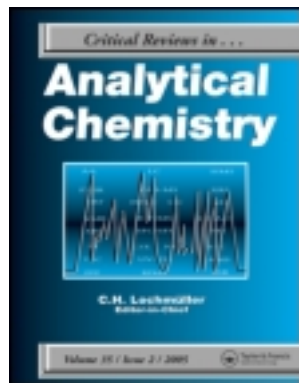


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Analysis of Airport Runoff Waters

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A particularly important facet of airport operation is the impact of the pollution caused by runoff waters. Runoff waters at an airport may contain relatively high concentrations of different contaminants resulting from the various aspects of its operation: de/anti-icing operations, washing and cleaning operations, spills of fuel and lubricants, exhaust fumes, and weed removal. The pollution caused by airport operations affects soil, surface waters, and groundwater. This issue is important to various stakeholders, particularly those residing in communities near airports, whose health, property values, and quality of life can be affected by such environmental impacts. The authors' intention is to present a critical review of literature data concerning (1) the types of pollution generated at airports, (2) methods for sampling runoff waters, (3) the analytical methods available for sample preparation, and (4) the analytical methods available for determining contaminants produced during airport operations. In addition, the article supplies literature information on the analytes contained in samples of runoff water from airports in different parts of the world.

Keywords Airport runoff waters, pollutants, sources of emission, sample collection, analytical procedures

INTRODUCTION

Despite the many positives ensuing from the rapid expansion of the air transport sector, airport operations as a whole are a substantial source of environmental pollution (Wensveen, 1999). These operations are connected with movement of aircraft and airport infrastructure. They include the deicing and anti-icing of aircraft and airfields; the movement of passenger vehicles and airport ground service equipment; the cleaning and maintenance of aircraft, ground service equipment, and motor vehicles; airport facility operations and maintenance; and removal of weeds and other vegetation from the airport apron (Luther, 2007). Very large amounts of harmful chemicals are utilized during these operations: for example, the quantities of de/anti-icing sprays range from as few as 11 gallons/plane to as many as 3,900 gallons/plane for large aircraft during bad weather (Zitomer, 2001).

All airport operations have adverse effects on water, air, soil, and animals, and further environmental problems are associated with climate change and aircraft noise (Douglas and Lawson, 2003; Kijewski, 2001).

A particularly important aspect is the contamination caused by airport runoff waters, which are produced when rain or other

precipitation washes the chemicals used during aircraft and airfield deicing and anti-icing, refueling aircraft, vehicle cleaning and maintenance, etc., off the airport apron. This runoff gets into the soil, surface waters, and even groundwater. Figure 1 shows a diagram of the effects of an airport on the environment.

This issue is important to various stakeholders, particularly those residing in communities near airports, whose health, property values, and quality of life can be affected by such environmental impacts (Luther, 2007).

Continuous monitoring of the airport waste stream is key in controlling runoff. The airport operator should understand the potential for and establish measures and procedures to address situations where there would be adverse consequences following the discharge of airport industrial waste waters. This may include reviewing spill prevention and countermeasure plans and notifying facility authorities of potential problems (O'Donnell, 2008).

The authors' intention is to present a critical review of literature data concerning:

- The types of pollution generated at airports;
- Methods for sampling runoff waters;
- The analytical methods available for sample preparation;
- The analytical methods available for determining contaminants produced during airport operations.

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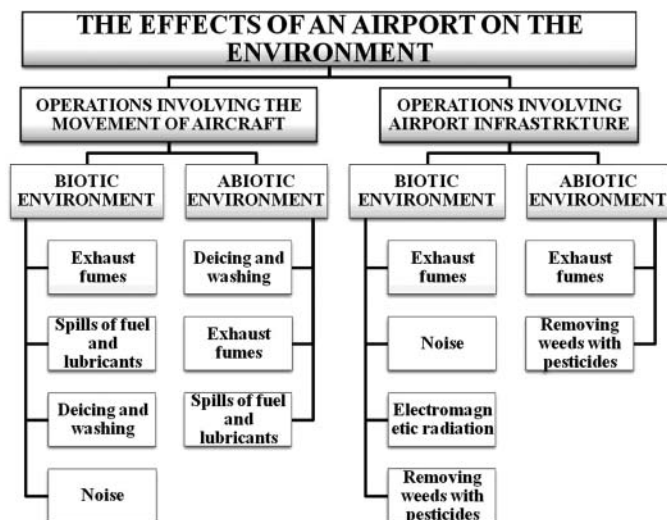


FIG. 1. The effects of an airport on the environment.

In addition, the article supplies literature information on the analysis of samples of runoff water from airports in different parts of the world.

TYPES OF AIRPORT INDUSTRIAL WASTE

The various activities of an airport may affect the abiotic and biotic environments. We will now discuss the pollution produced by different airport operations. Airport industrial wastewaters are generated during the de/anti-icing of aircraft, aircraft maintenance and repair work, aircraft and ground vehicle washing, aircraft and ground vehicle movement, and weed removal. A proper classification of airport-generated industrial wastes includes industrial wastewaters, hazardous wastes, and nonhazardous wastes. Industrial wastewaters are generally characterized in terms of conventional pollutants (oil and grease, total suspended solids, pH change of water, and 5-day biochemical oxygen demand) and priority pollutants (hydrocarbons, polycyclic aromatic hydrocarbon (PAHs), pesticides, polychlorinated biphenyls (PCBs), heavy metals, cyanides). Hazardous wastes may be inflammable, corrosive, reactive, and toxic. Nonhazardous wastes include oily rags or sludge that may be packaged for disposal in plastic bags or shipped to be recycled. Table 1 lists the main types of airport pollutants and their effects on the environment (Luther, 2007; McConnell et al., 1999; O'Donnell, 2008).

ANALYTICAL PROCEDURES USED IN STUDIES OF SAMPLES OF AIRPORT RUNOFF WATERS

In order to evaluate the threat to surface waters and soils, and, by extension, to ground and abyssal waters, the identification of the chemical compounds contaminating airport runoff is essential. Moreover, to assess the rate at which and the means by which runoff contaminants migrate through the environment, we need to know their physical and chemical properties, since it is

these that govern the extent to which chemical, biochemical, and photochemical processes participate in their environmental effects (Leško and Pasek, 1997). Waste composition varies greatly from airport to airport. This variability reflects the numerous aspects of aircraft deicing, cleaning, solvents used, and storm water management, including variable meteorological events, different deicing/anti-icing and cleaning practices, and collection systems (Switzenbaum et al., 1999).

The analysis of airport runoff samples presents a big challenge to analysts. The main problems in this respect are:

- The low, and even very low, levels of a wide range of contaminants;
- The very considerable variability in the concentrations of particular contaminants in runoff samples from different airports;
- The difficulties in standardizing measurement results because of:
 - ✓ The varying intensity of air traffic;
 - ✓ Changes in weather conditions;
 - ✓ The geographical locations of airports;
- The possibility that samples contain analytes with very similar physical and chemical properties but which are of widely differing toxicities with respect to both biota and abiota;
- The lack of standard techniques for sampling runoff waters, which can significantly affect the reliability of measurements;
- The limited availability of suitable standard solutions and the lack of reference materials (with different metrological values), which are essential for:
 - ✓ Calibrating monitoring instruments;
 - ✓ Validating the various steps in the analytical procedure as well as whole analytical methodologies, which are the tools for obtaining reliable information on the content of and processes occurring in samples of runoff waters during their transport, storage, and preparation for analysis.

Figure 2 shows a general scheme of the analytical procedures used for investigating samples of airport runoff.

Sampling and preparing samples for analysis are key stages in every procedure designed to analyze a particular group of constituents. It is most important that the sample is representative. However, accurate sampling necessary for the correct analysis of airport industrial wastes can be difficult because such wastes are seldom homogeneous, e.g., their composition may vary widely within a period of minutes.

For continuously flowing wastewater streams, the flow rates of both individual and combined streams should be measured at representative points and expressed in standard units such as gallons per minute (gpm), gallons per hour (gph), or gallons per day (gpd). The method used by the airport operator to

TABLE 1
Types of airport industrial wastes and a short description of their impact on the environment.

Type of pollutant	Origin of pollution	Compounds emitted	Environmental impact	References
Combustion gases	• Combustion of aviation fuels • Combustion of engine fuels	CO ₂ ; NO _x ; H ₂ O; sulfate ions; particulate matter components (PM ₁₀ , PM _{2.5-10} , PM _{2.5} , PM _{1.0}); polycyclic aromatic hydrocarbons (PAHs); CO; SO ₂ , aldehydes, aliphatic hydrocarbons, volatile organic compounds (VOCs)	• Present a direct threat to employees' health and safety • Resistance of trees to pests and disease • Impair growth of biomass and lower its quality • Adverse effect on fungi, algae, lichens • Damage to buildings • PAHs are mutagenic	Amato et al., 2010; Fang et al., 2007; Hoffman et al., 1984; Kesgin, 2006; Pison, 2004; Polidori et al., 2010; Ray et al., 2008; Takada et al., 1990; Unal et al., 2005; Westerdahlet al., 2008; Winter et al., 2006
Fuel, oil, grease	• Vehicle maintenance shop operations • Fuelling operations • Engine test cell operations	• Hydrocarbons: aliphatic (n-heptane, pentane, hexane, pentene), olefins, aromatic (benzene, ethylbenzene, toluene, phenol, o-xylene) • PAHs (benzo(a)pyrene, benzo(a)anthracene, naphthalene, phenanthrene) • Polychlorinated biphenyls (PCBs)	• Contamination of surface waters and soils • Consumption of dissolved oxygen • Hinder re-oxygenation of streams • May form sludge deposits that could interfere with stream self-purification processes • Aromatic hydrocarbons are carcinogenic • Volatile aliphatic hydrocarbons can give rise to an explosion or fire	Lesko and Pasek, 1997
Detergents	• Cleaning of aircraft and ground vehicles • Repairs to aircraft engines and ground vehicles, • Cleaning airport aprons	• Aqueous-neutral detergent • Aqueous-nonionic detergent • Aqueous-alkaline/hydroxide • Aqueous-alkaline with detergent • Semi-aqueous • Semi-aqueous/terpene • Semi-aqueous/glycol ether • Semi-aqueous/abrasive • Semi-aqueous/hydroxide • Aliphatic/terpene • Aliphatic/glycol ether • Glycol ether blends	• May cause foaming in aeration basins • May cause partial sludge flotation through release of carbon dioxide	O'Donnell, 2008; Sierra et al., 1981

De/anti-icing chemical wastes	De/anti-icing operations	• Propylene glycol • Ethylene glycol • Diethylene glycol • Urea • Salts, i.e., sodium and potassium acetates; sodium formate; calcium and sodium chlorides	• May interfere with biological activity • Consumption of dissolved oxygen • May form sludge deposits that could interfere with stream self-purification processes • Toxic and endocrine disrupting effects	Breedveld et al., 2003; Corsi et al., 2006a, Espey and Legarreta, 1993; U.S. EPA, Office of Water, 2000; Switzenbaum et al., 1999
Toxic metals and chromium compounds	• Bright dipping • Chromium plating • Copper stripping • Anodizing operations • Corrosion of aircraft parts and ground vehicles • Use of road salt • Removal of exterior paint	• Cd, Cr, Cu, Pb, Ni, Zn • Chromium compounds	• May interfere with biological activity and may complicate sludge disposal • Toxic to human beings, livestock, and aquatic life	O'Donnell, 2008; Walker et al., 1999
Alkalies and acids	Pickling and cleaning operations	• Generally, acids and alkalis	• May corrode pipes, pumps, and treatment units and may interfere with settling and biological activity	O'Donnell, 2008
Specific organic compounds	• Cleaning of aircraft and ground vehicles • Paint application and removal	• Benzotriazole, totyltriazole • Phenols	• Consumption of dissolved oxygen • May form sludge deposits that could interfere with stream self-purification processes • May interfere with biological activity	O'Donnell, 2008; Gigger et al., 2006
Batteries	Spent lead-acid, lithium, and nickel-cadmium batteries are generated from routine ground vehicle maintenance	• Cadmium, nickel, and lead compounds	• Contaminate surface waters, groundwater • Dissolve metals and other materials • Burn human skin • Toxic to human beings, livestock, and aquatic life.	O'Donnell, 2008
Cyanides	• Steel hardening • Metal plating • Rust prevention • Stain removal operations	HCN, KCN		O'Donnell, 2008

(Continued on next page)

TABLE 1
Types of airport industrial wastes and a short description of their impact on the environment. (*Continued*)

Type of pollutant	Origin of pollution	Compounds emitted	Environmental impact	References
Pesticides	Removing weeds and other vegetation from the airport apron	Organophosphate insecticides	• Chronic health effects: reduced cholinesterase (adults at work can have reduced attention)	McConnell et al., 1990
Electromagnetic radiation	Formation of electromagnetic radiation by the transmitting, telecommunication, radiolocation, and radionavigation equipment used in aviation		• Can lead to dysfunction of: the central nervous, reproductive, cardiovascular, or hormonal system, organs of sight and hearing	Amato et al., 2010
Noise	Large-scale operation of jet-powered aircraft, both military and civil		Differing effects on people: • Impaired sight and/or hearing • Apathy • Headache • Difficulties with concentration • Problems with social relations • May adversely affect the nervous system	Kil and Podciborski, 2008

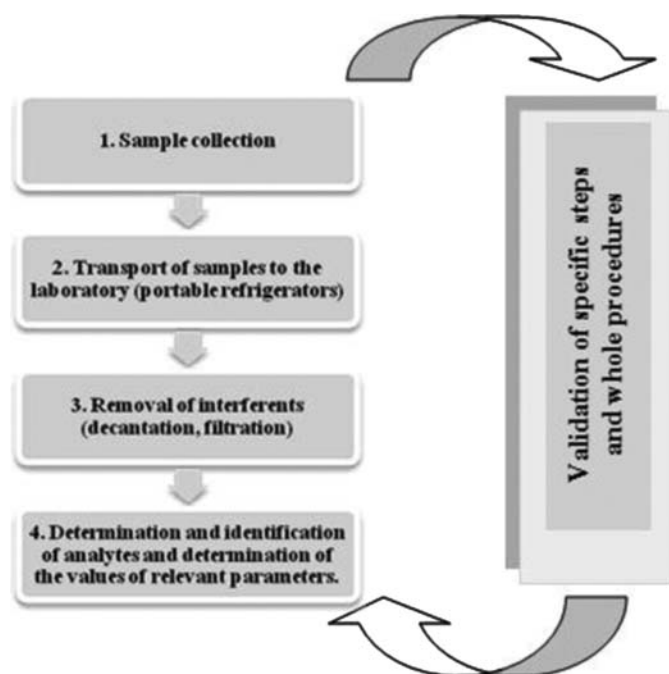


FIG. 2. General scheme of the analytical procedures used in studies of samples of airport runoff samples.

determine the flow rate will depend upon the magnitude of flow. Common metering devices include weirs, nozzles, flumes, and flow meters. For wastes that are generated on an intermittent basis, such as spent process baths, certain hazardous wastes, and deicing runoff, generation rates can be determined from the disposal volumes and dates. Flow proportional samples are recommended where applicable (O'Donnell, 2008).

The intensification of air traffic at airports and the expanding number of airports means that this particular human activity is putting significant pressure on the environment. Clearly, then, the monitoring of airport runoff content should cover the broadest possible range of substances. Only monitoring carried out on such a scale is capable of supplying data that will form a reliable basis for assessing the effects of airports on the abiotic and biotic compartments of the environment.

There should be two types of periods during which runoff samples collected from various parts of an airport are analyzed:

- An initial monitoring period, when samples are collected as often as possible, e.g., several samples should be collected each time contaminants are produced;
- A long-term monitoring period, when samples are collected, say, every month or every year, in order to confirm that the properties of airport runoff waters have not changed.

Collection of Airport Runoff Samples

An ideal sampling event may be one that takes advantage of special weather conditions, such as following a major snow-storm, which requires blowing and deicing. The post-storm

temperature rise will produce runoff from which it is possible to determine the seasonal and annual fluctuations in constituent concentration (Halm, 1996).

It is important to choose the appropriate location of storm water monitoring sites (Kent et al., 1999). These should be placed near sites where the largest amounts of pollutants are emitted, e.g., deicing pads, fuel distribution pads, fuel tanks, transshipment points, fuel pipelines, and repair shops, or at an upstream reference site, the primary and secondary airport outfalls, and the receiving stream site (Leško and Pasek, 1997).

Airport runoff waters are usually collected from storm drains, each serving a different land use: the main terminal area, the primary deicing and anti-icing area, taxiways and runways. Moreover, at most airports, aircraft deicing operations are performed on special pads (aircraft parking ramps or at the passenger terminal gates) (U.S. EPA, Office of Water, 2000). To collect the wastewaters generated at these locations, some airports have installed new collection systems or modified existing storm water drainage systems. The typical collection system consists of graded concrete pavement with trench or square drains connected to a wastewater storage facility via a diversion box (O'Donnell, 2008). The storage facility may consist of detention ponds (covered or uncovered), tanks, or underground concrete basins. The diversion box allows uncontaminated storm water to be diverted to storm water outfalls. The construction or modification of drainage collection systems with their associated underground piping, diversion boxes, and storage facilities can be extremely expensive, especially for larger airports that have several passenger terminals and a large number of gates. Sometimes aircraft deicing and anti-icing fluid (ADAF) formulations can be collected directly from storage tanks and deicing/anti-icing vehicles (glycol recovery vehicles), which remove the deicing and anti-icing agents directly from the airport apron (Corsi et al., 2006a).

Sample bottles for different analytes should be of the appropriate type (e.g., tubing model 3700R, Isco Industries, Lincoln, Neb.; refrigerated glass amber containers for analyzing deicing and anti-icing agents) (Budavari, 1996; Corsi et al., 2006b) and size (e.g., 500 mL) (Corsi et al., 2003) and contain the appropriate preservative. To minimize the possibility of sample contamination, containers must be thoroughly rinsed with deionized water prior to use (Knott et al., 1995).

For airport industrial wastewaters, grab or composite samples should be taken and properly preserved before analysis. The sampling operation should be as frequent as situation-specific requirements dictate (O'Donnell, 2008).

Airport runoff samples may be collected manually or by use of an automated sampler (for example, Isco, Lincoln, Neb.) from the drain outfall (Corsi et al., 2003; Saito et al., 2004).

Techniques of Sample Preparation

The preparation of samples for analysis is often an essential step in analytical procedure, especially when constituents present at trace or ultra-trace levels are to be determined. Runoff

TABLE 2
Literature information on the analytical procedures used and the concentration ranges of various types of xenobiotics in samples of airport runoff.

Analyte	Sample collection, preservation, and handling	Final determination technique	Validation parameters	References
pH Conductivity	• Calibration in a buffered solution of pH 4.01 • Before each measurement the electrode should be rinsed in demineralized water and dried with absorbent paper	Liquid samples—water Physicochemical parameters Electrochemical technique Electrochemical technique	No data available	U.S. EPA, Office of Water, 2000
BOD ₅	• Keep samples at or below 4°C during composting; limit composting period to 24 h • pH 6.5 to 7.5 • Incubation at the standard test temperature (20°C) for 5 days • Samples should be preserved with sulfuric acid at pH < 2 and maintained at 4°C until analysis • Interferences: chlorides, mercuric sulfate is added to the digestion tubes to complex the chlorides	Summary parameters Standard Methods for the Examination of Water and Wastewater 5210 B, 5-Day BOD Test	No data available	U.S. EPA Method 5210
COD		EPA Method 410.4 Titrimetric	• Measurement range: 3–900 mg/L • Precision and accuracy: 86 analysts in 58 laboratories analyzed a distilled water solution containing oxidizable organic material equivalent to 270 mg/L COD • Standard deviation (SD) was 17.76 mg/L COD with an accuracy, expressed as percentage relative error (bias), of –4.7%. (EPA Method Research Study)	EPA Method 410.4

TOC	• Collection and preparation of samples in glass bottles (the best way) • Samples should be kept cool (4°C) and protected from sunlight and atmospheric oxygen • In instances where analysis cannot be performed within 2 h of sampling, the sample should be acidified to pH 2 with HCl or H₂SO₄ • Carbonates and bicarbonates should be removed	EPA Method 415.1 Combustion Or Oxidation	• Measurement range: the method is best applicable to the measurement of organic carbon above 1 mg/L • Precision and accuracy: ✓ Increment as TOC (mg/L) = 4.9; - Precision as Standard Deviation TOC, mg/L = 3.93; Accuracy as Bias(%) = +15.27; Bias(mg/L) = +0.75 ✓ Increment as TOC (mg/L) = 107; - Precision as Standard Deviation TOC, mg/l = 8.32; Accuracy as Bias(%) = +1.01; Bias(mg/L) = +1.0 • Measurement range: 4–20 000 mg/L • Precision data are not available at this time	U.S. EPA Method 415.1
TSS	• Preservation of the sample is not practicable; analysis should begin as soon as possible • Refrigeration or icing to 4°C • A well-mixed sample is passed through a glass fiber filter, and the residue retained on the filter is dried to constant weight at 103–105°C	EPA Method 160.2 Gravimetric	• Measurement range: 4–20 000 mg/L • Precision data are not available at this time	U. S. EPA Method 160.2
TKN (total Kjehldahl nitrogen)	• Chloride anion-exchange resins should be used to remove nitrates prior to analysis • Maximum holding time: 28 days • Sample handling and preservation: may be preserved by the addition of conc. H₂SO₄, stored at 4°C	Groups of compounds EPA Method 351.3 • Colorimetric • Titrimetric • Potentiometric	Measurement range: 0.05–1400 mg/L	U.S. EPA Method 351.3
Ammonia as nitrogen (NH ₃)	• Samples may be preserved with H₂SO₄ and stored at 4°C, pH 9 • Distillation is used prior to analysis to reduce/eliminate interferences (e.g., formaldehyde) • Remove residual chlorine using sodium thiosulfate	EPA Method 350.1 • Colorimetric • Titrimetric • The electrode method • FIA	Measurement range: • FIA method: 1.0–1000 mg/L NH₄⁺; 1.2–1,200 mg/L NH₃ • 0.05–1.0 mg NH₃-N/L (colorimetric procedure) • 1.0–25 mg/L (titrimetric procedure) • 0.05–1400 mg/L (electrode method)	U.S. EPA, Office of Water, 2000; U.S. EPA Method 350.2

(Continued on next page)

TABLE 2

Literature information on the analytical procedures used and the concentration ranges of various types of xenobiotics in samples of airport runoff. (*Continued*)

Analyte	Sample collection, preservation, and handling	Final determination technique	Validation parameters	References
Phosphorus, all forms (P): total orthophosphate, hydrolyzable phosphate, and phosphorus	If the analysis cannot be performed on the day of collection, the sample should be preserved by the addition of conc. H_2SO_4 and refrigeration at 4°C	EPA Method 365.3 Colorimetric	• Measurement range: 0.01–0.5 mg P/L range • Precision and accuracy: ✓ Water samples at concentrations of 0.04, 0.19, 0.35, and 0.84 mg P/L, standard deviations were ± 0.005, ± 0.000, ± 0.003, and ± 0.000 respectively; ✓ In a single laboratory (EMSL), using surface water samples at concentrations of 0.07 and 0.76 mg P/L, recoveries were 99% and 100% respectively	U.S. EPA Method 365.3
TPH (total petroleum hydrocarbons)	• A representative sample of 1 L volume should be collected in a glass bottle • A delay between sampling and analysis of greater than 4 h requires sample preservation by the addition of HCl. A delay of greater than 48 h also requires refrigeration for sample preservation	EPA Method 418.1 Spectrophotometric	• The method is sensitive to levels of 1 mg/L and less, and may be extended to ambient monitoring	U.S. EPA Method 418.1
Semi-volatile organic compounds (including tolyltriazoles)	• Sample storage at 0–4°C • Any residual chlorine in the sample should be removed by the addition of sodium thiosulfate • Begin sample extraction within 7 days of collection, and analyze all extracts within 40 days of extraction • Extraction: ✓ Samples containing 1% solids or less: liquid/liquid extraction techniques;	EPA Method 1625C • Gas chromatography-mass spectroscopy (GC-MS) • Gel permeation chromatography (GPC)	No data available	Wan et al., 1996

✓ Samples containing 1–30% solids are diluted to 1% with reagent water and, extracted using continuous liquid/liquid extraction techniques ✓ Samples containing more than 30% solids are extracted using ultrasonic techniques • Clean up with GPC • Glassware should be cleaned with great care (wash with detergents, rinse with solvent) • Acidification to pH < 2 • Extraction ✓ Solid phase extraction (SPE) ✓ Liquid-liquid extraction (LLE) • The sample is acidified to a low pH (< 2) • Serially extracted with fluorocarbon-113 in a separating funnel • Maximum holding time: 28 days	Hexane EPA Method 1664 Gravimetry • Measurement range: 5–1000 mg/L • Method detection limit (MDL) = 1.4 mg/L • Minimum level of quantitation (ML) = 5.0 mg/L U.S. EPA, Office of Water, 2000; Vlaming et al., 2000
Fats, oils, and grease EPA Method 413.1 Gravimetry • Measurement range: 5–1000 mg/L of extractable material • Precision: the method determined the oil and grease level in sewage to be 12.6 mg/L. When 1 L portions of sewage were dosed with 14.0 mg of a mixture of #2 fuel oil and Wesson oil, the recovery was 93% with a standard deviation of +/– 0.9 mg/L U. S. EPA Method 413.1	Glycols (PG, EG) De-icing agents • EPA Method 624 • EPA Method 8015b • GC-MS • Gas chromatography-flame ionization detector (GC-FID) • All samples must be iced or refrigerated from the time of collection until analysis • If the sample contains residual chlorine, add sodium thiosulfate preservative to the empty sample bottle just prior to shipping to the sampling site • Grab samples must be collected in glass containers GC-FID • Measurement range: ✓ 36–640 mg/L for ethylene glycol ✓ 40–1150 mg/L for propylene glycol • Precision: 5–600 mg/L; • Average recoveries: ✓ 93.9% for ethylene glycol ✓ 100% for propylene glycol Fries, 2009; U.S. EPA, Office of Water, 1999; 2000; U.S EPA Method 624

(Continued on next page)

TABLE 2

Literature information on the analytical procedures used and the concentration ranges of various types of xenobiotics in samples of airport runoff. (*Continued*)

Analyte	Sample collection, preservation, and handling	Final determination technique	Validation parameters	References
Nonylphenol ethoxylates, alkylphenol ethoxylates	- Seal the bottle so that no air bubbles are entrapped in it - If preservative has been added, shake vigorously for one minute		- Standard deviations: ✓ 8.8% for ethylene glycol ✓ 8.9% for propylene glycol - Limit of detection (LOD) = 18 mg/L (ethylene glycol and propylene glycol) - Relative percent differences (RPD) = 11% for ethylene glycol, RPD = 7.9% for propylene glycol	
	- SPE (for GC/MS, HPLC-MS, HPLC-MS/MS, LC/MS) - Soxhlet (for HPLC fluorescence) - Gas-stripping (HPLC-UV) - Liquid/liquid (for HPLC fluorescence) - NPnEOs were isolated by passing samples through a mixed-bed ion exchange column and then through a column containing octadecylsilica (C18) to remove NPnEOs - The NPnEOs were extracted from the C18 using warm methanol, which was subsequently evaporated to dryness under a stream of nitrogen - The residue was dissolved in chloromethane/hexane	Surfactants - High-pressure liquid chromatography (HPLC) - GC/MS - HPLC fluorescence - HPLC-UV - HPLC-MS - HPLC-MS/MS - liquid chromatograph-hyass spectrometry (LC/MS)	- Measurement range for HPLC: - MDLs = 0.005, relative standard deviation (RSD) = 0.16% for NP1EO - MDLs = 0.236 μg/L, RSD = 8.4% for NP15EO - RSD for the entire method (extraction + analysis) was 24% for total NPnEO ($n = 1-15$) - Measurement range for LC/MS: ✓ NPEs = 20 ng/mL (NP1EO) and 0.07 ng/mL ✓ NP17EO in the lowest standard to 3,000 ng/mL (NP1EO) and 11 ng/mL (NP17EO) in the highest concentration standard - OPE concentrations ranged from 1.3 ng/mL (OPE1EO) and 0.017 ng/mL (OP17EO) in the lowest concentration standard to 2,000 ng/mL (OPE1EO) and 13 ng/mL (OP17EO) in the highest concentration standard	U.S. EPA, Office of Water, 1999

Benzotriazoles 1H-benzotriazole Tolyltriazole-4-methyl- 1H-benzotriazole (4-MeBT)-5-methyl- 1H-benzotriazole (5-MeBT)	• Samples stored in glass bottles • Transported to the lab in ice chests • Passed through 0.45 μm membrane filters and stored in a refrigerator at 4°C pending analysis, usually within the next 24 h • Samples were acidified to pH 7 fortified to a 1.0 mg/L concentration of 5,6-dimethylbenzotriazole as a surrogate (for 4-methyl-1H benzotriazole) • SPE	• GC/MS, usually • GC/FID • HPLC-UV • Ultraviolet-visible spectrophotometry (UV-Vis) in soil extracts • LC/MS/MS	• Measurement range = 0.03–2900 $\mu\text{g/L}$ • GC/MS; LOD = 0.1 $\mu\text{g/L}$–0.08 mg/L • HPLC-UV; LOD = 0.5–1.0 ppm • UV-VIS; LOD = 0.1 mg/L for both BT and TT • LC/MS/MS: ✓ LOD = 8 ng/L BT (water samples) ✓ LOD = 3 ng/L TT; ✓ LOD = 0.1 mg/kg BT (for soil extracts) ✓ Limit of quantification (LOQ) = 0.15 mg/L benzotriazoles (surface water) ✓ LOQ = 0.25 mg/L benzotriazoles (wastewater) ✓ LOQ = 0.05 mg/L benzotriazoles after SPE (groundwater) • MDL = 0.08 mg/L (for both 4- and 5-methyl-1H benzotriazole) • LOD = 1.2 mg/L • Calibration was linear in the range of 0.5–208.5 mg/L • Detection limit of the method was 0.176 mg/L	U.S. EPA, Office of Water, 1999
Potassium formate (K+CHOO–)	• All samples were acidified to shift dissociation equilibrium of potassium formate and sodium formate-d solutions towards the free acids formic acid and [2H] formic acid (formic acid-d) • Derivatization to methyl formate and methyl formate-d • Headspace solid-phase microextraction (HS-SPME) • Containers were thoroughly rinsed with methanol and deionized water prior to use • Samples were stored at room temperature (22°C) prior to analysis • SPE	GC-MS	• LOD = 0.06 ng/L • LOQ = 0.1 ng/L	Knott et al., 1995
Perfluorooctanoate (PFOA)		LC-MS		

(Continued on next page)

TABLE 2

Literature information on the analytical procedures used and the concentration ranges of xenobiotics in samples of airport runoff. (*Continued*)

Analyte	Sample collection, preservation, and handling	Final determination technique	Validation parameters	References
Perfluorooctane (PFOS)	- Containers were thoroughly rinsed with methanol and deionized water prior to use - Samples were stored at room temperature (22°C) prior to analysis - SPE	LC-MS	- LOD = 0.04 ng/L - LOQ = 0.1 ng/L	Knott et al., 1995
Polycyclic aromatic hydrocarbons (PAHs), polychlorinated biphenyls (PCBs)	This description applies to HPLC - Transport of samples in polyethylene bags - Storage of samples at 4°C pending analysis - Samples dried in the dark, twigs and stones removed - Grab samples mixed thoroughly to make a composite sample - After homogenization, soil samples were passed through a 2 mm sieve - Extracted by ultrasonication (toluene, later the solvent was replaced with acetonitrile)	Solid samples—soil - GC-MS - GC-FID - MS - HPLC - Size-exclusion HPLC - Infra-red spectroscopy (IR) - Supercritical fluid chromatography (SFC) - Thin-layer chromatography (TLC) - UV - Fluorescence spectroscopy - Isotope ratio mass spectrometry - Gravimetric methods	Sum of 12 PAHs ranged from 2.39 to 7.53 ng g⁻¹ with a mean concentration of 4.43 ± 1.45 ng g⁻¹	Tzovolou et al., 2009
VOCs	- Grab samples are collected in glass containers - Samples are maintained at 0–4°C from the time of collection until analysis - If an aqueous sample contains residual chlorine, add sodium thiosulfate preservative	GC-MS	Detection limits of the method are usually dependent on the level of interference rather than instrumental limitations	U.S. EPA, Office of Water, 2000; U.S. EPA Method 1624f

- Water samples containing aromatic compounds (e.g., benzene, toluene) to be stored for no longer than 7 days
- The medium can also be water
- For aqueous samples, experimental evidence indicates that some aromatic compounds (benzene, toluene, and ethyl benzene) should be acidified with HCl to pH ca 2
- Pure
 - ✓ Samples containing 1% or more solids and low to moderate levels of pollutants are analyzed by purging a known weight of sample added to reagent water;
- Samples containing 1% or more solids and high levels of pollutants are extracted with methanol; an aliquot of the methanol extract is then added to reagent water and purged

Heavy metals

- The sample bottle, whether of borosilicate glass, polyethylene, polypropylene, or teflon, should be thoroughly washed with detergent and tap water, rinsed with 1:1 nitric acid, tap water, 1:1 hydrochloric acid, tap water, and finally deionized distilled water, in that order
- For the determination of total metals the sample is acidified with 1:1 redistilled HNO_3 to a pH of less than 2 at the time of collection

Metals

- AAS
- Inductively coupled plasma mass spectrometry (ICP/MS)

Cadmium (EPA Method 213.2):

- Optimum concentration range: 0.5–10 $\mu\text{g/L}$
- Detection limit: 0.1 $\mu\text{g/L}$
- Precision and accuracy:
 - $\text{SD} \pm 0.10$; recoveries = 96%, $c = 2.5 \mu\text{g Cd/L}$
 - $\text{SD} \pm 0.16$; recoveries = 99%, $c = 5.0 \mu\text{g Cd/L}$
 - $\text{SD} \pm 0.33$; recoveries = 98%, $c = 10.0 \mu\text{g Cd/L}$ Chromium (EPA Method 218.2):

Knott et al., 1995;
Nabizadeh et al., 2005;
U.S. EPA, Office of
Water, 2000; U.S. EPA
Method 1620;
600/4-79-020; 213.2;
218.2; 220.2; 239.2;
289.2

TABLE 2
Literature information on the analytical procedures used and the concentration ranges of xenobiotics in samples of airport runoff. (*Continued*)

Analyte	Sample collection, preservation, and handling	Final determination technique	Validation parameters	References
	• Sample is not filtered before processing • Choose a volume of sample appropriate to the expected level of metals • If much suspended material is present, as little as 50–100 mL of well-mixed sample will most probably be sufficient. (The sample volume required may also vary proportionally with the number of metals to be determined.)		• Optimum concentration range: 5–100 $\mu\text{g/L}$ • Detection limit: 1 $\mu\text{g/L}$; • Precision and accuracy: • $\text{SD} \pm 0.1$; recoveries = 97%; c = 19 $\mu\text{g Cr/L}$ • $\text{SD} \pm 0.2$; recoveries = 101%; c = 48 $\mu\text{g Cr/L}$ • $\text{SD} \pm 0.8$; recoveries = 102%; c = 77 $\mu\text{g Cr/L}$ Copper (EPA Method 220.2); • Optimum concentration range: 5–100 $\mu\text{g/L}$ • Detection limit: 1 $\mu\text{g/L}$ • Precision and accuracy: data not available Lead (EPA Method 239.2); • Optimum concentration range: 5–100 $\mu\text{g/L}$ • Detection limit: 1 $\mu\text{g/L}$ • Precision and accuracy: • $\text{SD} \pm 1.3$; recoveries = 885%; c = 25 $\mu\text{gPb/L}$ • $\text{SD} \pm 1.6$; recoveries = 92%; c = 50 $\mu\text{gPb/L}$ • $\text{SD} \pm 3.7$; recoveries = 95%; c = 100 $\mu\text{gPb/L}$ Nickel (EPA Method 249.2); • Optimum concentration range: 5–50 $\mu\text{g/L}$ • Detection limit: 1 $\mu\text{g/L}$ Precision and accuracy: data not available Zinc (EPA Method 289.2); • Optimum concentration range: 0.2–4 g/L • Detection limit: 0.05 $\mu\text{g/L}$ • Precision and accuracy: data not available at this time.	

TABLE 3

CHRONIC										ACUTE								
Invertebrates			Others				Vertebrates			Invertebrates								
Vertebrates	Rainbow trout	Ciliated protozoan	Rotifer	Ceriodaphnia dubia	S. costatum	P. phosphoreum	C. paramecium	S. capricornutum	Fathead minnow	Rainbow trout	Bluegill sunfish	Sheepshead	Clawed toad	Daphnia magna	Ceriodaphnia dubia	Crayfish	Mysidopsis	Species
	12-14 d	12-14 d	24-h	7-d	7-d LOEC	96-h EC50	5/30 min LC50	7-d LOEC	24-336 h-EC50	96-h LC50	96-h LC50	96-h LC50	48-h LC50	24/48-h LC50	96-h LC50	96-h LC50	96-h LC50	Toxicity end-point
	LOEC	EC50	LOEC	LOEC	LOEC	EC50	LC50	LOEC	h-EC50	LC50	LC50	LC50	LC50	h LC50	LC50	LC50	LC50	
	7-d	7-d	7-d	7-d	7-d	7-d	7-d	7-d	7-d	7-d	7-d	7-d	7-d	7-d	7-d	7-d	7-d	
	LOEC	LOEC	LOEC	LOEC	LOEC	LOEC	LOEC	LOEC	LOEC	LOEC	LOEC	LOEC	LOEC	LOEC	LOEC	LOEC	LOEC	

TABLE 4
Literature information on the analyses of samples of runoff water from airports in different geographical regions

Sampling location	Parameter	Range of concentration (mean value)[mg/L]*	References
International Airport, Warsaw, Poland	TOC	(446.5)	Krzemieniowski et al., 2006
	TN (total nitrogen)	(142.1)	
	Oil	(26.4)	
Airport, Gdańsk, Poland	TSS (total suspended solids)	(159)	Siedlecka, and Downar, 2004
	BOD ₅	4.1–130.0	
	COD	31–807	
	Petroleum ether extract	5.6–553.7	
	Glycols	5.6–553.7	
	TKN	9.2–356.6	
	BOD ₅	6.9–47.9	
	COD	42–227	
	Petroleum ether extract	6.1–208.5	
	Glycols	3.1–50.4	
	TKN	2.21–7.1	
	pH	6.4–8.1 (7.2)	
Heathrow International Airport London, England	COD	0.2–18.4	Revitt et al., 1997
	Specific conductivity [$\mu\text{S}/\text{cm}$]	347–890 (628)	
	Benzotriazole [$\mu\text{g}/\text{L}$]	0.16–5.44	
Glatt River near Zurich International Airport, Switzerland	Totyltriazole [$\mu\text{g}/\text{L}$]	0.04–0.91	Gigger et al., 2006
Fornebu International Airport, Oslo, Norway (sampling done 2 years after airport closure)	Aircraft de-icing point	Benzotriazole [$\mu\text{g}/\text{l}$]	Breedveld et al., 2003
		PG (Propylene glycol)	
		DOC (dissolved organic carbon)	
		Fe	
		SO ₄ ²⁻	
	Drainage ditch	Benzotriazole [$\mu\text{g}/\text{L}$]	
		PG	
		pH	
		BOD ₅	
		PG	
Westchester County Airport, N.Y., USA	Taxiway, drainage system	Oil and grease	U.S. EPA, Office of Water, 2000
		pH	
		BOD ₅	
		PG	
		Oil and grease	
	Buildings, hangars	pH	
		BOD ₅	
		PG	
		Oil and grease	
	Ponds	pH	
		BOD ₅	
		PG	
		Oil and grease	
		pH	
		COD	
Newark International Airport, N.J., USA	Runway	TOC	U.S. EPA, Office of Water, 2000
		TSS	
		Hydrocarbons	
		pH	
		TOC	
		TSS	
		Hydrocarbons	
	Terminal	pH	
		TOC	
		TSS	

(Continued on next page)

TABLE 4

Literature information on the analyses of samples of runoff water from airports in different geographical regions (*Continued*)

Sampling location	Parameter	Range of concentration (mean value)[mg/L]*	References
Salt Lake City International Airport, USA	pH	6.6–9.5	U.S. EPA, Office of Water, 2000
	BOD ₅	11–1,050 (332)	
	COD	104–3,880 (835)	
	Nitrate/nitrite	0.9–9 (4.73)	
Baltimore-Washington International Airport, USA	Main terminal area ^a Conductivity [μ mho/cm]	120–540	U.S. EPA, Office of Water, 2000
	pH	6.36–8.61	
	Alkalinity	38–520	
	Hardness	28–800	
	BOD	3–64	
	BOD ₅	23–2,510 (1,010)	
	COD	11,000–270,000	
	TSS	11–31	
	NH ₃	ND–23	
	TKN	1,1–400	
	EG (ethylene glycol)	< 10	
	P	0.2–60	
	Oil	ND–29	
	TPH	ND	
	Cadmium [μ g/L]	ND–3	
	Chromium [μ g/L]	ND–1	
	Copper [μ g/L]	ND–45	
	Lead [μ g/L]	ND–16	
	Nickel [μ g/L]	ND–5	
	Zinc [μ g/L]	240–1,430	
	Commuter terminal ^a Conductivity [μ mho/cm]	120–2,000	
	pH	5.91–8.48	
	Alkalinity	30–158	
	Hardness	46–168	
	BOD	1,900–138	
	BOD ₅	197–769 (412)	
	EG	< 10	
	TPH	1	
	COD	700–2,700	
	TSS	14–31	
	NH ₃	ND–5	
	TKN	ND–26	
	P	ND–6	
	Oil	ND–14	
	TPH	ND–28	
	Cadmium [μ g/L]	ND–1	
	Chromium [μ g/L]	ND	
	Copper [μ g/L]	3–9	
	Lead [μ g/L]	1–11	
	Nickel [μ g/L]	5	
	Zinc [μ g/L]	35–60	

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TABLE 4

Literature information on the analyses of samples of runoff water from airports in different geographical regions (*Continued*)

Sampling location		Parameter	Range of concentration (mean value)[mg/L]*	References
Muddy Bridge Branch (receives runoff directly from Baltimore-Washington International Airport), USA		Conductivity [$\mu\text{mho/cm}$]	210.5–328.0	Hartwell et al., 1995
		DO (dissolved oxygen)	4.5–6.4	
		pH	7.0–7.4	
		Alkalinity	69.0–84.2	
		NH ₃	0.003–0.042	
		EG	> 25 ^b	
Dallas/Fort Worth International Airport, Tex., USA	Upstream reference	PG	> 50 ^b	Corsi et al., 2006c
		BOD ₅	2–3.1	
		COD	14–80	
	Airport drainage	PG	< 18	
		EG	< 18	
		BOD ₅	93–9,500	
		COD	10–37,900	
	Receiving stream	PG	< 18–3,800	
		EG	< 18–20,000	
		BOD ₅	< 2 – > 100	
		COD	< 9–1,600	
		PG	< 18–39	
Sites near General Mitchell International Airport, Milwaukee, Wisc., USA	Upstream site	EG	< 18–230	Budavari, 1996; Corsi et al., 2009
		NPnEO [$\mu\text{g/L}$]	(0.89)	
		PG	< 18	
		EG	< 18	
		COD	47–84	
		BOD ₅	(16.6)	
		4-MeBT	< 0.08	
		5-MeBT	< 0.08	
		Acetate	< 5.0	
		Formate	< 2.5	
	Primary outfall	Potassium	(6.25)	
		NPnEO [$\mu\text{g/L}$]	(776)	
		PG	1,900–4,400	
		EG	32–280	
		COD	5,600–10,200	
		BOD ₅	738	
		4-MeBT	< 0.08–0.6	
		5-MeBT	< 0.08–0.8	
		Acetate	(120)	
		Formate	(11)	
	Receiving stream	Potassium	(59.1)	
		NPnEO [$\mu\text{g/L}$]	(16.9)	
		PG	130–150	
		EG	< 18	
		COD	330–480	
		BOD ₅	(201)	
		4-MeBT	< 0.08	
		5-MeBT	< 0.08	
		Acetate	(8.75)	
		Formate	< 2.5	
		Potassium	(15.5)	

(Continued on next page)

waters have a complex matrix composition, and the metrological parameters of most analytical techniques preclude the determination of the majority of compounds present in such waters. In order to analyze samples of such waters quantitatively, the analytes they contain must first be isolated and/or preconcentrated before the relevant extracts can be analyzed using the appropriate instrumentation.

The following analyte extraction techniques are routinely used:

- Solid phase extraction (SPE) (Knott et al., 1995; U.S. EPA, Office of Water, 1999, 2000; Vlaming et al., 2000);
- Liquid-liquid extraction (LLE) (U.S. EPA, Office of Water, 1999, 2000; Vlaming et al., 2000; Wan et al., 1996);
- Headspace solid-phase microextraction (HS-SPME) (U.S. EPA, Office of Science and Technology, 1989).

TABLE 4

Literature information on the analyses of samples of runoff water from airports in different geographical regions (*Continued*)

Sampling location	Parameter	Range of concentration (mean value)[mg/L]*	References
Kansas City International Airport, Mo., USA	BOD ₅	(5,100)	U.S. EPA, Office of Water, 2000
	TOC	(3,000)	
	5-methylbenzotriazole [$\mu\text{g/L}$]	(17,000)	
	NH ₃	(3.9)	
	Zinc	(140)	
	Phenol [$\mu\text{g/L}$]	(93)	
	Diethylene glycol	> 20,000	
	EG	(3,200)	
	PG	(16,000)	
	Aluminum	(860)	
	Magnesium [$\mu\text{g/L}$]	(2,500)	
	Manganese [$\mu\text{g/L}$]	(170)	
	Copper [mg/L]	(14)	
	n-tetradecane [$\mu\text{g/L}$]	ND	
	Lead	(15)	
	Potassium [$\mu\text{g/L}$]	(13,000)	
	Sodium [$\mu\text{g/L}$]	(1,100)	
	Calcium [$\mu\text{g/L}$]	(3,400)	
Bradley International Airport, Conn., USA	BOD ₅	(39,000)	U.S. EPA, Office of Water, 2000
	TOC	(3,500)	
	5-methylbenzotriazole [$\mu\text{g/L}$]	(90,000)	
	NH ₃	(23)	
	Zinc	(340)	
	Phenol [$\mu\text{g/L}$]	(280)	
	Diethylene glycol	(15,000)	
	PE	(3,000)	
	PG	(160,000)	
	Aluminum	(1,100)	
	Magnesium [$\mu\text{g/L}$]	(2,000)	
	Manganese [$\mu\text{g/L}$]	(140)	
	Copper	(44)	
	n-tetradecane [$\mu\text{g/L}$]	(140)	
	Lead	(50)	
	Potassium [$\mu\text{g/L}$]	ND	
	Sodium [$\mu\text{g/L}$]	(10,000)	
	Calcium [$\mu\text{g/L}$]	(33,000)	

(Continued on next page)

TABLE 4

Literature information on the analyses of samples of runoff water from airports in different geographical regions (*Continued*)

Sampling location	Parameter	Range of concentration (mean value)[mg/L]*	References
Greater Rockford International Airport, Ill, USA	BOD ₅	> 7.3	U.S. EPA, Office of Water, 2000
	TOC	(12)	
	5-methylbenzotriazole [$\mu\text{g/L}$]	(120)	
	NH ₃	(46)	
	Zinc	(45)	
	Phenol [$\mu\text{g/L}$]	ND	
	Diethylene glycol	ND	
	PE	ND	
	PG	ND	
	Aluminum [$\mu\text{g/L}$]	(270)	
	Magnesium [$\mu\text{g/L}$]	(3,000)	
	Manganese [$\mu\text{g/L}$]	(360)	
	Copper [mg/L]	(9.2)	
	n-tetradecane [$\mu\text{g/L}$]	ND	
	n-tetradecane [$\mu\text{g/L}$]	(4.3)	
	Lead [$\mu\text{g/L}$]	(7,900)	
	Potassium [$\mu\text{g/L}$] (64,000)	(14,000)	
	Sodium [$\mu\text{g/L}$]		
	Calcium [$\mu\text{g/L}$]		
Louisville International Airport, Ky., USA	pH	7–9.1	U.S. EPA, Office of Water, 2000
	BOD ₅	3–1,250	
	TSS	2–3,530	
	DO	0.270–13.0	
	Benzene [$\mu\text{g/L}$]	< 0.03–171	
	Ethylbenzene [$\mu\text{g/L}$]	< 5–97	
		5–127	
	Naphthalene [$\mu\text{g/L}$]	5–361	
	Toluene [$\mu\text{g/L}$]	< 5	
	Xylene [$\mu\text{g/L}$]	< 5	
Nashville International Airport, Tenn., USA	pH	7.2–8.6	U.S. EPA, Office of Water, 2000
	BOD ₅	3–98	
	COD	< 20–130	
	TSS	18–55	
	DO	6.4–11.9	

*Unless otherwise noted.

^aData sample type: peak, composite, and grab.^bBelow detectable concentrations.

The use of ultrasound as an assisting agent will accelerate the extraction of hydrocarbons (PAHs, PCBs).

Table 2 lists information on techniques for preparing samples of runoff waters for analysis.

Analytical Procedures for Determining Physicochemical Parameters and Contents of Analytes in Airport Runoff Samples

To date, not many results of analyses of runoff waters have been published. This situation is changing, however; interest

in this type of data is growing, as it provides a source of information on the potentially negative effects of airports on the environment. The physicochemical parameters of airport runoff and the analytes it contains can be determined using the appropriate analytical procedures.

Usually, airport runoff water samples are analyzed in order to determine the level of pH, chemical oxygen demand (COD), 5-day biochemical oxygen demand (BOD₅), total organic carbon (TOC), glycol content, total suspended solids (TSS), total phosphorus, and total Kjeldahl nitrogen (TKN). Apart from

summary parameters, other groups of compounds are also analyzed (petroleum compounds, surfactants, glycols, benzotriazoles, metals, and other inorganic compounds). The following analytical techniques are used to determine target analytes in suitably prepared samples of airport runoff: GC-MS, GC-FID, HPLC, GPC, HPLC-MS, HPLC-UV, HPLC-MS/MS, LC/MS, TLC, AAS, ICP/MS (Fries and Klasmeier, 2009; U.S. EPA, Office of Water, 1996, 1999, 2000; Wan et al., 1996). Table 2 also lists literature information on the analytical procedures used and the concentration ranges of various types of xenobiotics in samples of airport runoff.

In tandem with the chemical analysis of airport runoff waters, their toxicity is also evaluated. Tests using biological material are crucial, since they constitute the basis for assessing the overall degree of contamination of particular compartments of the environment (U.S. EPA, 1996; Namieśnik et al., 2003). Table 3 summarizes the most commonly used tests for assessing the acute and chronic toxicity of airport runoff waters (Kent et al., 1999).

LITERATURE DATA ON THE LEVEL OF POLLUTION DUE TO DIFFERENT XENOBIOTICS IN AIRPORT RUNOFF WATERS FROM DIFFERENT GEOGRAPHICAL REGIONS

Most cases of pollution by airport runoff waters are defined using total parameters such as chemical oxygen demand, 5-day biological oxygen demand, total organic carbon, total suspended solids, and certain specific compounds like hydrocarbons, propylene, and ethylene glycol. Table 4 lists literature information on the analyses of samples of runoff water from airports in different parts of the world. The waste content of airport runoff waters varies greatly from airport to airport because the types of chemical agents used in airport operations vary widely. Consequently, the characteristics of airport runoff waters generated by different airports do so, too (Switzenbaum et al., 1999).

CONCLUSIONS

Airport daily activities, such as fueling operations and ground vehicle maintenance, aircraft de/anti-icing, ground vehicle washing and cleaning, aircraft maintenance and repair work, engine test cell operations, and weeding the airport apron, are all sources of environmental wastes. Airports sometimes have their own sewage treatment plants, but many smaller airports do not even have facilities for the pretreatment of wastewater. In these cases most of the substances from airport operations left on the apron eventually enter the airport storm water and are subsequently transported to the receiving waters. The large discharge of airport wastes to runoff waters may have numerous adverse consequences, especially potential drinking water contamination.

The levels of contaminants determined in airport runoff waters varies greatly from airport to airport. The waste content of airport runoff waters varies greatly from airport to airport be-

cause the types of chemical agents used in airport operations vary widely.

This issue is important to various stakeholders, particularly those living in communities near airports, whose health, property values, and quality of life can be affected by such environmental impacts. The runoff waters generated by airport activities thus require special management procedures, e.g., source reduction, the use of alternative de/anti-icing agents, recycling of materials, and remediation technologies.

A very important aspect is the changes that should be made to the standards applicable to airport operations (e.g., deicing operations and oil spills), pollution prevention procedures, as well as state and local agency directives for monitoring and controlling runoff water pollution, particularly toxic water pollutants.

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REFERENCES

- Amato, F.; Moreno, T.; Pandolfi, M.; Querol, X.; Alastuey, A.; Delgado, A.; Pedrero, M.; Cots, N. Concentrations, Sources and Geochemistry of Airborne Particulate Matter at a Major European Airport. *J. Environ. Monit.* **2010**, *12*, 854–862.
- Breedveld, G. D.; Roseth, R. R.; Sparrevik, M.; Hartnik, T.; Hem, L. H. Persistence of the De-icing Additive Benzotriazole at an Abandoned Airport. *Water Air Soil Pollut.* **2003**, *3*, 91–101.
- Budavari, S., Ed. *The Merck Index*, 12th ed. Merck: Whitehouse Station, N.J., 1996.
- Corsi, S. R.; Zitomer, D. J.; Field, J.; Cancilla, D. A. Nonylphenol Ethoxylates and Other Additives in Aircraft Deicers, Antiicers, and Waters Receiving Airport Runoff. *Environ. Sci. Technol.* **2003**, *37*, 4031–4037.
- Corsi, S. R.; Geis, S. W.; Loyo-Rosales, J.; Rice, C. P. Aquatic Toxicity of Nine Aircraft Deicer and Anti-Icer Formulations and Relative Toxicity of Additive Package Ingredients Alkylphenol Ethoxylates and 4,5-Methyl-1H-benzotriazoles. *Environ. Sci. Technol.* **2006a**, *40*, 7409–7415.
- Corsi, S. R.; Geis, S. W.; Loyo-Rosales, J.; Rice, C. P.; Sheesley, R. J.; Failey, G. G.; Cancilla, D. Characterization of Aircraft Deicer and Anti-Icer Components and Toxicity in Airport Snowbanks and Snowmelt Runoff. *Environ. Sci. Technol.* **2006b**, *40*, 3195–3202.
- Corsi, T. R.; Glenn, R. H.; Geis, S. W.; Bergman, D. Impacts of Aircraft Deicer and Anti-Icer Runoff on Receiving Waters from Dallas/Fort Worth International Airport, Texas, USA. *Environ. Toxicol. Chem.* **2006c**, *25*, 2890–2900.
- Corsi, S. R.; Geis, S. W.; Bowman, G.; Failey, G. G.; Rutter, T. D. Aquatic Toxicity of Airfield-Pavement Deicer Materials and Implications for Airport Runoff. *Environ. Sci. Technol.* **2009**, *43*, 4340–4346.
- Douglas, I.; Lawson, N. Airport Construction: Materials Use and Geomorphic Change. *J. Air Transp. Manag.* **2003**, *9*, 177–185.
- Espey, W.; Legarreta, G. Storm Water Regulations Aircraft Deicer/Anti-Icers Operations. Paper presented at National Conference on

- Hydraulic Engineering, San Francisco, Calif., USA, 25–30 July, 1993.
- Fang, G. Ch.; Wu, Y. S.; Lee, W. J.; Chou, T. Y.; Lin, I. Ambient Air Particulates, Metallic Elements, Dry Deposition and Concentrations at Taichung Airport, Taiwan. *Atmos. Res.* **2007**, *84*, 280–289.
- Fries, E.; Klasmeier, J. Analysis of Potassium Formate in Airport Storm Water Runoff by Headspace Solid-Phase Microextraction and Gas Chromatography–Mass Spectrometry. *J. Chromatogr. A* **2009**, *1216*, 879–881.
- Gigger, W.; Schaffner, Ch.; Kohler, H. P. Benzotriazole and Tylosotriazole as Aquatic Contaminants. 1. Input and Occurrence in Rivers and Lakes. *Environ. Sci. Technol.* **2006**, *40*, 7186–7192.
- Halm, M. J. Managing Storm Water at Airports. *Pollut. Eng.* **1996**, *28*, 62–64.
- Hartwell, S. I.; Jordahl, D. M.; May, E. B. Toxicity of Aircraft De-Icer and Anti-Icer Solutions to Aquatic Organisms. *Environ. Toxicol. Chem.* **1995**, *14*, 1375–1386.
- Hoffman, E. J.; Mills, G. L.; Latimer, J. S.; Quinn, J. G. Urban Runoff as a Source of Polycyclic Aromatic Hydrocarbons to Coastal Waters. *Environ. Sci. Technol.* **1984**, *18*, 580–587.
- Kent, R. A.; Andersen, D.; Caux, P. Y.; Teed, S. Canadian Water Quality Guidelines for Glycols—An Ecotoxicological Review of Glycols and Associated Aircraft Anti-icing and Deicing Fluids. *Environ. Toxicol.* **1999**, *14*, 481–522.
- Kesgin, U. Aircraft Emissions at Turkish Airports. *Energy* **2006**, *31*, 372–384.
- Kijewski, T. Presented at the Diploma Seminar on Samolot a środowisko-hałas i emisja spalin Rzeszów, Poland, 2001/2011. (in Polish).
- Kil, J.; Podciborski, T. *Dźwięk w krajobrazie jako przedmiot badań interdyscyplinarnych*; Komisja Krajobrazu Kulturowego PTG: Lublin, 2008. (in Polish).
- Knott, M. H.; Turley, S. D.; Turkley, B. S.; Yonkos, L. T.; Ziegler, G. P. The Acute Whole Effluent Toxicity of Storm Water from an International Airport. *Environ. Toxicol. Chem.* **1995**, *14*, 1103–1111.
- Krzemieniowski, M.; Białowiec, A.; Zieliński, M. The Effectiveness of the Storm Water Treatment Plant at Warsaw Frederick Chopin Airport. *Arch. Environ. Prot.* **2006**, *32*, 25–33.
- Leško, M.; Pasek, M. *Porty Lotnicze: wybrane zagadnienia inżynierii ekologicznej*. WPS: Gliwice 1997. (in Polish).
- Luther, L. *Environmental Impacts of Airport Operations, Maintenance, and Expansion*; Congressional Research Service Report for Congress, April 5, 2007.
- McConnell, R.; Pacheco Antón, A. F.; Magnotti, R. Crop Duster Aviation Mechanics: High Risk for Pesticide Poisoning. *Am. J. Publ. Health* **1990**, *80*, 1236–1239.
- McConnell, R.; Pacheco, F.; Wahlberg, K.; Klein, W.; Malespin, O.; Magnotti, R.; Åkeblom, M.; Murray, D. Subclinical Health Effects of Environmental Pesticide Contamination in a Developing Country: Cholinesterase Depression in Children. *Environ. Res. Sec. A* **1999**, *81*, 87–91.
- Nabizadeh, R.; Mahvi, A.; Mardani, G.; Yunesian, M. Study of Heavy Metals in Urban Runoff. *Int. J. Environ. Sci. Technol.* **2005**, *1*, 325–333.
- Namieśnik, J.; Chrzanowski, W.; Żmijewska, P. *New Horizons and Challenges in Environmental Analytics*. Centre of Excellence in Environmental Analysis and Monitoring: Gdańsk, 2003.
- O'Donnell, M. J. *Management of Airport Industrial Waste*; No150/5320-15A Advisory Circular, U.S. Department of Transportation, Federal Aviation Administration, AAS, 2008.
- Pison, I. Quantification of the Impact of Aircraft Traffic Emissions on Tropospheric Ozone over the Paris Area. *Atmos. Environ.* **2004**, *38*, 971–983.
- Polidori, A.; Kwon, J.; Turpin, B. J.; Weisel, C. Source Proximity and Residential Outdoor Concentrations of PM_{2.5}, OC, EC, and PAHs. *J. Expo. Sci. Environ. Epidemiol.* **2010**, *20*, 457–468.
- Ray, S.; Khillare, P. S.; Agarwal, T.; Shridhar, V. Assessment of PAHs in Soil around the International Airport in Delhi, India Sharmila. *J. Hazard. Mater.* **2008**, *156*, 9–16.
- Revitt, M.; Shutes, R. B. E.; Llewellyn, N. R.; Worrall, P. Experimental Reedbed Systems for the Treatment of Airport Runoff. *Water Sci. Technol.* **1997**, *36*, 385–390.
- Saito, N.; Harada, K.; Inoue, K.; Sasaki, K.; Yoshinga, T.; Koizumi, A. Perfluorooctanoate and Perfluorooctane Sulfonate Concentrations in Surface Water in Japan. *J. Occup. Health* **2004**, *46*, 49–59.
- Siedlecka, E. M.; Downar, D. Quality of Water from Airport Gdańsk-Trójmiasto Region. *Ecol. Chem. Eng.* **2004**, *11*, 557–567.
- Sierra, E.; Shahane, A. N.; Villate, J. T. Airport Wastes and Water Quality. *J. Am. Water Resour. Assoc.* **1981**, *17*, 190–196.
- Switzenbaum, M. S.; Veltman, S.; Schoenberg, T.; Durand, C. M.; Waggoner, M. *Best Management Practices for Airport Deicing Stormwater*, Publication 173, Environmental Engineering Program, Water Resources Research Center, July, 1999.
- Takada, H.; Onda, T.; Ogura, N. Determination of Polycyclic Aromatic Hydrocarbons in Urban Street Dust and Their Source Material by Capillary Gas Chromatography. *Environ. Sci. Technol.* **1990**, *24*, 1179–1186.
- Tzovolou, D. N.; Benoit, Y.; Haeseler, F.; Klint, K. E.; Tsakiroglou, C. D. Spatial Distribution of Jet Fuel in the Vadose Zone of a Heterogeneous and Fractured Soil. *Sci. Total Environ.* **2009**, *407*, 3044–3054.
- U.S. Environmental Protection Agency. *Chemical Analysis of Water and Wastes. Oil And Grease (Gravimetric, Separatory Funnel Extraction)*, EPA Method 413.1.; Editorial Revision 1978.
- U.S. Environmental Protection Agency. Sample Handling and Preservation. In *Methods for Chemical Analysis of Water and Wastes*, EPA-600/4-79-020; U.S. EPA: Cincinnati, 1983.
- U.S. Environmental Protection Agency. *Nonhalogenated Organics Using GC/FID*, EPA Method 8015B; 1996.
- U.S. Environmental Protection Agency. *Petroleum Hydrocarbons (Spectrophotometric, Infrared)*, EPA Method 418.1; 2004.
- U.S. Environmental Protection Agency, National Environmental Monitoring and Support Laboratory. *Cadmium (Atomic Absorption, Furnace Technique)*. EPA Method 213.2; U.S. EPA: Cincinnati, 1986a.
- U.S. Environmental Protection Agency, National Environmental Monitoring and Support Laboratory. *Chromium (Atomic Absorption, Furnace Technique)*. EPA Method 218.2; U.S. EPA: Cincinnati, 1986b.
- U.S. Environmental Protection Agency, National Environmental Monitoring and Support Laboratory. *Copper (Atomic Absorption, Furnace Technique)*. EPA Method 220.2; U.S. EPA: Cincinnati, 1986c.
- U.S. Environmental Protection Agency, National Environmental Monitoring and Support Laboratory. *Lead (Atomic Absorption, Furnace Technique)*. EPA Method 239.2; U.S. EPA: Cincinnati, 1986d.
- U.S. Environmental Protection Agency, National Environmental Monitoring and Support Laboratory. *Nickel (Atomic Absorption, Furnace Technique)*. EPA Method 249.2; U.S. EPA: Cincinnati, 1986e.

- U.S. Environmental Protection Agency, National Environmental Monitoring and Support Laboratory. *Zinc (Atomic Absorption, Furnace Technique)*. EPA Method 289.2; U.S. EPA: Cincinnati, 1986f.
- U.S. Environmental Protection Agency, National Pollutant Discharge Elimination System. *Nitrogen, Ammonia (Colorimetric, Titrimetric, Potentiometric Distillation Procedure)*, EPA Method 350.2; (Editorial Revision) 1974.
- U.S. Environmental Protection Agency, National Pollutant Discharge Elimination System. *Phosphorus, All Forms (Colorimetric, Ascorbic Acid, Two Reagent)*, EPA Method 365.3; (Editorial Revision) 1978a.
- U.S. Environmental Protection Agency, National Pollutant Discharge Elimination System. *Residue, Non-Filterable (Gravimetric, Dried at 103–105°C)*, EPA Method 351.3; (Editorial Revision) 1978b.
- U.S. Environmental Protection Agency, National Pollutant Discharge Elimination System. *Residue, Non-Filterable (Gravimetric, Dried at 103–105°C)*, EPA Method 160.2; 1979.
- U.S. Environmental Protection Agency, National Pollutant Discharge Elimination System. *Organic Compounds by Isotope Dilution GC/MS*, EPA Method 1624; 1989.
- U.S. Environmental Protection Agency, National Pollutant Discharge Elimination System. *Total Organic Carbon in Water Total (Combustion Or Oxidation)*, EPA Method 415.1; 1999.
- U.S. Environmental Protection Agency, Office of Research and Development. *The Determination of Chemical Oxygen Demand by Semi-Atomated Clorimetry*, EPA Method 410.4; U.S. EPA: Cincinnati, 1993.
- U.S. Environmental Protection Agency, Office of Research and Development. *Biochemical Oxygen Demand (BOD)*, EPA 5210; 2001.
- U.S. Environmental Protection Agency, Office of Science and Technology. *Engineering and Analysis. Method 1625C Semivolatile Organic Compounds by Isotope Dilution GCMS*. U.S. EPA: Washington, D.C., 1989.
- U.S. Environmental Protection Agency, Office of Water. *Methods for Organic Chemical Analysis of Multicicipal and Industrial Wastewater Method 624*, EPA-821-B-96-005; U.S. EPA: Washington, D.C., 1996.
- U.S. Environmental Protection Agency, Office of Water. *N-Hexane Extractable Material (HEM; Oil and Grease) and Silica Gel Treated N-Hexane Extractable Material (SGTHEM; Non-polar Material) by Extraction and Gravimetry*, EPA-821-R-98-002 PB99-121949; U.S. EPA: Washington, D.C., 1999.
- U.S. Environmental Protection Agency, Office of Water. *Preliminary Data Summary Airport De-icing Operations (Revised)*, EPA-821-R-00-016; U.S. EPA: Washington, D.C., 2000.
- U.S. Environmental Protection Agency, Office of Water. *EPA-Methods 8015b*; U.S. EPA: Coachella, 2003.
- U.S. Environmental Protection Agency, Office of Water Regulations and Standards. *Metals by Inductively Coupled Plasma Atomic Emission Spectrography and Atomic Absorption Spectroscopy*, EPA Method 1620; 1989.
- Unal, A.; Hu, Y.; Chang, M. E.; Odman, M. T.; Russell, A. G. Airport Related Emissions and Impacts on Air Quality: Application to the Atlanta International Airport. *Atmos. Environ.* **2005**, *39*, 5787–5798.
- Vlaming, V. de; Connor, V.; DiGiorgio, C.; Bailey, H. C.; Deanovic, L. A.; Hinton, D. E. Application of Whole Effluent Toxicity Test Procedures to Ambient Water Quality Assessment. *Environ. Toxicol. Chem.* **2000**, *19*, 42–62.
- Walker, W. J.; McNutt, R. P.; Maslanka, C. A. The Potential Contribution of Urban Runoff to Surface Sediments of the Passaic River: Sources and Chemical Characteristics. *Chemosphere* **1999**, *38*, 363–377.
- Wan, H. B.; Wong, M. K.; Mok, C. Y. Pesticides in Golf Course Waters Associated with Golf Course Runoff. *Bull. Environ. Contam. Toxicol.* **1996**, *56*, 205–209.
- Wensveen, J. G. Paper presented at Airports Council International, Greenport '99 Conference and Exhibition – Thinking Green – Strategies for Balanced Airport Development in the Next Century, Amsterdam, The Netherlands, 21–23 April, 1999.
- Westerdahl, D.; Fruin, S. A.; Fine, P. L. The Los Angeles International Airport as a Source of Ultrafine Particles and Other Pollutants to Nearby Communities. *Atmos. Environ.* **2008**, *42*, 3143–3155.
- Winter, M.; Kousgaard, U.; Oxbøl, A. Study of Heavy Metals in Urban Runoff. *Sci. Total Environ.* **2006**, *366*, 218–232.
- Zitomer, D. H. Waste Aircraft Deicing Fluid: Management and Conversion to Methane. Marquette University, June 2001. <http://w.doa.state.wi.us/ocs.view2.asp?ocid=817> (accessed 4/4/11).