# Monitoring of POPs in the ambient air



Jana Klánová

# The objectives of the POPs Global Monitoring Plan

To evaluate whether the POPs actually were reduced or eliminated as requested in Articles 3 and 5 of the Convention, information on environmental levels of the chemicals listed in the annexes should enable detection of *trends* overtime.

Therefore focus is upon monitoring of background levels of POPs at locations not influenced by local sources.

Reliable identification of trends will require that statistical evaluation is carried out on the design of each national monitoring programme contributing to the Global Monitoring Plan, to ensure that it is powerful enough to detect trends in time.

# Air sampling

Air is a key medium - responds quickly to sources

Air concentrations fluctuate widely in the space and time

Various concentrations in the gas/particulate phases - compromise over the sample time/volume/technique

Short-term sampling/bulking

# Air sampling

- ambient air permanent gases
  - volatile/semivolatile compounds
  - particules
- indoor air
- working environment
- emissions
- imissions

# Sampling and sampling preparation methodology

Experimental design

Sample matrices - choice of equipment

All sampling procedures have to be agreed upon and documented

Representativeness and integrity of the sample during the entire sampling process has to be assured

Quality requirements (equipment, transportation, pre-analytical treatment, storage, GPS referencing, standardization, documentation)

Personal protection, waste management

Sample handling

# High volume samplers for active POPs sampling





# **Dust aerosols samplers**





# Dust aerosols samplers

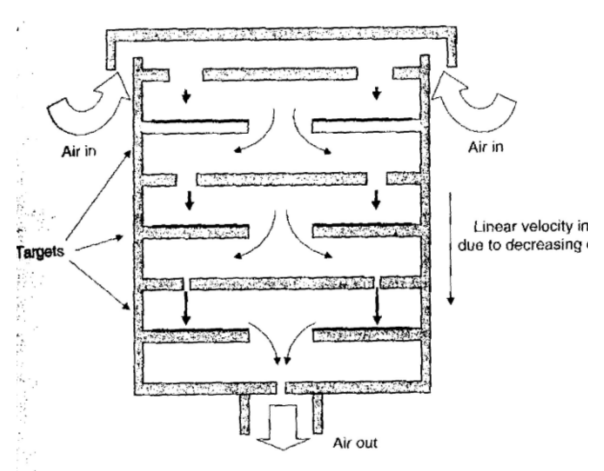
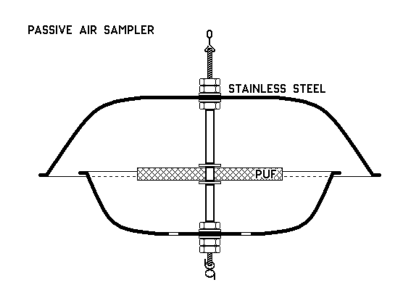


Figure 7.3 Schematic showing the operation of a cascade impactor.

# Passive Air Samplers





- filter (polyurethane foam) captures pollutants from the surrounding air
- sampler body filter protecting chamber (wind, rainwater, solar radiation)

# Analytical methodology

Extraction and clean-up

POPs analysis

Organization of quality control

Data treatment

# Basic requirements

Competence for infrastructure, instrumentation, and trained staff

- Validation of the analytical methods including in-house methods
- Standard operating procedures (SOPs) for the validated methods
- Quality criteria for quality assurance and quality control (QA/QC)

## Recommended analytes

#### Chemical Parent POPs

Aldrin
Chlordane *cis*- and *trans*-chlordane

DDT 4,4'-DDT, 2,4'-DDT

Dieldrin

Endrin

**HCB** 

Heptachlor

Mirex

Polychlorinated PCBs

PCB with TEFs\*

Polychlorinated 2,3,7,8-PCDD/PCDF

Toxaphene Congeners

#### Transformation products

(cis- and trans-nonachlor, oxychlordane)

(4,4'-DDE, 2,4'-DDE, 4,4'-DDD, 2,4'-DDD)

(Heptachlor, heptachlorepoxide)

(7 congeners: 28, 52, 101, 118, 138, 153, and 180)

(12 congeners: 77, 81, 105, 114, 118, 123, 126,

156, 157, 167, 169, and 189)

(17 congeners)

P26, P50, P62

### Extraction

The appropriately prepared sample can be extracted by any of a number of techniques (Soxhlet, automatic extraction, pressurized extractions, MAE, SFE).

The main points to consider are to allow adequate time of exposure of the solvent system in the sample matrix.

Cross contamination from residues left behind by high levels of POPs in other samples is a concern at this stage and equipment must be thoroughly cleaned and checked from batch to batch.

Purity of extraction solvents is also a major consideration. Only high purity glass distilled solvents should be used.

Standards should be added to the sample as early as possible in the process.

If the results are reported on a lipid weight basis, the determination of the lipid content in the sample is critical.

### **Standards**

should be added before extraction to control the extraction efficiency (the recoveries differ with POP and matrix, for PCB and pesticides: 80 % - 120 %, for tetra- and penta-chlorinated PCB, recoveries down to 60 % can be accepted, for PCDD/PCDF: 50 % - 130 % (for hepta- and octa-chlorinated PCDD/PCDF 40 % - 150 % can be accepted).

For PCB analysis and ECD detection, a minimum of two standards - one eluting at the beginning and one at the end of the chromatogram – should be used.

Concentration (evaporation under vacuum or with nitrogen – control of temperature, flow of nitrogen, drying of the extract should be avoided - keeper)

Elimination of water (sodium sulphate), lipids (sulphuric acid or permeation in gels), proteins (denaturation with oxalate), and sulfur (activated copper)

Clean-up is performed with various combinations of adsorbents and solvents depending on selectivity, conditioning and column flow in order to remove interfering substances/materials from the analyte. Sample clean-up procedures should be efficient to prevent contamination of the detector.

## Separation of POPs

is conducted using gas chromatography with electron capture detector (ECD), mass selective detector (MS detector) or, if available, high-resolution mass spectrometry (HRMS).

Other separation techniques, such as high pressure liquid chromatography (HPLC), have not been found adequate.

An appropriate stationary phase has to be selected and enough peak separation must be achieved to allow accurate quantification (capillary columns lengths of 30-60 m, internal diameters of 0.15-0.25 mm, a film thickness of 0.1-0.3 µm, helium or hydrogen as a carrier gas, cleanliness of injector - deactivated glass insert).

Verification of chromatographic conditions include resolution, symmetric peak shape, reproducibility of retention times; verification of the linear range of the instrument.

### Identification

The information available to identify the compounds eluted from the gas Chromatographic column depends on the type of detector being used.

Retention time should match between sample and internal standard;

Confirmation of peaks can be performed on a second column with different polarity;

Matrix spikes are recommended to verify components;

For GC-MS detection combinations, positive identification should be done on isotopic ratios within 20 % of theoretical value;

The retention time of the labeled internal standard to the native compound should be within 3 seconds;

The use of MS libraries is useful if using full scan.

## Quantification

In general, quantification of the analyte should be done according to the internal standard methodology;

At least one standard representative for the POPs analyte group analyzed should be added at the normal level of quantification; Verification that the concentration of blanks is significantly lower than the samples (recommendation < 10%).

#### Calibration:

Multi-point calibrations should be carried out, labeled internal standards are an added value;

Daily calibration checks in connection with analyzing a series of samples should be done;

Suitable laboratory reference material should be used to verify the performance. The maintenance of the analytical equipment is considered as one of the most important aspects in POPs analysis.

## Quality assurance and quality control

are important factors in sampling and analysis;

Internal and recovery stds have to be used;

Any method performance must be verified through control tables where optimal operational ranges are defined;

Periodical analysis of blank samples and certified reference materials;

Own laboratory reference materials, and blind or divided samples should be included in routine QA/QC;

The inter-calibration exercises are an essential component in quality assurance of the results and are deemed indispensable in the implementation of a regional laboratory network;

A recommendation would be that at least once a year such an intercalibration study is performed for each matrix and persistent organic pollutant of interest to the Region.

#### Data treatment

There are a number of parameters that have to be reported together with the analytical results.

These include the efficiency of the extraction and clean-up, and the blank values, but the results should not be compensated for these parameters.

The uncertainty of the results should also be at least estimated, but preferably determined, using results from inter- or intralaboratory comparisons.

## **Terminology**

Calibration

Linearity

Sensitivity and specificity

Robustnes and repeatibility

Limit of detection and quantification

Accuracy, trueness, precision

Standard deviation 
$$\sigma^2 = \frac{\sum R_i^2}{2n}$$
 RSD =  $\frac{100 \sigma}{X}$ 

Regulation diagram – Warning interval 
$$X \pm 2 \sigma$$
 (95.5% results)

Regulation interval  $X \pm 3 \sigma$  (99.7% results)

Confidence interval 
$$R = 2.8 \sigma$$

$$R = 2,8 \sigma$$

**Z-score** 
$$Z = \sigma$$
  $-2 < z < 2$ 

$$-2 < z < 2$$

RSZ- rescaled sum of z-scores RSZ = 
$$\frac{\sum z}{z}$$

$$\mathbf{RSZ} = \frac{\Sigma \mathbf{Z}}{\sqrt{\mathsf{m}}}$$

SSZ - sum of squared z-scores 
$$SSZ = \Sigma z^2$$

$$SSZ = \Sigma z^2$$

## LOD, LOQ

The lowest concentration at which a compound can be detected (limit of detection, LOD) is defined as that corresponding to a signal three times the noise.

The lowest concentration that can quantitatively be determined (limit of quantification LOQ) is three times higher than LOD.

Compounds found at levels between LOD and LOQ can be reported as present, or possibly as being present at an estimated concentration, but in the latter case the result has to be clearly marked as being below LOQ ("LOD-LOQ"), data below LOD as "<LOD";

Results for sum parameters where one or several individual compounds are <LOQ should be reported as intervals with a lower bound limit calculated with the <LOQ set to 0, and the upper bond limit with <LOQ set equal to LOQ.

There are, however, several statistical techniques for treating censored data when the true detection limit is known, e.g. by using a robust statistic such as the median which is unaffected by small numbers reported as below LOD. <"LOD" can be substituted by ½ LOD

#### Data treatment

There are a number of parameters that have to be reported together with the analytical results. These include the efficiency of the extraction and clean-up, and the blank values.

Recovery efficiency should be reported but reporting values should not be corrected for percentage of recovery;

It should be demonstrated that the blank is 10-times lower than the value that is reported but reporting values should not be corrected by laboratory blanks;

The uncertainty of the results should also be at least estimated, but preferably determined, using results from inter- or intralaboratory comparisons.

There are two methods available to provide information on uncertainty:

- Quantification of uncertainty for each step;
- Overall uncertainty derived from inter- and intra-laboratory results.

# **Terminology**

Primary GMP data: are the results of measurements made on samples GMP meta-data: are any other data or information that describe the primary GMP data in some way. This can include information on the methodologies employed (e.g., for sampling and analysis) and the laboratories responsible for a particular set of analyses.

Supplementary data: Are any other data or information that may be accepted for use in the Stockholm Convention evaluation process. This might include relevant information and/or data from published sources (e.g. the peer reviewed scientific literature, existing assessment, etc), results of modelling activities.

*Un-aggregated data:* individual sample measurement values (concentration of PCB153 in the liver tissue of a specific individual fish, sampled at location x at time y).

Aggregated data: (statistically) summarised data, e.g. averaged values that summarise the measurements on a number of individual samples.

The main goal of the Global Monitoring Plan data strategy is to compile unaggregated - primary GMP data.

Un-aggregated data permit data to be treated in a transparent and consistent manner.

If these methodologies are modified or further developed at some point in the future, the availability of un-aggregated – primary GMP data provides the best possibilities for re-calculation or for repeating previous data treatment.

To the greatest extent possible, data should be reported for individual compounds or congeners or isomers.

Data on contaminant concentrations should be reported together with a clear indication of both the units and the basis of determination (wet weight, lipid weight).

Mokrá - půdy 2002 - 4														vyhodnoceno: 25.4.2003		
							Koncentr	ace ng/g								
Číslo vzorku	toluen	02-753	02-752	02-740	02-741	02-742	02-743	02-744	02-745	02-746	02-747	02-748	02-749	02-750	02-751	
Lokalita	GC blank	Lab. blank	RM	454	Čihálky	332	Velká	Velká	Prostřed	420Vel	Chlumek	Chlumek	Horák	Nové pole	jižní CVM	LOQ
				Hosten		Vodojem	Bata1	Bata2	kopec	Bata	1	2	mysl.			
Číslo zadava				303S	304S	305S	306S	307S	308S	309S	310S	311S	312S	313S	314S	
Datum odbě				14.11.02	14.11.02	14.11.02	14.11.02	14.11.02	14.11.02	14.11.02	14.11.02	14.11.02	14.11.02	14.11.02	14.11.02	KALIB30
Naváž ka (g)	5,0	5,0	5,0	5,0	5,0	5,0	5,0	5,0	5,0	5,0	5,0	5,0	5,0	5,0	5,0	5,0
Ředění	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1
Naftalen	0,10	1,86	26,74	12,5	6,6	7,5	5,2	5,5	11,8	13,5	7,1	6,6	8,6	5,9	8,5	0,10
Acenaftyle	-	0,02	0,58	0,8	0,3	0,7	0,4	0,5	2,2	1,8	0,6	0,5	1,2	2,4	0,8	0,10
Acenaften	-	0,04	1,22	1,4	0,3	1,4	1,6	0,6	5,3	3,4	2,5	0,8	2,0	5,4	1,2	0,10
Fluoren	-	0,04	2,26	1,7	0,6	1,4	1,3	0,7	4,9	3,8	2,0	1,0	2,2	4,7	1,5	0,10
Fenantren	-	0,12	23,96	24,9	6,4	20,5	18,8	8,4	69,1	59,4	14,2	13,6	29,5	109,3	16,8	0,10
Antracen	-	-	1,12	2,0	0,4	1,9	3,4	1,1	6,1	5,2	2,1	1,4	2,9	16,9	1,8	0,10
Fluoranten	-	-	27,78	68,2	13,7	58,0	42,0	24,2	213,0	162,5	40,7	37,6	82,5	450,2	42,9	0,10
Pyren	-	-	19,38	50,5	9,7	45,6	35,4	20,2	159,3	123,6	32,0	28,6	63,8	377,2	33,0	0,10
Benz(a)ant	-	-	4,60	17,9	2,9	14,4	14,7	9,1	61,5	49,3	18,3	13,1	26,3	206,3	13,6	0,10
Chrysen	-	-	11,50	32,4	7,3	25,6	18,4	12,2	102,6	75,9	22,3	16,8	41,2	204,2	20,0	0,10
Benzo(b)flu	-	-	18,30	61,0	11,7	32,2	23,6	20,4	169,5	128,2	28,0	29,4	67,7	261,1	31,2	0,10
Benzo(k)flu	-	-	6,04	18,1	3,8	14,4	11,0	7,9	56,4	41,9	13,0	11,2	22,4	134,8	11,6	0,10
Benzo(a)p	-	-	8,34	27,6	3,5	23,6	20,3	13,3	92,8	71,6	24,2	18,4	38,4	285,9	21,3	0,10
Indeno(123	-	-	8,22	33,1	6,4	21,4	14,8	11,1	98,7	72,0	22,6	19,6	41,0	216,1	20,7	0,10
Dibenz(ah)	-	-	0,82	2,7	0,6	2,4	1,6	0,9	7,1	8,3	1,8	2,3	4,1	25,8	1,8	0,10
Benzo(ghi)	-	-	11,26	29,7	5,3	20,6	14,8	11,4	83,9	61,4	19,4	16,3	36,0	181,8	18,5	0,10
Suma PAI	0,10	2,08	172,12	384,5	79,5	291,6	227,3	147,5	1144,2	881,8	250,8	217,2	469,8	2488,0	245,2	1,60
100% D-P/	2 000	2 000	2 000	2 000	2 000	2 000	2 000	2 000	2 000	2 000	2 000	2 000	2 000	2 000	2 000	
ředění	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	
D8-naftaler	0%	0%	88%	72%	79%	66%	65%	80%	62%	66%	21%	61%	75%	81%	81%	
D10-fenant	0%	0%	90%	77%	91%	68%	72%	86%	77%	79%	88%	79%	85%	94%	92%	
D12-peryle	0%	0%	86%	74%	34%	67%	73%	86%	83%	83%	89%	82%	93%	101%	96%	

GC blank ..... slepý vzorek přístroje GC-MS - nástřik čistého rozpouštědla do plynového chromatografu

Lab. blank ..... laboratorní slepý vzorek - analyzovaný celým analytickým postupem s čistými rozpouštědly a všemi použitými materiály

GPC blank ..... slepý vzorek GPC chromatografu

blank, GF blank ..... terénní slepé vzorky - pasivní odběr na polyuretanovou pěnu a skleněné vlákno

CRM ..... analýza certifikovaného referenčního materiálu

RM ..... analýza laboratorního referenčního materiálu

NQ ..... nekvantifikováno - analyt byl překryt interferentem

LOQ ..... meze stanovitelnosti

# Regional monitoring reports

Introduction, background

Description of the region

Regional strategy for information gathering

Arrangements to address global and regional transport

Methodology for sampling, analysis and handling of data

Results

Summary of findings and discussion

# GMP should in each region strive for at least

Three to five stations with active high-volume sampling

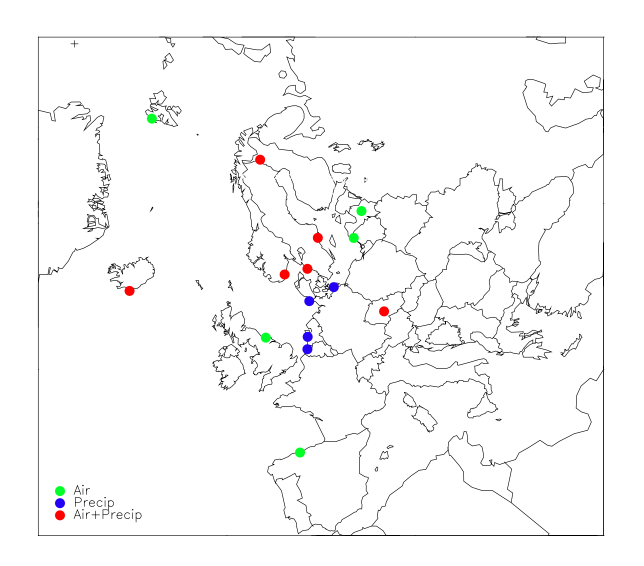
A network of 10 to 15 passive sampling stations arranged in a grid with spacing of approximately  $200 \times 200$  km for enhancing geographical coverage .

Passive samplers should be co-located at the high volume sites for comparison purposes.

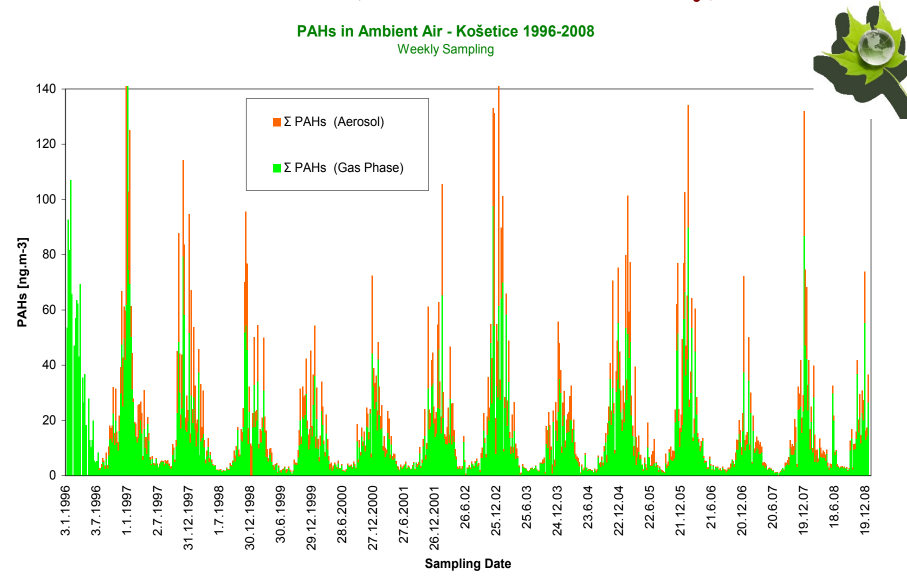
Cumulative sampling (for 1 to 2 days every week or continuously over periods of 1 to 2 weeks) by active high volume sampling (~0.5-1 m3/min. flow rate) at a few sites in each region. These samples would be separated into particulate and gaseous fractions.

Continuous, cumulative passive (diffusive) sampling for integration periods of 3 months to 1 year using passive samplers deployed at a large number of sites, including the high volume sampling sites.

# POP monitoring EMEP network, 2000



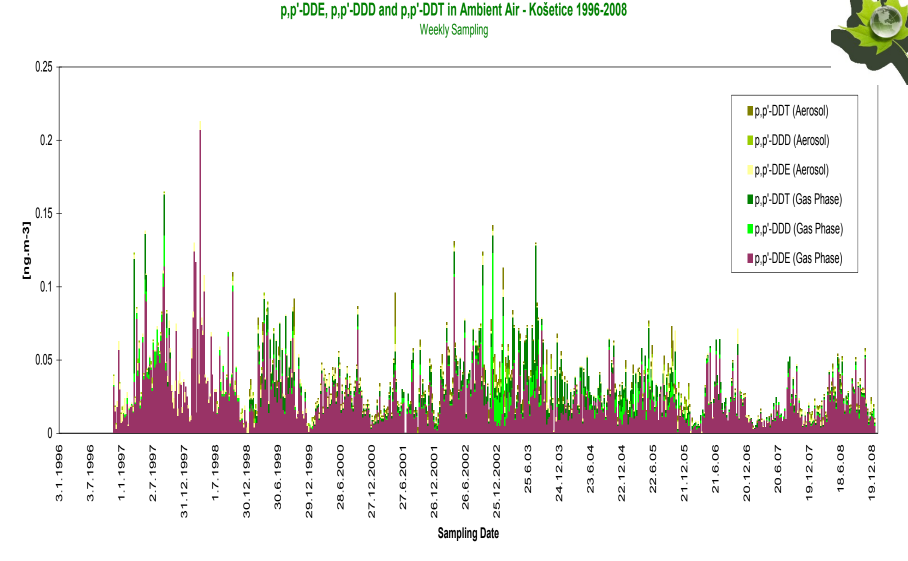
# PAHs in the ambient air, Kosetice observatory, 1996-2008



Holoubek, I., Klánová, J., Jarkovský, J., Kohoutek, J.: Trends in background levels of persistent organic pollutants at Kosetice observatory,

Czech Republic. Part I. Ambient air and wet deposition 1988-2005. Journal of Environmental Monitoring 9 (6), 557 – 563 (2007)

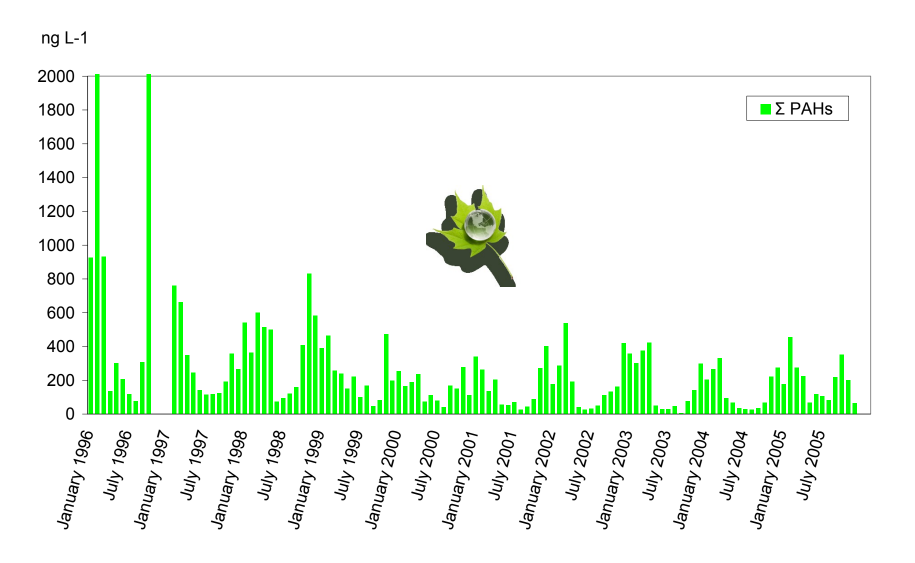
# DDTs in the ambient air, Kosetice observatory, 1996-2008



Holoubek, I., Klánová, J., Jarkovský, J., Kohoutek, J.: Trends in background levels of persistent organic pollutants at Kosetice observatory,

Czech Republic. Part I. Ambient air and wet deposition 1988-2005. Journal of Environmental Monitoring 9 (6), 557 – 563 (2007)

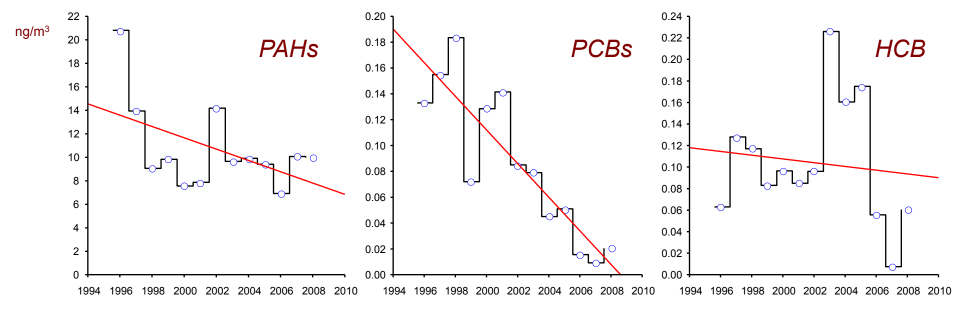
## PAHs in rain (monthly means), Kosetice, 1996-08

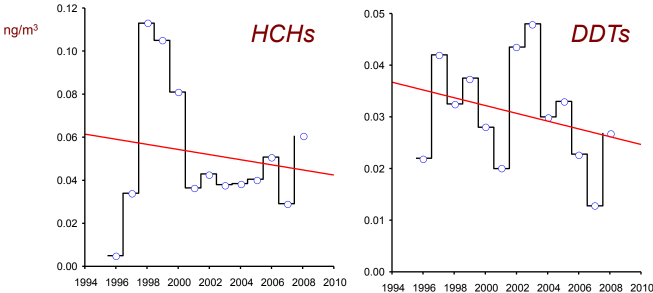


Holoubek, I., Klánová, J., Jarkovský, J., Kohoutek, J.: Trends in background levels of persistent organic pollutants at Kosetice observatory,

Czech Republic. Part I. Ambient air and wet deposition 1988-2005. *Journal of Environmental Monitoring 9 (6)*, 557 – 563 (2007)

# Time related trends of POPs in the air, gas and particle phase.





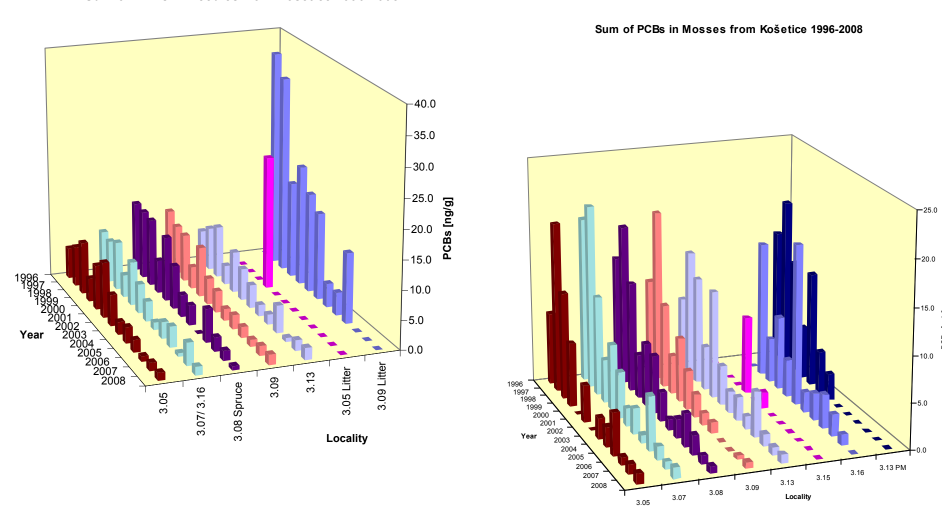
Holoubek, I., Klánová, J., Jarkovský, J., Kohoutek, J.: Trends in background levels of persistent organic pollutants at Kosetice observatory, Czech Republic. Part I. Ambient air and wet deposition 1988-2005. *Journal of Environmental Monitoring* 9 (6), 557 – 563 (2007)

Map of the sampling sites



# Spatial and temporal variations of PCB concentrations in needles and moss, Kosetice observatory, 1996-2005

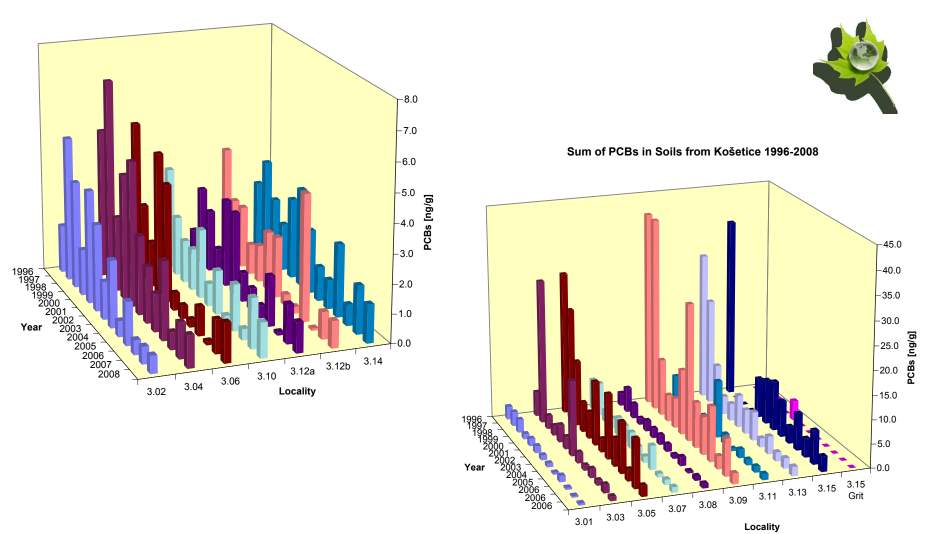
Sum of PCBs in Needles from Košetice 1996-2008



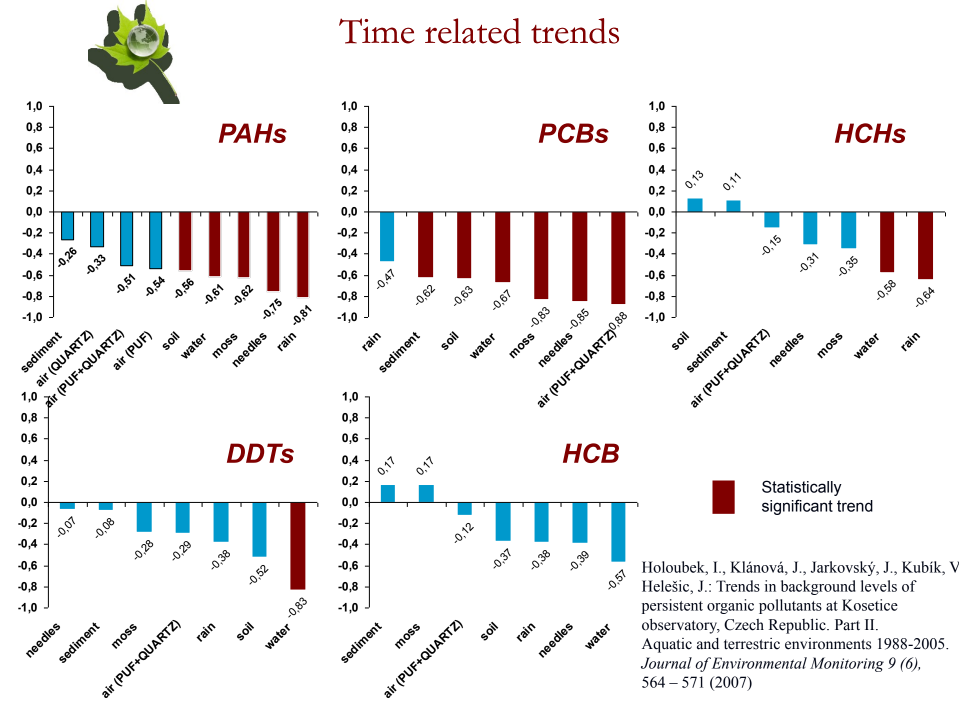
Holoubek, I., Klánová, J., Jarkovský, J., Kubík, V., Helešic, J.: Trends in background levels of persistent organic pollutants at Kosetice observatory, Czech Republic. Part II. Aquatic and terrestric environments 1988-2005. *Journal of Environmental Monitoring 9 (6)*, 564 – 571 (2007)

### Spatial and temporal variations of PCB concentrations in soil and sediment, Kosetice observatory, 1996-2005

Sum of PCBs in Sediments from Košetice 1996-2008

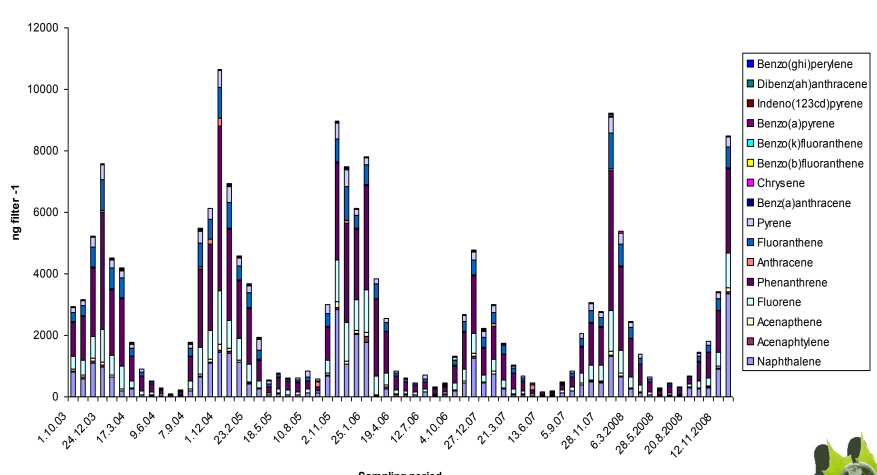


Holoubek, I., Klánová, J., Jarkovský, J., Kubík, V., Helešic, J.: Trends in background levels of persistent organic pollutants at Kosetice observatory, Czech Republic. Part II. Aquatic and terrestric environments 1988-2005. *Journal of Environmental Monitoring 9 (6)*, 564 – 571 (2007)



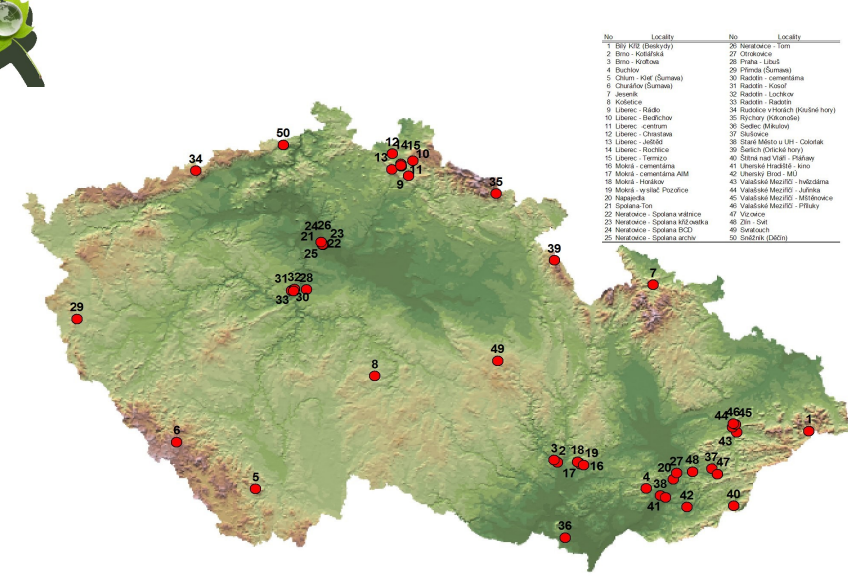
### Seasonal variation of PAH concentrations in the ambient air, Košetice observatory, 2003-2006

PAS - PAHs



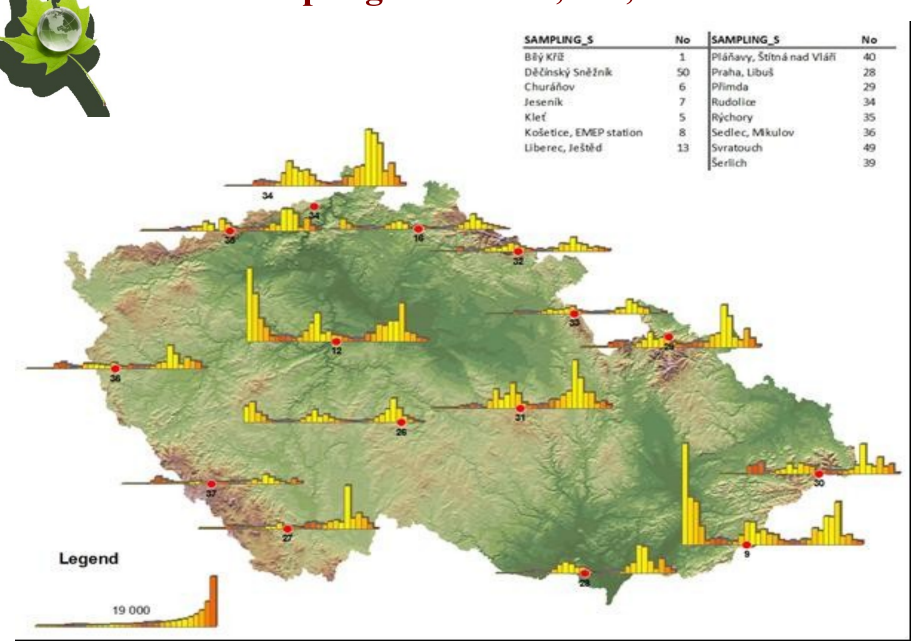
Sampling period

#### Passive sampling 2006-2008, CZ



Klanova, J., Cupr, P., Borůvková, J., Kohoutek, J., Kareš, R., Přibylová, P., Prokeš, R., Holoubek, I.: Application of passive sampler for monitoring of POPs in ambient air. IV. Model monitoring network in the Czech Republic (MONET\_CZ 2007), 2008. Masaryk Univerzity, Brno, Czech Republic. ISBN 978-80-210-4696-2

#### Passive sampling 2006-2008, CZ, PAHs



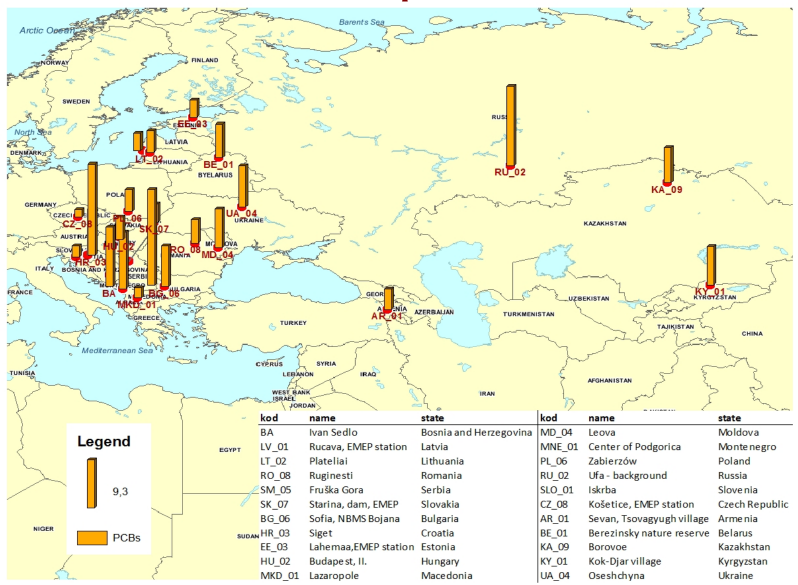
#### Regional strategy for information gathering

A regional organisation group will be established in each region to be responsible for implementing the global guidance document and the Global Monitoring Plan implementation plan within that region, taking into account regional realities.

The duties of the regional organisation groups would include

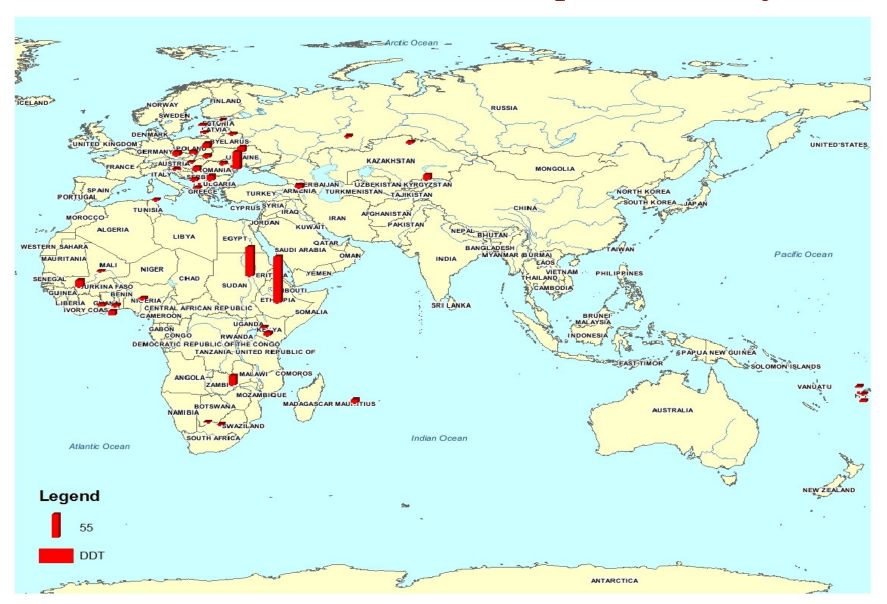
- identifying where existing suitable monitoring data are and are not available,
- developing a regional strategy for implementation of the GMP,
- establishing regional, sub-regional and inter-regional monitoring networks,
- coordinating sampling and analytical arrangements,
- ensuring compliance with protocols for QA/QC,
- data archiving and accessibility,
- maintaining the interaction with other regional organization groups,
- developing elements to encourage capacity building,
- preparing regional reports

#### Central and Eastern European Network 2006-2008



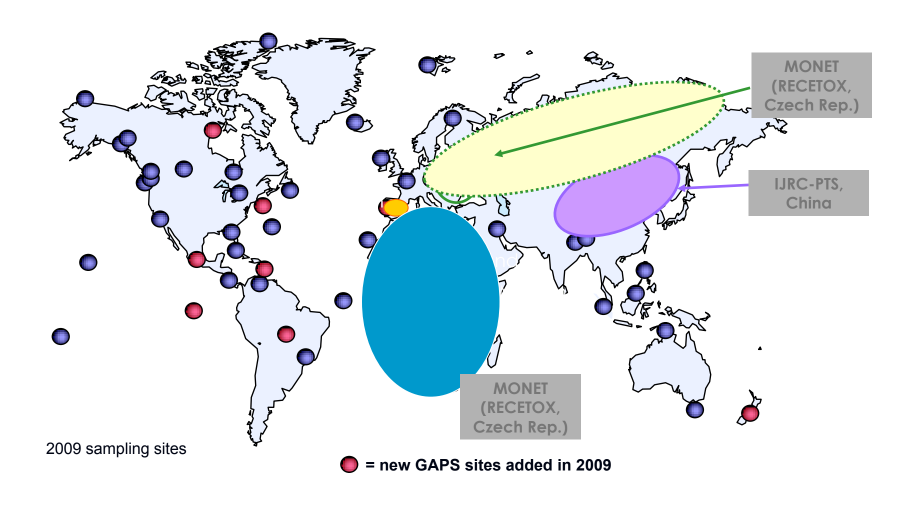
Klanova, J., Cupr, P., Holoubek, I., Borůvková, J., Přibylová, P., Kareš, R., Kohoutek, J.: Application of passive sampler for monitoring of POPs in ambient air. V. Pilot study for development of the monitoring network in the Central and Eastern Europe (MONET\_CEEC 2007), 2008. Masarykova Univerzita, Brno, Czech Republic. ISBN 978-80-210-4697-9

#### Central and Eastern Europe, Africa, Fiji



Klanova, J., Cupr, P., Holoubek, I., Borůvková, J., Přibylová, P., Kareš, R., Kohoutek, J., Dvorská, A,.Komprda, J.: Towards the global monitoring of POPs), 2009. Masarykova Univerzita, Brno, Czech Republic. ISBN 978-80-210-4853-9

### Global Passive Air Sampling Programs



## GMP Report on the first phase of the Effectiveness Evaluation of the Stockholm Convention: Conclusions and Recommendations.

#### Long-range transport

- The coordination group concluded that knowledge on long-range transport was key to assessing changes in POPs levels over time and to assess the effectiveness of the Convention.
- A plan or process to develop a coordinated cross-regional approach to assess long-range transport is needed.
- Long-range transport is also linked to variable climate and meteorology.

# GMP Report on the first phase of the Effectiveness Evaluation of the Stockholm Convention: Conclusions and Recommendations.

#### Comparability issues

- To interpret POPs concentrations in air, it was required that programs would remain consistent in their methods over time and thus ensure that the data collected within a program remained comparable and thus suitable for assessing changes in levels over time.
- It was noted that it would be very difficult to achieve comparability between various programs due to many sources of variability including the use of different laboratories, different sampling methods or analytical protocols.

# GMP Report on the first phase of the Effectiveness Evaluation of the Stockholm Convention: Conclusions and Recommendations.

#### New POPs

When the Conference of the Parties decides to add new substances to the Annexes of the Convention, it would be necessary to include those new POPs into the effectiveness evaluations and monitoring of those {new} POPs would have to be initiated as soon as possible so that effective baselines could be established. The inclusion of additional POPs would be likely to require modification or amendments to the current guidelines for global monitoring and the implementation plan. Meeting such requirements would undoubtedly increase the cost of the monitoring programs.

### Future of Global Air Monitoring" – Workshop Summary (TF HTAP Workshop)

March 31st, 2009, St. Petersburg, Russia.

The timing of the workshop coincided well with the recently completed first implementation of the Global Monitoring Plan, a mechanism for the effectiveness evaluation (EE) of the SC. Expert opinion from this group on several issues that were agreed to be relevant to the future of global air monitoring was summarized.

- 1. Data comparability issues.
- 2. Particle-bound compounds.
- 3. New POPs.
- 4. LRT, climate variability and meteorological variability.
- 5. Existing and new air programs.
- 6. Data Availability.

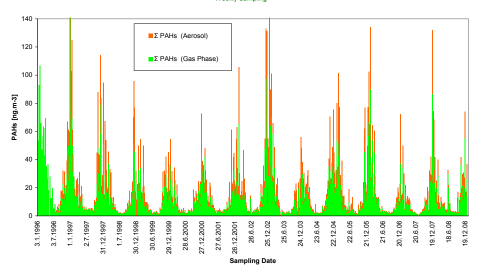


#### Data comparability issues.

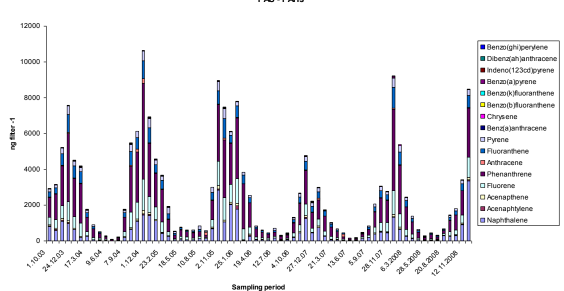
- within-program comparability is crucial for producing reliable temporal trends .
- intercomparison exercises and improving comparability between programs (overlaps)
- passive vs active air sampling (co-location of passive/active at one site in each region)
- selection of common/comparable reporting format

#### Temporal variations of PAH concentrations in air, Kosetice observatory, 1996-2005





#### PAS - PAHs



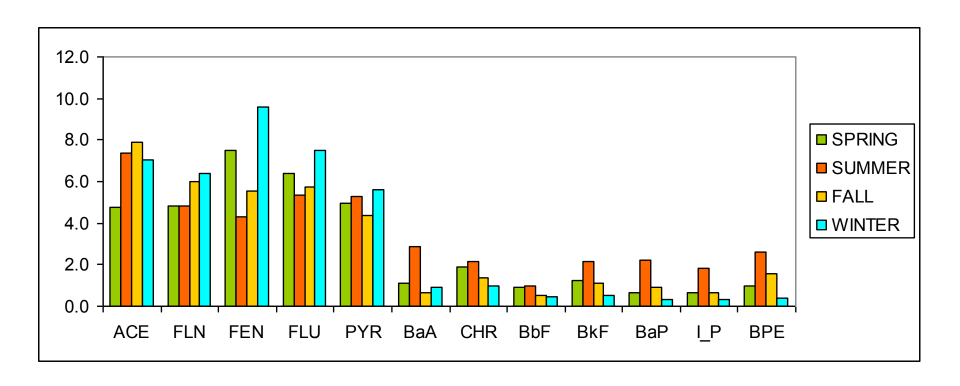


#### Particle-bound compounds

Difficulties exist in interpreting passive sampler data for chemicals that partition between gas and particles (e.g. high molecular weight components of: PAHs, PCDD/Fs, and BFRs). So far, the air monitoring of such chemicals requires active air sampling techniques.

Although some groups have begun to investigate particle-phase sampling by PAS, this is an area that requires further study.

## Seasonal variability of the sampling rates (m<sup>3</sup> day<sup>-1</sup>) for selected compounds in 5 years.



Klanova, J., Čupr, P., Kohoutek, J., Harner, T., 2008. Assessing the influence of meteorological parameters on the performance of polyurethane foam-based passive air samplers. ES&T 42, 550-555.



#### **New POPs**

It was recognized that in the near future new POPs maybe added to the existing POPs listed under the SC on POPs and CLRTAP POPs protocol. This will add to air monitoring obligations and challenges of existing programs.

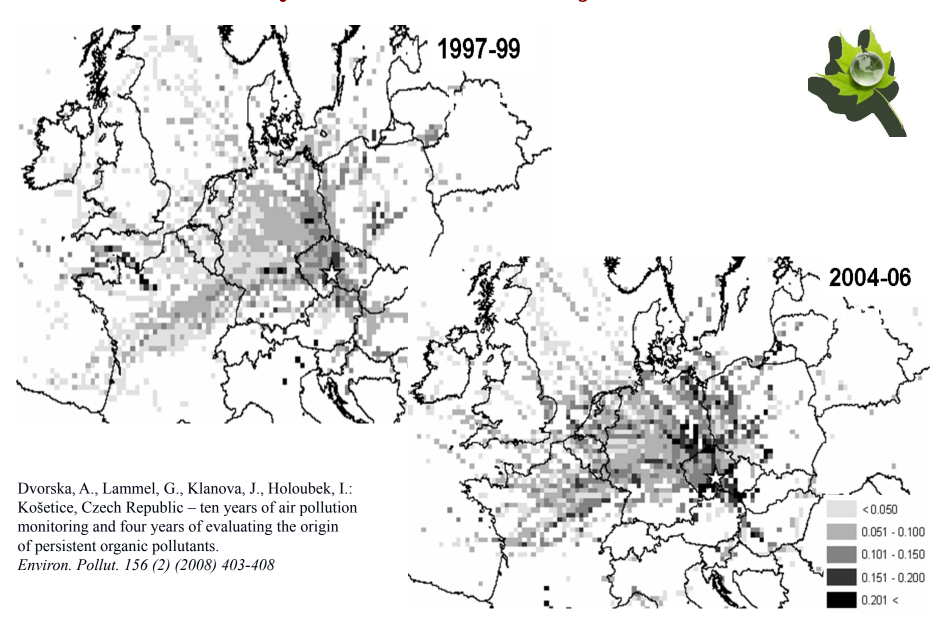


## Long range transport, climate variability and meteorological variability

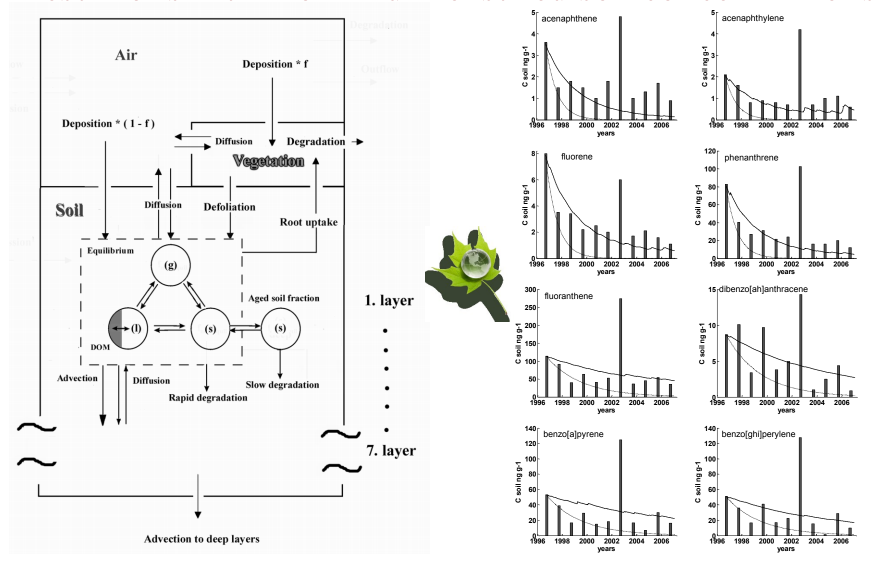
Understanding data is more important than just reporting levels and trends, especially for the purpose of assessing effectiveness of regulatory efforts on POPs.

It is important to develop tools (e.g. back trajectory techniques, multimedia and transport models and investigations of meteorological and climate variability) to better interpret monitoring data.

#### Analysis of the back trajectories

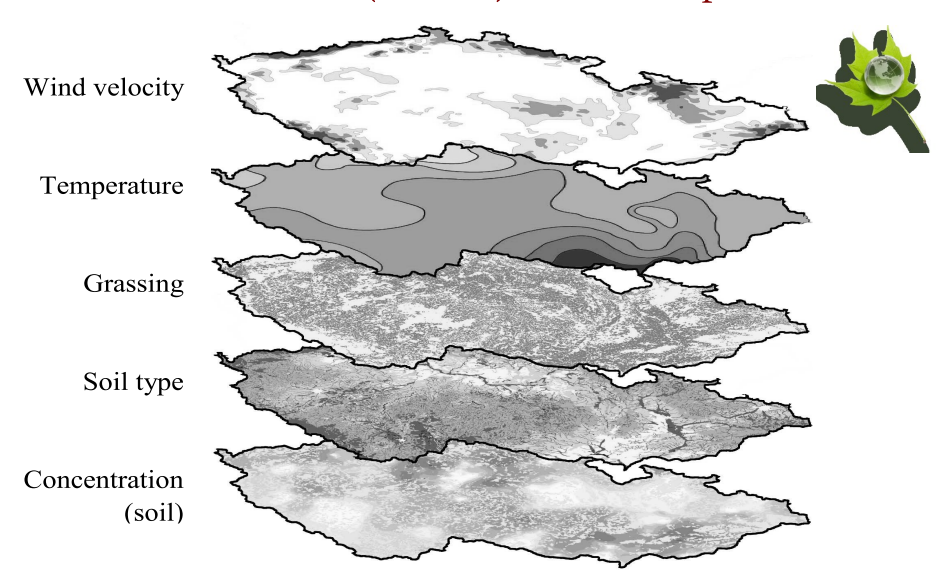


### Long term temporal trends of soil concentrations result of simulation and measured soil concentrations



Komprda, J., Kubošová, K., Dvorská, A., Scheringer, M., Klánová, J, and Holoubek, I.: Application of an unsteady state environmental distribution model to a decadal time series of PAH concentrations in the Central Europe. *Journal of Environmental Monitoring*, 2009 (doi: 10.1039/B815719G).

#### GIS based models (1x1 km) of various parameters



Kobližková, M., Růžičková, P., Čupr, P., Komprda, J., Holoubek I., Klánová, J.: Soil burdens of persistent organic pollutants – their levels, fate and risks. Part IV. Quantification of volatilization fluxes of organochlorine pesticides and polychlorinated biphenyls from contaminated soil surfaces. Environmental Science & Technology. Accepted.

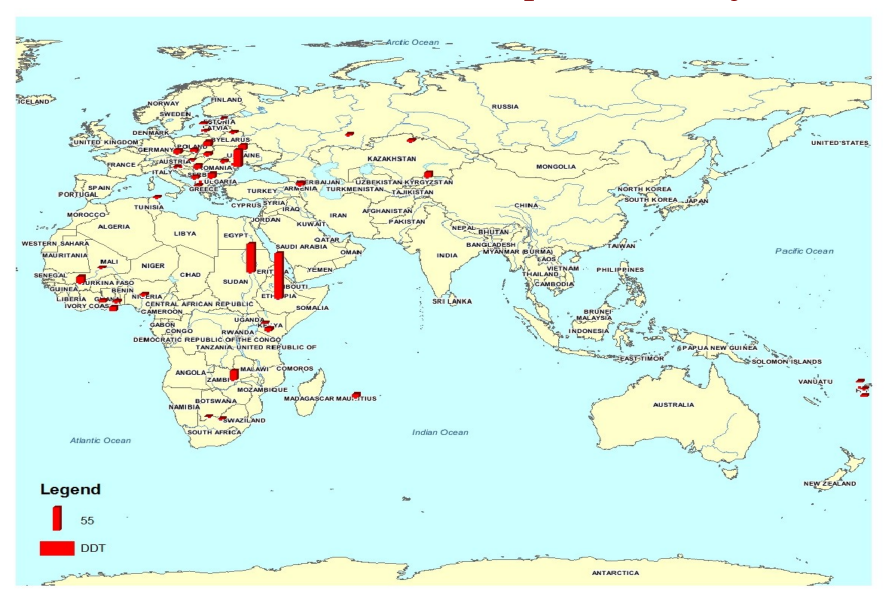


#### Existing and new air programs

To help ensure long term sustainability and comparability of results, it is recommended that new programs grow from strategic partnerships with existing programs, and benefit from their experiences.

Training visits for the purpose of capacity building/technology transfer are encouraged and should be supported.

#### Central and Eastern Europe, Africa, Fiji



Klanova, J., Cupr, P., Holoubek, I., Borůvková, J., Přibylová, P., Kareš, R., Kohoutek, J., Dvorská, A,.Komprda, J.: Towards the global monitoring of POPs), 2009. Masarykova Univerzita, Brno, Czech Republic. ISBN 978-80-210-4853-9



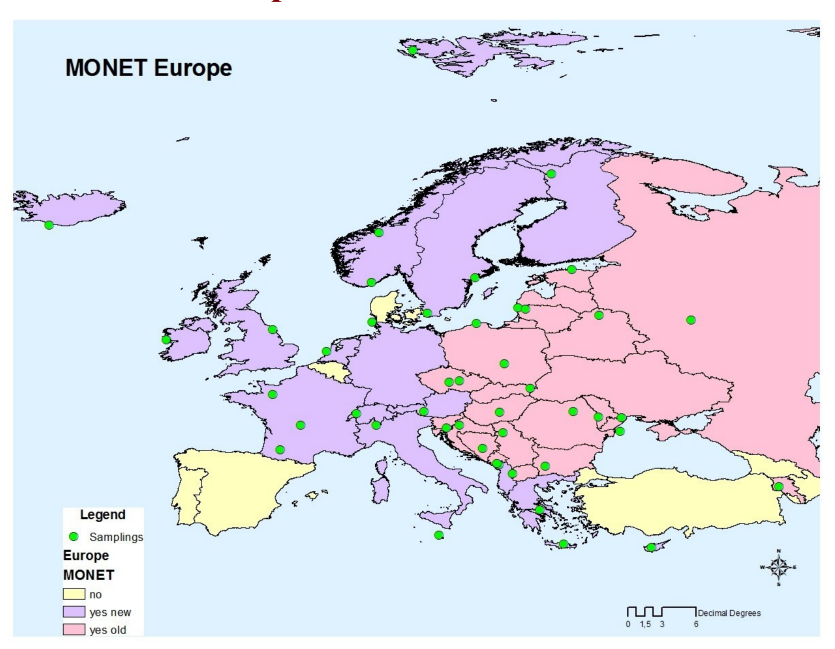
#### **Data Availability**

The group agreed that better access to data would be useful for several purposes e.g. model development.

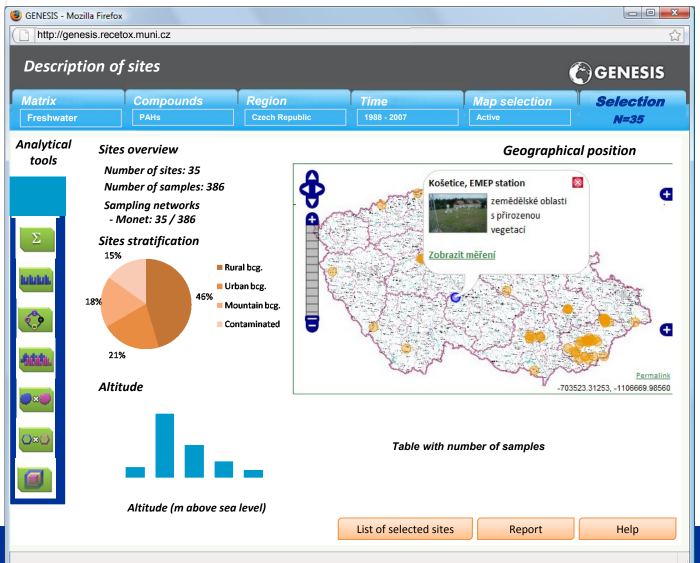
Existing and new programs are strongly encouraged to incorporate data management in their programs, i.e. employ sustainable databases.

Consideration should be given to the most suitable format and interface for end users of the data e.g. modelers.

#### **European PAS Network**



#### Description of more than one site



- Description of sites set includes:
- Number of sites and samplings and their affiliation to monitoring networks. Display of a table with sites list is also possible.
- Range of samplings in individual years
- Map with position (interactive map with options of zoom and turning on/off different layers)
- Distribution of sites altitude