



Alternate models for acute fish toxicity testing: a survey

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Agency for Toxic Substances and Disease Registry • Consumer Product Safety Commission • Department of Agriculture • Department of Defense

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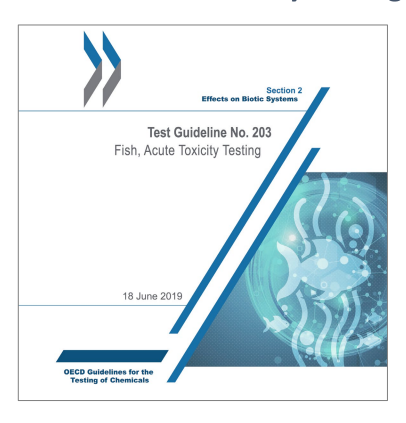


Summary

- Aquatic toxicity testing assesses the adverse effects of chemicals and other environmental stressors on aquatic organisms.
- It involves different media or environments (i.e., marine, freshwater), and a large number of species.
- Common tests include acute and chronic exposures, often standardized by agencies and international bodies such as the US EPA or OECD.
- There is interest and social pressure worldwide to develop alternative methods.
- Here, we summarize the use of the Acute Fish Toxicity Test, which is often requested as part of the registration of new substances and has lethality as an endpoint.
- We then explore the potential of alternate models that could provide hazard information and potentially replace that test.



Acute Fish Toxicity Testing





Fish Acute Toxicity Test

INTRODUCTION

- OECD Guidelines for Testing of Chemicals are periodically reviewed to incorporate scientific progress, changing regulatory needs, and animal welfare considerations. The revision of this Guideline (originally adopted in 1981, updated in 1984, 1992), reflects also updates on a series of recommendations from the OECD Fish Toxicity Testing Framework 2011 (OECD, 2012), and includes:
 - Alternative methods: in the interest of animal welfare and efficient use of resources, it is important to avoid/reduce the use of animals whenever possible and appropriate. Therefore, before carrying out a fish acute toxicity test according to this guideline, it should be considered whether reliable information on fish acute toxicity could be derived with alternative methods in a weight-of-evidence approach, such as the use of QSAR, read-across, fish embryos (OECD 2013), fish cell lines and others. Alternatively, the use of the threshold approach (OECD, 2010) or the limit test as described in § 30 of this guideline may be sufficient. Where testing on fish is required (i.e., alternative methods currently may not be sufficient for all jurisdictions and testing needs. Therefore; make sure the tests fulfil the regulatory requirements), alternative methods such as those listed above can be considered for range finding.
 - A specification that testing the minimum concentration causing 100% and the maximum concentration causing 0% mortality are not mandatory requirements (e.g. no need to test additional concentrations just to demonstrate 0 and/or 100% mortality).
 - guidance on the circumstances under which a water control is required when solvent is used (OECD, 2018).
 - the introduction of estuarine and marine fish species in the recommended species list.
 - the enhanced recording of visible abnormalities (also referred to as sublethal clinical signs) that fish may display during the exposure in order to improve our

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PRINCIPLE OF THE TEST

3. The fish are exposed to the test chemical for a period of 96 hours, under either static, semi-static or flow-through conditions. Mortalities and visible abnormalities related to appearance and behaviour are recorded. Where possible, the concentrations to kill 50% of the fish (LC₅₀) are determined.

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ability to predict chemical toxicity and minimise suffering of animals in the future analogously to those described in Guidance Document No. 19 on the recognition, assessment, and use of clinical signs as humane endpoints for experimental animals used in safety evaluation for mammalian studies (OECD, 2000).

Definitions used in this Test Guideline are given in Annex 1.



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ANNEX 2

TABLE 1: RECOMMENDED FISH SPECIES, TOTAL LENGTHS AND TEST CONDITIONS

Species ⁶	Temperature ⁷ (°C)	Salinity ⁸ (‰)	pН	Hardness (mg/L CaCO ₃)	Photoperiod (hours light)	Recommended length range ⁹ (cm)
Danio rerio						
Zebrafish	21-25	<0.2	6.0-8.5	40- 250, preferably <180	12-16	1-2
Pimephales promelas						
Fathead minnow	21-25	<0.2	6.0-8.5	40-250, preferably <180	12-16	1-3
Cyprinus carpio						
Carp	20-24	<0.2	6.0-8.5	40-250, preferably <180	12-16	2-4
Oryzias latipes						
Japanese Medaka	23-27	<0.2	6.0-8.5	40-250, preferably <180	12-16	1-2
Poecilia reticulata						
Guppy	21-25	<0.2	6.0-8.5	40-250, preferably <180	12-16	1-2
Lepomis macrochirus				1		
Bluegill	21-25	<0.2.	6.0-8.5	40-250, preferably <180	12-16	1-3

⁶ If other species are used, the rationale for the selection of the species must be reported together with any adaptations to the test guideline's recommendations. It is suggested that the species is selected on the basis of their ready availability, ease of maintenance, and historical use in safety testing.

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		_				
Oncorhynchus mykiss Rainbow trout	10-1410	<0.2	6.0-8.5	40-250, preferably <180	12-16	3-6
Gasterosteus aculeatus Three-spined stickleback	13-19	0-35	6.0-8.5	40-7500	12-16	1-2
Cyprinodon variegatus Sheepshead minnow	23-27	15-35	6.0-8.5	3000-7500	12-16	1-2
Dicentrarchus labrax European sea bass	18-22	15-35	6.0-8.5	3000-7500	12-16	4-8
Pagrus major Red sea bream	18-22	30-35	6.0-8.5	5000-7500	12-16	2-4

Where culture temperature differs from the recommended range, the acclimatization period should be used to acclimatize the fish to the desired test temperature.

⁸ For any given test this shall be performed to ± 2‰, e.g. 17±2 =15-19‰, 31±2 =29-33‰.

⁹ Test fish must be juveniles when used in this test (before reaching sexual maturity). If fish of sizes other than those recommended are used, this should be reported together with developmental stage (juvenile, sub-adult, adult stage) and the rationale.



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TABLE 1: Clinical signs observed in fish, compiled from publications (CCAC, 20015; Rufli, 2012; Drummond et al, 1986 and Middlyng et al, 2011) and TG203 score sheets provided by individual laboratories. Non-shaded rows are the major categories of visible abnormality for which recording has been mandatory in TG203 since 1992. Shaded rows are optional explanatory sub-categories.

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Aggression, direct attack, domination of choice tank locations	Mucus secretion	Excess mucus production	
	Faecal (anal) casts	String of faeces hanging from anus or on tank floor	
	Aggression and/or cannibalism		Aggression, direct attack, domination of choice tank locations, pick at or eat bodies of dead fish

Industry sector	Region	Example legislation	Acute in vivo test required for active substances (yes/no)	Species required/recommended, and product/formulation testing requirements
Biocides	EU	EU Biocidal Products Regulation (Regulation EU 528/2012)	Yes	 One freshwater (+marine species, if relevant) Testing of products and ingredients may be required in some circumstances depending on the use pattern, relative sensitivity of other taxa compared to fish, and if the risk cannot be predicted/resolved based on the ingredients. Exemptions can apply to active substance data, if 1) valid chronic (long-term) fish toxicity data are available or 2) in some rare cases, if negligible exposure is expected (attributable to the use pattern or properties of the active substance)

Industry sector	Region	Example legislation	Acute in vivo test required for active substances (yes/no)	Species required/recommended, and product/formulation testing requirements
Biocides	North America		Yes	 Cold freshwater, warm freshwater, marine; requirements can potentially be reduced dependent on use or expected exposure Testing of ingredients
	Asia Pacific		Yes	Cold freshwater; requirements can potentially be reduced dependent on use or expected exposure or country Testing of products and ingredients
	Notes	For some product categorie product registrations (e.g., o	-	available from plant protection insecticides).



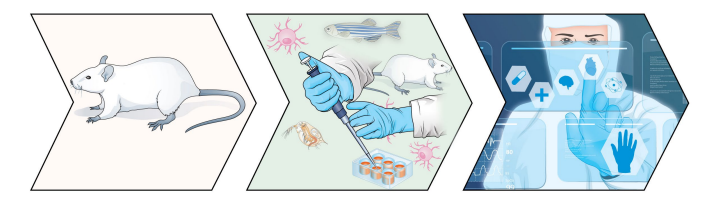
Industry sector	Region	Example legislation	Acute in vivo test required for active substances (yes/no)	Species required/recommended, and product/formulation testing requirements
Human pharmaceuticals	EU	EU Human Pharmaceuticals (Regulation EC 726/2004)	No; considered not relevant because of long- term, low-level exposure	n/a
	North America	US Food and Drug Administration Center for Drug Evaluation and Research	Yes; action limit at expected environmental concentration >100 ng/L (if not an endocrine-disrupting compound), then a tiered approach if Daphnia or algae risk quotient <1000	Not specified

Industry sector	Region	Example legislation	Acute in vivo test required for active substances (yes/no)	Species required/recommended, and product/formulation testing requirements
Cosmetics	EU		No, although information on fish may be required on ingredients covered under REACH (>10 tonnes/yr); see Notes	
	Rest of world		Country-specific	Dependent on country; requirement in China for ingredients imported >1 tonne/yr to test with a native species
	Notes	November 2009 on cosmet finished cosmetic products	tic products sets out a and cosmetic ingredic ing finished cosmetic	arliament and of the Council of 30 testing ban—prohibition on testing ents on animals and a marketing products and ingredients in the



Alternative Approaches: the 3R-Principle

- REDUCE the number of animals used in testing
- REFINE any procedures to minimize pain, suffering, and distress
- REPLACE the use of animals whenever possible



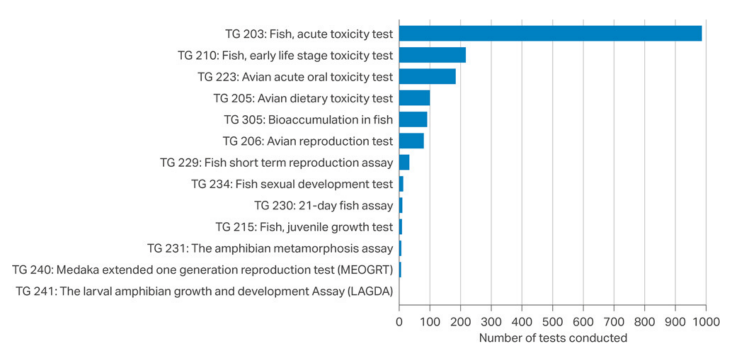


Moving towards the 6R-principle



- » Reproducible and Reliable
- » Relevant
- » Regulatory accepted

OECD test guideline vertebrate ecotoxicology studies conducted across 15 contract research organizations from 2014 through 2017





Considerations for in vitro methods

- Chemical space coverage
- Performance metrics:
 - Assay controls
 - Factors affecting assay results
- Proprietary information



Chemical space coverage

- The chemical space for fish acute toxicity testing is large (number of chemicals) and broad (variety of chemical properties), reflecting the wide range of regulatory (e.g., pesticides, pharmaceuticals, and industrial chemicals) and associated research needs for toxicity data on compounds that may reach aquatic environments.
- NAM coverage will be limited by factors including but not limited to physiochemical properties and mode of action (MOA) of the chemical of interest and those used to develop an approach.
- For regulatory or research acceptance, NAMs approaches must establish the range of their "fit for purpose" to be applied in lieu of or as a supplement to whole-fish acute toxicity studies.



Performance metrics

- There are numerous factors to consider when evaluating the quality of results from a NAM for estimating the acute toxicity of a chemical to fish using *in vitro* laboratory methods
- It is critical to test in-process control measurements (e.g., positive chemical control, dilution water control [or negative control without test chemicals]), solvent control, and, in the case of *in vitro*-NAMs, a no-cells control with only the assay reagents) to measure key sources of variability each time the assay is performed to ensure consistent performance



Performance metrics: Key control measurements in acute fish toxicity tests and NAMs

		_	
Guideline name	Control	Variation permitted	Citation
	Difference in cytotoxicity between solvent and negative		
RTgill-W1	controls	≤10%	(ISO, 2019)
Freshwater Alga and Cyanobacteria, Growth Inhibition Test	Average specific growth rate in replicate control cultures	≤ 7%	(OECD, 2011a)
Algal Toxicity Test	Average specific growth rate in replicate control cultures	< 15	(U.S. EPA, 2012a)
Daphnia Acute Immobilisation Test	Immobilisation in dilution water and solvent controls	≤10%	(OECD, 2004a)
Aquatic Invertebrate Acute Toxicity Test, Freshwater <i>Daphnids</i>	Immobilisation in dilution water and solvent controls	≤10%	(U.S. EPA, 2016a)
Fish Acute Toxicity Test	Mortality of dilution water and solvent controls	≤10%	(OECD, 2019a)
Freshwater and Saltwater Fish Acute Toxicity Test	Mortality of dilution water and solvent controls	≤10%	(U.S. EPA, 2016a)
Fish Embryo Acute Toxicity (FET) Test	Mortality of dilution water and solvent controls	≤10%	(OECD, 2013a)
Fish Embryo Acute Toxicity (FET) Test	Positive control mortality	>30%	(OECD, 2013a)
Fish Early Life Stage (FELS) Toxicity Test	Hatching success of control groups	>66-80% ^a	(U.S. EPA, 2016b)
Fish Early Life Stage (FELS) Toxicity Test	Post-hatch success of control groups	>60-80% ^a	(U.S. EPA, 2016b)



Proprietary information

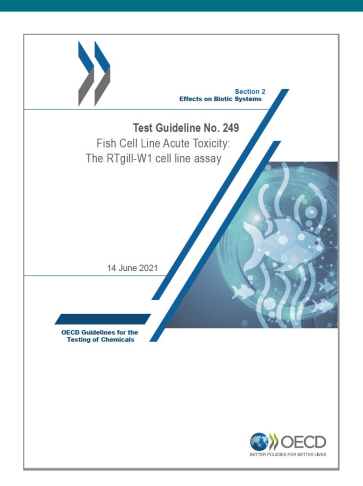
- A critical issue when trying to validate and accept in vitro and in silico methods for potential replacement of fish acute toxicity testing is the presence of proprietary information.
- As many commercially available assays were originally intended for pharmaceutical candidate screening, most are at least partially proprietary.
- This might interfere with standard validation approaches, and the development of specific testing guidelines.



Good news

OECD recently released a formal guidance document for the use of a 24-well plate formatted RTGill viability assay as part of:

- Predictor of acute fish toxicity
- Range-finding and pre-screening before conducting the acute fish toxicity test or other fish-based testing
- Part of a WoE for hazard assessment.





Potential strategies and suggestions

- To expedite the development and use of NAMs, ICCVAM established a generalized framework for regulators and stakeholders that is used to enable development and establish confidence in the use of NAMs through coordinated efforts that address three strategic goals:
 - Connect end users with the developers of NAMs
 - Foster the use of efficient, flexible, and robust practices to establish confidence in new methods
 - Encourage the adoption and use of new methods and approaches by U.S. federal agencies and regulated industries
- However, each federal Agency and program must evaluate NAM
 approaches in the context of its own regulatory needs to determine if it is
 fit for purpose and whether adequate environmental protection can be
 maintained using the new tools within their specific framework.



Acknowledgements

ICCVAM EcoWG

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