A *Lymnaea stagnalis* Embryo Test for Toxicity Bioindication of Acidification and Ammonia Pollution in Water

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Abstract: The paper presents a study leading to a new acute toxicity test on embryonic and juvenile organisms of the great pond snail (*Lymnaea stagnalis* Linnaeus). Sulfuric acid, nitric acid, and ammonium hydroxide were used as waterborne toxicants in laboratory experiments. The exposure time was 24 h. Tests were conducted in 5–10 replications for each toxicant. The toxicity of the substances was classified according to different scales and the test’s sensitivity was compared to that of the commonly used bioindicator *Daphnia magna* Straus. The assessment of toxicity impact was supported by microscopic observations. The probit method was used as a parametric statistical procedure to estimate LC50 and the associated 95% confidence interval. Our study showed that the early developmental stages of *Lymnaea stagnalis* are very sensitive bioindicators, making it possible to detect even very low levels of the above-mentioned water toxicants. The highest toxicity is shown by ammonium hydroxide with LC50/24h values, respectively, 24.27 for embryos and 24.72 for juvenile forms, and the lowest is shown by nitric acid ions with LC50/24h values, respectively, 105.19 for embryos and 170.47 for juvenile forms. It is highly cost-effective due to simple and efficient breeding and the small size of the organisms in the bioassay population. Compared with *Daphnia magna*, relatively low concentrations of toxicants caused a lethal effect on embryonic and juvenile organisms of the great pond snail. Owing to their common occurrence and sensitivity, early developmental forms of *Lymnaea stagnalis* can be a valuable new tool in biomonitoring of the freshwater environment.

Keywords: acute toxicity test; bioassay; bioindicators; water pollution; aquatic molluscs

1. Introduction

Acidification and ammonia contamination of water resources are currently recognized as serious environmental problems. Water acidification, in most cases, is associated with anthropogenic emissions of sulfur and nitrogen oxides and is addressed from a global perspective [1]. Ammonia content in water is increasingly being recognized as a pollutant of key environmental concern. This toxicant contributes to several environmental problems, including direct toxic effects on water communities leading to eutrophication and acidification of sensitive ecosystems, which affect the health of human beings as the final consumers [2]. Run-off and atmospheric emissions of this toxicant are increasing rapidly in
many parts of the world, so these environmental concerns are expected to intensify in the future [1,2].
The effects on aquatic ecosystems are difficult to predict and highly variable, depending on many
factors. The use of appropriate bioindication methods facilitates more comprehensive determination
of their negative impact on the aquatic community.

The effect of acidification has not been universally recognized as dangerous [3,4]; however, the
unfavorable environmental effects of acid rain on surface waters have recently been described in
a comprehensive manner by many authors [5–11]. Reports of the growing threat associated with
the environmental effects of acid deposition have focused mainly on monitoring of changes in
freshwater communities. The negative impact of acidification on different aquatic organisms—plants
and phytoplankton (producers), invertebrates, and vertebrates—has been widely reported [4,12–22].
All of these studies focused on the analysis of aquatic communities and transformations in populations
due to the decrease in pH caused by acidification of waters from various sources.

Rapid pH changes and significant content of NOx, as well as SO\textsubscript{4}^{2–} and SO\textsubscript{3}^{2–}, may lead to
the extinction of many aquatic communities. Depending on pH, the rate of withdrawal of different
species varies and additionally depends on their tolerance to environmental stress factors, as well as
their developmental stage. Ubiquitous species show tolerance for low pH values and can survive;
however, a decrease in biodiversity can be seen in ecosystems where highly-tolerant organisms are
predominant [23–25].

Pollution with ammonia compounds entails the opposite of acidification, i.e., alkalization. It poses
a threat because it is highly toxic for most aquatic organisms and is unfortunately quite common, often
coming from anthropogenic sources.

At increased concentrations ammonia halts the development and reproduction of various species.
It strongly affects internal organs. At low concentrations ammonia ions increase eutrophication of
water, and acts as a source of nitrogen for algae [26–29].

The physiological effects of acidification and alkalization on water organisms have been
extensively studied in various research centers. Among the toxic effects of decreased pH, at least four
main physiological functions are disturbed by high concentrations of H\textsuperscript{+} ions: (Figure 1).

1. Calcium regulation: very important, particularly for invertebrates—crustaceans and mollusks.
   At low pH calcium uptake is particularly difficult, which slows down and disturbs the formation
   of the outer skeleton [19,30].
2. Sodium regulation: vital for most aquatic animals because it plays a key role in ion balance and
   osmoregulation processes [4].
3. Respiration: particularly at the cellular level, pH change inhibits galactosidase. By applying
   enzymatic tests using fluorescence, we can evaluate the degree of disturbance [4,19].
4. Acid-base balance: closely related to other processes, its disturbance can secondarily disturb the
   basic physiological functions of organisms [4,19].

Changes in pH are also linked to toxicity of trace metals, which depending on pH may become
more bioavailable and, thus, dangerous for organisms. A slight variation in pH is not likely to have a
direct impact on life in a water ecosystem, but significantly influences the availability and solubility of
all chemical forms in fresh water and may aggravate nutrient problems [25,31,32]. At pH = 5 a high
concentration of aluminum in the water becomes toxic for many invertebrates and fish by disturbing
gill function [33]. A group of aquatic mollusks belongs to organisms particularly sensitive to pH
changes, mainly its fall below 7. Surface waters of low pH are also characterized with low calcium
concentrations. Tolerance of various mollusk species to pH in acidic waters is changeable and ranges
from 6.5 to 4.0. On average, in water of pH below 5.4 populations of aquatic snails die out as a result
of reproduction disturbance and disturbance in shell development. Temporary (short lasting) pH
changes can be tolerated in a wider range—up to 3.5 [19].
Adult developmental stages are more tolerant to decreases in pH than embryonic and juvenile forms. In some cases embryos develop in eggs, sometimes additionally covered with cocoons; however, their environment is rapidly altered and acidified during hatching. The shock causes high mortality in larvae and juveniles [34,35]. Various amphibian species are particularly sensitive to acidification of water bodies; thus, at the embryonic stage, an increased mortality rate is observed as a result of fungal infestation of eggs and a lack of oxygen, as well as disturbed osmoregulation [36]. A pH level of 5–6 or lower has been considered to be directly toxic to fish [37].

Water acidification and ammonia contamination are among the most common forms of water pollution worldwide. Their concentrations are variable in surface waters and their negative effects on water ecosystems are difficult to predict. Hence, new sensitive bioindication methods are needed. We have conducted a study on the use of the aquatic invertebrate *Lymnaea stagnalis* as a widely-distributed organism in freshwater ecosystems all over the world. The main objective of the study was to develop innovative bioindication methods which can be widely introduced in water biomonitoring. We proposed the use of embryonic and early juvenile stages of *Lymnaea stagnalis*. This method was applied in this study for bioindication of both acidification and ammonia contamination in water. The sulfuric and nitric acid solutions selected for testing are major acid rain components [38,39].

2. Materials and Methods

2.1. Selected Bioindicators

Juvenile and embryonic forms of *Lymnaea stagnalis* were selected for the analyses, and parallel tests on *Daphnia magna* were conducted for the same pollutants (Table 1). The response of *Lymnaea stagnalis* was compared with that of *Daphnia magna*.

*Lymnaea stagnalis*, the great pond snail (Figure 2), is a cosmopolitan species, common in all types of freshwater bodies [40]. The snails were obtained from breeding stock in small ponds. Under natural conditions they are found in ponds and lakes and are very common in our climate zone. Full embryonic development in this species takes place in eggs laid in cocoons on leaves or other
surfaces. In its early developmental stages (embryos and hatchlings) *Lymnaea stagnalis* is sensitive to changes in the chemical conditions of the water. Thus, these stages make it a good bioindicator for many chemical substances [40]. In the first stage of the experiment the specimens were adapted to laboratory conditions, until the third generation. After the adaptation period pilot tests were performed for a specific group of toxins.

**Table 1. Lymnaea stagnalis and Daphnia magna classification.**

| Kingdom   | Animalia          | Animalia          |
|-----------|-------------------|-------------------|
| Phylum    | Mollusca          | Arthropoda        |
| Subphylum | -                 | Crustacea         |
| Class:    | Gastropoda        |                   |
| (unranked): | clade Heterobranchia | Branchiopoda     |
|           | clade Euthyneura  |                   |
|           | clade Panpulmonata|                   |
|           | clade Hygrophila  |                   |
| Order:    | Hygrophila        | Cladocera         |
| Superfamily: | Lymnaeoidea  | -                 |
| Family:   | Lymnaeidae        | Daphniidae        |
| Subfamily: | Lymnaeinae      | -                 |
| Genus:    | Lymnaea           | Daphnia           |

**Figure 2. Lymnaea stagnalis.** (a) Juveniles (2–3 mm long); (b) adult (3–5 cm); and (c) a cocoon with embryos developing in the eggs (length of cocoon: 1–4 cm; diameter of egg: around 1 mm).

*Daphnia magna* is the most widely used organism in water bioindication. The selected forms of *Daphnia magna* were juvenile, up to the age of 24 h, with no signs of morbidity or developmental abnormalities.

2.2. Culture Techniques and Experimental Procedures on Lymnaea stagnalis

A laboratory culture of the *Lymnaea stagnalis* population was conducted at the Laboratory of Environmental Biotechnology and Ecology, AGH University of Science and Technology in Krakow. Adult forms of *Lymnaea stagnalis* were kept in covered rectangular 21-L aquariums. Constant parameters were maintained in the snail culture.

Water temperature ranged from 22 °C to 18 °C. Each aquarium was supplied with continuously aerated de-chlorinated tap water filtered through an active carbon reactor.

The pH value was about 7.0 ± 0.5. The light period was controlled (12 h light and 12 h darkness).

The snails were fed green lettuce leaves every three days. Each aquarium was cleaned by siphoning every week to remove feces. All of the snails in the tanks exhibited high reproductive potential. Adult snails laid their cocoons with eggs, attached to the walls of the aquarium.
The average number of eggs laid in a cocoon was 70 ± 30, but only cocoons with 60–90 eggs were selected to experiment. All of the embryos in cocoons selected for tests were in good condition. Selection was preceded by microscopic evaluation of vital parameters. Unfertilized eggs or those with immobilized embryos were removed.

Sulfuric acid (VI), nitric acid (V), and ammonium hydroxide were applied as toxicants. The concentration of toxicants covered a wide range, from lethal (100% mortality) to total survival (100% survival) over 24 h.

**Embryo toxicity bioassay** procedure: snail eggs were collected and cocoon envelopes were macerated to improve penetration of water with toxins to the inside, but no egg was damaged. Mechanically pre-treated embryos in the cocoons were immersed in the toxicants and exposed to them for 24 h. It took about 15–20 s to open cocoons, but the entire maceration operation was provided in the tested solution, so no other factors have an impact on the developed embryos. For each experimental population, control group, and suitable number of concentrations of toxicant solution were used to determine EC50 (Tables 2–4) and the experiments were repeated in five replications for each toxicant concentration.

**Table 2.** Mortality of tested organisms in a sulfuric acid solution (24 h) (AVG ± SD).

| Stock Solution H₂SO₄ (mg dm⁻³) | pH   | Lymnaea stagnalis Embryos |  | Lymnaea stagnalis Hatchlings |  | Daphnia magna Neonates |  |
|-------------------------------|------|---------------------------|---|------------------------------|---|------------------------|---|
|                               |      | Mortality (%) | SD (%) | Mortality (%) | SD (%) | Mortality (%) | SD (%) |
| 288.1 2.23 ± 0.02 |  | 100.00 | 0.00 | 100.00 | 0.00 | 100.00 | 0.00 |
| 228.8 2.33 ± 0.05 |  | 100.00 | 0.00 | 100.00 | 0.00 | 100.00 | 0.00 |
| 183 2.43 ± 0.03 |  | 100.00 | 0.00 | 100.00 | 0.00 | 100.00 | 0.00 |
| 146 2.53 ± 0.04 |  | 100.00 | 0.00 | 74.07 | 0.07 | 62.82 | 0.08 |
| 117 2.62 ± 0.02 |  | 100.00 | 0.00 | 37.82 | 0.06 | 21.88 | 0.04 |
| 93 2.72 ± 0.04 |  | 100.00 | 0.00 | 19.07 | 0.04 | 14.07 | 0.04 |
| 75 2.82 ± 0.06 |  | 41.90 | 1.15 | 6.88 | 0.03 | 8.75 | 0.02 |
| 60 2.91 ± 0.03 |  | 6.20 | 1.18 | 3.13 | 0.03 | 7.82 | 0.03 |
| 48 3.01 ± 0.02 |  | 1.60 | 1.11 | 1.25 | 0.02 | 3.44 | 0.03 |
| 38.4 3.11 ± 0.04 |  | 0.20 | 2.18 | 0.00 | 0.00 | 0.00 | 0.00 |
| 3.0 3.21 ± 0.04 |  | 0.20 | 2.11 | 0.63 | 0.01 | 0.31 | 0.01 |
| 24 3.31 ± 0.03 |  | 0.20 | 0.00 | 0.31 | 0.01 | 0.31 | 0.01 |
| Control 7.00 ± 0.01 |  | 0.30 | 0.00 | 0.00 | 0.00 | 0.31 | 0.01 |

**Table 3.** Mortality of tested organisms in a nitric acid solution (24 h) (AVG ± SD).

| Stock Solution HNO₃ (mg dm⁻³) | pH   | Lymnaea stagnalis Embryos |  | Lymnaea stagnalis Hatchlings |  | Daphnia magna Neonates |  |
|-------------------------------|------|---------------------------|---|------------------------------|---|------------------------|---|
|                               |      | Mortality (%) | SD (%) | Mortality (%) | SD (%) | Mortality (%) | SD (%) |
| 228 2.44 ± 0.03 |  | 100.00 | 0.00 | 100.00 | 0.00 | 100.00 | 0.00 |
| 182 2.54 ± 0.05 |  | 89.30 | 1.98 | 61.57 | 0.05 | 32.82 | 0.06 |
| 145 2.64 ± 0.02 |  | 61.70 | 1.63 | 27.19 | 0.05 | 5.63 | 0.02 |
| 116 2.73 ± 0.03 |  | 33.00 | 1.68 | 4.69 | 0.02 | 0.31 | 0.01 |
| 93.3 2.83 ± 0.06 |  | 8.10 | 0.93 | 0.63 | 0.01 | 0.31 | 0.01 |
| 74.7 2.93 ± 0.04 |  | 4.00 | 1.05 | 0.00 | 0.00 | 0.00 | 0.00 |
| 59.7 3.02 ± 0.05 |  | 0.40 | 0.61 | 0.00 | 0.00 | 0.63 | 0.01 |
| Control 7.00 ± 0.02 |  | 0.50 | 1.21 | 0.31 | 0.01 | 0.31 | 0.01 |

**Juvenile toxicity bioassay** (after hatching): young individuals were collected and immersed in toxicant solutions in 100 mL plastic containers, 32 specimens to each polystyrene container. The exposure time was 24 h. The toxicity test algorithm was similar as in the case of Daphnia magna.
Table 4. Mortality of tested organisms in an ammonium hydroxide (24 h) (AVG ± SD).

| Stock Solution NH₄OH (mg dm⁻³) | pH     | Lymnaea stagnalis Embryos | Lymnaea stagnalis Hatchlings | Daphnia magna Neonates |
|-------------------------------|--------|--------------------------|----------------------------|------------------------|
|                               |        | Mortality (%) SD (%)     | Mortality (%) SD (%)        | Mortality (%) SD (%)   |
| 91.5                          | 11.42 ± 0.04 | 100.00 0.00            | 100.00 0.00               | 100.00 0.00           |
| 73.21                         | 11.32 ± 0.05 | 100.00 0.00            | 100.00 0.00               | 100.00 0.00           |
| 58.5                          | 11.22 ± 0.04 | 100.00 0.00            | 100.00 0.00               | 100.00 0.00           |
| 46.8                          | 11.13 ± 0.03 | 100.00 0.00            | 100.00 0.00               | 100.00 0.00           |
| 37.5                          | 11.03 ± 0.04 | 94.33 1.44             | 100.00 0.00               | 100.00 0.00           |
| 30                            | 10.93 ± 0.07 | 78.5 1.87              | 97.50 2.46                | 92.82 3.62            |
| 25                            | 10.85 ± 0.03 | 40.49 5.23             | 45.94 7.66                | 30.63 6.72            |
| 18                            | 10.71 ± 0.02 | 18.11 12.43            | 6.25 2.55                 | 1.25 1.62             |
| 14.4                          | 10.61 ± 0.04 | 4.64 1.44              | 0.63 1.32                 | 0.31 0.99             |
| Control                       | 7.00 ± 0.02 | 0.51 1.14              | 0.31 0.99                 | 0.31 0.99             |

2.3. Culture Techniques and Experimental Procedures on Daphnia magna

*Daphnia* were cultured in glass tanks filled with 2 L of aerated, reconstituted, moderately hard water. The colony was maintained at 21–23 °C in a chamber with a photoperiod of 12 h light/12 h dark. *Daphnia* were fed 10 mL of *Scenedesmus* sp. algae (with a density between 32–36 × 10⁶ cell ml⁻¹ of algae). Neonates were separated from adults based on their size (<1.5 mm) by sieving. The bioassays were conducted in 120 mL polystyrene containers, with 100 mL of toxicant solution in each. Thirty-two neonates were transferred to containers containing different concentrations of the test pollutants. There was no feeding and no aeration during the tests. In OECD standards it is recommended to apply at least five, but generally 10, organisms per test vessel, and there are no obstacles to use 32 organisms for better statistical analysis. At least 2 mL of test solution should be provided for each animal (i.e., a volume of 10 mL for five daphnids per test vessel). In our experiment, for each individual, more than 3 mL test solution was used.

Dead individuals were identified visually after 24 h. For each tested population, control group and suitable number of toxicant solution concentrations were used to determine EC₅₀ and the experiments were repeated in 10 replications for each toxicant concentration [41,42].

The parameters of the water used in the preparation of experimental solutions are enclosed in Appendix Table A1.

2.4. Evaluation of Toxicity Tests and Statistical Analysis

The toxicity of the tested substances was classified according to three widely used scales: Dockal and Sold [43], European Union, and US EPA (The Environmental Protection Agency). The assessment of toxicity impact was supported by microscopic observations, using a Nikon Eclipse 1200 Research Light Microscope (LM) with a digital camera (Nikon Instruments Europe BV, Amsterdam, The Netherlands).

The STATISTICA software (version. 12, StatSoft Polska, Krakow, Poland) package was used to analyze the data. The test applied was ANOVA; two or more of the observed proportion mortalities had to be between zero and one. The probit method was used as a parametric statistical procedure to estimate LC₅₀ and the associated 95% confidence interval [44]. I was assumed a standard level of significance 0.05 to determine statistical significance.

3. Results

3.1. Results of Acute Toxicity Tests

The 24 h acute toxicity tests were conducted to assess the response of the species studied to acidification and ammonium hydroxide. Mortality of *Lymnaea stagnalis* (embryonic and juvenile forms) in comparison to *Daphnia magna* (neonates) was tested for different concentrations of sulfuric acid (Table 2), nitric acid (Table 3), and ammonium hydroxide solution (Table 4). These tables show concentrations causing a total lethal effect (LC₁₀₀, mortality rate of 100%) and concentrations which
exhibited no toxic effect within 24 h (mortality at the control level). Sensitivity of *Daphnia magna* was generally lower than that of *Lymnaea stagnalis* subjected to the stress of all of the tested substances. Nevertheless, the rank of toxicity of the three substances was analogous for *Daphnia magna* and *Lymnaea stagnalis*; the most toxic of the three was ammonium hydroxide, followed by sulfuric acid and nitric acid (Tables 2–4).

The LC$_{50}$ values confirmed that ammonium hydroxide was the most toxic substance for *Lymnaea stagnalis*; the LC$_{50}$/24 h was 24.27 mg·dm$^{-3}$ (pH = 10.84) for embryos and 24.72 (pH = 10.85) for juveniles (Tables 5–7, Figure 3). The toxicity of both acids was lower for both forms of *Lymnaea stagnalis*. Embryonic forms were more sensitive than juvenile ones. The LC$_{50}$/24 h for *Lymnaea stagnalis* embryos was 81.17 (pH = 2.78) mg·dm$^{-3}$ in the sulfuric acid solution and 105.19 mg·dm$^{-3}$ (pH = 2.78) in the nitric acid solution (Table 5). For juvenile *Lymnaea stagnalis* the LC$_{50}$/24 h of the sulfuric acid was 126.49 mg·dm$^{-3}$ (pH = 2.59) and that of the nitric acid was 170.47 mg·dm$^{-3}$ (pH = 2.57) (Table 6).

### Table 5. Assessment of acute toxicity of pollutants on *Lymnaea stagnalis* embryos (LC$_{50}$/24 h) *Lymnaea stagnalis* juveniles after hatching.

| Substance | LC$_{50}$ Value (*Lymnaea stagnalis* Embryos) Mean ± SD | Toxicity Class According to: |
|-----------|----------------------------------------------------|------------------------------|
|           |                                                    | Dockal and Sold | European Union | US EPA          |
| H$_2$SO$_4$ | 81.17 ± 0.76 mg·dm$^{-3}$ | Average toxicity | Harmful | Moderate toxicity |
| HNO$_3$ | 105.19 ± 1.54 mg·dm$^{-3}$ | Low toxicity | - | Low toxicity |
| NH$_4$OH | 24.27 ± 0.68 mg·dm$^{-3}$ | Average toxicity | Harmful | Moderate toxicity |
Daphnia magna

All of the bioindicators tested showed a similar reaction leading to disintegration and denaturation (Tables 5 and 6) and compared with Daphnia magna.

3.2. Comparison of the Sensitivity of the Organisms to the Toxicants

The results of one-way analysis of variance did not allow us to reject the hypothesis that there are no differences between the organisms tested exposed to the substances applied (Table 8). All of the organisms showed similar degrees of sensitivity to the substances. Thus, they can be used interchangeably in toxicity tests for these substances.
Post hoc test—Tukey’s HSD test (Honest Significant Difference); significance of differences between test organisms was assessed (Table 5).

The post hoc (Tukey’s) tests confirmed that all the bioindicators responded similarly to the substances tested. The statistical analysis revealed no statistically significant differences between them (Table 9).

**Table 9. Results of Tukey’s HSD test for significance of differences between the organisms tested.**

| Bioindicators   | *L. stagnalis* (Juveniles) | *L. stagnalis* (Embryos) | *D. magna* |
|-----------------|-----------------------------|---------------------------|------------|
| *L. stagnalis* (juveniles) | -                           | 0.49                      | 0.78       |
| *L. stagnalis* (embryos)    | 0.49                        | -                         | 0.19       |
| *D. magna*             | 0.78                        | 0.19                      | -          |

**4. Discussion**

The completed experiment using developmental stages of *Lymnaea stagnalis* revealed their high potential as a new bioassay for common water pollutants. Recent decades have seen continual progress in the testing of biomonitoring methods (especially acute toxicity tests) for surface water samples. Nevertheless, use of these methods in practice is still limited [45-50] *Lymnaea stagnalis* is a widespread species found throughout Europe, Northern Asia and North America [41]. It is noteworthy that this species can be found in still or slow-moving waters [51,52]. Features such as its common occurrence and tolerance for different types of habitats suggest that *Lymnaea stagnalis* is exceptionally well-suited for environmental monitoring.

The proposed new toxicity bioassay can be widely used to evaluate water degradation—both acidification and alkalization. As *Lymnaea stagnalis* showed great sensitivity to acid, it can be extensively used as a bioindicator in toxicological examination of water with low pH. Its response to acidification is also pronounced, and it is more sensitive to a sulfuric acid solution than to nitric acid. The value of pH was additionally measured by the parameters, together with the concentration of used toxicants, so the primary toxicity action is strongly dependent on the ion form of the used toxicant and pH value. For the
used chemical compounds, HNO$_3$ pKa = −2, Ka = 100%, H$_2$SO$_4$ pKa = −6.62, Ka = 4.17 $\times$ 10$^6$, and for NH$_4$OH is 100% ionization in water. This means that almost 100% of the molecule is broken down into its constituent ions. We observed toxic action of these ions depending on their concentrations, and the rate of pH indirectly affects osmosis. Hence, it can be widely used for biomonitoring of surface water quality in areas threatened by acid deposition from anthropogenic sources, caused by release of NO$_x$ and SO$_2$ into the atmosphere. These emissions are currently regarded as one of the most widespread global threats, and a major implication of them is acidification of the aquatic environment [38,39].

The use of early developmental stages of Lymnaea stagnalis increases the choice of acute toxicity bioassay methods for quick and quantified assessment of surface water quality. The most important symptoms of toxicity include delayed development in low concentrations of the water pollutants tested. This process has been observed before, e.g., [43,53,54] stress caused by environmental pollution reduces the energy budget of the organism. Exposed organisms spend a relatively large portion of their energy resources on decontamination and defense against the toxic influence of pollutants. In this situation the amount of energy that can be spent on growth and development is reduced. Higher concentrations of all the tested toxins have a similar influence on aquatic organisms. The aggressive action of strong acids and hydroxides on biochemical structures causes protein denaturation. Changes in the properties of cellular membranes occurring at high concentrations of these substances disintegrate the plasmalemma. The final effect of stress includes damage to the organisms and their death [19]. The great pond snail is a very sensitive bioindicator, facilitating detection of even very low levels of environmental stress factors. Compared with Daphnia magna, relatively low concentrations of toxicants caused a lethal effect in early developmental stages of Lymnaea stagnalis. In an increased range of concentrations of acid anions and ammonia cations, the process of immobilization is very rapid, with individuals dying within 10–45 min. After 24 h very severe disintegration and destruction can be observed.

Toxic action can also be detected as changes in the surface morphology of Lymnaea stagnalis that can be observed microscopically (Figure 6). Maceration of cocoon envelopes accelerates the direct exposure of embryos developing in eggs to selected toxicants. The cocoon envelopes reduce embryos exposed to selected water xenobiotics. They stabilize the internal environment of developing embryos. Proper maceration technique by scalpel eliminates the cocoon’s barrier effect and increases bioassay sensitivity. All of the above-mentioned substances act aggressively and cause serious damage in the form of progressive tissue disintegration. Damage caused by acids, and that caused by ammonia, are distinguishable by their specific appearance, and can be distinguished by microscopic observations from the effects of other substances, such as detergents [43]. The response of immobilized organisms (both snails and daphnids) during the experiments was very characteristic. Their bodies became opaquely white, and juvenile forms showed progressive disintegration of tissues (depending on the concentration and exposure time).

The experiment showed that Lymnaea stagnalis has numerous advantages as a bioindicator, during both embryonic and juvenile development. The embryonic form of this snail is more sensitive to pollution than the juvenile form. Scientific evidence clearly indicates the need to use embryonic stages and juveniles of various aquatic organisms after hatching in water bioindication, as highly-sensitive bioindicators of the toxicity of various ecotoxins [55].

It is noteworthy that Lymnaea stagnalis showed an analogous response to that of Daphnia magna. The Daphnia magna bioassay is one of the most commonly used bioassays worldwide for acute toxicity testing of various chemical substances [56]. Moreover, toxicity values between Daphnia and many other aquatic organisms are known to be strongly correlated, especially in the case of fish. Toxicity evaluation of the toxicants discussed above can be used interchangeably with other bioassays or together in a battery of bioassays for comprehensive bioindication [57].

Laboratory studies have shown that Lymnaea stagnalis (embryonic and juvenile forms (immediately after hatching) is a good bioindicator fulfilling the 5R criterion [58]: it is relevant, reliable, robust, responsive, and reproducible. The bioassay with embryos and juveniles of Lymnaea stagnalis gives very similar results, and we can base on them in the interpretation of toxic effects caused by water toxicants.
Moreover, our results are an answer to the long search for sensitive and low-cost bioassays for testing surface water toxicity [59,60]. Since *Lymnaea stagnalis* is common and easy to breed and raise, its use is not costly. We plan to conduct further tests on this organism to develop a detailed technique for its use in laboratories carrying out environmental monitoring.

The research on the new objective and sensitive biotest with the application of embryonic and juvenile *Lymnaea stagnalis* [61] continues. In the future it will be validated and directives will be established for the laboratory procedures of the biotest (OECD, ISO etc.).

Moreover, the bioassays use embryonic and juvenile forms which are small. The bioassay can be performed even in small containers (120 mL) on a large group of individuals, enabling statistical testing with 3–5 organisms per 10 mL of test solution. Due to their high breeding rate, *Lymnaea stagnalis* specimens are available for laboratory bioassays throughout the year.

The proposed organisms can be used in bioassay batteries to detect environmental effects of selected chemicals and their mixtures. This approach is recommended by the US EPA and APHA (American Public Health Association) [62], as well as the EU. The biotests presented in this study can be applied to examine water samples both taken from the water bodies suspected of acidification or ammonia intoxication, as well as in testing the quality of treated wastewater [53,54]. In water monitoring one of the most important criteria is biological assessment. Thus, the biotests designed here can be included into the system of biomonitoring as a tool for better assessment of water quality.

5. Conclusions

*Lymnaea stagnalis* is a very sensitive bioindicator capable of detecting even very low levels of environmental stressors. The test procedures are simple and inexpensive, and make it possible to detect negative effects of both acidification and alkalization. These organisms can be used in a battery of bioassays.

The experiment revealed that *Lymnaea stagnalis*, during both embryonic and juvenile development, has numerous advantages as a bioindicator.

*Lymnaea stagnalis* shows a similar response to that of *Daphnia magna*, with the values of LC50 were on a similar levels.
The toxic action can also be detected as changes in the surface morphology of Lymnaea stagnalis which can be observed microscopically.

The study showed that the early developmental stages of Lymnaea stagnalis are good bioindicators that may be used in monitoring surface water quality. Lymnaea stagnalis is a species that is easy to obtain, representative for the majority of fresh water areas and, thus, meets the requirements for bioindicators.

The easiness of breeding in laboratory, high reproduction, and large amount of embryonic and juvenile forms for the tests allows the statistical assessment of toxic effects in large statistical groups.

The proposed biotest is based on organisms from the group of aquatic invertebrates (mollusks) and increases the number of bioindicators, which can be widely applied in the batteries of biotests for better assessment of negative impact of different water pollutants.

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Conflicts of Interest: The authors declare no conflict of interest.

Appendix A

Table A1. Chemical parameters of water used for experiment media preparation.

| No. | Water Quality Indicators | Unit | Value | HAC                  |
|-----|--------------------------|------|-------|----------------------|
|     |                          |      |       | PI ¹  EU ²  WHO ³    |
| 1   | The color                | mg/dm³ | 1     | NAC ⁴  NAC ⁴    | 15     |
| 2   | Turbidity                | NTU   | 0.07  | 1       | accepted  5 |
| 3   | pH                       | -     | 7.0   | 6.5–9.5 6.5–9.5 - |
| 4   | Conductivity 25 °C       | µS/cm | 322   | 2500 2500 - |
| 5   | Oxidizability in KMN₄    | mg/dm³ | 0.7   | 5 5 - |
| 6   | Chlorides                | mg/dm³ | 7.3   | 250 250 250 |
| 7   | Ammonium ions            | mg/dm³ | 0.018 | 0.5 0.5 1.5 |
| 8   | Nitrates                 | mg/dm³ | <0.01 | 0.5 0.5 3 |
| 9   | Nitrates                 | mg/dm³ | 2.2   | 50 50 50 |
| 10  | Sulfates                 | mg/dm³ | 12    | 250 250 205 |
| 11  | Total hardness           | mg/dm³ | 90    | 60–500 - - |
| 12  | Calcium                  | mg/dm³ | 7     | - - - |
| 13  | Magnesium                | mg/dm³ | 5.5   | 125 - - |
| 14  | Total iron               | mg/dm³ | <0.025 | 0.2 0.2 0.3 |
| 15  | Mangan                   | mg/dm³ | <0.002 | 0.05 0.05 0.5 |
| 16  | Copper                   | mg/dm³ | <0.003 | 2 2 2 |
| 17  | Chromium                 | mg/dm³ | <0.002 | 0.05 0.05 0.05 |
| 18  | Nickel                   | mg/dm³ | <0.0025 | 0.02 0.02 0.02 |
| 19  | Cadmium                  | mg/dm³ | <0.00045 | 0.005 0.005 0.003 |
| 20  | The SUM of 4 THMs⁵       | µg/dm³ | <0.3 | 100 100 - |
| 21  | Chloroform               | µg/dm³ | <0.3 | 30 30 - |
| 22  | The SUM of 4 PAHS⁵       | µg/dm³ | <0.003 | 0.1 0.1 - |
| 23  | Benzo pyrene             | µg/dm³ | <0.003 | 0.01 0.01 - |
| 24  | Coliform Bacteria        | cfu b /100mL | 0 | 0 0 0 |
| 25  | Escherichia coli         | cfu b /100mL | 0 | 0 0 0 |
| 26  | Enterococcus faecalis    | cfu b /100mL | 0 | 0 0 - |
| 27  | Clostridium perfringens  (with spores) | cfu b /100mL | 0 | 0 0 - |
| 28  | The total number of micro-organisms on the agar medium in temp. 22°C | cfu b /100mL | 1 | NAC ⁴  NAC ⁴ |
| 29  | Free chlorines in dechlorinated tap water | mg/dm³ | 0.02 | 0.3 - |

¹, ², ³ HAC: the highest acceptable concentration according to standards (¹—PI, ²—EU, ³—WHO); ⁴—No abnormal change; ⁵—The total concentration of four common trihalomethanes: bromodichloromethane, trichloromethane, tribromomethane, and dibromochloromethane; ⁶—Colony-forming units.
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