



Norwegian State Pollution
Monitoring Programme

Report 537/93

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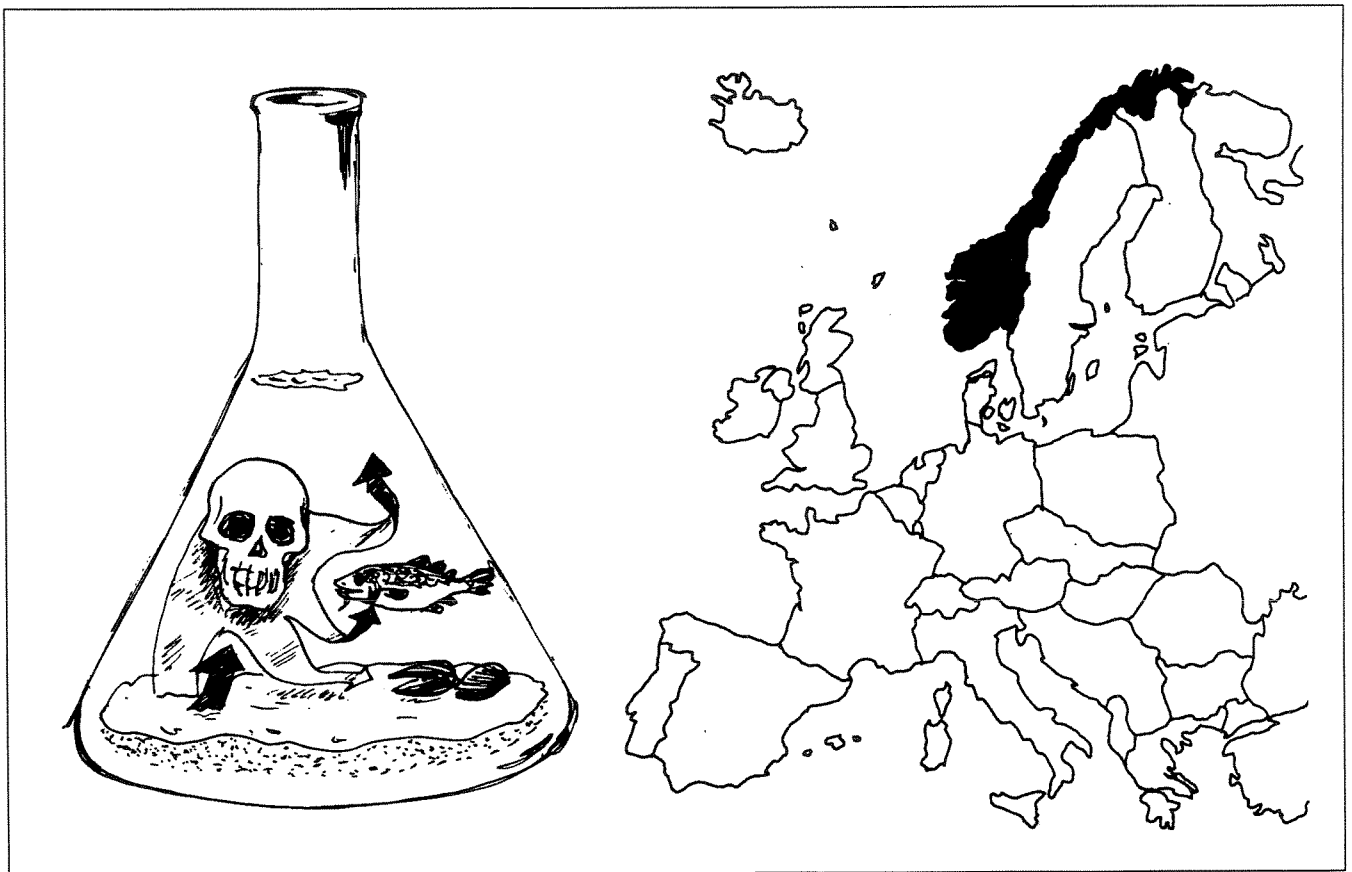
State Pollution Control Authority

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NIVA

Joint Monitoring Programme - JMP

Overview of Analytical Methods Employed by JMP in Norway 1981-92



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Abstract:
This report is a compilation of analytical method codes and descriptions used in the Norwegian contribution to the Joint Monitoring Programme monitoring of contaminants (mainly: selected metal, organochlorines, polycyclic aromatic hydrocarbons) in sea water, sea bed sediment and marine biota collected 1981-1992 and analyzed by June 1993. The method descriptions are brief and focus on the principles involved.

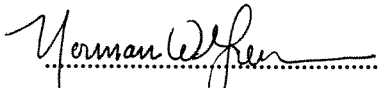
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1. Analytiske metoder
2. Mijøgifter
3. Sediment
4. Organismer


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1. Analytical methods
2. Contaminants
3. Sediment
4. Organisms

Project manager


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O-80106

Joint Monitoring Programme - JMP

**Overview of analytical methods employed by JMP
in Norway 1981-92**

Project coordinator;

Norman W.Green

3.11.93

FOREWORD

This report presents the Norwegian method overview and brief descriptions of chemical analyses used for 1981-92 investigations for the Joint Monitoring Programme (JMP). JMP is administered by the Oslo and Paris Commissions (OSPARCOM) under the guidance of the International Council for the Exploration of the Seas (ICES). The programme is implemented by participating members comprising the Joint Monitoring Group (JMG).

Information for this report was compiled by the Norwegian Institute for Water Research (NIVA) by contract from the Norwegian State Pollution Control Authority (SFT) (NIVA contract 80106) The report is an updated version of information reported earlier (OSPAR, 1984; Green, 1988; see Section 1)

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Oslo, 3.November 1993

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1. Introduction

This report outlines the analytical methods employed under the Joint Monitoring Programme (JMP) in Norway 1981-1992 for the analyses of contaminants in sea water, sea bed sediment and marine biota samples (collected 1981-92). It is largely based on information filed at the International Council for the Exploration of the Sea (ICES) and in particular the Norwegian contribution to the report:

- OSPAR, 1984. "Automatic data processing codes and descriptions of the sampling procedures and methods of analysis used in the Joint Monitoring Programme". Section 8 (Norway), 58/A/1-5/84 and 58/B/1/84. Oslo and Paris Commissions, October, 1984.

This report is largely a revision of OSPAR (1984) and Green (1988) and includes updated and extended information but retains the same codes and abbreviations. Section 2 summarizes the analytical methods employed sorted in various ways for convenient reference. Section 3 gives a brief description of each method.

2. Method overview

2.1. Medium and method code

Overview of analytical methods for trace metals, chlorinated hydrocarbons and other substances in the marine environment.(abbreviations are defined in section 2.4)

Medium	method code	institute code	contaminant(s) etc. (JMP code)	sampling year
Sea water				
	320	SIIF	Hg	84
	322	SERI	Hg	84-86
	330	SIIF	Cd	83
	331	NIVA	Cu,Zn	85-90
	331	NIVA	Cd,Pb	84-90
	331	NIVA	Fe,Ni,Co	86
	332	NIVA	Hg	86-90
Sea bed sediment				
	350	NIVA	Hg	86,90,92
	351	NIVA	Cu,Zn	86,90,92
	352	NIVA	Cd,Pb	86,90,92
	360	NIVA	PCB ¹	86,92
	369	NIVA	PAH ²	92
	390	NIVA	NTOT,CORG	86,90,92
	760	IMRN	PCB ¹	90
	769	IMRN	PAH ²	90

.../...

Method overview

Medium and method code overview (cont'd)

Medium	method code	institute code	contaminant(s) etc. (JMP code)	sampling year
Marinebiota				
	120	SIIF	Hg	81-85
	121	SIIF	Hg ³	83-85
	130	SIIF	Ni	83
	130	SIIF	Cu,Pb	84
	130	SIIF	Cd	81-85
	130	SIIF	Pb ³	83-85
	131	SIIF	Zn	83
	132	SIIF	Mn,Zn	84-85
	220	VETN	Hg	82-85
	220	VETN	Se ³	85
	230	VETN	Cd	82-85
	240	VETN	Se	82
	310	NIVA	Hg	86-92
	311	NIVA	Cu,Zn	86-92
	312	NIVA	Cd,Pb	86-92
	401	FIER	Hg	84,876
	402	FIER	Cd	84,87
	403	FIER	Pb	87
	404	FIER	Cu	87
	405	FIER	Zn	87
	406	FIER	As ³	83
	110	SIIF	PCB ¹	81
	111	SIIF	PCB ¹ ,	82-91
	210	VETN	PCB,HCB,DDEPP	82-85
	211	VETN	PCB	82-85
	309	NIVA	PAH ²	87,92
	340	NIVA	PCB ¹	87,90-92
	341	NIVA	PCB ¹	90-92
	510	NACE	PCB ¹	86-89
	511	NACE	PCB	86-89
	605	SIIF	EPOCI	86-91
	610	NACE	EPOCI	86-89
	615	NIVA	EPOCI	90-92

¹) selected individual chlorobiphenyls and pesticides (cf., Section 2.4).

²) selected PAHs (cf., Section 2.4).

³) not in data base

2.2. Seawater and sediment

Overview of chemical analyses employed by JMP 1981-1991 in Norway for trace metals, chlorinated hydrocarbons and other substances in sea water and sea bed sediment. Detection limits (detec.) are listed. Intercalibration codes (and in some cases laboratory codes, see Table 4) are given to distinguish different ICES exercises. "x" indicates data is registered at ICES but that no intercalibration exercise was filed. "-" indicates that no data are registered at ICES.

medium	parameter	detection limit ppb ¹	institute	sample year	method code	Intercal- ibration
Sea water						
	Cr	200	NIVA	87	331	-
	Mn	500	NIVA	87	331	-
	Fe	50	NIVA	87	331	-
	Co	5	NIVA	87	331	-
	Ni	10	NIVA	87	331	-
	Cu	10	NIVA	86-90	331	4I
		50	NIVA	85	331	4Z
	Zn	70	NIVA	85	331	4Z
		10	NIVA	86-90	331	4I
	Cd	1	NIVA	85	331	4Z
		0.5	NIVA	86-90	331	4I
		5	NIVA	84	331	4Z
		100	SIIF	83	330	4F
	Hg	0.02	SERI	84-85	322	4H
		0.02	SERI	86-90	322	4I
		2	NIVA	86	332	4I
		10	SIIF	84	320	4F
	Pb	6	NIVA	86-90	331	4Z
Sea bed sediment						
	Cu	10	NIVA	86,87,90	351	7E
	Zn	100	NIVA	86,87,90	351	7E
	Cd	1	NIVA	86,87,90	352	7E
	Hg	10	NIVA	86,87,90	350	7E
	Pb	50	NIVA	86,87,90	352	7E
	PCB	5	NIVA	86	360	-
	PCB	0.05	NIVA	92	360	8C-
	PAH	1	NIVA	92	369	-
	NTOT	0.1*	NIVA	86,87,90	390	-
	CORG	0.2*	NIVA	86,87,90	390	-
	PCB ²	0.05	IMRN	90	760	8B
	PAH ³	1	IMRN	90	769	-

*) mg/g

¹) note definition in section 2.6.

²) all individual chlorobiphenyls and pesticides (cf., section 2.4).

³) all PAHs (cf., section 2.4).

2.3. Biota

Overview of chemical analyses employed by JMP 1981-1992 in Norway for trace metals, chlorinated hydrocarbons and other substances in biota. Detection limit is the general limit for the method and not specific for a particular sample. "Count" (n) refers to number of values registered in JMP database at NIVA. "N (<)" indicates number of values with less-than qualifier above detection limit. Note qualifier for detection limit in section 2.6. Code explanations for intercalibration exercises are listed in Section 2.5

This section is sorted in three ways for easy reference:

- | | |
|----------------|---|
| Section 2.3.1. | Method, laboratory and tissue |
| Section 2.3.2 | Tissue, method and laboratory |
| Section 2.3.3. | Contaminant, year-laboratory and tissue |

Method overview

2.3.1. Method, laboratory and tissue

Analytical overview sorted by METHOD, LAB. and TISSUE.

Method	Lab.	Tissue=	Sample Year	Contaminants
110	SIIF	Fish fillet	81	PCB
		Fish liver	81	PCB
111	SIIF	Mussel	81	PCB
		Mussel	82-91	PCB
		Mussel	83-91	DDTEP, HCB
		Mussel	86-87,89-91	HCHG
		Mussel	87-91	CB101, CB180, CB52
		Mussel	88-91	CB138, CB153, CB28
		Mussel	89-91	CB118
		Shrimp tail	82,84,86,88,90	PCB
		Shrimp tail	84,86,88,90	DDTEP, HCB
		Shrimp tail	86,90	HCHG
		Shrimp tail	88,90	CB101, CB138, CB153, CB180, CB28, CB52
		Shrimp tail	90	CB118
		Other	88	CB101, CB138, CB153, CB180, CB28, CB52, DDTEP, HCB, PCB
120	SIIF	Fish fillet	81	HG
		Fish liver	81	HG
		Mussel	81-85	HG
		Shrimp tail	82,84	HG
130	SIIF	Fish fillet	81	CD
		Fish liver	81	CD
		Mussel	81-85	CD
		Mussel	83	NI
		Mussel	83-84	CU
		Mussel	83-85	PB
		Shrimp tail	82,84	CD
		Shrimp tail	84	CU, PB
131	SIIF	Mussel	83	ZN
132	SIIF	Mussel	84-85	MN, ZN
		Shrimp tail	84	MN, ZN
210	VETN	Fish fillet	83	DDEPP, HCB
		Fish liver	82-85	DDEPP, PCB
		Fish liver	83-85	HCB
211	VETN	Fish fillet	82-85	PCB
		Fish fillet	83	DDEPP, HCB
220	VETN	Fish fillet	82-85	HG
		Fish liver	82	HG
230	VETN	Fish liver	82-85	CD
240	VETN	Fish fillet	82	SE
		Fish liver	82	SE
309	NIVA	Fish fillet	92-93	ACNE, ACNLE, ANT, BAA, BAP, BBF, BEP, BGHIP, BIPN, BJKF, CHR, COR, DBA3A, DBP, FLE, FLU, ICDP, NAP, NAP1M, NAP2M, NAPDI, NAPTM, PA, PAM1, PER, PYR
		Fish liver	87	PAH
		Fish liver	92-93	ACNE, ACNLE, ANT, BAA, BAP, BBF, BEP, BGHIP, BIPN, BJKF, CHR, COR, DBA3A, DBP, FLE, FLU, ICDP, NAP, NAP1M, NAP2M, NAPDI, NAPTM, PA, PAM1, PER, PYR
		Mussel	92	ACNE, ACNLE, ANT, BAA, BAP, BBF, BEP, BGHIP, BIPN, BJKF, CHR, COR, DBA3A, DBP, FLE, FLU, ICDP, NAP, NAP1M, NAP2M, NAPDI, NAPTM, PA, PAM1, PER, PYR
		Shrimp tail	92	ACNE, ACNLE, ANT, BAA, BAP, BBF, BEP, BGHIP, BIPN, BJKF, CHR, COR, DBA3A, DBP, FLE, FLU, ICDP, NAP, NAP1M, NAP2M, NAPDI, NAPTM, PA, PAM1, PER, PYR
310	NIVA	Fish fillet	86-93	HG
		Mussel	86-92	HG
		Shrimp tail	86,88,90,92	HG
		Other	88	HG
311	NIVA	Fish liver	86-93	CU, ZN
		Mussel	86-92	CU, ZN
		Shrimp tail	86,88,90,92	CU, ZN
		Other	88	CU, ZN
312	NIVA	Fish liver	86-93	CD, PB
		Mussel	86-92	CD, PB
		Mussel	92	CR, NI
		Shrimp tail	86,88,90,92	CD, PB
		Other	88	CD, PB
340	NIVA	Fish liver	87	PCB
		Fish liver	90-93	CB101, CB118, CB138, CB153, CB180, CB209, CB28, CB52, DDEPP, HCB, HCHA, HCHG, OCS, QCB, TDEPP
341	NIVA	Fish liver	91-93	CB105, CB156
		Fish fillet	90-93	CB101, CB118, CB138, CB153, CB180, CB209, CB28, CB52, DDEPP, HCB, HCHA, HCHG, OCS, QCB, TDEPP
		Fish fillet	91-93	CB105, CB156
		Mussel	92	CB101, CB105, CB118, CB138, CB153, CB156, CB180, CB209, CB28, CB52, DDEPP, HCB, HCHA, HCHG, OCS, QCB, TDEPP
		Shrimp tail	92	CB101, CB105, CB118, CB138, CB153, CB156, CB180, CB209, CB28, CB52, DDEPP, HCB, HCHA, HCHG, OCS, QCB, TDEPP
401	FIER	Fish fillet	84,87	HG
402	FIER	Fish liver	84,87	CD
403	FIER	Fish liver	87	PB
404	FIER	Fish liver	87	CU
405	FIER	Fish liver	87	ZN
510	NACE	Fish liver	86-89	DDEPP, DDTTP, HCB, HCHG, PCB
		Fish liver	89	CB101, CB118, CB138, CB153, CB180, CB28, CB52
511	NACE	Fish fillet	86-89	PCB
		Fish liver	86	PCB
		Fish liver	86-91	PCB
605	SIIF	Mussel	89	EPOCL
		Mussel	89	EOCL
		Shrimp tail	86,88,90	EPOCL
		Other	88	EPOCL
610	NACE	Fish liver	86-89	EPOCL
615	NIVA	Fish liver	90-91	EPOCL

2.3.2. Tissue, method and laboratory

Analytical overview sorted by TISSUE, METHOD and LAB.

Tissue	Method	Lab.=	Sample Year	Contaminants
Fish fillet	110	SIIF	81	PCB
	120	SIIF	81	HG
	130	SIIF	81	CD
	210	VETN	83	DDEPP, HCB
	211	VETN	82-85	PCB
	211	VETN	83	DDEPP, HCB
	220	VETN	82-85	HG
	240	VETN	82	SE
	309	NIVA	92-93	ACNE, ACNLE, ANT, BAA, BAP, BBF, BEP, BGHIP, BIPN, BJKF, CHR, COR, DBA3A, DBP, FLE, FLU, ICDP, NAP, NAP1M, NAP2M, NAPDI, NAPTM, PA, PAM1, PER, PYR
	310	NIVA	86-93	HG
	341	NIVA	90-93	CB101, CB118, CB138, CB153, CB180, CB209, CB28, CB52, DDEPP, HCB, HCHA, HCHG, OCS, QCB, TDEPP
	341	NIVA	91-93	CB105, CB156
	401	FIER	84,87	HG
	511	NACE	86-89	PCB
	Fish liver	110	SIIF	81
120		SIIF	81	HG
130		SIIF	81	CD
210		VETN	82-85	DDEPP, PCB
210		VETN	83-85	HCB
220		VETN	82	HG
230		VETN	82-85	CD
240		VETN	82	SE
309		NIVA	87	PAH
309		NIVA	92-93	ACNE, ACNLE, ANT, BAA, BAP, BBF, BEP, BGHIP, BIPN, BJKF, CHR, COR, DBA3A, DBP, FLE, FLU, ICDP, NAP, NAP1M, NAP2M, NAPDI, NAPTM, PA, PAM1, PER, PYR
311		NIVA	86-93	CU, ZN
312		NIVA	86-93	CD, PB
340		NIVA	87	PCB
340		NIVA	90-93	CB101, CB118, CB138, CB153, CB180, CB209, CB28, CB52, DDEPP, HCB, HCHA, HCHG, OCS, QCB, TDEPP
340		NIVA	91-93	CB105, CB156
402	FIER	84,87	CD	
403	FIER	87	PB	
404	FIER	87	CU	
405	FIER	87	ZN	
510	NACE	86-89	DDEPP, DDTTP, HCB, HCHG, PCB	
510	NACE	89	CB101, CB118, CB138, CB153, CB180, CB28, CB52	
511	NACE	86	PCB	
610	NACE	86-89	EPOCL	
615	NIVA	90-91	EPOCL	
Mussel	110	SIIF	81	PCB
	111	SIIF	82-91	PCB
	111	SIIF	83-91	DDTEP, HCB
	111	SIIF	86-87, 89-91	HCHG
	111	SIIF	87-91	CB101, CB180, CB52
	111	SIIF	88-91	CB138, CB153, CB28
	111	SIIF	89-91	CB118
	120	SIIF	81-85	HG
	130	SIIF	81-85	CD
	130	SIIF	83	NI
	130	SIIF	83-84	CU
	130	SIIF	83-85	PB
	131	SIIF	83	ZN
	132	SIIF	84-85	MN, ZN
	309	NIVA	92	ACNE, ACNLE, ANT, BAA, BAP, BBF, BEP, BGHIP, BIPN, BJKF, CHR, COR, DBA3A, DBP, FLE, FLU, ICDP, NAP, NAP1M, NAP2M, NAPDI, NAPTM, PA, PAM1, PER, PYR
310	NIVA	86-92	HG	
311	NIVA	86-92	CU, ZN	
312	NIVA	86-92	CD, PB	
312	NIVA	92	CR, NI	
341	NIVA	92	CB101, CB105, CB118, CB138, CB153, CB156, CB180, CB209, CB28, CB52, DDEPP, HCB, HCHA, HCHG, OCS, QCB, TDEPP	
Shrimp tail	605	SIIF	86-91	EPOCL
	605	SIIF	89	EOCL
	111	SIIF	82,84,86,88,90	PCB
	111	SIIF	84,86,88,90	DDTEP, HCB
	111	SIIF	86,90	HCHG
	111	SIIF	88,90	CB101, CB138, CB153, CB180, CB28, CB52
	111	SIIF	90	CB118
	120	SIIF	82,84	HG
	130	SIIF	82,84	CD
	130	SIIF	84	CU, PB
	132	SIIF	84	MN, ZN
	309	NIVA	92	ACNE, ACNLE, ANT, BAA, BAP, BBF, BEP, BGHIP, BIPN, BJKF, CHR, COR, DBA3A, DBP, FLE, FLU, ICDP, NAP, NAP1M, NAP2M, NAPDI, NAPTM, PA, PAM1, PER, PYR
	310	NIVA	86,88,90,92	HG
	311	NIVA	86,88,90,92	CU, ZN
	312	NIVA	86,88,90,92	CD, PB
341	NIVA	92	CB101, CB105, CB118, CB138, CB153, CB156, CB180, CB209, CB28, CB52, DDEPP, HCB, HCHA, HCHG, OCS, QCB, TDEPP	
Other	605	SIIF	86,88,90	EPOCL
	111	SIIF	88	CB101, CB138, CB153, CB180, CB28, CB52, DDTEP, HCB, PCB
	310	NIVA	88	HG
	311	NIVA	88	CU, ZN
	605	SIIF	88	CD, PB

2.3.3. Contaminant, year-laboratory and tissue

Analytical overview sorted by CONTAMINANT, Year&Lab, Intercalibration+Basis and ordered by TISSUE.

Tissue	Fish liver					Fish fillet, Shrimptail, Mussel, Other						
	Contam.	Year& Lab.	Inter-calibr.+Basis	Analys Method Code	Detect Limit (ppb)	Total Value Count	Count Below D.Lim	N (<) Above D.Lim	Analys Method Code	Detect Limit (ppb)	Total Value Count	Count Below D.Lim
ACNE	92-NIVA	W	309	0.10	5		2	309	0.10	43		15
	93-NIVA	W	309	0.10	3			309	0.10	3		3
ACNLE	92-NIVA	W	309	0.10	5		2	309	0.10	43		19
	93-NIVA	W	309	0.10	3			309	0.10	3		3
ANT	92-NIVA	W	309	0.10	5			309	0.10	42		20
	93-NIVA	W	309	0.10	3			309	0.10	3		3
BAA	92-NIVA	W	309	0.10	5		5	309	0.10	41		3
	93-NIVA	W	309	0.10	3			309	0.10	3		3
BAP	92-NIVA	W	309	0.10	5		5	309	0.10	42		3
	93-NIVA	W	309	0.10	3		3	309	0.10	3		19
BBF	92-NIVA	W	309	0.10	5			309	0.10	42		6
	93-NIVA	W	309	0.10	3			309	0.10	3		3
BEP	92-NIVA	W	309	0.10	5			309	0.10	42		3
	93-NIVA	W	309	0.10	3			309	0.10	3		5
BGHIP	92-NIVA	W	309	0.10	5		1	309	0.10	3		3
	93-NIVA	W	309	0.10	3		2	309	0.10	43		17
BIPN	92-NIVA	W	309	0.10	5			309	0.10	3		3
	93-NIVA	W	309	0.10	3		2	309	0.10	43		6
BJKF	92-NIVA	W	309	0.10	5			309	0.10	3		3
	93-NIVA	W	309	0.10	3		3	309	0.10	42		21
CB101	87-SIIF	W	309	0.10	3		3	309	0.10	3		3
	88-SIIF	D						111	0.20	21	1	
	88-SIIF	W						111	0.10	6		
	89-NACE	W	510	20.00	93			111	0.10	22		
	89-SIIF	W						111	0.10	36		
	90-NIVA	2G W	340	1.00	169	1		341	0.05	58		
	90-SIIF	2G W						111	0.40	41	6	
	91-NIVA	2H W	340	1.00	179		8	341	0.05	62		
	91-SIIF	2H W						111	0.20	35		1
	92-NIVA	2H W	340	5.00	173	3		341	0.10	137		
	93-NIVA	2H W	340	5.00	19			341	0.10	3		
CB105	91-NIVA	2H W	340	1.00	87		1	341	0.05	47		
	92-NIVA	W	340	5.00	173	3		341	0.10	137		
	93-NIVA	W	340	5.00	19			341	0.10	3		
CB118	89-NACE	W	510	20.00	93							
	89-SIIF	W						111	0.10	36		
	90-NIVA	2G W	340	1.00	169			341	0.05	58		
	90-SIIF	2G W						111	0.20	41	1	
	91-NIVA	2H W	340	1.00	179			341	0.05	62		
	91-SIIF	2H W						111	0.20	35		1
	92-NIVA	2H W	340	5.00	173	2		341	0.10	137		
	93-NIVA	2H W	340	5.00	19			341	0.10	3		
CB138	88-SIIF	D						111	0.10	6		
	88-SIIF	W						111	0.10	21		
	89-NACE	W	510	20.00	93							
	89-SIIF	W						111	0.10	36		
	90-NIVA	2G W	340	1.00	169			341	0.05	58		
	90-SIIF	2G W						111	0.30	41		
	91-NIVA	2H W	340	1.00	179			341	0.05	62		
	91-SIIF	2H W						111	0.30	35		1
	92-NIVA	2H W	340	5.00	173			341	0.10	134		
	93-NIVA	2H W	340	5.00	19			341	0.10	3		
CB153	88-SIIF	D						111	0.10	6		
	88-SIIF	W						111	0.10	22		
	89-NACE	W	510	20.00	93							
	89-SIIF	W						111	0.10	36		
	90-NIVA	2G W	340	1.00	169			341	0.05	58		
	90-SIIF	2G W						111	0.30	41		
	91-NIVA	2H W	340	1.00	179			341	0.05	62		
	91-SIIF	2H W						111	0.50	35		1
	92-NIVA	2H W	340	5.00	173			341	0.10	137		
	93-NIVA	2H W	340	5.00	19			341	0.10	3		
CB156	91-NIVA	2H W	340	1.00	87		15	341	0.05	47		5
	92-NIVA	W	340	5.00	173	3		341	0.10	137		
	93-NIVA	W	340	5.00	19			341	0.10	3		
CB180	87-SIIF	W						111	0.20	21	6	
	88-SIIF	D						111	0.10	6		
	88-SIIF	W						111	0.10	22		
	89-NACE	W	510	20.00	93	1						
	89-SIIF	W						111	0.10	36		
	90-NIVA	2G W	340	1.00	169			341	0.05	58		
	90-SIIF	2G W						111	0.20	41	8	
	91-NIVA	2H W	340	1.00	179			341	0.05	62		
	91-SIIF	2H W						111	0.20	35		
	92-NIVA	2H W	340	5.00	173	3		341	0.10	137		
	93-NIVA	2H W	340	5.00	19			341	0.10	3		
CB209	90-NIVA	W	340	2.00	169	24	11	341	0.05	58		
	91-NIVA	W	340	2.00	179	11	88	341	0.05	62	5	7
	92-NIVA	W	340	5.00	173	3		341	0.10	137		1
	93-NIVA	W	340	5.00	19			341	0.10	3		
CB28	88-SIIF	D						111	0.10	6		
	88-SIIF	W						111	0.10	22		
	89-NACE	W	510	20.00	93							
	89-SIIF	W						111	0.10	36		1
	90-NIVA	2G W	340	1.00	169	2	2	341	0.05	58		
	90-SIIF	2G W						111	0.20	41	7	

Contaminant, year-laboratory and tissue (cont.)

Tissue	Fish liver							Fish fillet, Shrimptail, Mussel, Other					
	Contam.	Year & Lab.	Inter-calibr. Basis	Analys Method Code	Detect Limit (ppb)	Total Value Count	Count Below D.Lim	N (<) Above D.Lim	Analys Method Code	Detect Limit (ppb)	Total Value Count	Count Below D.Lim	N (<) Above D.Lim
CB52	91-NIVA	2H	W	340	1.00	179	2	52	341	0.05	62	5	1
	91-SIIF	2H	W						111	0.30	35		
	92-NIVA	2H	W	340	5.00	173	3		341	0.10	134		
	93-NIVA	2H	W	340	5.00	19			341	0.10	3		
	87-SIIF		W						111	0.20	20	1	
	88-SIIF		D						111	0.10	6		
	88-SIIF		W						111	0.10	22		
	89-NACE		W	510	20.00	93							
	89-SIIF		W						111	0.10	36		
	90-NIVA	2G	W	340	1.00	169	2	6	341	0.05	58		
	90-SIIF	2G	W						111	0.40	41	7	
	91-NIVA	2H	W	340	1.00	179	1	37	341	0.05	62	5	1
	91-SIIF	2H	W						111	0.30	35		
	CD	92-NIVA	2H	W	340	5.00	173	3		341	0.10	134	
93-NIVA		2H	W	340	5.00	19			341	0.10	3		
81-SIIF		1E	W	130	10.00	28			130	5.00	27		
81-SIIF		1F	W						130	10.00	7		
82-SIIF		1F	W						130	10.00	18		
82-VETN			W	230	10.00	54							
83-SIIF		1F	W						130	10.00	17		
83-VETN		1Z	W	230	10.00	46							
84-FIER		1H	W	402	1.00	23							
84-SIIF		1G	W						130	10.00	27		
84-VETN		1Z	W	230	10.00	66							
85-SIIF		1G	D						130	10.00	35		
85-VETN		1Z	W	230	10.00	45		3					
86-NIVA		1H	D	312	30.00	29	1		312	30.00	20		
87-FIER		1G	W	402	1.00	37							
87-NIVA		1H	D	312	30.00	59		1	312	30.00	37		
88-NIVA		1H	D	312	30.00	61	2	4	312	30.00	43		
89-NIVA		1H	D	312	30.00	160	20	8					
89-NIVA		1H	W						312	30.00	36		
90-NIVA		1H	W	312	10.00	189	9	2	312	30.00	77	5	
91-NIVA		1H	W	312	10.00	190	29	2	312	10.00	67		
92-NIVA		1H	W	312	10.00	172	4		312	10.00	111		
93-NIVA		1H	W	312	10.00	19							
CHR		92-NIVA		W	309	0.10	5		309	0.10	41		5
	93-NIVA		W	309	0.10	3		309	0.10	3		3	
	92-NIVA		W	309	0.10	5		309	0.10	43		3	
COR	93-NIVA		W	309	0.10	3		309	0.10	3			
	92-NIVA		W	309	0.10	5		309	0.10	43			
CR	92-NIVA		W					312	10.00	6			
	83-SIIF	1G	W					130	10.00	12			
CU	84-SIIF	1G	W					130	10.00	27			
	86-NIVA	1H	D	311	150.00	29		311	150.00	20			
DBA3A	87-FIER	1G	W	404	50.00	37							
	87-NIVA	1H	D	311	150.00	59		311	150.00	37			
	88-NIVA	1H	D	311	150.00	61		311	150.00	43			
	89-NIVA	1H	D	311	150.00	160							
	89-NIVA	1H	W					311	150.00	36			
	90-NIVA	1H	W	311	150.00	189		311	150.00	77			
	91-NIVA	1H	W	311	50.00	193	2	311	50.00	67			
	92-NIVA	1H	W	311	10.00	172		311	10.00	111			
	93-NIVA	1H	W	311	10.00	19							
	92-NIVA		W	309	0.10	5		309	0.10	43		35	
	93-NIVA		W	309	0.10	3		309	0.10	3		3	
	DBP	92-NIVA		W	309	0.10	5		309	0.10	43		
		93-NIVA		W	309	0.10	3		309	0.10	3		
	DDEPP	82-VETN		W	210	0.05	53						
		83-VETN	2E	W	210	50.00	48		211a	50.00	48		
		84-VETN	2E	W	210	50.00	66						
		85-VETN	2E	W	210	50.00	45						
		86-NACE	2Z	W	510	20.00	31						
		87-NACE	2Z	W	510	40.00a	54						
		88-NACE	2Z	W	510	40.00	60						
		89-NACE	2Z	W	510	40.00a	118						
		90-NIVA		W	340	1.00	169		341	0.05	58		
		91-NIVA		W	340	1.00	179		341	0.05	62		
		92-NIVA		W	340	5.00	173		341	0.10	137		
93-NIVA			W	340	5.00	19	2	341	0.10	3			
DDTEP		83-SIIF		W					111	0.50	12		
		84-SIIF		W					111	0.50	24		
	85-SIIF		W					111	0.50	27	1	5	
	86-SIIF		W					111	0.50	21			
	87-SIIF		W					111	0.50	21	1		
	88-SIIF		D					111	0.50	6			
	88-SIIF		W					111	0.50	22	1		
	89-SIIF		W					111	0.50	36	1		
	90-SIIF		W					111	0.20	41	1		
	91-SIIF		W					111	0.30	35			
	DDTPP	86-NACE		W	510	40.00	31						
87-NACE			W	510	40.00	54							
88-NACE			W	510	40.00	60							
89-NACE			W	510	40.00a	118							
EOCL	89-SIIF		W					605	170.00	5			
	86-NACE		W	610	800.00	31		605	5000.00	21	21		
EPOCL	86-SIIF		W					605	40.00	20			
	87-NACE		W	610	800.00	54							
	88-NACE		W	610	800.00	59							

Method overview

Contaminant, year-laboratory and tissue (cont.)

Tissue			Fish liver				Fish fillet, Shrimptail, Mussel, Other					
Contam.	Year & Lab.	Inter-calibr. Basis	Analys Method Code	Detect Limit (ppb)	Total Value Count	Count Below D.Lim	N (<) Above D.Lim	Analys Method Code	Detect Limit (ppb)	Total Value Count	Count Below D.Lim	N (<) Above D.Lim
FLE	88-SIIF	W	610	800.00	114	1		605	40.00	27		
	89-NACE	W										
	89-SIIF	W						605	40.00	35		
	90-NIVA	W	615	40.00	117		3					
	90-SIIF	W						605	40.00	41		
	91-NIVA	W	615	40.00	116		12					
	91-SIIF	W						605	130.00	35		
	92-NIVA	W	309	0.10	5		2	309	0.10	42		9
	93-NIVA	W	309	0.10	3			309	0.10	3		3
	FLU	92-NIVA	W	309	0.10	5			309	0.10	41	
93-NIVA		W	309	0.10	3			309	0.10	3		3
HCB	83-SIIF	W						111	0.50	12		
	83-VETN	22 W	210	10.00	48			211a	10.00	48		
	84-SIIF	W						111	0.20	24		1
	84-VETN	22 W	210	10.00	66							
	85-SIIF	W						111	0.20	30	6	2
	85-VETN	22 W	210	10.00	45		4					
	86-NACE	22 W	510	10.00	31							
	86-SIIF	22 W						111	0.20	21		3
	87-NACE	22 W	510	40.00a	54			111	0.20	21		4
	87-SIIF	22 W						111	0.20	6		
HCHA	88-SIIF	22 D						111	0.20	22		2
	88-SIIF	22 W						111	0.20	22		2
	89-NACE	22 W	510	40.00a	118							
	89-SIIF	22 W						111	0.05	36		
	90-NIVA	W	340	1.00	169	2		341	0.05	58		
	90-SIIF	22 W						111	0.05	41	3	
	91-NIVA	W	340	1.00	179	4	13	341	0.05	62	5	
	91-SIIF	22 W						111	0.10	35		
	92-NIVA	W	340	5.00	170	3		341	0.10	137		
	93-NIVA	W	340	5.00	19			341	0.10	3		
HCHG	90-NIVA	W	340	1.00	168			341	0.05	58		
	91-NIVA	W	340	1.00	179	2	111	341	0.05	62	5	10
	92-NIVA	W	340	5.00	173	3		341	0.10	137		
	93-NIVA	W	340	5.00	19			341	0.10	3		
	86-NACE	W	510	30.00	31							
	86-SIIF	W						111	3.00	21		
	87-NACE	W	510	40.00a	54	1		111	5.00	21		1
	87-SIIF	W										
	88-NACE	W	510	40.00	60							
	89-NACE	W	510	40.00a	118							
HG	89-SIIF	W						111	50.00	36		
	90-NIVA	W	340	1.00	169	1	9	341	0.05	58		
	90-SIIF	W						111	0.10	41		
	91-NIVA	W	340	1.00	179	3	18	341	0.05	62	5	1
	91-SIIF	W						111	0.30	35		
	92-NIVA	W	340	5.00	173	3		341	0.10	137		
	93-NIVA	W	340	5.00	19			341	0.10	3		
	81-SIIF	1E W	120	10.00	15		1	120	10.00	35		
	82-SIIF	1E W						120	10.00	18		
	82-VETN	W	220	10.00	51			220	10.00	54		
ICDP	83-SIIF	1E W						120	10.00	17		
	83-VETN	1Z W						220	10.00	48		
	84-FIER	1G W						401	10.00	39		
	84-SIIF	1G W						120	10.00	27	6	
	84-VETN	1Z W						220	10.00	66		
	85-SIIF	1G D						120	10.00	30		
	85-VETN	1Z W						220	10.00	90		
	86-NIVA	1H D						310	10.00	49		
	87-FIER	1G W						401	10.00	38		
	87-NIVA	1H D						310	10.00	93		14
MN	88-NIVA	1H D						310	10.00	104		
	89-NIVA	1H D						310	100.00a	159		
	89-NIVA	1H W						310	10.00	36	5	
	90-NIVA	1H W						310	10.00	266		
	91-NIVA	1H W						310	100.00a	264	126	
	92-NIVA	2H W						310	100.00a	283	109	
	93-NIVA	2H W						310	100.00	19	13	
	92-NIVA	W	309	0.10	5		3	309	0.10	43		13
	93-NIVA	W	309	0.10	3			309	0.10	3		3
	NAP	84-SIIF	W						132	40.00	27	
85-SIIF		D						132	40.00	35		
92-NIVA		W	309	0.10	5		5	309	0.10	43		5
93-NIVA		W	309	0.10	3		3	309	0.10	3		
92-NIVA		W	309	0.10	5		1	309	0.10	43		5
93-NIVA		W	309	0.10	3			309	0.10	3		2
92-NIVA		W	309	0.10	5		1	309	0.10	43		5
93-NIVA		W	309	0.10	3			309	0.10	3		1
92-NIVA		W	309	0.10	5		2	309	0.10	43		7
93-NIVA		W	309	0.10	3			309	0.10	3		3
NAP1M	92-NIVA	W	309	0.10	5		2	309	0.10	43		3
	93-NIVA	W	309	0.10	3			309	0.10	3		3
	92-NIVA	W	309	0.10	5		2	309	0.10	43		7
	93-NIVA	W	309	0.10	3			309	0.10	3		3
	92-NIVA	W	309	0.10	5		2	309	0.10	43		11
	93-NIVA	W	309	0.10	3			309	0.10	3		3
	92-NIVA	W	309	0.10	5			130	20.00	12		
	93-NIVA	W	309	0.10	3			312	10.00	6		
	90-NIVA	W	340	2.00	169	31	24	341	0.05	58		1
	91-NIVA	W	340	2.00	179	14	81	341	0.05	62	5	8
OCS	92-NIVA	W	340	5.00	173	3		341	0.10	137		
	93-NIVA	W	340	5.00	19			341	0.10	3		

Method overview

Contaminant, year-laboratory and tissue (cont.)

Tissue	Fish Liver						Fish fillet, Shrimptail, Mussel, Other							
	Contam.	Year & Lab.	Inter-calibr. Basis	Analys Method Code	Detect Limit (ppb)	Total Value Count	Count Below D.Lim	N (<) Above D.Lim	Analys Method Code	Detect Limit (ppb)	Total Value Count	Count Below D.Lim	N (<) Above D.Lim	
PA	92-NIVA	W	309	0.10	5			1	309	0.10	42		5	
	93-NIVA	W	309	0.10	3			2	309	0.10	3		3	
PAH	87-NIVA	W	309	0.02	1									
PAM1	92-NIVA	W	309	0.10	5			1	309	0.10	42		7	
	93-NIVA	W	309	0.10	3				309	0.10	3		3	
PB	83-SIIF	1G W							130	20.00	12			
	84-SIIF	1G W							130	20.00	27		2	
	85-SIIF	1G D							130	20.00	35			
	86-NIVA	12 D	312	150.00	29	4			312	150.00	20			
	87-FIER	1G W	403	10.00	37	1								
	87-NIVA	12 D	312	150.00	59		8		312	150.00	37			
	88-NIVA	12 D	312	150.00	61	14	7		312	150.00	43			
	89-NIVA	12 D	312	150.00	160	12	9							
	89-NIVA	12 W							312	150.00	36			
	90-NIVA	12 W	312	50.00	187	3	1		312	150.00	77		3	
	91-NIVA	12 W	312	50.00	193	14			312	50.00	67			
	92-NIVA	12 W	312	50.00	172	108			312	50.00	111		2	
	93-NIVA	12 W	312	50.00	19	11								
	PCB	81-SIIF	2D W	110	10.00	27				110	10.00	35		
		82-SIIF	2D W							111	5.00	17		
		82-VETN	W	210	50.00	53				211	50.00	54		
		83-SIIF	2E W							111	5.00	14		
83-VETN		2E W							211	50.00	48			
83-VETN		2Z W	210	50.00	48									
84-SIIF		2E W							111	5.00	24			
84-VETN		2E W							211	50.00	66			
84-VETN		2Z W	210	50.00	66									
85-SIIF		2E W							111	5.00	32		6	
85-VETN		2E W							211	50.00	90		1	
85-VETN		2Z W	210	50.00	45									
86-NACE		2Z W	511a	40.00a	31				511	20.00	31			
86-SIIF		2E W							111	5.00	21			
87-NACE		2Z W	510	40.00	54				511	20.00	54			
87-NIVA		W	340	0.10	2									
87-SIIF		2E W							111	5.00	21			
88-NACE		2Z W	510	40.00	60				511	20.00	37			
88-SIIF		2E D							111	5.00	6			
88-SIIF		2E W							111	5.00	22		4	
89-NACE	2Z W	510	40.00a	118				511	20.00	18				
89-SIIF	2E W							111	5.00	36		6		
90-SIIF	2E W							111	5.00	41				
91-SIIF	2E W							111	5.00	35				
PER	92-NIVA	W	309	0.10	5		5	309	0.10	43		31		
	93-NIVA	W	309	0.10	3		3	309	0.10	3		3		
PYR	92-NIVA	W	309	0.10	5			309	0.10	41		6		
	93-NIVA	W	309	0.10	3			309	0.10	3		3		
QCB	90-NIVA	W	340	2.00	169	33	39	341	0.05	58				
	91-NIVA	W	340	2.00	178	13	97	341	0.05	57	5	7		
	92-NIVA	W	340	5.00	173	3		341	0.10	122				
	93-NIVA	W	340	5.00	19			341	0.10	3				
SE	82-VETN	W	240	10.00	46			240	10.00	54				
TDEPP	91-NIVA	W	340	1.00	138		1	341	0.05	62				
	92-NIVA	W	340	5.00	172	3		341	0.10	137				
	93-NIVA	W	340	5.00	19			341	0.10	3				
ZN	83-SIIF	1G W							131	400.00	12			
	84-SIIF	1G W							132	400.00	27			
	85-SIIF	1G D							132	400.00	35			
	86-NIVA	1H D	311	3000.00	29			311	3000.00	20				
	87-FIER	1G W	405	20.00	37									
	87-NIVA	1H D	311	3000.00	59			311	3000.00	37				
	88-NIVA	1H D	311	3000.00	61			311	3000.00	43				
	89-NIVA	1H D	311	3000.00	160		1							
	89-NIVA	1H W						311	3000.00	36				
	90-NIVA	1H W	311	3000.00	189			311	3000.00	77				
	91-NIVA	1H W	311	1000.00	193			311	1000.00	67				
	92-NIVA	1H W	311	1000.00	172			311	1000.00	111				
	93-NIVA	1H W	311	1000.00	19									
Sum of Counts					15765	426	746			11766	400	434		

a(15) > Ambiguous value in cell (Maximum value displayed).

2.4. Abbreviations

Abbreviation ¹	English	Norwegian
ELEMENTS		
Al	aluminium	<i>aluminium</i>
As	arsenic	<i>arsenikk</i>
Cd	cadmium	<i>kadmium</i>
Co	cobalt	<i>kobolt</i>
Cr	chromium	<i>krom</i>
Cu	copper	<i>kobber</i>
Fe	iron	<i>jern</i>
Hg	mercury	<i>kvikksølv</i>
Li	lithium	<i>litium</i>
Mn	manganese	<i>mangan</i>
Ni	nickel	<i>nikkel</i>
Pb	lead	<i>bly</i>
Pb210	lead-210	<i>bly-210</i>
Se	selenium	<i>selen</i>
Ti	titanium	<i>titan</i>
Zn	zinc	<i>sink</i>
PAHs		
PAH	polycyclic aromatic hydrocarbons	<i>polysykliske aromatiske hydrokarboner</i>
ACNE	acenaphthene	<i>acenaften</i>
ACNLE	acenaphthylene	<i>acenaftylen</i>
ANT	anthracene	<i>antracen</i>
BAA	benzo(a)anthracene	<i>benzo(a)antracen</i>
BAP	benzo(a)pyrene	<i>benzo(a)pyren</i>
BBF	benzo(b)fluoranthene	<i>benzo(b)fluoranten</i>
BBKF	benzo(b+k)fluoranthene	<i>benzo(b+k)fluoranten</i>
BEP	benzo(e)pyrene	<i>benzo(e)pyren</i>
BGHIP	benzo(ghi)perylene	<i>benzo(ghi)perylen</i>
BIPN	biphenyl	<i>bifenyl</i>
BJKF	benzo(j,k)fluoranthrene	<i>benzo(j,k)fluorantren</i>
CHR	chrysene	<i>chrysen</i>
COR	coronen	<i>coronen</i>
DBAHA	(see DBA3A)	<i>(se DBA3A)</i>
DBA3A	dibenzo(a,c/a,h)anthracene	<i>dibenzo(a,c/a,h)antracen</i>
DBP	dibenzopyrener	<i>dibenzopyren</i>
DBT	dibenzothiophene	<i>dibenzotiofen</i>
DBTC1	C ₁ -dibenzothiophenes	<i>C₁-dibenzotiofen</i>
DBTC2	C ₂ -dibenzothiophenes	<i>C₂-dibenzotiofen</i>
DBTC3	C ₃ -dibenzothiophenes	<i>C₃-dibenzotiofen</i>
FLE	fluorene	<i>fluoren</i>
FLU	fluoranthene	<i>fluoranten</i>

Abbreviations (cont'd.)

Abbreviation ¹	English	Norwegian
PAHs (cont.)		
ICDP	indeno(1,2,3-cd)pyrene	<i>indeno(1,2,3-cd)pyren</i>
NAMTM	2,3,5-trimethylnaphthalene	<i>2,3,5-trimetylnaftalen</i>
NAP	naphthalene	<i>naftalen</i>
NAP1M	1-methylnaphthalene	<i>1-metylnaftalen</i>
NAP2M	2-methylnaphthalene	<i>2-metylnaftalen</i>
NAPC1	C ₁ -naphthalenes	<i>C₁-naftalen</i>
NAPC2	C ₂ -naphthalenes	<i>C₂-naftalen</i>
NAPC3	C ₃ -naphthalenes	<i>C₃naftalen</i>
NAPDI	2,6-dimethylnaphthalene	<i>2,6-dimetylnaftalen</i>
PA	phenanthrene	<i>fenantren</i>
PAC1	C ₁ -phenanthrenes	<i>C₁-fenantren</i>
PAC2	C ₂ -phenanthrenes	<i>C₂-fenantren</i>
PAM1	1-methylphenanthrene	<i>1-metylfenantren</i>
PER	perylene	<i>perylen</i>
PYR	pyrene	<i>pyren</i>
DI-Σn	sum of "n" dicyclic "PAH"s	<i>sum "n" disykliske "PAH"</i>
P-Σn	sum "n" PAH	<i>sum "n" PAH</i>
PK-Σn	sum carcinogen PAH's	<i>sum kreftfremkallende PAH</i>
PAHΣΣ	DI-Σn + P-Σn etc.	<i>DI-Σ n + P-Σ n mm..</i>
SPA H	= PAHΣΣ	<i>= PAHΣ Σ</i>
PCBs		
PCB	polychlorinated biphenyls	<i>polyklorerte bifenyler</i>
CB	individual chlorobiphenyls (CB)	<i>enkelte klorobifenyl</i>
CB28	CB28 (IUPAC)	<i>CB28 (IUPAC)</i>
CB31	CB31 (IUPAC)	<i>CB31 (IUPAC)</i>
CB44	CB44 (IUPAC)	<i>CB44 (IUPAC)</i>
CB52	CB52 (IUPAC)	<i>CB52 (IUPAC)</i>
CB95	CB95 (IUPAC)	<i>CB95 (IUPAC)</i>
CB101	CB101 (IUPAC)	<i>CB101 (IUPAC)</i>
CB105	CB105 (IUPAC)	<i>CB105 (IUPAC)</i>
CB110	CB110 (IUPAC)	<i>CB110 (IUPAC)</i>
CB118	CB118 (IUPAC)	<i>CB118 (IUPAC)</i>
CB128	CB128 (IUPAC)	<i>CB128 (IUPAC)</i>
CB138	CB138 (IUPAC)	<i>CB138 (IUPAC)</i>
CB149	CB149 (IUPAC)	<i>CB149 (IUPAC)</i>
CB153	CB153 (IUPAC)	<i>CB153 (IUPAC)</i>
CB156	CB156 (IUPAC)	<i>CB156 (IUPAC)</i>

Abbreviations (cont'd.)

Abbreviation ¹	English	Norwegian
PCBs (cont.)		
CB170	CB170 (IUPAC)	<i>CB170 (IUPAC)</i>
CB180	CB180 (IUPAC)	<i>CB180 (IUPAC)</i>
CB194	CB194 (IUPAC)	<i>CB194 (IUPAC)</i>
CB209	CB209 (IUPAC)	<i>CB209 (IUPAC)</i>
CB-Σ7	CB: 28+52+101+118+138+153+180	<i>CB: 28+52+101+118+138+153+180</i>
CB-Σn	sum of CBs, n = number of compounds	<i>sum CBer, n = antall forbindelser</i>
ALD	aldrin	<i>aldrin</i>
DIELD	dieldrin	<i>dieldrin</i>
ENDA	endrin	<i>endrin</i>
CCDAN	cis-chlordane	<i>cis-chlordane</i>
ACDAN	α-chlordane	<i>α -chlordan</i>
GCDAN	γ-chlordane	<i>γ -chlordan</i>
OCDAN	oxy-chlordane	<i>oxy-chlordane</i>
TNONC	trans-nonachlor	<i>trans-nonaklor</i>
TCDAN	trans-chlordane	<i>trans-chlordane</i>
OCS	octachlorostyrene	<i>octaklorstyren</i>
QCB	pentachlorobenzene	<i>pentaklorbenzen</i>
DDD	dichlorodiphenyldichloroethane 1,1-dichloro-2,2-bis-(4-chlorophenyl)ethane	<i>diklordifenyldikloretan</i> <i>1,1-dikloro-2,2-bis-(4-klorofenyl)etan</i>
DDE	dichlorodiphenylethylene (principle metabolite of DDT) 1,1-dichloro-2,2-bis-(4-chlorophenyl)ethylene*	<i>diklordifenyletylen</i> <i>(hovedmetabolitt av DDT)</i> <i>1,1-dikloro-2,2-bis-(4-klorofenyl)etylen</i>
DDT	dichlorodiphenyltrichloroethane 1,1,1-trichloro-2,2-bis-(4-chlorophenyl)ethane	<i>diklordifenyltrikloretan</i> <i>1,1,1-trikloro-2,2-bis-(4-klorofenyl)etan</i>
DDEOP	o,p'-DDE	<i>o,p'-DDE</i>
DDEPP	p,p'-DDE	<i>p,p'-DDE</i>
DDTOP	o,p'-DDT	<i>o,p'-DDT</i>
DDTPP	p,p'-DDT	<i>p,p'-DDT</i>
TDEOP	o,p'-DDD	<i>o,p'-DDD</i>
TDEPP	p,p'-DDD	<i>p,p'-DDD</i>
DDTEP	p,p'-DDE + p,p'-DDT	<i>p,p'-DDE + p,p'-DDT</i>
DD-nΣ	sum of DDT and metabolites, n = number of compounds	<i>sum DDT og metaboliter,</i> <i>n = antall forbindelser</i>

Method overview

Abbreviations (cont'd.)

Abbreviation ¹	English	Norwegian
HCB	hexachlorobenzene	<i>heksaklorbenzen</i>
HCHG	lindane γ HCH = gamma hexachlorocyclohexane (γ BHC = gamma benzenehexachloride, outdated synonym)	<i>lindan</i> γ HCH = <i>gamma heksaklorsyκλοheksan</i> (γ BHC = <i>gamma benzenheksaklorid</i> , <i>foreldret navn</i>)
HCHA	α HCH = alpha HCH	α HCH = <i>alpha HCH</i>
HCHB	β HCH = beta HCH	β HCH = <i>beta HCH</i>
HC-nΣ	sum of HCHs, n = count	<i>sum av HCHs, n = antall</i>
EOCI	extractable organically bound chlorine	<i>ekstraerbart organisk bundet klor</i>
EPOCI	extractable persistent organically bound chlorine	<i>ekstraerbart persistent organisk bundet klor</i>
NTOT	total organic nitrogen	<i>total organisk nitrogen</i>
CORG	organic carbon	<i>organisk karbon</i>
GSAMT	grain size	<i>kornfordeling</i>
MOCON	moisture content	<i>vanninnhold</i>

Abbreviations (cont'd.)

Abbreviation ¹	English	Norwegian
INSTITUTES		
FIER	Institute for Nutrition, Fisheries Directorate	<i>Fiskeridirektoratets Ernæringsinstitutt</i>
FORC	FORCE Institutes, Div. for Isotope Technique and Analysis [DK]	<i>FORCE Institutterne, Div. for Isotopteknik og Analyse [DK]</i>
IMRN	Institute of Marine Research (IMR)	<i>Havforskningsinstituttet</i>
NACE	Nordic Analytical Center	<i>Nordisk Analyse Center</i>
NIVA	Norwegian Institute for Water Research	<i>Norsk institutt for vannforskning</i>
SERI	Swedish Environmental Research Institute	<i>Institutionen för vatten- och luftvårdsforskning</i>
VETN	Norwegian Veterinary Institute	<i>Veterinærinstituttet</i>
SIIF	Fondation for Scientific and Industrial Research at the Norwegian Institute of Technology - SINTEF-SI (previously: Senter for Industrial Research SI)	<i>Stiftelsen for industriell og teknisk forskning ved Norges tekniske høgskole-SINTEF-SI (tidligere: Senter for industriforskning)</i>

1) After: ICES Environmental Data Reporting Formats. International Council for the Exploration of the Sea. January 1992.

*) The Pesticide Index, second edition. The Royal Society of Chemistry, 1991.

2.5. Intercalibration exercises

Sea water:

- 4H - ICES/JMG Fifth Round Intercalibration on Trace Metals in Sea Water - Section 4, analysis for Hg - 1983 - (5/TM/SW:4).
- 4I - JMG Sixth Intercalibration on Trace Metals in Estuarine Waters - 1986 - (6/TM/SW).
- 4Z - Intercalibration exercise for SIIF/SERI (Cd) and NIVA/IAMK (IAMK=Chalmers Inst., Göteborg) - 1985.

Seabed sediment:

- 7E ICES, First Intercalibration Exercise on Trace metals in Marine Sediments - 1984 - (1/TM/MS).
- 8B ICES/OSPARCOM, First Intercomparison Exercise on Organochlorines (individual chlorobiphenyl congeners) in Marine Sediments - Phase 1, analysis of standard solutions - 1989 - (1/OC/MS:1).
- 8C ICES/OSPARCOM, First Intercomparison Exercise on Organochlorines (individual chlorobiphenyl congeners) in Marine Sediments - Phase 2, analysis of standard solutions - 1991 - (1/OC/MS:2).

Marine biota:

- 1E ICES, Fifth Intercalibration Exercise on Trace Metals in Biological Tissues - 1978 - (5/TM/BT).
- 1F ICES, Sixth Intercalibration Exercise on Trace Metals (Cadmium and Lead only) in Biological Tissues - 1979 - (6/TM/BT).
- 1G ICES, Seventh Intercalibration Exercise on Trace Metals in Biological Tissues - Part A - 1983 - (7/TM/BT).
- 1H ICES, Seventh Intercalibration Exercise on Trace Metals in Biological Tissues - Part B - 1985 - (7/TM/BT) (preliminary report 1987).
- 1Z VETN Interlabcalibration exercise with VETN and SIIF 1983, mercury and cadmium in cod filet and liver.
- 1Z NIVA Interlabcalibration exercise with VETN, NACE and NIVA 1986 (Hg, Cd, Cu, Pb and Zn in 6 samples).
- 2D ICES Fourth Intercalibration Exercise on Organochlorines (mainly PCBs) in Biological Tissues (Sample No.5) - 1979 - (4/OC/BT).
- 2E ICES Fifth Intercalibration Exercise on Organochlorines (PCBs only) in Biological Tissues - 1982 - (5/OC/BT).

Method overview

- 2G ICES/IOC Seventh Intercomparison Exercise on Organochlorines (individual chlorobiphenyl congeners) in Biological Tissue - Phase 1, analysis of standard solutions - 1989 - (7/OC7BT:1).
- 2H ICES/IOC Seventh Intercomparison Exercise on Organochlorines (individual chlorobiphenyl congeners) in Biological Tissue - Phase 2 - 1991 - (7/OC7BT:2).
- 2Z VETN Interlabcalibration exercise with VETN among others, 1983, PCB and HCB in cod liver.
- 2Z NACE Interlabcalibration exercise with NACE, VETN and SIIF 1986 (PCB (all labs), DDE, OCS, HCB and DCB (NACE and VETN)).

2.6. Comment on detection limit

The detection limits given here are approximations based on 3 times the standard deviation of the 'blank' or near zero concentration of a solution. However, day-to-day variations in the analytical instrument may lead to different detection limits.

3. Method descriptions

The following descriptions focus on the principles involved and hence are not intended as detailed specifications. The descriptions may vary arbitrarily in detail and may be coupled to specific time periods (cf., Section 2). Hence, they may not necessarily reflect methods currently practiced by the contributing institutes.

3.1. Analyses of sea water

3.1.1. Sampling method codes

code *description*

Samplers for sea water

- | | |
|---|--|
| 1 | Hydrobios hydrographical water sampler |
| 2 | Ruttner industrial water sampler |
| 3 | Ultracleaned polyethylene flask |
| 4 | Ultracleaned glass Erlenmeyer flask |

Sampler deployment for sea water

- | | |
|---|-----------------------------|
| 1 | 2 nylon lines |
| 2 | Nylon line, brass messenger |
| 3 | By plastic-gloved hands |

Methods of pretreatment of sea water samples

- | | |
|---|-----------------|
| 0 | None |
| 1 | Membrane filtre |

Methods of preservation of sea water samples

- | | |
|---|-----------------------------------|
| 0 | None |
| 1 | Nitric acid addition |
| 2 | Freezing |
| 3 | Nitric acid addition and freezing |

Method descriptions

Sea water - inorganic

3.1.2. Inorganic determinations

code *description*

320 **Mercury in sea water (SIIF)**

Reference: S.H. Omang, 1971. Determination of mercury in natural waters and effluents by flameless atomic absorption spectrophotometry. *Anal. Chim. Acta* (1971) 53: 415-420.

322 **Mercury in sea water (SERI)**

Reference: Iverfeldt, Å, 1984. Structural, thermodynamic and kinetic studies of mercury compounds; applications within the environmental mercury cycle. PhD thesis. Department of Inorganic Chemistry, Chalmers University of Technology and University of Göteborg. 48pp. + 7 appendices.

Abstract (Iverfeldt, 1984)

The 0.5-liter glass Erlenmeyer sampling bottles are specially cleaned: first filled with 6M HCl for 7 days, then 7M HNO₃ for 7 days and finally with deionized water with 1ml conc. HNO₃/liter sample for 7 days. The bottles are rinsed extensively between each stage. After sampling, »1ml conc. HNO₃/liter sample is added to achieve a pH of 1-2.

Mercury is preconcentrated on a gold trap after being reduced and volatilized by NaBH₄ from an all quartz glass reduction vessel. Mercury free nitrogen gas is used for purging.

The gold trap is constructed as gold grains in layers separated by SiO₂ glass pieces. All gold traps used are individually calibrated and give the same response.

The gold traps are analysed by a double amalgamation step; i.e. the mercury is transferred by heating, to about 800⁰C, to a second gold trap. This gold trap is analyzed using a helium direct current-plasma emission spectrometer (DCPAES). A Keithley 427 Current Amplifier and a Shimadzu Chromatopac C-R2AX Integrator are used. The system is optimized for the mercury line at 253.65 nm using a Hg(0) diffusion tube.

The stable sensitivity of the DCPAES instrument is ensured by a check before and after every sample determination.

Standard solutions are prepared from commercial stock solutions of CH₃HgCl (1000 ppm, Alfa Products) and mercuric nitrate (1ml = 1mg Hg, BDH Spectrosol).

This combination resulted in the extremely low detection limit of 0.02ng/litre with 5% reproducibility (Cossa & Courau, 1984.)

Reference: Cossa, D, Courau, P., 1984. ICES Fifth round intercalibration for trace metals in seawater (Intercalibrations 5/TM/SW). Report of Section 4 - Round Robin Intercalibration for total Mercury in seawater. International Council for the Exploration of the Seas, May 1984.

Method descriptions

Sea water - inorganic

330 Cadmium in sea water (SIIF)

Reference: P.E. Paus, 1973. Determination of Heavy Metals in Seawater by AAS. J. Anal. Chem. 118-122 (1973).

331 Cadmium, copper, zinc, lead, iron, nickel, cobalt in sea water (NIVA)

The 1-liter polyethylene sampling bottles are specially cleaned: first filled with 6M HCl for 7 days, then 7M HNO₃ for 7 days and finally with deionized water with 1ml conc. HNO₃/liter sample for 7 days. The bottles are rinsed extensively between each stage. After sampling, »1ml conc. HNO₃/liter sample is added to achieve a pH of 1-2.

The analysis uses chelation with APDC (ammonium-pyrrolidine dithio-carbamate) and DDTC (diethylammonium-N,N-diethyl- dithio-carbamate) extraction with freon, reversed extraction back into water and reading using a graphite furnace atomic absorption electrothermal spectrometry or GFAAS.

50-250ml of the acidified sample is transferred to a separatory funnel, buffer is added to pH=4.75 and 20ml of freon. The mixture is vigorously shaken for 120 seconds. After separation of the phases, the organic one is transferred to a 50ml plastic bottle. The extraction is repeated with further 10ml of freon, and the organic phase added to the first 20ml after separation. 0.1ml concentrated nitric acid is added to the organic phase. The bottle is vigorously shaken and let to stand at least 5 minutes. 4.9ml deionized water is added and the solution is shaken; this process is repeated after 30 minutes. More acid and water are used for back-extraction if a greater volume of the solution is necessary for the determination.

Apparatus: Perkin Elmer (P-E) 2380 AAS, HGA 500 (P-E graphite furnace), AS 40 (P-E autosampler), HCL (hollow cathode lamp).

332 Mercury in sea water (NIVA)

Sample-bottle preparation and sampling is the same procedure as 331.

SnCl₂ is added to 250ml of acidified sample. The liberated mercury is driven off with air as carrier gas through a gold trap onto which the mercury is amalgamated. CVAAS (cold-vapour atomic absorption spectrometry) is used to quantify the sample. When all the mercury is trapped, the gold is heated to at least 500⁰C, and the mercury is driven off by the carrier gas into a quartz cell where the atomic absorption signal is measured at 253.7nm. Apparatus: Perkin Elmer (P-E) 300SG AAS converted with gold trap.

Since 1988: a maximum of 100ml sample used, diluted if Hg >50ng/l; P-E 1100 B with gold trap used, helium replaced air as carrier gas and lowest signal was 2.5ng/l.

Method descriptions

Sea bed sediment - sampling

3.2. Analyses of sea bed sediments

3.2.1. Sampling methods

code *description*

Sampling of sediment

GC Reference: Niemistö (1974) gravity corer, inner diameter 50mm. (Niemistö, H., 1974. A gravity corer for studies of soft sediment. Havforskningsinst. Skr. Helsinki, 238:33-38)

GS Grab sampler

Methods of storage/preservation of sea bed sediment samples

01 Frozen (prior to inorganic analyses) and freeze dried (prior to organic analyses)

Methods of grain size analysis of sea bed sediment

01 Dry sieving

Methods of structural analysis of sea bed sediment

01 Visual observation through clear plastic cores

Methods of sea bed sediment extraction

HFO 'Total' digestion with mineral acids including hydrofluoric acid (HF), in open vessels, evaporation of excess HF before analysis.

HNO (outdated code = HNO1 3)
Extraction with 1:1 HNO₃ (suprapur) for inorganic analyses on "fresh" (i.e., frozen) material.

EXN (outdated code = EXN1 2)
Extraction of (organic) contaminants by shaking with non-polar solvents cyclohexane/isopropanol (1:1 v/v) on freeze dried material.

3.2.2. Inorganic determinations

code *description*

350 Mercury in sea bed sediment (NIVA)

Drying procedure

An accurately weighed sample of approximately 1g is dried at 105°C for one hour. The sample is cooled in a desiccator for one hour before weighing. Normally, determinations are on wet samples and the water content is determined of a subsample.

Extraction (oxidation)

Approximately 1g of the sample is accurately weighed in pyrex flasks, 20ml 7N (concentrated) nitric acid (suprapur) is added and the solution heated 120°C for 30min in an autoclave. The solution is transferred to a 100ml volumetric flask and diluted to the mark with deionized water.

Determination

Prior to 1988: Mercury is determined by CVAAS (cold-vapour atomic absorption spectrometry), using the instrument Coleman Model MAS-50. 50ml of the sample solution is transferred to the aeration flask. The lowest signal detectable is corresponding to 0,03µg mercury.

Since 1988: a maximum of 100ml sample used, diluted if Hg >50ng/l; P-E 1100 B with gold trap used, helium replaced air as carrier gas and lowest signal was 2.5ng/l.

351 Chromium, copper, iron, manganese, nikkol and zinc in sea bed sediment (NIVA)

Same procedure as 350: #1-2, Drying and nitric acid Extraction.

Prior to 1992 (1990-91 JMP samples) 'total' extraction (HFO): Approximately 0.1g of the sample is accurately weighed in, 2ml of hydrofluoric acid and 2+2ml of concentrated nitric acid ('aqua regia') is added and the solution heated in a microwave oven. The solution is transferred to a 100ml volumetric flask and diluted to the mark with deionized water.

Since 1992 'total' extraction (HFO): 0.2g of freeze dried homogenated sample is digested in teflon vessels with 1ml 'aqua regia' plus 6ml hydrofluoric acid neutralized with boric acid and diluted to 100ml. (cf., Loring D.H., Rantala, R.T.T., 1992. ICES manual for the geochemical analyses of marine sediment and suspended particulate matter).

Method descriptions

Sea bed sediment - inorganic

Determinations by **flame atomic absorption spectrometry** using acetylene/air flame. Instrument: *Prior to 1986* a Perkin Elmer model 2380 was used and *since 1986* the P-E 560 has been used. For determinations of low concentrations (below detection limits) the flameless method (352) is used. The following are elements often analyzed by flame and their respective detection limits of extract solution:

<u>Element</u>		<u>µg/l</u>
Al	aluminium	1000
Cr	chromium	50
Cu	copper	100
Fe	iron	200
Li	lithium	10
Mn	manganese	50
Ni	nickel	100
Zn	zinc	10

352 Aluminium, cadmium, chromium, cobalt, copper, iron, lead, lithium, manganese, nickel, and zinc in seabed sediment (NIVA)

Same procedure as 350: #1-2, Drying and Extraction.

'Total' extraction (HFO): Approximately 0.1g of the sample is accurately weighed in, 2ml of hydrofluoric acid and 2+2ml of concentrated nitric acid (suprapur) is added and the solution heated in a microwave oven. The solution is transferred to a 100ml volumetric flask and diluted to the mark with deionized water.

Concentrations are determined by **graphite furnace atomic absorption** electrothermal spectrometry or GFAAS using a hollow cathode lamp (HCL) or an electrodeless discharge lamp (EDL) as a light source. *Prior to 1986* a Perkin-Elmer model 560 with HGA-500 graphite furnace was used and *since 1986* the P-E 2380 has been used instead of the P-E 560.

A 20µl portion of extract, treated with HNO₃, is injected into graphite tube. The sample is then heated electrothermally in a stepwise manner through drying, ashing and atomization by a programme designed for each element. The programme which controls the ramp time, holding time and temperature for each phase is often adjusted to achieve optimal results.

Method descriptions

Sea bed sediment - inorganic

The elements analyzed and approximated limit of detection for the extract are:

<u>Element</u>		<u>µg/l</u>
Al	aluminium	5
Cd	cadmium	0.1
Co	cobalt	5
Cr	chromium	0.5
Cu	copper	0.5
Fe	iron	5
Li	lithium	10
Mn	manganese	0.5
Ni	nickel	5
Pb	lead	0.5
Zn	zinc	10

353 Cadmium, chromium, nickel and lead in seabed sediment (NIVA)

As 352 but *since 1992* the L'vov platform technique is used for these metals.

650 Pb-210 dating (FORC)

reference: Pfeiffer Madsen, P., Sørensen, J., 1979. Validation of the Lead-210 dating method. *Journal of Radioanalysis and Chemistry* 54:39-48.

Excerpt (Larsen, B., & Jensen, A., 1989. *Marine Pollution Bulletin* 20(11):556-560.): "The determination of time- dependent sediment parameters is based on the vertical distribution of the natural radioactive isotope lead-210 [= ²¹⁰ Pb] ... The content of unsupported lead-210, that lead-210 not produced in the sediment) decreases regularly downwards in undisturbed and steadily deposited sediment owing to radioactive decay. Departures from this predictable lead-210 profile in the topmost sediment column permit an assessment of mixing and/or intermittent erosion."

Dried slices of sediment are employed.

3.2.3. Organic determinations

code *description*

360 PCB in sea bed sediment (NIVA)

Prior to 1990: the method is quite similar to SIIF method JMG code 110.

Cleaning of chemicals and equipment

The equipment is washed with soap and water, rinsed in first in water, then in distilled water and then with acetone. Finally, the glass equipment is heated to 550°C.

The equipment is washed with soap and water, then rinsed in 1:5 mixture of HNO₃ and H₂SO₄, respectively. Finally, the equipment is rinsed with acetone and cyclohexane.

All solvents are distilled in a special room for this purpose. Distilled water is shaken twice with distilled cyclohexane before use. Sodium sulphate is washed twice with distilled cyclohexane and heated to 550°C.

Extraction

10g freeze dried, homogenized material is extracted twice with a mixture of non-polar solvents cyclohexane/isopropanol (1:1 v/v). The two hour extraction is performed in a continuously shaken Erlenmeyer flask with 200ml solvent mixture. The extraction is repeated with half solvent volume. The two extracts are combined and the cyclohexane phase separated by addition of 150+ml distilled water (excess water relative to isopropanol). The isopropanol/water phase is decanted and the cyclohexane phase washed once with distilled water (several times if also determination of total persistent organic chlorine). The extract is dried over sodium sulphate and then weighed.

Clean-up of extract

2ml cyclohexane extract is shaken vigorously with 2ml concentrated sulphuric acid and then centrifuged. This process is repeated.

Gas chromatographic condition

Carlo Erba 2350 with electron-capture detector (ECD). Splitless injection at 70°C and then programmed temperature raise with 7°/min to 230°C. Column: 30m x 0.25mm (inner diameter), 0.25µm DB-5 fused silica capillary column. Carrier gas: H₂, 0.8 bar.

Identification and quantification

The sample is quantified using 4-5 dominant peaks in the Clophen A60 standard.

Since 1990 the principle is the same but details have been altered as followed:

Method descriptions

Sea bed sediment - organic

Cleaning of chemicals and equipment

The equipment is kept in soap and water overnight then rinsed first in water, then in distilled water and finally with acetone. The air-dried glass equipment is heated to 550°C.

All solvents are either distilled in a special room for this purpose or if the quality is satisfactory commercial solutions are used. Distilled water is shaken twice with distilled cyclohexane before use. Sodium sulphate is washed twice with distilled cyclohexane and heated to 550°C.

Extraction

10g freeze dried, homogenized material is disintegrated/extracted twice with ultrasonic disintegrator with acetone and cyclohexane (1:1).

Clean-up of extract

2ml cyclohexane extract is shaken by whirl mixer with 6ml concentrated sulphuric acid and then centrifuged. This process is repeated.

Gas chromatographic condition

Gas chromatograph HP 5890 II is used with electron-capture detector (ECD). Splitless injection at 90°C and then programmed temperature raise with 3°/min to 280°C. Column: 60mx0.25mm (inner diameter), 0.25µ Rtx 05 fused silica capillary column. Carrier gas: H₂, 36 cm/sec.

Identification and quantification

The individual PCB-congeners are identified by their retention times relative for the internal standard, PCB-53. The PCB's are quantified by using internal standard and eight-level calibration in the concentration range of the CBs in the solution to be analyzed.

369 PAH in sea bed sediment (NIVA)

Extraction

Deuterated internal standards are added to about 0.5-5g of dried sediment and the sample is extracted in Soxhlet with cyclohexane. The extract is clean first by partition with DMF:water, and then by silicagel. Finally, the sample is evaporated to a small volume before GC analysis.

Determination by GC

PAHs are determined on GC coupled to mass-selective detector. The compounds are detected by their respective molecular ions and retention time and quantified by using National Institute for Standards and Technology (NIST) Certified Reference Material (CRM) numbers 1491 and 1941. Coronene and Dibenzopyrene are quantified with the help of in house standards.

Method descriptions

Sea bed sediment - organic

760 PCB in sea bed sediment (IMRN)

PCB in total sediment (50g) were extracted 3 times by acetone and hexane:Acetone (3:1) using repeated ultrasonication and agitation (Jensen et al., 1977).

Sulphur was removed with metallic mercury.

A florisil column (100-230 mesh, 30 cm x 6 mm ID) was used for the separation of the extract into 3 fractions. The first fraction eluted with 2 ml pentane was discarded; the second fraction eluted with 6.5 ml pentane contained PCB, HCB, aldrin, o,p-DDE, p,p-DDE and o,p-DDT; and the third fraction eluted with 10ml pentane:acetone (9:1) contained, alpha-HCH, beta-HCH, gamma-HCH (Lindane), o,p-DDD, p,p-DDD, o,p-DDT (20%) and p,p-DDT.

The third fraction needed further clean up on a neutral alumina column (30 cm x 6 mm ID; deactivated with 6% water). The chlorinated pesticides were eluted with 18 ml pentane. Beta-HCH was not eluted using this method.

A few samples (1990 sediment stations 15S-67S) were cleaned up before separation on the florisil column. A short silica column (10 cm x 6 mm ID) was used, followed by a alumina column (10 cm x 6 mm ID, acidic Al₂O₃). Pentane:dichloromethane (4:1) was used for elution of the compounds.

The chlorinated compounds were quantified on GC (ECD) using two different columns: SE-54 CB, fused silica, 50 m x 0.20 mm, 0.11 µm; SP-2330, fused silica, 60 m x 0.25 mm, 0.20 µm.

Reference: Jensen, S., Renberg, L., Reutergårdh, L., 1977. Residue analysis of sediment and sewage sludge for organochlorines in the presence of elemental sulfur. Anal. Chem. 49:316-318.

769 PAH in sea bed sediment (IMRN)

Ca.50 g of total sediment (< 2mm) were extracted three times with acetone and hexane:acetone (3:1) using ultrasonication and agitation.

The clean-up of the extract was carried out on a short silica column (10 cm x 6 mm ID) using pentane:dichloromethane (9:1) as eluent. GC/MS equipped with a SE-54 fused silica capillary column (50 m x 0.20 mm ID, 0.11 µm film thickness) was used for the analysis of 2-6 ring aromatic hydrocarbons.

3.2.4. Organic carbon determinations

code *description*

390 Total organic nitrogen and organic carbon (CORG) in sea bed sediment (NIVA)

5-8mg of freeze dried sample is weighed in a tin-foiled capsule and heated to over 1800°C in an oven. The carbon in the gas is analyzed in a C-N 1106 Carlo-Erba element analyzer. Detection limit for C is 1 µg/mg and N is 1 µg/mg.

3.3. Analyses of marine biota

3.3.1. Inorganic determinations

code *description*

120 **Mercury in biota (SIIF)**

Representative samples are homogenized in a whirlmixer.

1.0g sample is weighed into a special digestion apparatus with reflux (Bethge apparatus).

10ml conc. HNO₃ and 1ml 47% HBr is added and the solution boiled for approximately 30min. under reflux.

The solution is cooled down to room temperature and diluted to volume into a 50 ml volumetric flask with distilled water.

Mercury is determined with CVAAS (cold-vapour atomic absorption spectrometry). Mercury is reduced with SnCl₂.

121 **Mercury in shellfish (SIIF)**

Same procedure as 120 but bombe digestion (pressurized decomposition) with HNO₃ at 160°C is used instead of pretreatment with HNO₃ and HBr.

130 **Cadmium, lead, copper and nickel in biota (SIIF)**

Representative samples are homogenized in a whirlmixer. 1g freeze dried sample is weighed into a vitrosil vessel and dried at 110°C to constant weight to determine the total water content.

The vessel is then placed in a cold muffle furnace and the temperature increased slowly to 450°C. The vessel is removed from the furnace and cooled down to room temperature. After wetting the ash with 1ml conc. HNO₃ and approximately 2ml H₂O, gentle heating on a hot plate is performed.

The final solution is diluted to volume into a 50 ml volumetric flask with distilled water.

Cadmium is determined after extraction with APDC and MIBK (ammonium-pyrrolidine-dithio-carbamate and methylisobutylketon) with flame atomic absorption spectrometry.

131 **Zinc in biota (SIIF)**

Same procedure as 130 (cadmium) but without extraction with APDC/MIBK.

Method descriptions

Biota - inorganic

132 Zinc and manganese in biota (SIIF)

Same procedure as 131 but quantified by ICP.

220 Mercury and selenium in fish (VETN)

Samples are digested in a mixture of nitric and perchloric acid and the mercury content is determined by CVAAS (cold- vapour atomic absorption spectrometry).

Reference: Haugen, A., Høle, R., and Norheim, G., 1985. Automated hydride generator determination of selenium and mercury in biochemical material. Proc. from 10th. Nordic Atomic Spectroscopy and Trace Element Conference, August 6-9. 1985, Turku, Finland.

Abstract (Haugen et al., 1985)

Tissue samples are digested in a mixture of nitric and perchloric acid in a temperature programmable aluminium block. Maximum temperatures for the mercury and selenium determinations are 180 and 225°C, respectively. After reduction of hexavalent selenium with hydrochloric acid and dilution, the samples are transferred to a programmable sample changer. Both elements are determined with hydride generator producing a continuous, integratable signal. The precision at an absorbance reading of 0.4 is better than 1% and the quantification limit is better than 0.02 µg/g, when using a 1.0g sample. Good agreement was obtained with other methods. Seven determinations of selenium in NBS bovine liver (1577a) gave an average of 0.71µg Se/g, which is equivalent to the certified value.

230 Cadmium in fish (VETN)

Samples are digested by boiling with concentrated nitric acid (Suprapur) during several hours. The metal content is recorded by graphite furnace atomic absorption spectrometry. Quantification is based on standard addition to the digested samples.

240 Selenium in biota (VETN)

Reference: Norheim, G. and Nymoen, U.K., 1981. Fluorimetric determination of selenium in biological material using automatic digestion. Lecture presented at 8th. Nordic trace element and microchemistry conference, Sandefjord, Norway, 10-13, June 1981.

Abstract (Norheim & Nymoen, 1981)

The fluorimetric method is used, employing 2,3- diaminonaphtalene (DAN) as a complexing agent. The method uses 5g of material in an automatic wet digestion procedure with 17ml of 3+7 mixture of perchloric and nitric acid. The solution is heated slowly (225°C) in a thermostatically controlled aluminium block to distil off the nitric acid without charring. After digestion the hexavalent selenium is reduced with hydrochloric acid. EDTA (ethylenediaminetetraacetic acid) is added and aminoacetic acid is used as buffer. The pH is adjusted to 2.4 using a 35cm long electrode.

Method descriptions

Biota - inorganic

DAN is added and the solution is heated at 60°C for 1hr. Finally, the solution is extracted with cyclohexane and the selenium content is measured fluorometrically on a Perkin-Elmer Model 1000 filter instrument. The detection limit is 10ppb wet weight.

310 Mercury in biota (NIVA)

Homogenizing of large samples (eg., fish fillet) by Tedal Quick Foodmaster Holberth silent cutter commercial use. Stainless steel blades are used. For smaller samples (eg., liver) a Silverson 4R Homogenizer is used.

Drying procedure

An accurately weighed sample of approximately 1g is freeze-dried until constant weight. If the sample has excessive fat content (e.g., fish liver, and therefore, can not be freeze-dried) the sample is dried at 105°C for one hour. The samples are cooled in a desiccator for one hour before weighing. Normally mercury is determined on wet samples and the water content is determined of a subsample.

Since 1991 (1990 JMP samples) extracts have been made from wet (fresh) samples.

Extraction (oxidation)

Prior to 1991: 50-200mg freeze-dried sample is weighed in teflon vessels, 2ml concentrated nitric acid (suprapur) is added and capped loosely. The solution is heated for about 2hrs. at 50°C in a thermostatically controlled aluminium block until foaming ceases. The temperature is raised to 110°C and kept there for 6-8hrs. The solution is then cooled. For samples with high fat content (e.g., liver) 2ml of 30% H₂O₂ is added and the solution is heated again to 110°C for 3-4hrs. After cooling the solution is diluted to 25ml. For mercury samples approximately 200mg material is used and the solution is diluted to 100ml.

Since 1991: extracts are made from 0.2-0.5g dried or 2-2.5g wet sample. When teflon vessels are used 10ml concentrated HNO₃ is added and digested for 30 min in autoclave and then diluted to 100ml. If there is excessive fat in sample 2 ml 30% hydrogen peroxide is added.

Determination

Prior to 1988: Mercury is determined by CVAAS (cold-vapour atomic absorption spectrometry), using the instrument Coleman Model MAS-50. 50ml of the sample solution is transferred to the aeration flask. Tin chloride is added as a reducing agent.

Since 1988: a maximum of 30ml sample is used, up to concentrations 1.5µg/l, and diluted if Hg in the solution is more than 1.5µg/l. A P-E 1100 B with gold trap used with helium as carrier gas.

Method descriptions

Biota - inorganic

311 Copper, iron and zinc in biota (NIVA)

(Same homogenizing, drying and extraction procedure as 310.)

Determinations by flame atomic absorption spectrometry using acetylene/air flame. Instrument: *Prior to 1986* a Perkin Elmer model 2380 was used and *since 1986* the P-E 560 has been used. For determinations of low concentrations (below detection limits) the flameless method (352) is used. The following are elements often analyzed by flame and their respective detection limits of extract solution:

Element		$\mu\text{g/l}$
Cu	copper	50
Fe	iron	200
Zn	zinc	10

312 Cadmium, chromium, cobalt, copper, iron, lead, manganese, nickel, and zinc in biota (NIVA)

(Same homogenizing, drying and extraction procedure as 310.)

Determination

Concentrations are determined by graphite furnace atomic absorption electrothermal spectrometry or GFAAS using a hollow cathode lamp (HCL) or an electrodeless discharge lamp (EDL) as a light source. *Prior to 1986* a Perkin-Elmer model 560 with HGA-500 graphite furnace was used and *since 1986* the P-E 2380 has been used instead of the P-E 560.

A 20 μl portion of extract, treated with HNO_3 , is injected into graphite tube. The sample is then heated electrothermally in a stepwise manner through drying, ashing and atomization by a programme designed for each element. The programme which controls the ramp time, holding time and temperature for each phase is often adjusted to achieve optimal results.

Since 1992 the L'vov platform techniques has been used for determination of cadmium and lead

The elements analyzed and approximated limit of detection for the extract are:

Element		$\mu\text{g/l}$
Al	aluminium	5
Cd	cadmium	0.1
Co	cobalt	5
Cr	chromium	0.5
Cu	copper	0.5
Fe	iron	5
Mn	manganese	0.5
Ni	nickel	5
Pb	lead	0.5
Zn	zinc	10

Method descriptions

Biota - inorganic

401 Mercury in biota (FIER)

Reference: Eliann E. & Julshamn, K., 1978. A method for the determination of selenium and mercury in fish products using the same digestion procedure. Atomic Absorption Newsletter, 17(6):135-138 (November - December 1978).

Reference: Julshamn, K., Ringdal, O., & Braekkan, O.R., 1982. Mercury concentration in liver and muscle of cod (*Gadus morhua*) as an evidence of migration between waters with different levels of mercury. Bull. Environm. Contam. Toxicol. 29:544-549 (1982).

Abstract (Eliann & Julshamn, 1978; Julshamn et al., 1982)

Representative samples are homogenized in a whirlmixer. About 1g of sample tissue is dried at 95°C for 24 hrs. 0.5g sample is weighed into a special digestion apparatus with reflux (Bethge apparatus).

10 ml conc. HNO₃/H₂SO₄ (1+1) + V₂O₅ (0.1% w/v) are added. The solution is boiled for approximately 30 min under reflux.

The solution is cooled down to room temperature and diluted to volume into a 50ml volumetric flask with distilled water and MnO₄.

Mercury is determined by CVAAS (cold-vapour atomic absorption spectrometry), EDL (electrodeless discharge lamp), wavelength = 253.6nm, Perkin Elmer (P-E) 370 AAS, P-E mercury analysis system (303-0830 and 303-0832). Mercury is reduced with SnCl₂ to avoid interference with iodine.

Quantification is based on standard curves. Detection limit: dry weight sample. 5 ng/g.

402 Cadmium in biota (FIER)

Reference: Julshamn, K. & Brækkan, O.R., 1975. Determination of trace elements in fish tissues by the standard addition method. Atomic Absorption Newsletter, 11(3):49-52. (May-June 1975). (Concerns: Mn, Fe, Cu, Zn, Cd and Pb).

Reference: Julshamn, K., 1977. Inhibition of response by perchloric acid in flameless atomic absorption. Atomic Absorption Newsletter, 16(6):149-150 (November - December 1977). (Concerns: Mg, Al, Cr, Mn, Fe, Co, Ni, Cu, Zn, Cd and Pb).

Reference: Julshamn, K. & Andersen, K.-J., 1983. Subcellular distribution of major and minor elements in unexposed molluscs in Western Norway - II. The distribution and binding of cadmium, zinc, copper, magnesium, manganese and iron in the kidney and the digestive system of the common mussel *Mytilus edulis*. Comp. Biochem. Physiol. 75A(1):13-16 (1983). Pergamon Press.

Abstract (Julshamn & Brækkan, 1975; Julshamn, 1977; Julshamn & Andersen, 1983)

Representative samples are homogenized in a whirlmixer. 1g of sample tissue is dried at 95°C for 24hr. 0.1g sample is weighed into a Sovirel test-tube (20ml). 2ml conc. HNO₃/HClO₄ (9:1) is added and is boiled under pressure.

Method descriptions

Biota - inorganic

The solution is cooled down to room temperature and diluted to volume into a 10ml volumetric flask with distilled water.

Concentrations are determined by graphite furnace atomic absorption electrothermal spectrometry or GFAAS, hollow cathode lamp (HCL), Perkin Elmer (P-E) 5000 AAS, HGA 500 (P-E graphite furnace), AS 50 (P-E autosampler) and wavelength = 228.8nm. Matrix modification reagent is $(\text{NH}_4)_2\text{HPO}_4$.

Quantification is based on standard addition to the digested samples: Amount added in the analyses: Cd 1ng/ml. Detection limit for dry tissue sample: 0.5ng/g.

403 Lead in biota (FIER)

(Same procedure as 402: par.#1 and #3-5).

Concentrations are determined by graphite furnace atomic absorption electrothermal spectrometry or GFAAS, EDL (electrodeless discharge lamp) Perkin Elmer (P-E) 5000 AAS, HGA 500 (P-E graphite furnace), AS 50 (P-E autosampler) and wavelength = 283.3nm. Matrix modification reagent is $(\text{NH}_4)_2\text{HPO}_4$.

Quantification is based on standard addition to the digested samples: amount added in the analysis Pb 10ng/ml. Detection limit for wet tissue sample: 10ng/g.

404 Copper in biota (FIER)

with high Cu content (> 1.5 ppm d.w.)

(Same procedure as 402: par.#1-5).

Concentrations are determined by flame AAS (atomic absorption spectrometry), HCL (hollow-cathode lamp, Perkin Elmer (P-E) 370 AAS and wavelength = 324.7nm without background correction.

with low Cu content (< 1.5 ppm d.w.)

(Same procedure as 402: par.#1-5).

Concentrations are determined by graphite furnace atomic absorption electrothermal spectrometry or GFAAS, HCL (hollow cathode lamp, Perkin Elmer (P-E) 5000 AAS, HGA 500 (P-E graphite furnace), AS 50 (P-E autosampler) and wavelength = 324.7nm. No matrix modification reagent is used.

Quantification is based on standard addition to the digested samples: 10 ng/ml.

Method descriptions

Biota - inorganic

405 Zn in biota (FIER)

(Same procedure as 402: par.#1-5).

Concentrations are determined by flame AAS (atomic absorption spectrometry), HCL (hollow-cathode lamp, Perkin Elmer (P-E) 370 AAS and wavelength = 213.9 nm without background correction.

Quantification is based on standard curves.

406 Arsenic in biota (FIER)

(Same procedure as 402: par.#4-5).

Concentrations are determined by graphite furnace atomic absorption electrothermal spectrometry or GFAAS, HCL (hollow- cathode lamp) Perkin Elmer (P-E) 5000 AAS, HGA 500 (P-E graphite furnace) with a conventional tube without platform, AS 50 (P-E autosampler) and wavelength = 193.7 nm. Matrix modification reagent is a Ni-solution.

Method descriptions

Biota - organic

3.3.2. Organic determinations

<i>code</i>	<i>description</i>
110	PCB HCB DDTEP (p,p'DDE + p,p'DDT) HCHG (gHCH = gBHC) in fish and shellfish (SIIF)

Cleaning of chemicals and equipment

All solvents are distilled in a special room for this purpose. Distilled water is shaken twice with distilled cyclohexane before use. Sodium sulphate is washed twice with distilled cyclohexane and heated in the same way as the glass equipment.

Extraction

10g wet, homogenized material is extracted twice with a mixture of cyclohexane/isopropanol (1:1 v/v). The two hour extraction is performed in a continuously shaken Erlenmeyer flask with 200 ml solvent mixture. The extraction is repeated with half solvent volume. The two extracts are combined and the cyclohexane phase separated by addition of 150+ ml distilled water (excess water relative to isopropanol). The isopropanol/water phase is decanted and the cyclohexane phase washed once with distilled water (several times if also determination of total persistent organic chlorine). The extract is dried over sodium sulphate and then weighed.

Determination of fat

A part of the cleaned and dried cyclohexane extract is evaporated in an oven to constant weight at 100°C.

Alternatively, the extract is evaporated to constant weight in a flask with Vigreux column in oil bath at 1 atm., 110°C and reflux or in vacuum, 50°C, reflux and weak N₂-stream.

The results are corrected for loss of cyclohexane. Cyclohexane output is calculated as sum cyclohexane extracts from first and second extraction (grams) minus fat in the same extract.

The precision of the fat determinations is roughly ±10%. Determinations are given to 0.1% fat. The two methods give comparable results.

Clean-up of extract

0.2g fat is dissolved in 2ml cyclohexane, shaken vigorously with 2ml concentrated sulphuric acid and then centrifuged. For further clean-up about 1ml of the sulphuric acid treated extract is treated with a solution of KOH.

Gas chromatographic condition

Hewlett-Packard 5730 A with ECD. Splitless injection at 60°C and then programmed temperature raise with 8°/min to 230°C. Column: 50m x 0.3mm (inner diameter), 0.15µ SE-54 glass capillary column. Carrier gas: He, 20 psi.

Method descriptions

Biota - organic

Identification and quantification

By comparing the whole pattern with various commercial standard mixtures, it was found that Clophen A60 was in best accordance with the sample types. Some of the isomers in the PCB pattern were selected by comparison with standards of specific isomers and these were used for quantification:

SIIF code	CB code	Structure (-biphenyl)	name
3	95	2,3,6,2',5'	Pentachlorine
4	101	2,4,5,2',5'	Pentachlorine
9/	149/	2,4,5,2',3',6'/	Hexachlorine
10	118	2,4,5,3',4,	Pentachlorine
14	138	2,3,4,2',4',5'	Hexachlorine
15	128	2,3,4,2',3',4'	Hexachlorine
16	180	2,3,4,5,2',4',5'	Heptachlorine

By the GC conditions used it was not possible to separate isomers 9 and 10.

HCB, HCHG (gHCH=gBHC) and DDTEP (p,p'DDE + p,p'DDT) is determined by multi-level calibration curve. HCHG is identified and quantified by the breakdown product of HCHG (three trichlorobenzene peaks) after treatment with sodium hydroxide (NaOH)

111 PCB HCB DDTEP (p,p'DDE + p,p'DDT) HCHG (gHCH = gBHC) in fish and shellfish (SIIF)

Same procedure as 110, except that the organochlorine standard was Arochlor 1254 instead of Clophen A60 which is used earlier. The detection limit given for 1982 JMP data for this method is erroneously low. Corrected limits are given in the 1983 JMP data submitted

The detection limit is dependant on sample quantity. For example the detection limit for HCHG is approximately: 0.1ppb with 10g dry weight material, 0.03ppb with 25g and 0.01 with 80g.

Since 1991 CB204 has been used as an internal standard.

210 PCB, HCB, DDEPP in fish liver samples (VETN)

References: Bjerk, J.E., & Sundby, R., 1970. Residues of organochlorine insecticides and polychlorinated biphenys in terrestrial and aquatic test-organisms. Norwegian Part of OECD program 1967-68. This was modified by Norheim, G., 1978. The composition and distribution of PCB in arctic fox (Alopex lagopus) caught near Longyearbyen on Svalbard. Acta pharmacol. et toxicol. 42:7-13.

Abstract (Norheim, 1978)

Extraction: 0.5g of sample is homogenized in a mortar with 2.5g of anhydrous sodium sulphate and 2.5g purified sand and allowed to stand overnight in the dark. After being mixed with 2g magnesium sulphate, the dry powder is transferred to a short chromatographic column (20mmx10cm) equipped with ground glass stoppers and elutriated with 2x10ml diethyl ether. The column is carefully rotated to release air bubbles and the ether is allowed to stand for 2hrs in the column before elutriation. The ether is evaporated in centrifuge tubes and the residue dissolved in 1.0ml n-heptane. The extract is finally treated with 2.0ml concentrated sulphuric acid for about 1 hour. 5 μ l n-heptane is injected into the gas chromatograph.

Quantification: Phenoclor DP6 is used as standard. The peak height of 2,4,5-2',4',5' hexachlorbiphenyl is used to quantify PCB.

211 PCB in fish filet samples (VETN)

Reference: Norheim, G., Økland, E.M., 1980. Rapid extraction of some persistent chlorinated hydrocarbons from biological material with low fat content. Analyst (Okt.1980)105:990-992.

Abstract (Norheim & Økland, 1980)

Apparatus: A Carlo Erba 2100 gas chromatograph equipped with a nickel-63 electron-capture detector and a 2m x 3mm (inner diameter) glass column is used. The column material is 1.5% SP-2250 - 15.9% SP-2401 on 100-120-mesh Supelcon AW DMCS. The column, injector and detector temperatures are 200, 250 and 275⁰C, respectively. Argon-methane (95+5) is used as the carrier gas, the flow-rate being 55ml·min⁻¹. The electrometer attenuation is x128.

Reagents: Sulphuric acid, 95-97%. Pro analysi grade (Merck). Heptane. Pro analysi grade (Merck). Hexachlorobenzene. Pract. grade (Fluka). Octachlorostyrene. Obtained as a gift from Norsk Hydro.

Standard solutions: Amounts of 100mg each of hexachlorobenzene and octachlorostyrene are dissolved in 100ml of heptane and the mixture is diluted 1 + 50 000 with heptane.

Procedure: A 0.5g amount of sample is accurately weighed into a 10ml Soveril glass tube fitted with a screw-cap, and 6ml of concentrated sulphuric acid are measured into the tube. The tube is placed in a thermostatically controlled oven at 60°C for 4hr, during which time it is shaken lightly a few times to ensure complete solubilisation of the sample. After cooling, 1.0ml of heptane is pipetted into the tube, the screw-cap put on and the tube shaken for about 3min. Finally, the tube is centrifuged with the screw-cap on, after which the sample is ready for gas chromatography. An injection volume of 5 μ l is used.

Method descriptions

Biota - organic

Quantification: The same standard and isomer as in 210 are used to quantify the sample.

309 PAH in biota (NIVA)

Extraction

Deuterated internal standards are added to about 20-30g of homogenized wet sample and the sample is then saponified with KOH/methanol. After filtering through a glass filter the solution is extracted with n-pentan. The extract is clean first by partition with DMF:water, and then by silicagel. An aliquot of the homogenized sample is used for dry weight determination.

Determination by GC

PAHs are determined on GC coupled to mass-selective detector. The compounds are detected by their respective molecular ions and retention time and quantified by using National Institute for Standards and Technology (NIST) Certified Reference Material (CRM) number 1491 and number 1974 for blue mussel samples. Coronen and Dibenzopyrener are quantified with the help of in house standards.

340 PCB, QCB, OCS, HCB, DDTEP (p,p'DDE + p,p'DDT), HCHB, (b HCH), HCHG (g HCH = g BHC) in fish liver (NIVA)

Prior to 1991 (1987 JMP NIVA samples): Equivalent to SIIF method 111 with the following exception:

Determination of fat

The extract is evaporated to constant weight in a flask with Vigreux column in oil bath at 1 atm., 110°C and reflux or in vacuum, 50°C, reflux and weak N₂-stream.

The results are corrected for loss of cyclohexane. Cyclohexane output is calculated as sum cyclohexane extracts from first and second extraction (grams) minus fat in the same extract.

Since 1991 (1990 JMP samples):

Cleaning of chemicals and equipment

The equipment is kept in soap (3% RBS/Deconex) and water overnight then rinsed first in water, then in distilled water and finally with acetone. The air-dried glass equipment is heated to 550°C.

All solvents are distilled in a special room for this purpose or if the quality is satisfactory commercial solutions are used. Sodium sulphate is washed twice with distilled cyclohexane and heated to 550°C.

Method descriptions

Biota - organic

Extraction

About 5g (dependant on species/tissue) of wet homogenized material is extracted twice by ultra sonic desintegration with a mixture of cyclohexane/acetone (1:1 v/v). *Since June 1991* PCB 53 has been added as internal standard and sodium sulphate step has been omitted.

Determination of fat

A part of the cleaned and dried cyclohexane extract is evaporated in an oven to constant weight at 105°C.

The precision of the fat determinations is roughly $\pm 10\%$. Determinations are given at 0.1%. The two methods give comparable results.

Clean-up of extract

About 0.1g fat is dissolved in 2ml cyclohexane, shaken with 6ml concentrated sulphuric acid and then centrifuged. For further clean-up about 1ml of the sulphuric acid treated extract may be treated with a solution of KOH:

Gas chromatographic condition

Since 1992 the following has been used: HP 5890 series II with ECD; splitless injection at 90°C and then programmed temperature raise with 3°/min. to 280°C; column: 60m x 0.25mm (inner diameter), 0.25µ Rtx 05 silica capillary column; carrier gas: H at 37 cm/s.

Identification and quantification

HCB, HCHG (γ HCH= γ BHC) and DDTEP (p,p'DDE + p,p'DDT) and the individual PCB-congeners are identified by their retention times relative for the internal standard, PCB-53. The PCB's are quantified by using internal standard and eight-level calibration in the concentration range of the CHC's in the solution to be analyzed.

341 PCB, QCB, OCS, HCB, DDTEP (p,p'DDE + p,p'DDT), HCHB, (b HCH), HCHG (g HCH = g BHC) in shellfish and fish fillet (NIVA)

Same procedure as 340: except for detection limits:

Detection limits: the minimum detectable amount corresponds to 0.001µg/g wet weight for liver samples. This gives minimum quantification limits of 0.002µg/g.

510 PCB, HCB, DDEPP (p,p'DDE), DDTTP (p,p'DDT), HCHG (g HCH = g BHC) in fish liver (NACE)

Pretreatment and fat determinations: Samples are homogenized in a Waring blender. Homogenised liver samples are ground in a mortar with sea sand and anhydrous sodium sulphate and allowed to stand overnight. The samples are mixed with magnesium-

Method descriptions

Biota - organic

sulphate, transferred to a glass column with sintered glass frit and extracted with diethyl ether. The ether is collected in pre-weighed tubes, evaporated and the amount of fat determined by weighing.

The fat extract is dissolved in hexane for pesticide analyses and treated with concentrated sulphuric acid with gentle agitation. After centrifugation the hexane phase is used for gas chromatography (GC) analysis for pesticides and PCBs. An aliquot of the hexane phase is also treated with sodium alcoholate to convert p,p'DDT to p,p'DDE for the determination of DDT by the increase in DDE.

Moisture content: samples are dried in an oven overnight (16hr) at 105⁰C, equilibrated in a desiccator for 1hr and re-weighed. Filet samples are also dried for 72hr at 45°C for later determination of mercury.

GC analyses: a Perkin Elmer 8500 GC equipped with an auto-sampler and an electron capture detector (Ni-63) and connected to a 7500 computer with Chrom 3 software is used. The column is a glass 2mx1/4", 2mm (inner diameter) packed with 1.5% SP-2250/1.95% SP-2400 on Suplecoport 100/120. The carrier gas is argon with 5% methane at a flow rate of 40ml/min. The oven temperature is 210⁰C, with the injector at 250°C and the detector at 300°C. The amount of sample injected is 2µl and the analysis takes 40min.

Reference standards: commercially available Aroclor 1242, Aroclor 1254, Aroclor 1260 and Supelco's CP pesticide mix are used in addition to a special mixture containing Phenoclor DPG (60% chlorination), hexachlorobenzene (HCB), octachlorostyrene (OCS), p,p'-DDE and decachlorobiphenyl (DCB).

Quantification: response factors are calculated from the integrated areas for each component and the amount injected. The corresponding peaks for the samples are integrated and the concentrations calculated from the area and the response factor. A simplified method for the calculation of the concentrations of PCBs is used. This is based on using the area for the peak in the Phenoclor standard corresponding to 2,4,5,2',4',5'-hexachlorobiphenyl and the total amount of PCB components injected. The concentrations for samples are calculated from the area of the peak corresponding to that used in the standard. This requires that the pattern of PCB components in the sample corresponds to that of the Phenoclor standard.

Detection limits: the minimum detectable amount corresponds to 0.01µg/g wet weight for liver samples. This gives minimum quantification limits of 0.04µg/g for PCB.

Method descriptions

Biota - organic

511 PCB in fish filet (NACE)

Pretreatment: Homogenised filet samples are treated with concentrated sulphuric acid for 4hr at 60°C and PCBs extracted with hexane. After centrifugation the hexane phase is used for gas chromatography (GC) analysis.

(Same procedure as 510: par.#3-6).

Detection limits: the minium detectable amount corresponds to 0.005µg/g wet weight for liver samples. This gives minimum quantification limits of 0.02µg/g.

605 EPOCl in shellfish (SIIF)

The cyclohexane extract from chlorinated hydrocarbon analysis is reduced in volume (by evaporation) and treated with concentrated H₂SO₄ until the extract is clear. An aliquot is sent to the Institue for Energy Technology (Kjeller, Norway) to be exposed to neutron bombardment in a JEEP II atomic reactor. The radioactivity of the persistant chlorine isotope is measured and quantified against a complete procedural blank.

The detection limit is 5 ppb wet weight.

610 EPOCl in fish liver (NACE)

Same procedure as 605 but higher detection limit.

The detection limit is 800 ppb wet weight.

615 EPOCl in fish liver (NIVA)

Same procedure as 605.

The detection limit is 40 ppb wet weight.

3.3.3. Fat determinations

<i>code</i>	<i>description</i>
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A	Weight of extracted solids during chlorinated hydrocarbon determinations: procedure 110 (SIIF).
B	Weight of extracted solids during chlorinated hydrocarbon determinations: procedure 510 (NACE and VETN).
C	Weight of extracted solids from freeze dried material using ethyl acetate (FIER).
D	Weight of extracted solids during chlorinated hydrocarbon determinations: procedure 340 (NIVA).



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