Neurochemical outcomes ↓ ChE activity (larvae) ↑ GST activity (larvae)
Fitzmaurice et al. ⁸³	Benomyl	0.29	120 h	Embryo/Larvae	Locomotor behavior ↓ Distance traveled
Mu et al. ⁸⁴	Difenoconazole	0.5, 1,1.5, 2, 2.5, 3	24 h	Embryo/Embryo	Locomotor behavior ↑ Spontaneous movements (1.5 mg/L) ↓ Spontaneous movements (2.5, 3 mg/L) ↑ Reversal rate behavior
Andrade et al. ⁸⁵	Carbendazim	1.1, 1.19, 1.3, 1.41, 1.53, 1.66, 1.8	120 h	Embryo/Larvae	Locomotor behavior ↓ Distance in the light ↓ % Small distance in the light ↓ % Small distance in the dark ↑ % Long distance in the light ↑ % Long distance in the light ↑ Swimming time in the light ↑ Swimming time in the dark Neurochemical outcomes ↑ ChE activity ↑ GST activity = CAT activity
Jin et al. ⁸⁶	Imazalil	0.01, 0.03, 0.1, 0.3	96 h	Embryo/Larvae	Locomotor behavior ↓ Distance traveled ↓ Distance in the dark ↓ Distance in the light ↓ VelocityNeurochemical outcomes↓ AChE levels ↓ AChE activity = DA levels
Lulla et al. ⁸⁷	Ziram	0.0003—0.305	7 days	Embryo/Larvae	Locomotor behavior ↓ Distance in the dark = Distance in the light ↓ Velocity
Mu et al. ⁸⁸	Difenoconazole	0.5, 2	96 h	Embryo/Embryo	Locomotor behavior = Spontaneous movements
Yang et al. ⁶⁸	Thifluzamide	2.66, 2.76, 2.85, 2.95, 3.04, 3.23 & 2.66, 2.76, 2.85, 2.95, 3.04 & 2.66, 2.85, 3.04, 3.23, 3.42, 3.61	96 h & 144 h & 96 h	Embryo/Embryo & Embryo/Larvae & Larvae/Larvae	Locomotor behavior ↓ Spontaneous movements (embryo) ↓ Swimming rate (larvae)
Yang et al. ⁸⁹	Flutolanil	1.5, 1.8, 2.16, 2.59, 3.1	24 h	Embryo/Embryo	Locomotor behavior ↑ Spontaneous movements
Altenhofen et al. ⁹⁰	Tebuconazole	1, 2, 4 & 1, 4, 6	120 h & 96 h	Embryo/Larvae & Adult/Adult	Locomotor behavior ↓ Distance traveled ↓ Absolut turn angle (larvae) = Crossings Anxiety/fear-related behavior ↓ Time in the periphery = Time in the upper zone Aggressive behavior ↑ Time in the bottomNeuro- chemical outcomes↓ AChE activity
De la Paz et al. ⁹¹	Triadimefon	16	8 h	Larvae/Larvae	Locomotor behavior ↑ Locomotor activity
Costa-Silva et al. ⁹²	Mancozeb	1	23 h & 43 h	Embryo/Embryo & Embryo/Embryo dechorionated	Locomotor behavior ↑ Spontaneous movements ↑ Number of stimuli (embryo dechorionated) ↑ Response to touch (embryo dechorionated) Neurochemical outcomes = GST activity ↓ GSH levels = GPx activity
Fan et al. ⁹³	Hymexazol	417, 480, 554, 639, 738	48 h	Embryo/Larvae	Locomotor behavior ↓ Swimming rate
Li et al. ⁹⁴	Pyraoxystrobin	2.03, 2.44, 2.9, 3.51, 4.22, 5.08	24 h	Embryo/Embryo	Locomotor behavior = Spontaneous movements
Continued					

Study ID	Fungicide	Concentration (mg/L)	Duration of exposure	Developmental stage during exposure/outcome assessment	Main findings
Qian et al. ⁹⁵	Boscalid	0.7, 2, 2.3, 2.6, 2.9, 3.2	22 h	Embryo/Embryo	Locomotor behavior ↑ Spontaneous movements
Teng et al. ⁹⁶	Difenoconazole	0.0005, 0.005, 0.05, 0.5	24 h	Embryo/Embryo	Locomotor behavior ↑ Spontaneous movements
Teng et al. ⁹⁷	Difenoconazole	0.0005, 0.005, 0.05, 0.5	24 h	Embryo/Embryo	Locomotor behavior ↑ Spontaneous movements
Wang et al. ⁹⁸	Fluazinam	0.04, 0.09, 0.13	6 days	Embryo/Larvae	Locomotor behavior ↑ Swimming activity in the dark (0.04 mg/L) ↓ Swimming activity in the dark (0.09, 0.13 mg/L) = Swimming activity in the light
Cao et al. ⁶⁵	Ziram	0.0003, 0.003	7 days	Embryo/Larvae	Locomotor behavior ↑ Swimming activity ↑ Distance traveled (0.003 mg/L) = Distance in the dark = Distance in the light = Total velocity ↑ Velocity in light Anxiety/fear-related behavior ↓ Time in the dark = Frequency in the dark
			24 h &	Embryo/Embryo	Locomotor behavior
Cao et al. ⁹⁹	Cyproconazole	2.9, 7.2, 14.5, 29.1, 72.9, 145.8	48 h & 7 days	& Embryo/Larvae	↓ Swimming activity in the dark = Swimming activity in the light
Cao et al. ¹⁰⁰	Maneb	0.02, 0.13, 0.26	7 days	Embryo/Larvae	Locomotor behavior ↓ Swimming activity in the dark ↓ Swimming activity in the light
Li et al. ⁶⁷	Pyraclostrobin	0.009, 0.018, 0.36	4 days	Larvae/Larvae	Locomotor behavior ↓ Distance traveled ↓ VelocityNeurochemical outcomes↑ Glutamate receptor activity
Paredes-Zúñiga et al. ⁶³	Triadimefon	5, 20, 35	10 h & 11 min	Larvae/Larvae & Adult/Adult	Locomotor behavior ↓ Swimming activity ↓ Distance traveled (larvae) ↑ Distance traveled (adult) ↑ Velocity Anxiety/fear-related behavior ↓ Time in the periphery ↑ Time in the bottom zone ↓ Time in the upper zone Aggressive behavior ↑ Number of bitesNeurochemi- cal outcomes↑ DA levels ↓ 5-HT levels
Perez-Rodriguez et al. ¹⁰¹	Tebuconazole	0.03, 0.3, 3	6 days	Embryo/Larvae	Locomotor behavior ↓ Distance in the dark = Distance in the light = Velocity Anxiety/fear-related behavior ↑ Mean time in the dark = Cumulative time in the dark ↑ Frequency in the dark zone
Qian et al. ¹⁰²	Penthiopyrad	2.3, 2.4, 2.5, 2.6, 2.7, 2.8, 2.9 & 0.3, 0.6, 1.2	1 day & 5–8 days	Embryo/Embryo & Embryo/Larvae	Locomotor behavior ↑ Spontaneous movements (2.5, 2.6, 2.7 mg/L) ↓ Spontaneous movements (2.9 mg/L) ↓ Swimming activity ↓ Distance traveled ↓ Velocity ↓ Acceleration
Souders et al. ¹⁰³	Propiconazole	0.03, 0.3, 3.4	144 h	Embryo/Larvae	Locomotor behavior ↓ Distance traveled ↓ Distance in the dark
Teng et al. ¹⁰⁴	Propiconazole	0.5, 2.5, 4.5	24 h & 120 h	Embryo/Embryo & Embryo/Larvae	Locomotor behavior ↑ Spontaneous movements ↓ Distance traveled ↓ Velocity ↓ Swimming activity ↓ Acceleration
Continued					

Study ID	Fungicide	Concentration (mg/L)	Duration of exposure	Developmental stage during exposure/outcome assessment	Main findings
Tian et al. ¹⁰⁵	Prothioconazole	0.0375, 0.075, 0.15	24 h	Embryo/Embryo	Locomotor behavior = Spontaneous movements Neurochemical outcomes ↓ GSH levels = SOD activity = CAT activity ↑ MDA levels
Valadas et al. ¹⁰⁶	Propiconazole	0.000425, 0.00085, 0.0017, 0.0085	96 h	Adult/Adult	Locomotor behavior = Distance traveled ↓ Crossings Anxiety/fear-related behavior ↑ Time in the upper zone ↑ Time in the upper zone ↓ Entries in the upper zone = Entries in the bottom zone Neurochemical outcomes ↑ SOD activity ↑ CAT activity = MDA levels = SH levels = NPSH levels
Wang et al. ¹⁰⁷	Oxine-copper	0.01, 0.02, 0.04	24 h	Embryo/Embryo & Larvae/Larvae	Locomotor behavior ↓ Number of tail coiling ↓ Distance traveled ↓ Swimming activity ↓ Velocity ↑ Absolut turn angleNeuro- chemical outcomes↓ AChE activity(embryo) ↑ SOD activity (embryo) ↑ CAT activity (embryo) ↑ MDA levels (embryo) ↑ ROS levels (embryo)
Yang ¹⁰⁸	Flutolanil	0.125, 0.5, 2	24 h & 96 h	Embryo/Embryo & Embryo/Larvae	Locomotor behavior ↓ Spontaneous movements = Distance traveled Neurochemical outcomes ↑ DA levels
Yang ¹⁰⁹	Thifluzamide	0.19, 1.9, 2.85	24 h & 96 h	Embryo/Embryo & Embryo/Larvae	Locomotor behavior = Spontaneous movements = Distance traveled Neurochemical outcomes ↓ DA levels
Zhou et al. ¹¹⁰	Captan	0.58, 0.66, 0.75, 0.86, 1.00, 1.16	24 h	Embryo/Embryo	Locomotor behavior = Spontaneous movements
Hussain et al. ¹¹¹	Tebuconazole & Dimethomorph & Difenoconazole	0.3 & 0.3 & 0.4	24 h	Larvae/Larvae	Locomotor behavior ↑ Distance in the dark (tebucona- zole, dimethomorph) ↓ Distance in the dark (difeno- conazole) ↑ Distance in the light ↑ Burst movement count in the light ↑ Burst movement count in the dark (dimethomorph, difeno- conazole) ↑ Rotation count in the light (dimethomorph, difenoconazole) ↑ Rotation count in the dark (dimethomorph, difenoconazole)
Jia et al. ²³	Penconazole (+) & Penconazole (-)	1, 2	24 h & 96 h	Embryo/Embryo & Embryo/Larvae	Locomotor behavior ↑ Spontaneous movements ((+)-penconazole) ↓ Velocity ((+)-penconazole) Neurochemical outcomes ↓ AChE activity ((+)-pencona- zole) ↓ DA levels ((+)-penconazole) ↓ 5-HT levels ((+)-penconazole) = Glycine levels = Norepinephrine levels
Kumar et al. ¹¹²	Azoxystrobin & Pyraclostrobin	0.00001, 0.0001, 0.01, 0.1, 1	5 days	Embryo/Larvae	Locomotor behavior ↓ Distance traveled Neurochemical outcomes = MDA levels
Continued					

Study ID	Fungicide	Concentration (mg/L)	Duration of exposure	Developmental stage during exposure/outcome assessment	Main findings
Liu et al. ¹¹³	Propamocarb	0.01, 0.1, 1	7 days	Embryo/Larvae	Locomotor behavior ↑ Distance traveled ↑ Distance in the dark = Distance in the light ↑ Velocity Neurochemical outcomes ↓ AChE activity = MDA levels ↓ DA levels = SOD activity ↑ CAT activity ↓ GST activity
Pang et al. ¹¹⁴	Myclobutanil	4, 6, 8, 10, 12, 14, 16	24 h	Embryo/Embryo	Locomotor behavior ↑ Spontaneous movements (4, 6, 8, 10, 12 mg/L) ↓ Spontaneous movements (16 mg/L)
Shen et al. ²⁶	Mepanipyrim	0.0001, 0.001, 0.01, 0.1	7 days	Embryo/Larvae	Locomotor behavior ↑ Distance traveled (7, 14 dpf) ↓ Distance traveled (14 dpf) ↑ Velocity (7, 14 dpf) ↑ Velocity (14 dpf) ↑ Acceleration = Absolut turn angle ↓ Immobile time (7 dpf) Neurochemical outcomes = AChE activity ↑ GABA levels
Souders et al. ¹¹⁵	Triticonazole	0.3, 3.1, 31.7	6 days	Embryo/Larvae	Locomotor behavior ↑ Distance in the dark = Distance in the light
Tang et al. ⁶⁶	Cyprodinil	0.0001, 0.001, 0.01, 0.1	24 h	Embryo/Embryo	Locomotor behavior ↓ Spontaneous movements
Teng et al. ²⁷	Flutolanil	0.00025, 0.05, 1	60 days	Adult/Embryo (offspring)	Locomotor behavior = Spontaneous movements
Vasamsetti ¹¹⁶	Etridiazole	3.75, 7.5, 15, 30, 60	96 h	Embryo/Larvae	Locomotor behavior ↑ Immobile time
Wang et al. ²⁸	Boscalid	5, 15, 25	24 h	Embryo/Embryo & Larvae/Larvae	Locomotor behavior ↓ Number of tail coiling ↓ Distance traveled ↓ Distance in the dark ↓ Distance in the light ↓ Velocity ↑ Absolut turn angle ↑ Immobile timeNeurochemical outcomes = AChE activity ↑ MDA levels ↓ SOD activity ↑ CAT activity ↑ ROS levels
Zhang et al. ¹¹⁷	Zoxamide	0.16, 0.33, 0.84, 1.68	24 h & 6 days	Embryo/Larvae & Embryo/Larvae	Locomotor behavior ↑ Distance in the dark (24 h, 6 days exposure) ↓ Distance in the dark (6 days exposure) ↑ Distance in the light (6 days exposure)
Barreto et al. ¹¹⁸	Fosetyl-al	0.02, 0.2, 2, 20, 200	120 h	Embryo/Larvae	Locomotor behavior ↓ Distance traveled = Swimming time ↑ Velocity ↑ Acceleration ↑ Absolut turn angleNeurochem- ical outcomes = ChE activity = CAT activity ↑ GST activity
Brenet et al. ¹¹⁹	Bixafen	0.08, 0.2	96 h	Embryo/Larvae	Locomotor behavior ↓ Distance traveled
Fan et al. ¹²⁰	Carbendazim	0.52, 0.65, 0.82, 1.02, 1.28, 1.6	24 h	Embryo/Embryo	Locomotor behavior ↑ Spontaneous movements
Forner-Piquer et al. ⁶⁴ Continued	Boscalid & Captan & Thiophanate & Ziram	0.00001, 0.00005, 0.001, 0.01, 0.1, 1, 10	120 h	Embryo/Larvae	Locomotor behavior ↓ Distance traveled ↓ Velocity

Study ID	Fungicide	Concentration (mg/L)	Duration of exposure	Developmental stage during exposure/outcome assessment	Main findings
Huang et al. ¹²¹	Fenamidone	0.03, 0.3, 0.4, 0.6	144 h	Embryo/Larvae	Locomotor behavior = Distance traveled (light-dark test) ↓ Distance in the dark (visual motor response test) = Distance in the light Anxiety/fear-related behavior = Frequency in the dark = Time in the dark
Leandro et al. ¹²²	Mancozeb	0.005, 0.01, 0.02	24 h & 68 h & 164 h	Embryo/Embryo & Embryo/Larvae	Locomotor behavior ↓ Spontaneous movements (28 hpf) ↑ Number of stimuli (72 hpf) ↓ Response to touch (72 hpf) ↑ Distance traveled (0.005 mg/L, 168 hpf) ↑ Absolut turn angle (0.005 mg/L, 168 hpf) ↑ Absolut turn angle (0.002 mg/L, 168 hpf) ↑ Immobile episodes (0.02 mg/L, 168 hpf) ↑ Immobile episodes (0.02 mg/L, 168 hpf) ↑ Immobile time (0.02 mg/L, 168 hpf) ↓ Immobile time (0.02 mg/L, 168 hpf) ↓ Immobile time (0.005 mg/L, 168 hpf) ↓ Immobile time (0.005 mg/L, 168 hpf) ↓ Immobile time (0.005 mg/L, 168 hpf) ↑ Entries in the periphery (168 hpf) ↑ Entries in the periphery (168 hpf) ↓ SOD activity (24 hpf) ↓ SOD activity (24 hpf) ↓ SOD activity (24 hpf) ↓ SOD activity (72 hpf) ↓ CAT activity (168 hpf) ↑ GST activity (168 hpf) ↑ GSS activity (72, 168 hpf) ↑ ROS levels (72, 168 hpf)
Li et al. ¹²³	Azoxystrobin & Kresoxim-methyl & Pyraclostrobin & Trifloxystrobin	0.02027 & 0.01567 & 0.01939 & 0.02042	5 days	Embryo/Larvae	Locomotor behavior ↑ Distance in the dark (kresoxim- methyl, pyraclostrobin, triflox- ystrobin) ↑ Distance in the light (trifloxystrobin)Neurochemical outcomes↑ MDA levels (pyra- clostrobin, trifloxystrobin) ↑ SOD activity (pyraclostrobin, trifloxystrobin) ↑ CAT activity (trifloxystrobin) ↑ ROS levels (pyraclostrobin, trifloxystrobin)
Lin et al. ¹²⁴	Fluxapyroxad	1.1, 1.2, 1.3, 1.4, 1.5, 1.6	24 h	Embryo/Embryo	Locomotor behavior ↑ Spontaneous movements Neurochemical outcomes ↑ MDA levels = SOD activity = CAT activity ↑ GPx activity (0.174 mg/L) ↓ GPx activity (0.694 mg/L)
Paredes-Zúñiga et al. ¹²⁵	Triadimefon	5, 15	3 days	Adult/Adult	Locomotor behavior ↑ Time in the drug-paired zone (5 mg/L) ↓ Time in the drug-paired zone (15 mg/L) ↑ Circling behavior (days 1, 2)
Pompermaier et al. ¹²⁶	Copper	0.105	48 h	Adult/Adult	Locomotor behavior = Distance traveled = Absolut turn angle = Crossings Anxiety/fear-related behavior = Time in the upper zone = Time in the middle zone = Time in the bottom zone

Study ID	Fungicide	Concentration (mg/L)	Duration of exposure	Developmental stage during exposure/outcome assessment	Main findings
Qian et al. ¹²⁷	Boscalid	0.3, 0.6, 1.2 & 0.01, 0.1, 1.0	8 days & 21 days	Embryo/Larvae & Adult/Adult	Locomotor behavior ↓ Distance traveled (larvae) ↑ Distance traveled (adult) ↓ Distance in the dark (larvae) ↓ Distance in the light (larvae) ↓ Velocity ↓ Acceleration ↓ Active time (larvae) ↑ Active time (larvae, adult) Neurochemical outcomes ↑ AChE levels (larvae) ↓ AChE activity (larvae)
Tang et al. ¹²⁸	Cyprodinil	0.0001, 0.001, 0.01	209-211 days & 215-217 days	Embryo/Adult	Locomotor behavior ↓ Distance traveled ↓ Velocity = Acceleration ↓ Absolut turn angle Aggressive behavior ↑ Time in the interaction zone
Wu et al. ¹²⁹	Procymidone	0.001, 0.01, 0.1	4 days & 7 days	Embryo/Larvae	Locomotor behavior ↑ Distance in the dark (4 days) ↑ Distance in the light (4 days) ↓ Distance in the dark (7 days) ↓ Distance in the light (7 days)
Yang ¹³⁰	Azoxystrobin & Pyraclostrobin & Trifloxystrobin	0.0002, 0.001, 0.005 & 0.77, 1.54, 2.32 & 0.51, 1, 2	6 days	Embryo/Larvae	Locomotor behavior ↑ Distance in the dark (azox- ystrobin, pyraclostrobin) ↓ Distance in the dark (triflox- ystrobin) = Distance in the light
Yang ¹³¹	Thifluzamide	0.19, 1.9, 2.85	96 h & 144 h	Embryo/Larvae & Embryo/Larvae	Locomotor behavior ↓ Distance traveled ↓ Velocity ↓ Swimming activity ↓ Rotating frequency (144 hpf) Neurochemical outcomes ↓ AChE activity ↑ 5-HT levels ↑ Norepinephrine levels

Table 1. Qualitative description of studies reporting effects of fungicide exposure on neurobehavioral and neurochemical outcomes in zebrafish. The indicated concentrations of exposure were used to assess the behavioral outcomes. The main findings were described as: ↑, higher when compared to the control group; ↓, lower when compared to the control group; =, no difference when compared to the control group. AChE = acetylcholinesterase, CAT = catalase, ChE = cholinesterase, DA = dopamine, GABA = gamma-aminobutyric acid, GPx = glutathione peroxidase, GSH = glutathione, GST = glutathione S-transferase, MDA = malondialdehyde, NPSH = non-protein thiols, ROS = reactive oxygen species, SH = thiols, SOD = superoxide dismutase, 5-HT = serotonin.

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Development (OECD) to assess acute fish toxicity in protocols 203 (adults) and 236 (embryos)^{61,62}. Regarding the developmental stage during the exposure, the embryonic was the most common (n = 52, 86.7%). Subsequently, the larval stage was observed in 7 studies (11.6%) and the adult stage in 8 (13.3%). Some articles used more than one stage for the exposure.

The outcome assessment was mostly performed in larvae (n = 41, 68.33%) and embryos (n = 25, 41.7%). Some studies assessed the outcomes in more than one developmental stage.

The sex of the adult animals was mainly reported as an equal proportion of male and female (F:M), except for one in which it was not reported (unclear).

Regarding the authors included in this review, co-authorship network analysis identified 24 clusters of researchers investigating the neurobehavioral effects of fungicides globally (Fig. S1). An interactive version of the co-authorship network is available at https://tinyurl.com/239thp6t.

Reporting quality

The summary plot of the reporting quality evaluation is shown in Fig. 2. Randomization process was not cited in 17 studies (28.33%). Only 3 articles described methods for sample size estimation (5%), and none of the authors explicitly stated the data inclusion or exclusion criteria. Blinding was reported in 38 papers (63.33%). Individualized scores for each study included are available at https://osf.io/pgrhq.

Meta-analysis

Distance

The meta-analysis included 61 comparisons from 12 independent studies. The total of animals used as controls was 1112, whereas the exposure individuals counted 2045. The highest concentration of fungicide in the



Figure 2. Reporting quality assessment of the included studies. The reporting quality assessment was performed by two independent investigators based on the criteria by⁴¹. Each item was scored as yes or no, meaning that the item is either reported or not, respectively. Classification is given as the percentage of assessed studies (n = 60) presenting each score.

meta-analysis was 20 mg/L for triadimefon⁶³, while the lowest was 0.0001 mg/L for mepanipyrim, ziram, tiophanate, captan, and boscalid^{26,64}.

The overall analysis showed that exposed animals present a lower distance traveled as compared to controls (SMD -0.44 [-0.74; -0.13], p=0.0055, Fig. 3). The estimated heterogeneity was considered high, with an I²=80%, a τ^2 =0.88, and a Q=300.1 (*df*=60, p<0.01). When calculating strictly for the developmental stage of the larvae, there was a significant effect of the fungicides on decreasing the distance traveled (SMD -0.44 [-0.83; -0.05], p=0.03, Fig. 3). The heterogeneity was still considered high for this subgroup, with an I²=84%, a τ^2 =1.21, and a Q=284.48 (p<0.01). Similarly, analyzing the adults subgroup, there was a significant effect of the exposure to fungicides on decreasing the distance traveled (SMD -0.55 [-0.89; -0.21], p<0.01, Fig. 3). Unlike the larvae, the heterogeneity was considered low, with an I²=5%, a τ^2 =0.07, and a Q=13.72 (p=0.39). The difference between subgroups was not significant (p=0.68), indicating that the developmental stage is not a direct moderator for this outcome.

The result from the meta-analysis of distance using only fungicides of the triazole group was similar, and a decrease in distance was observed (Fig. S2).

Spontaneous movements

The meta-analysis comprised 64 comparisons from 13 independent studies. The total of embryos used as controls was 190, and the exposure individuals counted 670. The highest fungicide concentration in the meta-analysis was 145.89 mg/L for cyproconazole⁶⁵, while the lowest was 0.0001 mg/L for cyprodinil⁶⁶. All the experiments performed the outcome assessment at 24 h of exposure, except for one (48 h).

The overall analysis showed that fungicide exposure had no significant effect on the number of spontaneous movements (SMD -0.16 [-0.67; 0.34], p = 0.5265, Fig. 4). The estimated heterogeneity was considered moderate, with an I² = 74%, a τ^2 = 1.86, and a Q = 243.19 (df = 63, p < 0.01).

The result from the meta-analysis of spontaneous movements using only fungicides of the anilide or triazole groups was similar, and no significant effects were observed (Fig. S3 and S4, respectively).

The meta-regression of both outcomes showed no significant correlation of the concentration with the effects (Fig. S5 and S6). Meta-regressions excluding studies from⁶⁷ (distance)⁶⁵, and⁶⁸ (spontaneous movements), maintained no significant correlation (Fig. S7 and S8).

Additional information regarding the meta-analysis can be found at https://osf.io/hdu5c/.

Publication bias

Visual inspection of the funnel plot for the distance outcome showed an asymmetrical distribution of the studies (Fig. 5a). Trim and fill analysis for distance imputed 4 studies to the meta-analysis. The overall effect of the fungicide exposure was no longer significant for this outcome when imputing potentially unpublished data (SMD -0.29 [-0.66, 0.08], p = 0.1252).

For spontaneous movements, the funnel plot also demonstrated an asymmetrical distribution (Fig. 5b). Trim and fill analysis for this outcome imputed 20 studies to the meta-analysis, and the overall effect of fungicide exposure remained not significant (SMD 0.64 [-0.02, 1.29], p=0.0568).

Egger's regression test indicated publication bias only for spontaneous movements, which showed a p < 0.0001 (for distance, p = 0.4120) (Table S1).

Sensitivity analysis

The leave-one-out analysis for distance revealed that none of the comparisons significantly modified the metaanalysis result (Fig. 6a). The overall effect and heterogeneity remained close to the original value. However, to confirm that any isolated study is skewing the results, we performed another meta-analysis, excluding all the comparisons from the study by⁶⁷. This study showed unusually high SMD in the forest plot, and the omission of their experiments in the leave-one-out analysis altered the overall effect direction. The significant overall effect was sustained (SMD -0.31 [-0.54; -0.08] (Fig. S9).

The leave-on-out analysis for spontaneous movements showed that omitting comparisons did not significantly modify the meta-analysis original result (Fig. 6b). We also ran the meta-analysis without 2 studies:^{65,68}. In the forest plot, these studies showed atypically high SMD, and omitting their experiments in the leave-one-out

Study	Fungicide	Concentration (mg/L)	SMD	SMD	95%-CI	Weight
Larva						
Shen_2020.1	Mepanipyrim	0.0001		0.97	[0.04; 1.90]	1.7%
Shen_2020.5	Mepanipyrim	0.0001		0.22	[-0.68; 1.11]	1.7%
Forner-Piquer_2021.1	Thiophonoto	0.0001	I	0.00	[-0.34, 0.34]	2.0%
Forner-Piquer_2021.12	Cantan	0.0001	Ţ	-0.05	[-0.33, 0.47]	2.0%
Forner-Piquer 2021.17	Boscalid	0.0001	-	-0.31	[-0.74: 0.13]	2.0%
Forner-Piquer 2021.2	Ziram	0.0005		-0.88	[-1.28; -0.47]	2.0%
Forner-Piquer_2021.6	Thiophanate	0.0005		-0.11	[-0.60; 0.39]	1.9%
Forner-Piquer_2021.13	Captan	0.0005		-0.13	[-0.54; 0.28]	2.0%
Forner-Piquer_2021.18	Boscalid	0.0005	-	-0.71	[-1.19; -0.22]	2.0%
Shen_2020.2	Mepanipyrim	0.001		1.38	[0.41; 2.35]	1.6%
Shen_2020.6	Mepanipyrim	0.001		0.48	[-0.42; 1.39]	1.7%
Forner-Piquer_2021.3	Ziram	0.001	•	-0.80	[-1.16; -0.45]	2.0%
Forner-Piquer_2021.7	Thiophanate	0.001		-0.34	[-0.77; 0.10]	2.0%
Forner-Piquer_2021.14	Captan	0.001	+	0.05	[-0.32; 0.42]	2.0%
Forner-Piquer_2021.19	Boscalid	0.001	-	-0.50	[-0.94; -0.05]	2.0%
Li_2019.1	Pyraclostrobin	0.009		-1.85	[-2.91; -0.78]	1.6%
Shen_2020.3	Mepanipyrim	0.01		1.30	[0.34; 2.26]	1.6%
Shen_2020.7	Mepanipyrim	0.01		1.84	[0.82; 2.86]	1.6%
Forner-Piquer_2021.4	Ziram	0.01	-	-0.68	[-1.10; -0.25]	2.0%
Forner-Piquer_2021.8	Thiophanate	0.01		-0.86	[-1.38; -0.34]	1.9%
Forner-Piquer_2021.15	Captan	0.01		-1.46	[-2.08; -0.85]	1.9%
Forner-Piquer_2021.20	Boscalid	0.01	-	0.40	[-0.07; 0.87]	2.0%
LI_2019.2	Pyraciostropin	0.018		-4.90	[-0.00; -3.19]	1.1%
LI_2019.3	Pyraciostrobin	0.038		-0.30	[-0.37; -4.24]	0.9%
Brenet_2021.1	Dixaten	0.0828		-0.39	[-0.76; -0.01]	2.0%
Shen 2020.4	Mepanipyrim	0.1		1.07	[1.30, 4.40]	1.470
Eorner-Piquer 2021.9	Thionhanate	0.1		-0.99	[-2.01, -0.13]	1.0%
Forner-Piquer_2021.16	Cantan	0.1	-	-0.00	[-1.02, -0.47] [-0.97: 0.05]	1.9%
Forner-Piquer_2021.10	Boscalid	0.1		0.34	[-0.37, 0.03]	1.9%
Yang 2021b 1	Thifluzamide	0.19		-1 79	[-2.84: -0.73]	1.6%
Yang 2019b.1	Thifluzamide	0.19		0.93	[-0.43: 2.28]	1.4%
Brenet 2021.2	Bixafen	0.2071	-	-0.91	[-1.30: -0.52]	2.0%
Teng 2019.1	Propiconazole	0.5		-0.11	[-1.71; 1.49]	1.2%
Altenhofen_2017.1	Tebuconazole	1		-0.60	[-1.47; 0.27]	1.7%
- Forner-Piquer_2021.10	Thiophanate	1	+	0.00	[-0.49; 0.49]	1.9%
Forner-Piquer_2021.22	Boscalid	1		0.45	[-0.06; 0.96]	1.9%
Yang_2021b.2	Thifluzamide	1.9		-1.30	[-2.29; -0.32]	1.6%
Yang_2019b.2	Thifluzamide	1.9		0.25	[-1.05; 1.55]	1.4%
Altenhofen_2017.2	Tebuconazole	2		-0.65	[-1.52; 0.22]	1.7%
Teng_2019.2	Propiconazole	2.5		-2.75	[-5.19; -0.31]	0.8%
Yang_2019b.3	Thifluzamide	2.85		-0.67	[-1.99; 0.66]	1.4%
Yang_2021b.3	Thifluzamide	2.85		-1.03	[-1.99; -0.07]	1.6%
Altenhofen_2017.3	Tebuconazole	4		-1.32	[-2.25; -0.40]	1.7%
Teng_2019.3	Propiconazole	4.5		-2.96	[-5.51; -0.42]	0.7%
Forner-Piquer_2021.11	Thiophanate	10		-1.10	[-1.63; -0.57]	1.9%
Random effects model			♦	-0.44	[-0.83; -0.05]	80.5%
Prediction interval					[-2.68; 1.80]	
Heterogeneity: $I^2 = 84\%$, τ^2	= 1.21, <i>p</i> < 0.01					
Test for effect in subgroup:	$t_{46} = -2.27 \ (p = 0.0)$	03)				
Adult	0					0.001
Tang_2021.1	Cyprodinii	0.0001		-0.29	[-2.62; 2.04]	0.8%
Valadas_2019.1	Propiconazole	0.0004		-0.09	[-1.19; 1.00]	1.5%
valauas_2019.2	Cyprodiail	0.0008		-0.14	[-1.24; U.96]	0.7%
Valadae 2019 3	Proniconazola	0.0017		-0.31	[-2.30, 1.37]	1 5%
Valadas 2019.4	Propiconazole	0.0085		-1.12	[-1.40, 0.72]	1.5%
Tang 2021 3	Cyprodinil	0.01		-0.57	[-3.10: 1.95]	0.7%
Pompermaier 2021.1	Copper	0.105		-0.36	[-1.27: 0.54]	1.7%
Pompermaier 2021.2	Copper	0.105		-0.39	[-1.41: 0.64]	1.6%
Altenhofen 2017.4	Tebuconazole	1		-0.52	[-1.33; 0.29]	1.8%
Altenhofen_2017.5	Tebuconazole	2		-1.23	[-2.09; -0.38]	1.7%
Altenhofen_2017.6	Tebuconazole	4	_ _ _	-1.53	[-2.42; -0.64]	1.7%
Paredes-Zúñiga_2019.1	Triadimefon	5		0.24	[-1.12; 1.60]	1.3%
Paredes-Zúñiga_2019.2	Triadimefon	20		0.63	[-0.77; 2.03]	1.3%
Random effects model			↓	-0.55	[-0.89; -0.21]	19.5%
Prediction interval			<u></u>		[-1.25; 0.16]	
Heterogeneity: $I^2 = 5\%$, $\tau^2 =$	0.07, <i>p</i> = 0.39					
Test for effect in subgroup:	$t_{13} = -3.46 \ (p < 0.0)$	01)				
Random effects model			•	-0.44	[-0.74; -0.13]	100.0%
Prediction interval					[-2.33; 1.46]	
Heterogeneity: $I^2 = 80\%$, τ^2	= 0.88, p < 0.01		-5 0 5			
rest for subgroup difference	es: χ ₁ = 0.17, df = 1	1 (p = 0.68)	Higher in control group Higher in exposed group			

Figure 3. The effect of exposure to fungicides on distance traveled in zebrafish. Subgroup analyses were based on the developmental stage (either larva or adult). Data are presented as Hedges' G standardized mean differences (SMD) and 95% confidence intervals.

analysis changed the overall effect direction. Although the direction of the effect changed, it was still not significant (SMD 0.22 [-021; 066]) (Fig. S10).

Study	Fungicide	Concentration (mg/L)	SMD	SMD	95%-CI	Weight
Tang_2020.1	Cyprodinil	0.0001		-0.14	[-2.29; 2.01]	1.5%
da Costa-Silva_2018.1	Mancozeb	0.001	•	1.84	[1.37; 2.31]	2.4%
Tang_2020.2	Cyprodinil	0.001		-0.78	[-3.04; 1.48]	1.5%
Li_2018.1	Pyraoxystrobin	0.002		2.05	[0.78; 3.32]	2.0%
Li_2018.2	Pyraoxystrobin	0.002		-0.01	[-1.09; 1.08]	2.1%
Li_2018.3	Pyraoxystrobin	0.002	—	0.61	[-0.49; 1.71]	2.1%
Li_2018.4	Pyraoxystrobin	0.003	-	2.50	[1.15; 3.86]	2.0%
Teng_2018.2	Difenoconazole	0.005	🛓	0.00	[-0.82; 0.82]	2.3%
Tang_2020.3	Cyprodinil	0.01	_	-0.35	[-2.52; 1.82]	1.5%
Teng_2018.3	Difenoconazole	0.05	🛓	0.00	[-0.82; 0.82]	2.3%
Tian_2019.2	Prothioconazole	0.075	- <u>+</u> -	-0.42	[-2.05; 1.20]	1.8%
Tang_2020.4	Cyprodinil	0.1		-2.25	[-5.21; 0.71]	1.1%
Tian 2019.3	Prothioconazole	0.15		0.86	[-0.84; 2.56]	1.8%
	Prothioconazole	0.375		-0.36	[-1.98; 1.26]	1.8%
Teng 2018.1	Difenoconazole	0.5	1	0.00	[-0.82; 0.82]	2.3%
Teng 2018.4	Difenoconazole	0.5	1	0.00	[-0.82; 0.82]	2.3%
Teng 2019.1	Propiconazole	0.5	-	0.11	[-1.49; 1.71]	1.8%
Mu 2013.1	Difenoconazole	0.5		0.09	[-1.96; 2.15]	1.6%
- Zhou 2019.1	Captan	0.58	-	-0.50	[-2.04: 1.04]	1.9%
Zhou 2019.2	Captan	0.66	-	-1.38	[-3.04: 0.28]	1.8%
Zhou 2019.3	Captan	0.76		-0.68	[-2.24: 0.87]	1.9%
Zhou 2019.4	Captan	0.87		-1.12	[-2.73: 0.49]	1.8%
Mu 2013 2	Difenoconazole	1		0.43	[-1.64: 2.50]	1.6%
Zhou 2019 5	Cantan	1		-0.68	[-2.23: 0.87]	1.9%
Lin 2021 1	Fluxapyroxad	1.1		-0.27	[-2.43: 1.89]	1.5%
Lin_2021.1	Eluxapyroxad	1.1	1	0.31	[-1.85; 2.48]	1.5%
Lin_2021.2	Fluxapyroxad	1.2		0.90	[-1.00, 2.40]	1.0%
Lin_2021.3	Fluxapyroxad	1.3		1.57	[-1.01: 4.15]	1 296
Lin_2021.4	Elutologil	1.4		1.57	[2 12: 0 22]	1.0%
Tang_20100.1	Diferencementele	1.5		-1.45	[-3.12, 0.22]	1.0%
Wu_2013.3	Eluveryreved	1.5		1.07	[-0.60, 3.54]	1.3%
Lin_2021.5	Fluxapyroxad	1.5		1.05	[-1.30, 3.40]	1.490
Ciap. 2018 1	Pluxapyroxau	1.0		0.12	[-1.14, 3.79]	0.4%
Qiai1_2016.1	Buscaliu	1.7		0.13	[-1.09, 1.34]	2.170
Cian 2018 2	Flutolanii	1.0		-1.57	[-3.20; 0.13]	0.4%
Qian_2018.2	Boscalid	2		0.39	[-0.83; 1.61]	2.1%
Mu_2013.4	Difenoconazole	2		-0.52	[-2.59; 1.55]	1.6%
Yang_2016b.3	Flutolanii	2.16		-2.46	[-4.38; -0.53]	1.6%
Qian_2018.3	Boscalid	2.3		1.74	[0.41; 3.08]	2.0%
Teng_2019.2	Propiconazole	2.5		1.87	[-0.16; 3.91]	1.6%
Mu_2013.5	Difenoconazole	2.5		-1.74	[-3.98; 0.50]	1.5%
Yang_2016b.4	Flutolanii	2.59		-2.19	[-4.04; -0.34]	1.7%
Qian_2018.4	Boscalid	2.6		3.18	[1.61; 4.76]	1.9%
Yang_2016.1	Thifluzamide	2.66		-1.19	[-3.33; 0.96]	1.5%
Yang_2016.2	Thifluzamide	2.76		-2.16	[-4.50; 0.18]	1.4%
Yang_2016.3	Inifluzamide	2.85		-2.36	[-4.75; 0.03]	1.4%
Qian_2018.5	Boscalid	2.9		3.26	[1.67; 4.86]	1.8%
Cao_2019b.1	Cyproconazole	2.9178		0.96	[-1.35; 3.28]	1.4%
Cao_2019b.7	Cyproconazole	2.9178		0.87	[-1.42; 3.16]	1.4%
Yang_2016.4	Thifluzamide	2.95		-4.41	[-7.47; -1.35]	1.1%
Mu_2013.6	Difenoconazole	3		-2.35	[-4.73; 0.04]	1.4%
Yang_2016.5	Inifiuzamide	3.04		-10.95	[-16.94; -4.96]	0.4%
Yang_2016b.5	Flutolanil	3.10		-2.19	[-4.04; -0.34]	1.7%
Qian_2018.6	Boscalid	3.2		3.67	[1.99; 5.35]	1.8%
Yang_2016.6	Thifluzamide	3.23		-11.14	[-17.21; -5.06]	0.4%
Teng_2019.3	Propiconazole	4.5	1 -	1.67	[-0.28; 3.63]	1.6%
Cao_2019b.2	Cyproconazole	7.2945		0.07	[-2.08; 2.22]	1.5%
Cao_2019b.8	Cyproconazole	7.2945		-0.21	[-2.37; 1.95]	1.5%
Cao_2019b.3	Cyproconazole	14.5829		-0.68	[-2.92; 1.55]	1.5%
Cao_2019b.9	Cyproconazole	14.5829		-0.86	[-3.14; 1.43]	1.4%
Cao_2019b.4	Cyproconazole	29.178		-9.58	[-18.53; -0.62]	0.2%
Cao_2019b.10	Cyproconazole	29.178		-4.39	[-8.91; 0.14]	0.6%
Cao_2019b.5	Cyproconazole	72.945		-6.49	[-12.76; -0.22]	0.4%
Cao_2019b.11	Cyproconazole	72.945		-4.87	[-9.78; 0.05]	0.6%
Cao_2019b.6	Cyproconazole	145.89	•	-10.17	[-19.65; -0.69]	0.2%
Random effects model			•	-0.16	[-0.67; 0.34]	100.0%
Prediction interval					[-2.92; 2.60]	
Heterogeneity: $I^2 = 74\%$, τ^2	= 1.86, <i>p</i> < 0.01		-10 0 10			
			Higher in control group Higher in exposed group			

Figure 4. The effect of exposure to fungicides on spontaneous movements in zebrafish. Data are presented as Hedges' G standardized mean differences (SMD) and 95% confidence intervals.

Discussion

This work aimed to evaluate and synthesize the neurobehavioral effects of fungicide exposure in zebrafish through a systematic review and meta-analysis. As main findings, we can highlight that fungicides cause a decrease in distance traveled by larval and adult zebrafish; no effect was observed on spontaneous movements of embryos. The locomotor behavior was the category most frequently assessed in the included studies. Along with distance traveled, velocity was also commonly reported. It is important to emphasize that a decreased distance



Figure 5. Funnel plot including studies analyzed within distance (**a**) and spontaneous movements (**b**) outcomes. Each gray circle represents a single comparison. Hollow circles represent imputed studies in the trim and fill analysis. The vertical line represents the overall effect size, and the triangular region represents the 95% confidence interval. Shaded areas represent the interval for statistically significant effects.

traveled or velocity does not necessarily imply toxicity, as a substance may have a sedative effect. However, even if not directly related to toxicity or locomotor damage, altered locomotion poses a risk to organisms as it impacts their ability to forage, reproduce, and escape predators⁶⁹. These data should be observed together with the neurochemical outcomes, which were also consistently investigated and are linked to behavioral variation. The included studies frequently reported altered outcomes related to enzymatic activity, some involved in locomotion (AChE) and oxidative status (GST, SOD, GPx, among others), which are possible mechanisms for reduced locomotor behavior. Few included studies reported investigations of behavioral domains other than locomotor (9), and even so, it was limited to anxiety-fear-related and aggressive behavior, revealing a gap in the literature. The lack of standardized protocols or unpublished negative results could explain this observation⁷⁰.

The overall high heterogeneity observed in the meta-analysis for distance traveled can be attributed to several sources. The experimental conditions, from rearing until exposure and tests, are extremely variable between laboratories. The researchers employed many protocols, including distinct durations of exposure, frequency of solution renewal, number of coexposed animals, age of the fish, type, and test apparatus. When considering the subgroup analysis, studies with adults had a lower heterogeneity than those performed at the larval stage. Even though fewer adult studies were included, we can indeed verify more uniformity between the protocols of these



Figure 6. Sensitivity analyses for studies for distance (**a**) and spontaneous movements (**b**) outcomes. Data are presented as Hedges' G standardized mean differences (SMD) and 95% confidence intervals.

experiments, mostly during the outcome assessment. Therefore, this similarity can explain the low heterogeneity of this subgroup.

Interestingly, there was no significant difference between the subgroups, indicating that the developmental stage of the animals does not significantly impact the effect of fungicides on the distance traveled. Despite the different locomotor mechanisms exhibited by adults and larvae⁶⁹, it suggests that fungicide exposure consistently affects both subgroups.

On the other hand, the heterogeneity of the outcome of spontaneous movements was considered moderate. Unlike the distance traveled, the spontaneous movements can be measured in a single developmental stage: the embryo, generally at 24 h post-fertilization (hpf). Consequently, the age of the animals can be excluded as a potential source of heterogeneity, which helps to explain why the heterogeneity did not reach the highest level.

The reporting quality analysis showed a high percentage of negative answers, especially regarding "sample size estimation" and "inclusion or exclusion criteria". None of the authors explicitly stated previously determined parameters for the eligibility of the data. The result from this evaluation indicates that the conclusions of this review should be interpreted with caution since the report of the included studies presents considerable uncertainty. This lack of methodological information has been recognized as one of the main reasons behind the reproducibility crisis in preclinical research⁷¹. Aiming to improve the quality of the studies, guidelines for the research report with animals have been developed in the last years⁷²; however, it is a multifaceted problem that demands complex and long-term solutions⁷³.

Trim and fill analysis for distance imputed 4 studies into the meta-analysis, resulting in no overall significant effect. This fact suggests the presence of missing studies with null and/or significant results⁷⁴. The unpublished data may have influenced the previously observed significant effect, revealing a potential bias towards the publication of studies only with significant findings in which fungicide exposure decreases locomotion. However, Egger's test suggests no evidence of publication bias.

Despite the input of 20 studies in the trim and fill analysis for spontaneous movements, it did not alter the non-significant overall effect found in the meta-analysis. This indicates that publication bias may not explain the observed non-significance. However, it is important to note that the significant result obtained from Egger's test indicates the presence of potential publication bias. The Egger's test suggests a tendency to publish studies with significant results, which could skew the meta-analysis. Although the trim and fill analysis did not change the overall effect, the imputed studies may impact the precision and confidence interval of the effect estimate. There is an important role of selective publishing in the misinterpretation of a meta-analysis⁷⁵, highlighting the need for new practices regarding the publication of non-significant results. Even if this represents a complex, deep-rooted issue that requires a change in the whole culture of publishing scientific data, some authors have been raising this discussion and proposing alternatives⁷⁶⁻⁷⁹. However, the results of our publication bias analysis should be interpreted with caution, as our funnel plots were based on SMD versus standard error (SE). Although this method is standard practice in the field, it may introduce distortion and overestimate the existence of publication bias, as demonstrated empirically by Zwetsloot et al.⁸⁰.

The sensitivity analysis indicated that the meta-analysis results were not significantly influenced by any particular study or set of studies, suggesting that the overall effect size is robust and reliable. This finding supports the validity of the meta-analytic conclusions and can increase the confidence in the reliability of the results. However, the reliability of each comparison could not be determined due to poor reporting practices and a general lack of protocol preregistration.

One limitation of this study was the inclusion of only studies that used analytical-grade fungicides while excluding those involving commercial formulations and fungicide mixtures. This exclusion was necessary to isolate the specific effects of individual chemicals and ensure more accurate conclusions. Although this approach may be less realistic, it enhances the precision of the findings. Additionally, we did not restrict the inclusion criteria to studies involving exposure to environmentally relevant concentrations, as this would severely reduce the number of eligible articles, making it impossible to conduct a comprehensive meta-analysis. Another significant limitation worth highlighting is the potential inclusion of fraudulent data, which becomes evident when implausible results are observed. While various tools and techniques exist to perform statistical checks and verify data integrity, it is important to note that there is currently no foolproof method to confirm whether a study is fraudulent or not definitively. This task becomes even more challenging without direct access to the data.

Our results reinforce the effects of these chemicals, with their misuse representing a threat to the ecosystems. Since we depend on the affected environment, its contamination is an alert to public health. Besides that, we confirm the demand for well-designed studies with greater clarity of report on this topic. The authors should clearly state key elements such as sample size, sample size estimation, data inclusion or exclusion criteria, and blinding. Some available tools, like preregistration of study protocols and adherence to animal studies reporting guidelines such as the ARRIVE⁷², could be useful. Compliance with specific reporting guidelines for ecotoxicological studies as the "Criteria for Reporting and evaluating Ecotoxicity Data" (CRED)⁸¹ is also highly encouraged. In addition, standardization of behavioral tests could enable more comprehensive meta-analyses. These recommendations can lead to more reliable conclusions and contribute to effectively monitoring environmental pollution.

Data availability

All data are available at Open Science Framework (https://osf.io/hdu5c/).

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Competing interests

The authors declare no competing interests.

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