Relationships between in vitro lymphoproliferative responses and levels of contaminants in blood of free-ranging adult harbour seals (Phoca vitulina) from the North Sea

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ABSTRACT

In vitro culture of peripheral blood leucocytes (PBLs) is currently used in toxicological studies of marine mammals. However, blood cells of wild individuals are exposed in vivo to environmental contaminants before being isolated and exposed to contaminants in vitro. The aim of this study was to highlight potential relationships between blood contaminant levels and in vitro peripheral blood lymphocyte proliferation in free-ranging adult harbour seals (Phoca vitulina) from the North Sea. Blood samples of 18 individuals were analyzed for trace elements (Fe, Zn, Se, Cu, Hg, Pb, Cd) and persistent organic contaminants and metabolites (ΣPCBs, ΣHOPCBs, ΣPBDEs, 2-MeO-BDE68 and 6-MeO-BDE47, ΣDDXs, hexachlorobenzene, oxychlordane, trans-nonachlor, pentachlorophenol and tribromoisocyanurate). The same samples were used to determine the haematology profiles, cell numbers and viability, as well as the in vitro ConA-induced lymphocyte proliferation expressed as a stimulation index (SI). Correlation tests (Bravais-Pearson) and Principal Component Analysis with multiple regression revealed no statistically significant relationship between the lymphocyte SI and the contaminants studied. However, the number of lymphocytes per millilitre of whole blood appeared to be negatively correlated to pentachlorophenol (r = −0.63, p = 0.005). In adult harbour seals, the interindividual variations of in vitro lymphocyte proliferation did not appear to be directly linked to pollutant levels present in the blood, and it is likely that other factors such as age, life history, or physiological parameters have an influence. In a general manner, experiments with in vitro immune cell cultures of wild marine mammals should be designed so as to minimize confounding factors in which case they remain a valuable tool to study pollutant effects in vitro.

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1. Introduction

During these last decades, large-scale mortalities of marine mammals have been increasingly documented (Greenland and Limpus, 2007; Osinga et al., 2012). Epizootics and mass strandings events have impacted numerous cetacean and pinniped species (Colegrove et al., 2005; de Swart et al., 1995a; Duignan et al., 1995; Greenland and Limpus, 2007; Hanni et al., 1997; Lipscomb et al., 1996; Rogan et al., 1997). In 1988 and 2002, two major epizootics due to the phocine distemper virus (PDV) affected the harbour seal (Phoca vitulina vitulina) population inhabiting the North Sea (Härkönen et al., 2006; Müller et al., 2004). The population of the Wadden Sea situated along the east and south-east coasts of the North Sea, suffered losses up to approximately 57% and 47% in 1988 and 2002, respectively (Dietz et al., 1989; Härkönen et al., 2006). As a consequence, these mass die-offs have initiated numerous pathological and toxicological studies (Das et al., 2008; de Swart et al., 1996; Ross et al., 1996b; Siebert et al., 2007, 2012).
that notably discussed the impact of environmental pollutants on marine mammal health. Indeed, seals are exposed to a variety of anthropogenic contaminants mainly through their diet. As top predators, they are particularly at risk for bioaccumulating contaminants that biomagnify up the trophic food chain (De Guise et al., 2003). In fact, the chemical compounds known to bioaccumulate in harbour seal tissues are toxic trace elements including mercury (Hg) (Das et al., 2003) and persistent organic pollutants (POPs) such as organohalogenated compounds (polychlorinated biphenyls – PCBs), the dichlorodiphenyltrichloroethanes (DDTs), polybrominated diphenyl ethers (PBDEs), hexachlorobenzene (HCB) and chlordanes (O’Shea and Tanabe, 2003; Reijnders, 1980).

Evidence suggests that various immunotoxic contaminants (e.g. PCBs and metabolites) might have contributed at least partly to the severity and extent of the epizootics (Ross, 2002). Pollutants may indeed have toxic effects on peripheral blood leucocytes (PBLs), including lymphocytes, monocytes and granulocytes, thereby diminishing defenses against invading pathogens (Lahvis et al., 1995; Levin et al., 2007; Mos et al., 2006; Nakatsuru et al., 1985). Exposure to pollutants was hypothesized to be notably related to the high levels of parasitic or bacterial infections observed in seals found dead or killed due to severe illness before and after the 2002 epizootic (Siebert et al., 2007).

Blood sampling of marine mammals is considered as a convenient and minimally invasive method, relevant for the analyses of pollutants and biomarkers (Habran et al., 2011, 2013; Vanden Berghe et al., 2012), and for in vitro assays (Ross et al., 2003a). In vitro cultures of PBLs are indeed routinely used to assess the relationship between environmental contaminants and immune alterations in free-ranging marine mammals (Cámara Pellissós et al., 2008; De Guise et al., 1998; Kakuschke et al., 2008; Levin et al., 2005, 2007).

Organic and inorganic pollutants were analyzed recently in harbour seals from the North Sea showing high concentrations compared to other species or locations (Das et al., 2008; Habran et al., 2013; Vanden Berghe et al., 2012; Weijs et al., 2009a,b,c). PCBs often exceed the threshold values of 20 ng PCBs/g wet weight for seal blood (Weijs et al., 2009a) and 17 μg PCBs/g lipid weight for marine mammal blubber (Jepson et al., 2005; Siebert et al., 2012; Weijs et al., 2009b) above which deleterious effects might be observed (Kannan et al., 2000). No threshold has been documented for other compounds such as pesticides, brominated compounds or toxic metals in this species, but deleterious effects of POPs and metallic trace elements on the immune system have been documented in both in vivo and in vitro exposures in laboratory animals and wildlife (Cámara Pellissós et al., 2008; Dufresne et al., 2010; Kakuschke and Prange, 2007; Kakuschke et al., 2005; Luebke et al., 1997; Ross et al., 1996c). Associations have been established between environmental contaminants and immunotoxicity in free-ranging seal populations (Das et al., 2008; de Swart et al., 1996; Dufresne et al., 2010; Ross et al., 1996a,b;c; van Loveren et al., 2000) as demonstrated for instance by significant impairment of natural killer cell activity and specific T-lymphocyte responses in vitro (Ross et al., 1996b). Nevertheless, the current knowledge of those cellular effects remains fragmentary. Moreover, the influence that the blood contaminants present in free-ranging harbour seals may have on in vitro immune responses once PBLs are isolated and cultured in controlled conditions is unknown. This point is highly relevant when cell viability and proliferation are evaluated in the framework of in vitro exposure to contaminants.

The present study aims to highlight potential relationships between blood pollutant levels and in vitro immune responses of free-ranging adult harbour seal from the North Sea. We determined the in vitro mitogen-induced lymphocyte proliferation and a broad range of blood parameters including haematology profiles, lymphocyte numbers and viability, essential element concentrations as well as non-essential element, persistent organic pollutant (POP) and naturally-produced methoxylated polybrominated diphenyl ether (MeO-PBDE) levels.

2. Material and methods

2.1. Blood sample collection and conservation

Blood samples were collected from 18 free-ranging harbour seals caught on sandbanks of the German part of the Wadden Sea: 3 from Kolumbus Loch (geographical coordinates 54° 23′ N and 8° 35′ E) in April (spring) 2010; and 15 on Lorenzenplate (54° 25′ N and 8° 38′ E) in September (autumn) 2010 and April (spring) 2011 (Fig. 1). Seals were captured in seine nets at inter-tidal haul-out sites. They were transferred to individual tube nets and restrained manually for clinical examination. Sex, length and weight were determined before blood samplings (Table 1). Collection procedures were designed to preserve sample integrity and minimize sample contamination during processing and handling. All sampled animals were considered older than 3 years (adults) based on length and weight (Weijs et al., 2009b). During the sampling procedure, the animals were continuously under care of two experienced veterinarians. All individuals were released at the site of capture after completing the investigations. The maximum detention time, between the transfer of all animals in tube nets and the release of the last one, was one hour. All animals appeared healthy by veterinary check. They were in normal nutritional status and showed no sign of disease. The field studies were carried out under the relevant permits of the National Park Office Schleswig-Holstein and the animal experiment permit (AZ 312-72241.121-19).

Blood was drawn from the extradural venous sinus into sterile evacuated blood collection tubes using a 20 ml syringe and a 1.2 mm × 100 mm needle (Carromco, Hamburg-Norderstedt, Germany). Blood was transferred into serum tubes S-Monovette® for POP analysis and in 7.5 ml S-Monovette® tubes treated with Lithium-Heparin (LH) for trace metal analysis (Sarstedt, Essen, Belgium). Serum for POP analysis was isolated by centrifugation at 1500 × g for 20 min at 21°C (Multifuge 3 S-R, Kendro, Hamburg, Germany). Whole blood (2 ml) and serum samples (2 ml) were stored at −20°C until analysis.

Blood samplings were also used to test the in vitro lymphocyte proliferation. Blood (15–30 ml) was collected in Cell Preparation Tubes with Sodium Heparin (CPT™, BD Vacutainer®, Plymouth, UK) that were gently inverted 10 times and stored upright at room temperature until further processing, within maximum 5 h of blood collection. Additional blood samples were taken to determine the clinical blood chemistry and haematology profiles, and other health parameters as part of an ongoing health monitoring.

2.2. Haematological profiles

The haematology profiles (white blood cells [WBC], red blood cells [RBC], haemoglobin [HbG], hematocrit [HCT], platelets [PLT]) were determined for all whole blood samples (before PBL isolation) using a scil Vet abc haematology analyzer (scil Animal Care Company GmbH, D-68519 Viernheim, Germany) which was calibrated for seal blood (Hasselmeier et al., 2008).

2.3. PBL isolation and conservation

PBLs were isolated according to the principle of density gradient centrifugation on Ficoll Hypaque™ (density: 1.077 g/ml) contained in CPT™ (BD Vacutainer®), using a modified protocol described by de Swart et al. (1993). The blood samples contained in CPT™ (BD Vacutainer®) were centrifuged at 1800 × g for 30 min at room temperature (20°C). Cells were then resuspended into the plasma
by 10 gentle inversions to improve cell viability during transport-
ination at room temperature, as advised by the manufacturer
(BD Benelux N.V., Erembodegem, Belgium). The Ficoll Hypaque™
density fluid contained in CPTubes™ (BD Vacutainer®) is origi-
nally used to separate peripheral blood mononuclear cells (PBMCs,
cluding lymphocytes and monocytes) and plasma from granu-
ocytes and erythrocytes. However, flow cytometry scatter plots
of cell suspensions indicated that all leucocyte subpopulations
(i.e., lymphocytes, monocytes and granulocytes) were present (see
point 2.4, Fig. 2). After transportation to the cell culture labora-
tory within 18 h, the PBLs were pipetted into separated vessels and
washed 3 times in phosphate buffered saline (PBS, sterile filtered,
BioWhittaker®, Lonza, Verviers, Belgium). Cells were suspended in
RPMI 1640 medium (BioWhittaker®, Lonza, Verviers, Belgium) sup-
plemented with 10% foetal calf serum (Gibco®, Invitrogen, Paisley,
UK), 0.33% L-α-aminola-glutamine (GlutaMAX™, Gibco®, Invitrogen,
Paisley, UK), 1% non-essential amino-acids (BioWhittaker, Lonza,
Verviers, Belgium), 1% Na pyruvate (BioWhittaker, Lonza, Verviers,
Belgium) and 1% penicillin-streptomycin (100 IU and 100 μg/ml,
BioWhittaker, Lonza, Verviers, Belgium). This complete culture
medium was further referred to as culture medium. The viability
and number of cells were determined with a Nucleocounter® NC-
100™ (Chemometec, Allerød, Denmark). The NucleoCounter® is an
integrated fluorescence microscope designed to detect signals from
the fluorescent dye, propidium iodide (PI) bound to DNA of cells
with compromised cell membranes. The cell viability of samples
was determined using the total cell count (following a treatment
rendering all the cell nuclei susceptible to staining) and the count
of non-viable cells (Chemometec A/S, 2013). After counting, cells
were centrifuged, and the pellet was resuspended in a determined
volume of culture medium added with 10% DMSO (dimethyl sulfox-
ide, Sigma–Aldrich Chemie GmbH, Steinheim, Germany) at 4 °C to
adjust their concentration to 10 × 10⁶ viable cells/ml. The cell sus-
ension was transferred into cryovials (1 ml/vial) and placed into
a freezing container (Mr Frosty, Nalgene) at −80 °C during 24 h,
before being transferred in liquid nitrogen.

2.4. PBL in vitro cultures

PBL cryovials were quickly thawed and transferred into 50 ml
Falcon tubes containing 40 ml of complete culture medium pre-
warmed at 37 °C. They were centrifuged at 300 × g during 10 min,
neutrophils, (FSC) centrifugation bodegem, populations Fig. 2. Flow cytometry scatter plot of cell populations obtained after whole blood centrifugation in CPT®. Four subpopulations were gated based on forward scatter (FSC – size) and side scatter (SSC – internal complexity) characteristics. The gate A most likely includes lymphocytes. Gates B and C probably include monocytes and neutrophils, respectively. Gate D predominantly includes early apoptotic and dead cells, and the square in the left corner includes cell debris.

the supernatant was discarded and the cell pellet was resuspended in warm culture medium. Monocytes were partially depleted by plastic adherence in tissue culture flasks (Vented Cap, Sarstedt, Newton, NC, USA) incubated during 1 h at 37 °C in a humidified atmosphere enriched with 5% CO₂. The cell suspension was harvested, the living cells were counted with a nucleocounter® (Chemometec, Allered, Denmark) and the concentration was adjusted to 1 × 10⁶ cells/ml (final concentration). Cells were also co-stained with propidium iodide (PI) and Annexin V-fluorescein isothiocyanate (FITC) conjugate according to the manufacturer’s instructions (Annexin V: FITC Apoptosis Detection Kit I, BD, Erembodegem, Belgium) and analyzed by flow cytometry in order to determine the proportions of viable, early apoptotic and dead cells (in late apoptosis and necrosis). The Annexin V assay was used to detect phosphatidylserine translocation on the membrane surface of cells undergoing apoptosis (Vermes et al., 1995). Cells were displayed as scatter plots based on their sizes, estimated by forward scatter (FSC), and internal complexity (granulosity), estimated by side scatter (SSC). The analysis was conducted with a FACSCanto™ II (BD, Erembodegem, Belgium) flow cytometer. The light source used was an air-cooled laser tuned to 488 nm. FITC-fluorescence (FL1) was collected through a 502 nm long pass (LP) filter and then through a 530/30 nm bandpass filter, and PI fluorescence through a 585/42 nm LP filter (FL2), and through 655 and 670 nm LP filters (680 nm, FL3). Five-parameter (FSC-A, SSC-A, FL1, FL2, and FL3) data of minimum 50,000 events were collected and analyzed for each sample using FACS Diva software version 6.1.2. Scatter plots obtained by flow cytometry showed that all leucocyte subpopulations (i.e., lymphocytes, monocytes and granulocytes) were present, and that cell suspensions were composed of lymphocyte-enriched leucocytes (Fig. 2). All cells were analyzed for AV-FITC and PI labelling. In all likelihood, the gate A included lymphocytes (mostly but not exclusively) as they are characterized by small sizes and low complexity compared to the other leucocyte types (Fig. 2). However, the exact position and quantification of leucocyte subpopulations in scatter plots was not possible as no specific or suitable marker was available for harbour seal at the time of experiment. Monocyte and neutrophil (most abundant type of granulocyte) gates were thus hypothetically drawn (gates B and C in Fig. 2) based on sizes and complexity characteristics as well as on literature data (Pillet et al., 2002; Zharinov et al., 2006). The presence of monocytes (despite depletion) and granulocytes was confirmed by microscopy analysis of cells stained with Giemsa (Modified Solution, Fluka Chemika, Sigma–Aldrich, Steinheim, Germany) (data not shown). Lymphocyte-enriched leucocytes were further referred to as PBLs.

Proliferation to Concanavalin A (ConA, Sigma–Aldrich Chemie GmbH, Steinheim, Germany), a T-cell-specific mitogen in harbour seals (de Swart et al., 1993), was assessed by culturing PBLs in a flat bottom 96-well plate (transparent, Sarstedt, Newton, NC, USA) at 2 × 10⁵ cells in 200 μl culture medium per well with an optimal concentration of 5 μg/ml ConA in an incubator at 37 °C under humidified atmosphere enriched with 5% CO₂. Control PBLs were cultured in medium alone and each culture condition was in triplicate. The actively proliferating lymphocytes were assayed with bromodeoxyuridine (BrdU) incorporation into newly synthesized DNA (BrdU Proliferation Assay, Calbiochem®, Overijse, Belgium). Two types of controls were used for the BrdU tests: blanks (culture medium alone) and background controls (cells without BrdU).

Each culture condition was in triplicate. BrdU label was added to wells of the microtiter plate for the last 24 h of culture. After 72 h of incubation, cells were fixed, permeabilised and the DNA denatured to enable antibody (Ab) binding to the incorporated BrdU. Detector anti-BrdU monoclonal Ab was pipetted into the wells and allowed to incubate for 1 h, during which time it binds to any incorporated BrdU. Unbound Ab was then washed away and horseradish peroxidase-conjugated goat anti-mouse was added to bind to the detector Ab. The horseradish peroxidase catalyzed the conversion of the substrate to a blue product, which is then converted to yellow after the addition of the solution which stops the reaction. The intensity of the coloured reaction product, which is proportional to the amount of incorporated BrdU in the cells, was quantified by measuring the absorbance (optical density, OD) in each well using a spectrophotometric plate reader (PowerWave X, Bio-Tek Instruments, Inc., Winooski, VT, USA) at dual wavelength of 450–540 nm. The signals of proliferating cells were significantly higher than the corresponding blanks and background controls. For each well OD, the average blank absorbance was subtracted. Lymphocyte proliferation was expressed as a stimulation index (SI), calculated as the ratio of the proliferation of mitogen-induced stimulated cells to that of cells cultured without mitogen (Levin et al., 2005).

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SI = \frac{OD \text{ of stimulated cells}}{OD \text{ of corresponding control}}
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In addition, to confirm the presence of lymphoblasts in stimulated conditions and the lack of cytotoxicity in control wells, the corresponding cell suspensions were stained with PI and Annexin V (Annexin V: FITC Apoptosis Detection Kit I, BD, Erembodegem, Belgium) and analyzed by flow cytometry as described above.

2.5. Chemical analysis

2.5.1. Trace elements

After being lyophilized, approximately 200 mg of homogenized whole blood powder were digested in Teflon jar with concentrated HNO₃, H₂O₂ and ultrapure water (Milli-Q® ultrapure water for trace element analysis, Millipore, Billerica, MA, USA) in a microwave oven (20 min between 0 and 600 W, ETHOS, Milestones). After cooling, samples were diluted to 50 ml with Milli-Q water in a volumetric flask. The trace elements Cd, Fe, Ni, Cu, Se, Pb, and Zn were analyzed by Inductively Coupled Plasma-Mass Spectrometry (ICPMS, Elan DRC II, PerkinElmer, Shelton, USA). Total mercury (T-Hg) was determined by atomic absorption spectrophotometry with the Direct
Mercury Analyzer (DMA80, Milestones, Sorisole, Italy). Concentrations are expressed in μg/g dry weight (dw). Concurrent analyses of a set of certified reference materials (CRM) and of procedural blanks were repeated throughout each set to ensure method’s accuracy and precision. The control materials used were SERONORM™ Human, Trace Element Whole Blood Level-3 (SERO AS, Billingstad, Norway) and DOLT-3, Dogfish Liver Certified Reference Material for Trace Elements (National Research Council Canada, Ontario, Canada). The handling and preparation procedures were identical for all samples and reference materials analyzed.

2.5.2. Organic contaminants

In all serum samples, 29 PCBs (ΣPCBs), 18 HO-PCBs (ΣHO-PCBs), 7 PBDEs (ΣPBDES), 2 MeO-PBDEs (2-MeO-BDE68 and 6-MeO-BDE47), ΣDDXs (o,p’-DDE, p,p’-DDE and metabolites o,p’-DDD, p,p’-DDD and p,p’-DDE), hexachlorobenzene (HCB), oxychlordane (OxCl), trans-nonachlor (TN), pentachlorophenol (PCP) and 2,4,6-tribromoanisole (TBA) were targeted. More precisely, investigated PCBs were IUPAC numbers (HCB) 28, 52, 74, 87, 95, 99, 101, 105, 110, 118, 128, 138, 146, 149, 153, 156, 170, 171, 172, 174, 177, 180, 183, 187, 194, 199, 196/203, 206, 209. Targeted HO-PCB congeners were 3HO-PCBs (# 118, 138, 153 and 180), 4HO-PCBs (# 79, 107, 120, 127, 130, 146, 162, 163, 172, 177, 187, 193 and 208) and di-4HO-CB202. Analyzed PBDEs were IUPAC # 28, 47, 99, 100, 153, 154 and 183.

The method for serum analysis was slightly adapted from the methods described previously (Weijis et al., 2009a). A volume of serum (1.5 ml) was spiked with internal standards (PCB 143, BDE 77 and 4’-HO-BC 195), diluted 1:1 with Milli-Q water, mixed with formic acid, sonicated for 20 min and extracted using solid-phase extraction (SPE) cartridges (6 ml/500 mg Oasis HLB, Waters). Elution was done by 10 ml of MeOH: dichloromethane (DCM, 1:1). After evaporation to near dryness, the analytes were reconstituted in 500 μl hexane:DCM (1:1) and fractionated on silica SPE cartridges (3 ml/500 mg, Varian) topped with 200 mg acid silica (44%, w/w). A first fraction containing PCBs and PBDEs was eluted with 5 ml hexane, while the phenolic compounds were eluted in a second fraction with 6 ml DCM (1:1, v/v). Both fractions were evaporated to dryness. The first fraction was reconstituted in 100 μl iso-octane, while the second fraction was derivatized for 30 min with diazomethane to form MeO-PCBs and MeO-PBDEs after which it was evaporated to dryness and reconstituted in 75 μl iso-octane.

For the analysis of PBDEs and MeO-derivatives, a GC-MS operating in electron capture negative ionization (ECNI) mode was equipped with a 30 m × 0.25 mm × 0.25 μm DB-5 capillary column (J&W Scientific, Folsom, CA, USA). The MS was used in the SIM mode with two ions monitored for each MeO-PBDE congeners in specific windows, while ions m/z = 79 and 81 were monitored for MeO-PCBs and for PBDEs during the entire run. For the PCB and organochlorine pesticides (OCP) analysis, a GC-MS operating in electron impact ionization (EI) mode was equipped with a 25 m × 0.22 mm × 0.25 μm HT-8 capillary column (SGE). The MS was used in the SIM mode with two ions monitored for each PCB homologue group or individual OCP in specific windows. Multi-level calibration curves (r² > 0.999) in the linear response interval of the detector were created for the quantification. Quality control (QC) was performed by regular analyses of procedural blanks, by random injection of standards, spiked samples and solvent blanks. The QC scheme is also assessed through regular participation to interlaboratory comparison exercises organized by AMAP (POPs in human serum). Obtained values were deviating with <20% from consensus values. For analytes detected in procedural blanks, the mean procedural blank value was used for subtraction. After blank subtraction, the limit of quantification (LOQ) was set at 3 times the standard deviation of the procedural blank. For analytes not detected in procedural blanks (HO-PCBs and MeO-PBDEs), LOQs were calculated for a signal-to-noise ratio = 10. Results were expressed on a per serum volume basis as the lipid fraction of serum is composed of more polar lipids when compared to subcutaneous fat or other organs (Henderson et al., 1994), and contains non-polar lipids (i.e. hexane-extractable fat) at lower levels (Debier et al., 2003). Expressing POP concentrations in serum on a lipid weight basis (i.e. hexane-extractable fat) would thus lead to an overestimation (Debier et al., 2003).

2.6. Data analysis

Statistical analysis was performed using Statistica 9.1. (StatSoft, Maisons-Alfort, France). Each culture condition was in triplicate for each individual. Optical density values (BrDU test) and cell counts were averaged for each individual, giving one mean value per culture condition. Coefficients of variations (CV) were calculated as the ratio of the standard deviation (SD) to the mean. Comparisons of the OD means of control and stimulated PBLs after the 72 h of incubation was assessed by the Wilcoxon test for paired samples. The normality of all data distribution was assessed by visual inspection and using the Shapiro–Wilk test. All data included length, weight, haematology values, cell numbers and viability, SI, essential and non-essential element levels, and persistent organic pollutant (POP) and naturally produced methoxylated polybrominated diphenyl ether (MeO-PBDE) levels. All values were log-transformed (log10) to achieve normal distributions. Comparisons of the means of lymphocyte stimulation indexes between sexes, locations, seasons and years of sampling were performed using the Student’s t test (two-tailed), respecting the normality of distribution and equality of variances (Bartlett, p > 0.05). When variances were unequal, the Satterthwaite’s approximate t test was used. The level of statistical significance was defined at p < 0.05. When normality of distribution could not be achieved despite the log-transformation, the non-parametric Mann–Whitney U test was used. As sample sizes were small (n < 20 for each group), the obtained U values were compared to respective critical U values (Appalachian State University, 2013).

For chemical element measurements, each value below LOQ was replaced with a value of LOQ/2 and used in the statistical tests. In the other tests performed (correlation tests, Principal Component Analysis and regressions), pollutants with more than 50% of values < LOQ were suppressed as most of their values were identical (LOQ/2), thus presenting a low variability. After log-transformation, none of the considered distributions differed significantly from normality as shown by the Shapiro–Wilk tests (all W > 0.897 and p > 0.05). Correlation tests (Bravais-Pearson) were used to investigate potential relationships between in vitro lymphoproliferative responses and peripheral blood contaminant and trace element concentrations two-by-two. Then, a more global approach was foreseen, considering pollutants as combined compounds as they can act as a mixture with synergic and antagonistic behaviours. A Principal Component Analysis (PCA) was realized in order to reduce the number of original pollutant variables. Then, the two first factor scores revealed by the PCA were used in a multiple regression test as the independent variables, with SI and lymphocyte numbers as dependant variables.

3. Results

3.1. Levels of trace elements in whole blood

Trace element levels were all above the detection limit except that for Cd, for which seven seals sampled in 2011 exhibited non-detectable concentrations (Table 2). A general LOQ mean value was calculated for all individuals, and when some individuals presented concentrations below their own LOQ, their number was added
between parentheses. For example, Cd presented 12 values below its LOD, of which 7 below its LOD. The essential trace elements exhibited minor variations in concentration levels, with relative standard deviation (RSD) values lower than 20% for Fe, Cu, and Zn, and 28% for Se. Their RSD expressed greater interindividual variations with 32% for T-Hg, 55% for Pb and 142% for Cd. The median Se/T-Hg molar ratio was 7.9 (min. – max. = 4.7–17.6).

No significant difference of medians (Mann–Whitney U test) was observed between genders, seasons of sampling (spring-autumn) except for Cd and Fe (p < 0.05), years of sampling (2010–2011) except for Cd (p < 0.01), and places of sampling (KL-LP) except for Zn (p < 0.01).

2.2. Levels of POPs in serum

Table 3 summarizes ΣPCBs, ΣHO-PCBs, ΣDDXs, PCP, ΣPBDEs, TBA, TN, OxC and HCB serum concentrations (pg/ml). The contaminants found at the highest concentrations were PCBs, constituting 86% of the mean total POPs, followed by their hydroxylated metabolites HO-PCBs, and total DDT (ΣDDXs) of which the contribution of p,p’-DDE was up to 93%. The PCB profile of the 18 animals was CB 153 > CB 138 > CB 187 > CB 146 > CB 99. The HO-PCB profile was 4-HO-PCB > 107 > 162 > 164 > 187. The 4-HO-PCB 107, one of the lowest chlorinated compounds, indeed accounted for 49% of the sum in average (min. 24–max. 58%). Its median concentration was two times higher than that of 4-HO-PCB 162 and ten times that of 4-HO-PCB 187. Some HO-PCBs (3-HO-PCB 118, 4-HO-PCB 127 and 3-HO-PCB 180) were not found in any sample, while 4-HO-PCB 79 and 4-HO-PCB 193 were detected in less than 25% of all samples. The mean ratio ΣHO-PCBs/ΣPCBs equaled 0.10 and there was a good correlation between the two groups of pollutants (r² = 0.76).

For PBDEs, BDE 47 was the predominant congener followed by BDE 100. Trans-nonachlor and oxychlordane levels appeared variable among individuals. Among extreme values, 6 individuals presented concentrations below or just above the LOD, and 3 animals sampled in September 2010 presented concentrations 10 times higher than the LOD. HCB was <LOQ in more than 65% of the samples analyzed. o,p’-DDT, o,p’-DDE, 4,4’-DDD, 2-MeO-BDE 47 and 2’-MeO-BDE 68 were below LOQ for all individuals. One individual (harbour seal no. 7 in Table 1) displayed higher pollutant loads than the other individuals. Its ΣPCBs concentration was on average 5.5 times higher than the one of the other animals, while its ΣPBDEs level was on average 3 times higher than in the others.

No statistical difference of POP medians (Mann–Whitney U test) was observed between males and females for PCB28 (trichlorobiphenyl), ΣtetraCBs, ΣpentaCBs, ΣoctaCBs, PCB 206 and 209 (nona- and deca-CB, respectively), HCB, ΣHO-CB and PCP (p > 0.05). On the contrary, ΣhexaCBs (p = 0.030), ΣheptaCBs (p = 0.046), ΣPCBs (p = 0.046), OxC, TN, ΣDDXs, ΣPBDEs and TBA had p-values < 0.05.

No statistically significant difference was observed between the season and year of sampling, except for PCP (p < 0.01) and HCB (p < 0.05).

3.3. Haematological profiles

Median values were calculated taking the season of sampling into account as some haematological values vary with it (Table 4: Hasselmeier et al., 2008). Indeed, the Mann–Whitney U test showed that the numbers of lymphocytes, monocytes, RBC and HCT were significantly different between the two seasons considered (U < 16, p < 0.05). The median values for spring were 7900 neutrophils per blood millilitre (min. 6400–max. 10,500), 1600 lymphocytes/ml (min. 900–max. 2300) and 500 monocytes/ml (min. 300–max. 900). In autumn, the median values were 6800 neutrophils per blood ml (min. 4400–max. 9400), 3000 lymphocytes/ml (min. 2500–max. 5400) and 200 monocytes/ml (min. 100–max. 200).

This distinction between seasons was helpful to adequately evaluate the health parameters linked to haematological profile, as compared to literature data (Hasselmeier et al., 2008).

3.4. PBL in vitro cultures

The lymphocyte proliferation responses to in vitro ConA stimulation were characterized for 18 harbour seals from the same population after 72 h of incubation. The viability of cells isolated by density gradient centrifugation was always 96.7% (median 98.5%) after thawing. After 72 h of incubation, lymphocyte proliferation in presence of ConA was demonstrated by the BrdU test analysis. The flow cytometry AV-PI labelling tests showed that the majority of the lymphocytes were out of the apoptotic and necrotic pathways before and after the incubation periods.

As regards the stimulation index, no correlation between the latter and body length or weight was detected (p > 0.05). All data were pooled as there was no significant difference in stimulation indexes between genders, seasons, years and places of samplings. Despite a strong interindividual variability, lymphocytes of all individuals showed strong stimulation responses, with mean values reaching at least 3 times the one obtained for the corresponding control (Fig. 3). Optical density values were found to be 0.09 ± 0.01 (mean ± SE; median: 0.07, min. – max.: 0.01–0.24) in unstimulated control, compared to ConA-stimulated conditions which average value was 0.9 ± 0.08 (mean ± SE; median: 0.9, min. – max.: 0.3–1.5).
Table 4

<table>
<thead>
<tr>
<th>Blood cell parameters</th>
<th>Spring (n = 11)</th>
<th>Autumn (n = 7)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Median</td>
<td>Range</td>
</tr>
<tr>
<td>Lymphocytes (G/l)</td>
<td>1.6</td>
<td>0.9–2.3</td>
</tr>
<tr>
<td>Monocytes (G/l)</td>
<td>0.5</td>
<td>0.3–0.9</td>
</tr>
<tr>
<td>Neutrophils (G/l)</td>
<td>7.9</td>
<td>6.4–10.5</td>
</tr>
<tr>
<td>RBC (million/mm³)</td>
<td>5.0</td>
<td>4.6–5.3</td>
</tr>
<tr>
<td>HGB (g/dl)</td>
<td>20.9</td>
<td>17.4–24.8</td>
</tr>
<tr>
<td>HCT (%)</td>
<td>59.2</td>
<td>55.6–64.3</td>
</tr>
<tr>
<td>PLT (thousand/mm³)</td>
<td>358</td>
<td>324–437</td>
</tr>
</tbody>
</table>

* Significant difference: U < 16, and p < 0.05.
* Significant difference: U < 10, and p < 0.01.

Fig. 3. Stimulation index (SI) mean values of triplicates with coefficient of variation bars for the BrdU proliferation assay of harbour seal lymphocytes after 72 h of incubation (n = 18).

3.5. Relationships between lymphocyte parameters and contaminant levels

Analyses of in vitro lymphocyte SI suggested no statistically significant correlation with the different trace elements, organic contaminant species and sums studied (p > 0.05, e.g., correlation coefficient of SI and ΣPCBs = 0.14, p = 0.58). On the other hand, the number of lymphocytes per ml of whole blood appeared negatively correlated to PCB 209 (r = −0.51, p = 0.03), 4-HO-CB208 (r = −0.53, p = 0.02) and PCP (r = −0.63, p = 0.005).

To further investigate these relationships, a PCA based on the correlation matrix was realized with the metallic pollutant T-Hg and all sums of POP, except Pb and HCB for which at least 50% of values were <LOQ, and were not normally distributed even after log-transformation. The correlation-based PCA revealed two principal components explaining 81.0% of the total variance. The two first component loads are summarized in Table 5. The variables

<table>
<thead>
<tr>
<th>Pollutant loads for the two first components of the PCA.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Component 1</td>
</tr>
<tr>
<td>T-Hg (log)</td>
</tr>
<tr>
<td>ΣPCBs (log)</td>
</tr>
<tr>
<td>OxC (log)</td>
</tr>
<tr>
<td>TN (log)</td>
</tr>
<tr>
<td>ΣDDXs (log)</td>
</tr>
<tr>
<td>ΣPBDEs (log)</td>
</tr>
<tr>
<td>ΣHO-PCBs (log)</td>
</tr>
<tr>
<td>PCP (log)</td>
</tr>
<tr>
<td>TBA (log)</td>
</tr>
<tr>
<td>Variance explained (cumulative)</td>
</tr>
</tbody>
</table>

Fig. 4. PC1 and PC2 loading plot. The nine pollutant vectors of log-transformed concentrations represent the Bravais-Pearson correlations with the first and second principal components.

ΣPCBs, OxC, TN, ΣDDXs, ΣPBDE and ΣHO-PCBs loaded principally in PC1 (all loadings positive and at least 0.86), whereas TBA loaded mostly on PC2 (loading = 0.68). T-Hg loaded mostly on PC2 but presented a negative value. The loading plot (Fig. 4) highlighted distinctive features of Hg-T and TBA compared to the other POPs of which blood levels were correlated with one another, indicating multiple exposure.

Multiple regressions of the two principal components with the lymphocyte stimulation index and number of lymphocytes per ml of whole blood revealed a significant positive correlation only between PC2 and lymphocyte numbers (Table 6, p-value < 0.05).

A correlation test was also performed grouping the 3 dioxin-like PCB congeners (IUPAC # 105, 118, 156) and the 26 non-dioxin-like PCB congeners (IUPAC # 28, 52, 74, 87, 95, 99, 101, 110, 128, 138, 146, 149, 153, 170, 171, 172, 174, 177, 180, 183, 187, 194, 196/203, 199, 209) together. No significant correlation was found between the SI and one of those two groups (p = 0.98 and 0.58, respectively), even when considering males and females separately.

Table 5

<table>
<thead>
<tr>
<th>Pollutant</th>
<th>Component 1</th>
<th>Component 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>T-Hg (log)</td>
<td>−0.49</td>
<td>−0.71</td>
</tr>
<tr>
<td>ΣPCBs (log)</td>
<td>0.94</td>
<td>−0.13</td>
</tr>
<tr>
<td>OxC (log)</td>
<td>0.94</td>
<td>−0.04</td>
</tr>
<tr>
<td>TN (log)</td>
<td>0.97</td>
<td>0.07</td>
</tr>
<tr>
<td>ΣDDXs (log)</td>
<td>0.97</td>
<td>0.03</td>
</tr>
<tr>
<td>ΣPBDEs (log)</td>
<td>0.97</td>
<td>−0.02</td>
</tr>
<tr>
<td>ΣHO-PCBs (log)</td>
<td>0.86</td>
<td>−0.46</td>
</tr>
<tr>
<td>PCP (log)</td>
<td>0.30</td>
<td>−0.50</td>
</tr>
<tr>
<td>TBA (log)</td>
<td>0.41</td>
<td>0.68</td>
</tr>
<tr>
<td>Variance explained (cumulative)</td>
<td>64.7%</td>
<td>81.0%</td>
</tr>
</tbody>
</table>

Table 6

<table>
<thead>
<tr>
<th>Pollutant</th>
<th>Component 1</th>
<th>Component 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>SI (log)</td>
<td>0.64</td>
<td>0.53</td>
</tr>
<tr>
<td>lymphocyte number (log)</td>
<td>0.69</td>
<td>0.50</td>
</tr>
</tbody>
</table>

r-Value | p-Value
---|---
SI (log) | Component 1 | 0.64 | 0.53 |
| Component 2 | −0.84 | 0.41 |
| lymphocyte number | Component 1 | 0.69 | 0.50 |
| Component 2 | 4.43 | 0.00 |
3.6. Relationships between body weight and contaminant levels

The body weight was significantly correlated to PCB 99 representing 65% of pentacB (r = −0.48, p = 0.044), ΣhexaCB, ΣheptaCB, ΣoctaCB, PCB 206 (nonaCB) and 209 (decaCB), and ΣPCBs (r = −0.57, p = 0.015), ΣDDXs (r = −0.53, p = 0.025), BDE 100 and ΣHO-PCBs (r = 0.5, p = 0.04). Body weight was not significantly correlated to the SI (p < 0.05).

4. Discussion

During these last decades, the harbour seal population from the North Sea has been depleted by two epizootics caused by phocine distemper virus (Härkönen et al., 2006; Müller et al., 2004). Several studies have therefore investigated the health and presence of contaminants in harbour seals from this area (Reijnders, 1980, 1981; Siebert et al., 2012; Wejs et al., 2009b,c). Concomitant development of health monitoring programmes and in vitro assays circumvented the lack of experimental data about contaminant exposure and brought information on potential effects, as well as on biochemical pathways of toxic compounds (de Swart et al., 1995b,c; Ross, 2002; Ross et al., 2003a;b; Vos et al., 2003). The use of in vitro peripheral blood leucocytes offers a wide range of applications but one drawback is that these cells have already been exposed to a mixture of organic and inorganic contaminants in the blood of marine mammals. Indeed, harbour seals of the Wadden Sea displayed high blood levels of contaminants in the present study (ΣPCBs, ΣHO-PCBs, ΣDDXs and T-Hg) (Tables 2 and 3). Concentrations of organic contaminants (ΣPCBs, ΣHO-PCBs, ΣDDXs, ΣPBDs, MeO-PBDs) and trace elements (Hg, Cd, Pb, Zn, Cu, Fe) analyzed in the blood of these 18 harbour seals are from the same order of magnitude than those found in the last few years in individuals from the same seal population (Das et al., 2008; Griesel et al., 2006, 2008; Kakuschke et al., 2008; Wejs et al., 2009a). The profile of PCBs (CB 153 > CB 138 > CB 187 > CB 146 > CB 99) analyzed in the serum was similar to that described previously for harbour seals collected between 2006 and 2008 in the same geographical area (Wejs et al., 2009a), and for grey seals from the North Sea where PCB 153 and 138 were the main contributors (Vanden Berghe et al., 2012).

Blood samples provide a window into an animal’s complex network of immune function (De Guise, 2002). Haematology profiles of the 18 seals were comparable to a previous study on free-ranging and captive harbour seals from the Wadden Sea (Hasseleimer et al., 2008). However, the lymphocyte numbers in autumn were higher compared to the data given by Hasseleimer et al. (2008), but stay in the range given for spring. One individual (harbour seal number 4 – P4) presented a much higher autumn concentration with 5400 lymphocytes/ml. In all animals of this study, the erythrocyte haematology profile was generally more homogeneous than the leucocyte profile, which is more influenced by physiological changes (e.g., bacterial or viral diseases, stress) (Hasseleimer et al., 2008). The life histories of these individuals are unknown as they are living in the wild but it is known from pathological investigations that these free-ranging animals have mild infectious diseases, which explains the variability observed in blood leucocyte profiles (Siebert et al., 2007).

Lymphoproliferative response, as a specific T-cell response, can also give valuable information on the adaptive immune response. Despite the variability related to samples obtained from free-ranging harbour seals, a clear stimulation of the T lymphocytes associated with a significant proliferation could be observed (Fig. 3).

The in vitro stimulation index (SI), currently used in cell culture, showed significant interindividual variation. Nevertheless, high inter- and intra-individual variations of lymphocyte proliferation also occur in humans in which the whole life history is generally known, and in which homogeneous groups were tested (e.g., Stone et al., 2009). Our results did not highlight any clear correlation between the stimulation indices and the pollutants or class of pollutants studied (component 1, Table 5). In contrast, a previous study showed that in vitro Con A-induced peripheral blood lymphocyte proliferation was inversely correlated to whole blood levels of tetra- to octa-CBs, p,p′-DDT, o,p′-DDE and p,p′-DDE in a small group (n = 5) of bottlenose dolphins (Tursiops truncatus) (Lahvis et al., 1995). However, the very small sample size and large age range of those animals (3–32 years old, approximate age) precluded drawing extensive conclusions. Studies on harbour seal pups highlighted modulating effects of PCBs from the blubber (Levin et al., 2005; Mos et al., 2006). Mos et al. (2006) showed that the proliferative responses of T lymphocytes, but not of B lymphocytes (mimicked by stimulation with the polyclonal activators ConA and lipopolysaccharide (LPS), respectively), were significantly decreased in harbour seal pups presenting higher PCB concentrations in their blubber. On the other hand, Levin et al. (2005) reported a significant body weight–independent positive correlation between both T-cell mitogen (phytohemagglutinin [PHA])- and B-cell mitogen (LPS)-induced lymphocyte proliferation and the blubber concentrations of Σ11 PCBs, but no significant correlations were found between ConA-induced lymphocyte proliferation and total PCB concentration (Levin et al., 2005). However, the correlations observed between lymphocyte proliferation indices and blubber or blood PCB levels are different as these two tissues differ in OC affinities and retention capacities. This is because their lipid composition is different and because contaminants have different lipophilic properties (Addison and Brodie, 1987; Sinjari et al., 1996), resulting in a selective retention of the more lipophilic compounds in the blubber and the more polar compounds in the blood (Lydersen et al., 2002).

In the framework of in vitro exposure to contaminants, Neale et al. (2002) showed no significant depression in harbour seal lymphoproliferation when exposed to 10 µM CB-156 and CB-80. Other studies of in vitro exposure to some OC mixtures (PCBs and TCDD) have revealed significant positive correlations with the ConA-induced T cell proliferation in the harbour seal, Northern fur seal, Steller sea lion, beluga whale, Commerson’s dolphin and sea otter (Levin et al., 2007; Mori et al., 2006). On the contrary, significant decreases in lymphocyte proliferation were seen after mercury exposure in harbour seals (Das et al., 2008; Dufesne et al., 2010) and after OC mixture exposure in mice (Mori et al., 2006), for example. The in vitro exposure of harbour seal granulocytes to 12 µM of BDE-47, -99 and -153 had also been shown to induce a thiol depletion and phagocytosis inhibition (Fraun et al., 2010).

In the present study, the number of activated lymphocytes appeared significantly negatively correlated to only a few blood pollutants (PCB 209, 4-OCB-208 and PCP) but this cannot be clearly discussed for PCB 209 and 4-OCB-208 as they are minor compounds. In contrast, PCP, a widely used pesticide ubiquitous in the environment, had been demonstrated to have suppressive effects on human lymphocytes in vitro (Lang and Mueller-Ruchholtz, 1991) as well as on diverse cellular and humoral immune parameters (Daniel et al., 1995, 2001). Serum levels are not available for comparison in marine mammals, but humans usually present much lower concentrations (Butte and Heinzow, 1995). PCP should thus be monitored in marine mammals in order to find out more about its biological effects.

The fact that no clear correlation could be found between the in vitro SI and the contaminants present in blood of free-ranging adult harbour seals at that time of their annual cycle does not mean that chemicals have no toxic effect on PBls. In all likelihood, there are more complex interactions between confounding factors explaining immunological variations observed in vitro. In any case, the
results obtained in in vitro assays should be interpreted cautiously as PBls are isolated from environmentally contaminated harbour seals. Generally speaking, factors such as precise age, nutritional status, genetic factors, stress, disease, nursing, or fasting, may have additional influences on immune functions, thereby limiting the direct ability to detect the effects of contaminants on the immune system (Levin et al., 2005). Those factors should thus be taken into consideration, within the realms of possibility. In this study, all individuals were adults in a good body condition, presenting no sign of disease. Yet minor injuries or infections and stress may have influenced the lymphocyte responses.

Furthermore, contaminants were measured in serum (organic pollutants) or whole blood (metals) and not in blubber or in liver, both tissues known to accumulate these contaminants over a lifetime (Wejs et al., 2011) in contrast to blood (Habran et al., 2013). However, correlations could be made between levels of organochlorines found in blood and blubber (Lydersen et al., 2002). Phocid seals regularly go through fasting or reduced food intake periods and most of their mass loss during these periods is derived from catabolism of lipids from their inner blubber layer (Nordoy and Blix, 1985). In harbour seals, the mobilization of lipids from the blubber notably occurs in individuals having lower food intake during the breeding and moulting seasons (Burns, 2008), in pups during the postweaning fast (Muehlebert and Bowen, 1993) and in females producing milk (Boness et al., 1994). Lactation is indeed particularly demanding, leading to important loss of body fat and energy (Bowen et al., 1992). Blood samples from a seal which is in a state of positive energy balance would thus have POP levels reflecting those in the diet, while blood levels during fasting and lactating periods are reflecting inner blubber levels of bioaccumulated lipophilic contaminants. Moreover, seals in a prime condition generally have lower concentrations than animals in a poor condition (Lydersen et al., 2002). Harbour seals of this study were sampled while in a good condition before the breeding season and after the moulting season, and not while in a poor condition as during moulting for example. Their blood levels could thus reflect low contaminant concentrations compared to those of fasting and milk production periods.

Results obtained in our study highlighted the fact that variations in proliferation could not be directly linked to the blood pollutants investigated in adult harbour seals in a good body condition, but that other factors could most likely have an influence. Field studies do thus allow the opportunity to study the direct and indirect effects of real-world exposures to POPs on the immune system when confounding factors are minimized (Levin et al., 2005).

5. Conclusions

We provide here a relevant approach to get a better understanding of the relationships between the immune function and several organic and inorganic contaminants present in the blood of free-ranging adult harbour seals from the North Sea. This is the first study reporting the simultaneous measurement of blood haematology parameters, trace elements, PCBs, PBDEs, naturally produced MeO-PBDEs, DDXs, OXc, TN, HCB and the metabolites HO-PCBs, in blood of healthy harbour seals, concomitantly to the study of lymphocyte proliferation in vitro. Significant lymphocyte proliferation was observed and results highlighted a clear correlation between the number of lymphocytes per blood millilitre and PCP, but not between the lymphocyte stimulation indices and blood contaminant levels. This corroborates the fact that peripheral blood lymphocytes isolated from wild harbour seals and cultured in controlled conditions constitute a valuable tool for in vitro studies, as long as confounding factors are limited. Nevertheless, a study carried out on a bigger sample size would be necessary to confirm those results.

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