Ecotoxicology TP Course

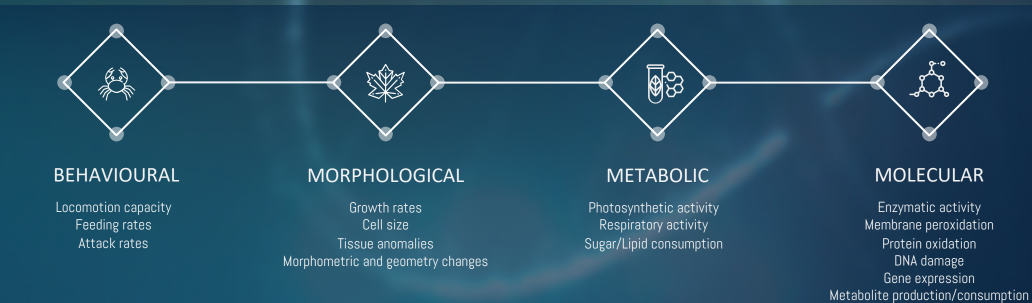
Concepts, Tests & Biomarkers

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PRATICAL COURSE

Activities

PRATICAL COURSE PROGRAM



PRATICAL COURSE PROGRAM

CELL DENSITY AND ECOTOXICOLOGICAL VARIABLES

Effects of the test substance in the growth traits and determination of the ecotoxicological doses.

PHOTOCHEMISTRY

Use of remote sensing optical techniques to evaluate the energetic metabolism of the cells under different concentrations of the test dose.

PIGMENTS

Quantification of the pigment profile of the cells as potencial source of biomarkers.

NON-ENZYMATIC BIOMARKERS

Determination of effect biomarkers such as lipid peroxidation or DNA damage

ENZYMATIC BIOMARKERS

Analysis of enzymatic activity of enzymes related to oxidative stress.

TESTS

Trial preparation, setups and typologies

01 TESTS: ECOTOXICOLOGY



ECOLOGY

Ecology is a branch of biology concerning **interactions among organisms and their biophysical environment**, which includes both biotic and abiotic components.

ECOTOXICOLOGY

The study of the effects of toxic chemicals on biological organisms, especially at the population, community, ecosystem, and biosphere levels. Ecotoxicology is a multidisciplinary field, which integrates toxicology and ecology.

In Ecotoxicology the concentration of the test substance in the target organisms should reflect the environmentally relevant or expected concentrations.

TOXICOLOGY

Toxicology is a scientific discipline, overlapping with biology, chemistry, pharmacology, and medicine, that involves **the study of the adverse effects of chemical substances on living organisms** and the practice of diagnosing and treating exposures to toxins and toxicants.

01 TESTS: Typologies

Typologies

- Acute: 1-4 days (at least 10% of the organism life cycle)
- Subacute: standard 28 days test
- Subchronic: standardized to 90 days
- **Chronic**: more than 90 days (should allow a complete life cycle)
- **Transgenerational**: Allows production of a new generation and evaluates the effects on the offspring.

ORGANIMS

- Bacteria
- Micro- and macro- algae
- Plants
- Invertebrates
- Fishes
- Mammals



Typology

The exposure typology should reflect the environmental exposure time typically observed or to ansewer to the target scientific question (for e.g. what is the effect at the reproduction level).

ORGANISM

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The selected organism should be cosmopolitan and representative of a certain group/environment.

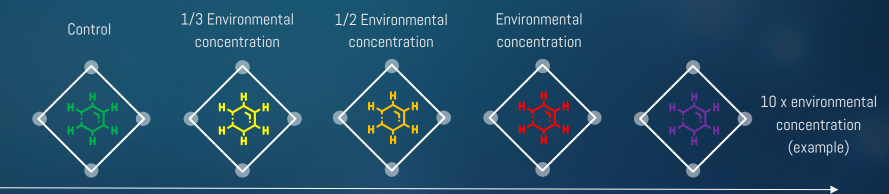
01 TESTS: DESIGN

INDIVIDUALS

- Clonal or with the same genetic background (lab cultured/maintained organisms);
- Similar age or life cycle stage;
- Similar sex (or grouped by sex if sex is a variable to analyse);
- Similar morphometric characteristics (height, weight, volume);
- Similar life history (maintained or reproduced under the same abiotic conditions);

DESIGN

- Control and test groups exposed simultaneously and under the same environmental conditions (light, temperature, etc);
- A consistent number of replicates must be ensure in all exposure mesocosmos;
- Increasing concentrations should follow a mathematical and logic succession or increase rate;



Increasing concentration

01 TESTS: MATERIAL PREPARATION

DECONTAMINATION

- In Ecotoxicology
 Decontamination refers to the
 cleaning of laboratory materials
 (glass and plasticware) from
 contaminants and chemical
 agents, avoiding these to
 contaminate the exposure trial.
- Typically uses acid or alkaline detergents, followed by acid bath and acetone (or other polar solvent) washing.
- Removes all adsorbed ions and molecules.

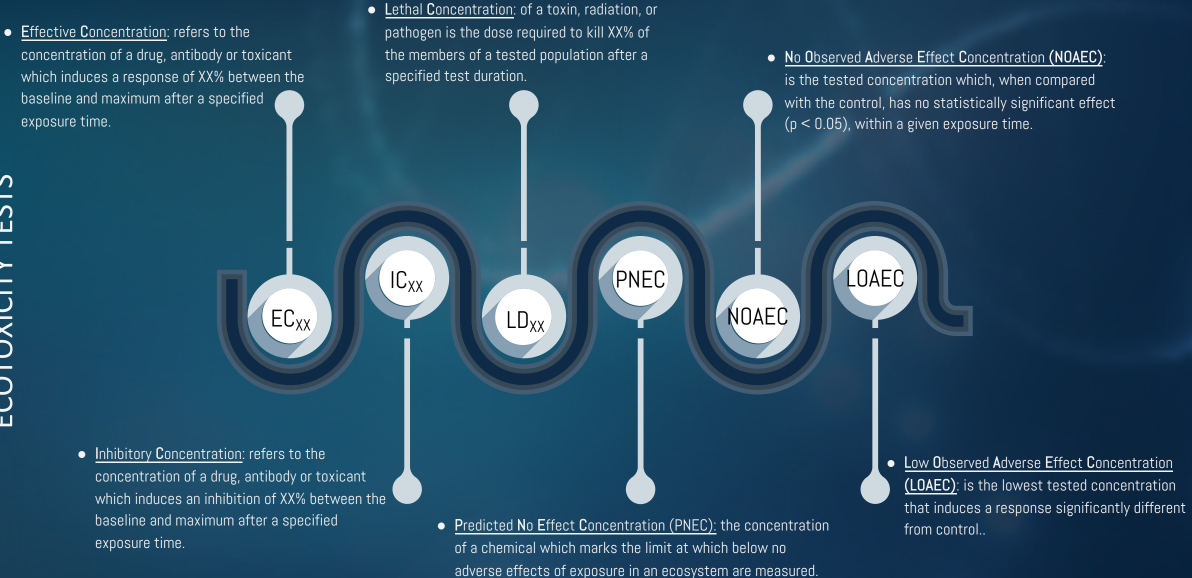
STYERILIZATION

- Any process that eliminates, removes, kills, or deactivates all forms of life.
- For most ecotoxicological trials is not a requirement although it is advisable.

PARAMETERIZATION

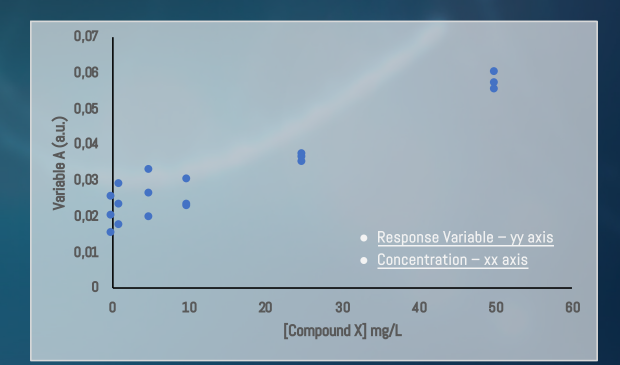
Ecotoxicity parameters, calculations and significance

02 PARAMETERIZATION

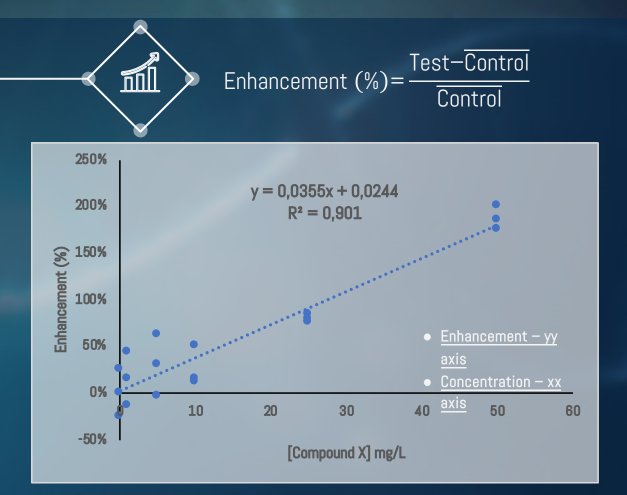


02 PARAMETERIZATION : Effective Concentration

[Compound X] mg/L	Variable A (a.u.)	
0	0,02	
0	0,015	
0	0,025	
1	0,023	
1	0,01725	
1	0,02875	
5	0,026	
5	0,0195	
5	0,0325	
10	0,03	
10	0,0225	
10	0,023	
25	0,035	
25	0,037	
25	0,036	
50	0,06	
50	0,057	
50	0,055	

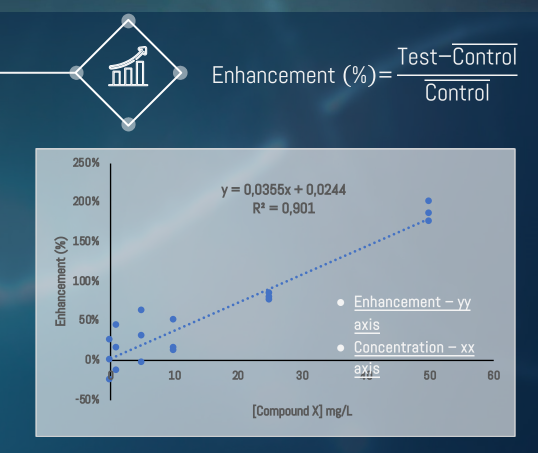


02 PARAMETERIZATION : Effective Concentration



[Compound X] mg/L	Variable A (a.u.)	Enhancement (%)
0	0,02	0%
0	0,015	-25%
0	0,025	25%
1	0,023	15%
1	0,01725	-14%
1	0,02875	44%
5	0,026	30%
5	0,0195	-3%
5	0,0325	63%
10	0,03	50%
10	0,0225	13%
10	0,023	15%
25	0,035	75%
25	0,037	85%
25	0,036	80%
50	0,06	200%
50	0,057	185%
50	0,055	175%

02 PARAMETERIZATION : Effective Concentration



Using the linear regression equation calculate the concentration at which the enhancement is 50% (EC_{50})

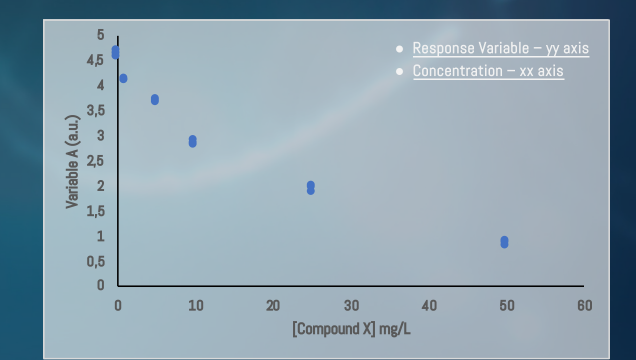
 $50\% = 0.0355x + 0.0244 \Leftrightarrow 0.5 = 0.0355x + 0.0244 \Leftrightarrow 0.5 - 0.0244 = 0.0355x$ x = 13.40 mg/L = EC₅₀

Upon the application of 13.40 mg/L the variable A suffers a 50% increase relative to the control.

[Compound X] mg/L	Variable A (a.u.)	Enhancement (%)
	0,02	0%
0	0,015	-25%
0	0,025	25%
1	0,023	15%
1	0,01725	-14%
1	0,02875	44%
5	0,026	30%
5	0,0195	-3%
5	0,0325	63%
10	0,03	50%
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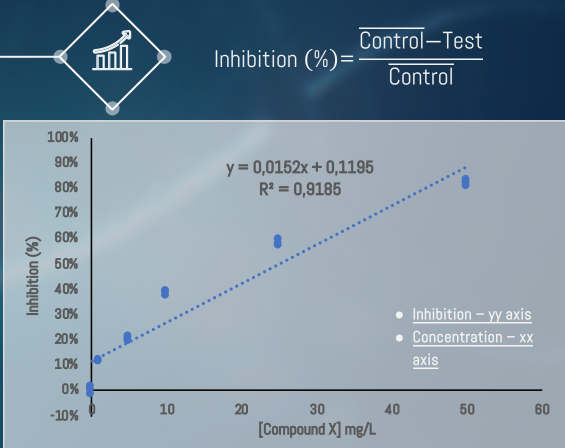
Concentration

[Compound X] mg/L	Variable A (a.u.)
0	4,69
0	4,57
0	4,63
1	4,08
1	4,11
1	4,095
5	3,72
5	3,65
5	3,685
10	2,89
10	2,81
10	2,85
25	1,99
25	1,87
25	1,95
50	0,9
50	0,79
50	0,845



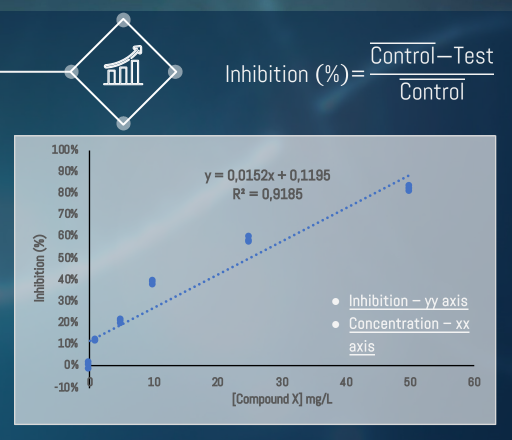
Concentration

[Compound X] mg/L	Variable A (a.u.)	Inhibition (%)		
0	4,69	-1%		
0	4,57	1%		
0	4,63	0%		100%
1	4,08	12%		90%
1	4,11	11%		80%
1	4,095	12%		70%
5	3,72	20%		S 60%
5	3,65	21%	the second se	(%) 20% 20%
5	3,685	20%		
10	2,89	38%		20%
10	2,81	39%		10%
10	2,85	38%		0%
25	1,99	57%		-10%
25	1,87	60%		
25	1,95	58%		
50	0,9	81%		
50	0,79	83%		
50	0,845	82%		



Concentration

[Compound X] mg/L	Variable A (a.u.)	Inhibition (%)	
0	0,02	0%	
0	0,015	-25%	
0	0,025	25%	
1	0,023	15%	
1	0,01725	-14%	
1	0,02875	44%	
5	0,026	30%	
5	0,0195	-3%	
5	0,0325	63%	
10	0,03	50%	
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25	0,037	85%	Using the linear re
25	0,036	80%	
50	0,06	200%	50% = 0.0152x +
50	0,057	185%	x = 13.40 mg/L =
50	0,055	175%	llnon the applicat



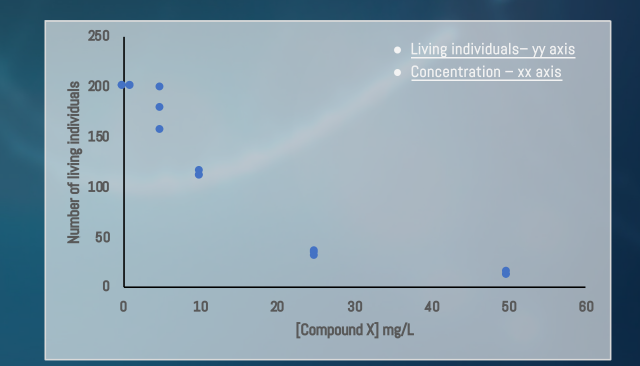
Using the linear regression equation calculate the concentration at which the inhibition was 50% (IC_{50})

 $50\% = 0.0152x + 0.1195 \Leftrightarrow 0.5 = 0.0152x + 0.1195 \Leftrightarrow 0.5 - 0.1195 = 0.0152x$ x = 13.40 mg/L = IC₅₀

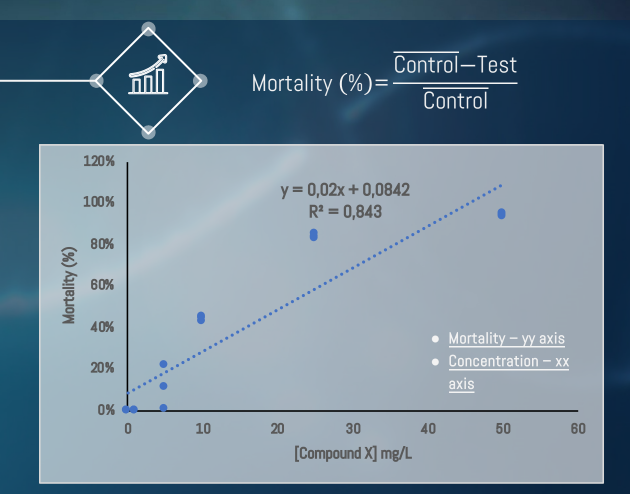
Upon the application of 13.40 mg/L the variable A suffers a 50% inhibition relative to the control.

02 PARAMETERIZATION : LETHAL Concentration

[Compound X] mg/L	Number of living individuals
0	200
0	200
0	200
1	200
1	200
1	200
5	156
5	178
5	198
10	110
10	115
10	111
25	30
25	33
25	35
50	11
50	12
50	14

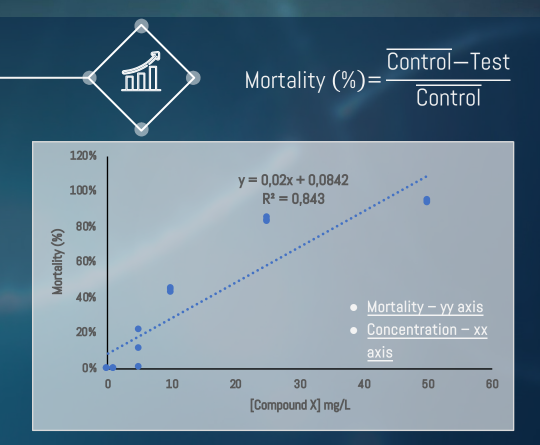


02 PARAMETERIZATION : LETHAL Concentration



[Compound X] mg/L	Variable A (a.u.)	Mortality (%)
0	4,69	0%
0	4,57	0%
0	4,63	0%
1	4,08	0%
1	4,11	0%
1	4,095	0%
5	3,72	22%
5	3,65	11%
5	3,685	1%
10	2,89	45%
10	2,81	43%
10	2,85	45%
25	1,99	85%
25	1,87	84%
25	1,95	83%
50	0,9	95%
50	0,79	94%
50	0,845	93%

02 PARAMETERIZATION : LETHAL Concentration



Using the linear regression equation calculate the concentration at which half the number of initial individuals is dead (LC_{50})

 $50\% = 0.02x + 0.0842 \Leftrightarrow 0.5 = 0.02x + 0.0842 \Leftrightarrow 0.5 - 0.0842 = 0.02x$ x = 15.79 mg/L = LC₅₀

Upon the application of 15.79 mg/L 50% of the individual die.

[Compound X] mg/L	Variable A (a.u.)	Mortality (%)
0	0,02	0%
0	0,015	-25%
0	0,025	25%
1	0,023	15%
1	0,01725	-14%
1	0,02875	44%
5	0,026	30%
5	0,0195	-3%
5	0,0325	63%
10	0,03	50%
10	0,0225	13%
10	0,023	15%
25	0,035	75%
25	0,037	85%
25	0,036	80%
50	0,06	200%
50	0,057	185%
50	0,055	175%

• EC, IC and LC refer to effective, inhibitory and lethal **concentration** respectively.

EC

IC

LC

- It refers to an external or exogenous concentration applied to a certain organism.
- Expressed as a mass per volume unit.

• ED, ID and LD refer to effective, inhibitory and lethal **dose** respectively.

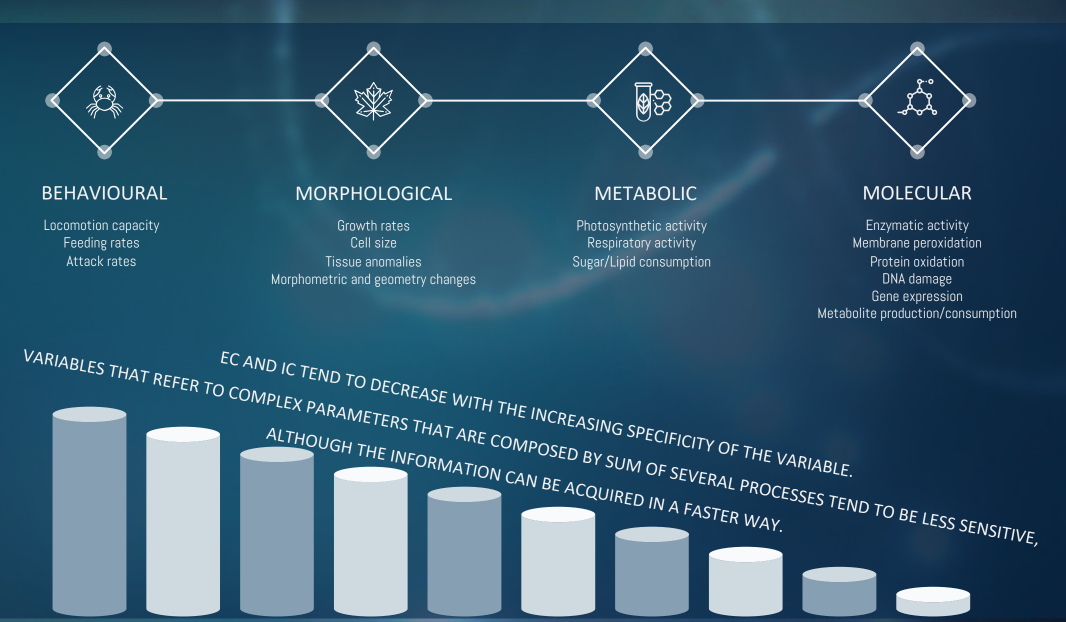
ED

ID

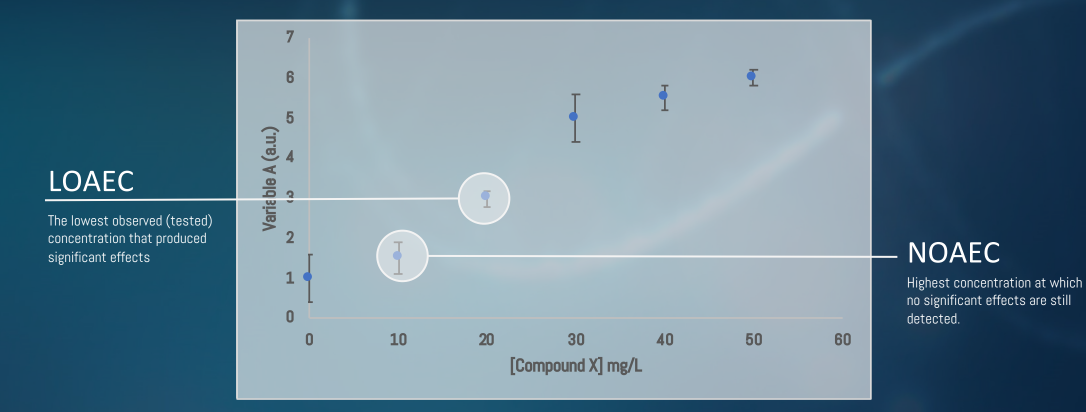
LD

- It refers to measured concentration inside an organism or to an external or exogenous concentration applied to a certain organism calculated proportionally to the organism body mass
- Expressed as mass of target substance per organism fresh weight.

02 PARAMETERIZATION : VARIABLE TYPOLOGY



02 PARAMETERIZATION : NOAEC & LOAEC



NOAEC/LOAEC versus IC/EC/LC

- NOAEC and LOAEC depend on the concentrations tested and defined by the user; if the range of concentrations tested has a low resolution power the NOAEC and LOAEC assessed can be deceiving.
- IC, EC and LC are obtained by linear regression analysis and thus even if the concentration correspondent to each of these parameters was not tested, it can be calculated.

PREDICTED NO EFFECT CONCENTRATION (PNEC):

- The concentration of a chemical which marks the limit at which below no adverse effects of exposure in an ecosystem are measured.
- Conservative values and predict the concentration at which a chemical will likely have no toxic effect.
- Do not intended to predict the upper limit of concentration of a chemical that has a toxic effect
- PNEC values are often used in environmental risk assessment as a tool in ecotoxicology.
- A PNEC for a chemical can be calculated with acute toxicity or chronic toxicity single-species data, species sensitivity distribution (SSD) multi-species data, field data or model ecosystems data. depending of the type of data used, an **ASSESSMENT FACTOR** is used to account for the confidence of the toxicity data being extrapolated to an entire ecosystem.

Available test result	Assessment factor
One long-term test (NOEC or EC10)	100
Two long-term tests (NOEC or EC10) with species representing different living and feeding conditions	50
Three long-term tests (NOEC or EC10) with species representing different living and feeding conditions	10

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- The concentration of a chemical which marks the limit at which below no adverse effects of exposure in an ecosystem are measured.
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- The ASSESSMENT FACTOR can be determined by the available bibliography or pre-defined taking into account the type of assay.

	Available test result	Assessment factor
⊆z≻	One long-term test (NOEC or EC10)	100
RAPH ATIC BILIT	Two long-term tests (NOEC or EC10) with species representing different living and feeding conditions	50
BIBLIOGRAPHIC INFORMATION AVAILABILITY	Three long-term tests (NOEC or EC10) with species representing different living and feeding conditions	10
BIBL INF(AV	Available test result	Assessment factor
PRE-DEFINED BY THE TEST TYPE	Acute Toxicity Data	The lowest LC50 in the compiled database is then divided by the assessment factor to calculate the PNEC for that data. The assessment factor applied to acute toxicity data is typically 1000.
	Chronic Toxicity Data	The lowest NOEC value in the test dataset is divided by an assessment factor between 10 and 100 dependent on the diversity of test organisms and the amount of data available. If there are more species or data, the assessment factor is lower.