RESEARCH ARTICLE

Spatial variability of metallic and organic contamination of anguilliform fish in New Caledonia

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Abstract New Caledonia is one of the main hot spots of biodiversity on the planet. Large amounts of contaminants are discharged into the lagoon as a result of increasing anthropogenic activities such as intense mining, urbanization, and industrialization. Concentrations of 14 trace elements and 26 persistent organic pollutants (POPs: PCBs and pesticides) were measured in the muscles of two anguilliform fish species, over a coast to barrier reef gradient in two lagoon areas differently exposed to anthropic disturbances. This study emphasizes the high trace element contamination status of anguilliform fish and also highlights slight but perceptible organic pollution. The contamination extends throughout the lagoon, from coast to barrier reef, even in areas remote from emission points. High levels of trace elements, especially those linked to mining activities (i.e., Co, Cr, Fe, Mn, and

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E. Wafo Laboratoire de Chimie Analytique, IMBE UMR 7263 CNRS, 237 IRD/UMR S476 UMR A1260, AMU, Marseille, France Ni), were detected in coastal sites. Furthermore, the large dispersion of most POPs throughout the entire lagoon poses the question of their potential toxicity on marine organisms from numerous habitats. Our results underline the need for long-term monitoring of various contaminants over large spatial and time scales.

Keywords Trace elements · PCBs · Pesticides · POPs · Carnivorous fish · Coral reefs · SW Pacific Ocean

Introduction

Coral reefs are threatened by a wide range of natural and anthropic disturbances, including chemical pollution (Richmond 1993). Three main contaminant classes are particularly involved: polychlorinated biphenyls (PCBs), pesticides (both classes are part of persistent organic pollutants or POPs), and metallic trace elements. These ubiquitous contaminants can be toxic, even at low concentrations; they are resistant to degradation, transported over long distances, and can bioaccumulate in marine organisms (e.g., Phillips 1995; Robertson and Hansen 2001). The half-life of contaminants in the marine environment is estimated to be at least a decade for the most persistent PCBs (Sinkkonen and Paasivirta 2000; Robertson and Hansen 2001), several years to a decade for trace elements such as Hg (Lodenius 1991) and several months to years for pesticides (Hellawell 1988). After their release into the marine environment, some contaminants can be strongly accumulated by organisms of different trophic levels (Van Ael et al. 2012; Dummee et al. 2012).

Southwest Pacific coral reefs are generally in good health. However, for a few decades, toxic wastes released by human activities threaten some of them (Richmond 1993). New Caledonia's lagoon, the largest in the world, is no exception. This hot spot of biodiversity is subjected to increasing



contamination pressure resulting from industrialization and urbanization. New Caledonia is the third producer of nickel ore in the world (Dalvi et al. 2004). Since the end of the nineteenth century, many open-cast mines and three metallurgical factories have been opened. Mining activities and natural soil erosion due to rainfall (Ambastian et al. 1997) generate massive sediment deposits and discharge of associated trace elements (Co, Cr, Fe, Mn, and Ni,) into the lagoon. Furthermore, with the expansion of industrial factories and urban development, the lack of efficient treatment of wastewater, and the use of pesticides for agriculture, the contamination by POPs is also an important issue. Nearly 250 pesticides are officially authorized in New Caledonia (http://davar.gouv.nc/). Twenty-six of these are banned but in Europe are still extensively used in New Caledonia.

Several studies focused on trace metal contaminants in coastal waters, in particular around Nouméa (the main urban region) (Fernandez et al. 2006; Metian et al. 2008a, b; Hédouin et al. 2008, 2009, 2011). Bioaccumulation of mining (Co, Cr, Fe, Mn, and Ni) and urban trace elements (Ag, As, Cd, Cu, Hg, Pb, Se, V, and Zn) have been investigated in various taxa (crustaceans, molluscs, ascidians) (Bustamante et al. 2000; Hédouin et al. 2006, 2007, 2009, 2010, 2011; Pernice et al. 2009; Metian et al. 2008a, 2010). One study recently revealed high contamination levels by trace elements in fish from the south-western lagoon, extending from the coast to the barrier reef (Bonnet et al. 2014). However, there is still little information available about trace element contamination of fish across the whole lagoon in New Caledonia (Chouvelon et al. 2009; Fernandez and Breau 2011; Metian et al. 2013). Furthermore, to our knowledge, there is no information available to date on PCB and pesticide concentrations in marine organisms in New Caledonia.

Analyses of spatial patterns of contamination allow the identification of their environmental sources. They also lead to a better understanding of the physical, chemical, and biological processes involved in contaminant accumulation (Robertson and Hansen 2001; Johnson et al. 2005). However, with the exception of the coastal region (especially in Nouméa), contamination by trace elements is insufficiently documented and its impact on marine ecosystems remains unclear. Considering the immense surface of the south-western lagoon of New Caledonia, it is necessary to extend investigations to a large spatial scale and to other contaminants such as POPs.

Marine top predators, mainly long-living species, bioaccumulate high levels of metallic elements (e.g., Cd and Hg) or organochlorine pollutants in their tissues (e.g., Adams and McMichael 1999; Wafo et al. 2012). Fish are particularly useful to assess contamination status and distribution of pollutants, especially if they are site-attached or even territorial, living in benthic habitats and have a high trophic level (carnivorous) (e.g., Kojadinovic et al. 2007; Dierking et al. 2009). The sedentary nature of some fish is a crucial aspect in

spatial variability studies of contamination. Anguilliform fish (morays, congers, and snake eels), which are ubiquitous organisms highly represented in tropical marine waters, fulfill the conditions of good bioindicators in contamination monitoring (Bonnet et al. 2014). These sedentary benthic predators, which mainly live in coral reef matrices of shallow waters, are good candidates for studying the impact and spatial variability of contamination in marine environments. Surprisingly, to date, they have received little interest, which could be due to their cryptic lifestyle and the difficulty to collect them (Ineich et al. 2007).

In this general context, the main purposes of this study were (1) to determine the level of contamination of two anguilliform fish species by trace elements and POPs, (2) to study the spatial variability of these contaminants along two coast-barrier reef gradients which are differently exposed to human activities. As the influence of ecological and biological factors on the concentration of contaminants is important, three biological parameters (body size, trophic position, and lipid content in tissue) and their link to the bioaccumulation processes were also investigated.

Materials and methods

Fish sampling and study areas

Two anguilliform species were studied, Gymnothorax chilospilus (Muraenidae) and Conger sp. (Congridae). Capture of specimens was carried out using sea kraits (Laticauda spp.) because the prey of these predators is almost exclusively anguilliform fish. This method has been successfully used in previous studies (e.g., Reed et al. 2002; Ineich et al. 2007; Brischoux et al. 2011). The prey items were obtained by a gentle massage of the sea krait abdomen. Because the eels consumed are non-spiny, the snake readily regurgitates their prey (Brischoux and Bonnet 2009) without any effect on their survival (Fauvel et al. 2012). The spatial accuracy of sea snake sampling and their use as sentinels was described in previous studies (Brischoux et al. 2007a, 2009; Brischoux and Bonnet 2008; Bonnet 2012). Sea kraits probe the surrounding seafloors within a radius of 10-15 km, and anguilliform fish are extremely sedentary. Considering the large spatial scale of the current study, this accuracy was sufficient to examine possible local influences along a coastbarrier reef gradient.

The study was realized in the South Western lagoon of New Caledonia. Both fish species were caught in two main areas: Grand Nouméa (GN) and Grand Lagon Sud (GS) (Fig. 1). Grand Nouméa, which is close to the main city of Nouméa, is subject to a variety of anthropic pollution such as industrial activities, farming industries, and wastewater. In particular, a nickel factory has functioned since 1880 (SLN,



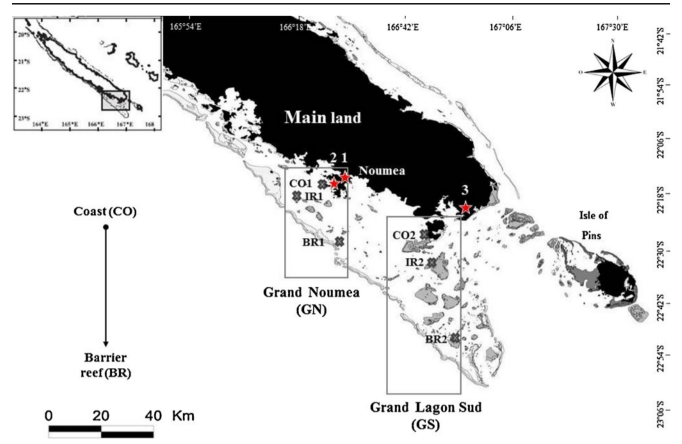


Fig. 1 Location of the sampled sites distributed on two coast-barrier reef gradients in New Caledonia: Kuendu (CO1), Signal (IR1), and Amédée (BR1) in Grand Nouméa (GN) and Ouen (CO2), Grand Mato (IR2) and N'Da (BR2) in Grand Lagon Sud. CO=coastal sites; IR=intermediate reef sites; BR=barrier reef sites. Stars indicate areas of human activities,

which are the industrial area of Ducos (I), the Nickel factory « SLN » (2), and the Nickel factory « Goro-Nickel » (3). The emerged land is indicated in black; grey areas represent coral reefs; the dark grey line represents the slope of the barrier reef

Société Le Nickel, Fig. 1). Grand Lagon Sud is less influenced by industrial and urban pollution. Nevertheless, since 1950, mining activity has developed near the Bay of Prony (Goro-Nickel, Vale Inco, Fig. 1), increasing the erosion of lateritic soil in this region. For each area, three stations located on a coast-barrier reef gradient inside the lagoon were sampled: Kuendu beach (coast; CO1), Signal islet (intermediate-reef; IR1), and Amédée islet (barrier-reef; BR1) inside GN and Ouen island (coast; CO2), Mato islet (intermediate-reef; IR2), and N'da islet (barrier-reef; BR2) inside GS (Fig. 1). Sampling was carried out from January to April 2011 and from August to September 2011, corresponding respectively to the hot and wet versus cool and dry seasons in New Caledonia.

Each fish was identified, measured, and weighed prior to dissection (Table 1). As some fish collected were partially digested, their total length (TL) was estimated applying allometric equations using either the snout vent length or the tail length (Brischoux et al. 2007b). For each individual of both species, a piece of white muscle was sampled on the non-digested part and immediately frozen at -30 °C for subsequent

analyses. Muscle tissues were freeze-dried and ground to powder with a porcelain mortar and pestle.

Trace element analyses

The total Hg concentration in the powder obtained from the tissues was determined by analyzing the Hg, with an advanced mercury analyzer (ALTEC AMA 254), directly on aliquots ranging from 5 to 50 mg of dry sample weighed to the nearest 0.01 mg (Bustamante et al. 2006). From 150 to 300 mg of each sample were digested using a 3:1 ν/ν nitric–hydrochloric acid mixture with 65 % ultrapure HNO₃ and ultrapure 37 % HCl. The acidic digestion was performed overnight under ambient temperature and then heated in a microwave for 30 min, increasing the temperature up to 105 °C, and 15 min at 105 °C (1200 W). After the mineralization process, each sample was diluted to 30 or 50 ml with milli-Q quality water, according to the volume of acid added to the mineralization (3 and 4.5 ml, respectively).

The analysis of Ag, As, Cd, Co, Cr, Cu, Fe, Mn, Ni, Pb, Se, V, and Zn required an extra step in the preparation protocol. These elements were analyzed by inductively coupled plasma



Table 1 Number (N) of fish analyzed from the coastal (CO), intermediate (IR), and barrier reef (BR) sites in Grand Nouméa (GN) and Grand Lagon Sud (GS)

Species	Site	N	N (GN)	N (GS)	TL (cm) (min-max)	$\delta^{15}N$ (‰) (min–max)	Lp (%) (min-max)
G. chilospilus	СО	8	3	5	21.2±4.3 (14.7–28.4)	8.0±0.8 (7.1–9.3)	1.2±1.3 (0.2–3.6)
	IR	20	10	10	25.3±4.8 (14.0-37.2)	$7.9\pm0.7~(6.7-9.5)$	0.8±0.8 (0.2-3.3)
	BR	20	10	10	25.1±2.3 (21.4-30.8)	$7.8\pm0.5\ (7.1-8.8)$	0.7±0.6 (0.2-1.8)
	Total	48	23	25	24.5±4.1 (21.7-31.2)	$7.9\pm0.6~(6.7-9.5)$	0.8±0.8 (0.2-3.6)
Conger sp.	CO	14	9	5	29.0±4.6 (23.2-38.3)	8.6±1.4 (6.6–10.5)	1.0±0.7 (0.2-1.9)
	IR	10	0	10	34.4±8.9 (20.3-48.7)	8.5±0.8 (7.7-9.9)	1.5±1.6 (0.3-5.5)
	BR	13	8	5	35.7±6.3 (27.7-51.7)	8.5±0.9 (7.1-10.3)	1.0±1.2 (0.3-3.6)
	Total	37	17	20	32.7±7.1 (20.3–51.7)	$8.6 \pm 1.0 \ (6.6 – 10.5)$	1.2±1.2 (0.2–5.5)

Mean (\pm SD) fish total length (TL, centimeters), trophic position (express with δ^{15} N, %), and lipid content (Lp, %) are given, with minimal and maximal values (min–max) indicated in brackets. No variation of TL, δ^{15} N, and Lp between sites was observed ($p_{\rm ANOVA}$ >0.05)

atomic emission spectrometry (Varian Vista-Pro ICP-OES) and mass spectrometry (ICP-MS II Series Thermo Fisher Scientific). Reference tissues—dogfish liver DOLT-4 (NRCC), lobster hepatopancreas TORT-2 (NRCC)—were treated and analyzed in the same way as the samples. Results were in line with the certified values, and the standard deviations were low, proving good repeatability of the method. The results for standard reference materials displayed recovery of the elements ranging from 73 % to 116 %. For each set of analyses, blanks were included in each analytical batch. The detection limits (micrograms per gram dry weight) were 0.005 (Hg), 0.015 (Cd), 0.017 (Ag), 0.02 (Cr, Co, Pb), 0.03 (Ni), 0.08 (Mn), 0.1 (Cu, Se), 0.2 (As), 0.3 (V), and 3.3 (Fe and Zn). All trace element concentrations are given on a dry weight basis (micrograms per gram dry weight).

POP analyses

Choice of PCB congeners and pesticides

The concentrations of 15 individual congeners (IUPAC Nos. 18, 20, 28, 31, 44, 52, 101, 105, 118, 138, 149, 153, 170, 180, 194; Ballschmiter and Zell 1980) were determined in each sample. This list contains the seven target congeners (PCBs 28, 52, 101, 118, 138, 153, 180) proposed by the International Council for the Exploration of the Sea (ICES) as indicators of PCB contamination (Valoppi et al. 2000).

The samples and blank were analyzed for the following pesticides: aldrin, diazinon, dieldrin, endosulfan I, endosulfan II, endrin, heptachlor, heptachlor-epoxide A, heptachlor-epoxide B, lindane, and, pp'-DDD, pp'-DDE, pp'-DDT.

Sample extraction and quantification

Compounds were extracted and quantified following the procedures described by Dierking et al. (2009). About 1 g of lyophilized sample was extracted and concentrated to 2 ml.

About 100 μL of this extract was reserved to determine the content of lipid (Lp, %) of each sample (see next paragraph). The remaining fraction was purified with concentrated sulfuric acid, followed by additional purification by liquid chromatography on a silica-alumina column (Murphy 1972; Wells et al. 1985). Four fractions were eluted: fraction Ia (aldrin, pp'-DDT, heptachlor (50 %)), fraction Ib (heptachlor (50 %), pp'-DDT), fraction II (endosulfan I, endosulfan II, lindane, pp'-DDD) and fraction III (dieldrin, endrin, heptachlor-epox A and B).

PCBs and pesticides were analyzed by gas chromatography coupled to mass spectrometry. Calibration was done using fifteen individual standard solutions for PCBs and a standard mixture containing all pesticides was used for pesticides. All organic pollutant concentrations are given on a dry weight basis (nanograms per gram dry weight). Detection limits were 0.01 ng g⁻¹ for PBC congeners, 0.1 ng g⁻¹ for DDT, and its metabolites, 0.01 ng.g⁻¹ for heptachlor-epoxide A and B, 0.1 ng.g⁻¹ for lindane, diazinon, heptachlor, aldrin, endosulfan II, endrin, and 0.2 ng g⁻¹ for endosulfan I and dieldrin.

δ^{15} N and lipid quantification

The nitrogen stable-isotope ratio (δ^{15} N) was used to give an approximate trophic position. Analyses were carried out on dorsal white muscle, as this tissue gives the most reliable values (Pinnegar and Polunin 1999). Then 1 ± 0.1 mg of powdered freeze-dried samples was weighed out and sealed in tin capsules for analyses. 15 N/ 14 N ratios were determined by continuous-flow isotope-ratio mass spectrometry with a Thermo Scientific Delta V Advantage mass spectrometer coupled to a Thermo Scientific Flash EA1112 elemental analyzer (Chouvelon et al. 2011). Results are expressed as isotope ratios δ^{15} N (‰) relative to international standard (atmospheric N₂ for nitrogen), according to the formula:

$$\delta^{15}N = \left[\left(R_{\text{sample}} / R_{\text{standard}} \right) - 1 \right]^* 10^3$$



where $R = {}^{15}N/{}^{14}N$ (Peterson and Fry 1987). Replicate measurements of internal laboratory standards (acetanilide) indicated a precision of approximately 0.2‰ for δ^{15} N values.

To evaluate the lipid content (Lp, %) in each fish, $100~\mu L$ of the evaporated extraction residue was introduced in an initially tarred (tablet) container; then the whole was deposited into a desiccator. After drying to constant mass, the lipid content is determined by gravimetry.

Statistical analysis

Several indices were used to characterize the POP contamination. Our PCB profiles revealed that PCBs detected in the environment mainly came from capacitors (pyralene: DP3, DP4 or Arochlor1230, 1242). Consequently, it was decided to express the global footprint of PCB contamination (PCBTot) using the estimation proposed by Annema et al. (1995): $PCBTot = \Sigma$ (28+52+101+138+ 153+180)*5. $\Sigma ICES$ (sum of concentrations of the seven ICES congeners), and the proportional contribution of each PCB class (tri- to octa-chlorinated) was also calculated using all of the PCBs analyzed. The total DDT concentration, ΣDDT , was calculated as the sum of pp'-DDD, pp'-DDE, and pp'-DDT. An indication of the timing of PCB inputs was estimated through the ratio of the two congeners CB153/CB138 (Monod et al. 1995; Wafo et al. 2005: A high ratio (>1) indicates continuous inputs or persistence of PCBs in the environment, whereas a low ratio (<1) reflects a decrease or sporadicity of PCB inputs or nonpersistence. Finally, the ratio of the total concentration of pesticides to total concentration of PCBs analyzed ($\Sigma Pest$ / ΣPCB) was used as an approximate indication of the relative importance of agricultural versus industrial sources of organic pollutants in the environment (de Mora et al. 2004); a ratio higher than 1 indicates prevalence of agricultural sources, whereas a ratio less than 1 indicates prevalence of industrial sources.

Data were log-transformed to satisfy conditions of normality (Shapiro-Wilk test) and homogeneity of variance (Bartlett test). However, for some compounds (aldrin, diazinon, dieldrin, endrin, pp'-DDD, and pp'-DDT), transformation was not sufficient. When conditions were satisfied, parametric tests were used. In other cases, non-parametric analogues were applied.

Principal component analysis (PCA) was used to obtain an overview of the contribution of each contaminant according to sites. Trace elements and POP concentrations were analyzed in separate PCAs, and as no clear patterns were revealed from POP analysis, only trace element results were presented. Spatial variability of contaminant concentrations in each species was also tested using two-way ANOVA and Tukey's post hoc multiple comparisons. For all contaminants, Pearson or

Spearman correlations were used to explore the relationship between both contaminant concentration and fish size (TL) and contaminant concentration and trophic level (δ^{15} N). For organic pollutants, the relationship between contaminant concentration and the percentage of lipids in muscle (Lp) was also examined. When a correlation existed, ANCOVA, instead of ANOVA was applied, using TL, δ^{15} N or Lp as a covariate. R project (R version 2.13.2) was the statistical program used for all analyses. The levels of significance for statistical analyses was always set at α =0.05.

Results

Accumulation levels in anguilliform fish

All trace elements measured were detected in anguilliform fish, except Ag and V, which were below the detection limit. Mean concentrations and ranges of detected values of contaminants (trace elements and POPs) in anguilliform fish muscle are presented in Table 2. Several elements showed a high inter-individual variability (i.e., Cd, Cr, Ni, Pb; coefficient of variation CV>100 %). However, it was most often the same individuals that accumulated high concentrations of several trace elements linked to their origin (e.g., a *G. chilospilus* specimen from the coastal site of Grand Nouméa; 35.7 μ g g⁻¹ dw of Cr and 17.4 μ g g⁻¹ dw of Ni).

Overall, few differences in mean contaminant concentrations were found between species. However, significantly higher concentrations of As, Hg, and Se were measured in *Conger* sp., whereas *G. chilospilus* exhibited higher concentrations of Cd and Zn ($p_{\rm ANOVA}{<}0.001$). PCBs had been similarly accumulated by both species, albeit weak interspecific differences were detected for pesticides: *Conger* sp. presented significantly higher concentrations of heptachlor, heptachlor-epoxy B ($p_{\rm ANOVA}{<}0.01$), while slightly higher concentrations of endosulfan I and aldrin were measured in *G. chilospilus* ($p_{\rm ANOVA}{<}0.05$).

Spatial patterns in contaminants

Trace elements

A significant spatial variability of several trace elements, in particular, those linked with mining activity (Co, Cr, Fe, Mn, and Ni), was detected along the coast-barrier reef gradient (Fig. 2 and Table 3). The PCAs in *G. chilospilus* indicated that the coastal sites were characterized by Co, Mn, and Ni elements, in contrast to the barrier reef sites which were characterized by As and Cd concentrations (the plot of the first two PCA dimensions explained respectively 51.1 % and 59.0 % of data variability in *G. chilospilus* and *Conger* sp.,



Table 2 Trace element and organic pollutant concentrations (mean±SD and min-max values; micrograms per gram dry weight) measured in muscle of *G. chilospilus* and *Conger* sp., all sites confounded

		G. chilospilus	Conger sp.
Trace elements	As	11±7.4 (0.9–31.9)	25.7±25.6 (3.0-117.9)
	Cd	$0.41\pm0.47~(0.01-2.91)$	0.04±0.04 (0.01-0.16)
	Co	$0.11\pm0.08~(0.020.45)$	0.13±0.15 (0.02-0.68)
	Cr	4.53±7.15 (0.86–35.71)	4.03±5.36 (0.45-23.60)
	Cu	$1.6\pm0.7~(0.7-3.4)$	1.9±0.9 (0.7-4.5)
	Fe	40.2±36.9 (12.2-196.8)	34.3±31.1 (7.9–134.4)
	Hg	$0.049\pm0.028~(0.02-0.13)$	0.117±0.112 (0.03-0.6)
	Mn	2.3±1.4 (0.6-7.2)	2.7±2.3 (0.4-10.1)
	Ni	1.95±3.19 (0.30-17.38)	1.44±1.77 (0.10-7.74)
	Pb	$0.05\pm0.04~(0.010.2)$	0.04±0.09 (12.2-196.8)
	Se	1.2±0.4 (0.7-2.6)	1.8±0.8 (0.9-4.6)
	Zn	56.5±16.4 (26.5-112.3)	45.4±23.8 (21.6-112.1)
Organic pollutants	PCBTot	59.1±52.7 (9.3–271.5)	63.2±54.8 (9.1–211.4)
	$\Sigma ICES$	14.0±12.2 (2.0-64.3)	16.5±14.3 (2.1-60.9)
	Aldrin	$0.1\pm0.06~(0.08-0.3)$	0.1 ± 0.05 (0.04-0.2)
	Diazinon	1.8±1.9 (<0.1–10.2)	2.4±3.3 (<0.1-15.3)
	Dieldrin	0.4±0.3 (<0.2-1.8)	0.5±0.4 (<0.2-2.2)
	Endosulfan I	1.3±1.3 (<0.2-6.1)	1.0±1.7 (0.09-9.7)
	Endosulfan II	2.1±1.7 (<0.1-7.3)	2.9±2.3 (<0.1-8.9)
	Endrin	0.3±0.8 (<0.1-4.5)	$0.8\pm2.0~(0.07-9.9)$
	Heptachlor	0.4±0.4 (<0.1-1.9)	0.5±0.4 (<0.1-1.9)
	Heptachlor-epox. A	1.5±1.2 (0.03-5.6)	1.8±1.9 (0.1–9.1)
	Heptachlor-epox. B	0.3±0.5 (<0.01-2.6)	0.5±0.6 (<0.01-2.7)
	Lindane	1.3±0.6 (0.3–3.1)	1.7±1.2 (0.2–5.1)
	ΣDDT	1.6±1.5 (0.3–6.9)	1.6±1.5 (0.3–7.2)

Fig. 2). In *Conger* sp., the difference between sites was even stronger, with most trace elements (As, Co, Cr, Fe, Ni, Se, and Zn) characterizing coastal sites. These general trends were confirmed by the significantly higher concentrations of mine elements at coastal sites for both species ($p_{\rm ANOVA}$ or $p_{\rm ANCOVA}$ < 0.05, Table 3).

The same spatial pattern (coast vs. barrier reef) was revealed in the two studied areas (GN and GS, Table 3 and S1). The coastal site of Grand Nouméa showed significantly higher concentrations of Ni in G. chilospilus (p_{ANOVA} =0.02) and Cr and Fe in Conger sp. (p_{ANOVA} <0.05, Table 3 and S1 of the Electronic supplementary material), with the same trends followed by several other trace elements (Cr, Fe in G. chilospilus, As, Co, Cu, Ni, Se, and Zn in Conger sp., results of PCA not shown). However, in Grand Lagon Sud, the spatial gradient was less pronounced. The coastal site was characterized by higher Co and Mn concentrations in both species (results of PCA not shown), but spatial analyses only suggested significantly lower concentrations of As and Cd in G. chilospilus (p_{ANCOVA} =0.001, Tables 3 and S1of the Electronic supplementary material).

The coastal site of Grand Nouméa was characterized by significantly higher levels of several trace elements compared with Grand Lagon Sud (e.g., As, Cd, and Ni in *G. chilospilus*, and Cr, Fe in *Conger* sp.; $p_{\text{ANCOVA}} < 0.05$, Tables 3 and S1of the Electronic supplementary material).

Organic pollutants

The PCAs did not reveal any clear spatial pattern for POPs in either species (results not shown). Analyses suggested a low spatial variability along the coast-barrier reef gradient (Table 4).

The PCB contamination in *G. chilospilus* did not vary significantly between sites (see *PCBTot* index; $p_{ANOVA} > 0.05$), in spite of a slight increase from the coast to the barrier reef. However, in *Conger* sp., the intermediate sites were characterized by high concentrations ($p_{ANCOVA} < 0.05$, Table 4). The *CB*153/*CB*138 ratio, which was generally below 1, showed few spatial variations and only *G. chilospilus* from coastal sites showed a significantly higher ratio ($p_{ANCOVA} < 0.05$, Tables 4 and S2 of the Electronic supplementary material). The PCB chlorination classes differed slightly along the coast-barrier reef gradient and between studied areas, with a low variability detected in each case (<10 %, p_{ANCOVA} or $p_{ANOVA} < 0.05$, Table 4). However, the



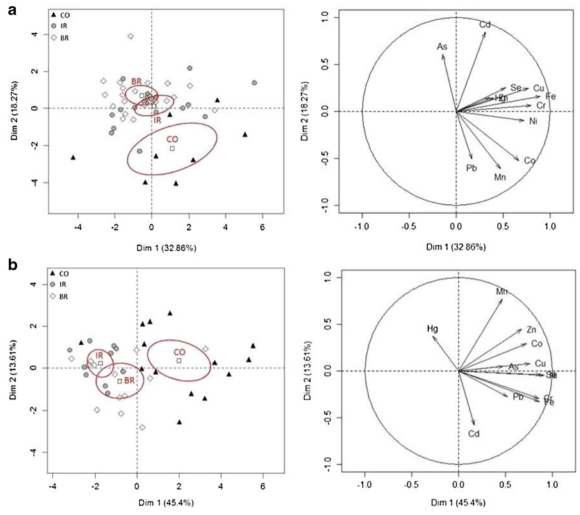


Fig. 2 Plot of the principal component analysis (PCA) assessing spatial patterns of anguilliform fish contamination by trace elements on the coast-barrier reef gradient (*left panel*) and the contribution of the contaminants analyzed to site differentiation (*right panel*) in *G. chilospilus* (**a**) and *Conger* sp. (**b**). *CO*=coastal sites; *IR*=intermediate reef sites;

BR=barrier reef sites. The most contributing variables in both species were Cr, Cu, Fe, and Ni concentrations for the first axis and Cd and Mn concentrations for the second axis. The significant differences between sites are illustrated by confidence ellipses of centroids at 95 %

results highlighted a remarkable increase in the proportion of the most toxic chlorinated classes in coastal sites (i.e., 8Cl in GN for *G. chilospilus* and 7Cl in GS for *Conger* sp., $p_{\rm ANOVA}{<}0.05$).

The results for pesticides indicated few spatial variations (Tables 4 and S2 of the Electronic supplementary material). The coastal sites highlighted significantly higher concentrations of some pesticides in *G. chilospilus* (aldrin and heptachlor, $p_{\rm ANCOVA}$ or $p_{\rm KW}{<}0.05$), whereas the intermediate reef sites revealed significantly higher rates of several other pesticides in *Conger* sp. (e.g., diazinon, heptachlor epoxide A and pp'-DDT; $p_{\rm ANCOVA}$ or $p_{\rm KW}{<}0.05$).

Finally, spatial analyses of POP contamination according to sources of pollution (agriculture vs. industrial) reinforce previous results (see $\Sigma Pest/\Sigma PCBs$; Table 4). Along the gradient, *G. chilospilus* from the coastal sites were significantly different from other sites, with quite homogeneous sources of contamination (ratio ≥ 1). In comparison industrial

sources dominated in other sites (ratio \cong 0.50, $p_{\rm ANOVA}$ <0.05, Table 4). A strong influence of industrial pollution in intermediate sites (ratio \cong 0.50, $p_{\rm ANCOVA}$ <0.05) was also highlighted in *Conger* sp.

Influence of size, trophic position, and lipid content on contamination level

Several trace elements (Co, Cr, Ni, Pb, Se, and Zn in *G. chilospilus*, Cd and Mn in *Conger* sp.) were not correlated with either the size or the trophic position of fish ($p_{Pearson} > 0.05$), but some trace elements were significantly correlated with fish size (Table 5). Only As and Hg highlighted a positive relationship with the trophic position of fish (δ^{15} N) in *Conger* sp., whereas no correlation was found for *G. chilospilus*.

Organic pollutant concentrations were linked to size, trophic position, or lipid content, depending on the contaminant.



Table 3 Trace element concentrations (mean (SD); micrograms per gram dry weight) in *G. chilospilus* and *Conger* sp. along the coastal-barrier reef gradient

		Gradient					Spatial pattern	ns		
		СО	IR	BR	Stat test	p	Gradient	GN	GS	GN vs. GS
G. chilospilus	As	6.77 (10.27)	7.84 (5.75)	14.33 (5.85)	ACV (δ ¹⁵ N)	***	BR>CO/IR	BR1>IR1	IR2/BR2>CO2	CO1>CO2
	Cd	0.16 (0.23)	0.44 (0.63)	0.43 (0.31)	ACV (TĹ)	***	IR/BR>CO		IR2/BR2>CO2	CO1>CO2
	Co	0.19 (0.12)	0.10 (0.05)	0.07 (0.05)	ANV	***	CO>IR>BR			
	Cr	6.59 (11.93)	3.57 (5.98)	3.97 (5.81)	ANV	NS				
	Cu	1.62 (0.76)	1.66 (0.78)	1.37 (0.40)	ACV (TL)	NS				
	Fe	46.5 (50.5)	38.2 (39.9)	33.8 (27.4)	ACV (TL)	NS				
	Hg	0.05 (0.02)	0.05 (0.03)	0.04 (0.03)	ACV (TL)	NS				
	Mn	3.77 (2.18)	1.84 (0.91)	1.76 (0.76)	ACV (TL)	***	CO>IR/BR			
	Ni	3.72 (5.65)	1.51 (2.33)	1.34 (2.32)	ANV	*	CO>BR	CO1>IR1/BR1		CO1>CO2
	Pb	0.06 (0.05)	0.04 (0.03)	0.04 (0.03)	ANV	NS				
	Se	0.98 (0.27)	1.20 (0.38)	1.07 (0.42)	ANV	NS				
	Zn	42.9 (9.9)	55.8 (17.3)	55.1 (17.1)	ANV	NS				
Conger sp.	As	37.96 (31.44)	20.57 (21.08)	16.45 (16.60)	ACV $(\delta^{15} N)$	*	CO>BR			
	Cd	0.04 (0.04)	0.03 (0.04)	0.06 (0.05)	ANV	ns				
	Co	0.23 (0.19)	0.10 (0.06)	0.05 (0.03)	ACV (TL)	***	CO/IR>BR			
	Cr	7.88 (7.09)	0.95 (0.31)	2.26 (1.68)	ACV (TL)	**	CO/IR>BR	CO1>BR1		CO1>CO2
	Cu	2.37 (1.05)	1.32 (0.43)	1.88 (0.79)	ACV (TL)	ns				
	Fe	57.4 (40.2)	15.3 (4.0)	24.2 (9.7)	ACV (TL)	**	CO>IR/BR	CO1>BR1		CO1>CO2
	Hg	0.11 (0.07)	0.07 (0.03)	0.15 (0.17)	ACV $(\delta^{15} N)$	**	CO/BR>IR			
	Mn	3.7 (2.30)	1.76 (0.95)	2.28 (2.66)	ANV	ns				
	Ni	2.78 (2.25)	0.35 (0.14)	0.83 (0.67)	ACV (TL)	***	CO>IR /BR			
	Pb	0.07 (0.05)	0.03 (0.01)	0.09 (0.14)	ACV (TL)	ns				
	Se	2.31 (1.06)	1.37 (0.21)	1.51 (0.43)	ACV (TL)	**	CO>BR			
	Zn	59.5 (26.2)	33.8 (6.3)	39.0 (23.10)	ACV (TL)	*	CO>IR/BR			

ANOVA (ANV) or ANCOVA (ACV) with Tukey's pairwise comparisons were used to analyze spatial patterns

GN Grand Nouméa, GS Grand Lagon Sud, CO coastal sites, IR intermediate reef sites, BR barrier reef sites, ns not significant (p>0.05)*p<0.05, **p<0.01, ***p<0.01

Most of them were mainly significantly correlated with the lipid contents in fish (Table 5). For example, PCBs were only correlated with lipids in both species. Pesticides were also correlated with Lp, and to a lesser extent with δ^{15} N and TL (Table 5).

Discussion

This study confirms, reinforces, and extends the high trace element contamination status of anguilliform fish in New Caledonia (Bonnet et al. 2014). It also constitutes the first substantial baseline on organic pollutant contamination of fish in New Caledonia. Importantly, our results show that large scale contamination reaches remote parts of the lagoon, more than 30 km offshore (SW lagoon), and thus well beyond the urbanized Nouméa region.

Contaminant levels and spatial variability

Trace element patterns

Anguilliform fish concentrate both significant levels of trace elements associated with mining exploitation (Cr, Fe, Mn, and Ni for both species), and also some elements linked to urban activities (e.g., Cd, Cu, and Zn, especially in *G. chilospilus*) (Table 2). This elevated typical mining signature detected in sedentary benthic species contrasts with other studies which highlight low contaminant levels in some more mobile fish in both the lagoons of New Caledonia and other tropical regions (Eilser 2010; Bonnet et al. 2014; Table 6).

Although coastal sites in New Caledonia are highly contaminated by trace elements, directly, or indirectly linked to mining activity (Co, Cr, Fe, Mn, and Ni), the contamination



IR												
G. chilotopalia 1. Cross (9.21) 6.864 (67.30) 6.062 (43.53) ANV B. S. Chadicay I. N. Childhopalia 1. S. Chi				Gradient					Spatial patterns	so		
G. chilotophilax PCB index (tig g ⁻¹ dw) PCB index (tig g ⁻¹ dw) PCB index (tig g ⁻¹ dw) PCB divinitation classes (% of C1.016) (2.16 (a) 13.74 (b).4) ANV is 3.7(4.04)				00	R	BR	Stat test	d	Gradient	RN	GS	GN vs. GS
PCB index (tag g-1 dw)		G. chilospilus										
Fig. 26.96 (9.21) 68.64 (67.30) 60.62 (43.55) ANV Bis Fig. ANV Bis Fig. ANV Bis Fig. ANV Bis ANV Bis ANV ANV Bis ANV A	4)	PCB index (ng g ⁻¹	(wp									
ES 602 (2.16) 17.00 (16.02) 13.74 (9.04) ANV ns CO>IRBR CO2-IRCJBR 1388 11.7 (052) 0.76 (0.13) 0.80 (0.10) ACV (Lp) *** CO>IRBR CO2-IRCJBR 11 2.0.39 (7.04) 16.64 (5.30) 14.12 (3.21) ACV (Lp) ns CO>BR CO2-IRCJBR 11 2.0.39 (4.87) 14.84 (7.34) 18.12 (5.31) ACV (Lp) ns CO>BR CO2-IRCJBR 11 2.26 (4.87) 3.43 (6.83) 30.48 (5.83) ANV ** CO>BR CO2-IRCJBR 11 3.55 (1.12) 3.82 (1.99) 4.33 (2.74) ANV ** CO1-BRI 11 3.55 (1.12) 3.82 (1.99) 4.33 (2.74) ANV ** CO1-BRI 12 2.25 (3.12) 0.82 (0.24) 0.50 (0.75) ANV ** CO1-BRI 1.1 0.11 (0.02) 0.06 (0.024) 0.37 (0.39) KW ** CO2-BR 1.1 1.05 (1.04) 1.25 (1.04) 1.35 (0.025) KW **<			PCBTot	26.96 (9.21)	68.64 (67.30)	60.62 (43.55)	ANV	ns				
438 1.17 (6.52) 6.76 (0.13) 0.80 (0.10) ACV (4.p) **** CO-IR/BR CO2-IR/BR 1.0 (3.05 of 9.93) 1.48 (7.94) 14.12 (3.22) ANV * CO-BR CO2-IR/BR2 1.1 2.60 (9.93) 1.48 (7.94) 18.12 (5.31) ACV (4.p) ns CO-BR CO2/IR/BR2 1.1 2.60 (9.93) 1.48 (7.94) 18.12 (5.31) ACV (4.p) ns CO-BR CO2/IR/BR2 1.1 2.807 (6.89) 2.9.56 (8.73) 3.48 (5.83) ANV ** RP>CO CO2/IR/BR2 1.1 3.55 (1.15) 3.82 (1.99) 4.33 (2.72) AVV ** CO1>BR CO2/IR/BR2 1.1 3.55 (1.15) 3.82 (1.90) ANV ** CO1>BR CO1 CO1 1.1 3.55 (1.15) 3.82 (1.90) ANV ** CO1>BR CO1>BR <td></td> <td></td> <td>$\Sigma ICES$</td> <td>6.02 (2.16)</td> <td>17.00 (16.02)</td> <td>13.74 (9.04)</td> <td>ANV</td> <td>ns</td> <td></td> <td></td> <td></td> <td></td>			$\Sigma ICES$	6.02 (2.16)	17.00 (16.02)	13.74 (9.04)	ANV	ns				
cs (% of CTotal PCBs) cc (2.39 (7.57) 16.64 (5.30) 14.12 (3.23) ANV * CO> BR CO> IIII 1 1 2.0.09 (9.33) 14.84 (7.34) 18.12 (5.33) 3.0.48 (5.85) ANV * IR>CO R CO> IIII IIII IIII AS (1.53) 3.246 (7.44) ANV ** R COI > BR COI > BR R COI > BR COI > BR R COI > BR COI > BR R COI > BR COI > BR R COI > BR R COI > BR COI			153/138	1.17 (0.52)	0.76 (0.13)	0.80 (0.10)	ACV (Lp)	* * *	CO>IR/BR		CO2>IR2/BR2	CO2>CO1
1		PCB chlorination	classes (% of CTo	tal PCBs)								
1			3 CI	20.39 (7.67)	16.64 (5.30)	14.12 (3.22)	ANV	*	CO>BR		CO2/IR2>BR2	CO2>CO1
1 25.65 (4.87) 34.32 (6.53) 30.48 (5.85) ANV * IR>CO 1 28.07 (6.89) 29.56 (8.73) 32.46 (7.44) ANV ns IR>CO 1 3.55 (1.15) 3.82 (1.99) 4.33 (2.72) ACV (Lp) ns CO1>BRI 1 2.25 (3.12) 0.82 (1.17) 0.50 (0.75) ANV ** CO1>BRI 1 0.16 (0.11) 0.11 (0.03) 0.06 (0.04) KW ** CO1>BRI 1 0.41 (0.26) 0.36 (0.24) 1.35 (0.39) KW ns CO2>BR 0.1 1.09 (1.06) 1.27 (1.04) 1.35 (1.80) KW ns CO3-BR 0.1 1.09 (1.06) 1.27 (1.04) 1.36 (1.80) KW ns CO3-BR 0.1 1.09 (1.06) 1.27 (1.04) 1.36 (1.09) KW ns CO3-BR 0.1 0.10 (0.09) 0.23 (0.24) 0.34 (0.39) KW ns CO3-BR 0.1 0.10 (0.09) 0.24 (0.14) 1.38 (1.09) A			4 CI	20.09 (9.93)	14.84 (7.94)	18.12 (5.31)	ACV (Lp)	ns				
1			5 CI	25.65 (4.87)	34.32 (6.53)	30.48 (5.85)	ANV	*	IR>CO			
$ \begin{array}{llllllllllllllllllllllllllllllllllll$			6 CI	28.07 (6.89)	29.56 (8.73)	32.46 (7.44)	ANV	ns				
1			7 Cl	3.55 (1.15)	3.82 (1.99)	4.33 (2.72)	ACV (Lp)	ns				
0.16 (0.11) 0.11 (0.03) 0.06 (0.04) KW *** CO>BR 3.91 (3.41) 1.47 (1.68) 1.35 (0.92) KW ns CO>BR 0.41 (0.26) 0.36 (0.24) 0.37 (0.39) KW ns CO>BR o.1 1.09 (1.06) 1.27 (1.04) 1.36 (1.66) ACV ns CO>III o.1 1.53 (0.86) 2.38 (2.16) 2.00 (1.54) ANV ns CO>IR/BR r. 0.16 (0.09) 0.28 (0.62) 0.34 (0.97) KW ns CO>IR/BR r. 0.16 (0.09) 0.28 (0.62) 0.34 (0.97) KW ns CO>IR/BR r. 0.16 (0.09) 0.28 (0.17) ACV (Lp) *** CO>IR/BR r. 0.16 (0.02) 0.23 (0.23) 0.27 (0.60) ANV ms CO>IR/BR r. 0.20 (0.23) 0.23 (0.23) 0.20 (0.77) ANV ms CO>IR/BR g 0.23 (0.23) 0.24 (0.21) 0.41 (0.63) KW <td< td=""><td></td><td></td><td>8 CI</td><td>2.25 (3.12)</td><td>0.82 (1.17)</td><td>0.50 (0.75)</td><td>ANV</td><td>* *</td><td></td><td>CO1>BR1</td><td></td><td>GN>GS</td></td<>			8 CI	2.25 (3.12)	0.82 (1.17)	0.50 (0.75)	ANV	* *		CO1>BR1		GN>GS
$\begin{array}{cccccccccccccccccccccccccccccccccccc$		Pesticides (ng g ⁻¹)	dw)									
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$			Ald.	0.16 (0.11)	0.11 (0.03)	0.06 (0.04)	KW	*	CO>BR			
$\begin{array}{llllllllllllllllllllllllllllllllllll$			Dia.	3.91 (3.41)	1.47 (1.68)	1.35 (0.92)	KW	ns				
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$			Diel.	0.41 (0.26)	0.36 (0.24)	0.37 (0.39)	KW	ns				
1.53 (0.86) 2.38 (2.16) 2.00 (1.54) ANV ns ns 0.16 (0.09) 0.28 (0.62) 0.34 (0.97) KW ns 0.83 (0.51) 0.43 (0.48) 0.20 (0.17) ACV (Lp) *** CO>IR/BR 0.66 (0.22) 1.69 (1.49) 1.58 (1.09) ANV ns 0.30 (0.23) 0.23 (0.33) 0.27 (0.60) ANV ns 0.30 (0.23) 0.34 (0.36) 0.50 (0.77) ANV ns 0.23 (0.24) 0.34 (0.36) 0.50 (0.77) ANV ns 0.23 (0.27) 0.41 (0.52) KW ns 0.29 (0.17) 0.24 (0.21) 0.41 (0.52) KW ns 0.29 (0.17) 0.44 (1.58) 1.38 (1.36) 0.50 (0.12) 0.51 (0.12) ANV ** CO>IR/BR			Endo.I	1.09 (1.06)	1.27 (1.04)	1.36 (1.66)	ACV	ns				
1.53 (0.86) 2.38 (2.16) 2.00 (1.54) ANV IIS 0.16 (0.09) 0.28 (0.62) 0.34 (0.97) KW IIS 0.66 (0.22) 1.69 (1.49) 1.58 (1.09) ANV IIS 0.30 (0.23) 0.23 (0.33) 0.27 (0.60) ANV IIS 1.32 (0.47) 1.50 (0.85) 1.22 (0.46) ANV IIS 0.23 (0.17) 0.24 (0.21) 0.41 (0.52) KW IIS 1.88 (1.58) 0.86 (1.19) 0.47 (0.63) KW IIS 2.40 (1.70) 1.44 (1.58) 1.38 (1.36) ANV IIS 0.51 (0.20) 0.5 (0.12) 0.51 (0.12) ANV IIS 0.52 (0.13) 0.5 (0.12) 0.51 (0.12) ANV IIS 0.54 (0.13) 0.5 (0.12) 0.51 (0.12) ANV IIS 0.55 (0.15) 0.51 (0.12) ANV IIS 0.55 (0.15) 0.51 (0.12) ANV IIS 0.55 (0.15) 0.51 (0.12) ANV IIS						3	(N :: 8)					
0.16 (0.09) 0.28 (0.62) 0.34 (0.97) KW ns ns 0.83 (0.51) 0.43 (0.48) 0.20 (0.17) ACV (Lp) *** CO>IR/BR 0.66 (0.22) 1.69 (1.49) 1.58 (1.09) ANV ns 0.30 (0.23) 0.23 (0.33) 0.27 (0.60) ANV ns 0.23 (0.47) 1.50 (0.85) 1.22 (0.46) ANV ns 0.23 (0.23) 0.34 (0.36) 0.50 (0.77) ANV ns 0.29 (0.17) 0.24 (0.21) 0.41 (0.52) KW ns 0.29 (0.17) 0.41 (0.52) KW ns 0.29 (0.17) 0.44 (1.58) 1.38 (1.36) 0.5 (0.12) 0.51 (0.12) ANV ** CO>IR/BR			Endo. II	1.55 (0.86)	2.38 (2.16)	2.00 (1.54)	ANA	ns				
0.83 (0.51) 0.43 (0.48) 0.20 (0.17) ACV (Lp) *** CO>IR/BR 0.66 (0.22) 1.69 (1.49) 1.58 (1.09) ANV ns 0.30 (0.23) 0.23 (0.33) 0.27 (0.60) ANV ** 1.32 (0.47) 1.50 (0.85) 1.22 (0.46) ANV ns 0.23 (0.23) 0.34 (0.36) 0.50 (0.77) ANV ns 0.29 (0.17) 0.24 (0.21) 0.41 (0.52) KW ns 1.88 (1.58) 0.86 (1.19) 0.47 (0.63) KW ns 2.40 (1.70) 1.44 (1.58) 1.38 (1.36) ANV *** CO>IR/BR			Endr.	0.16(0.09)	0.28 (0.62)	0.34 (0.97)	KW	ns				
0.66 (0.22) 1.69 (1.49) 1.58 (1.09) ANV ns CO2>IR2/BR2 0.30 (0.23) 0.23 (0.33) 0.27 (0.60) ANV * CO2>IR2/BR2 1.32 (0.47) 1.50 (0.85) 1.22 (0.46) ANV ns CO2>IR2/BR2 0.23 (0.23) 0.34 (0.36) 0.50 (0.77) ANV ns Ns 0.29 (0.17) 0.24 (0.21) 0.41 (0.52) KW ns Ns 1.88 (1.58) 0.86 (1.19) 0.47 (0.63) KW ns Ns 2.40 (1.70) 1.44 (1.58) 1.38 (1.36) ANV *** CO>IR/BR			Hept.	0.83 (0.51)	0.43 (0.48)	0.20 (0.17)	ACV (Lp)	* * *	CO>IR/BR			
B 0.30 (0.23) 0.23 (0.33) 0.27 (0.60) ANV * CO2>IR2/BR2 1.32 (0.47) 1.50 (0.85) 1.22 (0.46) ANV ns CO2>IR2/BR2 0.23 (0.23) 0.34 (0.36) 0.50 (0.77) ANV ns RW ns 0.29 (0.17) 0.24 (0.21) 0.41 (0.52) KW ns RW ns 0.7 1.88 (1.58) 0.86 (1.19) 0.47 (0.63) KW ns RW ns 0.7 2.40 (1.70) 1.44 (1.58) 1.38 (1.36) ANV *** CO>IR/BR			Hept. A	0.66 (0.22)	1.69 (1.49)	1.58 (1.09)	ANV	su				
1.32 (0.47) 1.50 (0.85) 1.22 (0.46) ANV ns 0.23 (0.23) 0.34 (0.36) 0.50 (0.77) ANV ns 1.88 (1.58) 0.86 (1.19) 0.41 (0.52) KW ns 1.88 (1.58) 0.86 (1.19) 0.47 (0.63) KW ns 1.44 (1.58) 1.38 (1.36) ANV ns 1.38 (1.36) 0.5 (0.12) 0.51 (0.12) ANV ****			Hept. B	0.30 (0.23)	0.23 (0.33)	0.27 (0.60)	ANV	*			CO2>IR2/BR2	IR1>IR2
0.23 (0.23) 0.34 (0.36) 0.50 (0.77) ANV ns 0.29 (0.17) 0.24 (0.21) 0.41 (0.52) KW ns 1.88 (1.58) 0.86 (1.19) 0.47 (0.63) KW ns 7T 2.40 (1.70) 1.44 (1.58) 1.38 (1.36) ANV ns 1.45 \(\subseteq \text{TOPCB}\) 1.13 (0.30) 0.5 (0.12) 0.51 (0.12) ANV ***			Lind.	1.32 (0.47)	1.50 (0.85)	1.22 (0.46)	ANV	ns				
0.29 (0.17) 0.24 (0.21) 0.41 (0.52) KW ns 1.88 (1.58) 0.86 (1.19) 0.47 (0.63) KW ns 0.7 2.40 (1.70) 1.44 (1.58) 1.38 (1.36) ANV ns π/ΣΡCΒ 1.13 (0.30) 0.5 (0.12) 0.51 (0.12) ANV ****			DDE	0.23 (0.23)	0.34 (0.36)	0.50 (0.77)	ANV	ns				
1.88 (1.58) 0.86 (1.19) 0.47 (0.63) KW ns 0.7 2.40 (1.70) 1.44 (1.58) 1.38 (1.36) ANV ns 1.13 (0.30) 0.5 (0.12) 0.51 (0.12) ANV ***			DDD	0.29 (0.17)	0.24 (0.21)	0.41 (0.52)	KW	su				
2.40 (1.70) 1.44 (1.58) 1.38 (1.36) ANV ns ΣPCB 1.13 (0.30) 0.5 (0.12) 0.51 (0.12) ANV ***			DDT	1.88 (1.58)	0.86 (1.19)	0.47 (0.63)	KW	ns				
1.13 (0.30) 0.5 (0.12) 0.51 (0.12) ANV ***			ΣDDT	2.40 (1.70)	1.44 (1.58)	1.38 (1.36)	ANV	ns				
			$\Sigma Pest/\Sigma PCB$	1.13 (0.30)	0.5 (0.12)	0.51 (0.12)	ANV	* * *	CO>IR/BR			



Table 4 (continued)

			Gradient					Spatial patterns	St		
			00	IR	BR	Stat test	D	Gradient	CN	CS	GN vs. GS
	Conger sp.										
B)	PCBs (ng g ⁻¹ dw)	(A									
	PCBTot	67.66 (54.60)	99.47 (67.68)	31.91 (16.39)	ACV (Lp)	* *	IR>BR				
	$\Sigma ICES$	17.17 (12.78)	26.32 (18.94)	8.58 (4.55)	ACV (Lp)	**	IR>BR				
	153/138	0.82 (0.19)	0.88 (0.36)	1.0 (0.47)	ACV (Lp)	ns					
	PCB chlorination	PCB chlorination classes (% of CTotal PCBs)	otal PCBs)								
	3 CI	15.75 (7.20)	19.94 (6.08)	16.09 (7.62)	ACV (Lp)	ns					
	4 CI	17.71 (6.99)	11.07 (4.59)	21.74 (8.23)	ACV (Lp)	*	IR <br< td=""><td></td><td></td><td></td><td></td></br<>				
	5 CI	32.02 (8.23)	32.74 (7.86)	34.84 (5.25)	ACV (Lp)	ns					
	6 CI	26.5 (5.15)	32.37 (7.73)	22.28 (4.51)	ACV (Lp)	* * *	IR>CO>BR				
	7 Cl	6.50 (5.97)	3.62 (1.38)	3.70 (2.12)	ANV	*			CO2>BR2		
	8 CI	1.53 (1.38)	0.26 (0.24)	1.36 (1.03)	ACV (Lp)	ns					
	Pesticides (ng g ⁻¹ dw)	⁻¹ dw)									
	Ald.	0.13 (0.06)	0.12 (0.04)	0.10 (0.03)	KW	ns					
	Dia.	2.02 (4.74)	3.99 (2.31)	1.54 (1.74)	KW	* * *	IR>CO			CO2>CO1	
	Diel.	0.77 (0.64)	0.25 (0.09)	0.36 (0.17)	KW	*	CO>IR				
	Endo. I	1.06 (0.89)	0.61 (0.73)	1.30 (2.69)	ACV (Lp)	ns					
	Endo. II	3.78 (2.46)	2.61 (2.60)	2.26 (1.83)	ACV (Lp)	ns					
	Endr.	0.24 (0.31)	1.69 (3.19)	0.56 (1.55)	KW	ns					
	Hept.	0.59(0.40)	0.51 (0.55)	0.52 (0.31)	ACV (TL)	ns					
	Hept. A	2.37 (2.58)	2.47 (1.77)	0.88 (0.56)	ACV (Lp)	*	IR>BR				
	Hept. B	0.79 (0.80)	0.30 (0.48)	0.33 (0.19)	ACV (Lp)	ns					
	Lind.	2.56 (1.51)	1.25 (0.93)	1.23 (0.62)	ACV (Lp)	ns					
	DDE	0.44 (0.61)	0.37 (0.21)	0.36 (0.29)	ANV	ns					
	DDD	0.29 (0.30)	0.65 (0.22)	0.15 (0.09)	KW	* * *	IR>CO/BR		IR2>BR2		
	DDT	0.82 (1.07)	0.40 (0.61)	1.19 (1.90)	KW	ns					
	ΣDDT	1.56 (1.19)	1.42 (0.71)	1.70 (2.12)	ANV	ns					
	$\Sigma Pest/\Sigma PCB$	0.92 (1.29)	0.51 (0.21)	0.94 (1.02)	ACV (Lp)	*	CO/BR>IR				

GN Grand Nouméa; GS Grand Lagon Sud, CO coastal sites, IR intermediate reef sites, BR barrier reef sites, ns not significant (p>0.05)ANOVA or ANCOVA with Tukey's pairwise comparisons were used to analyze spatial patterns

^a Only one value

 $^*p < 0.05, \ ^**p < 0.01, \ ^***p < 0.001/(+)$ positive and (-) negative correlation



Table 5 Summary of correlations performed on trace elements and organic pollutants (PCBs and pesticides) concentrations versus total length (TL), trophic position (δ^{15} N), and lipid content (Lp) in *G. chilospilus* and *Conger* sp

	Trace elements	PCBs	Pesticides
G. chilospilus			
TL	As*, Cd**, Cu*, Fe*, Hg*, Mn*		Ald.**, Dia.**
$\delta^{15}N$			Endo.*
Lp		CB153/138**, 4 Cl**, 8 Cl*	Hept.**
Conger sp.			
TL	Co**, Cr*, Cu**, Fe*, Ni*, Pb*, Se*, Zn*		Hept.**
$\delta^{15}N$	As***, Hg***		Ald.*
Lp		CTotal PCBs**, <i>SICES</i> *, <i>CB153/138</i> *, 3 Cl*, 4 Cl***, 5 Cl**, 6 Cl*, 8 Cl**	Dia.*, Diel.*, Endo.I**, Endo.II**, Endr.**, Hept.A*, Hep.B*, Lind.**, DDT**

Positive correlations are underlined

has spread throughout the entire lagoon as far as the remote barrier reef of the extremely vast south-western lagoon. This is probably due to dispersal by local currents (Fichez et al. 2008). However, it should be noted that concentrations decrease significantly from the coast to the barrier reef.

Unsurprisingly, fish from the coastal site in Grand Nouméa present important concentrations in trace elements, especially Cr, Fe, and Ni (Table 3). This urbanized and industrialized region is subjected to clear impact from the metallurgic industry (Hédouin et al. 2009, 2011; Metian et al. 2008, 2013). In contrast, the Grand Lagon Sud is less impacted by trace element inputs. Even if coastal sites in this area seem to be less threatened, trends still reveal important concentrations of Co, Mn, and Ni (Table 3). This strong contamination level likely results from severe soil erosion, due to deforestation and mining exploitation in this region over several decades. Given the numerous mines in New Caledonia and the intensive development of mining activities, assessing to what extent other areas of the New Caledonian lagoon have been affected is timely.

We expected a strong impact of urban pollution at least in Grand Nouméa sites, given the intense activity in the coastal industrial area of Ducos. However, while spatial patterns are obvious for trace elements associated with mining activities, those associated with urban pollution (As, Cd, Cu, Pb, and Zn) are less clear. Whereas Se and Zn are mostly concentrated in coastal sites for *Conger* sp., Cu and Pb are homogeneously distributed along the coast-barrier reef gradient. Furthermore, the cases of As and Cd are more complex with sometimes higher concentrations (e.g., As in *Conger* sp.) and sometimes lower concentrations in coastal sites (e.g., As and Cd in *G. chilospilus*). Finally, elements issued from urban pollution concern the entire lagoon, maybe due to their wide dispersal by local currents (Fichez et al. 2008). Another possible explanation is that in the past some islets were used as rubbish tips for

decades (e.g., Amédée islet in Nouméa region, Bonnet et al. 2014). Further information on these issues is thus needed.

Organic pollutant patterns

Contamination by POPs detected in anguilliform fish was rather low. However, considering the high toxicity of some POPs, even at very low doses, this contamination should be considered as important. Levels of PCB concentrations recorded in New Caledonia are generally lower than in fish from other tropical or temperate regions (e.g., $\Sigma ICES$ index in New Caledonia, 14.0± 12.2 ng g⁻¹ dw (*G. chilospilus*) and 16.5 \pm 14.3 ng g⁻¹ dw (Conger sp.); in Wallis, Central Pacific, 31.3±9.6 ng g⁻¹ dw (Cephalopholis argus), and 49.7±45.3 ng g⁻¹ dw (Epinephelus merra) (Letourneur et al., unpublished data); in the Mediterranean Sea, 100.6 ng g⁻¹ dw (Solea solea) (Dierking et al. 2009)). Exposure profiles of PCBs showed a contribution of chlorination classes in both species, with the prevalence of moderately chlorinated classes (5Cl and 6Cl) and the scarcity of highly chlorinated classes (7Cl and 8Cl). These results emphasize a recent or even current use of PCBs in New Caledonia, otherwise only the more persistent classes (highly chlorinated) should be detected. Further studies are needed to confirm this hypothesis.

In New Caledonian fish, pesticide concentrations are equivalent or lower than in other regions (e.g., lindane in New Caledonia, 1.3±0.6 ng g⁻¹ dw (*G. chilospilus*) and 1.7±1.6 (*Conger* sp.); in Wallis, Central Pacific, 3.5±1.03 ng g⁻¹ dw (*C. argus*) and 3.6±2.07 ng g⁻¹ dw (*E. merra*) (Letourneur et al., unpublished data); in French Polynesia, 73.3±34.5 ng g⁻¹ dw (*Epinephelus hexagonatus*) (Salvat et al. 2012); in the Mediterranean Sea, 35.4 ng g⁻¹ dw (*S. solea*) (Dierking et al. 2009)). Despite the quantities and diversity of pesticides imported into New Caledonia, these low levels detected reflect



³ Cl to 8 Cl classes of chlorination, Lind. lindane, Dia. diazinon, Hept. heptachlor, Ald. Aldrin, Hept. B heptachlor-epox. B, Hept. A heptachlor-epox. A, Endo. I endosulfan I, Diel dieldrin, Endr. endrin, DDT pp'-DDT

p < 0.05, p < 0.01, p < 0.001, significance levels

Table 6 Comparison of several trace element concentrations (mean±SD and range of values; micrograms per gram dry weight) measured in muscle of *G. chilospilus*, *Conger* sp., and other lagoon fish species in New Caledonia

Trace element	Concentration ($\mu g.g^{-1}dw$)	Species	Reference
As	11±7.4 (0.9–31.9)	G. chilospilus	This study
As	25.7±25.6 (3.0-117.9)	Conger sp.	This study
As	13.7±7.57 (<7.87-19.1)	Lethrinus laticaudis	Metian et al. (2013)
As	9.47±1.47 (<7.85-10.7)	Priacanthus hamrur	Metian et al. (2013)
As	<6.01-7.77	Cymbacephalus beauforti	Metian et al. (2013)
As	1.2-52.5	Mix	Fernandez and Breau (2011)
Cd	$0.41\pm0.47\;(0.01-2.91)$	G. chilospilus	This study
Cd	$0.04\pm0.04~(0.010.16)$	Conger sp.	This study
Cd	< 0.06	Mix	Fernandez and Breau (2011)
Co	$0.11\pm0.08~(0.02-0.45)$	G. chilospilus	This study
Co	0.13±0.15 (0.02-0.68)	Conger sp.	This study
Co	< 0.03	L. laticaudis	Metian et al. (2013)
Co	< 0.03	P. hamrur	Metian et al. (2013)
Co	< 0.02	C. beauforti	Metian et al. (2013)
Co	<0.1-1.1	Mix	Fernandez and Breau (2011)
Cr	4.53±7.15 (0.86–35.71)	G. chilospilus	This study
Cr	4.03±5.36 (0.45-23.60)	Conger sp.	This study
Cr	<0.79	L. laticaudis	Metian et al. (2013)
Cr	1.39 (<0.70–1.39)	P. hamrur	Metian et al. (2013)
Cr	<0.60	C. beauforti	Metian et al. (2013)
Cr	<0.1-5.7	Mix	Fernandez and Breau (2011)
Cu	1.6±0.7 (0.7–3.4)	G. chilospilus	This study
Cu	1.9±0.9 (0.7–4.5)	Conger sp.	This study
Cu	< 0.79	L. laticaudis	Metian et al. (2013)
Cu	< 0.83	P. hamrur	Metian et al. (2013)
Cu	<0.53-0.64	C. beauforti	Metian et al. (2013)
Cu	0.25-3.0	Mix	Fernandez and Breau (2011)
Fe	40.2±36.9 (12.2-196.8)	G. chilospilus	This study
Fe	34.3±31.1 (7.9–134.4)	Conger sp.	This study
Fe	9.19±1.42 (7.56–10.1)	L. laticaudis	Metian et al. (2013)
Fe	10.4±3.56 (7.82–16.9)	P. hamrur	Metian et al. (2013)
Fe	7.01–7.60	C. beauforti	Metian et al. (2013)
Fe	4.0-211	Mix	Fernandez and Breau (2011)
Mn	2.3±1.4 (0.6-7.2)	G. chilospilus	This study
Mn	2.7±2.3 (0.4–10.1)	Conger sp.	This study
Mn	< 0.79	L. laticaudis	Metian et al. (2013)
Mn	< 0.83	P. hamrur	Metian et al. (2013)
Mn	< 0.60	C. beauforti	Metian et al. (2013)
Mn	<0.03-3.3	Mix	Fernandez and Breau (2011)
Ni	1.95±3.19 (0.30-17.38)	G. chilospilus	This study
Ni	1.44±1.77 (0.10–7.74)	Conger sp.	This study
Ni	<0.06–1.35	Mix	Fernandez and Breau (2011)

Mix=144 fish from 27 species and 10 families (Acanthuridae, Carangidae, Haemulidae, Labridae, Lethrinidae, Lutjanidae, Mugilidae, Scaridae, Serranidae, and Siganidae)

their relatively moderate use, which probably comes from local and non-professional activities rather than from the agricultural industry. In addition, Nouméa is located far from the most important agricultural areas. This type of non-professional use could also explain a non-negligible detection of pesticides such as DDT and its metabolites (i.e., ΣDDT , 1.55±1.50 ng g⁻¹ dw

in *G. chilospilus* and 1.57 ± 1.48 in *Conger* sp.), which are supposed to be prohibited by international regulations and not homologated in New Caledonia (http://davar.gouv.nc/).

Overall, POP concentrations are homogeneous throughout the lagoon, without any clear spatial pattern along the coastbarrier reef gradient, or between studied areas. Nevertheless,



some high pesticide concentrations in *G. chilospilus* were observed in several coastal sites and in intermediate reefs, with high PCB and some high pesticide levels in *Conger* sp. (Tables 4 and S2). Storage organs (i.e., liver, kidney, or digestive gland) usually accumulate higher concentrations than muscle (e.g., Bustamante et al. 2003; Chouvelon et al. 2009; Metian et al. 2013; Eilser 2010). Further research is required to test this notion in anguilliforms and thus to better assess spatial contamination gradients, especially for POPs.

The results underlined the complexity in identifying the agricultural and/or industrial origin of organic contamination for each site (see $\Sigma Pest/\Sigma PCBs$ ratio, Table 4). Trends seem to emphasize a slight prevalence of agricultural contamination sources at coastal sites, in comparison with other sites dominated by industrial pollution. Furthermore, the Nouméa area generally seems to be more exposed to industrial contamination, whereas Grand Lagon Sud has revealed strong local agricultural pollution (except at intermediate reefs, Table S2). In Grand Nouméa, anguilliform fish have not been exposed to PCB inputs in recent times in view of their low CB153/CB138 ratio (i.e., ratio < 0.8, \approx 0.75 in G. chilospilus and \approx 0.78 in Conger sp., Table S2). On the other hand, the PCB inputs in Grand Lagon Sud seem to be more recent mainly in coastal sites, as detected in both species (ratio >1, 1.64 in G. chilospilus and 1.04 in Conger sp., Table S2). Industrial activities in this area, including the recent construction of a mining factory, could explain in part the continuing inputs of PCBs in this area.

Influence of biological parameters on contamination level

As mentioned in several studies, concentrations of contaminants can also vary according to ecological and biological factors such as fish size (or age), trophic position, and feeding habits or lipid content (Monteiro et al. 1991; Geyer et al. 2000; Penedo de Pinho et al. 2002). These complex relationships can induce interspecific differences in bioaccumulation processes and so emphasize the relevance in detecting the contamination rate at a specific level.

Our results indicated the influence of the three factors tested (body size, trophic position, and lipid content) on several contaminants in both species. The link between body size and bioaccumulation is complex, depending on both the contaminants and the fish species considered. In some cases, trace elements accumulated with increasing fish size (e.g., As, Cd, and Hg in *G. chilospilus*, Table 5), which is consistent with continuous bioaccumulation in muscle during life span (Braune 1987; Burger et al. 2007; Bloom 1992). In some other cases, trace element concentration is negatively correlated with size (e.g., Cu and Fe in both species, or Co, Cr, Ni, Pb, Se, and Zn in *Conger* sp.). This trend could be explained by a decrease in assimilation or by more efficient elimination processes with increasing fish size (Braune 1987; Swaileh and

Adelung 1995; Warnau et al. 1995). Such a relationship can also suggest that an ontogenetic diet shift might occur, implying a variation in the exposure to contaminants through the food pathway, as found in other species (Chouvelon et al. 2011, 2014). However, with the exception of Hg, and to a lesser extent Se, biomagnification of trace elements in trophic webs is still unclear and difficult to identify (Wang 2002). For anguilliform fish from New Caledonia, the trophic position influenced As and Hg concentrations only in *Conger* sp. The POP results are in accordance with their hydrophobic nature, which allows their sequestration in fat tissues during the process of organochlorine accumulation (Robertson and Hansen 2001). Most of them were indeed positively linked to the lipid content of fish, in particular in *Conger* sp., where lipid variation ranged between 0.2 and 5.5 % (Table 1).

Measures of δ^{15} N in fish confirm the high trophic level of anguilliform fish (Brischoux et al. 2011). When this is associated with other life history traits (size, age, longevity, philopatry, etc.), the detection of high trace element levels appears consistent. These benthic predators are also probably good candidates for monitoring organic pollution because of the strong relationship of POPs with lipid content (even if the flesh of these fish is not very fatty). The specific biological characteristics of each species may well explain interspecific differences in bioaccumulation of contaminants.

Moreover, contamination of anguilliform fish (high trophic level predators) suggests that trophic networks in general are contaminated (Monniot et al. 1994; Hédouin et al. 2006, 2007; Metian et al. 2008b). As a consequence, it is important to carry out contaminant analysis in prey, top predators (e.g., sea snakes, Bonnet 2012), and in the global trophic web. This will provide a more complete assessment of contamination of the environment and lead to better understanding of the mechanisms and processes involved (e.g., bioaccumulation, biomagnification, Bryan 1984; Rainbow and Phillips 1993). A good understanding of both the organisms (i.e., biology, ecology and metabolic capacities) and the trophic network structures, plus environmental conditions, is necessary to understand the dynamics of bioaccumulation and biomagnification throughout the trophic webs (Harmelin-Vivien et al. 2009).

Conclusion

This study strongly confirms a clear accumulation of trace metal contaminants in the muscles of anguilliform fish and the obvious coastal contamination by trace elements linked to mining activities. It is also shown that contamination by POPs occurs in the whole lagoon.

These results raise an alert about complex contaminations throughout the lagoon on a large spatial scale. Various contaminants may show different behavior in abiotic and biotic



environments. Due to their lipophilic nature, POPs have different targets compared with trace elements. This characteristic can explain in part their divergences in mobility in comparison to trace elements. Thus, the more widespread repartition of POPs within the lagoon is probably due to their great capacity of dispersion, through atmospheric transport and current-driven dilution during tropical rain events. This highlights the importance of considering multiple sources of contamination. Moreover, the possibility of a cocktail effect of these various contaminants is not known and could increase the threat. Therefore, in the future, long-term monitoring studies and experiments on toxicity are necessary.

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