Protocol Article

The submerged plus maze as an assay for studying anxiety-like behaviour in fish

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A B S T R A C T

The elevated plus maze is a commonly used and well-validated test of anxiety-related behaviour in rodents. The use of fish in behavioural neuroscience paradigms is increasing, necessitating an equivalent test for studying anxiety-like behaviour in fish. Because behaviour in the elevated plus maze is driven by aversion to open space, the submerged plus maze described here uses transparent walls to elicit similar behaviour in fish. The tendency of fish to explore or avoid the sections of the maze containing transparent walls is used as proxy for anxiety level. This submerged plus maze was designed and validated for convict cichlid (Amatitlania nigrofasciata) fish.

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A R T I C L E   I N F O

Protocol name: Submerged plus maze
Keywords: Elevated plus maze, Anxiety-testing, Behavioural testing, Fish behaviour, Anxiolytic, Anxiogenic
Article history: Received 24 February 2019; Accepted 3 July 2019; Available online 8 July 2019

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https://doi.org/10.1016/j.mex.2019.07.002
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**Value of the Protocol**

- Fish are increasingly more prevalent in anxiety research.
- The submerged plus maze is an aquatic adaptation of the rodent elevated plus maze.
- The submerged plus maze can be used to identify anxiolytic or anxiogenic drugs in fish.

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**Specifications Table**

| Subject Area: | Neuroscience |
|---------------|--------------|
| More specific subject area: | Animal behaviour |
| Protocol name: | Submerged Plus Maze |
| Reagents/tools: | Fish, dip net, apparatus, acclimation chamber, tank water |
| Experimental design: | Anxiety-like behaviour examined in an aquatic plus maze, where transparent walls elicit behaviours as a result of aversion to open spaces. |
| Trial registration: | Protocols were approved by the University of Alberta Biological Sciences Animal Policy and Welfare Committee (protocol number 00000055) and adhere to the guidelines of the Canadian Council for Animal Care. |

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**Description of protocol**

The elevated plus maze is used to study anxiety-like behaviour in rodents [1,2] by examining aversion to open spaces (i.e., open arms relative to closed arms) [2]. Studying anxiety using fish is becoming more popular and, as such, an equivalent test should be available. To date, there have been two aquatic mazes using a plus maze structure with four arms [3–5]. Walsh-Monteiro and colleagues [3] constructed a maze with two arms containing ramps that decreased the height of the water column with increasing distance from the center of the maze. In a second maze, Sackerman et al. [4] alternated the arms with white and dark walls, similar to a light/dark preference test [4,5]. Although both of these mazes use a ‘plus’ format, neither are analogous to an elevated plus maze in rodents because of its dependence on aversion to open spaces [2]. Most fish, like rodents, display a preference for dark areas vs. light [6], either for cover [7] or to have a dark background to hide against [8], likely for its potential to provide more protection from predators [9]. The submerged plus maze, as described here, uses aversion to open spaces as an indication of anxiety-like behaviour in fish [3]. This apparatus uses a plus maze format, which includes alternating arms constructed of black or transparent plexiglass walls (each arm 12 cm × 4.5 cm × 13 cm, see Figs. 1 and 2). This arrangement results in two visually closed arms, two visually open arms, and a center area similar to that of the elevated plus maze. The arms are marked in 1.5 cm increments to quantify travel in each area. A transparent plastic cylinder (4.5 cm × 4.5 cm × 13 cm) serves as an acclimation chamber and is placed in the centre area at the start of each trial. A video recording device views the maze from above to record the movements of the fish in the maze for later behavioural scoring. The present study used convict cichlids (Amatitlania nigrofasciata) to validate the test, however, the submerged plus maze is also amenable to testing other fish species. Individual fish of the same species may display strong biases for making turns to either the right or left depending on the individual, and these biases can be sensitive to a fish’s shy-vs-bold personality and to the fish’s perceived probability of impending positive or negative outcomes [10,11]. This apparatus has the advantage of being symmetrical, an arm of the opposing type to that currently occupied is immediately to the right and left when exiting the current arm. This should minimize cerebral lateralization biases, but researchers may benefit by recording whether turns are biased to either the right or left when exiting each type of arm.
Methods

Behaviour testing

1 Set up plus maze apparatus with a video recorder mounted above the maze so that all areas can be seen clearly. Ensure all areas of the maze are well-lit while reducing glare.
   a Experimenter should be occluded from view.
2 Fill maze with aerated stock tank water or clean water from the fish habitat to a depth of 10 cm and place clear acclimation chamber in the centre area of the maze. Ensure the water matches the physical-chemical properties (temperature, pH, etc.) of habitat water to reduce the risk of climate shock. Specifically, for convict cichlids, we recommend that the water is as follows: pH 7–8, temperature 25 ± 1 °C.
3 Remove fish from home tank using a dip net and deposit in acclimation chamber of the plus maze. Record handling time.
4 Leave fish in the acclimation chamber for two minutes.
   a Start the video recording before the two minutes are fully elapsed (i.e., at 1.5 min).
5 Remove the acclimation chamber, allowing the fish to move freely within the maze. Record behaviour for 5 min.
   a Do not make any loud noises that may disturb the fish.

Fig. 1. Submerged plus maze apparatus schematic. The apparatus is shaped as a plus symbol with alternating black (black fill) and transparent (white fill, dashed lines) arms. Arms (12 cm long × 4.5 cm wide) are marked to quantify travel within the maze. Fish were placed in an acclimation chamber in the centre area for two minutes before they were released to explore the maze for five minutes.
b Allow the recording to run slightly past 5 min in case water movement from the removal of the chamber prevents immediate detection of the fish.
6 Return the fish to its home tank.
7 If possible, water should be changed between fish. Otherwise, the temperature, pH, etc. of the water in the arena should be maintained and monitored throughout testing and aerated in between trials. If the temperature drops too low (Δ3 °C) or water chemistry changes too drastically, then the arena should be filled with new water or adjusted.
   a The arena should also be rotated after every trial to eliminate any spatial biases or any other uncontrolled auditory or visual stimuli.

Behaviour scoring

1 Score behaviour using an automated behaviour tracking software (e.g., EthoVision (Noldus), etc.) if possible.
   a This may be difficult if there is too much light glare on the surface of the water or if the fish does not contrast well enough with its background.
2 If the experimenter is scoring behaviour manually, take note of the time the fish enters each arm and how many lines the fish crosses within each arm.
3 Sum the amount of time the fish occupies each area and the number of lines crossed in each of the three areas (i.e., visually closed arms, visually open arms, and centre area).
4 Count the number of entries into new arms and into visually open arms.

Validation

This apparatus was validated using the anxiety-reducing (anxiolytic) benzodiazepine diazepam administered by immersion (Fig. 3) [12–15]. Immersion is the method of choice because injection of substances requires anesthesia with MS-222 or cold water, which can impair behavioural responses.
Table 1
Drug solutions for validation. Amount of reagents required for administration through immersion at typical drug concentrations. Vehicle for diazepam was 5% DMSO in tank water and diazepam is soluble in 100 mM DMSO. Dilute diazepam + vehicle solution to a final volume of 500 mL for administration.

| Concentration (mg/L) | Diazepam (mg) | DMSO (μL) | Tank Water (mL) | Final Volume (mL) |
|----------------------|---------------|-----------|-----------------|------------------|
| 2.5                  | 1.25          | 44        | 8.8             | →500             |
| 5                    | 2.5           | 88        | 17.6            | →500             |
| 10                   | 5.0           | 176       | 35.1            | →500             |

Immersion can quickly and reliably be used to validate anxiety-like behavioural tests with other anxiolytic substances (e.g., ethanol) [16], as well as anxiogenic substances (e.g., GABA\textsubscript{A} receptor antagonist, gabazine) [17,18]. Note that time of immersion in a drug solution can vary from 3 min (e.g., Diazepam) [12,15] up to thirty minutes [16]. If a longer duration of immersion is used, the dosing beaker should be kept under a heating mat to maintain water temperature [16]. To reduce the amount of handling the fish experienced, we used a slotted plastic cup to transport fish between each step. This likely minimizes netting stress and damage to the slime coat of the fish. Fish were housed individually in aquaria partitioned by transparent dividers to control for aggression and social isolation effects and to ensure handling stress was restricted to the fish of interest.

1. Dissolve diazepam (100 mM) in a 0.5% solution of dimethyl sulfoxide (DMSO) in aerated tank water. Then, dilute this solution with aerated tank water to a final volume of 500 mL while maintaining the desired concentration (2.5 mg/L was used in the current study, see Table 1) in the drug administration beaker. Agitate slightly to resuspend the solution.
   a. Note: 2.5 mg/L was the concentration used in the current study, determined after an initial pilot test to obtain a dose-response curve. Different sizes and species of fish may respond optimally to different drug concentrations and therefore any new studies should obtain an appropriate dose-response curve.
2. Prepare an additional holding container with aerated tank water.
3. Remove the fish from its home tank with a dip net and deposit fish into the slotted plastic cup.
4. Expose the fish to the drug by placing the cup holding the fish in the drug administration beaker for 3 min.
   a. Note: the drug solution remains in this beaker and is not transferred to any subsequent steps.
5. Lift the cup and fish and move them into the holding container for 5 min to allow the drug to take effect.
6. Lift the cup and fish and transfer fish into the acclimation chamber of the submerged plus maze by pouring it out of the cup and begin the test.

This validation method was used in a within-subjects design, so fish were exposed to the submerged plus maze with and without prior diazepam administration. In vehicle trials, fish underwent the same steps but with only DMSO and tank water in the drug administration beaker. Drug and vehicle trial order were randomized to ensure behavioural differences between trials was not due to testing order, and we imposed a 48-h inter-trial interval to ensure drug elimination between trials. Note that we used a within-subjects design because of low numbers of fish and high individual variability, however, a between-subjects design can also be used to validate this test with other fish species.

\[ \text{Drug Tank} \rightarrow \text{Delay Tank} \rightarrow \text{Behaviour Test} \]

**Fig. 3.** Anxiolytic validation procedure. A fish is first placed into a tank containing the drug treatment (diazepam + vehicle or vehicle only) for 3 min. Next, the fish was moved to a delay tank (tank water) for 5 min to allow the drug to take effect. Finally, the fish was moved to the submerged plus maze test, where it acclimated for 2 min before the 5 min testing period.
Additional information

The dimensions of this apparatus were designed to accommodate fish ranging between 1.0–4.5 cm in standard length. An alternate apparatus for fish ranging between 3.0–7.0 cm in standard length was constructed where each arm measured 24 cm × 9 cm × 16.5 cm and lines were marked in 3 cm increments. The maze dimensions should be adjusted accordingly to match target fish size.

A basin should be used to surround the maze and water should flow between the basin and the maze to alleviate pressure against the walls of the plus maze. The present maze employed small gaps at the far end of each arm to exchange water with the surrounding basin. A previous iteration of the maze used gaps too large for some of the smallest fish in our experiments and gauze had to be used to prevent the fish from wedging themselves in the gaps. Smaller gaps were used in a subsequent version, eliminating the issue.

Acknowledgements

This research was supported by a Natural Sciences and Engineering Research Council of Canada (NSERC) Discovery Grant to PLH and TJH. We thank Isaac Lank for constructing all iterations of the apparatus, Kennedy Fjellner for insight when adjusting to accommodate smaller fish, Katelyn Wonsiak for assistance in the early stages of validation, and Declan Ali for helpful comments and suggestions in validating the method.

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