INTRODUCTION
Polychlorinated biphenyls belong to a group of synthetic chlorine-containing compounds. They are a mixture of congeners built of biphenyl rings, saturated by different number of chlorine atoms, ranging from 1 to 10. These compounds are characterised by low vapour pressure and co-distil easily with water vapour. Due to their good heat conduction and dielectric properties, they were widely used in many industrial applications, mainly in electrical engineering. As a result of many years’ production and broad application, they are present in all elements of natural environment and in foods of vegetable- and animal origin (Hansen 1987, Falandysz et al. 2002).

Due to exceptional stability and concerns for its toxicity, PCB production was discontinued or their application was limited to closed cycles in many countries (Booker 2001). The PCBs, as lipophilic compounds, are characterised by high bioaccumulation coefficients in living organisms, and consequently in food items, which is the main source of their assimilation for most populations (Falandysz 1986, Anonymous 1999).

Considering the potentially high content of these compounds in seafood, their consumption may constitute a threat for the health of consumers. Thus, an important issue is to understand the effect of manufacturing processes and culinary procedures on concentration changes of PCBs.

Since smoked fish products are in great demand in Poland, the goal of this study was to determine changes in total PCB content in herring fillets during hot smoking and in hot and cold smoked mackerel fillets.
MATERIALS AND METHODS

The study material consisted of mackerel- and herring block fillets (= angel fillet or butterfly filet; a double fillet joined together along the back) from the fish caught in the Norwegian Sea in autumn 2001. After storing them at −18°C, the fillets were smoked. Supplementary raw materials were salt and a mixture of spicy condiment. Alder sawdust was used for smoke generation. The smoking was carried out under industrial conditions in a smoking chamber of the Atmos type (Kurko 1963), equipped with temperature- and humidity automatic control and forced-circulation warm-air heating system. The smoke generator was located outside the chamber. Prior to hot smoking, the mackerel and herring fillets were brined in 15% NaCl solutions for 3 min, whereas the mackerel fillets for cold smoking were brined (15% NaCl) with spicy condiment (laurel leaves, allspice, paprika, and mustard) for 3 h.

In order to determine changes occurring in PCB levels during the brining and smoking, 5 block fillets were split longitudinally into two full nape fillets (single fillets which included belly flap and rib bones) (later referred to as fillets), half of which constituted control samples and the other were brined and smoked.

For the hot smoking process, 5 brined block fillets were randomly sampled and split into two full nape fillets, for smoking under assumed time intervals (0.5; 1.0; 1.5; 2.0; and 2.5 h). During the first hour of smoking, the fillets were partially dried at 40°C, whereupon the smoke was supplied to the smoking chamber and the temperature was raised to 80°C. The process of hot smoking lasted for 2.5 h. In addition, 28 × 7 cm filtration paper strips immersed in soybean oil (with an average surface area of single mackerel fillets of 196 cm²), were hang up on upper smoking bars. The smoking chamber charge was 2 smoking trolleys, 60 kg of raw product each, and the consumption of alder sawdust amounted to 20 kg per smoking cycle.

The samples of mackerel fillets for the cold smoking process were prepared similarly. Fish fillets were removed from the smoking chamber after 2, 4, and 6 h of smoking, while the final product was collected after 8 h. The cold smoking was carried out at 27°C. The smoking chamber charge was 2 smoking trolleys, with 80 kg of fish each, and the consumption of sawdust amounted to 40 kg per smoking cycle.

To determine weight losses, the fillets were weighed before-and after the brining, as well as after successive smoking stages. For all samples, dry matter and lipid contents were determined. To determine which PCB congeners were present in the curing smoke, destructive wood distillation of alder sawdust was conducted under laboratory conditions. To determine recovery of analyzed PCB compounds, all samples were fortified with a known amount of surrogate Pesticides Surrogate Spike Mix (SUPELCO, USA), which was a solution of 2 compounds dissolved in acetone: decachlorobiphenyl and 2,4,5,6-tetrachloro-m-xylene. This was aimed at determining the recovery of total PCB, which amounted to 71%–88%.

“Chlorobiphenyls in mackerel oil” No. 350 (Promochem GmbH) was used as a reference material. In the analysis, a standard solution of 7 indicator congeners dissolved in isooctane (Promochem GmbH, D 46485 Wesel) was used. For determining the PCB total, a solution of 3 Aroclor standards (1242, 1254, 1260; N0132, N0135, and N0129, Promochem GmbH) was used.

For analysing the content of polychlorobiphenyls, the fillets were homogenised after skinning. Samples of 30 g of the muscle tissue were weighed and used for analysis in three replicates. The samples were dried by rubbing them with roasted Na2SO4 (at 400°C for 4 h) until uniform dry mass was obtained (according to Polish Standard PN-EN 15282-2). The extraction of the target compounds and lipid content was carried out in two stages: 50 mL of acetone and n-hexane solution (2.5 : 1), followed by 50 mL n-hexane and diethyl ether solution (9 : 1). After filtration, the combined extracts were concentrated in a rotary vacuum evaporator. The extract was transferred quantitatively into 10 mL weighed glass test-tubes, and evaporated under nitrogen atmosphere, and residues were desiccated at 60°C to a solid mass for determination of lipid content. The analysis for PCB compounds was continued by dissolving again the obtained lipid in n-hexane to 2 mL. Then, the sample was purified with 6 mL fuming H2SO4 (7% SO3 in concentrated H2SO4). After layer separation, the upper layer was rinsed three times with deionised water and dried with anhydrous Na2SO4. Next, the sample was concentrated in a rotary vacuum evaporator to 0.1 mL, and transferred to a tight vial.

The sample prepared this way was analysed with the method of gas chromatography coupled with mass spectrometry GC-MS (HP 6890/5973). A HP-5 column with 5% phenyl methyl siloxane (30 m, ID 250 µm, film thickness 0.25 µm) column was used. The analyses were made under the following chromatograph conditions: pulsed, splitless injection 2 µL; carrier gas-helium; column temperature-programme of oven’s column—140°C (hold 0.5 min), increase 30°C · min⁻¹, 280°C (hold 10 min), increase 10°C · min⁻¹, 280°C (hold 10 min), increase 30°C · min⁻¹, 300°C (hold 1 min).

Statistical processing of the results included analysis of variance with the aid of ANOVA tests (using STATISTICA 6.1 software package) and determination of appropriate coefficients of correlation and regression equations.

RESULTS

The average content of dry matter in the initial products was: 33.45% ± 0.63% (±standard deviation s) in herring fillets, 36.22% ± 0.62% in hot smoked mackerel and 36.88% ± 0.21% in cold smoked mackerel, and the lipid content was 14.42% ± 0.21%, 18.93% ± 0.99%, and 18.74% ± 0.21%, respectively.

Mean total PCB content in herring fillets before the brining was 64.35 ± 0.67 µg · kg⁻¹ of wet weight matter (w/w). In mackerel fillets assigned for hot smoking, mean PCB levels were 20.06 ± 0.43 µg · kg⁻¹ w/w, whereas in mackerel fillets for cold smoking mean PCB levels we-
re18.22 ± 0.45 µg · kg⁻¹ w/w. Lipid normalized data for PCBs was: 446.25 ± 11.77 µg · kg⁻¹ in herring fillets, 105.97 ± 2.47 µg · kg⁻¹ in mackerel fillets assigned for hot smoking, and 97.22 ± 0.41 µg · kg⁻¹ in mackerel fillets assigned for cold smoking (Fig. 1a–c).

Both in the mackerel and the herring fillets, no significant (P < 0.05) changes of total PCB were detected in wet weight or lipid normalized samples during the brining (Fig. 1).

During the hot smoking of mackerel fillets, a drop in total PCB content in wet weight was found in the preliminary smoking by 14.09 percentage points. During the proper smoking (after a preliminary drying, with smoke), the losses ranged from 22.12 percentage points after 1.5 h of smoking time.

**Fig. 1.** Total PCB (±s) content in the muscle tissue of hot smoked mackerel (a), hot smoked herring (b), and cold smoked mackerel (c)
to 23.32 percentage points in the final product. In lipids, the losses ranged from 25.57 percentage points after 0.5 h of preliminary drying to 38.71 percentage points after 2.5 h in the final product (Fig. 2). These changes were statistically significant ($P < 0.05$) in all examined time intervals. At the same time, an increase of dry weight and lipids was observed in the final product, by an average of 14.65 and 25.12 percentage points, respectively.

The hot smoking of herring fillets significantly decreased total PCB content by 13.32 percentage points after 0.5 h of preliminary drying, and by 30.17 percentage points in the final product after 2.5 h of smoking (Fig. 2). The content of lipids in the final product increased by 20.54 percentage points in relation to the initial content, with simultaneous increase of dry weight content by 15.87 percentage points, on the average.

During the cold smoking of mackerel fillets, significant increases in total PCB content, compared to initial levels, were observed up to 6 h (by 30.78 percentage points). However, after lipid normalization, these increases were much lower and statistically non-significant ($P < 0.05$) (Figs. 1, 2). At the same time, as the smoking time was prolonged, a repeated increase in lipid content in the final product (by 26.65 percentage points) was observed compared to the value after 2 h (by 3.35 percentage points) (Fig. 2), with simultaneous increase of dry matter in the final product by 44.78 percentage points.

While analysing individual factors affecting the changes of PCBs contents in the examined smoked fish flaps, a possible effect of the sawdust and the smoke obtained was taken into account. Sinkkonen et al. (1996) revealed the presence of PCB compounds in pine needles while Ciereszko et al. (2004) detected them in pine bark. However, the alder sawdust universally used in smoke-curing has not been analysed considering the contents of PCB compounds.

The total PCB content in alder sawdust amounted to $11.97 \pm 2.79 \mu g \cdot kg^{-1}$ dry matter. The smoke obtained from the alder sawdust during destructive wood distillation was $10.82 \pm 3.54 \mu g \cdot kg^{-1}$ dry matter of sawdust. The total PCB content in soybean oil on absorbent paper strips, which were hung above with the fish charge in the smoke-
ing chamber during hot smoking, increased in respective time intervals (Fig. 3). This experiment was provided in order to explain the changes of PCBs contents. These compounds were delivered with smoke to the oil on the filtration paper stripes thus to the fish flaps.

DISCUSSION

During the hot smoking process, more extensive losses in total PCB were observed in herring fillets when compared to the mackerel fillets (Figs. 1a–b, 2). This result is most likely from difference in the structure and consistency of the muscle tissue of these fish species. Both for the hot smoked mackerel and the herring fillets, negative coefficients of correlation between changes in total PCB content and smoking time \( r = -0.928; \) regression line: \( y = -10.216 - 2.236x \) and the herring fillets \( (r = -0.931; y = -4.660 - 1.393x) \).

The decrease in total PCB content in wet weight and when lipid normalized during the hot smoking process may be explained by the loss of these compounds in co-distillation with water vapour, which is suggested by negative rectilinear correlations between water loss and total PCB content change, both for the mackerel \( (r = -0.98; \) regression line: \( y = -10.216 - 2.236x \) and the herring fillets \( (r = -0.931; y = -4.660 - 1.393x) \).

According to Sherer and Price (1993), the temperature of heat processing is a significant determinate of the volatilization rate for these compounds, while their thermal degradation occurs at over 300°C (Morita et al. 1978). Zabik et al. (1996), Salama et al. (1998), and Ciereszko and Witczak (2003) also came to similar conclusions, when examining the heat processing methods used commonly for the fillets of different fish species. This may be also observed by the increase of total PCB content in soybean oil on filtration paper strips during the first hour of preliminary drying (Fig. 3).

The repeated increase in the content of these compounds in the hot smoked mackerel fillets results from the presence of PCB in the curing smoke, which penetrated into the smoked fish tissue (Fig. 4). This increase in the final product may be estimated at approximately 15 percentage points.

Based on these studies, total PCB levels in the hot smoked mackerel and herring fillets are a function of changes in the concentration of these compounds in lipids and changes of the percent content of lipids in wet matter of fish tissue during the whole smoking cycle. During the hot smoking of mackerel and herring fillets, strong negative correlations were found between total PCB changes and lipid contents \( r_{\text{mackerel}} = -0.98 \) and \( r_{\text{herring}} = -0.85 \). This suggests that such a significant loss of these compounds in lipids, despite the increase of PCBs on a per weight basis in the final product, determined conclusively the decrease of total PCBs. This finding is supported by the earlier studies of Sherer and Price (1993), who determined that PCB losses depended mainly on changes in lipid content, heat processing temperature and duration.

During the whole hot smoking process, dry matter content increased in the final products by 14.65 percentage points in mackerel fillets and by 16.67 percentage points in herring fillets on the average, whereas total PCB content decreased compared to the initial content by 23.32 and 30.17 percentage points, respectively. These results are similar to those obtained earlier by Zabik et al. (1996). They found that out of the heat processing methods examined, the most extensive loss of total PCB concentration (up to 39% of the final product), in the muscle tissue of lake trout occurred during hot smoking. Salama et al. (1998) concluded that the hot smoking process resulted in the highest decrease in total PCB content (65 percentage points) in the fillets of North Atlantic bluefish, Pomatomus saltatrix, compared to other methods such as baking or pan frying (27–39 percentage point losses).

When examining the cold smoked mackerel fillets, a 30 percentage point increase of total PCB content in the

![Fig. 4](image)

**Fig. 4.** The impact of curing smoke on changes in total PCB content in the lipids of hot smoked mackerel fillets; (A), total PCB loss curve; (B), hypothetic total PCB loss curve, after subtracting PCB content on filtration paper strips.
final product was observed (Fig. 2). Statistical analysis of the obtained results showed a rectilinear correlation between the increase of total PCB content in the muscle tissue and the length of smoking time \( (P < 0.05, r = 0.936) \). The increase of total PCB content in the wet matter of final product may be explained by lower losses of these compounds in co-distillation with water vapour due to low temperature of cold smoking \( (25–27^\circ C) \), and thereby much compact structure of muscle tissue when compared to the hot smoked fish and penetration of these compounds to the fillets from the curing smoke.

**CONCLUSIONS**

1. The hot smoking process resulted in decrease of total PCB content in the examined fish fillets, primarily from the loss of lipid contents.
2. The main factor determining the changes in the level of total PCB content in wet matter and lipids was their loss in co-distillation with water vapour.
3. The increased levels of PCBs in the final stage of hot smoking were most likely the result of contamination from curing smoke.
4. The cold smoking process of mackerel fillets contributed to an increase of PCB content in the final product.

**REFERENCES**


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