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## Pollutants in Airport Runoff Waters

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*Runoff waters from airport areas constitute a serious environmental problem. It is essential to monitor levels of pollutants emitted into the environment and measure their toxicity on a continuous basis. The authors' aim was to critically review data on pollution from aviation fuel combustion, aviation fuel spillage, the washing and cleaning of aircraft and airport service equipment, and the use of de-/anti-icing agents published in scientific journals. The authors also summarize information on the analytical methodologies available for the determination of pollutants emitted during airport operations and provide data on the toxicity of runoff containing compounds used in de-/anti-icing operations.*

**KEY WORDS:** airports, management of wastes, pollutants, runoff waters, toxicity

### 1. INTRODUCTION

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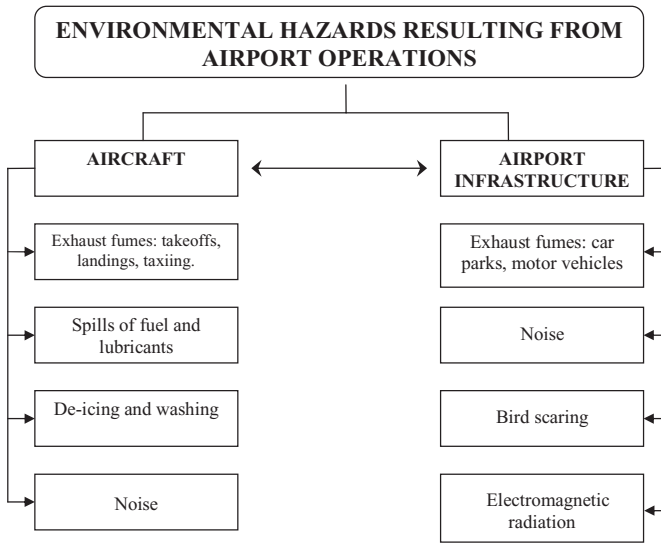
Air transport is the safest, quickest, and most convenient means of travelling long distances.<sup>1</sup> In addition, it facilitates economic and cultural exchanges and is a significant source of employment and growth in many regions.<sup>2</sup> Despite the positive aspects resulting from the intensive development of aviation, airports are large-scale polluters of the environment. The everyday activities at airports, such as aircraft and ground vehicle washing, cleaning, aircraft maintenance and repair work (including painting and metalwork), fuelling operations, engine test cell operations, de-/anti-icing operations, and ground vehicle maintenance are all sources of wastes (Figure 1).<sup>3</sup>

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Airport operations can contribute to the pollution of air, water, and soil; they can also adversely affect climate, the plant kingdom, and buildings (see Table 1).<sup>4</sup> However, their most destructive effects are on people and

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**FIGURE 1.** The environmental hazards resulting from airport operations.

animals.<sup>5</sup> The most effective way of reducing such pollution is, of course, to prevent it from occurring in the first place. So it is crucial to monitor levels of contaminants emitted into the environment and measure their levels of toxicity on a continuous basis.

One of the more important problems in this respect is the runoff waters that form when precipitation or atmospheric deposits flush the airport surface during its operation (runways, taxiways, washing and de-/anti-icing pads, aprons, transhipment points, fuel stores, repair shops, garages). The contaminants arising out of airport operation penetrate into all compartments of the environment.<sup>4</sup> In most cases airports do not possess their own effluent treatment plants, so all effluents carrying petroleum compounds, surfactants, the de-icing agents used in winter, and other organic and inorganic pollutants run off together with rain water or snow melt into drainage ditches, where they enter the soil, surface waters (rivers, lakes, ponds), and ultimately ground waters.

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## 2. THE MAIN SOURCES OF EMISSION AND ANALYTES PRODUCED DURING AIRPORT OPERATIONS

### 2.1 Washing and Cleaning of Aircraft and Airport Service Vehicles

Another important source of pollution at airports is the use of solvents for cleaning. Solvents are used extensively in the aerospace industry for cleaning processes that touch virtually all aspects of the manufacture, maintenance, operation, and repair of space, commercial, and military air vehicles. They

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**TABLE 1.** The principal contaminants produced during airport operations and their sources

Source	Xenobiotic	Reference(s)
Combustion of aviation fuels	CO <sub>2</sub> , CO, CxHy Volatile organic compounds (VOCs) Nitrogen oxides, NO <sub>x</sub> (NO <sub>x</sub> = NO <sub>2</sub> + NO+ HONO+HNO <sub>3</sub> ) Water vapor Particulate matter (PM <sub>10</sub> , PM <sub>2.5</sub> , PM <sub>1.0</sub> ), including airborne metals Sulfates Polycyclic aromatic hydrocarbons (PAHs) Sulfur dioxide, SO <sub>x</sub> Aldehydes Aliphatic hydrocarbons Benzene, ethylbenzene, toluene n-heptane, n-pentane, n-hexane, 1-pentene, o-xylene Phenol Benzo(a)anthracene, benzo(a)pyrene, naphthalene, phenanthrene	[1,6,7] [8] [1] [1,3,9] [1] [10,11] [5,6] [5] [5] [5]
Spillage of aviation fuels and greases	n-heptane, n-pentane, n-hexane, 1-pentene, o-xylene Phenol Anionic, cationic, nonionic detergents Glycols Methyl esters of fatty acids Propylene glycol (PG), ethylene glycol (EG) pH buffers (potassium hydroxide)	[12] [12] [13–16] [16]
Cleaning of aircraft and of airport service vehicles	Water Dyes 1,4-Dioxane Surfactants Triazole Sodium nitrite Sodium benzoate Borax Diethylene glycol Isopropyl alcohol (Isopropanol) Alkylphenol ethoxylates (APEO) Urea (NH <sub>2</sub> CONH <sub>2</sub> ) Ammonia (and conversion products—nitrites, nitrates)	[16] [14–16] [16] [17] [14,16] [16]
De-/anti-icing agents	CO Potassium acetate, calcium magnesium acetate (CMA), sodium acetate Formate (sodium, potassium) Potassium chloride, magnesium chloride, sodium chloride	[16] [16] [16] [14]

remove greases, oils, soot, and dust. Washing and polishing the body of an aircraft improves its performance by reducing friction. A clean aircraft consumes up to 3% less fuel and causes far less environmental contamination.<sup>18,19</sup> But these solvents are very harmful organic pollutants. The U.S. aerospace industry generates between 220,000 and 300,000 tons per year

of hazardous air pollutants (HAPs) and 145,000–200,000 tons per year of volatile organic compounds. Solvent cleaning operations account for 62% of these emissions. The three solvents that generate more than 90% of the total emissions from solvent cleaning operations are methyl ethyl ketone (MEK), 1,1,1-trichloroethane, and Freon 113. All three solvents are on the HAP list; they are used for general purpose cleaning processes.<sup>12</sup>

The federal aerospace regulations in the United States were developed by a team of representatives from government and civilian aerospace organizations, environmental organizations, and the U.S. Environmental Protection Agency (EPA). The aerospace National Emissions Standard for Hazardous Air Pollutants regulates cleaning solvent categories: hand-wipe cleaning (the removal of residue or contaminants from surfaces by rubbing these with rags, usually wetted with solvents), solvent flushing (involves pumping solvent from a reservoir through a pipe or hose or through equipment or assemblies to remove contaminants), and spraying (involves applying the solvent to a surface through a nozzle so that the solvent is under pressure).

Aerospace cleaning solvents can be grouped into four general categories: aqueous, organic, semiaqueous and halogenated solvents. The EPA has approved categories of certified solvents: aqueous-neutral detergent, aqueous-nonionic detergent, aqueous-alkaline/hydroxide, aqueous-alkaline with detergent, semiaqueous, semiaqueous/terpene, semiaqueous/glycol ether, semiaqueous/abrasive, semiaqueous/hydroxide, aliphatic/terpene, aliphatic/glycol ether, glycol ether blends, and solvent solutions  $\leq 5$  mm Hg at 20°C.<sup>12</sup> Examples of commercial preparations used for washing aircraft include the MrMuscle<sup>20</sup> and Aero 9738 aircraft shampoos.<sup>21</sup>

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Although solvent hand-wipe cleaning represents the largest pollutant source in the aerospace industry, difficulties are encountered in the formulation of regulations that would reduce pollutant emissions without compromising the operation being performed. The extreme variety of applications, required levels of cleanliness, and the location of the cleaning process (i.e., closed hangar vs. flight line) have complicated the process for identifying alternative solvent cleaning processes. The alternative solvents and processes used must be fully compatible with aircraft materials and not contribute to their degradation. A newly patented alternative cleaning agent is Spray Nine Corp (U.S. Patent No. 7,666,264 B2). This consists of at least one fatty acid methyl ester, and one or more surfactants in sufficient quantity to form an optically clear, stable, and effective emulsion.<sup>12</sup>

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## 2.2 Application of De-/Anti-Icing Agents

During periods of low temperatures, it is standard practice at airports to use de-icing agents and substances preventing the re-formation of ice on aircraft. De-icing involves removing ice, snow, and hoarfrost from the aircraft

95 surface, whereas the aim of anti-icing is to prevent ice from re-forming on the aeroplane.<sup>16</sup>

100 In the de-icing/anti-icing process, aircraft are usually sprayed with aircraft de-icing and antifreeze fluids (ADAFs) that contain chemical de-icing agents; however, nonchemical methods are also used. Numerous such operations may be in progress at airports: de-icing runways, de-icing and anti-icing aircraft, de-icing taxiways, aircraft maintenance, and salt de-icer application.<sup>21</sup>

105 Typically, the de-icing season lasts from October to April. In cooler climates this period may be longer, and in warmer ones it is usually shorter.<sup>16</sup>  
110 In winter, ice and snow on aircraft and runways impair the aerodynamic properties of planes and can compromise flight safety. The icing of aircraft increases their weight, makes it more difficult to achieve takeoff speed, and can cause mechanical equipment to become blocked.<sup>23</sup> Accumulating ice increases head resistance, adversely affects engine operation, and substantially reduces the stability and maneuverability of the aircraft. All larger aircraft are equipped with a battery of ice protection systems: heated leading edges and windowpanes, de-icing boots that can be inflated to break off ice, and a system for moistening propeller blades with alcohol. In most cases these arrangements are reasonably effective, but under certain conditions they may be insufficient, and in any case they use up a lot of energy.<sup>24</sup> That is why chemical de-icing and anti-icing agents are used at airports.

115 For removing hoarfrost and ice from an aircraft a mixture of chemical reagents and hot water is used, which is sprayed at high pressure onto the fuselage. De-icing mixtures consist mostly of glycol and urea, but they may also contain various additives.

120 Two types of de-icing are performed: wet-weather and dry-weather de-icing, depending on a number of climatic and operational factors. Wet-weather de-icing is performed during storm events that include precipitation such as snow, sleet, or freezing rain. Dry-weather de-icing is performed when changes in the ambient temperature cause frost or ice to form on aircraft but no precipitation is present.<sup>16</sup>

125 The quantities of spray range from as low as 40 l/plane to as high as 15,000 l/plane for large aircraft during bad weather.<sup>25</sup> Aircraft anti-icing fluids are applied in much smaller volumes than their deicing counterparts. A commercial jet requires approximately 133 l of fluid for anti-icing after deicing. Generally, dry-weather deicing requires 76–190 l of deicing fluid, depending on the size of the aircraft.<sup>16</sup> The amount of de-icing fluid required to de-ice a commercial aircraft depends on its size, the weather conditions, and airport operations.<sup>25</sup>

130 After use, the fluid typically mixes with storm water runoff and may enter lakes, rivers, and streams.<sup>25</sup>

### 2.3 Types of Aircraft De-/Anti-Icing Fluids

Aircraft de-/anti-icing fluid formulations consist of ethylene glycol, propylene glycol, or diethylene glycol as the freezing-point depressant; water; and numerous proprietary additives.<sup>26</sup> On one hand, de-icing agents must be environmentally friendly (biodegradable), and on the other they must be effective and not corrode the materials used in aircraft construction. For this reason, they also contain corrosion inhibitors, surfactants, antifoaming agents, and dyes.<sup>23</sup>

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Aircraft de-icers are categorized into four classes: Type I, Type II, Type III, and Type IV. Not all types are currently used.<sup>16</sup> Table 2 lists information characterizing the four types.

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### 2.4 Pavement De-/Anti-Icing

Pavement de-/anti-icing removes or prevents the accumulation of frost, snow, or ice on runways, taxiways, aprons, gates, and ramps. Because ice, sleet, and snow may be difficult to remove by mechanical methods alone, most airports use a combination of mechanical methods and chemical de-icing agents. Common pavement de-/anti-icing agents include urea, ethylene glycol, propylene glycol, an ethylene glycol-based fluid known as UCAR (50% ethylene glycol, 25% urea, and 25% water by weight), sodium formate, potassium acetate, calcium magnesium acetate, and sodium acetate. Sand may be sprinkled onto the pavement to increase friction but may mechanically damage the aircraft surface. Salts such as sodium chloride or potassium chloride cannot be used to de-ice taxiways or runways because of their corrosive properties; they are used only on airport car parks and roads.

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Pavement de-icing typically occurs during the same season as aircraft de-icing, but may be of shorter duration than the aircraft de-icing season.<sup>16</sup>

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The U.S. Environmental Protection Agency has estimated that 40 million liters of ADAF are discharged annually to receiving waters in the United States.<sup>26</sup> Though crucial for flight safety, ADAFs are the principal source of airport pollutants entering water, soil, and air: along with rainwater and snowmelt, they flow off the pavement, ultimately getting into surface waters, then ground waters, and the soil.<sup>5</sup> The principal environmental impacts associated with ADAF discharge to the environment include reduced dissolved oxygen in receiving waters and potential toxicity.<sup>27</sup> Glycols, which constitute the majority of ADAF formulations, have both a high biochemical oxygen demand (BOD) and a high chemical oxygen demand (COD).<sup>16,23,26,28</sup>

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The migration of these organic compounds to waters is also reflected in the ammonium nitrogen and total nitrogen factors. Nitrogen compounds are formed as a result of the biodegradation of urea and are responsible for

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**TABLE 2.** Characteristics of the four types of de-icing fluids used at airports<sup>[6,25,27]</sup>

Type of de-icing fluid	Type I	Type II	Type III	Type IV
Application	Removes ice and snow from aircraft surfaces, ensures short-term protection from refreezing	Prevent aircraft surfaces from refreezing and the accumulation of ice and snow on them		
Composition	8% water, 90% glycol, and less than 2% additives (dilution essential)	33% water, 65% glycol, and 2% additives (no additional dilution necessary)	No data	The same as Type II
Freezing temperature		From -27 °C to -60 °C		
Hold time	6–15 min	45 min	No data	70 min
Cost	Cheaper than types II, III, and IV		More expensive than Type I	
Additional information	<ul style="list-style-type: none"> <li>• the most commonly used de-icing agent used on aircraft</li> <li>• are diluted with water and heated before application</li> </ul>	<ul style="list-style-type: none"> <li>• are thicker and more viscous than type I fluids and contain a higher percentage of additives compared with type I fluids</li> <li>• are applied at full strength and ambient temperatures</li> </ul>	<ul style="list-style-type: none"> <li>• provide more effective protection against refreezing than Type I agents, but are less effective in depressing the freezing point of water</li> <li>• are thicker and more viscous than type I fluids and contain a higher percentage of additives compared with type I fluids</li> <li>• are applied at full strength and ambient temperatures</li> </ul>	

the accelerated eutrophication of waters.<sup>23</sup> The concentrations of these compounds in airport effluents vary widely, depending on climatic, hydrologic, and ADAF management conditions.<sup>16</sup> Table 3 sets out literature data on the action of de-icing agents on the environment.

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### 3. PROBLEMS AND CHALLENGES POSED BY SAMPLES OF AIRPORT RUNOFF

The analysis of contaminants arising from airport operations comes up against ever newer challenges, especially the need to determine a wide range of analytes in samples with a very complex (and often variable) matrix composition. The development of suitable analytical techniques is driven by ecotoxicological concerns and the desire to describe the state of the environment more accurately.<sup>29</sup>

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The basic problems faced by the analysis of airport runoff concern: the spatially different amounts of runoff (local conditions), the composition of the matrix, the presence of solid particles, and changes in composition during the various phases of the process.<sup>30</sup>

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The objectives of runoff analyses are to ascertain the source, properties, and quantities of the pollutants it contains. Once these characteristics are known, measures for the proper management of airport effluents can be drawn up (reduction in amount, recycling, modification of processes).<sup>3</sup>

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### 4. DETERMINATION OF POLLUTANTS PRESENT IN SAMPLES OF AIRPORT RUNOFF WATERS

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Pollutants emitted from airports can be analyzed by examining samples of runoff (i.e., the rainwater or snowmelt flushing contaminants off the pavement surface). These waters find their way to the soil, surface, and then ground waters, which may be sources of drinking water. Airport runoff waters usually contain large quantities of toxic chemicals and need to be monitored constantly. The analysis of selected contaminants provides a basis for assessing the effects of airport operations on the environment.

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#### 4.1 Sample Collection and Pretreatment

The collection and preparation of samples for analysis is a key stage in every analytical procedure for determining a particular group of constituents.<sup>31</sup> Because any errors committed at this stage will affect the remainder of the procedure, it is very important to ensure that samples are representative.<sup>32</sup> However, such samples may be difficult to collect sample because waters

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**TABLE 3.** Environmental impacts of de-icing agents

Compound	Characteristics	Effect on the environment	Reference(s)
PG	Aircraft de-icing fluid components <ul style="list-style-type: none"> <li>• synthetic clear liquid substances that absorb water</li> <li>• biodegradable (but more slowly than EG)</li> <li>• commonly used in small amounts as a food additive and in cosmetics and certain medicines to absorb moisture</li> <li>• synthetic clear liquid substances that absorb water</li> <li>• biodegradable</li> </ul>	<ul style="list-style-type: none"> <li>• adverse effect on soil and ground waters</li> <li>• less toxic than EG</li> <li>• high BOD</li> <li>• toxicity</li> </ul> <p>✓ 48-hr LC<sub>50</sub> = 43,500 mg/l (<i>Daphnia magna</i>)</p>	[13,14]
EG	<ul style="list-style-type: none"> <li>• aircraft de-icing fluid components</li> <li>• synthetic clear liquid substances that absorb water</li> <li>• biodegradable</li> </ul>	<ul style="list-style-type: none"> <li>• hazardous air pollutant—significant risks to humans and the environment</li> <li>• highly toxic to aquatic mammals and humans</li> <li>• oestrogenic action</li> <li>• 96-hr LC<sub>50</sub> = 27,540 mg/l (<i>Lepomis macrochirus</i>)</li> <li>• 48-hr LC<sub>50</sub> = 46,300–54,700 mg/l (<i>Daphnia magna</i>)</li> <li>• LC<sub>50</sub> 96 hr &gt;18,500 mg/l (Rainbow trout)</li> <li>• Toxicity:</li> </ul> <p>✓ Risk of death following consumption</p> <p>✓ LD<sub>50</sub> = 1,230 mg/kg (rats)</p>	[14–16]
pH buffers (potassium hydroxide)	highly corrosive	No data available	[16]
Water	used to color ADAFs		[16]
Dyes	moistening agent, dispersant, used in very low concentrations	<ul style="list-style-type: none"> <li>• carcinogenic, teratogenic</li> <li>• toxicity</li> </ul>	[16]
1,4-Dioxane		<ul style="list-style-type: none"> <li>• LD<sub>50</sub> = 5,700 mg/kg (mice)</li> <li>• LD<sub>50</sub> = 5,200 mg/kg (rats)</li> </ul>	[16]
Surfactants	reduce surface tension of liquids, improve adherence of liquids to the aircraft surface	May be very toxic toward aquatic organisms	[53]

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**TABLE 3.** Environmental impacts of de-icing agents (*Continued*)

Compound	Characteristics	Effect on the environment	Reference(s)
Triazole	• are generally used to reduce theflammability hazard posed by thecorrosion of metal components	Corrosion inhibitors	[14-17]
• Benzotriazoles	• high reactivity potential	• high biological toxicity	
1H-benzotriazole <sup>a</sup>	• highly reactive with glycols	• 5-methyl-1H-benzotriazole is significantlymore toxic than glycols:	
✓ Tolytriazole <sup>b</sup>		✓ 96-hr LC <sub>50</sub> = 31 mg/l ( <i>Lepomis macrochirus</i> )	
✓ 4-methyl-1Hbenzotriazole <sup>c</sup>		✓ 48-hr LC <sub>50</sub> = 71 mg/l ( <i>Daphnia magna</i> )	
• 5-methyl-1H-benzotriazole <sup>d</sup>	• 1H-benzo-1,2,3-triazole and its methylatedanalogues (tolyltriazole, TTri) arecorrosion inhibitors		
• 1H-benzo-1,2,3-triazole (BTri)	• high polarity		
Sodium nitrite			
Sodium benzoate			
Borax			
Diethylene glycol	• a clear, colourless liquid with theconsistency of syrup	Other freezing-point depressants	[16]
	• reaches higher eutectic temperatures	• toxicity comparable with that of EG, death canfollow swallowing; but not as toxic to humansand mammals	
		• Fathead minnow LC <sub>50</sub> = 75,200 mg/l (96 h)	
		• Oral administration of 1.34 ml diethyleneglycol per kilogram body weight damageskidneys and the central nervous system,causes inflammation of the liver and spleen,coma, and death	
		• Exposure leads to vomiting, nausea, headache,unconsciousness	
Isopropyl alcohol (isopropanol)	Colorless inflammable liquid with a slightsmell resembling that of ethanol andacetone	• Affects the airways, absorbed by the skin	[16]
		• Teratogenic, negative effects on thereproductive system (extreme cases)	

Alkyphenol ethoxylates (APEO)	APEO are low-cost nonionic surfactants that reduce surface tension and are characterized by a relatively low foaming potential	<ul style="list-style-type: none"> <li>potentially toxic to aquatic organisms</li> <li>have been shown to degrade into several by-products, including nonylphenol (NP), NP1EO, NP2EO, octylphenol (OP), OP1EO, and OP2EO, which are more toxic than their parent compounds</li> <li>considered to be endocrine disruptors</li> </ul>	[17,54]
Urea ( $\text{NH}_2\text{CONH}_2$ )	• applied in granular form	<ul style="list-style-type: none"> <li>nontoxic</li> <li>one of its degradation products is ammonia, which is highly toxic to aquatic organisms</li> <li>a source of nitrogen for algae, which may then consume large amounts of oxygen needed by other aquatic animals</li> <li>in humans may cause sneezing, irritation, sore throat, cough</li> </ul>	[14,16]
Ammonia	• product of urea hydrolysis	<ul style="list-style-type: none"> <li><math>\text{LD}_{50}(\text{intravenous}) = 4,600 \text{ mg/kg}</math> (mice)</li> <li>highly toxic to aquatic organisms, even at low concentrations</li> </ul>	[16]
$\text{CO}_2$	• may be converted to ionic forms: $\text{NH}_4^+$ , NO or $\text{N}_2$	<ul style="list-style-type: none"> <li><math>\text{LC}_{50} = 1\text{--}10 \text{ mg/l}</math></li> <li>highly toxic to aquatic organisms, even at low concentrations</li> </ul>	[16]
EG	see above	<ul style="list-style-type: none"> <li>toxicity relatively low</li> <li>may cause irritation of the eyes</li> <li><math>\text{LD}_{50} = 3,250 \text{ mg/kg}</math> (rats)</li> </ul>	[16,17]
Potassium acetate	• may damage the insulation materials used in electrical systems	<ul style="list-style-type: none"> <li>inflammable in combination with corrosion inhibitors</li> <li>corrosive</li> <li><math>\text{BOD}_5 = 140,000\text{--}300,000 \text{ mg/kg}</math></li> <li><math>\text{COD} = 315,000 \text{ mg/kg}</math></li> <li>Solid, granular texture</li> </ul>	[16]
Calcium magnesium acetate (CMA)	• $\text{BOD} = 580,000 \text{ mg/kg}$	<ul style="list-style-type: none"> <li>Relatively non-toxic to the environment</li> <li><math>\text{LD}_{50}</math> (oral) <math>&gt; 5,000 \text{ mg/l}</math></li> <li>may cause irritation on contact</li> <li>can be inhaled</li> </ul>	[16]
Sodium acetate	• $\text{COD} = 780,000 \text{ mg/kg}$	<ul style="list-style-type: none"> <li><math>\text{LD}_{50} = 3,530 \text{ mg/kg}</math> (rats; oral)</li> </ul>	

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**TABLE 3.** Environmental impacts of de-icing agents (*Continued*)

Compound	Characteristics	Effect on the environment	Reference(s)
Sodium formate	<ul style="list-style-type: none"> <li>• Highly biodegradable</li> <li>• BOD = 230,000 mg/kg (highly dependent on temperature)</li> <li>• COD = 242,000 mg/kg</li> </ul>	<ul style="list-style-type: none"> <li>• LD<sub>50</sub> (oral) = 11,200 mg/kg (mice)</li> <li>• LD<sub>50</sub> (intraperitoneal) = 807 mg/kg (mice)</li> <li>Significant exposure to sodium formate de-icer may adversely affect people suffering from chronic complains of the respiratory system, skin and/or eyes</li> </ul>	[16]
Potassium formate	<ul style="list-style-type: none"> <li>• Low BOD during biodegradation (BOD<sub>5</sub> = 40 mg/l)</li> <li>• does not contain toxic corrosion inhibitors</li> </ul>	No data available	
Chlorides	No data available	Alternative pavement de-icers	[14]
Potassium chloride		<ul style="list-style-type: none"> <li>• corrosive, can cause ground and surface water contamination</li> <li>• harmful to vegetation and streams</li> <li>• can damage cars and other manufactured products</li> <li>• that come in contact with them</li> <li>• large doses may adversely affect human health (irritation of the alimentary canal, weakness)</li> </ul>	
Magnesium chloride			
Sodium chloride			
Synthetic fire-fighting surfactants			
Perfluorooctanoate (PFOA)	No data available	<ul style="list-style-type: none"> <li>• significant increase in prostate cancer mortality</li> <li>• PFOA perturbs sex hormone homeostasis</li> <li>• an analogue chemical of PFOA; very similar action</li> </ul>	[55]
Perfluorooctane (PFOS)	No data available		

<sup>a</sup>Other names of 1H-benzotriazole: BT, 1H-BT, BTri, BTAH, BTA, BZT, Bz. <sup>b</sup>Tolytriazole: TT, TTri, MBT, MeBTA, MBz. <sup>c</sup>4-methyl-1H-benzotriazole: 4-MeBT, 4-TTri.  
<sup>d</sup>5-methyl-1H-benzotriazole: 5-MeBT, 5-TTri.

running off airport pavements are rarely homogeneous: their composition  
215 may change radically within the space of a few minutes.

In order to determine the properties of contaminants, initial sample collection  
215 should be frequent (e.g., a few samples should be collected each time pollutants are generated). Thereafter sampling should take place at regular intervals (e.g., every month, every year), to check whether the properties of the runoff waters have changed or not. Flow-proportional samples are recommended where applicable.<sup>3</sup>

All samples collected should be preserved in the proper manner according to the latest Standard Methods for the Examination of Water and Wastewater and thoroughly completed by the sampler.<sup>3</sup>

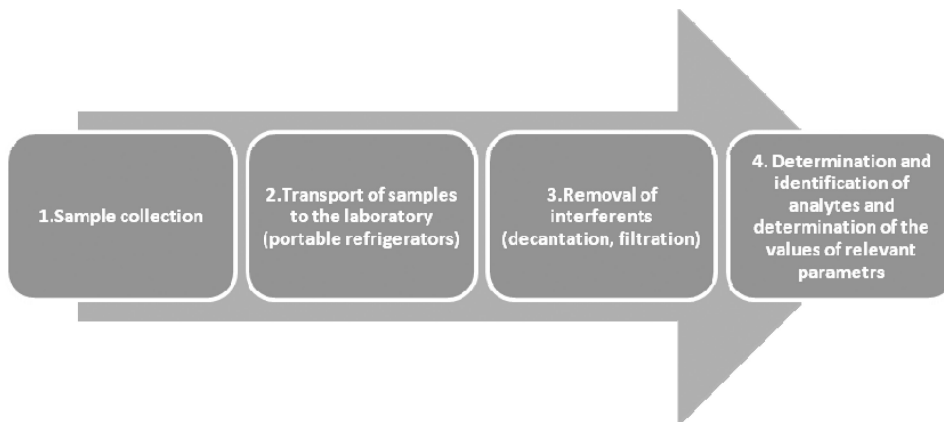
## 225 4.2 Collection and Preparation of Runoff Water Samples for Analysis

The usual means of collecting samples of runoff water is to install an appropriate drainage system. Then samples of runoff water can be scooped from a specially prepared erosion ditch and then taken to the laboratory.<sup>32</sup>

Another solution is to place the sampling container (a glass bottle) in an earthen embankment.<sup>33,34</sup> The bottle is placed in a plastic container with a metal cover to prevent damage and also to prevent rainwater from getting into the bottle. The bottle's neck should protrude 2 cm above the soil surface to prevent insects from falling into it.<sup>33</sup>

Runoff can also be sampled using fully automated equipment.<sup>35</sup> Apart from the actual sample collection, a number of parameters—the intensity of runoff, its temperature, conductivity, and pH—can all be measured using the system described in Gryniewicz et al.<sup>32</sup> The main part of this setup is a 10-l glass vessel connected by means of polyethylene tubing to measurement instruments operating on-line. The whole system is placed inside an erosion pipe lying 0.5 m deep in the soil. Runoff samples are pumped out of the glass bottle to other, smaller vessels (made from glass for samples to be analyzed for organic compounds and from polyethylene for samples analyzed for inorganic compounds) and are then analyzed for target compounds. Automated sampling systems such as this one are complex and expensive, which is why they are rarely used in practice.<sup>32</sup> The simplest and cheapest way of sampling runoff waters is hand sampling using hermetic polyethylene syringes fitted with rubber hoses. These samples are taken from drainage ditches, drainage installations, or places where water naturally collects and poured into sealable glass or plastic containers. Before use, every container and syringe is washed, and before sample collection flushed with the water to be sampled.<sup>16</sup>

Environmental samples are a very heterogeneous research material because of the different sampling sites, types of matrix, and the possible spectrum of analytes. The preparation of environmental samples for



**FIGURE 2.** The steps involved in preparing samples of airport runoff water for analysis.

analysis is thus a complicated process. The operations carried out at this stage may cause loss of analytes and may also be a source of additional contamination.<sup>36</sup>

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Samples should be stored at a low temperature (around 4°C). At the same time, they should be taken to the laboratory and analyzed as soon as possible after sampling. If this is not possible, appropriate measures should be taken to ensure sample stability during transport and storage.<sup>16</sup> If samples are highly contaminated with solid particles/bodies (sand, leaves, insects), they must be filtered or the liquid decanted before further analysis can be embarked upon.<sup>37</sup> Figure 2 shows a scheme for preparing samples for the determination of target analytes and the relevant parameters.

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#### 4.3 Techniques for Determining the Physicochemical Parameters and Analyte Contents in Airport Runoff Samples

In the runoff water samples collected from airport areas the following physicochemical parameters were determined: pH and conductivity<sup>16</sup>; summary parameters ( $BOD_5$ ,<sup>17</sup> COD,<sup>38</sup> total organic carbon,<sup>39</sup> total suspended solids,<sup>39</sup> total Kjeldahl nitrogen<sup>40</sup>); and different chemicals such as phosphorus (all forms),<sup>41</sup> petroleum hydrocarbons,<sup>42</sup> semivolatile organic compounds<sup>43</sup> (fats, oils, and grease<sup>44</sup>), de-icing agents (glycols,<sup>16,45</sup> nonylphenol ethoxylates,<sup>46</sup> benzotriazoles,<sup>17</sup> alkylphenol ethoxylates<sup>17</sup>), and metals.<sup>17,47,48</sup>

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The following techniques were used to determine the previously mentioned parameters: gas chromatography,<sup>16</sup> liquid chromatography,<sup>49</sup> gel permeation chromatography,<sup>43</sup> electrochemical technique,<sup>16</sup> spectrophotometry,<sup>42</sup> gravimetric techniques,<sup>44</sup> and potentiometric techniques.<sup>40</sup>

280        5. LITERATURE INFORMATION ON XENOBIOTICS DETERMINED  
IN SAMPLES OF AIRPORT RUNOFF WATERS

285        The world literature boasts a large number of articles dealing with the study of contaminants discharged at airports. The majority of them focus on the pollutants generated by the washing and cleaning of aircraft and airport pavements. Most (around 70%) of these studies were carried out in the United States over the last 15 years or so, their objective being to record the changes taking place in the environment when contaminants emitted by airport operations get into it. The following constituents/parameters of airport runoff waters were determined: ethylene glycol, propylene glycol, BOD (5-day), COD, total organic carbon, total nitrogen, total suspended solids, total Kjeldahl nitrogen, metals (magnesium, copper, lead, calcium, sodium, potassium, iron, manganese), and hydrocarbons.

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Table 4 reviews the results of analyses of runoff waters from airports in different parts of the world.

295        6. THE TOXICITY OF AIRPORT RUNOFF WATER SAMPLES

295        In tandem with the chemical analysis of airport runoff waters, their toxicity is also evaluated. Tests using biological material are crucial, as they constitute the basis for assessing the overall degree of contamination of particular compartments of the environment.<sup>29</sup>

300        6.1 Assessment of Toxic Effects

300        Determining the toxicity of runoff samples may provide solutions to a number of basic problems, for example, hazard assessment (the probability of a factor acting destructively on living organisms), evaluating the scale of toxicity (determining the dose eliciting a toxic effect), and estimating the long-term effects of exposure of organisms to toxic substances (mutagenic and/or carcinogenic effects, compounds with embryotoxic and/or teratogenic properties).<sup>29</sup>

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310        The quantitative measure of an analyte's toxicity is given by the values of indices determined on a dose-response basis (Table 5). This relationship is expressed by (a) effective concentration (EC) 25 or EC50 or (b) effective dose (ED) 25 or ED50. They define the concentration of a toxin in the environment or the maximum dose of it that elicits a given biological effect in 25% or 50% of the organisms tested. Inhibition concentration (IC) 50 or IC25 defines the concentration of an environmental toxin that impairs (inhibits) by 50% or 25% a given process (e.g., growth).

315        The lethal concentration (LC) 50 parameter presents the dose causing the death of 50% of individuals in a population after a given time. Often,

**TABLE 4.** Literature information on the analytes of samples of runoff water from airports in different parts of the world

Parameter/ Analyte	Sampling location	Range of concentration (mg/l; unless otherwise noted)	Mean value (mg/l)	Type of flow	Water flow (l/s)	Mass yield <sup>a</sup> (mg/l)	Sampling year	Reference(s)
COD	Airport, Gdańsk, Poland	runways, internal roads, aprons, car park, passenger terminal	31–807 32–202 45–287 44–177 38–145	— Storm water runoff and liquid drainage	401	—	1999 2000 2001 2002 2003	[23]
	commuter terminal	89	—			35,689	1999	
		227				91,027	2000	
		83				33,283	2001	
		42				16,842	2002	
		60	47.0	Liquid drainage	NA <sup>b</sup>	24,060	2003	
		8.0–150.0			—	—	September 1994	[57]
	Heathrow International Airport London, England						September 1995	
	Newark International Airport, New York, USA	49–338	189	Storm water	NA	—	—	[16]
	Salt Lake City International Airport, USA	104–3,880	835	Liquid drainage	NA	—	Winter 1997–1998	[16]
	Baltimore–Washington International Airport, USA	main terminal area <sup>c</sup> 11,000–270,000 700–2,700	—	Liquid drainage	NA	—	Winter 1997–1998	[16]
	Dallas/Fort Worth International Airport, Texas, USA	commuter terminal <sup>c</sup> upstream reference 14–80	—	Rainfall	NA	—	1993	[58]
		10–37,900 <9–1,600	—	Liquid drainage	NA	—	November 2002–April 2004	[26]
		47–84	26	Rainfall	NA	—	January 29, 2001–March 16, 2003	[46,54]
	Sites near General Mitchell International Airport, Milwaukee, USA	upstream site primary outfall receiving stream	5,600–10,200 330–480 <20–130	Drainage liquid Drainage liquid Effluent from treatment basin	NA	—	—	[16]
	Nashville International Airport, USA						Winter 1997–1998	

BOD <sub>5</sub>	Airport, Gdańsk, Poland	runways, internal roads, aprons, car park, passenger terminal	4.1–130.0 5.6–30.9 7.7–18.9 7.0–10.3 4.1–26.6	—	Storm water runoff and liquid drainage	401	—	1999 2000 2001 2002 2003
	commuter terminal	—	15.5	—	6215.5	1999	[23]	
			47.9	—	19207.9	2000		
			8.1	—	3248.1	2001		
			7.5	—	3007.5	2002		
			6.9	—	2766.9	2003		
Westchester County Airport, New York, USA	taxiway, drainage system	buildings, hangars	2–8.4	2.53	Storm water from taxiways and ditch drainage	NA	—	Winter 1997–1998
		ponds	2–37	4.9	Storm water from buildings and hangars	NA	—	[16]
Salt Lake City International Airport, USA	main terminal area	11–1,050	2–7.2	2.8	Storm water from ponds	NA	—	Winter 1997–1998
Baltimore–Washington International Airport, USA	commuter terminal upstream reference airport drainage receiving stream	23–2,510	1,010	332	Liquid drainage	NA	—	[16]
Dallas/Fort Worth International Airport, Texas, USA	upstream site primary outfall receiving stream	197–769 2–3.1 93–9,500 <2 to >100	412	1,010	Storm water runoff	NA	—	Winter 1997–1998
Sites near General Mitchell International Airport, Milwaukee, USA	upstream site primary outfall receiving stream	—	2.5	412	Liquid drainage	NA	—	[16]
Kansas City International Airport, USA	—	738	—	2.5	Rainfall	NA	—	November 1997–1998
Bradley International Airport, USA	—	—	—	>20	—	NA	—	[26]
Greater Rockford International Airport, USA	—	>7.3	—	16.6	Liquid drainage	NA	—	April 2002–April 2004
Louisville International Airport, USA	—	3–1250	—	201	Drainage liquid	NA	—	January 29, 2001–March 2003
Nashville International Airport, USA	—	3–98	—	5,100	Drainage liquid	NA	—	[16]
				39,000	Storm water	NA	—	Winter 1997–1998
					Liquid drainage	12.6 <sup>d</sup>	491,400	Winter 1997–1998
					—	NA	—	[16]
					—	NA	—	Winter 1997–1998
					—	NA	—	[16]
					—	NA	—	Winter 1997–1998
					—	NA	—	[16]
					Effluent from treatment basin	NA	—	Winter 1997–1998

(Continued on next page)

**TABLE 4.** Literature information on the analytes of samples of runoff water from airports in different parts of the world (*Continued*)

Parameter/ Analyte	Sampling location	Range of concentration (mg/l; unless otherwise noted)	Mean value (mg/l)	Type of flow	Water flow (l/s)	Mass yield <sup>a</sup> (mg/l)	Sampling year	Reference(s)
BOD	Baltimore—Washington International Airport, USA; commuter terminal <sup>c</sup> Heathrow International Airport London, England	3–64 138–1,900 0–18.4	— — 5.54	Liquid drainage Liquid drainage	NA NA	— —	1993 September 1994– 1995	[58] [61]
Dissolved oxygen (DO)	Muddy Bridge Branch (receives runoff directly from Baltimore—Washington International Airport), USA Louisville International Airport, USA Nashville International Airport, USA	— 0.270–13.0 6.4–11.9	4.5 6.4 —	Liquid drainage Storm water runoff Effluent from treatment basin	450 270 NA	2,025 1,728	March 1991 April 1991	[64]
TOC	International Airport, Warsaw, Poland Newark International Airport, runway terminal New York, USA Kansas City International Airport, USA Bradley International Airport, USA	— 7–1120 9–23 —	446.5 83.5 16 3,000	Storm water sewage Storm water Storm water	NA NA NA NA	— — — —	Winter 1997–1998 Winter 1997–1998 2002 1997–1998	[16] [16] [58] [16]
Total nitrogen-(TN)	Greater Rockford International Airport, USA International Airport, Warsaw, Poland Airport, Gdańsk, Poland	— — runways, internal roads, aprons, car park, passenger terminal	12 142.1 3.1–158.50 3.4–69.00 3.27–11.00 2.50–3.21 2.80–17.90	Storm water Storm water sewage	NA NA 401	44,100 44,100 — — —	Winter 1997–1998 Winter 1997–1998 1999 2000 2001 2002 2003	[16] [16] [62] [23]

	commuter terminal	—	7.10 4.60 2.40 2.21 2.91	2847.1 1844.6 962.4 886.21 1166.91	1999 2000 2001 2002 2003
	main terminal area commuter terminal	2–27 ND-26	— Storm water runoff Liquid drainage Liquid drainage	— NA	— Winter Winter 1997–1998 1993
Baltimore–Washington International Airport, USA	Newark International Airport, runway terminal	2–3 <2–64	2.25 <12.5	NA	— — 1997–1998 [16]
TSS [mg/l]	Baltimore–Washington International Airport, USA commuter terminal <sup>c</sup> Louisville International Airport, USA	3–38 11–31 14–31 2–3,530	11.3 — — Storm water from runway terminal Liquid drainage Storm water runoff	NA NA NA	— — 1997–1998 [16]
	Nashville International Airport, USA	18–55	— Effluent from treatment basin	NA	— —
Ammonia-NH <sub>3</sub> [mg/l]	Baltimore–Washington International Airport, USA commuter terminal <sup>c</sup> Muddy Bridge Branch (receives runoff directly from Baltimore–Washington International Airport), USA	ND-23 ND-5 —	— — 0.003	Liquid drainage Liquid drainage Liquid drainage	— — 450
	Kansas City International Airport, USA	—	3.9	Storm water	11.61 April 1991 — Winter
	Bradley International Airport, USA	—	23	Liquid drainage	12.6 <sup>d</sup> 289.8 Winter 1997–1998
	Louisville International Airport, USA	<0.03–171	— 46	Storm water runoff Storm water Liquid drainage	— — — 1997–1998 1993
Total phosphorus- P [mg/l]	Greater Rockford International Airport, USA Baltimore–Washington International Airport, USA	0.2–60 ND-6	— —	NA NA	— — 1997–1998 [16] [16]
	main terminal area <sup>b</sup> commuter terminal <sup>b</sup>				(Continued on next page)

**TABLE 4.** Literature information on the analytes of samples of runoff water from airports in different parts of the world (*Continued*)

Parameter/ Analyte	Sampling location	Range of concentration (mg/l; unless otherwise noted)	Mean value (mg/l)	Type of flow	Water flow (l/s)	Mass yield <sup>a</sup> (mg/l)	Sampling year	Reference(s)
Fe	Fornebu International Airport, Oslo, Norway (sampling done 2 years after airport closure)	aircraft de-icing point	0.2–6.5	Malats [µg/l]	Liquid drainage	100,000 m <sup>3</sup> / year <sup>f</sup>	—	2000
Cd	Baltimore–Washington International Airport, USA	main terminal area <sup>c</sup> commuter terminal <sup>c</sup>	ND-3 ND-1	—	Liquid drainage	NA	—	1993 [58]
Cr	Baltimore–Washington International Airport, USA	main terminal area <sup>c</sup> commuter terminal <sup>c</sup>	ND-1 ND	—	Liquid drainage	NA	—	1993 [58]
Cu	Baltimore–Washington International Airport, USA	main terminal area <sup>c</sup> commuter terminal <sup>c</sup>	ND-45 3-9	—	Liquid drainage	NA	—	1993 [58]
	Kansas City International Airport, USA					14	Storm water	NA
	Bradley International Airport, USA					44	Liquid drainage	12.6 <sup>d</sup>
	Greater Rockford International Airport, USA					9.2	Storm water	NA
Pb	Baltimore–Washington International Airport, USA	main terminal area <sup>c</sup>	ND-16	—	Liquid drainage	NA	—	1993 [58]
	Kansas City International Airport, USA	commuter terminal <sup>c</sup>	1-11	—	Storm water	NA	—	Winter 1997–1998 [16]
	Bradley International Airport, USA		—	15	Storm water	NA	—	Winter 1997–1998 [16]
	Greater Rockford International Airport, USA		—	50	Liquid drainage	12.6 <sup>d</sup>	630	Winter 1997–1998 [16]
			4.3	Storm water	NA	—	—	Winter 1997–1998 [16]

Ni	Baltimore-Washington International Airport, USA	main terminal area <sup>c</sup> commuter terminal <sup>c</sup>	ND-5 5	—	Liquid drainage	NA	—	1993	[58]
Zn	Baltimore-Washington International Airport, USA	main terminal area <sup>c</sup> commuter terminal <sup>c</sup>	240-1,430 35-60	—	Liquid drainage	NA	—	1993	[58]
	Kansas City International Airport, USA	—	140	Storm water	NA	—	Winter 1997-1998	[16]	[16]
	Bradley International Airport, USA	—	340	Liquid drainage	12.6 <sup>d</sup>	4,284	Winter 1997-1998	[16]	[16]
	Greater Rockford International Airport, USA	—	45	Storm water	NA	—	Winter 1997-1998	[16]	[16]
Potassium-K	Sites near General Mitchell upstream site, International Airport, Milwaukee, USA	—	6,25	Drainage liquid	17	106,25	2000-2007	[67]	[67]
	Primary outfall receiving stream	—	59,1	Drainage liquid	62	2,664			
	Kansas City International Airport, USA	—	15,5	Drainage liquid	309	4,789			
	—	13,000	Storm water	NA	—		Winter 1997-1998	[16]	[16]
	ND (900)	—	ND (900)	Liquid drainage	12.6 <sup>d</sup>	—	Winter 1997-1998	[16]	[16]
	Bradley International Airport, USA	—	64,000	Storm water	NA	—	Winter 1997-1998	[16]	[16]
	Greater Rockford International Airport, USA	—	860	Storm water	NA	—	Winter 1997-1998	[16]	[16]
Aluminum-Al	Kansas City International Airport, USA	—	1,100	Liquid drainage	12.6 <sup>d</sup>	13,860	Winter 1997-1998	[16]	[16]
	Bradley International Airport, USA	—	270	Storm water	NA	—	Winter 1997-1998	[16]	[16]
	Greater Rockford International Airport, USA	—	2,500	Storm water	NA	—	Winter 1997-1998	[16]	[16]
Magnesium-Mg	Kansas City International Airport, USA	—	2,000	Liquid drainage	12.6 <sup>d,e</sup>	25,200	Winter 1997-1998	[16]	[16]
	Bradley International Airport, USA	—	3,000	Storm water	NA	—	Winter 1997-1998	[16]	[16]
	Greater Rockford International Airport, USA	—	170	Storm water	NA	—	Winter 1997-1998	[16]	[16]
Manganese-Mn	Kansas City International Airport, USA	—							

(Continued on next page)

**TABLE 4.** Literature information on the analytes of samples of runoff water from airports in different parts of the world (*Continued*)

Parameter/ Analyte	Sampling location	Range of concentration (mg/l; unless otherwise noted)	Mean value (mg/l)	Type of flow	Water flow (l/s)	Mass yield <sup>a</sup> (mg/l)	Sampling year	Reference(s)
Sodium-Na	Bradley International Airport, USA	—	140	Liquid drainage	12.6 <sup>d</sup>	1,764	Winter 1997-1998	[16]
	Greater Rockford International Airport, USA	—	360	Storm water	NA	—	Winter 1997-1998	[16]
	Kansas City International Airport, USA	—	1,100	Storm water	NA	—	Winter 1997-1998	[16]
	Bradley International Airport, USA	—	10,000	Liquid drainage	12.6 <sup>d</sup>	126,000	Winter 1997-1998	[16]
	Greater Rockford International Airport, USA	—	7,900	Storm water	NA	—	Winter 1997-1998	[16]
Calcium-Ca	Kansas City International Airport, USA	—	3,400	Storm water	NA	—	Winter 1997-1998	[16]
	Bradley International Airport, USA	—	33,000	Liquid drainage	12.6 <sup>d</sup>	415,800	Winter 1997-1998	[16]
	Greater Rockford International Airport, USA	—	14,000	Storm water	NA	—	Winter 1997-1998	[16]
Crude oil extract	International Airport, Warsaw, Poland	Specific organics	26.4	Storm water	NA	—	2002	[62]
Petroleum ether extract	Airport, Gdańsk, Poland	14.4-553.7 11.2-40.6 7.6-26.1 11.5-13.4 5.6-10.4	—	Storm water runoff and liquid drainage	401	—	1999 2000 2001 2002 2003	[23]
	commuter terminal	—	19.1 208.5 7.7 8.1 6.1		7,659 83,609 3,088 3,248 2,446	1999 2000 2001 2002 2003		

Oil and grease	Westchester County Airport, New York, USA	taxiway, drainage system	5	5	Storm water from taxiways and ditch drainage	NA	—	Winter 1997–1998	[16]
	buildings, hangars		5	5	Storm water from buildings and hangars				
	ponds		5	5	Storm water from ponds				
Oil	Baltimore-Washington International Airport, Gdańsk, Poland	main terminal area <sup>c</sup> commuter terminal <sup>c</sup> runways, internal roads, aprons, car park, passenger terminal	ND-29 ND-14 16.2–356.6 11.2–161.8 9.2–25.5 13.3–20.6 9.6–10.2	— — — — — — —	Liquid drainage Storm water runoff and liquid drainage	NA 401	— —	1993 1999 2000 2001 2002 2003 2010	[58] [23]
Glycols	Westchester County Airport, New York, USA	aircraft de-icing point drainage ditch	—	<1	Liquid drainage	100,000 m <sup>3</sup> /year <sup>f</sup>	—	2000	[14]
Propylene glycol	Førnebu International Airport, Oslo, Norway (sampling done 2 years after airport closure)	aircraft de-icing point drainage ditch	—	<2					
	Westchester County Airport, New York, USA	taxiway, drainage system	0.05–1.3	0.213	Storm water from taxiways and ditch drainage	NA	—	1997–1998	[16]
	buildings, hangars		0.05–0.82 0.05–220	0.13 32.8 <50	Storm water Liquid drainage				
Muddy Bridge Branch (receives runoff directly from Baltimore-Washington International Airport), USA	Dallas/Fort Worth International Airport, Texas, USA	upstream reference	<18	—	Rainfall	NA	—	November 2002–April 2004	[26]

(Continued on next page)

**TABLE 4.** Literature information on the analytes of samples of runoff water from airports in different parts of the world (*Continued*)

Parameter/ Analyte	Sampling location	Range of concentration (mg/l; unless otherwise noted)	Mean value (mg/l)	Type of flow	Water flow (l/s)	Mass yield <sup>a</sup> (mg/l)	Sampling year	Reference(s)
Sites near General Mitchell International Airport, Milwaukee, USA	airport drainage receiving stream upstream site primary outfall	<18–3,800 <18 1,900–4,400 130–150	<18 — 150	Liquid drainage Rainfall Liquid drainage Liquid drainage	NA	—	January 29, 2001–March 16, 2003	[46,54]
Kansas City International Airport, USA	receiving stream	—	16,000	Storm water	NA	—	Winter 1997–1998	[16]
Bradley International Airport, USA	—	160,000	Liquid drainage	12.6 <sup>d</sup>	2,016 × 10 <sup>6</sup>	Winter 1997–1998	[16]	
Greater Rockford International Airport, USA	—	ND (5.0)	Storm water	NA	—	Winter 1997–1998	[16]	
Ethylene glycol	Baltimore–Washington International Airport, USA main terminal area commuter terminal Muddy Bridge Branch (receives runoff directly from Baltimore–Washington International Airport), USA	<10 — — —	<10 2,100 <25 <sup>c</sup>	Storm water runoff Liquid drainage Liquid drainage	NA 450 270	— 945 —	Winter 1997–1998 March 1991 April 1991	[16]
Dallas/Fort Worth International Airport, Texas, USA	upstream reference airport drainage receiving stream	<18 <18–20,000 <18	— — —	Rainfall Liquid drainage Rainfall	NA	—	November 2002–April 2004	[26]
Sites near General Mitchell International Airport, Milwaukee, USA	upstream site primary outfall	32–280	—	Drainage liquid Drainage liquid	NA	—	January 29, 2001–March 16, 2030	[46,54]
Kansas City International Airport, USA	receiving stream	—	<18	Drainage liquid	NA	—	Winter 1997–1998	[16]
	—	3,200	3,200	Storm water	NA	—		

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Bradley International Airport, USA	—	3,000	Liquid drainage	12.6 <sup>d</sup> [16]	
Greater Rockford International Airport, USA	—	ND (10)	Storm water	NA — Winter 1997–1998 [16]	
Kansas City International Airport, USA	—	>20 000	Storm water	NA — Winter 1997–1998 [16]	
Diethylene glycol	Bradley International Airport, USA	—	15,000	Liquid drainage	12.6 <sup>d</sup> [16]
Greater Rockford International Airport, USA	—	ND (5.0)	Storm water	NA — Winter 1997–1998 [16]	
Dissolved organic carbon (DOC)	Fonnebu International Airport, aircraft de-icing point Oslo, Norway (sampling point done 2 years after airport closure)	21–79	—	Liquid drainage 100,000 m <sup>3</sup> /year <sup>f</sup>	— 2000 [14]
Hydrocarbons	Newark International Airport,runway terminal New York, USA	<0.4–8.8	<1.98	Storm water from runway terminal	NA — Winter 1997–1998 [16]
Baltimore–Washington International Airport, USA	main terminal area <sup>c</sup> commuter terminal <sup>c</sup>	ND 1	— 1	Liquid drainage	NA — Winter 1993–1998 [16]
Acetate	Sites near General Mitchell International Airport, Milwaukee, USA	>5.0 —	—	Drainage liquid	17 — 2005–2007 [67]
	upstream site primary outfall receiving stream	—	120	Drainage liquid	62 7,440 [67]
Formate	Sites near General Mitchell International Airport, Milwaukee, USA	>2.5 —	8.75	Drainage liquid	309 2,703 — 2005–2007 [67]
	upstream site primary outfall receiving stream	—	—	Drainage liquid	17 682 — 2005–2007 [67]
Phenol <sup>e</sup>	Kansas City International Airport, USA	—	<2.5	Drainage liquid	62 — Winter 1997–1998 [16]
	Bradley International Airport, USA	—	93	Storm water	309 NA — Winter 1997–1998 [16]
	Greater Rockford International Airport, USA	—	340	Liquid drainage	12.6 <sup>d</sup> 4,284 Winter 1997–1998 [16]
	ND (10) Storm water	—	ND (10)	Storm water	NA — Winter 1997–1998 [16]

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**TABLE 4.** Literature information on the analytes of samples of runoff water from airports in different parts of the world (*Continued*)

Parameter/ Analyte	Sampling location	Range of concentration (mg/l; unless otherwise noted)	Mean value (mg/l)	Type of flow	Water flow (l/s)	Mass yield <sup>a</sup> (mg/l)	Sampling year	Reference(s)
n-tetradecone <sup>e</sup>	Kansas City International Airport, USA	—	ND (10)	Storm water	NA	—	Winter 1997–1998	[16]
	Bradley International Airport, USA	—	140	Liquid drainage	12.6 <sup>d</sup>	1,764	Winter 1997–1998	[16]
	Greater Rockford International Airport, USA	—	ND (10)	Storm water	NA	—	Winter 1997–1998	[16]
Benzene <sup>e</sup>	Louisville International Airport, USA	<5–97	—	Storm water runoff	NA	—	1997–1998	[16]
Ethylbenzene <sup>e</sup>	Louisville International Airport, USA	5–127	—	Storm water runoff	NA	—	Winter 1997–1998	[16]
Naphthalene <sup>e</sup>	Louisville International Airport, USA	5–361	—	Storm water runoff	NA	—	Winter 1997–1998	[16]
Toluene <sup>e</sup>	Louisville International Airport, USA	<5	—	Storm water runoff	NA	—	Winter 1997–1998	[16]
Xylene <sup>e</sup>	Louisville International Airport, USA	<5	—	Storm water runoff	NA	—	Winter 1997–1998	[16]
Benzotriazole <sup>e</sup>	Glatt River near Zurich International Airport, Switzerland	0.16–5.44	—	Surface water	7000m <sup>3</sup> /f	—	Winter 2003/2004	[59]
	Forsenbu International Airport, aircraft de-icing Oslo, Norway (sampling point done 2 years after airport closure)	1.2–1100	—	Liquid drainage	100,000 m <sup>3</sup> / year <sup>g</sup>	—	2000	[14]
4-MeBT <sup>e</sup>	Sites near General Mitchell International Airport, Milwaukee, USA	<0.08 <0.08–0.6	—	Liquid drainage	NA	—	29.01. 2001– 16.03.2003	[46,54]
5-MeBT <sup>d</sup>	Sites near General Mitchell International Airport, Milwaukee, USA	<0.08 <0.08–0.8	—	Liquid drainage	17	—	2005–2007	[67]
		—	<0.08	Liquid drainage	NA	—	29.01.2001 29.01.2001– 16.03.2003	[46] [46,54]

	Kansas City International Airport, USA	—	17,000	Storm water	NA	—	Winter 1997–1998	[16]
	Bradley International Airport, USA	—	90,000	Liquid drainage	12.6 <sup>d</sup>	$1,134 \times 10^6$ Winter	1997–1998	[16]
	Greater Rockford International Airport, USA	—	120	Storm water	NA	—	Winter 1997–1998	[16]
Totyltriazole <sup>e</sup>	Glatt River near Zurich International Airport, Switzerland	0.04–0.91	—	Surface water	7000m <sup>3</sup> /f	—	Winter 2003/2004	[63]
Nonylphenol ethoxylates-NpnEO <sub>e</sub>	Sites near General Mitchell International Airport, Milwaukee, USA	—	776	Liquid drainage	NA	—	January 29, 2001–March 16, 2003	[46,54]
	Inorganics	0.9–9	4.73	Liquid drainage	NA	—	Winter 1997–1998	[16]
Nitrate/Nitrite SO <sub>4</sub> <sup>2–</sup>	Salt Lake City International Airport, USA	87–1600	—	Liquid drainage	100,000 m <sup>3</sup> / year <sup>g</sup>	—	2000	[14]
Conductivity [μS/cm]	Fornebu International Airport, Oslo, Norway (sampling done 2 years after airport closure)	347–890	628	Liquid drainage	NA	—	September 1994– September 1995	[61]
	Heathrow International Airport London, England	120–540 120–2000 —	—	Liquid drainage	NA	—	1993	[58]
	Baltimore–Washington International Airport, USA (commuter terminal)	120–540	—	Liquid drainage	450 270	2,025 1,728	March 1991	[64]
	Muddy Bridge Branch (receives runoff directly from Baltimore–Washington International Airport), USA	210.5	—	Liquid drainage	—	—	—	—
pH	Heathrow International Airport London, England	6.4–8.1	328.0 7.2	Liquid drainage	NA	—	April 1991 September 1994– September 1995	[61]
	Westchester County Airport, taxiway, drainage system New York, USA	6–8	—	Storm water from taxiways and ditch drainage	NA	—	Winter 1997–1998	[16]
	buildings, hangars	6.3–8.8	—	Storm water from buildings and hangars	—	—	—	—

(Continued on next page)

**TABLE 4.** Literature information on the analytes of samples of runoff water from airports in different parts of the world (*Continued*)

Parameter/ Analyte	Sampling location	Range of concentration (mg/l; unless otherwise noted)	Mean value (mg/l)	Type of flow	Water flow (l/s)	Mass yield <sup>a</sup> (mg/l)	Sampling year	Reference(s)
	ponds	6.9–8.6		Storm water from ponds				
Newark International Airport, runway New York, USA	terminal	5.1–7.5	—	Storm water	NA	—	Winter 1997–1998	[16]
Salt Lake City International Airport, USA	main terminal area	6.1–7 6.6–9.5	—	Liquid drainage	NA	—	Winter 1997–1998	[16]
Baltimore–Washington International Airport, USA	commuter terminal Muddy Bridge Branch (receives runoff directly from Baltimore–Washington International Airport), USA	6.7–7.5	—			—	Winter 1997–1998	[16]
		6.7–7.1	7.0	Liquid drainage	450	945	March 1991	[64]
Louisville International Airport, USA		7–9.1	7.4	Liquid drainage	270	—	April 1991 Winter 1997–1998	[16]
Nashville International Airport, USA		7.2–8.6	—	Storm water runoff	NA	—	—	[16]
				Effluent from treatment basin	NA	—	Winter 1997–1998	[16]
Alkalinity	Baltimore–Washington International Airport, USA	main terminal area commuter terminal Muddy Bridge Branch (receives runoff directly from Baltimore–Washington International Airport), USA	6.7–7.5 30–158 69.0	—	Liquid drainage	NA	—	1993
						450	945	March 1991
Hardness	Baltimore–Washington International Airport, USA	main terminal area commuter terminal	84.2 28–800 46–168	—	Liquid drainage	270	—	April 1991 1993
						NA	—	[58]

<sup>a</sup>Mass yield = flow\*concentration. <sup>b</sup>NA = Data not available. <sup>c</sup>Data sample type: peak, composite, and grab. <sup>d</sup>Maximum flow rate. <sup>e</sup>Concentration in sample [μg/l].  
<sup>f</sup>Only data about flow of surface water. <sup>g</sup>Approximate total amount of storm water for one year.

**TABLE 5.** Literature data on acute and chronic toxicity information on aircraft de-icing and anti-icing fluids as pure compounds

Organism	Test type/ Parameter	Literature review of mean values of toxicity indices	Reference
Ethylene glycol			
<i>Pimephales promelas</i>	Acute toxicity tests: • 48-hr LC <sub>50</sub>	81,950 mg/l 9.82 ml glycol/l	[59] [60]
	• LC <sub>50</sub> <sup>a</sup>	24,700 mg/l (Type I <sup>b</sup> ) 371 mg/l (Type IV <sup>c</sup> )	[26]
	Acute toxicity tests: • 96-hr LC <sub>50</sub>	72,860 mg/l 9.82 ml glycol/l	[57] [64]
	• 7-day LC <sub>50</sub>	49,000–57,000 <sup>d</sup> mg/l	[61]
	• NOAEC	9.82 ml glycol/l 39,140 mg/l	[64] [57]
	Chronic toxicity test: • NOEC	32,000	[57]
	✓ Mortality	15,380 mg/l	
	✓ Reprod. or growth	22,520 mg/l	
	• IC <sub>25</sub>	4,400 mg/l (Type I)	[26]
	• IC <sub>25</sub>	179 mg/l (Type IV)	
<i>Ceriodaphnia dubia</i>	Acute toxicity tests: • 48-h LC <sub>50</sub>	34,440 mg/l	[57]
	• LC <sub>50</sub>	15,700 mg/l (Type I) 449 mg/l (Type IV)	[26]
	• NOAEC	24,000 mg/l	[57]
	• 48-hr EC <sub>50</sub>	12.85 ml glycol/l	[64]
	• 96-hr EC <sub>50</sub>	8.95 ml glycol/l	
	• 7-day LC <sub>50</sub>	3.02 ml glycol/l	
	Chronic toxicity test: • NOEC :	24,000	[57]
	✓ Mortality	8,590 mg/l	
	✓ Reprod. or growth	12,310 mg/l	
	• IC <sub>25</sub>	5,500 mg/l (Type I)	[26]
	• IC <sub>25</sub>	113 mg/l (Type IV)	
<i>Daphnia magna</i>	Acute toxicity tests: • <48-hr neonate; 48-hr EC <sub>50</sub> (s, m) <sup>e</sup>	50,450 mg/l	[62]
	• <24-hr-neonate; 48-hr LC <sub>50</sub> (s, n) <sup>e</sup> 3 different samples	46,300–51,100 mg/l	[58]
	• Age not reported; 48-hr LC <sub>50</sub> (s, n)	>10,000 mg/l	[58]
<i>Daphnia pulex</i>	Acute toxicity tests: • 48-hr LC <sub>50</sub>	8.44 ml glycol/l	[64]
	• 96-hr LC <sub>50</sub>	4.25 ml glycol/l	
	• EC <sub>50</sub>	11,800 mg/l (Type I) 7,700 mg/l (Type IV)	[26]
<i>Vibrio fischeri</i>	Chronic toxicity tests	4,600 mg/l (Type I) 1,430 mg/l (Type IV)	[29]
<i>Selenastrum capricornutum</i>	• IC <sub>25</sub>		

(Continued on next page)

**TABLE 5.** Literature data on acute and chronic toxicity information on aircraft de-icing and anti-icing fluids as pure compounds (*Continued*)

Organism	Test type/ Parameter	Literature review of mean values of toxicity indices	Reference
Propylene glycol			
<i>Ceriodaphnia dubia</i>	Acute toxicity tests:		[57]
	<ul style="list-style-type: none"> <li>• 48-hr LC<sub>50</sub></li> <li>• 48-hr EC<sub>50</sub></li> <li>• 96-hr EC<sub>50</sub></li> <li>• 7-day EC<sub>50</sub></li> <li>• NOAEC</li> </ul>	18,340 mg/l 0.44 ml glycol/l 0.12 ml glycol/l 0.07 ml glycol/l 13,020 mg/l	
	Chronic toxicity test:		[57]
	<ul style="list-style-type: none"> <li>• NOEC</li> <li>✓ Mortality</li> <li>✓ Reprod. or growth</li> <li>• IC<sub>25</sub></li> </ul>	29,000 13,020 mg/l 13,470 mg/l	
<i>Pimephales promelas</i>	Acute toxicity tests:		[57]
	<ul style="list-style-type: none"> <li>• 48-hr LC<sub>50</sub></li> <li>• 96-hr LC<sub>50</sub></li> <li>• 7-day LC<sub>50</sub></li> <li>• NOAEC</li> </ul>	>62,000 mg/l 0.07 ml glycol/l 55,770 mg/l 0.03 ml glycol/l 0.03 ml glycol/l 52,930 mg/l	[64] [57] [64] [57]
	Chronic toxicity test:		[60]
	<ul style="list-style-type: none"> <li>• NOEC :</li> <li>✓ Mortality</li> <li>✓ Reprod. or growth</li> <li>• IC<sub>25</sub></li> </ul>	<11,530 <11,530 mg/l 6,940 mg/l	
<i>Daphnia magna</i>	Acute toxicity tests:		[64]
	<ul style="list-style-type: none"> <li>• 48-hr LC<sub>50</sub></li> <li>• 96-hr LC<sub>50</sub></li> </ul>	0.24 ml glycol/l 0.05 ml glycol/l	
<i>Daphnia pulex</i>	Acute toxicity tests:		[64]
	<ul style="list-style-type: none"> <li>• 48-hr LC<sub>50</sub></li> <li>• 96-hr LC<sub>50</sub></li> </ul>	0.27 ml glycol/l 0.06 ml glycol/l	
<i>Daphnia magna</i>	1,3-Propylene glycol		
	Acute toxicity tests		[58]
	<ul style="list-style-type: none"> <li>• 24-hr neonate, 24-hr LC<sub>50</sub> (s, n)</li> <li>• &lt;24-hr neonate, 48-hr LC<sub>50</sub> (s, n)</li> </ul>	8,285 mg/l 7,417 mg/l	
<i>Daphnia magna</i>	1,2-Propylene glycol		
	Acute toxicity tests:	>10,000 mg/l	[58]
	<ul style="list-style-type: none"> <li>• &lt;24-hr neonate, 48-hr LC<sub>50</sub> (s, n)</li> <li>• &lt;24-hr neonate, 24-hr LC<sub>50</sub> (s, n)</li> </ul>	>10,000 mg/l	
<i>Pimephales promelas</i>	Diethylene glycol		
	Acute toxicity tests:		[58]
	Juvenile (35-day); 96-hr LC <sub>50</sub> (s, n)	75,200 mg/l	
<i>Daphnia magna</i>	Acute toxicity tests:		[58]
	<ul style="list-style-type: none"> <li>• &lt;24-hr neonate, 24-hr LC<sub>50</sub> (s, n)</li> </ul>	>10,000 mg/l	

**TABLE 5.** Literature data on acute and chronic toxicity information on aircraft de-icing and anti-icing fluids as pure compounds (*Continued*)

Organism	Test type/ Parameter	Literature review of mean values of toxicity indices	Reference
Potassium acetate de-icing fluid <sup>f,g</sup>			
<i>Vibrio fischeri</i>	Acute toxicity test • Microtox (EC <sub>50</sub> )	6,560 mg/l	[63]
<i>Pimephales promelas</i>	Acute toxicity test • LC <sub>50</sub>	298 mg/l	[67]
	Chronic toxicity test • IC <sub>25</sub>	54.5 mg/l	
<i>Ceriodaphnia dubia</i>	Acute toxicity test • LC <sub>50</sub>	421 mg/l	[67]
	Chronic toxicity test • IC <sub>25</sub>	54.5 mg/l	
<i>Selenastrum capricornutum<sup>h</sup></i>	Chronic toxicity test • IC <sub>25</sub>	19.9 mg/l	[67]
Sodium formate de-icing fluids			
<i>Vibrio fischeri</i>	Acute toxicity test • Microtox (EC <sub>50</sub> )	1,780 mg/l	[67]
<i>Pimephales promelas</i>	Acute toxicity test • LC <sub>50</sub>	4,130 mg/l	[67]
	Chronic toxicity test • IC <sub>25</sub>	1,200 mg/l	
<i>Ceriodaphnia dubia</i>	Acute toxicity test • LC <sub>50</sub>	1,860 mg/l	[67]
	Chronic toxicity test • IC <sub>25</sub>	584 mg/l	
<i>Pseudokirchneriella subcapitata</i>	Chronic toxicity test • IC <sub>25</sub>	1,670 mg/l	[67]

<sup>a</sup>Toxicity test results for four different species to ethylene glycol-based type I and type IV aircraft de-/anti-icing fluids. <sup>b</sup>Ethylene glycol content in type I fluid is 92%. <sup>c</sup>Ethylene glycol content in type IV fluid is 64%. <sup>d</sup>The test organisms were of different ages (fry, juvenile, subadult). <sup>e</sup>s = static; n = nominal concentrations; m = measured concentrations. <sup>f</sup>Multiply by 0.66 for toxicity endpoint expressed as potassium concentration. <sup>g</sup>Multiply by 3.32 for toxicity endpoint expressed as original K-Ac de-icing product. <sup>h</sup>Previously name as *Pseudokirchneriella subcapitata*.

the LC<sub>50</sub> parameter is also used: this defines the lethal concentration of a substance in water, soil, or air, causing the death of 50% of individuals in a population. But the determination of the acute toxicity level is preceded by a step in which the effects a substance has on an organism are assessed—this then enables further toxicity tests to be specified more precisely.<sup>29</sup>

The dose-biological response relationship can also be used to predict the level of risk—a dose or exposure time for which the probability of toxic effects manifesting themselves is suitably low. Threshold (boundary/limiting) concentrations or doses are expressed using the following parameters: No Observed Effect Level/Concentration (NOEL or NOEC); Lowest Observed Effect Level/Concentration (LOEL or LOEC), NOAEL or NOAEC (No Observed

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Adverse Effect Level/Concentration), Lowest Observed Adverse Effect Level (LOAEL).<sup>29,50</sup>

## 6.2 Methods of Testing Toxicity

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The aquatic toxicity tests described here are the most commonly used ones; they are highly standardized.

### 6.2.1 ACUTE FRESHWATER TOXICITY TESTS

The following tests measure the lethality of water samples to freshwater organisms, indicating the toxicity of water samples. The *Daphnia pulex* or *Daphnia magna* (Daphnid) acute toxicity test evaluates the acute toxicity of a sample to a water flea belonging to the genus *Daphnia*. The test uses a static or static-renewal design and lasts 24, 48, or 96 hr. Observations at 24, 48, and 96 hr or 7 days permit the calculation of 24-, 48-, 96-hr and 7-day LC values.<sup>51,64</sup>

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The procedure using the fathead minnow (*Pimephales promelas*) and the water flea (*Ceriodaphnia dubia*) is also used to measure acute toxicity.<sup>50,52</sup>

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### 6.2.2 CHRONIC FRESHWATER TOXICITY TESTS

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These tests measure both lethal and sublethal effects over the life cycle or partial life cycle of freshwater organisms, providing information useful in assessing the potential long-term effects of contamination.<sup>57</sup>

The *Ceriodaphnia dubia* survival and reproduction test estimates the chronic toxicity of a sample to *C. dubia*, a water flea. The test uses the static-renewal design and lasts for seven days, monitoring both the survival of test organisms and the number of offspring they produce.<sup>57</sup>

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The fathead minnow (*Pimephales promelas*) larval survival and growth test uses the static-renewal design and lasts for seven days, tracking the survival of test organisms and their increase in weight.<sup>57</sup>

The fathead minnow (*Pimephales promelas*) embryolarval survival and teratogenicity test assesses the chronic toxicity of a sample to minnows, beginning as embryos and extending to the larval stage. The test uses the static-renewal design and lasts for seven days, noting both the survival of the fish and the induction of terata.<sup>57</sup>

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The algal (*Selenastrum capricornutum*) growth test identifies the bio-stimulatory and chronic toxic effects of a sample to a one-celled freshwater alga. The test uses the static design and lasts 96 hr, most commonly monitoring cell density (cells per ml). Alternative measures include biomass.<sup>57</sup>

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### 6.2.3 MICROBIAL TOXICITY TESTS

Microbial toxicity tests assess toxic effects on the microbial community and can serve as cost-effective and rapid screening indicators.<sup>57</sup>

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The Microtox test measures the toxicity of water samples or elutriates to *Photobacterium phosphoreum*, a species of bioluminescent marine bacteria.<sup>57</sup> Two features of *Vibrio fischeri* bacteria—rapid division and the triggering of luciferase (combined with cell respiration)—make these organisms ideal for measuring chronic toxicity.<sup>56</sup> Some contaminants inhibit the bacteria's metabolism, decreasing the intensity of light emitted; others stimulate the bacteria and cause an increase in luminescence.<sup>57</sup> Samples containing one or more toxic chemicals retard the reproductive cycle, growth, respiratory activity, and triggers luciferase, which can be detected at low levels by comparing the level of light in the control cuvette with that in the sample cuvettes after 22 hr of incubation.<sup>60</sup>

### 6.3 Literature Information on the Toxicity of Runoff Water Samples From Airports

The airport runoff waters may contain chemicals such as ethylene glycol, propylene glycol, 1,3-propylene glycol, 1,2-propylene glycol, diethylene glycol, potassium acetate, and sodium formate. These compounds are characterized with high toxicity (Table 5). For example, propylene glycol concentration levels in the runoff samples are very high and may reach even 160,000 mg/l with flow rate intensity 12.6 l/s (Bradley International Airport, Windsor Locks, CT, USA).<sup>16</sup> Similarly high concentration levels were determined in case of ethylene glycol and diethylene glycol, and reach levels exceeding 20,000 mg/l (Dallas/Fort Worth International Airport, Dallas, TX, USA; Kansas City International Airport, Kansas City, MO, USA).<sup>16</sup>

Toxicity of particular chemicals being present in the runoff water samples constitutes total toxicity of such kind of environmental samples (Table 6). Sometimes, the toxicity of runoff water samples may be very high and may reach LC50 values of 85%. If the impact of de-icing chemicals toxicity effects on aquatic organisms is to be concerned, several toxicity tests should be applied to assure compliance. The test species and conditions are specified in EPA guidelines for acute and chronic whole effluent toxicity tests. Freshwater species mentioned in most requirements described in legal system of United States include the fathead minnow (*Pimephales promelas*) and the daphnid *Ceridaphnia dubia* as presented in Table 5.<sup>17,54</sup>

## 7. MANAGEMENT OF AIRPORT INDUSTRIAL WASTE

Because of the possible adverse environmental effects of airport runoff and pending regulation, various technologies and management approaches are being investigated or implemented.<sup>25</sup> Many large airports, and airports that face severe weather conditions frequently through the winter months, are

**TABLE 6.** Toxicity studies conducted airport water

Toxicity test/organism	Sampling location	Values measured	Sampling time	Reference
48-hr LC <sub>50</sub> /Fathead minnow	Baltimore-Washington International Airport	LC <sub>50</sub> : NT <sup>a</sup> -70.0%	October 1990	[58]
Daphnid		LC <sub>50</sub> : NT-85.3%		
Fathead minnow and Daphnid	commuter terminal	NT		
Microtox	Muddy Bridge Branch (receives runoff directly from Baltimore—Washington International Airport)	<ul style="list-style-type: none"> <li>• EC-50 (5 min) = NR<sup>b</sup> [ml glycol/l]</li> <li>• EC-50 (15 min) = NR [ml glycol/l]</li> </ul>	March 1991	[64]
Microtox	A major North American international airport	<ul style="list-style-type: none"> <li>• 5 min IC<sub>50</sub><sup>c</sup> = 21–25%</li> <li>• 15 min IC<sub>50</sub> = 23–27%</li> </ul>	—	[64]

<sup>a</sup>NT = not toxic (no mortality in 100% storm water treatment). <sup>b</sup>NR = no response. <sup>c</sup>IC<sub>50</sub> = the inhibition concentration causing a 50% decrease in light production at 5 and 15 min.

**TABLE 7.** Airport runoff management techniques

Research site	Time of experiment	Technique	Plants/organisms/materials used in remediation techniques	Compounds, total parameters	Removal efficiencies (%; unless otherwise noted)	Reference(s)
Heathrow Airport, England	September 1994-September 1995	Reedbed systems: • Surface flow system (SF) • Sub-surface flow system (SSF)	Remediation techniques: Macrophytes: • Reedmace ( <i>Typha</i> species) • Common reed ( <i>Phragmites</i> species)	SSF system	SF system	[61]
			BOD, COD BOD Orthophosphate Nitrate Ammonia Metals Zinc Lead	31.0 <sup>a</sup> 23.0 63.5 52.6 46.5 — 43.8 12.4	18.0 40.4 37.4 27.5 — 46.9 —	
	• August-September 1995 • January, June and October 1996	Constructed wetland (soil surface flow, grab subsurface flow)	<i>Typha latifolia</i> <i>Typha angustifolia</i> <i>Phragmites australis</i> <i>Schoenoplectus lacustris</i> <i>Iris pseudacorus</i>	SSF system	SF system	[66]
			EG	99.4 <sup>b</sup> 71.6 <sup>c</sup> 76.6 <sup>d</sup> 1,2-Propylene glycol 99.6 <sup>b</sup> 59.2 <sup>c</sup> 95.3 <sup>d</sup> Diethylene glycol (DEG) 90.0 <sup>b</sup> 69.1 <sup>c</sup> 44.5 <sup>d</sup>	60.2 <sup>c</sup> — — 48.6 <sup>c</sup> 59.7 <sup>d</sup> — 61.3 <sup>c</sup> 44.5 —	

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**TABLE 7.** Airport runoff management techniques (*Continued*)

Research site	Time of experiment	Technique	Plants/organisms/materials used in remediation techniques	Compounds, total parameters	Removal efficiencies (%; unless otherwise noted)	Reference(s)
Moreppen (neighborhood of Oslo International Airport, Gardermoen), Norway	—	Microbial degradation in the vadose zone	• Microbial population	• Sodium formate • PG • Potassium acetate • Toluene	NA <sup>e</sup>	[67]
Gardermoen Airport, Norway	Snowmelt 1994/1995	Degradation in the unsaturated zone	Extracting soil water from 3 (PG or 40 suction cups placed at Potassium acetate depths between 0.4 and 2.4 m.  Degradation was analyzed by spatial moment calculations.	• Benzotriazole • Benzotriazole	0,015 day <sup>-1,f</sup> (1994) 0,047 day <sup>-1,f</sup> (1995) NA	[68]
Dane County Regional Airport, Wisconsin, USA	April-July	Aerobic bioremediation	• Bacteria ( <i>Pasteurella multocida</i> , <i>Acinetobacter</i> • COD <i>anitratus</i> , <i>Pseudomonas stutzeri</i> )  • Nutrient mixes • Enzyme mixes	• BOD • COD	The amount of sludge generated from this technique ranges 5% to 10% beginning volume.	[69]
Airport, Ohio, USA	—	Aerobic fixed-film biodegradation	• Ultrasonic stimulation Macronutrients, passively	EG	over 90%	[70]
Several north-eastern airports, USA	1966-1967	Anaerobic degradation	Anaerobic treatment	PG EG	3,5 day <sup>-1,f</sup> 5,2 day <sup>-1,f</sup>	[71]

Miami International Airport, USA	—	Thermal Treatment	Dewatering procedures	<ul style="list-style-type: none"> <li>• TSS</li> <li>• TOC</li> <li>• TDS (total dissolved solids)</li> <li>• Heavy metals</li> </ul>	NA	
General Mitchell International Airport, Wisconsin, USA	—	<ul style="list-style-type: none"> <li>• Recycling</li> <li>• Anaerobic biotreatment</li> <li>• Aerobic biotreatment</li> </ul>	<ul style="list-style-type: none"> <li>• De-icers</li> <li>• BOD</li> </ul>	Anaerobic biotreatment: degrade the waste sludge solids to produce biogas containing approximately 65% CH <sub>4</sub> and 35% CO <sub>2</sub>	[25]	
Airport, Texas, USA	—	Mineralization in root zone soil	<ul style="list-style-type: none"> <li>• Legumes: <i>Medicago</i> <i>sativa</i>, <i>Lotus</i> <i>corniculatus</i></li> <li>• Grasses: <i>Panicum</i> <i>virgatum</i>, <i>Poa pratensis</i> <i>L.</i>, <i>Lolium perenne L.</i>, <i>Sporobolus crypsandrurus</i>, <i>Festuca arundinacea</i></li> <li>• Mixtures: grass mix, grass mix + legumes</li> </ul>	NA	[69]	
Airport, USA	—	Waste de-icing fluid minimization	<ul style="list-style-type: none"> <li>• Microbial degradation</li> <li>• A microbial consortium (EG-C) especially: Gram-negative rod (EG-y)</li> </ul>	<ul style="list-style-type: none"> <li>• EG</li> <li>• PG</li> </ul>	NA	
General Mitchell International Airport, USA	—	Forced hot air de-icing and reduced fluid usage	Mixtures (more water and less glycol)	<ul style="list-style-type: none"> <li>• ADAF</li> </ul>	NA	[25]

(Continued on next page)

**TABLE 7.** Airport runoff management techniques (*Continued*)

Research site	Time of experiment	Technique	Plants/organisms/materials used in remediation techniques	Compounds, total parameters	Removal efficiencies (%; unless otherwise noted)	Reference(s)
Milwaukee Airport, Wisconsin, USA	—	Recycling	The runoff is distilled to • Glycol remove water and concentrate glycol.	• ADAF	NA	[25]
Buffalo International Airport, New York, USA	1997 and 1998	Infrared heating	Heating units	• ADAF	NA	[25]
Rhineland/Oneida County Airport, Wisconsin, USA	—	Three-stage steps	Filtration	Suspended solids (greater than 10 microns) chlorides, sulfates	NA	[74]
			Ion exchange or nanofiltration Distillation	PG EG	reduced from 10% to < 6% (within 7 days) Antifreeze agents (ethylene glycol-based)	

<sup>a</sup>For both raft systems. <sup>b</sup>Time of measurement: winter. <sup>c</sup>Time of measurement: summer. <sup>d</sup>Time of measurement: autumn. <sup>e</sup>Not available. <sup>f</sup>Degradation rate constant for PG and EG; percentage of data of removal efficient data was not available.

405 striving to reduce ADAF-laden wastewater runoff. Better weather forecasting and ice sensing instrumentation can reduce the amount of de-icing fluid needed to keep aircraft safe.<sup>65</sup> Technologies being developed include technologies remediation, forced hot air de-icing, appropriate use of less glycol, recycling and infrared de-icing (Table 7).<sup>25</sup>

## 8. SUMMARY

410 The need to describe more accurately the state of the environment and the processes taking place in it is a great challenge to analytical chemists where determining the concentrations of a wide range of analytes in samples with a complex matrix composition is concerned. Runoff waters from various surfaces are among the most interesting of such environmental samples.

415 A particular example of samples with a very complex composition are the waters due to rainfall and/or snowmelt running off the various surfaces to be found at airports. The greatest danger is caused by runoff from such surfaces as runways,<sup>16,23</sup> taxiways,<sup>16</sup> aircraft de-/anti-icing pads,<sup>14</sup> aprons,<sup>5,15</sup> transshipment points,<sup>5</sup> fuel stores and distribution points,<sup>5,16</sup> and repair shops and garages.<sup>5,16</sup>

420 Runoff waters generated by airport operations pose a serious hazard to the environment because they contribute to the contamination of the air, water, and soil. Airport operations also exert an adverse effect on the climate, plant kingdom, and above all, on human beings and animals. The principal pollutants discharged during airport operations are fuel, oil, grease, detergents, combustion gases, de-/anti-icing chemical wastes, toxic metals, alkalis, acids, and organic solvents; all these compounds are very often highly toxic.<sup>3,16,54,60</sup>

425 Analysis of the contaminants present in samples of airport runoff waters provides a clearer picture and understanding of the xenobiotic cycle in the environment. The worldwide increase in air traffic (more takeoffs and landings, more aircraft) means an increase in the amounts of pollutants getting into the environment. The runoff waters generated by these activities thus require special management procedures (e.g., technologies remediation, forced hot air de-icing, appropriate use of less glycol, recycling, infrared de-icing).<sup>25</sup>

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