Biomarkers of pollution effects in the bivalves *Mytilus edulis* and *Macoma balthica* collected from the southern coast of Finland (Baltic Sea)

Kari K. Lehtonen1,*, Sari Leiniö1, Rolf Schneider2, Mirja Leivuori1

1Finnish Institute of Marine Research, PO Box 2, 00561, Helsinki, Finland
2Baltic Sea Research Institute, Department of Biological Oceanography, Seestrasse 15, 18119 Rostock-Warnemünde, Germany

ABSTRACT: Levels of selected biomarkers in 2 bivalve species, the hard-bottom filter-feeder *Mytilus edulis* and the infaunal deposit/suspension-feeder *Macoma balthica* were measured in 2 areas (Tvärminne and Turku Archipelago) in the southern coast of Finland. The bivalves were collected for the measurement of acetylcholinesterase (AChE), glutathione-S-transferase (GST) and catalase (CAT) activities, and for the determination of metallothionein (MT) content. The content of organochlorines (8 PCB congeners, DDE, DDD, DDT, HCH, HCB, dieldrin) and selected heavy metals (Cd, Cu, Hg, Zn) were measured in the soft tissues. In *M. balthica* collected from the Turku Archipelago region, biomarker gradients were observed in MT, GST and CAT, mostly coinciding with the tissue concentrations of total PCBs, DDTs and selected metals. In *M. edulis* the biomarker responses were less consistent with regard to tissue pollutant concentrations. In the Tvärminne region no marked biomarker gradients were observed. The levels of all biomarkers varied significantly between the two species. The integrated biomarker response (IBR) index showed good accordance with the observed high tissue levels of organochlorines at the northern Airisto station in both species. This study is the first of its kind performed in the coastal waters of Finland, and the results clearly demonstrate the usefulness of the multibiomarker approach. Moreover, the use of 2 sessile species with markedly differing feeding modes and living habitats is considered a rational strategy for assessing the pollution status of coastal sea areas.

KEY WORDS: Biomarkers · Baltic Sea · Bivalves · *Macoma balthica* · *Mytilus edulis* · Organochlorines · Heavy metals

INTRODUCTION

Assessment and monitoring of the state of contamination of the marine environment has traditionally been almost entirely based on measurements of concentrations of well-known harmful substances in seawater, sediments and/or biota. However, with the currently enormous and continuously increasing number of potentially toxic substances present in and released into world seas, this approach alone can no longer be considered practical, cost-effective or even possible. Besides the obvious technical and economical constraints, the most important fact is that by using merely chemical measurements the biological effects of both single pollutants and the combined effect of a number of compounds in marine organisms remain unresolved; therefore, the true environmental hazards of contamination are insufficiently assessed. During the past decades, molecular, biochemical, cytological, immunological and physiological techniques for the detection of effects of pollutants in marine organisms, the so-called biomarkers, have been under dynamic develop-
ment. More recently, a strong emphasis on their further development and application in marine monitoring programmes has emerged within the European Union (e.g. OSPAR and MEDPOL related activities and the pan-European BEEP project [Biological Effects of Environmental Pollution on Marine Coastal Ecosystems, www.beep.u-bordeaux1.fr]). This topic needs urgent attention, particularly within the Baltic Sea area, where this approach has been largely neglected, e.g. in the HELCOM monitoring programme (Lehtonen & Schiedek 2006).

Sessile benthic organisms are universally accepted as very suitable bioindicators of environmental contamination (e.g. Rainbow & Phillips 1993). In the poor-diversity northern Baltic Sea, the hard-bottom filter-feeding mussel *Mytilus edulis* and the infaunal deposit/suspension-feeding clam *Macoma balthica* are among the very few benthic species available for environmental monitoring. The aims of the study were (1) to look at selected biomarkers in field conditions in bivalves from a northern Baltic Sea coastal marine environment with suspected pollution gradients; (2) to apply an integrated multibiomarker index; and (3) to compare the biomarker signals with concentrations of selected pollutants in the bivalve tissues. The nature of pollution in the investigated areas was largely unknown. Thus, biomarkers were chosen to represent different types of biological responses to elucidate the pollutant groups present in the environment that may cause effects. Of the selected biomarkers, acetylcholinesterase (AChE) activity is an indicator of synaptic neurotoxic effects (e.g. Day & Scott 1990, Bocquené & Galgani 1998); metallothionein (MT) content is used to indicate exposure to heavy metals (e.g. George & Olsson 1994, Viarengo et al. 1999); glutathione-S-transferase (GST) is a Phase II enzyme involved in the detoxification of organic xenobiotics (e.g. Habig et al. 1974, Lee et al. 1988); and an increase in catalase (CAT) activity signifies oxidative stress caused by excessive oxyradical formation in the metabolism of various compounds (Claiborne 1985, Di Giulio et al. 1989).

**MATERIALS AND METHODS**

**Sampling.** *Mytilus edulis* and *Macoma balthica* were collected in 2 regions on the southern coast of Finland. In the archipelago area close to Tvärminne Zoological Station of the University of Helsinki (hereafter called ‘Tvärminne region’), 4 sampling sites (A1 to A4) extending from the vicinity of the Koverhar steel factory towards the outer archipelago were sampled (Fig. 1). In the archipelago area close to the city of Turku (‘Turku region’) the 3-station sampling transect (B1 to B3) started from northern Airisto, close to a for-
mer dumping site of harbour dredging waste, and ended in a presumably less-polluted area near the Archipelago Sea Research Station (on Seili Island). Because of habitat differences, the exact sampling locations of *M. edulis* and *M. balthica* differed slightly at each station, but together they represented a distinct area. Samples were taken in spring 2001, in the Turku region on 8 May and in the Tvärminne region on 15 to 17 May. *M. edulis* were collected by scuba diving, and *M. balthica* with the van Veen grab or by bottom trawling.

Hydrography. At the *Macoma balthica* stations, a water sample was obtained from ca. 1 m above the bottom and measured for dissolved oxygen concentration using the Winkler method (Grasshoff 1983) (Table 1). From the same sample, near-bottom temperature and salinity were measured with a YSI instrument. In connection with the sampling of *Mytilus edulis*, temperature and salinity were determined from the surface water (1 m). Water transparency was recorded using a Secchi disk (Tvärminne region) or estimating diving visibility (Turku region).

Sample preparation. For biomarker analyses, the bivalves collected from the Tvärminne region were dissected the same day as sampled. The specimens collected from the Turku region were kept overnight in ambient water for gut purging. To minimise the differences in laboratory maintenance time, the dissecting of bivalve groups collected from different stations was always performed in the same order as the sampling in the field was done. Bivalves collected for pollutant analyses were kept overnight in a cold room (6ºC) in water buckets containing water collected from the Turku region were kept overnight in ambient water for gut purging. The specimens collected from the shells, all precautions were taken to avoid sample contamination. In 2 cases adequate numbers of bivalves could not be obtained for pollutant analyses.

Biomarker measurements. *AChE activity:* Analyses (6 replicates) were performed essentially as described in Bocquené & Galgani (1998). Tissues were homogenised in 1:2 w/v 0.02 M phosphate buffer (pH 7.0) with 0.1% Triton X. Homogenates were centrifuged at 10,000 × g for 20 min, and the supernatants (S9) were used for AChE measurement. A BioRad Benchmark 96-well microplate reader (BioRad) was used for the spectrophotometric determination of the Ellman reaction (Ellman et al. 1961). Tissue protein concentration was determined using the Bradford (1976) method, using bovine serum albumin (BSA) as the protein standard. AChE activity values are expressed as equivalents of acetylthiocholine (ACTC) hydrolysed (nmols ACTC min⁻¹ mg⁻¹ protein), with 1 unit change in optical density (ΔOD) corresponding to the hydrolysis of 75 nmol of ACTC.

*MT content:* Analyses (5 replicates) were carried out according to the method of Viarengo et al. (1997). Tissues were homogenised in reducing conditions (0.05 M sucrose TRIS buffer, pH 8.6, containing 0.01% β-mercaptoethanol). Homogenates were centrifuged at 30,000 × g for 20 min. S9 were collected, and ethanol/chloroform fractionation was used to obtain the partially purified metalloprotein fraction. The MT concentration was then measured by spectrophotometric determination of –SH groups using Ellman’s reagent (DTNB).

*GST and CAT activity:* Measurements were performed on 10 individuals from each study station, in liquid nitrogen immediately after dissection and stored at –80°C until analysis.

For pollutant analyses the bivalves were stored whole (with shells) at –80°C until analysis. For heavy metal analyses 19 to 35 individuals were taken, while for the measurement of organochlorine levels the amount of specimens was up to 100, depending on the availability of animals. During dissection of soft bodies from the shells, all precautions were taken to avoid sample contamination. In 2 cases adequate numbers of bivalves could not be obtained for pollutant analyses.

### Table 1. Hydrographic data recorded at the study stations during the sampling of *Mytilus edulis* and *Macoma balthica.*

<table>
<thead>
<tr>
<th>Station code</th>
<th>Station name: <em>M. balthica/M. edulis</em></th>
<th>Date (2001)</th>
<th>Depth (m):</th>
<th><em>M. balthica/M. edulis</em></th>
<th>Temp. (°C): bottom/surface</th>
<th>Salinity (psu): bottom/surface</th>
<th>Secchi depth/visibility (m) at bottom</th>
</tr>
</thead>
<tbody>
<tr>
<td>A1</td>
<td>Koverhar</td>
<td>15–17 May</td>
<td>30/4</td>
<td>4.2/7.7</td>
<td>4.2/5.1</td>
<td>3.5/–</td>
<td>6.55</td>
</tr>
<tr>
<td>A2</td>
<td>Kummeli</td>
<td>15–17 May</td>
<td>32/5</td>
<td>4.1/8.3</td>
<td>5.6/5.0</td>
<td>3.5/–</td>
<td>6.86</td>
</tr>
<tr>
<td>A3</td>
<td>Storfjärden/Sundholmen</td>
<td>15–16 May</td>
<td>33/5</td>
<td>4.4/7.6</td>
<td>4.9/5.1</td>
<td>4.0/–</td>
<td>6.24</td>
</tr>
<tr>
<td>A4</td>
<td>Långskär/Granbusken</td>
<td>15–16 May</td>
<td>36/4</td>
<td>4.4/8.1</td>
<td>6.2/5.1</td>
<td>6.0/–</td>
<td>6.96</td>
</tr>
<tr>
<td>B1</td>
<td>N. Airisto</td>
<td>8 May</td>
<td>23/2</td>
<td>4.7/11.5</td>
<td>5.7/4.9</td>
<td>–/0.5</td>
<td>9.27</td>
</tr>
<tr>
<td>B2</td>
<td>Grangrundet</td>
<td>8 May</td>
<td>22/5</td>
<td>5.2/9.1</td>
<td>5.0/5.1</td>
<td>–/1.0</td>
<td>9.86</td>
</tr>
<tr>
<td>B3</td>
<td>Högholmen</td>
<td>8 May</td>
<td>32/4</td>
<td>4.8/7.8</td>
<td>5.4/5.4</td>
<td>–/1.5</td>
<td>9.53</td>
</tr>
</tbody>
</table>

...
determining both enzymatic rates from the same homogenate. Pieces of digestive gland (>15 mg) were homogenised (1:3 w/v) for 2 min in cold 100 mM potassium phosphate buffer (pH 7.0) using a Kontes pellet pestle. The homogenate was centrifuged at 10,000 × g for 20 min. The S9 were diluted with cold buffer (see below) and kept on ice for analyses of enzyme activities, performed within 3 h. The measurements were performed in a 1 cm quartz cuvette with total reaction volumes of 1275 (GST) and 1350 µl (CAT) in buffer solution kept at room temperature (20 to 22°C).

The GST assay was performed using a modified method based on Habig et al. (1974). Dilutions of 1:10 (Mytilus edulis) and 1:40 (Macoma balthica) were prepared. 20 mM chlorodinitrobenzene (CDNB) was used as the substrate and 20 mM glutathione (GSH) as the co-substrate. GST activity was estimated by measuring the formation rate of conjugated substrate (CDNB-GSH). CAT activity was measured as the disappearance rate of hydrogen peroxide (H₂O₂) from the reaction solution according to Claiborne (1985). S9 dilutions of 1:10 (M. edulis) and 1:25 (M. balthica) were prepared for the test, and H₂O₂ was applied as a 60 mM solution. All the reagents (except the buffer solution) were prepared daily immediately prior to the measurements.

Following addition of sample (supernatant) enzymatic reactions were monitored for 60 s using a Perkin-Elmer Lambda 2 spectrophotometer using the Time-drive mode of WinLab software. GST conjugates were measured at 340 nm, and the reduction of H₂O₂ by catalase at 240 nm. The reaction speed was afterwards evaluated between 0 and 30 s, which showed a better linearity and slightly higher rates. Blank readings for spontaneous conjugate formation (GST) and H₂O₂ hydrolysis (CAT) were subtracted from the sample activity measurements. Protein in the S9 was determined on microplates using the Bradford (1976) method and a BSA standard.

**Pollutant levels in tissues. Organochlorines:** Before and after freeze-drying with a Christ-Alpha 1–4, the pooled samples of bivalve soft bodies were weighed to allow for calculations of wet-weight-based data. Samples were vigorously ground to a homogenous powder, from which lipids were extracted for 8 h in a Soxhlet apparatus using a mixture of 20% acetone in n-hexane. The extracts were concentrated in tared flasks using a rotary evaporator, and total lipids gravimetrically estimated. Clean-up and fractionation of organochlorine compounds were carried out by alumina and silica column chromatography within the guidelines of Holden & Marsden (1969), and as specified by Schneider (1982). The gas chromatograph employed for the estimation of organochlorines was a FISONS instrument, GC 8000, equipped with an AS 800 auto-sampler for automatic on-column sample injection and an ECD 800 electron capture detector. It was fitted with a J&W DB-35 30 m × 0.32 mm fused silica capillary column for separation and a J&W 5 m × 0.53 mm untreated fused silica capillary column as a retention gap (pre-column). The carrier gas, hydrogen, was adjusted to a flow rate of 2 ml min⁻¹ at 40°C. N₂ was used as detector make-up gas. The GC oven temperature programme was as follows: 1 min 40°C isothermal, 40°C min⁻¹ to 160°C, 10 min 160°C isothermal, 4°C min⁻¹ to 290°C and 10 min 290°C isothermal.

Mixed pesticide standard (α-, β-, γ-HCH, dieldrin, α,p'- and p,p'-isomers of DDE, DDD and DDT) and PCB congener standard (PCB 52, 101, 105, 118, 149, 153, and 180) solutions at (individual compound) concentrations of 5, 10, 15, 20, 50, 100 and 150 pg µl⁻¹ were applied for calibration. As an internal standard, 20 pg µl⁻¹ of PCB 198 was added to each standard and sample solution. The analytical quality of organochlorine estimation in bivalves was repeatedly checked by triplicate analysis of IAEA reference materials (IAEA 1980, 1988; copepod and mussel homogenates), resulting in typical standard deviations between 3 and 5%, as well as by frequent participation in QUASIMEME laboratory performance studies.

**Heavy metals:** One pooled sample from each species and station was analysed for selected heavy metals (Cd, Cu, Hg and Zn). The soft tissue was removed from the shells, refrozen and freeze-dried for 30 h with an Edwards Super Modulyo freeze dryer. Samples were homogenised in a Fritsch planetary mill (Pulverisette 5) with grinding bowls and balls made of zirconium dioxide. Aliquots (0.25 g) were digested in Teflon bombs with 2 ml H₂O and 5 ml HNO₃ in a microwave oven. Samples were heated with pressure regulation at 150 psi for 60 min. After cooling, samples were diluted with Milli-Q water up to the 25 ml mark in a volumetric plastic flask, transferred to plastic storage bottles and stored at room temperature until analysis. Heavy-metal determinations were done with an atomic adsorption spectrophotometer (Perkin-Elmer AAS 5100 ZL). These measurements were made with the flameless method, with matrix modification and platform techniques. Commercial certified sediment reference materials BCR-422 and DORM-2 (NRCC) were used to control analytical reliability. For the elements studied, nearly 100% recoveries (80 to 110%) were obtained for both reference materials.

**Statistical treatment of data.** Single-biomarker variability between the stations within each of the 2 study areas was investigated using a standard ANOVA and post-hoc Tukey’s test. To study the relations between biological responses and tissue contaminant levels jointly in both species and study areas, the biomarker
and tissue contaminant data from both species and study areas were combined by calculating normalised parameter values (between 0 and 1) for each species at each study station. The normalised values of single biomarkers or the sums of all 4 of them, now combined for both species, were analysed against normalised values of tissue levels of total organochlorines, ΣPCBs, ΣDDTs, single heavy metals or their sum recorded for both species. SYSTAT™ 9.0 software was used for all statistical analyses.

To assess the ‘pollution status’ of each location, an integrated biomarker response (IBR) index was calculated for each species using all the 4 biomarkers studied. A simple starplot graphic tool (Microsoft Excel) was applied to the data to visualise the results. The IBR method of Beliaeff & Burgeot (2002) is briefly described here. For each biomarker: (1) Calculation of mean and SD for each station. (2) Standardisation of data for each station: \[ x' = \frac{x - \text{mean}}{\text{SD}} \] Result: variance = 1, mean = 0. (3) Using standardised data, addition of the value obtained for each station to the absolute (= non-negative) value of the minimum value in the data set: \[ B = x' + |x_{min}| \] Result: adjusts the lowest value in the set to zero. For all the biomarkers treated this way: calculation of starplot areas by multiplication of the obtained value of each biomarker (Bi) with the value of the next biomarker, arranged as a set, dividing each calculation by 2 and summing-up of all values: \[ \left\{ \frac{(B_1 \times B_2)/2}{B_1} \right\} + \left\{ \frac{(B_2 \times B_3)/2}{B_2} \right\} + \ldots + \left\{ \frac{(B_{n-1} \times B_n)/2}{B_{n-1}} \right\} \] Result: IBR (average of different arrangements of biomarkers in the set).

Tissue concentrations of organochlorines and heavy metals were also standardised this way, to obtain indices for both total organochlorines and total heavy metal loads in the bivalve tissues at the different study stations. From all the standardised organochlorines and heavy metals, a mean value was calculated to represent the ‘general load’ of these pollutant groups at the different study stations.

RESULTS

Spatial variability in biomarker responses

AChE

In both bivalve species, AChE activities were relatively homogenous over the regions and stations, and no significant differences could be noted (Fig. 2, Table 2). In the gill tissue of Mytilus edulis the AChE activity ranged from 19.3 to 25.9 nmol ACTC min⁻¹ mg⁻¹ protein, being lower in the foot tissue of Macoma balthica with values between 17.7 and 21.8 nmol ACTC min⁻¹ mg⁻¹ protein.

MT

In the Tvärminne region, a gradient could be observed in the MT content of Mytilus edulis, with the highest values observed at Stn A1, closest to the Koverhar steel factory (mean and SE: 426 ± 43 µg g⁻¹ wet wt; Fig. 3, Table 2). The levels were significantly (p < 0.05) lower at Stns A3 (303 ± 21 µg g⁻¹ wet wt) and A4 (311 ± 25 µg g⁻¹ wet wt), furthest away from Koverhar. In Macoma balthica, no significant differences in MT levels between the stations in the Tvärminne region could be observed, with values ranging from 392 (A1) to 460 µg g⁻¹ wet wt (A4). In the Turku region, somewhat higher MT levels were recorded in M. edulis collected at the northern Airisto Stn B1 (405 ± 24 µg g⁻¹ wet wt) compared with the other (‘reference’) station sampled (B3; 345 ± 38 µg g⁻¹ wet wt), but difference was not statistically significant. In M. balthica a 3-station gradient was marked with the values observed at Stn B1 (542 ± 29 µg g⁻¹ wet wt) being significantly (p < 0.01) higher than those recorded at Stn B3 (414 ± 27 µg g⁻¹ wet wt). In general, the levels of MT were about 25% higher in M. balthica than in M. edulis.
In *Mytilus edulis*, the highest values for GST activity were recorded at the mid-transect Stn A3 in the Tvärminne region (472 ± 66 nmol min⁻¹ mg⁻¹ protein) and the reference Stn B3 in the Turku region (472 ± 19 nmol min⁻¹ mg⁻¹ protein; Fig. 4). The lowest activity was observed close to the Koverhar factory (A1; 339 ± 34 nmol min⁻¹ mg⁻¹ protein). No statistically significant differences could be detected between the stations and the

Table 2. *Mytilus edulis*, *Macoma balthica*. Compilation of statistical analyses (ANOVA, post-hoc Tukey) on variability in biomarker responses of bivalves at different stations within different regions. *p < 0.05; **p < 0.01; ns: not significant at p ≥ 0.05.

<table>
<thead>
<tr>
<th>Study area</th>
<th>Species</th>
<th>Biomarker</th>
<th>F</th>
<th>p</th>
<th>Differing stations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tvärminne region</td>
<td><em>M. edulis</em></td>
<td>AChE</td>
<td>2.410</td>
<td>0.097</td>
<td>ns</td>
</tr>
<tr>
<td></td>
<td></td>
<td>MT</td>
<td>4.552</td>
<td>0.022</td>
<td>* A1 vs A3 and A4: p &lt; 0.05</td>
</tr>
<tr>
<td></td>
<td></td>
<td>GST</td>
<td>1.385</td>
<td>0.264</td>
<td>ns</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CAT</td>
<td>0.880</td>
<td>0.461</td>
<td>ns</td>
</tr>
<tr>
<td>Turku region</td>
<td><em>M. edulis</em></td>
<td>AChE</td>
<td>0.144</td>
<td>0.866</td>
<td>ns</td>
</tr>
<tr>
<td></td>
<td></td>
<td>MT</td>
<td>1.219</td>
<td>0.311</td>
<td>ns</td>
</tr>
<tr>
<td></td>
<td></td>
<td>GST</td>
<td>1.205</td>
<td>0.315</td>
<td>ns</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CAT</td>
<td>0.117</td>
<td>0.890</td>
<td>ns</td>
</tr>
<tr>
<td>Tvärminne region</td>
<td><em>M. balthica</em></td>
<td>AChE</td>
<td>1.593</td>
<td>0.222</td>
<td>ns</td>
</tr>
<tr>
<td></td>
<td></td>
<td>MT</td>
<td>2.262</td>
<td>0.118</td>
<td>ns</td>
</tr>
<tr>
<td></td>
<td></td>
<td>GST</td>
<td>2.161</td>
<td>0.110</td>
<td>ns</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CAT</td>
<td>1.073</td>
<td>0.373</td>
<td>ns</td>
</tr>
<tr>
<td>Turku region</td>
<td><em>M. balthica</em></td>
<td>AChE</td>
<td>0.399</td>
<td>0.678</td>
<td>ns</td>
</tr>
<tr>
<td></td>
<td></td>
<td>MT</td>
<td>7.500</td>
<td>0.008</td>
<td>** B1 vs B3: p &lt; 0.01</td>
</tr>
<tr>
<td></td>
<td></td>
<td>GST</td>
<td>6.597</td>
<td>0.005</td>
<td>** B1 vs B3: p &lt; 0.01</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CAT</td>
<td>6.355</td>
<td>0.007</td>
<td>** B1 vs B2 and B3: p &lt; 0.01</td>
</tr>
</tbody>
</table>

Fig. 3. *Mytilus edulis*, *Macoma balthica*. Box plots of metallothionein (MT) content in the digestive gland of (A) *M. edulis* and (B) *M. balthica* at the study stations. Station codes as in Table 1. Each plot shows median, interquartile range, data range and mild (•) and extreme (○; >3.0 × interquartile range) outliers for each station shown.

Fig. 4. *Mytilus edulis*, *Macoma balthica*. Box plots of glutathione S-transferase (GST) in the digestive gland of (A) *M. edulis* and (B) *M. balthica* at the study stations. Station codes as in Table 1. Note different scales. Each plot shows median, interquartile range, data range and mild (•) and extreme (○; >3.0 × interquartile range) outliers for each station shown.
regions (Table 2). In *Macoma balthica*, the values were similar at all stations of the Tvärminne region transect (1852 to 1926 nmol min⁻¹ mg⁻¹ protein), except for the slightly lower activities recorded at Stn A2 (1467 ± 110 nmol min⁻¹ mg⁻¹ protein). In the Turku region, a distinct gradient in GST activity was observed, with enzyme rates recorded at Stn B1 (3078 ± 543 nmol min⁻¹ mg⁻¹ protein) twice as high compared with the reference Stn B3 (1416 ± 86 nmol min –¹ mg –¹ protein, p < 0.01). On average, the GST activity in *M. balthica* was ca. 5-fold higher compared with *M. edulis*.

**CAT**

In *Mytilus edulis*, no significant variability could be noted between the stations and the 2 study regions (Fig. 5, Table 2). Highest CAT activity was observed at the outermost station of the Tvärminne region (A4; 144 ± 24 µmol min⁻¹ mg⁻¹ protein), while at all the other stations the activity was 105 to 125 µmol min –¹ mg –¹ protein. In *Macoma balthica* collected from the Tvärminne region, no statistically significant gradient in CAT activity could be observed either, with the lowest mean value recorded at Koverhar (A1; 277 ± 36 µmol min⁻¹ mg⁻¹ protein) and highest in the open-sea region (A4; 379 ± 44 µmol min⁻¹ mg⁻¹ protein). In the Turku region, a significant (p < 0.01) gradient could be observed with the highest CAT activity at the northern Airisto station (B1; 412 ± 43 µmol min⁻¹ mg⁻¹ protein) compared with the 2 other stations (B2 and B3) showing equal mean values (270 ± 20 and 269 ± 22 µmol min⁻¹ mg⁻¹ protein, respectively). On average, CAT activity in *M. balthica* was ca. 2.5 times higher compared with *M. edulis*.

**Tissue pollutant levels**

**Organochlorines**

Analyses of organochlorines in the soft bodies of the bivalves showed marked gradients, especially in the Turku region (Table 3). The steepest gradient was observed in the case of ΣPCBs (8 congeners), with the clearly highest values at Stn B1, both in *Macoma balthica* and *Mytilus edulis* (881 and 133 ng g⁻¹ lipid, respectively), with congeners CB-153, CB-101, CB-149 and CB-138 forming ca. 90% of the ΣPCBs. At the ‘reference’ Stn B3 the respective ΣPCB concentrations in the tissues were 249 and 60 ng g⁻¹ lipid. A similar but less steep pattern could be observed in ΣDDT compounds with *M. balthica* and *M. edulis* at Stn B1 containing 217 and 83 ng g⁻¹ lipid, respectively, and 130 and 56 ng g⁻¹ lipid at Stn B3. In the Tvärminne region, homogeneous ΣDDT concentrations (99 to 109 ng g⁻¹ lipid) in the tissues of *M. balthica* were observed.

The DDT (α,α’-DDT + p,p’-DDT) to ΣDDTs (including metabolites) ratio showed marked variability, both between the species and the study regions (Fig. 6). In *Mytilus edulis* the ratios ranged from 7.5 to 12.0% in the Tvärminne region and between 39.1 and 42.3% in the Turku region, while in *Macoma balthica* the ranges were 2.3–3.8% and 8.1–13.2% in the respective areas.

If the organochlorine concentrations are expressed on a dry weight basis, the differences between the stations become somewhat smaller, due to the marked differences in tissue lipid levels (Table 4). In *Macoma balthica* the lowest lipid levels were recorded at Stns A4 and B1 (2.3% of wet wt), while the highest value was recorded at Stn B2 (3.3%). In *Mytilus edulis* a markedly low lipid level was observed at Stn A3 (1.0%), with the others being between 1.7 and 3.4%.

**Heavy metals**

Tissue heavy metal concentrations showed marked interspecies variability, with *Macoma balthica* containing an order of magnitude higher levels of Cu and Zn.
compared with *Mytilus edulis*, which in turn had significantly higher levels of Cd (Table 5). The levels of Hg were similar in both species. In *M. balthica*, the values for Cu and especially Zn were distinctly higher in the Tvärminne region (694 to 1410 µg g$^{-1}$ dry wt) compared with the Turku region (392 to 605 µg g$^{-1}$ dry wt). With regard to Zn, an apparent gradient was observed in both areas. In *M. edulis*, no significant variability between the main study areas could be noted.

**Integrated biomarker response index versus standardised pollutant levels**

In *Macoma balthica* the calculated IBR index indicated that the northern Airisto station (B1) was unquestionably the most contaminated site, with the IBR decreasing along the gradient (Fig. 7). *Mytilus edulis* at Stn B1 also displayed the highest IBR, but differences to the other stations were smaller.

Means of standardised organochlorines (PCBs + DDTs) in the tissues of *Macoma balthica* showed a clear gradient in the Turku region, with markedly higher values at Stn B1 (Fig. 7). In *Mytilus edulis* the picture was less clear, but the same station again emerged as the most polluted one.

In addition, normalised biomarker and tissue organochlorine levels in both species and sites combined showed significant linear regressions between total organochlorines and biomarkers ($n = 12$, $r^2 = 0.53$, $p =$
Lehtonen et al.: Biomarkers in Baltic Sea bivalves

0.008), ΣPCBs and biomarkers (n = 12, r² = 0.58, p = 0.004), total organochlorines and GST activity and ΣPCBs and GST activity (n = 13, r² = 0.64, p = 0.001, both cases). For heavy metals, Cd showed weaker but significant relationships with GST (n = 13, r² = 0.32, p = 0.045) and CAT activities (n = 13, r² = 0.33, p = 0.041).

### DISCUSSION

#### Chemical pollution and biomarker responses in the study regions

The high proportion of parent DDTs (DDT to ΣDDTs ratio) in tissues of both bivalve species collected from the Turku region indicates more recent inputs compared with the Tvärminne region. It is apparent that although not in active use anymore DDT is still circulating in the marine environment of this area and can therefore be found in unmetabolised and metabolised forms in both the sediments and the water phase, subsequently bioaccumulating in organisms. In the case of the Tvärminne region, the ratios of DDT to ΣDDTs show that, at present, DDT released into the environment decades ago is substantially ‘used up’ and increasingly found in the environment in metabolised forms, as DDDs and DDEs. However, with regard to biological effects, the poor correlations observed between biomarker responses and tissue levels of DDTs suggest that there is no direct relationship. With regard to PCBs, despite the fact that elevated levels were observed especially at Stn B1, some other, unmeasured anthropogenic substances potentially present at this station situated close to a former dumping site of harbour dredging waste can play a part in the impacts observed. Thus, the levels of organochlorines measured here should rather be taken as proxies of the level of general anthropogenic pollution at the different sites, not necessarily as the direct causes of the observed responses.

*Macoma balthica* from the Tvärminne region contains markedly higher amounts of Cd, Cu and Zn compared with those inhabiting the Turku region. The impact of the large Koverhar steel factory, present since the 1950s, although with significantly reduced pollutant release in recent years, can still be observed in terms of high levels of certain metals in the tissues of clams. Tissue concentration of Zn at the Koverhar station (1440 µg g⁻¹ dry wt) was strikingly high compared with other observations in the Baltic Sea (e.g.
Jankovski et al. (1996). Regoli et al. (1998) recorded levels of Cd between ca. 0.18 and 0.55 µg g⁻¹ dry wt, of Cu between ca. 13 and 37 µg g⁻¹ dry wt and of Zn between 200 and 700 µg g⁻¹ dry wt in *M. balthica* collected from several bays in the Russian Arctic. In the Tvärminne region the ranges for Cd and Zn are about twice as high, while the Cu levels are around 10-fold higher. In contrast to the other metals studied, the concentrations of Cd are markedly higher in *Mytilus edulis* compared with *M. balthica*. Studies by Lee et al. (1998) confirm that *M. balthica* is a good indicator of the accumulation of Zn, while Cd accumulates to a much lesser extent. In *M. edulis*, even higher Cd concentrations (4.05 µg g⁻¹ dry wt) have been recorded (Jankovski et al. 1996) in the Estonian part of the Gulf of Finland. In general, metal levels in bivalve tissues found here are within the ranges measured in these areas before (Kaitala 1981, Itkonen et al. 1998, Korhonen et al. 2001) and cannot be considered alarmingly high compared with observations in other sea areas.

However, *Macoma balthica* showed higher MT levels at Stn B1 in the Turku region compared with the stations in the Tvärminne region. In addition to binding by MTs, partitioning to metal-rich microgranules may play an important role in promoting metal tolerance of populations inhabiting sediments with elevated metal concentrations (Brown 1982, Klers & Weis 1987, Ballan-Dufresnais et al. 2001, Wallace et al. 2003). Apart from metals, there may also be other chemical stressors inducing MT formation at Stn B1 (see discussion below). In any case, when the 2 regions are compared, the content of MT in *M. balthica* does not appear to be always related to the tissue loads of metals.

Marked gradients observed in GST and CAT activity in *Macoma balthica* in the Turku region indicate that, at Stn B1, the presence of elevated chemical contamination intensifies phase II conjugation mechanisms and enhances hydrogen peroxide elimination related to the oxidative stress complex. The increased CAT signal may also be related to the elevated levels of MT, since these proteins are also a part of the antioxidant defence system (Viarengo et al. 2000). In vertebrates, the MT gene has been shown to be induced under pro-oxidant conditions (Dalton et al. 1996). In rainbow trout, activation of MT gene transcription by hydrogen peroxide, the substrate of CAT, has been demonstrated (Olsson et al. 1995). However, since the production of reactive oxygen species (ROS) is strongly linked to cytochrome P-450 induction, which in mussels (*Mytilus* spp.) is only about 1 % of that found in mammals, it has been argued that exposure to organic pollutants and MT induction through subsequent ROS production is not a significant process in mussels (Viarengo et al. 2000).
1999). Nevertheless, other ROS-producing detoxification pathways not mediated by P-450s may be in operation. In this study, the data on organic contaminants in the soft tissues of *M. balthica* with marked gradients in PCBs and DDTs in the Turku region corresponds well with the gradients observed in the levels of GST, CAT and MT in the same area. This implies that detoxification reactions and metabolic responses to oxidative stress may be caused by the catabolism of organic xenobiotics.

The observed interspecies differences, with *Macoma balthica* expressing 3- to 4-fold (CAT) and 7- to 8-fold (GST) higher enzyme activities in comparison with *Mytilus edulis*, most likely do not only represent species-characteristic ecological levels but also mirror the observed tissue loads of organic contaminants, which in the sediment-dwelling *M. balthica* are on average 2-fold (DDTs) or 3-fold (PCBs) higher than in *M. edulis*. Compared with *M. edulis*, *M. balthica* showed markedly higher tissue accumulation of Cu, Zn, PCBs and DDTs, whereas the levels of Cd were higher in *M. edulis*. Interspecies variability in MT levels was apparent, with 1.5-fold lower values in *M. edulis* compared with *M. balthica*, as observed by Lehtonen & Leiniö (2003). This is probably due to the fact that *M. balthica* lives within soft sediments and is mostly a deposit feeder, thus being constantly exposed to pollutants deposited in organic particles of sediments that it feeds on, as well as soluble pollutants in interstitial water. Being a suspension feeder, *M. edulis* is exposed to considerably lower levels of pollutants in the suspended particulate matter or soluble compounds present in seawater. Besides biological differences in uptake kinetics, the difference may be due to the sediment-dwelling lifestyle of *M. balthica*, which raises its exposure to, for example, bioavailable metals in both their physical environment (hypoxic/anoxic sediment layers, pore water) and food items. In addition, metabolic differences between the species could also play an important role in the observed differences in tissue pollutant levels between the species.

Finally, with regard to the observed homogeneity in the levels of AChE activity the results obtained here confirm the view that organophosphate/carbamate pesticides are not an environmental problem in Finnish coastal waters. Other chemical stressors that can cause AChE-coupled neurotoxic effects do not appear to be present in effective concentrations.

**Application of biomarkers in the Baltic Sea**

Among many other specific characteristics of the Baltic Sea, the constant low-salinity environment (<6.5 in Finnish coastal waters) creates physiological stress to marine species that may affect biomarker responses. Thus, basic information on the levels of biomarker responses and their natural variations is needed prior to the application of these techniques and use of these species in monitoring of pollutant-induced biological effects in the Baltic Sea ecosystem. Information is needed not only on the effects of potentially confounding factors such as eutrophication and the resulting hypoxic conditions in the near-bottom waters (commonly observed e.g. in the Gulf of Finland), but also on the physiological adaptations of species inhabiting these areas.

*Mytilus edulis* is a marine species able to penetrate into brackish environments; however, many of its physiological, morphological and growth characteristics are distinctively modified in the Baltic Sea (e.g. Kautsky et al. 1990, Tedengren et al. 1990). The low-salinity stress is extreme in the northern part of the Baltic, where the distribution of *M. edulis* comes to an end at salinities <4.5 (Lassig 1965). Therefore, concerning this species some precautions should be taken in the interpretation of biomarker data when comparing the responses observed in northern Baltic Sea populations to those inhabiting more saline southern regions (up to ca. 15) or fully marine environments. With regard to *Macoma balthica*, populations thrive in the Baltic Sea down to a salinity of 2 (Lassig 1965) if other main environmental factors such as sediment type, dissolved oxygen content of near-bottom water, food supply and water depth are suitable. Compared with mussels, *M. balthica* is physiologically much better adapted to the conditions prevailing, for example, in the Gulf of Finland, and the responses observed in the present study are thus more comparable with those measured in other areas of the species’ wide distribution.

Salinity can have an effect on the ‘baseline’ levels of biomarkers, as potentially is the case with MT. In *Mytilus edulis*, a gradient in MT levels was observed in the Tvärminne region, while the average levels did not differ from those recorded in the Turku region. No gradients could be observed in the tissue levels of metals, nor were the mean levels different between the areas. When examined together with the results obtained during the Baltic Sea component of the BEEP project, the mean MT level in *M. edulis* was observed to elevate by about 2-fold towards the north, seemingly along the salinity gradient (Leiniö & Lehtonen 2005, Barsiené et al. 2006, Kopecka et al. 2006, Schiedek et al. 2006, this study). Concerning *Macoma balthica*, the only data available on MT levels in the southern Baltic Sea also suggest salinity-related signal levels: a mean of ca. 300 µg g⁻¹ wet wt in spring in the Gulf of Gdańsk (BEEP project, unpubl. data) is markedly lower compared with the average value of 442 µg g⁻¹ wet wt obtained in the present study. The higher MT levels
might be connected to chemical speciation of certain metals resulting in higher bioavailability in low-saline environments (e.g. *M. balthica*: Lee et al. 1998). On the other hand, in the Russian Arctic no significant effect of salinity on the MT level of either species could be observed (Amiard-Triquet et al. 1998). In some cases, changes in general protein metabolism caused by natural factors have been shown to modulate MT content more than metal accumulation in tissues (Legras et al. 2000). Since MT levels exhibit a marked natural oscillation during the year in both species in the study region (Leiniö & Lehtonen 2005) and also elsewhere (e.g. Bordin et al. 1997), food conditions and reproductive stage must also be considered when interpreting the data (e.g. Mourgaud et al. 2002). The extensive seasonal variability in temperature in the northern Baltic Sea also affects the reproductive cycle of marine organisms, potentially having a marked effect on various biomarker responses (Leiniö & Lehtonen 2005). Regardless of the cause, the level of the MT induction was highest in the present study area compared with the other Baltic Sea areas studied.

### Integrated biomarker response approach

The use of integrated biomarker indices is a relatively new approach in the marine ecotoxicological context (Adams et al. 1993, Narbonne et al. 1999, Beliaeff & Burgeot 2002, Chèvre al. 2003a,b, Broeg & Lehtonen 2006). Integrating the numerical measures of a set of biological signals that reflect malfunctions or, potentially, a decreased performance in organisms into a single index has advantages considering the rough evaluation of the ‘pollution status’ of marine populations or environments which they inhabit. Chemical analyses can only show the concentrations of single contaminants in the environment, while an integrated biomarker index demonstrates also their bioavailability, interactive effect and combined effects. Since comparing areas with highly differing environmental characteristics that govern bioavailability and biological effects of different pollutants usually allows only very limited conclusions, a rational way is to treat the systems as a whole, i.e. using integrated response approaches that demonstrate more complex *in situ* signals in populations at each particular site. The IBR index used in this study merged the comparative information obtained from 4 biomarkers, and the value obtained this way related distinctly to the concentrations of tissue organochlorine residues measured at the most contaminated site in comparison with the other sites. The use of an integrated biological measure is of practical importance when examining, for example, temporal developments along different suspected pollution gradients or comparing different geographical regions. As our results show, this approach also has high potential in detecting biological effects along comparatively small-range contamination gradients.

### CONCLUSIONS

The results obtained indicate a pollution gradient in the Turku region, identified both by measurements of selected biomarker responses in bivalves and by concentrations of selected pollutants in their tissues. The gradient was clearly indicated in the sediment-dwelling *Macoma balthica*, while the suspension-feeding *Mytilus edulis* showed a more ambiguous distribution of tissue pollutants and biomarker responses. In the Tvärminne region such an obvious gradient was not detected, although variability in pollutant levels and biomarker signals between the stations was observed. Overall, the Turku region appears to be much more affected by organic anthropogenic pollutants, while the Tvärminne region is characterised by a higher load of heavy metals.

This study is the first of its kind performed in the coastal waters of Finland. The results obtained demonstrate the usefulness of the multibiomarker approach, including the integrated biomarker response index. Using 2 sessile species with markedly differing feeding modes, living habitats and, therefore, routes of exposure to environmental pollutants is considered a rational strategy for assessing the pollution status of coastal sea areas. Using an integrated response measure, one can detect possible synergistic effects of different compound groups (e.g. organochlorines and heavy metals) that might remain hidden if only one biomarker was used. Regarding the species studied, the infaunal, soft-sediment clam *Macoma balthica* appears an increasingly promising organism for biological-effects monitoring in the northern Baltic Sea.

**Acknowledgements.** Tvärminne Zoological Station (University of Helsinki) and Archipelago Research Station (University of Turku) are thanked for assistance in sampling. Dr. P. Mora (University of Bordeaux I) is thanked for advice on GST and CAT activity measurements and Ms. M. Kronholm for testing of the protocols in the early phase of the project. U. Karlström (MSc) assisted in the analyses. Ms. S. Lage and Mr. J. Riionen skilfully executed the organochlorine and metal analyses, respectively. We thank Dr. D. Schiedek for editorial suggestions on the final version of the manuscript.

**LITERATURE CITED**

response to metal contamination: validation in the field. Biomarkers 7:479–490
Schiedek D, Broeg K, Baršenò J, Lehtonen KK and 7 others (2006) Biomarker responses and indication of contaminant effects in blue mussel (Mytilus edulis) and eelpout (Zoarces viviparus) from the western Baltic Sea. Mar Pollut Bull (in press)

Editorial responsibility: Otto Kinne (Editor-in-Chief), Oldendorf/Luhe, Germany

Submitted: June 2, 2004; Accepted: March 2, 2006
Proofs received from author(s): August 30, 2006