In situ toxicity testing of Lake Orta sediments

Renato BAUDO*, Daria ROSSI and Monica BELTRAMI

CNR Istituto Italiano di Idrobiologia, Largo V. Tonolli 50, 28922 Verbania Pallanza, Italy
*e-mail corresponding author: r.baudo@iii.to.cnr.it

ABSTRACT

The use of in situ assays, which was initially proposed by Nebeker et al. (1984), is presently recognised as an effective tool to study the effects of contaminated sediments (Burton 1992). The placement of caged organisms to study short-term effects of exposure to contaminated environments has been employed for a number of aquatic species, both fish and invertebrate. This paper describes the application of this techniques to study the toxicity of Lake Orta sediments using both resident (Daphnia obtusa, D. longispina) and non-resident (D. magna, Echinogammarus stammeri) invertebrate species. In each of the selected stations, a group of four chambers, each containing 10 individuals, were deployed by scuba divers at 10 m depth and collected after 48 h. For each chamber, the surviving animals were counted, transferred to the laboratory and kept in Lake Orta water until their death. The number of neonates produced by each female in the laboratory was recorded daily in order to determine if the short exposure period could affect the reproductive behaviour of the animals. The technique and results reported here indicate the utility of in situ testing and suggest that, under certain conditions, this method of testing may yield results which are more representative of actual environmental conditions by avoiding the sample manipulation required for traditional laboratory toxicity tests.

Keywords: in situ toxicity testing, Lake Orta, Daphnia

1. INTRODUCTION

Sediments are generally recognised to play a central role in ecosystem processes, since the sediment surficial layers could be considered to be intimately linked to surface waters through physical, chemical, and biological processes. Studies of aquatic ecosystems should therefore not be limited to the water column, but must also consider the sediment quality and the reactions taking place at the sediment-water interface. This fact has been recognised recently and the integration of chemical, biological and toxicological data has become more commonly used in sediment quality assessment. A commonly cited example of this is the application of the sediment quality triad approach (Long & Chapman 1985; Chapman 1990; Chapman 1992). Toxicity may be predicted and/or detected in many ways, including through the use of biomarkers; laboratory toxicity tests; QSAR analyses; biological surveys; mesocosms; and mathematical models of ecological structure.

Probably, the most used technique is the traditional toxicity test in the lab, with single or multispecies experiments. A detailed review of the advantages and disadvantages of this approach is beyond the scope of this paper, and interested readers should refer to comprehensive reviews such as those provided by Cairns (1983), Cairns et al. (1992), Landis & Ho-Yu (1994), and Day et al. (1997). However, standard laboratory toxicity tests do have several limitations, including evaluation of the true extent of their ecological relevance (Day et al. 1997). This issue is related to the fact that exposure conditions may not mimic what occurs in nature, and the organisms are exposed to sediments that have been altered during collection and storage. As a consequence, the toxicity of contaminated sediments transferred to the laboratory may be either greater or less than the toxicity of sediments that are tested in situ (Sasson-Brickson & Burton 1991; Skalski 1991; Jacher & Burton 1993). Despite their limitations, in situ assays are presently recognised as an effective tool to study single-species effects of contaminated sediments (Burton 1992), and the placement of caged species to study short-term effects of exposure to contaminated environments has been employed for a number of organisms, including plankton (Munawar et al. 1989), mussels (Metcalfe & Hayton 1989), zooplankton (Sasson-Brickson & Burton 1991), leeches (Metcalfe & Hayton 1989), fish (Hartwell et al. 1987), and amphibians (Linder 1990).

Lake Orta has long been severely impacted by industrial pollution (see other papers in this monograph), and in 1989-1990 the lake was satisfactorily treated with calcium carbonate to neutralise the acidic pH and to lower the metal concentration in the water column. However, the treatment resulted also in the deposition of toxicants from the water column into the lakebed. Toxicity tests with Vibrio fischeri (Microtox test, 15 min) on pore waters from 10 samples collected in 1992 (Guzzella et al. 1993), indicated that sediments from most of the lake were highly toxic (Fig. 1). A further study (Burton et al. 2000) confirmed that the environment was still toxic for aquatic organisms in 1994. Therefore, as a part of the larger program of study to follow the lake recovery, it was decided to apply in situ techniques to evaluate the potential toxicity of the newly deposited sediments.
Fig. 1. Inhibition of bioluminescence of *Vibrio fischeri* (% luminescence vs controls = 100 %).

2. MATERIALS AND METHODS

In this study, specially devised benthic chambers (Fig. 2) were designed at the Italian Institute of Hydrobiology and manufactured by a commercial workshop (Rossi *et al.* 1998).

An *in situ* test vessel consists of a Plexiglas cylinder which is weighted at the bottom by a lead ring imbedded into the Plexiglas (Fig. 2). Both ends of the cylinder are equipped with screened disks, and can be sealed with threaded Plexiglas caps. When positioned onto the sediments by scuba divers, the lead keeps the chamber in a vertical position, while the screens ensure the free flow of water and particles smaller than the mesh size. In the present study the screens had a nominal pore size of 126 µm, although different screens could be used depending on the goal of the research. The 126 µm net permits exchange by diffusion alone. When a test vessel is filled with deionised water and placed into a water bath containing natural water, the concentration of the solutes inside and outside the chamber becomes homogeneous in less than 60 min. Since the chamber has an inner volume of about 600 ml, it can be used to accommodate an adequate number of animals of fairly large size.

Test vessels are prepared for deployment by installing the lower screen and the lower cap, and filling chambers with water. Test organisms are then inserted, and the upper net holder and cap are screwed on, thus sealing the chamber. At the selected station, the scuba divers carry the chambers (4 replicate chambers bound together) to the bottom, then gently remove both upper and lower caps before positioning the chambers onto the sediment.

At the end of the exposure period, the scuba diver replaces the caps before recovering the chamber and carrying it to the support boat. Upon reaching the boat, the upper cap and net holder are removed to permit recovery of the test organisms and the water from the chamber.

Fig. 2. *In situ* benthic exposure chamber.

2.1. *In situ* toxicity testing

The toxicity tests began in 1995 and continued till 1998, in different seasons, at the following stations (Fig. 3):
- Station n. 1 (Buccione) and Station 2 (Tortirogno) in October 1995; June 1996; November 1996; Septem-
ber 1997; July 1998; August 1998; and November 1998;
- Station n. 3 (Omegna) in October 1995. After this
time, Station 3 was no longer evaluated, since the
positioning of the benthic chambers was extremely
difficult on the steep slope, and the rocky substratun
prevented the collection of sediment samples.

To account for the missing Station 3, a new station
(N. 45) was added in 1997. However, this station is lo-
cated in the central and deepest part of the lake (140 m),
well beyond the access by scuba divers. Samples of
sediment and overlying water from this site were col-
lected by means of a Jenkin gravity corer. In this case,
the organisms were exposed to the sediment by placing
the benthic chambers within each undisturbed core (3
replicates for each sampling event).

![Sampling stations](image)

**Fig. 3.** Sampling stations.

### 2.2. Choice of the organisms

The experiments began with the cladoceran *Daphnia obtusa* Kurz (Crustacea, Cladocera), which has been
present in Lake Orta since at least 1986, before the
liming (Bonacina et al. 1988). Specimens were col-
lected from Lake Orta and reared in laboratory in Lake
Orta water to provide neonates (<24 h old) to be used
during the *in situ* tests.

Since *Daphnia obtusa* is adapted already to the pe-
culiar chemical conditions of Lake Orta, it is possibly
less sensitive to the toxicants present in the sediments.
To expand the data set, the non-native amphipod *Echi-
nogammarus stammeri* (S. Karaman 1931) was also
used twice in 1996. The source of the *E. stammeri* was
the River Ticino, and test organisms were reared in
Lake Maggiore water according to the procedure given
by de March (1981) for *Gammarus lacustris lacustris*.

In 1997 *D. obtusa* started to disappear from the lake,
so the non-native species *Daphnia magna* was used in-
stead. The *D. magna* were collected from laboratory
cultures, but again reared in Lake Orta water. In 1998,
the cladoceran *Daphnia longispina* O.F. Muller reap-
peared in Lake Orta. This species has previously been
reported as indigenous to the lake (Pavesi 1879; Monti
1929), and *D. longispina* individuals became increas-
ingly more abundant in our samples. Based on its natu-
ral presence, this species was used to continue the *in
situ* toxicity testing.

For the execution of *in situ* tests, a group of 4 ben-
thetic chambers containing test organisms were positioned
on the sediments at a depth of 10 m at each station. The
chambers were recovered after 48 h, an exposure period
which was selected to avoid the necessity of feeding the
test animals. After this time, the benthic chambers were
recovered, and the surviving organisms counted, trans-
ferred to the laboratory, and kept in Lake Orta water
until the death of all test organisms. Laboratory culture
conditions included a constant temperature of 20 ± 1 °C,
a photoperiod of 12 h light and 12 h dark, and a daily
feeding of 10 cal l⁻¹ as suspension of *Scenedesmus* sp.
cells plus yeast. During this period the number of young
were recorded daily in order to generate life tables.

For all species of *Daphnia*, 10 neonates (<24 h old)
were placed into each benthic chamber. For *E. stam-
meri*, 5 adults were used instead. During all experi-
ments, controls were maintained in Lake Orta water
(cladocerans), or in Lake Maggiore water (*E. stammeri*
only), and at a temperature close to that measured at the
exposure stations.

Obviously, the biological results obtained from ob-
servation of the exposed species in *in situ* experiments
alone does not allow inference of causality, unless fur-
ther chemical analyses and lab testing can substantiate
the causal relationships between ecological response
and anthropogenic stressors. Therefore, after recovery
of the benthic chambers in 1994, 1995, and 1996, the
scuba divers also collected sediment cores from the
same locations in order to provide information on the
chemical composition of the solid phase. The first 10
cm of each core were air dried and sieved (>200 µm),
then macro- and microelements (Si, Al, Fe, Ti, Ca, K,
Mg, Na, P, S, Pb, Zn, Cu, Ni, Mn, Cr) were determined
using X-Ray fluorescence spectrometry.

### 3. RESULTS AND DISCUSSION

The test organisms were reared in Lake Orta water
in order to acclimate them to the hydrochemical con-
ditions of the lake. In fact, the routine sampling of Lake
Orta plankton regularly showed healthy specimens of *D.
obtusa* and, later on, of *D. longispina* (Bonacina 2001).
The lake sediments themselves of course produced a
different medium in that sediments contained high con-
centrations of toxic metals (Tab. 1).

The dynamic equilibrium established at the sedi-
ment-water interface, which is controlled by back-diffu-
sion of the soluble species, very likely enhances the
toxicant levels in the water phase with respect to those detected in the water column itself (Calderoni & Tartari 2001). After 48 h exposure to sediments collected from Tortirogno in 1994, the survival of *D. obtusa* was 62.5% that of organisms in control treatments (Fig. 4). Survival of *D. obtusa* in sediments collected from Omegna and Bucino in 1994 was 67.5% and 77.5% of controls, respectively. The percentage of survivors increased in 1995 and 1996, but dropped again (to 80% at Buccione and 40% at Tortirogno) in 1997. These results may be due to different exposure conditions during the latter experiments, which were performed earlier (September) than the two previous years (October and November). In September 1997 the water temperature was higher (18.6°C) than in October 1995 (15.1 °C) or in November 1996 (12.7 °C). These temperature differences could potentially influence water - sediment exchange and metal toxicity. 1997 was also the year in which *D. obtusa* apparently disappeared from both Buccione and Tortirogno.

In the same experiments, *D. magna* seemed to respond better, with a survival even greater than that of the controls. In 1998, however, *D. magna* also displayed relatively high mortality, especially at St. 45, which has the greatest metal contamination. This station was also the most toxic for *D. longispina*, whereas Buccione and Tortirogno on average yielded better survival (albeit lower than in controls).

The results of the acute toxicity tests clearly show that sediments at all stations are still relatively toxic, resulting in a decreased survival for the three *Daphnia* species (Tab. 2). The amphipod *Echinogammarus*

---

**Tab. 1.** Chemical composition of the upper 10 cm sediment from cores at 4 stations in Lake Orta. Values represent mean concentrations.

<table>
<thead>
<tr>
<th>Station</th>
<th>Si (%)</th>
<th>Al (%)</th>
<th>Fe (%)</th>
<th>Ca (%)</th>
<th>Mg (%)</th>
<th>K (%)</th>
<th>Ti (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Buccione</td>
<td>28.9</td>
<td>8.0</td>
<td>4.6</td>
<td>0.90</td>
<td>1.22</td>
<td>2.04</td>
<td>0.44</td>
</tr>
<tr>
<td>Tortirogno</td>
<td>27.1</td>
<td>9.5</td>
<td>4.8</td>
<td>0.83</td>
<td>1.22</td>
<td>2.62</td>
<td>0.49</td>
</tr>
<tr>
<td>Omegna</td>
<td>26.6</td>
<td>8.4</td>
<td>3.6</td>
<td>1.65</td>
<td>1.70</td>
<td>2.71</td>
<td>0.40</td>
</tr>
<tr>
<td>St. 45</td>
<td>22.5</td>
<td>7.9</td>
<td>8.3</td>
<td>0.79</td>
<td>1.09</td>
<td>1.66</td>
<td>0.46</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Station</th>
<th>S (%)</th>
<th>P (%)</th>
<th>Na (%)</th>
<th>Cl (%)</th>
<th>C (%)</th>
<th>N (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Buccione</td>
<td>0.55</td>
<td>0.16</td>
<td>1.32</td>
<td>&lt; 0.05</td>
<td>8.7</td>
<td>1.0</td>
</tr>
<tr>
<td>Tortirogno</td>
<td>0.46</td>
<td>0.17</td>
<td>1.15</td>
<td>&lt; 0.05</td>
<td>3.4</td>
<td>0.3</td>
</tr>
<tr>
<td>Omegna</td>
<td>0.37</td>
<td>0.16</td>
<td>1.35</td>
<td>&lt; 0.05</td>
<td>10.1</td>
<td>0.6</td>
</tr>
<tr>
<td>St. 45</td>
<td>1.79</td>
<td>0.29</td>
<td>0.93</td>
<td></td>
<td>12.4</td>
<td>1.3</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Station</th>
<th>Pb (mg kg⁻¹)</th>
<th>Zn (mg kg⁻¹)</th>
<th>Cu (mg kg⁻¹)</th>
<th>Ni (mg kg⁻¹)</th>
<th>Mn (mg kg⁻¹)</th>
<th>Cr (mg kg⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Buccione</td>
<td>294</td>
<td>458</td>
<td>1176</td>
<td>113</td>
<td>489</td>
<td>436</td>
</tr>
<tr>
<td>Tortirogno</td>
<td>311</td>
<td>493</td>
<td>1273</td>
<td>122</td>
<td>718</td>
<td>485</td>
</tr>
<tr>
<td>Omegna</td>
<td>169</td>
<td>479</td>
<td>170</td>
<td>78</td>
<td>740</td>
<td>154</td>
</tr>
<tr>
<td>St. 45</td>
<td>334</td>
<td>500</td>
<td>1222</td>
<td>119</td>
<td>1080</td>
<td>1769</td>
</tr>
</tbody>
</table>

---

**Fig. 4.** Survival of test organisms after exposure (48 h) to Lake Orta sediments. Values represent % of control survival.
In situ toxicity testing of Lake Orta sediments

E. stammeri, appeared to be much less sensitive in response to sediments collected from both Buccione and Tortirogno in 1996. Survival for E. stammeri exposed to sediments from these sites was identical to control survival, and exposures with this organism were not repeated in subsequent years.

Demographic parameters were calculated from the resulting life-tables according to procedures suggested by Sieber (1973) and Margalef (1974). These parameters included life expectancy, net reproduction rate, and generation length.

The corresponding values of demographic parameters for controls are reported in table 3, whereas the results for the different experiments are shown in figures 5-7. Data in figures 5-7 are expressed in % of control values in order to facilitate comparisons. For all 3 species, the short exposure was sufficient to significantly lower the mean life expectancy (Fig. 5), with the exception of D. obtusa at Omegna. This result, however, refers only to a single test, which was conducted in 1994. St. 45 proved to be the most toxic, reducing by 75 % the life expectancy of both D. magna and D. longispina.

Similar results were also observed for the net reproduction rate (Fig. 6), and for the generation length (Fig. 7), indicating that these demographic parameters were also influenced by maternal exposure to a toxic environment.

For Echinogammarus stammeri (Tab. 4), different results were obtained in June and November. In the first experiment, a significant reduction in the number of neonates was observed, along with longer maternal survival compared to controls. On the other hand, the effects of contaminated sediments was greater in November for E. stammeri. Exposed individuals failed to reproduced and mean life expectancy decreased by 38% (Tortirogno) and 46% (Buccione). Therefore, at least for this species, the toxicants present into the sediments did not produce a direct and acute lethal effect, but still affected population parameters to a significant extent.

### Tab. 2. Survival, life expectancy, net reproduction rate, generation length (in % of control) after in situ exposures of 48 h.

<table>
<thead>
<tr>
<th></th>
<th>Survival</th>
<th>Life Expectancy</th>
<th>Net reproduction rate</th>
<th>Generation length</th>
</tr>
</thead>
<tbody>
<tr>
<td>D. obtusa</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Buccione</td>
<td>89.6</td>
<td>61.0</td>
<td>66.1</td>
<td>85.8</td>
</tr>
<tr>
<td>Tortirogno</td>
<td>80.8</td>
<td>82.0</td>
<td>110.9</td>
<td>108.3</td>
</tr>
<tr>
<td>Omegna</td>
<td>82.5</td>
<td>166.9</td>
<td>628.2</td>
<td>143.6</td>
</tr>
<tr>
<td>Mean</td>
<td>84.3</td>
<td>103.3</td>
<td>268.4</td>
<td>112.6</td>
</tr>
<tr>
<td>D. magna</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Buccione</td>
<td>88.6</td>
<td>67.7</td>
<td>95.9</td>
<td>91.7</td>
</tr>
<tr>
<td>Tortirogno</td>
<td>70.5</td>
<td>59.9</td>
<td>76.6</td>
<td>95.7</td>
</tr>
<tr>
<td>St. 45</td>
<td>31.0</td>
<td>26.4</td>
<td>40.0</td>
<td>81.2</td>
</tr>
<tr>
<td>Mean</td>
<td>63.4</td>
<td>51.3</td>
<td>70.8</td>
<td>89.5</td>
</tr>
<tr>
<td>D. longispina</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Buccione</td>
<td>93.0</td>
<td>105.5</td>
<td>58.1</td>
<td>112.3</td>
</tr>
<tr>
<td>Tortirogno</td>
<td>84.7</td>
<td>90.7</td>
<td>130.5</td>
<td>93.6</td>
</tr>
<tr>
<td>St. 45</td>
<td>21.4</td>
<td>25.3</td>
<td>42.4</td>
<td>70.3</td>
</tr>
<tr>
<td>Mean</td>
<td>66.4</td>
<td>73.8</td>
<td>77.0</td>
<td>92.1</td>
</tr>
</tbody>
</table>

In addition to direct lethal effect, it is reasonable to expect that exposure of populations to these sediments may elicit negative responses. To test this hypothesis, surviving animals were transferred to Lake Orta water and grown in optimal conditions until the death of the last individual.

### Tab. 3. Life-tables mean values for controls.

<table>
<thead>
<tr>
<th>Control mean</th>
<th>D. obtusa</th>
<th>D. magna</th>
<th>D. longispina</th>
</tr>
</thead>
<tbody>
<tr>
<td>Net reproductive rate (N° newborn)</td>
<td>16.2</td>
<td>7.4</td>
<td>8.1</td>
</tr>
<tr>
<td>Generation length (days)</td>
<td>23.8</td>
<td>28.8</td>
<td>43.2</td>
</tr>
<tr>
<td>Mean exp. Life (days)</td>
<td>17.1</td>
<td>21.7</td>
<td>27.9</td>
</tr>
</tbody>
</table>

For Echinogammarus stammeri (Tab. 4), different results were obtained in June and November. In the first experiment, a significant reduction in the number of neonates was observed, along with longer maternal survival compared to controls. On the other hand, the effects of contaminated sediments was greater in November for E. stammeri. Exposed individuals failed to reproduced and mean life expectancy decreased by 38% (Tortirogno) and 46% (Buccione). Therefore, at least for this species, the toxicants present into the sediments did not produce a direct and acute lethal effect, but still affected population parameters to a significant extent.

### Tab. 4. Demographic parameters (in % with respect to controls) for E. stammeri, after 48 h exposure to L. Orta sediments. * No neonates produced.

<table>
<thead>
<tr>
<th></th>
<th>Buccione</th>
<th>Tortirogno</th>
</tr>
</thead>
<tbody>
<tr>
<td>June 18, 1996</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Net reproductive rate</td>
<td>42</td>
<td>10</td>
</tr>
<tr>
<td>Generation length</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>Mean Life Expectancy</td>
<td>141</td>
<td>227</td>
</tr>
<tr>
<td>November 6, 1996</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>Net reproductive rate</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>Generation length</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>Mean exp. Life</td>
<td>54</td>
<td>62</td>
</tr>
</tbody>
</table>
Fig. 5. Mean life expectancy (in % with respect to the relative controls).

Fig. 6. Net reproduction rate (in % with respect to the relative controls).

Fig. 7. Generation length (in % with respect to the relative controls).
4. CONCLUSIONS

As suggested by Ingersoll et al. (1997), "field validation of endpoints is essential to reduce laboratory to field extrapolation error". The in situ approach may be a convenient tool to address this problem, especially as a first step for screening and monitoring polluted environments. At this stage, "the specific identity of the stressor may or may not be important to know (Day et al. 1997)". Only if and when a significant effect is detected should more expensive and time-consuming chemical and ecotoxicological investigations be initiated.

Current research on L. Orta sediments indicates that, after liming, the sediments are still contaminated enough to induce either a direct lethal effect after a short exposure, or sublethal effects influencing population parameters. These effects could potentially affect the whole trophic chain, since they reflect on the population structure of important components of the indigenous zooplankton.

By correlating the results of the tests with information on the chemical composition of cores collected at the same stations, possible causes for the lethal and sublethal effects can be identified. Primary contributors to toxicity appear to be: Zn (negatively correlated with survival of D. obtusa); Pb (negatively correlated with mean life expectancy and mean generation length in D. longispina, and with survival, mean life expectancy, and net reproduction rate in D. magna); Cr (negatively correlated with survival and mean life expectancy in D. longispina, and with mean life expectancy in D. magna); Mn (negatively correlated with mean generation length in D. longispina, and with survival and net reproduction rate in D. magna).

Unfortunately, chemical analyses on the cores did not cover organic micropollutants; but past and ongoing studies (e.g. Guzzella et al. 1993) indicate that PCBs, DDT, and PAHs may reach potentially harmful levels in sediments, at least in the central and northern basins. The toxic effects observed in this study could therefore also be due to such organic contaminants.

The results reported here are consistent with those obtained in toxicity tests with other organisms (Rossi & Beltrami 1998; Rossi et al. 1998; Beltrami et al. 1999; Burton et al. 2001; Barbero et al. 2001), and it can be concluded that in 1998, 8 years after liming, the sediments of Lake Orta are still contaminated to an extent which can pose some problems for the indigenous biota, possibly delaying the colonisation of the most contaminated areas (Nocentini et al. 2001).

REFERENCES


