NR 2009;43(7)

The Nordic Expert Group for Criteria Documentation of Health Risks from Chemicals

# 140. Sulphuric, hydrochloric, nitric and phosphoric acids

Marianne van der Hagen Jill Järnberg

ARBETE OCH HÄLSA

ISBN 978-91-85971-14-5



UNIVERSITY OF GOTHENBURG vetenskaplig skriftserie ISSN 0346-7821



#### Arbete och Hälsa

Arbete och Hälsa (Work and Health) is a scientific report series published by Occupational and Enviromental Medicine at Sahlgrenska Academy, University of Gothenburg. The series publishes scientific original work, review articles, criteria documents and dissertations. All articles are peer-reviewed.

Arbete och Hälsa has a broad target group and welcomes articles in different areas.

Instructions and templates for manuscript editing are available at http://www.amm.se/aoh Summaries in Swedish and English as well as the complete original texts from 1997 are also available online.

#### Arbete och Hälsa

Editor-in-chief: Kjell Torén

Co-editors: Maria Albin, Ewa Wigaeus Tornqvist, Marianne Törner, Wijnand Eduard, Lotta Dellve och Roger Persson Managing editor: Cina Holmer

© University of Gothenburg & authors 2009

Arbete och Hälsa, University of Gothenburg SE 405 30 Gothenburg, Sweden

ISBN 978-91-85971-14-5 ISSN 0346–7821 http://www.amm.se/aoh Printed at Reproservice, Chalmers

#### Editorial Board:

Tor Aasen, Bergen Kristina Alexanderson, Stockholm Berit Bakke, Oslo Lars Barregård, Göteborg Jens Peter Bonde, Köpenhamn Jörgen Eklund, Linköping Mats Eklöf, Göteborg Mats Hagberg, Göteborg Kari Heldal, Oslo Kristina Jakobsson, Lund Malin Josephson, Uppsala Bengt Järvholm, Umeå Anette Kærgaard, Herning Ann Kryger, Köpenhamn Carola Lidén. Stockholm Svend Erik Mathiassen, Gävle Gunnar D. Nielsen, Köpenhamn Catarina Nordander, Lund Gunnar Ahlborg, Göteborg Torben Sigsgaard, Århus Staffan Skerfving, Lund Kristin Svendsen, Trondheim Gerd Sällsten, Göteborg Allan Toomingas, Stockholm Ewa Wikström, Göteborg Eva Vingård, Uppsala

# Preface

The main task of the Nordic Expert Group for Criteria Documentation of Health Risks from Chemicals (NEG) is to produce criteria documents to be used by the regulatory authorities as the scientific basis for setting occupational exposure limits for chemical substances.

For each document, NEG appoints one or several authors. An evaluation is made of all relevant published, peer-reviewed original literature found. The document aims at establishing dose-response/dose-effect relationships and defining a critical effect. No numerical values for occupational exposure limits are proposed.

Whereas NEG adopts the documents by consensus procedures, thereby granting the quality and conclusions, the authors are responsible for the factual content of the document.

The evaluation of the literature and the drafting of this document were made by Cand. Scient Marianne van der Hagen at the Norwegian Pollution Control Authority, and Dr. Jill Järnberg at the Swedish Work Environment Authority. The draft document was discussed within the group and the final version was accepted by NEG on April 1, 2008 as its document. The following individuals participated in the elaboration of the document:

Gunnar Johanson	Institute of Environmental Medicine, Karolinska Institutet, Sweden (chairman)
Maria Albin	Department of Occupational and Environmental Medicine, University Hospital, Lund, Sweden (former NEG expert)
Karin Sørig Hougaard	National Research Centre for the Working Environment, Denmark (former NEG expert)
Kristina Kjærheim	Cancer Registry of Norway (NEG expert)
Tiina Santonen	Finnish Institute of Occupational Health, Finland (NEG expert)
Vidar Skaug	National Institute of Occupational Health, Norway (NEG expert)
Jill Järnberg and Anna-Karin Alexandrie	Swedish Work Environment Authority, Sweden (NEG secretariat)

Editorial work and technical editing were performed by the NEG secretariat. This work was financially supported by the former Swedish National Institute for Working Life, The Swedish Work Environment Authority, the Norwegian Ministry of Labour and Social Inclusion and the Nordic Council of Ministers.

All criteria document produced by NEG may be downloaded from *www.nordicexpertgroup.org*.

Gunnar Johanson, Chairman of NEG

# Contents

Preface	
Abbreviations and acronyms	
1. Introduction	1
2. Substance identification	1
3. Physical and chemical properties	2
4. Occurrence, production and use	4
5. Measurements and analysis of workplace exposure	6
6. Occupational exposure data	8
<ul> <li>7. Toxicokinetics</li> <li>7.1 Deposition</li> <li>7.2 Uptake</li> <li>7.3 Distribution</li> <li>7.4 Biotransformation and excretion</li> </ul>	14 14 15 15 15
8. Biological monitoring	15
9. Mechanisms of toxicity	16
<ul> <li>10. Effects in animals and <i>in vitro</i> studies</li> <li>10.1 Irritation, corrosion and sensitisation</li> <li>10.2 Effects of single exposure</li> <li>10.3 Effects of short-term exposure (up to 90 days)</li> <li>10.4 Mutagenicity and genotoxicity</li> <li>10.5 Effects of long-term exposure and carcinogenicity</li> <li>10.6 Reproductive and developmental effects</li> </ul>	17 17 20 32 39 39 47
11. Observations in man	47
<ul> <li>11.1 Irritation, corrosion and sensitisation</li> <li>11.2 Case reports</li> <li>11.3 Effects of single and short-term exposure</li> <li>11.4 Effects of long-term exposure</li> <li>11.5 Genotoxic effects</li> <li>11.6 Carcinogenic effects</li> <li>11.7 Reproductive and developmental effects</li> </ul>	47 48 50 63 66 70 72
12. Dose-effect and dose-response relationships	81
<ul> <li>12.1 Sulphuric acid</li> <li>12.1.1 Single/short-term exposure</li> <li>12.1.2 Long-term exposure</li> <li>12.2 Hydrochloric acid</li> <li>12.2.1 Single/short-term exposure</li> <li>12.2.2 Long-term exposure</li> </ul>	81 81 83 89 89 90
12.3 Nitric acid	91 91
12.3.1 Single/short-term exposure 12.3.2 Long-term exposure	91 93

12.4 Phosphoric acid	93
13. Previous evaluations by national and international bodies	93
13.1 Sulphuric acid	93
13.2 Hydrochloric acid	95
13.3 Nitric acid	95
13.4 Phosphoric acid	96
14. Evaluation of human health risks	96
14.1 Assessment of health risks	96
14.2 Groups at extra risk	100
14.3 Scientific basis for an occupational exposure limit	100
15. Research needs	102
16. Summary	103
17. Summary in Norwegian	104
18. References	105
19. Data bases used in the literature search	118
Appendix 1. Occupational exposure limits	119
Appendix 2. Previous NEG criteria documents	120

# Abbreviations and acronyms

ACGIH ARDS	American Conference of Governmental Industrial Hygienists
ARDS	acute (or adult) respiratory distress syndrome
CI	Agency for Toxic Substances and Disease Registry confidence interval
DECOS	
DECOS	Dutch Expert Committee on Occupational Safety
	Deutsche Forschungsgemeinschaft (German Research Foundation)
FEF <sub>x</sub>	forced expiratory flow at x % of FVC
$FEV_1$	forced expiratory volume in one second
FVC	forced vital capacity
IARC	International Agency for Research on Cancer
IOM	Institute of Occupational Medicine, Edinburgh, United Kingdom
IPCS	International Programme on Chemical Safety
$LC_{50}$	lethal concentration for 50 % of the exposed animals at single exposure
LOAEL	lowest observed adverse effect level
MD	median diameter
MMAD	mass median aerodynamic diameter
MMD	mass median diameter
NEG	Nordic Expert Group for Criteria Documentation of Health Risks from
	Chemicals
NIOSH	National Institute for Occupational Safety and Health (United States)
NOAEL	no observed adverse effect level
OR	odds ratio
$PM_x$	particulate matter with aerodynamic diameter up to x µm
RADS	reactive airways dysfunction syndrome
RD <sub>50</sub>	air concentration associated with a 50 % decrease in the respiratory rate
	of animals
RR	relative risk (risk ratio)
SCOEL	Scientific Committee on Occupational Exposure Limits (European
	Union)
SG <sub>aw</sub>	specific airway conductance
SIR	standardised incidence ratio
SMR	standard mortality ratio
STEL	short-term exposure limit
TNFα	tumour necrosis factor alpha
TWA	time-weighted average
VMD	volume median diameter
WHO	World Health Organization
,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	, one nouter organization

# 1. Introduction

Inorganic acids are of prime importance in the chemical and metal industries. They are used as raw materials in the manufacture of a wide range of chemicals, as well as in refining, electrolysis and extraction in chemical processes. Inorganic acids are widely used in the pickling processes of electroplating, in vehicle production plants and in steel producing plants.

The present document concerns the effects of four inorganic acids: sulphuric acid ( $H_2SO_4$ ), hydrochloric acid (HCl), nitric acid (HNO<sub>3</sub>) and phosphoric acid ( $H_3PO_4$ ). It does not consider sulphur dioxide, nitrous gases, or phosphorous compounds such as phosphorous pentoxide and red phosphorus, which may be converted to phosphoric acid.

A previous NEG document published in 1993 described the effects of aerosols of these four acids (142). The biological effects of  $H_2SO_4$  are relatively well investigated, whereas the documentation concerning the effects of the other three acids, discussed in this document, is limited.

Acid concentrations are given in  $mg/m^3$ . Data originally reported in units of ppm have been converted to units of  $mg/m^3$  by using the conversion factors given in Table 2. Occasionally, the originally reported ppm values are also given.

# 2. Substance identification

Substance identification data for the inorganic acids dealt with in this document are given in Table 1.

Although anhydrous *sulphuric* and *nitric acids* can be produced, the chemical names usually refer to aqueous solutions of the compounds.

*Hydrochloric acid* is the aqueous solution of hydrogen chloride gas. In this document, the chemical formula HCl may denote hydrochloric acid as well as hydrogen chloride gas.

Common name	CAS No.	Synonyms	Molecular formula	Molecular weight
Sulphuric acid	7664-93-9	Battery acid, dipping acid, electrolyte acid, fertiliser acid, hydrogen sulphate, matting acid, Nordhausen acid, oil of vitriol, spirit of sulphur, sulfuric acid	$H_2SO_4$	98.08
Hydrochloric acid	7647-01-0	Hydrogen chloride, chlorohydric acid, hydrochloride, muriatic acid, spirits of salt	HCl	36.46
Nitric acid	7697-37-2	Aqua fortis, azotic acid, hydrogen nitrate	HNO <sub>3</sub>	63.02
Phosphoric acid	7664-38-2	Orthophosphoric acid	$H_3PO_4$	98.00

Table 1. Substance identification data of the inorganic acids (142).

*Phosphoric acid,* when pure, is a solid at room temperature and normal pressure. However, an aqueous solution of  $85 \% H_3PO_4$  is a viscous liquid. In this document,  $H_3PO_4$  refers to aqueous solutions of the solid compound.

# 3. Physical and chemical properties

Data on physical and chemical properties are presented in Table 2.  $H_2SO_4$ , HCl and HNO<sub>3</sub> are all strong acids, and dissociate completely in water at moderate concentrations. HCl is the strongest acid (indicated by the lowest pK<sub>a</sub>) followed by  $H_2SO_4$  and HNO<sub>3</sub>.  $H_3PO_4$  is the weakest of the acids. All four acids are hygroscopic (attract water molecules from the surrounding environment) and corrosive to (destroy or irreversibly damage) living tissue. HNO<sub>3</sub> and concentrated  $H_2SO$  are strong oxidising agents.

Solutions of the acids are not flammable in themselves. However, contact with metals may release flammable and explosive hydrogen gas  $(H_2)$ .

The physical state of the acids in workplace air ranges from primarily liquid aerosols for the non-volatile  $H_2SO_4$  and  $H_3PO_4$  to liquid aerosols and vapours for the more volatile HCl and HNO<sub>3</sub> (37). There are a number of definitions of aerosols that are used in a wide variety of contexts. IUPAC defines an *aerosol* as a mixture of small (diameter 0.01-100 µm) solid or liquid particles and a carrier gas (usually air). A *droplet* is defined as a small liquid particle and a *mist* as a suspension of droplets in a gas. Also *fog* is used as a general term applied to a suspension of droplets in a gas (119). Other kinds of aerosols are *dust, fume, smoke, haze,* and *smog* (190).

In the present document the original wording regarding aerosols in the cited papers is retained.

#### Sulphuric acid

 $H_2SO_4$  is a colourless (when pure) to dark brown, oily liquid. It is odourless unless heated (then pungent). Cold  $H_2SO_4$  reacts with most metals and the reactivity increases upon heating. The acid can release flammable hydrogen gas when in contact with metals.  $H_2SO_4$  is highly corrosive, a property accentuated by its highly exothermic (generates heat) reaction with water.  $H_2SO_4$  is an excellent dehydrating agent. Its affinity for water is sufficiently strong to remove hydrogen and oxygen atoms from other compounds. Fuming  $H_2SO_4$  (oleum) is a solution of the anhydride sulphur trioxide (SO<sub>3</sub>) in anhydrous  $H_2SO_4$ . In air, sulphur trioxide vapour is rapidly converted to a stable mist of droplets of the acid by reacting with atmospheric moisture (6, 96, 110, 180).

#### Hydrochloric acid

HCl is a colourless fuming liquid with a pungent irritating odour. The mixture of HCl gas and water has a constant-boiling azeotrope at 20 % HCl. It is a very stable compound but decomposes at high temperatures into hydrogen and chlorine. HCl reacts violently with bases and is corrosive. It also reacts violently with oxidants

Property	Sulphuric acid (H <sub>2</sub> SO <sub>4</sub> )	Hydrochloric acid (HCl)	Nitric acid (HNO <sub>3</sub> )	Phosphoric acid (H <sub>3</sub> PO <sub>4</sub> )
Description	Colourless (pure) to dark brown, oily liquid; odourless unless heated, then pungent	Colourless to slightly yellow, fuming liquid; pungent irritating odour	Clear, colourless liquid; suffocating odour	Clear, colourless, syrupy liquid (<85 %); odourless
Boiling point (°C)	338 (98 %, aqueous solution)	110 (20 %) -85 (gas)	121 (70 %) 86 (100 %, decomposes)	158 (85 %) 261 (solid)
Melting point (°C)	10 (100 %)	- 85 (25 %) -114 (gas)	- 42 (70 %)	21 (85 %) 42 (solid)
Vapour pressure (kPa) at 20 °C	< 0.04	1.5	1.2	0.3
Vapour density (air=1)	3.4	1.3	1.4	3.4
Density (g/ml) at 25 °C	1.84 (100 %)	1.19 (38 %)	1.41 (70 %)	1.71 (85 % ) 1.86 (solid)
Solubility in water	Complete	Complete	Complete	Complete
Solubility in organic solvents	Soluble in ethanol	Very soluble in alcohols, soluble in ether and benzene, insoluble in hydrocarbons	No data on solubility in ethanol, soluble in ether	Soluble in ethanol
Octanol/water partition coefficient (log $P_{ow}$ ) <sup>a</sup>	-2.20 (estimated)	0.54 (estimated)	0.21 (estimated)	-0.77 (estimated)
pK <sub>a</sub> -value <sup>b, c</sup>	-3.0, 1.99	-8.0	-1.3	2.12, 7.21, 12.32
Odour threshold (mg/m <sup>3</sup> )	>1	Range 1-50	0.75-2.50	No odour
Conversion factors in air (25 °C, 101.3 kPa)	$1 \text{ mg/m}^3 = 0.25 \text{ ppm}$ 1 ppm = 4.0 mg/m <sup>3</sup>	$1 \text{ mg/m}^3 = 0.7 \text{ ppm}$ $1 \text{ ppm} = 1.4 \text{ mg/m}^3$	$1 \text{ mg/m}^3 = 0.4 \text{ ppm}$ $1 \text{ ppm} = 2.5 \text{ mg/m}^3$	$1 \text{ mg/m}^3 = 0.25 \text{ ppm}$ $1 \text{ ppm} = 4.0 \text{ mg/m}^3$

**Table 2.** Physical and chemical properties of the inorganic acids (3-6, 19, 74, 75, 129, 142, 143, 188).

5

<sup>a</sup> Data from Syracuse Research Corporation's LogKow (KowWin) Program. <sup>b</sup> Data from Harvard University, Evans Group, substrate H<sub>2</sub>O. Values < 0 were extrapolated. <sup>c</sup> H<sub>2</sub>SO<sub>4</sub>: Two pKa-values as it releases two hydrogen ions. H<sub>3</sub>PO<sub>4</sub>: Three pKa-values as it releases three hydrogen ions.

forming toxic chlorine. On contact with air it emits corrosive fumes. HCl attacks nearly all metals under release of hydrogen gas. A mixture of concentrated HCl and a strong oxidising agent dissolves gold, e.g. *aqua regia*, which is a 3:1 (v/v) mixture of concentrated HCl and HNO<sub>3</sub> (96, 110).

#### Nitric acid

Pure anhydrous HNO<sub>3</sub> is a transparent, colourless liquid with a characteristic choking odour and in moist air a white fuming liquid. In the presence of light or by heating, HNO<sub>3</sub> readily decomposes. The vapours formed are a mixture of HNO<sub>3</sub> and decomposition products such as nitrogen oxides (NO<sub>x</sub>), oxygen and water. Nitrogen dioxide (NO<sub>2</sub>) accounts for the red-brownish (yellow at low concentrations) colour that develops in the acid on standing. HNO<sub>3</sub> is highly corrosive and attacks most substances and all metals except the noble ones and certain alloys. Reactions with metals may produce nitrous gases and ammonia. Hydrogen gas is rarely formed. HNO<sub>3</sub> forms a constant boiling mixture with water. Concentrated HNO<sub>3</sub> (~70 %) is a powerful oxidising agent and reacts with combustible, organic and readily oxidisable materials. The reactions are often highly exothermic and explosive. Aqueous solutions of > 45 % HNO<sub>3</sub> may spontaneously ignite organic materials such as wood and straw (3, 96, 110, 129, 209).

#### Phosphoric acid

Pure anhydrous  $H_3PO_4$  is an odourless white solid that melts at 42 °C to form a clear, colourless, viscous liquid. At room temperature, an 85 % aqueous solution of  $H_3PO_4$  is likewise a clear, colourless, viscous liquid. Besides its acidic behaviour,  $H_3PO_4$  is relatively unreactive at room temperature (74).

## 4. Occurrence, production and use

The four inorganic acids in this document are all important industrial chemicals used in a wide variety of industries/trades and products.

#### Sulphuric acid

 $H_2SO_4$  is made as a by-product of other operations or directly from elemental sulphur, spent (contaminated and diluted)  $H_2SO_4$ , and hydrogen sulphide. Elemental sulphur is by far the most widely used raw material. Concentrated  $H_2SO_4$  is ca 96-98 % (w/w) (18 M). Battery acid used in lead-acid batteries is 33.5 %, chamber or fertiliser acid 62-68 %. Fuming  $H_2SO_4$ , also called oleum, is a solution of sulphur trioxide in anhydrous  $H_2SO_4$ .  $H_2SO_4$  is used in batteries, pH-regulation agents, electroplating agents, process regulators, cleaning/washing agents, complexing agents, and in surface treatment and laboratory chemicals.  $H_2SO_4$  is also used in the manufacture of chemicals (e.g. titanium dioxide pigments), fertilisers, basic metals, pulp and paper, metal products, food products and beverages, electrical machinery and apparatuses, and in purification of water and treatment of sewage.  $H_2SO_4$  is the main air contaminant at anodising plants (6, 59, 129, 222).

#### Hydrochloric acid

HCl is produced in various processes, such as in the reaction between NaCl and  $H_2SO_4$ , or by direct synthesis from  $H_2$  and  $Cl_2$ . HCl is also a major by-product in chemical processes when organic compounds are synthesised. It is produced in solutions up to 38 % (w/w) (concentrated grade) equivalent to 12 M (pH -1.1). HCl is used in the manufacture of chemicals, pulp and paper (although substituted in most enterprises by now), metal products, food products and beverages and in the extraction of crude petroleum and natural gas, in purification of water and treatment of sewage, and as a general disinfectant (5, 129, 222, 242).

#### Nitric acid

HNO<sub>3</sub> is difficult to manufacture as a pure substance due to its tendency to decompose. Virtually all HNO<sub>3</sub> manufacture is by oxidation of ammonia. The concentrated acid is an aqueous solution containing 70 % (w/w) (16 M) HNO<sub>3</sub> (azeotropic mixture). "White" fuming HNO<sub>3</sub> is a highly concentrated acid, typically > 90 %, containing 0.1-0.4 % nitrogen dioxide. "Red" fuming HNO<sub>3</sub> contains 8-17 % dissolved nitrogen dioxide. In practice, HNO<sub>3</sub> is usually found in conjunction with NO<sub>x</sub>, and vapours of HNO<sub>3</sub> are always a mixture of acid, NO<sub>x</sub>, oxygen and water whose composition is determined by factors such as temperature and humidity. HNO<sub>3</sub> is used in the manufacture of chemicals such as fertilisers and explosives, food products, beverages, metal products, in surface treatment and pH-regulation agents. It is also a component in laboratory chemicals and in cleaning/washing agents (3, 12, 66, 129, 209, 222).

#### Phosphoric acid

 $H_3PO_4$  is produced commercially by either the wet process or the electric furnace process. In the wet process,  $H_3PO_4$  is produced by reacting  $H_2SO_4$  with naturally occurring phosphate rock in a reactor. Reagent grade  $H_3PO_4$  is usually 85 % (w/w).  $H_3PO_4$  is used in fertilisers, cleaning/washing agents, surface treatment, laboratory chemicals, colouring agents, pH-regulation agents, process regulators, food/feedstuff, flavourings and nutrients, non-agricultural pesticides and preservatives, and corrosion inhibitors.  $H_3PO_4$  is also used in the manufacture of beverages, chemicals, metal products, pulp and paper, rubber and plastics, textiles, purification of water and treatment of sewage, and in the extraction of crude petroleum and natural gas (4, 114, 129, 222).

Acid/	× *	,	Year		
Country	2002	2003	2004	2005	2006
Sulphuric acid					
Denmark <sup>a</sup>	2	4	16	18	10
Finland	1 474	1 182	1 567	928	998
Norway	207	220	238	248	72
Sweden	240	253	240	316	324
Hydrochloric ac	id				
Denmark <sup>a</sup>	4	9	33	40	28
Finland <sup>b</sup>	<<1	23	58	<1	<1
Norway	34	41	34	39	40
Sweden	84	75	54	47	55
Nitric acid					
Denmark <sup>a</sup>	3	<1	25	26	27
Finland	273	212	513	277	279
Norway	1 217	1 254	1 260	1 253	1 176
Sweden	10	10	16	27	39
Phosphoric acid	,				
Denmark <sup>a</sup>	3	3	5	5	4
Finland	219	204	798	307	325
Norway	4	4	5	6	5
Sweden	49	49	53	41	50

Table 3. Annual total use (kilotonnes) of inorganic acids in the Nordic countries (222).

<sup>a</sup> The large increase in volume in 2004 is due to new regulations for declaration.

<sup>b</sup> The amounts are for HCl, water free (index no. 017-002-00-2).

## Uses in the Nordic countries

Table 3 shows the annual use of the four inorganic acids in question in Denmark, Finland, Norway, and Sweden in the years 2002-2006 (222). The large variations of tonnage between the countries are explained by national differences in type of industries. For example, Norway is a large producer of fertilisers, hence the large tonnage for  $HNO_3$ .

# 5. Measurements and analysis of workplace exposure

Data on particle size distribution of acid mists are limited, and sampling methods have generally not differentiated between liquid and gaseous forms of acid (180). The droplet size of acid aerosols is reported differently by various authors. Usually it is described as mass median aerodynamic diameter<sup>1</sup> (MMAD), mass median diameter (MMD) or volume median diameter (VMD). MMAD entails not only size and shape but also density of droplets. MMD describes the aerodynamic behaviour of spherical and unit density particles, so for such droplets MMAD equals MMAD. When the aerosol density is approaching 1, VMD equals MMAD.

<sup>&</sup>lt;sup>1</sup> Aerodynamic diameter: The diameter of a unit-density sphere having the same terminal settling velocity as the particle in question. It is used to predict where in the respiratory tract such particles will deposit.

The occupational exposure limits for the four acids in the Nordic countries (Appendix 1) applies to the *inhalable* fraction (particles with aerodynamic diameter  $\leq 100 \ \mu\text{m}$ ) of the airborne particles according to European standard EN 481:1993 (77). Poorly volatile acids such as H<sub>2</sub>SO<sub>4</sub> and H<sub>3</sub>PO<sub>4</sub> are typically sampled as inhalable aerosols on filters, whilst volatile acids such as HCl and HNO<sub>3</sub> are collected on sorbent tubes or alkali-impregnated filters. Analysis is usually carried out by ion chromatography (37, 92).

When comparing published exposure data, one must consider the sampling technique used. The IOM sampler<sup>1</sup>, the Respicon impactor and the Millipore sampler measure different particle size fractions. The IOM sampler is assumed to come closer to the health-related inhalable fraction of aerosols, especially for aerosols with an aerodynamic diameter above 20  $\mu$ m (40, 155). It is assumed that the Millipore sampler underestimates the H<sub>2</sub>SO<sub>4</sub> concentration by a factor 1.5 compared to the IOM cassette. The detection limit for the Occupational Safety and Health Administration (OSHA) method ID-113 corresponds to 0.5  $\mu$ g sulphate on the filter (40, 182).

In the year 2000, the former National Institute for Working Life in Sweden recommended sampling of these four inorganic acids on silica tubes and analysis by ion chromatography, with reference to the United States National Institute for Occupational Safety and Health (NIOSH) method no. 7903 (153). The NIOSH method provides total acid mist concentration, but no size-related information (113).

The Scientific Committee on Occupational Exposure Limits (SCOEL) of the European Union acknowledged in 2007 that the reliable measurement of  $H_2SO_4$  concentrations at and around the recommended limit values (8-hour time-weighted average (TWA) 0.05 mg/m<sup>3</sup>, short-term exposure limit (STEL) 0.1 mg/m<sup>3</sup>) was challenging. In some circumstances, there might be interference from sulphate salts also present in the atmosphere. It was concluded unproblematic to measure 8-hour shift concentrations in workplaces with exclusively  $H_2SO_4$  mist, but impossible to monitor a 15-minute period with the necessary limit of detection. However, from the most recent evidence presented to SCOEL, it appears that there are, or soon will be, measurement techniques available that are compatible with the proposed limits (210). Interference may also be from oxidisable inorganic compounds such as sulphur dioxide and sulphites, and organic sulphuric compounds (144).

Interference from salts may be a challenge also for measurements and analysis of the other three acids (179).

<sup>&</sup>lt;sup>1</sup> IOM sampler = personal sampler developed by the Institute of Occupational Medicine (IOM) in Edinburgh, United Kingdom. The sampler was developed to fit the curve for inhalable dust according to the European Committee for Standardization (CEN) standard EN 481:1993.

# 6. Occupational exposure data

Contrary to the wide industrial use of the four acids, published contemporary exposure data are limited and are almost exclusively air measurements of  $H_2SO_4$ (Table 4). The Norwegian occupational exposure database EXPO contains data from all samples analysed at the National Institute of Occupational Health in Norway since 1984 (Table 5). Most of these samples have been collected as a result of requests from different enterprises to control their exposures and are likely to represent "worst case measurements" in many cases (189). From published data and data in EXPO it is difficult to describe the exposure levels in Nordic industry today.

#### Sulphuric acid

Occupational exposure data from various occupational settings are summarised in Table 4. Droplet size distributions are usually not given but industrial aerosols can have MMADs as large as 14  $\mu$ m (160). In a study of lead acid battery plants, MMADs were in the range 2.6-10  $\mu$ m (122). Most of the available exposure data comes from the pickling and plating industries (116). In a review, pre 1970s H<sub>2</sub>SO<sub>4</sub> mist potential exposures were judged: a) high for workers in H<sub>2</sub>SO<sub>4</sub> and isopropanol production, and in metal pickling (>1 mg/m<sup>3</sup>, 8-hour TWA); b) judged moderate for workers in soap and detergent, HNO<sub>3</sub> and ethanol production (0.1-1 mg/m<sup>3</sup>); and c) low for workers in copper and zinc refining and in phosphate fertiliser and lead battery production (< 0.1 mg/m<sup>3</sup>) (197). Since then, substitution and other measures have reduced the levels considerably.

In 1990-1993, approximately 700 000 workers including 4 000 in Denmark, 2 000 in Finland and 8 000 in Sweden were exposed to strong inorganic mists containing  $H_2SO_4$ , as registered in the CAREX database (125). The CAREX database contains estimates of the numbers of workers occupationally exposed to carcinogens in the 15 member states of the European Union at the time (exposure data from 1990-1993) and in four of the ten countries that joined the European Union in 2004 (exposure data from 1997).

In conference proceedings from the Australian Institute of Occupational Hygienists (AIOH) 1996, Foster *et al* reported exposure levels of  $H_2SO_4$  mist in several branches of industry. All measurements were expressed as total sulphate. In some cases this included sulphate present as metal salts. The highest exposures were recorded in the lead-acid battery manufacturing industry (0.03-1.5 mg/m<sup>3</sup>). Relatively high exposures were found in the electrolytic refining of metals (0.04-0.5 mg/m<sup>3</sup>) whereas acid mist exposures in the other industries were low, e.g. in electroplating workplaces. Due to enclosure of the processes, also exposures in fertiliser production, wool carbonising, soap and margarine manufacture, tanning, catalytic alkylation of hydrocarbons, and  $H_2SO_4$  manufacture were low (81).

In a recent investigation, the maximum  $H_2SO_4$  concentration in phosphate fertiliser manufacture facilities in Florida measured as  $PM_{10}$  (particulate matter with an aerodynamic diameter up to 10  $\mu$ m), including fine and coarse mode,

was  $0.185 \text{ mg/m}^3$  obtained at the H<sub>2</sub>SO<sub>4</sub> pump tank area. Geometric mean concentrations obtained at the H<sub>2</sub>SO<sub>4</sub> pump tank area for PM<sub>23</sub>, PM<sub>10</sub> and PM<sub>2.5</sub> H<sub>2</sub>SO<sub>4</sub> were 0.042, 0.038 and 0.022 mg/m<sup>3</sup>, respectively, measured with a cascade impactor. By using the NIOSH method 7903, measurements were 1.5-229 times higher than those obtained by the cascade impactor. One possible explanation provided by the authors is interaction of SO<sub>2</sub> in the NIOSH method (113, 114).

For comparison, air levels from environmental pollution by inorganic acids are generally lower than occupational exposures.  $H_2SO_4$  concentrations ranging from 0.02 to 0.03 mg/m<sup>3</sup>, with peaks up to 0.1 mg/m<sup>3</sup>, have been determined in outdoor air in Europe and the United States (67, 210). Elevated environmental  $H_2SO_4$  concentrations of 0.027 mg/m<sup>3</sup> as a 12-hour average and > 0.1 mg/m<sup>3</sup> as a one-hour peak have been measured in the eastern United States and Canada. These concentrations were measured as H<sup>+</sup>-concentrations and subsequently converted to  $H_2SO_4$  (221). In Norway, the environmental concentrations of SO<sub>2</sub> in the centre of Oslo were in the range 4-6 µg/m<sup>3</sup> (as yearly averages) during 1997-2004. In comparison, the 24-hour environmental standard for SO<sub>2</sub> is 0.125 mg/m<sup>3</sup> (1). The concentration of  $H_2SO_4$  is not regularly measured in this programme, but SO<sub>2</sub> can be converted to  $H_2SO_4$  in the troposphere.

#### Hydrochloric acid

According to exposure data, picklers in a Dutch hot dip galvanising plant worked 27 % of their time in HCl concentrations above 7 mg/m<sup>3</sup>. In earlier studies, the HCl levels were slightly lower than 7 mg/m<sup>3</sup> in German galvanising plants (191). In measurements performed during the 1950s to 1970s in Finland and the United States, air concentrations of HCl in pickling, cleaning, plating and electrochemical drilling ranged from 0.001 to 14.5 mg/m<sup>3</sup>. In a German study, levels were 26.5-33.5 mg/m<sup>3</sup> in pickling industries (165).

#### Nitric acid

Few measurements are available of  $HNO_3$  exposure. Individual air concentrations in acid treatment of metals (cleaning, etching, electrolytic refining, plating and anodising), mostly performed 1975-1983, were in the range 0.01-2.8 mg/m<sup>3</sup> (116).

#### Phosphoric acid

The few individual air measurements available on  $H_3PO_4$  exposure from pickling, acid cleaning, and aluminium finishing operations performed in the 1970s and 1980s were below 0.67 mg/m<sup>3</sup>. Air concentrations during phosphate fertiliser manufacture ranged from below 0.005 to 3.43 mg/m<sup>3</sup> (116).

Processing method/job	No. of samples	Sampling type	Sampling time	Exposure level, mg/m <sup>3</sup>	Sampling method	Reference
Zink production/two cell		Personal	Full-shift,	GM (range)	Millipore (37 mm)	(40)
houses (inspectors,	59		5.5-7.5 h	0.07 (0.01-0.48)		
strippers and cleaners)	70			0.04 (0.01-0.15)		
		Stationary	4.5-8 h	GM (GSD)		
	6			0.11 (1.54)		
	4			0.11 (1.71)		
	6			0.09 (1.46)		
	4			0.09 (1.51)		
13 workplaces:	94	Personal	Not given	GM (range)	Measured and reported as	(81)
Lead-acid battery plants			-	0.03-1.45	total sulphate	
Metal refining				0.04-0.47	-	
Fertiliser manufacture				0.09 (single sample)		
Electroplating				0.03 (two samples, same level)		
Wool carbonising				0.04-0.05		
3 anodising plants	"Several"	Personal	8 h	GM (range)	Millipore	(95)
01				0.40 (0.24-0.87)	I · · · ·	()
				0.04 (0.03-0.05)		
				0.01 (0.005-0.03)		
		<b>D</b> 1				(100)
3 titanium dioxide	Plant A: 95	Personal	Mostly full-	<0.05 as means in all 3 plants except	Millipore (37 mm)	(108)
manufacturing plants	Plant B: 100		shift	in the Moore filtration workshops		
	Plant C: 39			(A: 0.38, B: 0.84 and C: 0.09).		

**Table 4.** Reported occupational exposure levels of sulphuric acid.

Processing method/job	No. of samples	Sampling type	Sampling time	Exposure level, mg/m <sup>3</sup>	Sampling method	Reference
5 lead-acid battery plants	245	Personal	Full-shift, 6-7 h	AM (range) 0.18 (ND-1.7)	Millipore (37 mm)	(122)
Paper and paper board production and recycling	10 5 6 10	Not described	> 1 h	AM (range) 4.1 (0.3-11.5) (pulping, refining) 0.89 (0.22-1.6) (paperboard machine) 0.11 (0-0.27) (paper/paperboard machine) ND (repulping)	Not described	(138)
8 phosphate fertiliser manufacturing plants	71	Stationary	12 h	GM 0.042 (PM <sub>23</sub> ) 0.038 (PM <sub>10</sub> ) 0.022 (PM <sub>2.5</sub> ) 0.185 (maximum)	Cascade impactor (Mark III) Dichotomous (SA241 CUM)	(113, 114)

Table 4. Reported occupational exposure levels of sulphuric acid.

AM: arithmetic mean, GM: geometric mean, GSD: geometric standard deviation, ND: not detectable, PM<sub>x</sub>: particulate matter with aerodynamic diameter up to x μm.

**Table 5.** Exposure levels of the inorganic acids measured in various branches in Norway, registered in the EXPO database in the years 2000-2006 and 1984-1999, respectively (189).

Acid/	Branch	NACE-	No. of		Exposure leve	el
year		code <sup>a</sup>	samples	Mean (mg/m <sup>3</sup> )	Standard deviation	Maximum (mg/m <sup>3</sup> )
Sulphuric acid						
2000-2006	Mining of non-ferrous metal ores, except uranium and thorium ores	13.200	36	0.14	0.68	4.1
	Operation of dairies and cheese making	15.510	2	0.0030	0.0007	0.0035
	Manufacture of dyes and pigments	24.120	12	0.075	0.030	0.13
	Manufacture of plastics in primary forms	24.160	3	0.034	0.043	0.083
	Manufacture of hollow glass	26.130	14	0.0037	0.0029	0.011
	Manufacture of basic iron, and steel and of ferro-alloys	27.100	3	3.50	5.80	10.2
	Production of first transformation of aluminium	27.422	31	0.076	0.076	0.3
	Forging, pressing, stamping and roll forming of metal, powder metallurgy	28.400	3	0.0088	0.0089	0.018
	Treatment and coating of metals	28.510	2	0.13	0.13	0.22
	General mechanical engineering	28.520	3	0.0012	0.0003	0.0015
	Manufacture of locks and hinges	28.630	5	0.089	0.13	0.3
	Manufacture of other fabricated metal products	28.750	3	0.026	0.029	0.06
	Manufacture of marine engines and parts	29.111	5	0.028	0.011	0.047
	Scheduled air transport	62.100	11	0.018	0.016	0.048
	Freight forwarding services	63.401	1	0.006		0.006
1984-1999	Manufacture of other organic basic chemicals	24.140	1	0.005		0.005
	Casting of steel	27.520	1	0.02		0.02
	Manufacture of other special purpose machinery	29.560	6	< DL <sup>b</sup>		< DL <sup>b</sup>
	Manufacture of computers and other information processing equipment	30.020	6	0.23	0.33	0.9
	Manufacture of accumulators, primary cells and primary batteries	31.400	32	1.23	1.24	4.77

Table 5. Exposure levels of the inorganic acids measured in various branches in Norway, registered in the EXPO database in the years 2000-2006 and 1984-1999, respectively (189).

Acid/	Branch		No. of		Exposure leve	el
year			samples	Mean $(mg/m^3)$	Standard deviation	Maximum (mg/m <sup>3</sup> )
Hydrochloric	acid					
2000-2006	Manufacture of basic iron, and steel and of ferro-alloys	27.100	3	0.028	0.0099	0.036
	Forging, pressing, stamping and roll forming of metal, powder metallurgy	28.400	3	0.060	0.044	0.09
	Manufacture of locks and hinges	28.630	3	0.0033	0.0003	0.0035
	Manufacture of television and radio transmitters and apparatus for line telephone	32.200	1	0.22		0.22
	Scheduled air transport	62.100	11	0.1	0.18	0.6
	Research and experimental development in natural sciences and engineering	73.100	12	0.23	0.39	1.1
1984-1999	Treatment and coating of metals	28.510	4	0.90	0.18	1.1
	Manufacture of other special purpose machinery	29.560	6	0.25	0	0.25
	Manufacture of aircraft and spacecraft	35.300	4	0.31	0.14	0.5
Nitric acid						
2000-2006	Manufacture of basic iron, and steel and of ferro-alloys	27.100	3	0.036	0.012	0.044
	Forging, pressing, stamping and roll forming of metal, powder metallurgy	28.400	3	0.013	0.0008	0.013
	General mechanical engineering	28.520	11	0.019	0.037	0.13
	Manufacture of other machine tools	29.430	3	0.013	0.0005	0.014
	Research and experimental development in natural sciences and engineering	73.100	12	0.061	0.052	0.17
1984-1999	Manufacture of other electrical equipment	31.620	4	< DL <sup>b</sup>	0	< DL <sup>b</sup>
Phosphoric ac	id					
2000-2006	Production of first transformation of aluminium	27.422	2	0.0100	0.000	0.01
	Forging, pressing, stamping and roll forming of metal, powder metallurgy	28.400	3	0.0003	0.0001	0.0004
	General mechanical engineering	28.520	3	0.0012	0.0003	0.0015
	Scheduled motor bus transport	60.211	5	0.74	0.85	2
1984-1999	Production of first transformation of aluminium	27.422	4	0.050	0.0061	0.057

<sup>a</sup> An international coding system for industrial classification used by the European Union.
 <sup>b</sup> Measurements below the detection limits.

# 7. Toxicokinetics

#### 7.1 Deposition

The four inorganic acids may be present in the workplace air as aerosols or vapours depending on their volatility and the air temperature. Vapours containing volatile acids may be transformed to aerosols in the airways, due to the humidity and the hygroscopic nature of the compounds (50). Aerosols containing inorganic acids will be deposited in the airways as liquid particles (droplets). Adsorption of the acid to solid particles is also important for their pathogenesis (150).

The site of deposition of acid aerosols in the respiratory tract depends on e.g. the droplet size (aerodynamic diameter), the hygroscopicity of the particles and breathing pattern. Droplet size is usually the critical factor that determines the region of deposition within the respiratory tract. The diameters often reported are MMD and MMAD (see Chapter 5), of which the latter takes into account both the density of the particles and the aerodynamic drag (19). The particle size distribution of the aerosol is an important difference among various studies.

The airways are usually divided in three functional regions, the nasopharynx, the tracheobronchial region and the pulmonary region. Inert particles with an aerodynamic diameter in the range 5-30  $\mu$ m generally deposit in the nasopharynx by impaction. Smaller particles with aerodynamic diameters of 1-5  $\mu$ m deposit in the tracheobronchial regions by sedimentation, and particles with an aerodynamic diameter less than 1  $\mu$ m are deposited in the alveoli by diffusion (245).

As compared to commonly used experimental animals, humans have larger airways and a more symmetrical upper bronchial airway branching pattern. In addition, humans do considerable oral breathing, thus bypassing the air cleaning capability of the nasal airways. These differences contribute to a greater amount of upper bronchial airway particle deposition in humans as well as a tendency to greater deposition near airway bifurcations (the point at which division into two branches occurs) (159).

All four acids are hygroscopic. Hygroscopic aerosols will take on water and grow in size within the respiratory tract. H<sub>2</sub>SO<sub>4</sub> aerosols in ambient air typically have an MMAD of 0.3-0.6  $\mu$ m, while industrial aerosols can have MMADs as large as 14  $\mu$ m (160). In another study, the MMADs of acid mists in an industrial setting averaged about 5  $\mu$ m (2.6-10  $\mu$ m) (122). The size of submicrosized (< 1  $\mu$ m) H<sub>2</sub>SO<sub>4</sub> droplets has been predicted to increase by a factor of 2-3 (128). Because of their hydrophilic properties, H<sub>2</sub>SO<sub>4</sub> aerosols are deposited mainly in the upper airways and the main target organ in man is the larynx. However, aerosol droplets of small diameters may reach the alveoli (67).

The growth of  $H_2SO_4$  aerosols in the airways tends to increase respiratory retention compared to that of inert particles of the same size as the *original* acid droplets (since exhalation from small airways is hindered after expansion) as well as compared to inert particles of the same size as the *enlarged*, humified droplets (because of deeper penetration) (19).

 $HNO_3$  vapour undergoes significant removal within the upper respiratory tract due to its high water solubility and reactivity, but when other particles act as vectors it can reach the lower respiratory tract (50).

After deposition in the respiratory tract, the high moisture content results in rapid dissociation and hydration of the acids. Inhaled inorganic acid aerosols are neutralised in the upper respiratory tract in a reaction with endogenous ammonia. A mathematical model developed to simulate the growth and endogenous ammonia neutralisation of sulphate containing aerosol particles in the human respiratory tract predicted substantial growth and neutralisation of smaller particles (below 0.1  $\mu$ m) but negligible neutralisation of larger particles (above 1  $\mu$ m) (196). The levels of oral ammonia in rabbits are 0.01-1.1 ppm (247), quite similar to the levels found in humans (146, 206).

#### 7.2 Uptake

#### Sulphuric acid

Once absorbed, the sulphate ions formed become indistinguishable from sulphate derived from dietary sources. There are no data describing the extent of dermal absorption of the aerosol or liquid. Its polarity suggests little significant absorption by this route unless the acidity causes skin damage and thereby breaches the skin barrier (67, 210).

*Hydrochloric, nitric and phosphoric acids* No information found.

#### 7.3 Distribution

The four inorganic acids will be protolysed, yielding protons  $(H^+)$  dissolved in the mucosa. The anions will enter the body pool. Unless exposure is excessive, the proton and anion contributions to the body pool will be negligible, except for a high local proton-concentration that may lead to effects. Clearance from the respiratory tract will occur via the mucociliary escalator and other mechanisms such as macrophage phagocytosis and removal with the blood and lymph flow (157).

#### 7.4 Biotransformation and excretion

Not applicable, see section 7.3.

# 8. Biological monitoring

Not applicable, see section 7.2.

# 9. Mechanisms of toxicity

Exposure to the four acids in the working atmosphere can lead to a number of effects, such as sensory irritation, dental erosion, skin corrosion, changes in mucociliary and alveolar clearance, decreased pulmonary function, and increased airway reactivity. Laryngeal cancer has been observed after exposure to strong inorganic mists containing  $H_2SO_4$ .

The toxic action of acid aerosols in the airways is determined by the site of deposition. The physical state and size of the droplets in the aerosol will therefore to a large extent determine the toxicological effects. The four acids are hygroscopic, and aerosol droplets of e.g.  $H_2SO_4$  increase in size as they absorb water during transit in the airways and deposit mainly proximally in the airways (Chapter 7) (127, 239). The anions are essential and enter the body pool; at relevant exposure levels probably without causing toxicity since the contribution to systemic levels will be low. However, excessive exposure may cause toxic effects such as hyperphosphataemia, and hypocalcaemia.

The toxicity from inhalation of these inorganic acids seems to arise mainly from the free hydrogen ions. Respiratory symptoms such as cough and phlegm are related to the aerosol acidity (hydrogen ion content) (128, 239). It is also widely believed that a low pH is a major factor in dental erosion (230, 244).

 $H_2SO_4$  causes skin corrosion due to generation of excessive heat and dehydration in the tissue. In small blood vessels, necrosis and thrombosis leading to eschars may occur. Contact is generally associated with considerable pain (79).

Chemosensory effects of chemicals including inorganic acids can either be irritating (trigeminal stimulation), odorous (olfactory stimulation) or both. For odorous irritants, a clear-cut distinction between odour and irritation is difficult to make (18). The challenge of identifying chemosensory thresholds for substances causing sensory irritation has been described in depth in several papers (18, 62, 130, 178, 216, 240).

Mucociliary clearance is a major respiratory tract defence mechanism. A large number of investigations suggest that mucociliary clearance is stimulated at short-term exposure to low/intermediate concentrations of gases or aerosols such as cigarette smoke, atmospheric pollutants, and oxygen. At higher exposure levels or long-term exposure, mucociliary clearance is impaired. Mucociliary impairment caused by  $H_2SO_4$  exposure occurs in the smaller airways, the major deposition site of submicrosized aerosols (i.e. MMAD < 1  $\mu$ m). Impaired mucociliary activity has also been observed to occur in the trachea (98, 199, 246). The mechanism by which mucociliary clearance is affected by exposure to acids is unknown but changes in pH, composition and viscosity of the mucus has been shown and may play a role (239). Increased alveolar clearance seen in experimental animals due to an inflammatory response takes place when there is an influx of macrophages in the lungs after exposure to acids (176). In some cases, pulmonary function decreases as a result of bronchoconstriction. Airway reactivity is increased because of airway inflammation.

Exposure to inorganic acid mists containing  $H_2SO_4$  has been associated with laryngeal cancer in some epidemiological studies (7, 80, 217, 218, 225, 227, 243). Limited information on possible carcinogenic mechanisms of the four acids is available. Carcinoma may develop due to regenerative cell proliferation as a reaction to the cytotoxicity and irritation. Cytotoxic induced epithelial hyperplasia has been observed in monkeys (8) and rabbits (90, 204) and epithelial metaplasia in rats (127). Inhalation of inorganic acid aerosols lowers the pH in the airway mucosa and causes local irritation. Reduced pH influences chromosomal integrity. Reviews of *in vitro* studies describe that acid pH can induce e.g. chromosomal aberrations such as sister chromatid exchanges and micronuclei (106, 211, 229). The mechanisms by which chromosomal aberrations are induced by low pH are unknown (171, 229).

# 10. Effects in animals and *in vitro* studies

#### 10.1 Irritation, corrosion and sensitisation

All four acids have corrosive and irritative properties and injure skin, eyes and the mucous membranes of the respiratory tract. Studies reporting on irritative effects on the respiratory tract are described below and are summarised in Tables 7-8 (Section 10.2).

Five *in vivo* studies of different acids including  $H_2SO_4$  (concentration up to 96 %) and HCl (36 %) investigating skin burns in animals were reviewed by Flammiger and Maibach (79). The studies show the importance of early treatment to prevent skin destruction.

There are no data on sensitisation of animals after exposure to the four acids.

#### Sulphuric acid

Non-specific airway hyperresponsiveness to acetylcholine or histamine *in vitro* was investigated after exposure of male rabbits to 0.05-0.5 mg/m<sup>3</sup> (MMD 0.3  $\mu$ m) for 3 hours *in vivo*. Bronchial hyperresponsiveness was seen in all groups exposed to 0.075 mg/m<sup>3</sup> and higher. No effect was seen in the 0.05 mg/m<sup>3</sup> group. Once a concentration threshold was reached (0.075 mg/m<sup>3</sup>), no further increase in reactivity was seen. Reactivity in the trachea was increased only at the highest exposure level. According to the authors, the higher concentration needed to alter reactivity in the trachea compared to those needed for similar changes in the bronchi likely reflect dosimetry differences, i.e. there would be greater deposition of the acid within the bronchi (70).

Lung function changes at low levels were reported by Amdur *et al* who observed that a 1-hour exposure to  $0.1-1.0 \text{ mg/m}^3 \text{ H}_2\text{SO}_4$  produced significant and dose-related increases in pulmonary flow resistance in guinea pigs (sex not given). A decrease in pulmonary compliance was also produced at these low levels. In all exposures but one ( $0.1 \text{ mg/m}^3$ , MMD 1 µm), values had not returned to normal 30 minutes post-exposure. Generally, 0.3-µm particles produced greater changes

than 1- $\mu$ m particles. The smaller particles produced significant changes in both resistance and compliance during as well as after exposure to 0.1 mg/m<sup>3</sup>, whereas the larger particles produced significantly increased resistance only during exposure. Tidal volume, respiratory frequency and minute volume were not affected. Each animal served as it own control (13).

Non-specific airway hyperresponsiveness *in vivo* was induced in male guinea pigs exposed for 1 hour to ultrafine H<sub>2</sub>SO<sub>4</sub>-particles at 0.2 mg/m<sup>3</sup> (mean diameter 0.06  $\mu$ m) (48).

Male guinea pigs exposed for 1 hour to  $0.3 \text{ mg/m}^3$  (MMD  $0.08 \mu$ m) had a significantly decreased single breath diffusion capacity for carbon monoxide compared with animals exposed to 3 % argon in filtered air. Vital capacity was not altered in the acid exposed animals (46).

Ventilatory pattern (respiratory frequency, inspiratory volume and pressure changes) was monitored in male guinea pigs, exposed in whole-body plethysmographs to  $H_2SO_4$  aerosols with co-exposure to 10 % CO<sub>2</sub> to stimulate ventilation. The guinea pigs were exposed nose-only to  $H_2SO_4$  ranging from 1.8 to 55 mg/m<sup>3</sup> (1.8, 4.6, 9.2, 18, 46 and 55 mg/m<sup>3</sup>, plus two additional intermediate concentrations not stated, aerodynamic diameter 0.6  $\mu$ m) for 30 minutes. Below 10 mg/m<sup>3</sup>, the response to H<sub>2</sub>SO<sub>4</sub> was transient. An initial decline in inspiratory volume faded as the exposures continued, and was sometimes followed by an increase in volume (shown only for 9.2  $mg/m^3$ ). Also at higher concentrations, inspiratory volume decreased shortly after initiation of exposure. At 18 mg/m<sup>3</sup>, inspiratory volume fluctuated somehow, but was still diminished at termination of exposure. At 46 mg/m<sup>3</sup>, inspiratory volume continued to decrease until termination of exposure. Mean respiratory frequency was unaffected by  $H_2SO_4$  up to 18 mg/m<sup>3</sup>, but seemed to decrease at the highest exposure level  $(55 \text{ mg/m}^3)$  (198). However, respiratory frequency was not reported for exposure levels between 18 and 55  $mg/m^3$ , hampering interpretation of effects at these levels.

Five groups of 4 female guinea pigs were exposed to 10 % CO<sub>2</sub> concurrently with  $H_2SO_4$  (24, 33, 40, 49, or 73 mg/m<sup>3</sup>, MMD 0.92-1.06 µm) for 1 hour. Concentration-dependent reductions in CO<sub>2</sub>-induced increases in tidal volume, respiratory frequency, and minute ventilation were seen. The CO<sub>2</sub>-induced increase in tidal volume was significantly reduced after 30 minutes exposure in all groups, except in those exposed to 24 and 33 mg/m<sup>3</sup>. Exposure to  $H_2SO_4$  mist at and above 40 mg/m<sup>3</sup> reduced respiratory frequency. Following exposure to 73 mg/m<sup>3</sup>, the animals showed definite signs of breathing difficulty, and significantly increased tidal volume. Recovery occurred in all animals during the 5 days after exposure (249).

Ventilation was measured in female guinea pigs in whole-body plethysmographs immediately following a 4-hour inhalation exposure to  $H_2SO_4$  aerosol. Baseline respiratory pattern was monitored prior to exposure. Groups of 6 guinea pigs were exposed to 14, 20, or 43 mg/m<sup>3</sup> (MMD 0.90-0.96 µm). Doseeffect relationships were seen regarding ventilatory responses. The highest concentration caused an increase in tidal volume and a decrease in breathing frequency immediately after exposure. At 24 hours post-exposure, frequency was increased, tidal volume decreased, and breathing was interspersed with short periods of apnoea. This rapid shallow breathing was associated with histological evidence of pulmonary oedema. By contrast, the lowest  $H_2SO_4$  concentration caused increased breathing frequency with no effect on tidal volume, and 24 hours later ventilatory parameters were similar to baseline values. The middle exposure group seemed divided into two populations, with respiratory characteristics resembling that of either the high- or low-dose group (192).

In a study on rabbits performed according to OECD guidelines,  $10 \% H_2SO_4$  in water was classified not irritating to eyes (120).

#### Hydrochloric acid

HCl is extremely irritating to the eyes, mucous membranes and exposed areas of skin. Evidence of corneal erosion and clouding has been reported (64, 142, 193).

Groups of 4 male mice were exposed to 56, 139, 343, 616 or 1 320 mg/m<sup>3</sup> (40, 99, 245, 440 or 943 ppm) HCl for 10 minutes to investigate the sensory irritation of the upper respiratory tract. Each animal was placed in a plethysmograph, allowing monitoring of inspiration and expiration. Sensory irritation was measured as the percentage decrease in the respiratory rate. The animals served as their own controls. The onset of the irritative response was rapid, with a plateau being reached within 5-7 minutes from the start of exposure. The decrease in respiratory rate was dose-related, i.e. at 56 mg/m<sup>3</sup> it decreased by approximately 10 % and at 1 320 mg/m<sup>3</sup> by 70 %. Return to control values was slow after exposure to concentrations of 343 mg/m<sup>3</sup> and above. The air concentration causing a 50 % decrease in the respiratory rate (RD<sub>50</sub>) was estimated to be 432 mg/m<sup>3</sup> (309 ppm) (23).

Airway irritation was also investigated in male guinea pigs. Four or eight animals/group were exposed to 448, 952, 1 456 or 1 932 mg/m<sup>3</sup> (320, 680, 1 040, or 1 380 ppm) HCl for 30 minutes. Signs of sensory irritation appeared 6 minutes after initiation of exposure to 448 mg/m<sup>3</sup>, but was evident in less than 1 minute at the higher exposure concentrations. Also pulmonary irritation was present, and the higher the concentration, the earlier the onset. Thus, pulmonary irritation was observed after 20 minutes at 448 mg/m<sup>3</sup>, and after less than 4 minutes at 1 932 mg/m<sup>3</sup>. Impairments of the respiratory function were also supported by evidence of morphological injury in both the airways and the alveolar region (42).

Baboons (12 males) were exposed head-only to 0, 500, 5 000 or 10 000 ppm HCl (700-14 000 mg/m<sup>3</sup>) for 15 minutes in an inductive plethysmograph. The acute respiratory response consisted of a concentration-related increase in respiratory frequency and minute volume, with a marked decrease in arterial blood partial pressure of oxygen at the two highest concentrations. Pulmonary function was normal in all groups 3 days and 3 months after exposure (123).

The same research group compared the effects of HCl in mice, rats, and guinea pigs with those in baboons. Male mice were exposed to 500 ppm (700 mg/m<sup>3</sup>) or 2 500 ppm (3 500 mg/m<sup>3</sup>) HCl in plethysmographs for 15 minutes. Female rats

were exposed to 4 200 ppm (5 880 mg/m<sup>3</sup>), and male guinea pigs to 500 and 4 200 ppm HCl under similar conditions. Respiratory function and arterial blood gases were monitored during exposure, and CO<sub>2</sub>-challenge response tests were conducted prior to, 3 days and 3 months following exposure. In mice, exposure to 500 ppm caused reduced respiratory frequency (133 breaths/minute versus 178 in controls). During the 3-month follow-up period, 4/6 mice died. In guinea pigs, the respiratory frequency was also lowered after exposure to 500 ppm (97 versus 113 breaths/ minute). Exposure to the highest levels of HCl showed species differences in respiratory response and sensitivity, the rat being the less sensitive. Results from the histopathological examination are described in Section 10.2. The combined results from the two studies showed that rodents were more sensitive than baboons regarding respiratory response, probably due to anatomical differences in the nose (124).

In a study on rabbits performed according to OECD guidelines, 10 % HCl in water was classified as a risk of serious damage to eyes (120).

#### Nitric acid

No bronchoconstriction (decreased post-exposure specific pulmonary flow resistance) was observed neither in normal nor in allergic sheep (sex not given) exposed head-only for 4 hours to 4.1 mg/m<sup>3</sup> (1.6 ppm) HNO<sub>3</sub> vapour. However, allergic sheep (animals that had a history of reacting with bronchospasm to inhalation challenge with *Ascaris suum* (large roundworm) antigen) showed increased airway hyperreactivity to aerosolised carbachol after exposure to HNO<sub>3</sub>. This response is characteristic of hyperresponsive airways (2).

*Phosphoric acid* No studies were found.

#### **10.2 Effects of single exposure**

Reported lethal concentrations for 50 % of the exposed animals at single inhalation exposures ( $LC_{50}s$ ) of 30 minutes-8 hours are given in Table 6. The variation in exposure time makes a direct comparison of  $LC_{50}$  data difficult but it seems that  $HNO_3$  and  $H_2SO_4$  have a higher acute toxicity than HCl and probably also than  $H_3PO_4$ .

Other effects of single inhalation exposure to  $H_2SO_4$  and HCl are summarised in Tables 7-8, respectively. The few single exposure studies on HNO<sub>3</sub> are summarised in Table 10 (Section 10.3).

#### Sulphuric acid

The guinea pig is the most sensitive animal species, with reported  $LC_{50}$ s in the range 18-50 mg/m<sup>3</sup> (Table 6). Reported pathological findings are distended lungs, probably from acute asphyxia due to laryngeal spasm.

A large number of animal studies of the effects from single exposure are published. The studies are listed in Table 7 and are also discussed in the text.

Acid/	Median	Exposure	LC <sub>50</sub>	LC <sub>50</sub>
Species	diameter <sup>a</sup> (µm)	duration (h)	$(mg/m^3)$	(mmol/m <sup>3</sup> ) <sup>b</sup>
Sulphuric acid				
Guinea pig	NG	8	18	0.2
Guinea pig	0.8	8	40	0.4
Guinea pig	1	8	50	0.5
Mouse	NG	2	320	3.3
Rat	NG	2	510	5.2
Hydrochloric acid				
Aerosol				
Mouse	< 1	0.5	3 200	88
Mouse	< 1	0.5	12 000 <sup>c</sup>	329
Rat	< 1	0.5	7 900 °	217
Rat	< 1	0.5	8 300	228
Vapour (gas)				
Mouse		1	780	21
Mouse		1	1 600 °	44
Mouse		0.5	3 700 °	102
Mouse		0.5	3 900	107
Rat		1	2 200	60
Rat		1	4 400 <sup>c</sup>	121
Rat		0.5	6 600 <sup>c</sup>	181
Rat		0.5	7 004	192
Nitric acid (physic	al state not given)			
Rat	NG	4	130	2.1
Rat	NG	0.5	260	4.1
Phosphoric acid				
Rat	NG	1	> 850	> 8.7

**Table 6.** Acute toxicity data ( $LC_{50}s$ ) after inhalation of the inorganic acids (64, 142, 193).

<sup>a</sup> Either mass median diameter (MMD), mass median aerodynamic diameter (MMAD) or volume median diameter (VMD).

<sup>b</sup> Converted from the  $LC_{50}$  (mg/m<sup>3</sup>).

<sup>c</sup> Converted from ppm-value given in reference.

LC<sub>50</sub>: lethal concentration for 50 % of the exposed animals at single exposure. NG: not given.

The effects of  $H_2SO_4$  on pulmonary defence mechanisms have been examined in a number of studies using insoluble particles or microbes. Respiratory tract clearance may occur by ciliary transport or by phagocytosis. Respiratory clearance has been monitored by measuring the retention of an insoluble aerosol labelled with a radioisotope, usually <sup>99m</sup>Tc Fe<sub>2</sub>O<sub>3</sub>- or <sup>85</sup>Sr latex-particles. Phagocytosis has usually been studied by determining the phagocytic index, which is the percentage of viable macrophages that have ingested at least one insoluble particle, or the phagocytic capacity, which is a measure of the number of ingested particles/ phagocytising cell (often quantified as the percentage of actively phagocytising macrophages that ingest at least 4 particles). Also the production and release of mediators participating in the pulmonary defence such as tumour necrosis factor alpha (TNF $\alpha$ ) and superoxide anions have been investigated.

Male rabbits were exposed nose-only for 3 hours to  $H_2SO_4$  at 0.05 or 0.125 mg/m<sup>3</sup> (MMD 0.3  $\mu$ m). The animals were sacrificed immediately after termination of exposure, and lung lavage was performed. In the high but not the low dose group, intracellular pH in pulmonary macrophages was significantly lowered.

Intracellular pH regulation measured as the ability of pH recovery (after pH lowering with sodium propionate) was decreased in the low dose group. This was not investigated in the high dose group (49).

Depression of the release/activity of TNF $\alpha$  and reduction of superoxide anion production by the pulmonary macrophages were seen in lungs from male rabbits exposed nose-only for 2 hours to 0.075-0.5 mg/m<sup>3</sup>. The effects were assumingly immunosuppressant. No effects were seen in the lowest dose group exposed to 0.05 mg/m<sup>3</sup> (250). In another study, male rabbits exposed for 3 hours to H<sub>2</sub>SO<sub>4</sub> aerosol at 0, 0.05, 0.075, or 0.125 mg/m<sup>3</sup> (MMD 0.3 µm), after which bronchopulmonary lavage was performed, exhibited similar effects, i.e. depressed phagocytic capacity of pulmonary macrophages, and depressed superoxide anion production and TNF $\alpha$  activity of stimulated macrophages at the two highest acid levels (205). In both studies, the reductions in TNF $\alpha$  activity were of borderline significance at 0.05 mg/m<sup>3</sup>.

In male guinea pigs exposed nose-only for 3 hours to 0.3 mg/m<sup>3</sup> aqueous H<sub>2</sub>SO<sub>4</sub> droplets (MMD 0.3 or 0.04  $\mu$ m), bronchoalveolar lavage revealed an effect on the phagocytic function of macrophages. The 0.3- $\mu$ m particles enhanced the phagocytic capacity 24 hours after exposure whereas the 0.04- $\mu$ m particles depressed it. As the same concentration of H<sub>2</sub>SO<sub>4</sub> (in mg/m<sup>3</sup>) was used in both exposures, the number of droplets in the ultrafine aerosol greatly exceeded the number of droplets in the ultrafine aerosol greatly exceeded the number of droplets after exposure to both aerosols but the depression persisted after 24 hours only after ultrafine acid exposure. There was an increased release of TNF $\alpha$  from macrophages after fine acid exposure. Small and transient alterations in biochemical parameters of cellular function and viability, seen as increases in  $\beta$ -glucuronidase, lactate dehydrogenase and total protein, were observed after exposure to aerosols of both sizes (47).

Tracheal mucous clearance of radiolabelled macroaggregated albumin was measured in 8 beagle dogs (5 females and 3 males) after 1-hour exposures to  $H_2SO_4$  mist with particle sizes of 0.3 and 0.9 µm (MMAD). Velocities were significantly depressed 30 minutes, 1 day and 1 week after exposure to 1 mg/m<sup>3</sup> of the aerosol with particle size 0.9 µm. At the lower level of 0.5 mg/m<sup>3</sup>, there was a significant depression only after 1 week. Values had returned to normal 5 weeks after exposures. There were no significant changes after exposures to the 0.3-µm aerosols at the concentrations tested (1 or 5 mg/m<sup>3</sup>) (246).

No inflammatory lesions in lung parenchyma or indications of cytotoxicity were registered in male rats exposed nose-only to 0.5 and 1.0 mg/m<sup>3</sup> H<sub>2</sub>SO<sub>4</sub> (MMD 0.3  $\mu$ m) for 4 hours (131).

In male rabbits exposed to  $1 \text{ mg/m}^3$  (MMD 0.3 µm) for 1 hour, alveolar clearance of latex particles (MMAD 3.5 µm) was accelerated up to 3 days post-exposure. The numbers of polymorphonuclear neutrophils were elevated at 1 hour post-exposure in both acid exposed and control animals. In acid exposed animals, levels were still elevated after 24 hours. Also, *in vivo* phagocytosis was enhanced in the acid exposed animals during 3 hours post-exposure (176).

Tracheal mucociliary impairment was reported in male hamsters after exposure to  $H_2SO_4$  mist at levels of 0.88 mg/m<sup>3</sup> for 2 hours (VMD 0.3 µm) and 1.1 mg/m<sup>3</sup> for 3 hours (mean size 0.12 µm) (98, 199).

In a comparative study in male rabbits and humans, a single inhalation exposure  $(1 \text{ mg/m}^3 \text{ H}_2\text{SO}_4 \text{ for 3 hours}, \text{MMAD } 0.88 \ \mu\text{m})$  reduced the activity/ability of recovered macrophages to attach to a solid substrate *in vitro*, as well as the serum-opsonised zymosan-stimulated superoxide anion production. The effects were observed in both rabbits and humans in response to  $\text{H}_2\text{SO}_4$  exposure. In rabbits, the ability of bacterial uptake and intracellular killing by the pulmonary macrophages was reduced and the phagocytic capacity increased (252).

No impairment in tracheal mucous velocity was seen in sheep (sex not given) exposed to 14 mg/m<sup>3</sup> H<sub>2</sub>SO<sub>4</sub> aerosol (VMD 0.1-0.2  $\mu$ m) for 20 minutes and examined immediately after and up to 10 days later. Likewise, no impairment was seen in sheep exposed to 4 mg/m<sup>3</sup> H<sub>2</sub>SO<sub>4</sub> aerosol for 4 hours and examined 0 and 2 hours post-exposure. Similar single H<sub>2</sub>SO<sub>4</sub> aerosol exposures of anaesthetised dogs (1 or 8 mg/m<sup>3</sup> for 7.5 minutes or 4 mg/m<sup>3</sup> for 4 hours) did not produce any immediate or delayed adverse effect on cardiopulmonary function (195).

Four to six female guinea pigs in each group were exposed to 14, 20, or 43 mg/m<sup>3</sup> (MMD 0.90-0.96  $\mu$ m) H<sub>2</sub>SO<sub>4</sub> for 4 hours. Breathing frequency increased in animals exposed to the lowest concentration. Animals exposed to the highest concentration showed increased tidal volume and decreased breathing frequency immediately after exposure (Section 10.1). Analysis of fluid from bronchoalveolar lavage performed only in the 43 mg/m<sup>3</sup> exposure group showed an increased proportion of eosinophils and neutrophils, a decreased recovery of macrophages, and increased protein content. Histology showed hyaline membranes and acute inflammatory cells in the proximal acinar region. At 24 hours post-exposure, histology revealed evidence of diffuse pulmonary oedema in the highest exposed animals (192).

Chamber exposure of female guinea pigs (43 mg/m<sup>3</sup>, MMD 0.93  $\mu$ m) and female rats (94 mg/m<sup>3</sup>, MMD 0.80  $\mu$ m) to high levels of H<sub>2</sub>SO<sub>4</sub> aerosol for 4 hours had an adverse effect on the biological properties of pulmonary surfactant in the guinea pig, but not in the rat. The number of macrophages was decreased and the number of neutrophils and eosinophils was increased in bronchoalveolar lavage fluid from guinea pigs (150).

In summary, single inhalation exposures of animals to  $H_2SO_4$  (Section 10.1-10.2) give rise to airway irritation and impaired pulmonary function. At lower exposure levels, airway hyperresponsiveness and effects on the lung defences are reported. A lowest observed adverse effect level (LOAEL) of 0.075 mg/m<sup>3</sup> is indicated from studies in rabbits showing airway hyperresponsiveness as well as effects on the TNF $\alpha$  release and lowered superoxide anion production *in vitro* after exposure for 2-3 hours *in vivo*. In guinea pigs, an increased pulmonary flow resistance was observed after exposure to 0.1 mg/m<sup>3</sup> for 1 hour. At single exposure, 0.05 mg/m<sup>3</sup> can be regarded as an overall no observed adverse effect level (NOAEL).

**Table 7.** Effects in animals of single inhalation exposure to sulphuric acid.

	Exposu	re	1		Effect	Reference	
Level, mg/m <sup>3</sup>	Droplet size MD <sup>a</sup> , µm	Duration		animal/ group			
0.05	0.3	2 h, nose-only	Rabbit	5	No effects observed on pulmonary macrophages. However, reduced release/activity of $TNF\alpha$ of borderline significance (p=0.06).	(250)	
0.05	0.03	3 h, nose-only	Rabbit	5	Intracellular pH in pulmonary macrophages not lowered by the exposure, but decreased ability to recover intracellular pH after addition of a pH-lowering agent in pulmonary macrophages.	(49)	
0.05	0.3	3 h, nose-only	Rabbit	5	No effect on bronchial and tracheal reactivity after <i>in vitro</i> assessment of the airways (bronchia and tracheal rings).	ıl (70)	
0.05	0.3	3 h, nose-only	Rabbit	5	No significant effect on phagocytic capacity of pulmonary macrophages, superoxide anion production and TNF $\alpha$ activity by stimulated macrophages <i>in vitro</i> . The latter, however, was of borderline significance (p=0.06).	(205)	
0.075	0.3	2 h, nose-only	Rabbit	5	Depression of the release of TNF $\alpha$ and lowered superoxide anion production by pulmonary macrophages <i>in vitro</i> after exposure <i>in vivo</i> . Also seen at higher exposure levels (0.125 and 0.5 mg/m <sup>3</sup> ) to a similar extent.	(250)	
0.075	0.3	3 h, nose-only	Rabbit	5	Depressed phagocytic capacity of pulmonary macrophages, depressed superoxide anion production and TNF $\alpha$ activity by stimulated macrophages <i>in vitro</i> . Also seen at 0.125 mg/m <sup>3</sup> to a similar extent.	(205)	
0.075	0.3	3 h, nose-only	Rabbit	5	Non-specific bronchial hyperresponsiveness <i>in vitro</i> after exposure <i>in vivo</i> . Also seen at higher exposure levels (0.125, 0.25 and 0.5 mg/m <sup>3</sup> ) to a similar extent.	(70)	
0.1	0.3	1 h, head-only	Guinea pig	20-25	Ca 40 % increase in pulmonary flow resistance at the end of exposure. Still above control value 0.5 h post-exposure. Decrease in compliance (-27 % at the end of exposure). No effect on tidal volume, respiratory frequency or minute volume.		
0.1	1	1 h, head-only	Guinea pig	20	Ca 10-15 % increase in pulmonary flow resistance at the end of exposure. Values returned to normal within 0.5 h post-exposure. No effect on compliance, tidal volume, respiratory frequence or minute volume.	(13) cy	

			inhalation			

	Exposure		Exposure		Species	No. of	Effect	Reference
Level, mg/m <sup>3</sup>	Droplet size MD <sup>a</sup> , µm	Duration	_	animal/ group				
0.125	0.03	3 h, nose-only	Rabbit	5	Lowered intracellular pH in pulmonary macrophages.	(49)		
0.2	0.06 (mean)	1 h	Guinea pig	6-8	Non-specific airway hyperresponsiveness assessed 1.5 h post-exposure.	(48)		
0.25	0.3	3 h, nose-only	Rabbit	5	No increase in tracheal reactivity after in vitro assessment of the airways (tracheal rings).	(70)		
0.3	0.08	1 h	Guinea pig	7-10	Significantly decreased single breath diffusion capacity for carbon monoxide. No effect on vital capacity.	(46)		
0.3	0.04	3 h, nose-only	Guinea pig	6	Small changes in biochemical parameters and depressed phagocytic capacity. Depression of intracellular pH of alveolar macrophages immediately and 24 hours after exposure.	(47)		
0.3	0.3	3 h, nose-only	Guinea pig	6	Small changes in biochemical parameters and enhanced phagocytic capacity. Increased $TNF\alpha$ release from alveolar macrophages. Depression of intracellular pH of macrophages immediately after exposure.	(47)		
0.4	1	1 h, head-only	Guinea pig	20	Ca 30 % increase in pulmonary flow resistance at the end of exposure. Still above control values 0.5 h post-exposure. No effect on compliance, respiratory frequency, tidal or minute volumes.	(13)		
0.5	0.3	1 h, head-only	Guinea pig	20	Ca 60 % increase in pulmonary flow resistance at the end of exposure. Still above control values 0.5 h post-exposure. Decrease in compliance (-33 % at the end of exposure), still below control values 0.5 h post-exposure. No effect on tidal volume, respiratory frequency or minute volume.	(13)		
0.5	0.3	3 h, nose-only	Rabbit	5	Increased tracheal reactivity after in vitro assessment of the airways (tracheal rings).	(70)		
0.5	0.9	1 h, nose-only	Dog	8	Significant depression of tracheal mucociliary clearance 1 week after exposure. Velocities had returned to normal 5 weeks after exposure. No significant effects 30 min or 1 day after exposure.	(246)		
0.7	1	1 h, head-only	Guinea pig	20	Ca 45 % increase in pulmonary flow resistance at the end of exposure. Still above control values 0.5 h post-exposure. Decrease in compliance. No effect on tidal volume, respiratory frequency or minute volume.	(13)		

Table 7. Effects	in	animals	of	singl	e inha	lation	exposure	to	sulphuric acid.	

	Exposu	re	Species	No. of	Effect	Reference
Level, mg/m <sup>3</sup>	Droplet size MD <sup>a</sup> , µm	Duration		animal/ group		
0.85	1	1 h, head-only	Guinea pig	20	Ca 60 % increase in pulmonary flow resistance at the end of exposure. Still above control value 0.5 h post-exposure. Decrease in compliance. No effect on tidal volume, respiratory frequency or minute volume.	s (13)
0.88	0.3	2 h	Hamster	10	Depressed ciliary beating in trachea.	(98)
1	0.3	4 h, nose-only	Rat	20	No inflammatory or cytotoxic effects seen at this exposure level, nor at 0.5 mg/m <sup>3</sup> .	(131)
1	0.3	1 h, head-only	Guinea pig	20	Ca 78 % increase in pulmonary flow resistance at the end of exposure. Still above control value 0.5 h post-exposure. Decrease in compliance (-40 % at end of exposure), still below control values 0.5 h post-exposure. No effect on respiratory frequency, tidal volume or minute volume.	s (13)
1	0.88	3 h	Rabbit	6	Reduced activity/ability of recovered macrophages to attach to a solid substrate <i>in vitro</i> , and serum-opsonised zymosan-stimulated superoxide anion production. Reduced ability of bacterial uptake and of intracellular killing by pulmonary macrophages. Increased phagocytic capacity.	(252)
1	0.3	1 h, nose-only	Dog	8	No significant depression of tracheal mucociliary clearance 30 min, 1 day or 1 week post- exposure.	(246)
1	0.9	1 h, nose-only	Dog	8	Significant depression of tracheal mucociliary clearance 30 min, 1 day and 1 week after exposure. Velocities had returned to normal 5 weeks after exposure.	(246)
1	0.3	1 h	Rabbit	30	Transiently increased alveolar clearance of tracer latex particles (MMAD 3.5 µm). Prolonged increase in PMN count. Enhanced <i>in vivo</i> phagocytosis during 3 h post-exposure.	(176)
1.1	0.12	3 h	Hamster	12	Depressed ciliary beating and damaged respiratory epithelium in the trachea.	(199)
4	0.1-0.2	4 h	Dog	5	No effect on cardiopulmonary function in anaesthetised animals.	(195)
4	0.1-0.2	4 h	Sheep	10	No mucociliary impairment in trachea up to 10 days after exposure.	(195)

<b>Table 7.</b> Effects in animals of single inhalation exposure to sulphuric acid.
---

	Exposu	re	Species	No. of	Effect	Reference	
Level, mg/m <sup>3</sup>	2		_	animal/ group			
5	0.3	1 h, nose-only	Dog	8	No significant depression of tracheal mucociliary clearance 30 min, 1 day or 1 week after exposure.	(246)	
8	0.1-0.2	7.5 min	Dog	5	No effect on cardiopulmonary function in anaesthetised animals at this or lower (1 mg/m <sup>3</sup> ) level	. (195)	
9.2 <sup>b</sup>	0.6	30 min	Guinea pig	4	No effect on respiratory frequency. Transient initial decline in inspiratory volume faded as the exposures continued, which was followed by an increase in volume. Same results obtained at 1.8 and 4.6 mg/m <sup>3</sup> (i.e. the slope in the exposure-response curve approximated zero at concentrations between 1 and 10 mg/m <sup>3</sup> ).	(198)	
14	0.1-0.2	20 min	Sheep	10	No mucociliary impairment in trachea up to 10 days after exposure.	(195)	
14	0.90-0.96	4 h	Guinea pig	6	Increased breathing frequency. No effect on tidal volume.	(192)	
18 <sup>b</sup>	0.6	30 min	Guinea pig	4	No effect on respiratory frequency. Inspiratory volume decline that remained at termination of exposure. At a higher exposure level (46 mg/m <sup>3</sup> ), the decline continued after exposure.	(198)	
20	0.90-0.96	4 h	Guinea pig	6	Signs of dyspnoea.	(192)	
33 <sup>b</sup>	1	1 h, head-only	Guinea pig	4	No significant effects on respiratory frequency, tidal volume or minute volume. Tendency to reduction of $CO_2$ -induced increase in tidal volume.	(249)	
40 <sup>b</sup>	1	1 h, head-only	Guinea pig	4	The $CO_2$ -induced increase in tidal volume and respiratory frequency was reduced. Also seen in a dose-dependent manner in groups exposed to 49 or 73 mg/m <sup>3</sup> .	(249)	
43	0.93	4 h	Guinea pig	10	Pulmonary surfactant adversely altered. Decreased number of macrophages and increased number of neutrophils and eosinophils.	(150)	
43	0.90-0.96	4 h	Guinea pig	6	Increase in tidal volume and decrease in breathing frequency immediately after exposure. Pulmonary oedema. Increased coughing and dyspnoea, changes in bronchoalveolar lavage (increased proportion of neutrophils and eosinophils, decreased recovery of macrophages) indicating lung injury.	(192)	

Table 7. Effects in animals of single inhalation exposure to sulphuric acid.

	Exposu	ire	Species	No. of	Effect	Reference
Level, mg/m <sup>3</sup>	Droplet size MD <sup>a</sup> , µm	Duration	_	animal/ group		
73 <sup>b</sup>	1	1 h, head-only	Guinea pig	4	Definite signs of breathing difficulty.	(249)
94	0.80	4 h	Rat	6	Pulmonary surfactant not adversely altered. No changes in the number of macrophages, neutrophils and eosinophils.	(150)

<sup>a</sup> Median diameter, either mass median diameter (MMD), mass median aerodynamic diameter (MMAD) or volume median diameter (VMD). <sup>b</sup> In 10 % CO<sub>2</sub>.

MD: median diameter, PMN: polymorphonuclear leukocytes, TNF: tumour necrosis factor.

#### Hydrochloric acid

Results from the  $LC_{50}$ -studies have shown that the respiratory tract is the primary target for HCl both as vapour and aerosol with effects such as emphysema and pulmonary oedema.

Mice seem to be more sensitive to HCl exposure than rats. In one study, the  $LC_{50}$  after 30 minutes of aerosol exposure was 8 300 mg/m<sup>3</sup> in rats and 3 200 mg/m<sup>3</sup> in mice, and the corresponding values for vapour were 6 600 mg/m<sup>3</sup> and 3 700 mg/m<sup>3</sup> (Table 6). The cause of death was the effects on the respiratory tract (64).

Male mice were exposed to 700 or 3 500 mg/m<sup>3</sup> HCl in plethysmographs for 15 minutes. Female rats were exposed to 5 880 mg/m<sup>3</sup>, and male guinea pigs to 700 and 5 880 mg/m<sup>3</sup> HCl under similar conditions. Histopathological examination was carried out 3 months post-exposure. Exposure to 3 500 mg/m<sup>3</sup> in mice and 5 880 mg/m<sup>3</sup> in guinea pigs caused severe damage to the respiratory tract and was lethal to some animals, while rats exposed to 5 880 mg/m<sup>3</sup> experienced eye damage, but only minimal morphological changes to the respiratory tract 3 months post-exposure. Exposure to 700 mg/m<sup>3</sup> HCl did not cause any significant morphological changes in either the mouse or the guinea pig (124).

Exposure of male rats to 1 300 ppm HCl (1 820 mg/m<sup>3</sup>) in plethysmographs via nasal breathing for 30 minutes caused marked toxicity after 24 hours in the nasal region, seen as epithelial and submucosal necrosis, accumulation of inflammatory cells, exudates and the extravasion of erythrocytes. Even higher toxicity (including an increased number of deaths) was seen after forced mouth breathing. In the airways, effects such as major tissue disruption in the trachea and accumulation of inflammatory cells and exudates were observed. Also more peripheral lung damage was demonstrated after forced mouth breathing than after nasal breathing. Breathing frequency was reduced by 4 %, minute ventilation by 6 %, and tidal volume by 7 % in exposed nose breathers compared to the controls. In the mouth breathing group, minute ventilation increased during exposure (223).

Acid instillation (0.1 ml HCl) into the left bronchus stimulated alveolar macrophages to produce TNF $\alpha$  in both lungs of male rats (145).

In summary, single exposure to HCl causes airway irritation seen as decreased respiratory rate, and at higher concentrations epithelial and submucousal necrosis and pulmonary congestion.

Details are given in Table 8.

Exposur	e level	Exposure	Species	No. of	Effect	Reference
mg/m <sup>3</sup>	ppm	duration	-	animal/ group		
56	39	10 min	Mouse	4	10 % respiratory depression.	(23)
432	309	10 min	Mouse	4	RD <sub>50</sub> .	(23)
448	320	30 min, head-only	Guinea pig	4	Sensory irritation onset after 6 min exposure. Pulmonary irritation onset after > 20 min exposure. Reduced breathing frequency.	(42)
700	500	15 min, head-only	Mouse	6	Reduced respiratory frequency 3 days and 3 months after exposure, also seen in the high dose group (3 500 mg/m <sup>3</sup> ). No morphological changes in the respiratory tract. 4 exposure-related deaths.	(124)
700	500	15 min, head-only	Guinea pig	9	Reduced respiratory frequency 3 days and 3 months after exposure, also seen in the high dose group (5 880 mg/m <sup>3</sup> ). No morphological changes in the respiratory tract. One death, most likely due to suffocation (20 min).	(124)
700	500	15 min, head-only	Baboon	3	No differences from control group regarding respiratory frequency 3 days and 3 months after exposure. Tendency to an increased respiratory rate and minute volume during exposure, which was significant at 7 000 and 14 000 mg/m <sup>3</sup> .	(123)
952	680	30 min, head-only	Guinea pig	4	Sensory irritation onset immediately. Pulmonary irritation after ca. 13 min exposure. Reduced breathing frequency.	(42)
1 456	1 040	30 min, head-only	Guinea pig	8	Sensory irritation onset immediately. Pulmonary irritation after ca. 9 min exposure. Persistent reduced breathing frequency. Alveolitis and congestion. Squamous metaplasia and loss of cilia in the larger conducting airways (only group examined lung morphologically). 2/8 died within the 16-day period. Corneal opacities in 4/6 survivors.	(42)
1 820	1 300	30 min, nose-only	Rat	5-8	Reduced breathing frequency, minute ventilation and tidal volume. Epithelial and submucousal necrosis, accumulation of inflammatory cells, exudates and extravasion of erythrocytes in the nasal region.	(223)

**Table 8**. Effects in animals of single inhalation exposure to hydrogen chloride.

Table 8. Effects in animal	s of single inhalation	exposure to hydrogen chloride.
----------------------------	------------------------	--------------------------------

		L L		I		
Exposur mg/m <sup>3</sup>	e level ppm	Exposure duration	Species	No. of animal/ group	Effect	Reference
1 820	1 300	30 min, mouth-only	Rat	5-8	Increased minute ventilation. Deaths and major tissue disruption in the trachea and accumulation of inflammatory cells and exudates. Peripheral lung damage.	(223)
1 932	1 380	30 min, head-only	Guinea pig	8	Sensory irritation onset immediately. Pulmonary irritation in less than 4 min after onset of exposure. Persistent reduced breathing frequency. 3/8 died within the 16-day period. Corneal opacities in all survivors.	(42)
3 500	2 500	15 min, head-only	Mouse	6	Reduced respiratory frequency 3 days and 3 months after exposure. Morphological changes: moderate to severe lung congestion, necrosis of the tracheal mucosa, exudate in the paranasal sinuses. Lung oedema in 1 animal. 5 exposure-related deaths.	(124)
5 880	4 200	15 min, head-only	Guinea pig	9	Reduced respiratory frequency 3 days and 3 months after exposure. Decreased pH in arterial blood. Morphological changes: severe pulmonary congestion with minimal oedema, congestion of the nasal turbinates, severe tracheitis and desquamation of the epithelia of the bronchi and bronchioles. Cloudy corneas. 3 exposure-related deaths.	(124)
5 880	4 200	15 min, head-only	Rat	9	Reduced respiratory frequency 3 days and 3 months after exposure. Increase in PaCO <sub>2</sub> -values in arterial blood. Eye damage, minimal histopathological changes to the respiratory tract. No exposure-related deaths.	(124)
7 000	5 000	15 min, head-only	Baboon	3	Increased respiratory frequency and minute volume and decreased blood PaO <sub>2</sub> .	(123)
14 000	10 000	15 min, head-only	Baboon	3	Same changes as at 7 000 mg/m <sup>3</sup> but more pronounced.	(123)

PaCO<sub>2</sub>: partial pressure of carbon dioxide, PaO<sub>2</sub>: partial pressure of oxygen, RD<sub>50</sub>: air concentration associated with a 50 % decrease in the respiratory rate; a measure of sensory irritation.

## Nitric acid

In rats, the LC<sub>50</sub> was 260 mg/m<sup>3</sup> after 30 minutes and 130 mg/m<sup>3</sup> after 4 hours (Table 6). Details of toxic effects other than death were not reported (193).

In an old study published 1907, rabbits and cats tolerated 15 ppm  $(38 \text{ mg/m}^3)$  for 3 hours without visible damage (Diem<sup>1</sup>, cited in (66)).

In male Fischer 344 rats, single exposure to  $1 \text{ mg/m}^3 \text{ HNO}_3$  vapour for 4 hours resulted in a modest but significant increase in elastase inhibitory capacity. Lavage fluid protein was unchanged. The same total dose was also given as repeated exposure (Table 10, Section 10.3) (175). The implications of the findings for human health are unclear.

Intratracheal instillation of 0.5 ml of 0.08 N HNO<sub>3</sub> in male hamsters caused secretory cell metaplasia in the bronchial airway epithelium and interstitial fibrosis, bronchiolectasis and bronchiolisation of alveoli (52). Similar results were obtained by Coalson *et al* (54). The relevance of these findings is unclear since the conversion of instilled doses to concentrations in inhaled air is difficult.

### Phosphoric acid

The  $LC_{50}$  was above 850 mg/m<sup>3</sup> in rats exposed for 1 hour (Table 6). Details of other toxic effects other than death were not reported.

# 10.3 Effects of short-term exposure (up to 90 days)

## Sulphuric acid

Details are given in Table 9.

Respiratory region clearance was studied in male rabbits exposed to clean air,  $H_2SO_4$  at 0.05 mg/m<sup>3</sup> (MMAD 0.3 µm) for 1, 2, or 4 hours/day for 14 days or at 0.1 mg/m<sup>3</sup> for 0.5, 1, or 2 hours/day for 14 days. Exposure to 0.05 mg/m<sup>3</sup> for 4 hours/day or 0.1 mg/m<sup>3</sup> at 2 hours/day (same daily dose) resulted in accelerated clearance of radiolabelled latex particles (MMAD 3.5 µm) (201).

Exposure of male rats to 0.02, 0.1, or 0.15 mg/m<sup>3</sup> of  $H_2SO_4$  aerosol (0.4-0.8 µm diameter) alone for up to 90 days did not affect body weight, lung lobe weight, or biochemical parameters and morphological measurements relating to pulmonary fibrogenesis (148).

No alterations in biochemical parameters except a transient increase in  $\beta$ -glucuronidase were seen in bronchoalveolar lavage from male guinea pigs exposed nose-only for 3 hours/day for 4 days to 0.3 mg/m<sup>3</sup> aqueous H<sub>2</sub>SO<sub>4</sub> droplets (MMD 0.3 or 0.04 µm). This contrasts findings in the same study after single exposure at the same concentration (Section 10.2). In agreement with the results from single exposure, the TNF $\alpha$  releases from alveolar macrophages were increased by aerosols of both sizes. The smallest, ultrafine droplets depressed the phagocytic capacity (ingestion of 3-µm latex particles) of the alveolar macrophages, while the largest particles enhanced it, both after single and multiple exposures. This finding

<sup>&</sup>lt;sup>1</sup> Diem L. Untersuchungen über die Einatmung von Saltpetersäure-dämpfen (thesis). Würzburg, 1907.

was regarded to be a result of the larger number of droplets being delivered to the macrophages from ultrafine acid aerosols. After repeated, but not single, exposure to both aerosols, the  $H_2O_2$ -production from macrophages was increased. Intracellular pH was depressed 24 hours after multiple exposures to ultrafine droplets (47).

The phagocytic activity of alveolar macrophages was studied in male rabbits exposed to 0.25, 0.5, 1 or 2 mg/m<sup>3</sup> H<sub>2</sub>SO<sub>4</sub> (MMAD 0.3  $\mu$ m) 1 hour/day for 5 days. The phagocytosis of latex spheres (3  $\mu$ m diameter) *in vitro* was reduced by exposure at and above 0.5 mg/m<sup>3</sup> in a dose-dependent manner (202).

In male rabbits exposed to  $H_2SO_4$  (MMD 0.3 µm) in concentrations of 0.25, 0.5 or 1 mg/m<sup>3</sup>, 1 hour/day for 5 days, the levels of the eicosanoids (arachidonic acid metabolites) prostaglandins E2 and F2(a) and thromboxane B2 in lung lavage fluid decreased in a dose-related way. Thromboxane B2 values were significantly different from controls already at the lowest concentration of  $H_2SO_4$ . Tracheal explants exposed to acidic environments *in vitro* also showed reduced production of eicosanoids. The sulphate ion (Na<sub>2</sub>SO<sub>4</sub>) did not elicit such an effect. The hydrogen ion was therefore assumed to be the causative agent. Eicosanoids are potent mediators of smooth muscle tone and the inflammatory response. According to the authors, modulation of their metabolism may be a possible factor in the pathogenesis of lung disease (203).

Significant histological (squamous metaplasia) and cell proliferative changes were seen in larynges of female rats after exposure to 1.38 and 5.52 mg H<sub>2</sub>SO<sub>4</sub>/m<sup>3</sup> (MMAD 0.83 and 0.94  $\mu$ m) for 6 hours/day for 5 days, or for 5 days/week over a 28-day period. Following exposure to the highest level, parakeratosis was seen in some animals. Partial recovery was observed after 4 weeks. No effects were observed at 0.3 mg/m<sup>3</sup> (MMAD 0.62  $\mu$ m) after 5 days and only minimal squamous metaplasia in 6/10 animals compared to 0/10 among controls, and no increased cell proliferation after 28 days. The response at the lowest concentration was considered to be an adaptive response. Statistics were not reported. No compoundrelated histopathological changes in the lung or nasal cavity were seen after any exposure (127).

Significant reductions in uptake and intracellular killing of bacteria, in superoxide anion radical production, and in TNF $\alpha$  activity by male rabbit pulmonary macrophages were seen *in vitro* after nose-only inhalation for 2 hours/day for 4 days to 0.75 and 1 mg/m<sup>3</sup> H<sub>2</sub>SO<sub>4</sub> aerosol (MMAD 0.3 µm). Exposure to 0.5 mg/m<sup>3</sup> stimulated the superoxide anion production. No clear dose-effect relationships were seen in the study. Once a critical exposure level was reached (0.75 mg/m<sup>3</sup>), no further effects were observed (251).

Antigen-induced histamine release from isolated lung mast cells *in vitro* of male guinea pigs was significantly enhanced by *in vivo* continuous exposure to 1 and  $3.2 \text{ mg/m}^3 \text{ H}_2\text{SO}_4$  aerosol (MMD < 0.8 µm) for 2 weeks. Also non-specifically induced (A23187) histamine release was enhanced by exposure for 2 weeks at 1 mg/m<sup>3</sup>. The effects were not seen in animals exposed continuously for 4 weeks. In contrast, A23187-induced histamine release was suppressed by exposure to 3.2

 $mg/m^3$ , but significantly only after 4 weeks. No changes in the number of lung mast cells were seen in any of the groups. Exposure to 0.3  $mg/m^3$  for 2 or 4 weeks elicited no changes in histamine release (84).

When groups of 10 male Sprague Dawley rats were exposed 4 hours/day for 2 days to  $0.5 \text{ mg/m}^3$  of H<sub>2</sub>SO<sub>4</sub> (two groups: MMD 0.3 µm or 0.06 µm), no morphological changes of the lungs, and no effects on cellular proliferation in the pulmonary parenchyma or in ventilatory parameters were seen compared with sham exposure except for a slight decrease in the minute volume on the second day in the group exposed to particles with MMD 0.06 µm. Co-exposure with ozone revealed that both acid aerosols increased ozone-induced morphologic effects (128).

Continuous exposure to 1 or  $3.2 \text{ mg/m}^3 \text{ H}_2\text{SO}_4$  aerosol (MMAD 0.54-0.56 µm) for 3, 7, 14, or 30 days in male guinea pigs had no effect on specific airway resistance. In the high dose group, the airway responsiveness to histamine treatment was transiently stimulated after 3 days of exposure and transiently inhibited after 14 days of exposure. It returned to normal levels after 7 and 30 days of exposure, respectively (133).

To conclude, repeated, short-term inhalation of  $H_2SO_4$  affected the defences of the rabbit lung. An accelerated respiratory clearance was reported in animals exposed to 0.05 mg/m<sup>3</sup>, 4 hours/day for 14 days (LOAEL) or to 0.1 mg/m<sup>3</sup>, 2 hours/day for 14 days. Effects on inflammatory response mediators appeared at 0.25 mg/m<sup>3</sup> (1 hour/day for 5 days). At 0.3 mg/m<sup>3</sup>, effects on TNF $\alpha$  release,  $H_2O_2$ production, phagocytosis, and intracellular pH (3 hours/day for 4 days) as well as minimal squamous metaplasia in ciliated epithelium of rats (6 hours/day, 5 days/ week for 28 days) were demonstrated. The effects in the studies were observed at the lowest levels tested.

#### Hydrochloric acid

In male guinea pigs exposed 2 hours/day, 5 days/week for 7 weeks, a NOAEL of 15 mg/m<sup>3</sup> was established for effects on pulmonary function and histological changes in the lungs and airways (181).

Groups of 16-24 male mice were exposed to HCl vapour for 6 hours/day at the  $RD_{50}$  concentration of 309 ppm (432 mg/m<sup>3</sup>). After 3 exposures, all animals were dead or moribund. Severe exfoliation, erosion, ulceration and necrosis, and mild inflammation of the respiratory epithelium were observed, as well as mild ulceration and necrosis of the olfactory epithelium, and serous exudate. No lesions were induced in the lower respiratory tract (41).

# Nitric acid

Details are given in Table 10 (including single inhalation exposures).

Male rabbits (New Zealand white) exposed nose-only 4 hours/day, 3 days/week for 4 weeks to HNO<sub>3</sub> vapour (0.05, 0.15 and 0.45 mg/m<sup>3</sup>) showed effects in the alveolar macrophages (lowered production of superoxide anions) already in the low dose group, and reduced *in vitro* bronchial responsivity in the groups exposed to  $\geq 0.15$  mg/m<sup>3</sup>. Tracheal responsivity was not affected. There was an apparent

concentration-related trend toward a reduction in TNF $\alpha$ -activity, which at  $\geq 0.15$  mg/m<sup>3</sup> departed significantly from those of controls. This study indicates that HNO<sub>3</sub> impacts both the conducting and respiratory airways. The reason for reduced bronchial responsiveness is not clear, but the authors speculate that formation of airway smooth muscle relaxants such as NO-containing chemical species or *S*-nitrosothiols is possible (207).

In male Fischer 344 rats, short-term exposure (4 hours/day, 4 days) to 0.25 mg/m<sup>3</sup> HNO<sub>3</sub> vapour decreased respiratory burst activity (superoxide anion production) and increased elastase inhibitory capacity as measured in lung lavage fluid. Elastase inhibitory capacity was increased also when the same total dose was given as a single exposure (1 mg/m<sup>3</sup>, 4 hours) (Section 10.2). Lavage fluid protein was unchanged by both exposures (175). The implications for humans of changes in elastase inhibitory capacity are unclear.

Seven dogs (sex not given) were exposed to nebulised 1 % HNO<sub>3</sub>, 3 days/week for 4 weeks (185). The concentration in air is not given in the paper but may be estimated from the delivered volume (approximately 13 ml corresponding to 130 mg HNO<sub>3</sub>), the exposure duration (approximately 2 hours), the body weight (15-19 kg), the tidal volume (15 ml/kg body weight corresponding to 0.23-0.29 l) and the breathing frequency (15-20 min<sup>-1</sup>). The inhaled air volume is thus between 0.4 and 0.7 m<sup>3</sup> and the average air level during each 2-hour exposure 190-320 mg/m<sup>3</sup> (80-130 ppm). Lung function tests revealed both restrictive and obstructive impairment. Histopathology showed central as well as peripheral damage with inflammation, epithelial damage, peribronchiolar fibrosis and an increase in smooth muscle. Further, hyperresponsiveness to histamine developed 3-5 months after exposure. Similar findings were made in another study of 7 mongrel dogs exposed to the same concentration of HNO<sub>3</sub> on alternate days for 4 weeks (85).

### Phosphoric acid

No peer-reviewed studies were found.

<b>Table 9.</b> Effects in animals of short-term exposure to sulphuric acid.
--

	Exposure		Species	No. of	Effect	Reference
Level, mg/m <sup>3</sup>			or out			
0.05	0.3	1, 2, or 4 h/day for 14 days	Rabbit	5	Accelerated respiratory clearance of latex particles (MMAD 3.5 $\mu$ m) after the 4-hour/day exposure.	(201)
0.1	0.3	0.5, 1, or 2 h/ day for 14 days	Rabbit	5	Accelerated respiratory clearance of latex particles (MMAD 3.5) after the 2-hour/day exposure.	(201)
0.15	0.4-0.8	23.5 h/day for up to 90 days	Rat	6	No effect on body weight, lung lobe weight or biochemical parameters and morphological measurements relating to pulmonary fibrogenesis. Same results at $0.02$ and $0.1 \text{ mg/m}^3$ .	(148)
0.25	0.3	1 h/day, 5 days	Rabbit	3	No effect on alveolar phagocytosis of latex spheres (diameter 3 µm) in vitro.	(202)
0.25	0.3	1 h/day for 5 days, nose-only	Rabbit	5	Decreased level of the eicosanoid <sup>b</sup> $TxB_2$ in bronchopulmonary lavage fluid. Also seen at 0.5 mg/m <sup>3</sup> for the eicosanoids $TxB_2$ and $PGE_2$ and at 1 mg/m <sup>3</sup> for $TxB_2$ , $PGE_2$ and $PGF_{2\alpha}$ .	(203)
0.3	0.04	3 h/day for 4 days, nose-only	Guinea pig	6	Increased TNF $\alpha$ release and H <sub>2</sub> O <sub>2</sub> -production from alveolar macrophages. Depressed <i>in vitro</i> phagocytic capacity. No changes in biochemical parameters. Depressed intracellular pH 24 hours after exposure.	(47)
0.3	0.3	3 h/day for 4 days, nose-only	Guinea pig	6	Increased TNF $\alpha$ release and H <sub>2</sub> O <sub>2</sub> -production from alveolar macrophages. Enhanced <i>in vitro</i> phagocytic activity. No changes in biochemical parameters.	(47)
0.3	0.62	6 h/day for 5 or 28 days	Rat	10	No histopathological changes in the lung or nasal cavity. No increase in cell proliferation in the larynx. At 28 days, minimal squamous metaplasia in ciliated epithelium in the larynx (6/10 compared to 0/10 in controls). Also seen at 1.38 and 5.52 mg/m <sup>3</sup> to a greater extent.	
0.3	0.65	Continuously for 2 or 4 weeks	Guinea pig	3-4	No changes in antigen- or non-specifically (A23187) induced histamine release in lung mast cells or in the number of lung mast cells.	(84)
0.5	0.06 or 0.3	4 h/day for 2 days, nose-only	Rat	10	No effect on morphology, cellular proliferation in the pulmonary parenchyma or ventilatory parameters except for a slight decrease in the minute volume on the 2nd day in the group exposed to particles with MMD $0.06 \mu$ m.	(128)

Table 9. Effects	in	animals	of	short-term	exposure	to sul	phuric ac	id.

	Expos	ure	Species	No. of	Effect	Reference	
Level, mg/m <sup>3</sup>				animal/ group			
0.5	0.3	1 h/day for 5 days	Rabbit	3	Reduced alveolar phagocytosis (phagocytic capacity) of latex spheres (3 $\mu$ m diameter) <i>in vitro</i> . The effect increased in a dose-dependent manner at 1 and 2 mg/m <sup>3</sup> .	(202)	
0.5	0.3	2 h/day for 4 days, nose-only	Rabbit	5	No immunosuppressive effects <i>in vitro</i> after exposure <i>in vivo</i> . Stimulated superoxide anion production by pulmonary macrophages.	(251)	
0.75	0.3	2 h/day for 4 days, nose-only	Rabbit	5	Immunosuppressive effects <i>in vitro</i> after exposure <i>in vivo</i> (reduction in uptake and intracellular killing of bacteria, in superoxide anion production and in TNF $\alpha$ activity). The same effects of similar magnitude were also seen at the highest exposure level (1 mg/m <sup>3</sup> ).	(251)	
1	0.5	Continuously, 3, 7, 14 or 30 days	Guinea pig	72	No effect on specific airway resistance and airway hyperresponsiveness to histamine.	(133)	
1	0.55	Continuously, 2 or 4 weeks	Guinea pig	3-4	Enhanced antigen as well as non-specifically (A23187) induced histamine release in the lung mast cells <i>in vitro</i> after 2 weeks exposure <i>in vivo</i> but not after 4 weeks. No changes in the number of lung mast cells.	(84)	
1.38	0.83	6 h/day for 5 days	Rat	10	Squamous metaplasia in ciliated epithelium of the larynx. Increased cell proliferation in larynx. No histopathological changes in the lung or nasal cavity. Effects were dose-dependent and seen also in rats exposed to 5.52 mg/m <sup>3</sup> (parakeratosis in some animals).	(127)	
3.2	0.5	Continuously, 3, 7, 14 or 30 days	Guinea pig	72	Airway responsiveness to histamine treatment transiently stimulated and later inhibited during exposure. No effect on specific airway resistance.	(133)	
3.2	0.73	Continuously, 2 or 4 weeks	Guinea pig	3-4	Enhanced antigen-induced histamine release from isolated lung mast cells <i>in vitro</i> after exposure <i>in vivo</i> after 2 weeks but not after 4 weeks. Reduced non-specifically induced (A23187) histamine release, significant after 4 weeks but not after 2 weeks. No changes in the number of lung mast cells.	(84)	

<sup>a</sup> Median diameter, either mass median diameter (MMD), mass median aerodynamic diameter (MMAD) or volume median diameter (VMD).
 <sup>b</sup> Eicosanoids are potent mediators of smooth muscle tone and the inflammatory response.
 MD: median diameter, PG: prostaglandin, TNF: tumour necrosis factor, Tx: thromboxan.

Exposure mg/m <sup>3</sup>	level ppm	Exposure duration	Species	No. of animals /group	Effect	Reference
0.05	0.02	4 h/day, 3 days/ week, 4 weeks, nose-only	Rabbit	6	Reduced production of superoxide anions in the alveolar macrophages, seen also at $0.15$ and $0.45 \text{ mg/m}^3$ . No effect on viability or numbers of cells.	(207)
0.15	0.06	4 h/day, 3 days/ week, 4 weeks, nose-only	Rabbit	6	Reduced bronchial reactivity to smooth muscle constrictor challenge (acetylcholine and histamine) and a reduction in TNF $\alpha$ activity in alveolar macrophages. The same effects were seen in the higher exposed group (0.45 mg/m <sup>3</sup> ) but there were no clear dose-effect relationships.	(207)
).25	0.1	4 h/day for 4 days, nose-only	Rat	10	Decreased spontaneous and stimulated respiratory burst activity (superoxide anion production) in isolated pulmonary macrophages. Increase in elastase inhibitory capacity of lung lavage fluid. No change in lavage fluid protein.	(175)
1	0.4	4 h, nose-only	Rat	10	Increase in elastase inhibitory capacity of lung lavage fluid. No change in lavage fluid protein content.	(175)
4.1	1.6	4 h, head-only	Sheep, normal and allergic	7	No bronchoconstriction (decreased specific pulmonary flow resistance post-exposure). Allergic sheep showed increased airway hyperreactivity to aerosolised carbachol.	(2)
190-320 <sup>a</sup>	80-130 <sup>a</sup>	3 days/week for 4 weeks	Dog	7	Bronchial injury was induced, with airway obstruction and chronic inflammation of the small airways.	(85)
190-320 <sup>a</sup>	80-130 <sup>a</sup>	Alternate days for 4 weeks	Dog	7	Bronchial injury was induced, with airway obstruction and chronic inflammation of the small airways.	(185)

**Table 10.** Effects in animals of single and short-term inhalation exposure of nitric acid vapour.

<sup>a</sup> Administered as spray, exposure level estimated by the Nordic Expert Group, for details see the text. TNF: tumour necrosis factor.

# **10.4 Mutagenicity and genotoxicity**

No data were available on the genetic and related effects of exposure to  $H_2SO_4$  mists *in vitro*. However the effects of pH-reduction have been investigated.

Low pH did not affect the frequency of point mutations in various bacteria strains, yeast and fungi but induced gene conversion in *Saccharomyces cerevisiae* (116).

Low pH led to sister chromatid exchanges and chromosomal aberrations in Chinese hamster cells treated *in vitro* in media over the pH range 5.4-7.2 during 24 hours continuous or 3 hours pulse treatments (172). Similar clastogenic effects of low pH have been reported in other studies of cultured mammalian cells, including human lymphocytes and epithelioid carcinoma (HeLa) cells (39, 169-171).

Sublethal pH decrease also caused developmental and mitotic abnormalities to embryos and sperm from eukaryotic sea urchins but failed to induce any changes in reversion rates in *Salmonella typhimurium* (53).

HCl, HNO<sub>3</sub> or H<sub>3</sub>PO<sub>4</sub> did not transform Syrian hamster embryo cells in a SA7/SHE-system *in vitro*. The concentrations ranged from 31 to 500 µg/ml for HCl and from 62 to 1 000 µg/ml for the two other acids (reviewed by Heidelberger *et al*, 1983 (107).

At relevant inhalation exposure levels, the local pH in the respiratory tract may be lowered by the acids although it can be assumed that systemic pH is not affected.

# 10.5 Effects of long-term exposure and carcinogenicity

Many of the published studies are based on combined exposure simulating environmental atmospheric pollution of  $H_2SO_4$  aerosol, ozone and  $NO_2$ . Here, mainly studies with isolated exposure to the acids are reported (Tables 11-12).

No animal inhalation studies according to modern standards on the carcinogenicity of strong inorganic acid aerosols were located. The few available studies have various weaknesses in study design such as inadequate exposure duration or an insufficient number of animals to address the carcinogenic potential of inorganic acid aerosols (Table 12). In a review by Swenberg and Beauchamp (1997), in which they assessed available animal studies on chronic toxicity and carcinogenicity from inorganic acid mists, it was concluded that the evidence from experimental animals neither strongly supports nor refutes the induction of cancer by inorganic acid mists (229).

## Sulphuric acid

The studies are compiled in Table 11. In a series of reports, Schlesinger and coworkers evaluated effects from  $H_2SO_4$  exposure on the lung defence in donkeys and rabbits. A sustained impairment of bronchial mucociliary clearance of radiolabelled Fe<sub>2</sub>O<sub>3</sub> (MMAD 5 µm) was observed in 2/4 donkeys exposed to approximately 0.1 mg/m<sup>3</sup> H<sub>2</sub>SO<sub>4</sub> (MMAD 0.5 µm), 1 hour/day, 5 days/week for 6 months (200). All 4 animals (1 female and 3 males) showed an increased variability in clearance rates.

In male rabbits exposed to  $0.125 \text{ mg/m}^3 \text{ H}_2\text{SO}_4$ , 2 hours/day, 5 days/week for up to 12 months, an accelerated tracheobronchial clearance of Fe<sub>2</sub>O<sub>3</sub> (MMAD 4.5  $\mu$ m) was observed after 4 and 8 months of exposure. A slowing trend was seen towards the end of the 12-month exposure. Clearance did not return to normal within the 6-month follow-up period but actually became slower. The animals developed increased secretory cell number in the small airways, an effect that was reversible (204). In contrast, male rabbits exposed to 0.25 mg/m<sup>3</sup> H<sub>2</sub>SO<sub>4</sub> (noseonly, MMD 0.3 µm), 1 hour/day, 5 days/week for 4, 8 or 12 months (the same daily dose as in the previous study) showed a retarded tracheobronchial clearance of radiolabelled Fe<sub>2</sub>O<sub>3</sub> (MMAD 4.5 µm) 18-20 hours after the preceding day's exposure. The lowering in clearance progressed with continued exposure. After cessation of exposure, clearance became extremely slow and did not return to normal during a 3-month follow-up period. Histological analysis revealed that acid-exposed animals had a narrowing of the bronchial airways and an increased lung epithelial secretory cell number. Significantly enhanced sensitivity to acetylcholine, a response characteristic of hyperresponsive airways, was seen in rabbits exposed for 4, 8, or 12 months. In addition, animals exposed for 8 or 12 months exhibited an increased airway reactivity, which had started to appear also in the 4-month exposure group (88-90). The authors concluded that both clearance and the morphometric endpoint were more dependent on inhaled acid concentration than on total dose.

Monkeys and guinea pigs were exposed to various combinations of SO<sub>2</sub>, H<sub>2</sub>SO<sub>4</sub> and fly ash. Guinea pigs (males and females) exposed to 0.08 (MMD 0.54 or 2.23  $\mu$ m) or 0.90 mg/m<sup>3</sup> (MMD 0.49  $\mu$ m) H<sub>2</sub>SO<sub>4</sub> for up to 52 weeks showed no signs of exposure-related toxicity. In contrast, monkeys (of both sexes) exposed to 0.88-0.99 mg/m<sup>3</sup> H<sub>2</sub>SO<sub>4</sub> for 78 weeks developed lesions in their lungs. These consisted of focal epithelial and goblet cell hypertrophy and hyperplasia, erosion, thinning and squamous metaplasia of the bronchiolar epithelium. H<sub>2</sub>SO<sub>4</sub> was considered responsible for the effects. No effects were demonstrable in 9 monkeys exposed to 0.09-0.11 mg/m<sup>3</sup> H<sub>2</sub>SO<sub>4</sub> (9).

Eight female dogs were exposed to  $0.9 \text{ mg/m}^3 \text{ H}_2\text{SO}_4$  for 21 hours/day for 620 days. Pulmonary function was seriously impaired and heart weights reduced, but no exposure-related histopathologic changes were demonstrated (154).

In guinea pigs (males and females) exposed to 0.08 or 0.1 mg/m<sup>3</sup> H<sub>2</sub>SO<sub>4</sub> (MMD 2.8 or 0.8  $\mu$ m) for 23 hours/day for 52 weeks, no exposure-related effects occurred. In monkeys of both sexes, chronic exposure to 2.4 and 4.8 mg/m<sup>3</sup> of H<sub>2</sub>SO<sub>4</sub> mists (MMD 3.6 or 0.73  $\mu$ m) for 23 hours/day for 78 weeks was associated with impaired lung function and histopathological changes. The lesions consisted of epithelial hypertrophy and hyperplasia, thickening of alveolar septa and bronchioles, and enlargement of air spaces. These effects were less pronounced or absent in the groups of monkeys exposed to lower concentrations (0.38 or 0.48 mg/m<sup>3</sup>). However, submicrometer particles (MMD 0.54  $\mu$ m) produced no alterations of

pulmonary structures at 0.48 mg/m<sup>3</sup> whereas particles with MMD 2.15  $\mu$ m at 0.38 mg/m<sup>3</sup> produced some histological changes (slight bronchiolar epithelial hyperplasia and slight thickening of walls of respiratory bronchioles) and an increased respiratory frequency. The results indicate that particle size was the important factor (8).

A lifetime study of the carcinogenic effect of  $H_2SO_4$  in rats and mice of both sexes is published, but the administration routes were not relevant to occupational exposure (intratracheal instillation and gastric intubation). The animals were exposed to maximum tolerated doses once a week for life. The concentrations were 0.3-0.5 ml 0.6 %  $H_2SO_4$  in distilled water administered both perorally and intratracheally to rats and 0.2 ml 0.2 %  $H_2SO_4$  for mice dosed perorally only. Tumours were seen in the second year of dosing in the organs where the acid acted directly.  $H_2SO_4$  was considered a weak, locally acting chemical carcinogen by the authors. The findings of tumours in the forestomach in mice were not significantly different from the controls (235).

There is only one study designed to evaluate pulmonary carcinogenicity in animals exposed chronically to aerosols. In male hamsters, no tumours were observed in the respiratory tract after inhalation exposure to  $100 \text{ mg/m}^3 \text{ H}_2\text{SO}_4$ , 6 hours/day, 5 days/week for life but laryngeal and tracheal epithelial hyperplasia was increased. No general toxic effects were noted (147) at this high level, which may indicate that the hamster is not a suitable animal model for assessing carcinogenicity.

In conclusion, the LOAEL after long-term exposure is  $0.1 \text{ mg/m}^3$  at which impaired mucociliary clearance in donkeys was reported. In rabbits, accelerated mucociliary clearance and secretory cells hyperplasia in small airways were observed at  $0.125 \text{ mg/m}^3$  and narrowing of the airways, airway hyperresponsiveness and decreased mucociliary clearance when the same daily dose was administered at a slightly higher level ( $0.25 \text{ mg/m}^3$ ). In monkeys exposed to larger particles (MMD 2.15 µm) at  $0.38 \text{ mg/m}^3$ , an increased respiratory rate and slight findings of hyperplasia of bronchiolar epithelium and thickening of the walls of respiratory bronchioles were reported. At higher levels, pulmonary function is impaired (LOAEL 0.9 mg/m<sup>3</sup> in dogs).

## Hydrochloric acid

In a 90-day inhalation study, male and female mice and rats were exposed to 0, 10, 20, or 50 ppm HCl (0, 15, 30, or 75 mg/m<sup>3</sup>), 6 hours/day, 5 days/week for 90 days. A slight but significant decrease in body weight gain was reported in male and female mice and in male rats in the highest dose group. No effects were observed in haematology, clinical chemistry and urinalysis parameters. Local irritative effects on the nose were observed in both rats and mice. In rats, concentration-and time-dependent minimal to slight rhinitis in the anterior nasal cavity were observed after 5 and 90 days. In mice, dose-dependently increasing lesions on skin and mucosa, particularly around the mouth and nose, were observed after 90 days at or above 30 mg/m<sup>3</sup>. In mice, histology revealed ulcerative dermatitis, necrotic

cells in the subcutis and increased phagocytised blood pigments in macrophages and intracytoplasmatically in sub-epithelial cells. The systemic NOAEL was found to be 30 mg/m<sup>3</sup>, and the LOAEL for irritation 15 mg/m<sup>3</sup> (industry report<sup>1</sup> cited in reference (68)).

No nasal cancer was observed after inhalation of 14 mg/m<sup>3</sup> HCl, 6 hours/day, 5 days/week for life in male Sprague Dawley rats. The exposure had no effect on body weight or mortality rate (10). In a study, with similar exposure regimen, no nasal cancer was observed, but the incidences of hyperplasia in larynx and trachea were increased (22 % and 26 % versus 2 % and 6 % among controls) (212). These studies were carried out as part of an investigation of the combined effect of formaldehyde and HCl. IARC pointed out (115) that as the studies were not designed to test the carcinogenicity of HCl, higher doses might have been tolerated. In neither study did HCl appreciably influence the tumourigenic effects of formaldehyde. No other experimental studies according to modern standards on the carcinogenicity of HCl were identified (Table 12).

## Nitric acid

In male rats, chronic exposure to low levels of HNO<sub>3</sub> vapour (0.05 mg/m<sup>3</sup>, noseonly for 4 hours/day, 3 days/week for 40 weeks) did not affect body weight or lung polyamine contents (essential for cell growth, multiplication, differentiation, and free radicals scavenging). Lung clearance of tracer microspheres was not statistically different from the control group. The level of stress inducible heatshock protein 70 was markedly elevated in lungs from the exposed animals (164, 215, 248). The relevance of this finding is uncertain. The results of the studies originate from the same animal experiment.

No experimental inhalation studies according to modern standards on the carcinogenicity of HNO<sub>3</sub> were located (Table 12).

### Phosphoric acid

No published studies on long-term exposure or carcinogenicity were identified.

<sup>&</sup>lt;sup>1</sup> Chemical Industry Institute of Toxicology (1984). 90-Day inhalation study of hydrogen chloride gas in B6C3F1 mice, Sprague-Dawley rats, and Fischer-344 rats. ToxiGenics, Inc for CIIT Research Triangle Park, NC, USA, CIIT Docket No. 20915.

	Exposure		Species	No. of	Effect	Reference	
Level, mg/m <sup>3</sup>	Droplet size, MD <sup>a</sup> , µm	Duration		animals			
0.08 or 0.1	2.8 and 0.8	23 h/d for 52 weeks	Guinea pig	50/group	No treatment-related histopathologic effects in sections of the lungs or trachea.	(8)	
0.09-0.11 and fly ash or $SO_2$	< 1 or 1-4	23 h/d for 78 weeks	Cynomolgus monkey	9/group	No changes in bronchial mucosa, pulmonary function or biochemical and haematological parameters.	(9)	
0.1	0.5	1 h/d, 5 d/week for 6 months	Donkey	4 totally	Increased variability in bronchial mucociliary clearance rates in all animals. A sustained impairment of mucociliary clearance in 2 animals (measured on tagged Fe <sub>2</sub> O <sub>3</sub> with MMAD 5 $\mu$ m).	(200)	
0.125	0.3	2 h/d, 5 d/week for up to 12 months	Rabbit	20 totally	6 months post-exposure, animals exposed for up to 8 months showed acceleration of tracheobronchial mucociliary clearance. Histological examination of animals sacrificed directly after 12 months of exposure revealed an increased secretory cell number in small airways and possibly mild focal epithelial hyperplasia.	(204)	
0.25	0.3	1 h/d, 5 d/week for 4, 8, or 12 months, nose-only	Rabbit	4-5/group	Development of airway hyperresponsiveness shown by: a) enhanced sensitivity to acetylcholine in all groups to a similar extent b) an increase in airway reactivity in animals exposed for 4 months (p < 0.075) and in animals exposed for 8 and 12 months (p < 0.05). Dynamic compliance was not affected in any group.	(88-90)	
					Decreased tracheobronchial mucociliary clearance of $Fe_2O_3$ particles (MMAD 4.5 µm) in all groups. Lowering of clearance progressed with continued exposure and were not normalised 6 months post-exposure. Decreased airway diameter, and secretory cell hyperplasia in the bronchial tree in all groups.		

	Exposure		Species	No. of	Effect	Reference
Level, mg/m <sup>3</sup>	Droplet size, MD <sup>a</sup> , µm	Duration		animals		
0.38	2.15	23 h/d for 78 weeks	Cynomolgus monkey	9/group	Slight findings of hyperplasia of bronchiolar epithelium and thickening of walls of respiratory bronchioles. Increased respiratory rate. No change in distribution of ventilation (nitrogen washout) or oxygen tension.	(8)
0.48	0.54	23 h/d for 78 weeks	Cynomolgus monkey	9/group	No exposure-related histological alterations in the lungs. No change in respiratory rate. Slight deterioration of the distribution of ventilation (nitrogen washout) at specific time intervals.	(8)
0.88-0.99 and fly ash or SO <sub>2</sub>	< 1 or 1-4	23 h/d for 78 weeks	Cynomolgus monkey	9/group	Changes in the bronchial mucosa (focal epithelial and goblet cell hypertrophy and hyperplasia, erosion, thinning and squamous metaplasia of epithelium). No changes in pulmonary function or biochemical and haematological parameters.	(9)
0.9	< 0.5	21 h/d for 620 d	Dog	8 totally	Impaired pulmonary function (reduced carbon monoxide diffusion capacity, residual volume, total lung volume, increased total expiratory resistance). No exposure-related histopathological changes. Reduced heart weight.	(154)
0.9	0.49	23 h/d for 52 weeks	Guinea pig	50/group	No exposure-related microscopic alterations in lungs. No effects on pulmonary function or biochemical parameters. Same results obtained also at exposure to 0.08 mg/m <sup>3</sup> (MMD 0.54 or 2.23 $\mu$ m) combined with fly ash.	(9)
2.4	3.6	23 h/d for 78 weeks	Cynomolgus monkey	9/group	Hyperplasia of bronchiolar epithelium and thickening of alveolar septa and bronchioles. Decreased pulmonary function (increased respiratory rate, decreased oxygen tension, deterioration of the distribution of ventilation). Same effects but more pronounced at 4.8 mg/m <sup>3</sup> (smaller particles, MMD 0.73).	(8)

# Table 11. Effects in animals of long-term exposure to sulphuric acid.

<sup>a</sup> Median diameter, either mass median diameter (MMD), mass median aerodynamic diameter (MMAD) or volume median diameter (VMD).

	Η	Exposure		Species	No. of	Effect/Comment	Reference	
Level	Route	Particle size	Duration		animals			
Sulphuric acid								
4-156 mg/m <sup>3</sup>	Inhalation	NG	6 h/d on alternate weekdays for 2 wk, 6 exposures	Rat	50/group	No significant increases in neoplasia. Animals were observed for lifetime. Inadequate reporting of study design, results, and short duration of exposure.	<sup>a, b</sup> cited in (229)	
100 mg/m <sup>3</sup>	Inhalation	2.6 µm	6 h/d, 5 d/wk, for life	Hamster	60/group	Laryngeal and tracheal epithelial hyperplasia, no tumours of the respiratory tract.	(147)	
0.2 ml 0.2 %, in distilled water	Oral (gastric intubation)		Once a week for life	Mouse	50-60/ group	Tumours in the forestomach, but not statistically significant from controls.	(235)	
0.3-0.5 ml 0.6 %, in distilled water (maximum tolerated doses)	Oral (gastric intubation) or intratracheal (instillation)		Once a week for life Twice a month for 12 months	Rat Rat	60/group 60/group	In the second year, various tumours in organs where $H_2SO_4$ acted directly (both benign and malign in the forestomach after intubation and in the trachea and lungs after instillation).	(235)	
Hydrochloric acid								
10 ppm vapour (14 mg/m <sup>3</sup> )	Inhalation		6 h/d, 5 d/wk for life	Rat	100/group	No nasal carcinomas or neoplasms of other sites.	(10)	
10 ppm vapour (14 mg/m <sup>3</sup> )	Inhalation		6 h/d, 5 d/wk for life	Rat	100/group	Increased incidence of hyperplasia in larynx and trachea (22 $\%$ and 26 $\%$ versus 2 $\%$ and 6 $\%$ among controls). No nasal cancer. No other observations reported.	(212)	

# Table 12. Carcinogenicity studies in animals exposed to the inorganic acids.

	E	Exposure		Species	No. of	Effect/Comment	Reference	
Level	Route	Particle size	Duration		animals			
5-22 mg/m <sup>3</sup> aerosols	Inhalation	NG	6 h/d on alternate weekdays for 2 wk, 6 exposures	Rat	50/group	No significant increases in neoplasia. Animals were observed for lifetime. Inadequate reporting of study design, results, and short duration of exposure.	<sup>a, b</sup> cited in (229)	
0.25 ml 0.5 %	Subcutaneous injections		6 times/wk for 10.5-16 months	Mouse	2 totally	Subcutaneous sarcomas at the site of injection.	<sup>c</sup> cited in (229)	
Nitric acid								
13-49 mg/m <sup>3</sup> aerosols	Inhalation	NG	6 h/d on alternate weekdays for 2 wk, 6 exposures	Rat	50/group	No significant increases in neoplasia. Animals were observed for lifetime. Inadequate reporting of study design, results, and short duration of exposure.	<sup>a, b</sup> cited in (229)	
Phosphoric acid								
No studies found.								

## Table 12. Carcinogenicity studies in animals exposed to the inorganic acids.

NG: Not given.

<sup>a</sup> Ballou JE, Gies RA, Dagle GE, Burton FG, Moss OR. Late effects of acid inhalation. In: *Pacific Northwest Laboratory Annual Report*. Richland, WA: Pacific Northwest Laboratory, 1978: 6.1-6.2.

<sup>b</sup> Ballou JE, Gies RA, Dagle GE, Burton FG, Moss OR. Late effects of acid inhalation. In: *Pacific Northwest Laboratory Annual Report*. Richland, WA: Pacific Northwest Laboratory, 1981: 223-225.

<sup>c</sup> Suntzeff V, Babcock RS, Loeb L. The development of sarcoma in mice following long continued injections of a buffered solution of hydrochloric acid. *Am J Cancer* 1940;39:56-60.

#### **10.6 Reproductive and developmental effects**

Pregnant CF-1 mice and New Zealand white rabbits were exposed to  $H_2SO_4$  aerosol (0, 5 or 20 mg/m<sup>3</sup>) in exposure chambers 7 hours/day, during organogenesis (day 6-15 in mice, 6-18 in rabbits). There were 35 mice and 20 rabbits in each dose group. No embryotoxic, foetotoxic or teratogenic effects from the exposures were observed. Slight maternal toxicity was seen in both species at 20 mg/m<sup>3</sup> (174).

No published studies on reproductive and developmental toxicity of HCl,  $HNO_3$ , or  $H_3PO_4$  were found.

Reproductive and developmental effects of the four acids are unlikely at relevant exposure levels since the contributions of protons and anions to the systemic levels will be low. However, effects secondary to lung damage cannot be excluded.

# 11. Observations in man

## 11.1 Irritation, corrosion and sensitisation

All four acids are classified as corrosive in the European legislation on classification and labelling of chemicals, with HNO<sub>3</sub> as the strongest and classified as corrosive in solutions containing more than 5 % (the most dilute solution classified).  $H_2SO_4$  is classified as corrosive in concentrations from 15 %, and HCl and  $H_3PO_4$  from 25 %; more dilute solutions are irritating to eyes, skin, and respiratory system. Thus,  $H_2SO_4$  is classified as irritating in the range 5-15 %, and HCl and  $H_3PO_4$  in the range 10-25 % (76).

Selected case-reports are described in Section 11.2. At lower air concentrations, the inorganic acid vapours or mists are respiratory tract and mucous membrane irritants (Section 11.3). Discolouration and erosion of the teeth have been reported after occupational exposure to the inorganic acids (Section 11.4) (110).

Repeated skin contact with the inorganic acids may lead to dermatitis (110). No sensitising effects have been reported after exposure to any of the four acids.

### Sulphuric acid

 $H_2SO_4$  is corrosive and highly irritating to skin. On direct contact,  $H_2SO_4$  causes violent dehydration, which can carbonise (char) the skin. In the reaction with water, heat is released in sufficient quantities to produce burns similar to thermal burns (110).  $H_2SO_4$  is rapidly injurious to the eyes and the mucous membranes of the respiratory tract and can etch teeth (6).

### Hydrochloric acid

Exposure to a high concentration of HCl gas or to a concentrated solution of HCl will cause burns of the skin and mucous membranes; repeated or prolonged

exposures of the skin to dilute solutions may cause dermatitis. Contact with the eyes may produce reduced vision or blindness (5, 110).

# Nitric acid

HNO<sub>3</sub> can cause severe burns in eyes and skin. Permanent corneal opacification leading to blindness has been reported. Dermal contact with concentrated solutions of HNO<sub>3</sub> may cause deep ulcers and yellowish staining of the skin. Dilute solutions of HNO<sub>3</sub> is irritating to the skin, eyes and mucous membranes. Acute occupational exposure to HNO<sub>3</sub> fumes (reported in cases with concurrent inhalation of NO<sub>2</sub> and NO) can elicit prompt irritation of the upper respiratory tract, leading to coughing, dyspnoea, cyanosis and acute pulmonary oedema. Inhalation of high (but unknown) concentrations of HNO<sub>3</sub> fumes has been responsible for numerous fatalities; death can be delayed several days (3).

## Phosphoric acid

 $H_3PO_4$  mist is a mild irritant of the eyes, upper respiratory tract and the skin; the dust is especially irritating to skin in the presence of moisture (4).

# 11.2 Case reports

Human case reports (exemplified below) describe severe reactions to the four acids when in contact with the skin, eyes and mucous membranes. These include corrosion and destruction of body tissue from chemical burns leading to ulcers, blindness and death. In a prospective study of 16 patients with acid ingestion including H<sub>2</sub>SO<sub>4</sub>, HCl or HNO<sub>3</sub>, all had oesophageal and gastric injuries. Two died of gastric perforation and one of bronchopneumonia. In surviving patients with severe injuries, late complications developed, e.g. oesophageal stricture (69). Life-threatening acute (or adult) respiratory distress syndrome (ARDS) as well as persistent reactive airways dysfunction syndrome (RADS) have been reported after single exposure to high acid levels.

# Sulphuric acid

The death of a man 5 days after he was splashed with concentrated  $H_2SO_4$  was attributed to extensive burns and chemical damage to the respiratory tract (208). In a survey of burns caused by chemicals including  $H_2SO_4$ , 3 deaths were identified in patients whose injuries exceeded 50 % of the total body surface (36). A case of accidental inhalation of fumes of strong  $H_2SO_4$  during application to blocked drainpipes with fatal outcome is reported. Autopsy revealed congestion of the respiratory passages and severe pulmonary oedema (26). A 40-year old man accidentally exposed to liquid fuming  $H_2SO_4$  and for 8 minutes to acid mist and fumes from the action of water (safety shower) on the liquid developed a disabling pulmonary fibrosis, residual bronchiectasis, and pulmonary emphysema that persisted 18 months (93). A 45-minute exposure to a cleaning compound containing 66 %  $H_2SO_4$  in an unventilated washroom caused cough, chest tightness, dyspnoea and mild rhinoconjunctivitis in a 45-year-old woman immediately after exposure

(33). A plumber developed ARDS, after accidental exposure in a manhole to unknown levels of a 95 %  $H_2SO_4$  mixture from a pipe (132).

## Hydrochloric acid

A case developing tracheobronchial stenosis from HCl ingestion and aspiration presenting as asthma has been reported (194). A man with a history of mild asthma that inhaled high concentrations of a product containing HCl for 1 hour developed rapidly progressive and severe bronchospasm. One year later, marked asthma symptoms remained (33). A 30-year-old woman presented with RADS after a single accidental inhalation of HCl (78). Among 9 pharmaceutical workers exposed to HCl fumes, 5 had severe symptoms, reduced peak expiratory flow rate or hypoxaemia. One patient developed long-term airway hyperreactivity, super-imposed on a background of chronic obstructive airways disease (35).

Following an accident with a container truck leaking HCl, burning and tearing eyes, burning throats, headache, chest pain, shortness of breath and flu-like symptoms were observed among the exposed residents. A follow-up on 45 exposed adults and 56 age-matched controls was performed 20 months later. The exposed differed significantly from controls in neurobehavioural tests for balance, reaction time, digit-symbol and perceptual motor speed. Respiratory, neurobehavioural, general and vegetative function symptoms were also more frequent among exposed than among controls. Furthermore, balance scores, reaction times and forced expiratory flow at 25-75 % (FEF  $_{25-75}$ ) of forced vital capacity (FVC) were poorer in subjects living closest to the accident site (126).

# Nitric acid

Headache together with nausea and/or vomiting, phono- and photophobia and lachrymation induced 5 minutes after inhalation of HNO<sub>3</sub> vapour from a metal polish, has been observed in a goldsmith. The headache was resistant to non-steroidal anti-inflammatory drug therapy but responded to triptans. The patient suffered from migraine with a monthly frequency when not exposed to HNO<sub>3</sub> but every day when exposed to HNO<sub>3</sub> vapour. The effects were assigned to the formation of nitrous gases and not to HNO<sub>3</sub> itself. No further details were given, e.g. on the level and duration of the exposure (94).

Symptoms and findings after accidental inhalation of HNO<sub>3</sub> fumes (and nitrous gases), including cases of deaths, were described in a paper by Hall and Cooper (1905). Few details were given on exposure levels. Symptoms listed were dyspnoea, cough, pain in the chest, stomach, lungs, throat, loins and head, dizziness, and nausea and vomiting. Pulmonary congestion (cough with bloody or voluminous expectoration) typically arose with a delay of 6-7 hours after the exposure to HNO<sub>3</sub> and developed into bronchitis (102).

Three cases of fatal pulmonary oedema caused by 10-15 minutes of accidental exposure to  $HNO_3$  fumes from an exploding tank containing 68 %  $HNO_3$  at a pulp mill have been reported (101).

A man who was accidentally exposed to  $NO_2$  fume from a reaction between  $HNO_3$  and the metal of a bucket for about 30 minutes, died 14 days after the exposure from extreme cyanosis with extensive pneumonia (63).

#### Phosphoric acid

A man exposed to high but unknown levels of  $H_3PO_4$  by accident for  $3 \times 20$  minutes on a tank ship during a storm developed RADS (34).

A 64-year-old man ingested  $H_3PO_4$  in a suicide attempt and developed hyperphosphataemia, hypocalcaemia and systemic metabolic acidosis. Local caustic effects were mild (43).

The death of one individual 19 days after ingestion of an unknown amount of  $H_3PO_4$  was reported by Hawkins *et al*. Death was a result of recurrent internal abdominal haemorrhage (105).

## 11.3 Effects of single and short-term exposure

#### Sulphuric acid

In several volunteer studies, subjects were exposed to clean air and different concentrations of  $H_2SO_4$ , and mixtures of  $H_2SO_4$  with ozone,  $NO_x$  or particulate matter, usually one week apart. As the effect of each chemical compound in a mixture is challenging to isolate and interpret, only the results of exposure to  $H_2SO_4$  alone is reported (Table 13). The studies vary with regard to exposure system, test concentrations, particle size, relative humidity, exposure duration, physical activity and choice of test persons; they include healthy as well as asthmatic volunteers, and include exposures in chambers, via mouthpieces or nasal masks. The latter has been used by one team to minimise *in situ* neutralisation of the acid by ammonia. Others have e.g. let the subjects gargle citrus juice before exposure or rinse with mouthwash to suppress airway ammonia. The droplet sizes vary considerably among the studies, the MMADs being in the range of 0.1-10  $\mu$ m. The pulmonary function has been measured before (baseline) and after exposure, each subject being its own control. Sex differences have, in general, not been addressed in the studies.

The few studies on occupationally exposed workers are presented below and are listed in Table 15.

## Sulphuric acid: Pulmonary function and symptoms

A self-administered questionnaire on respiratory and eye symptoms and their severity was completed by 75 controls and 82 workers exposed to  $H_2SO_4$  in 13 workplaces representing several industries. The exposed were divided in a low-exposure group (below 0.15 mg/m<sup>3</sup>) and a high-exposure group (0.15-0.50 mg/m<sup>3</sup>). The respondents had been exposed for at least 6 months. Highly significant relationships were found between symptom reporting and exposure level. Those with low exposure (< 0.15 mg/m<sup>3</sup>), typically reported 1 symptom, whereas those with exposure of about 0.5 mg/m<sup>3</sup> reported an average of 5 symptoms. The most

commonly reported symptoms were sneezing, irritated nose and cough. Runny nose increased most markedly with exposure. Younger workers (below 40 years) consistently reported more symptoms than the older workers. The results indicate that workers suffer symptoms of respiratory and eye irritation at exposures in the range 0.1-0.5 mg/m<sup>3</sup> (presented as conference proceedings) (81).

In a cross-sectional study, 225 workers (98 % white males) in 5 plants manufacturing lead acid storage batteries were administered a questionnaire on workrelated symptoms, underwent spirometry, and had personal samples for H<sub>2</sub>SO<sub>4</sub> taken over the shift. Most personal samples were below 1 mg/m<sup>3</sup>. MMADs were 2.6-10  $\mu$ m estimated from area samples. Workers with the higher exposure to acid (greater than 0.3 mg/m<sup>3</sup>, n=41) did not have an increased rate of acute workrelated symptoms of e.g. skin, eyes or respiratory airways as compared to the lowexposure group (less than 0.07 mg/m<sup>3</sup>, n=116). Changes in pulmonary function over the shift were not related to levels of airborne acid (86).

In healthy volunteers, one study demonstrated no significant decreases in specific airway conductance, forced expiratory volume in one second (FEV<sub>1</sub>), or maximum flow rates at 40 or 60 % of total lung capacity, but enhanced airway reactivity to inhaled carbachol after exposure to 1 mg/m<sup>3</sup> (237). Otherwise, single exposures of healthy volunteers at 0.1-0.47 and 1-2 mg/m<sup>3</sup> have not produced changes in pulmonary function (details below and in Table 13) (14, 20-22, 83, 149, 151, 152, 156, 195, 219, 220, 232).

In 14 adolescent asthmatic volunteers (9 males, 5 females, 13-18 years), however, a small but significant (6 %) fall in FEV<sub>1</sub> was seen after exposure for 45 minutes to concentrations as low as 0.035 mg/m<sup>3</sup>. A tendency to decreased FEV<sub>1</sub> (p = 0.08) was seen also after exposure for 45 minutes to 0.07 mg/m<sup>3</sup>. After a 90-minute exposure at both levels no such effect was seen. The changes after acid exposure were compared to the changes after air exposure (136).

Another study conducted in the same laboratory demonstrated small but significant changes in FEV<sub>1</sub> and FVC in 22 adolescent asthmatics (15 males, 7 females, 12-19 years) after exposure to 0.05-0.18 mg/m<sup>3</sup> (MMAD 0.72) for 40 or 45 minutes during intermittent moderate exercise. The changes were no longer significant 20 minutes after exposure (103). The study has weaknesses in reporting and design.

Small but significant reductions in FEV<sub>1</sub>, total respiratory resistance and maximal flow calculated at 50 % of FVC (FEF<sub>50</sub>) were seen in 10 adolescent asthmatics (4 males, 6 females, 12-17 years) exposed to 0.1 mg/m<sup>3</sup> (MMAD 0.6  $\mu$ m) for 40 minutes (30 minutes at rest followed by 10 minutes of moderate exercise) compared with changes obtained after a saline aerosol exposure. The effects appeared after exercise and were reversible. Symptom scores (nasal, respiratory and non-respiratory effects) were not affected by the acid exposure (134). In the three studies on adolescent asthmatics, the relative humidity was 65-75 % (103, 134, 136).

Pulmonary function and symptom scores were not affected in healthy or asthmatic adult men exposed for 2 hours with intermittent exercise to 0.1 mg/m<sup>3</sup> H<sub>2</sub>SO<sub>4</sub> aerosol (MMAD 0.5  $\mu$ m) at a relative humidity of 40 % (20), nor in healthy

and asthmatic volunteers of both sexes exposed at similar conditions for 3 hours (MMAD 0.64  $\mu$ m) (83).

Exposure of 45 adult volunteers (15 non-atopic or atopic, 30 asthmatic) to 0.1 mg/m<sup>3</sup> (MMAD 0.5  $\mu$ m) H<sub>2</sub>SO<sub>4</sub> caused no significant changes in lung function, symptom scores, or bronchial reactivity relative to clean air (156).

Utell *et al* reported no effects in neither healthy nor adult asthmatics (sex not given) after 16 minutes exposure at rest to 0.1 mg/m<sup>3</sup> (MMAD 0.6-1  $\mu$ m), but enhanced bronchoconstriction as well as increased reactivity to carbachol in asthmatics after similar exposures to 0.45 and 1 mg/m<sup>3</sup>. An increased reactivity to carbachol was demonstrated in healthy after exposure to 1 mg/m<sup>3</sup>. A saline aerosol served as control. Relative humidities were 25 % or less (236, 237).

Fifteen asthmatics (sex not given) inhaled 0.35 mg/m<sup>3</sup> (MMAD 0.8  $\mu$ m) of an H<sub>2</sub>SO<sub>4</sub> aerosol for 20 minutes at rest followed by 10 minutes of exercise at high and low oral ammonia levels. A saline aerosol at a low ammonia level served as control exposure. The relative humidity was 20-25 %. A significant reduction in FEV<sub>1</sub> following exercise was observed at acid inhalation combined with low oral ammonia levels as compared to the other exposures. Also maximum expiratory flow at 60 % of total lung capacity was reduced (238).

No increase in symptom scores, bronchial reactivity to methacholine or effects on lung function were observed in 15 healthy and 15 asthmatics volunteers of both sexes exposed to  $0.1 \text{ mg/m}^3$  (MMAD  $1.0 \mu \text{m}$ ) for 1 hour including exercise at a relative humidity of 50 % (14).

Framton *et al* exposed 12 healthy volunteers (10 men and 2 women) for 2 hours to 0 or 1 mg/m<sup>3</sup> H<sub>2</sub>SO<sub>4</sub> aerosols (MMAD 0.9  $\mu$ m, relative humidity 40 %) with intermittent exercise and performed bronchoalveolar lavage 18 hours post-exposure. Acid exposure did not result in alveolar inflammation, influx of plasma proteins into the alveolar space, or alterations in selected antiviral functions of alveolar macrophages (82).

No differences in symptom scores (ordinate scale), ventilation, lung function (FEV<sub>1</sub>, FEF<sub>50</sub>, FVC), antioxidant levels in nasal lavage or exhaled nitric oxide were observed in 12 healthy and 12 asthmatic volunteers (sex not given) exposed 1 hour to 0, 0.2 and 2 mg/m<sup>3</sup> aerosol (MMD 0.3  $\mu$ m) (232).

Sackner *et al* (1978) reported no effects on ventilation, lung function or cardiopulmonary function in 5 healthy and 5 asthmatic volunteers of both sexes exposed up to 1 mg/m<sup>3</sup> for 10 minutes (MMAD 0.1  $\mu$ m, relative humidity 20-30 %) (195).

After exposure of 22 adults of both sexes to larger  $H_2SO_4$ -particles (VMD 10  $\mu$ m) at 0, 0.5, 1 and 2 mg/m<sup>3</sup> for 1 hour including exercise at 10 °C and 100 % relative humidity, a significant dose-dependent increase in upper and lower respiratory symptom scores was observed over the whole exposure range among healthy volunteers and at 1 and 2 mg/m<sup>3</sup> in asthmatics. The healthy subjects showed only a small symptom increase at the lowest exposure level. A trend towards small decrements in pulmonary function (seen as a decrease in peak expiratory flow rate) was reported among asthmatics at the higher levels. The exposures lasted 1 hour and included 3×10 minutes of heavy exercise (22).

In a similarly designed study with  $0.9 \ \mu m H_2SO_4$  particles, healthy and asthmatic volunteers of both sexes were exposed to 0, 0.38, 1 and 1.5 mg/m<sup>3</sup>. At the two highest concentrations, asthmatics showed significant increases in lower respiratory symptoms as well as non-respiratory symptoms (headache, fatigue, eye irritation) and decrements in pulmonary function (FVC and FEV<sub>1</sub>) but no changes in airway reactivity. There were no significant effects among the healthy participants apart from increased cough with increasing acid concentration, which was scored "minimal" at the highest concentration (21). Taken together, these two studies suggest that symptoms were less pronounced with the 0.9- $\mu$ m aerosol although the physiologic response of asthmatics seemed greater. According to the authors, the rationale for this could be that symptoms are caused primarily by larger acid droplets deposited in the upper airway or proximal bronchi, whereas disturbances of pulmonary mechanics in asthmatics are caused primarily by smaller acid droplets deposited in more distal airways (99).

In 10 asthmatics exposed to 0, 0.1, 0.3 and 1 mg/m<sup>3</sup> (MMAD 0.5  $\mu$ m) for 1 hour, decrements in respiratory function was produced (specific airway conductance, FEV<sub>1</sub>/FVC, FEF<sub>25</sub>, and FEF<sub>50</sub>). The study also demonstrated effects on respiratory clearance and is described in more detail below (219).

In a series of exposures assessing the influence of particle size, osmolarity, relative humidity, liquid water content and physical exercise, 7 male and 4 female asthmatics were exposed to  $H_2SO_4$  aerosols at 3 mg/m<sup>3</sup> via mouthpiece for 16 minutes at rest and exercise. Ten of the subjects were also exposed to 1 and 1.4 mg/m<sup>3</sup> for 1 hour at intermittent exercise (100 W) in a chamber. No effects on specific airway resistance, lower respiratory or non-respiratory symptom ratings were observed. Throat irritation was significantly greater after exposure to acid aerosol with < 10 % relative humidity than after exposure to acid aerosol of 100 % relative humidity or to saline at a low relative humidity (15).

## Sulphuric acid: Respiratory clearance

A number of studies have examined the effect of  $H_2SO_4$  inhalation on the clearance of insoluble particles from the respiratory tract. In some studies, in which exposure lasted for 1 or 2 hours, bronchial mucociliary clearance was affected at exposure levels from 0.1 mg/m<sup>3</sup> H<sub>2</sub>SO<sub>4</sub> (MMAD 0.5 µm) (151, 152, 220). In the study by Leikauf *et al* in 10 healthy volunteers whereof 3 females, aerosol exposure via nasal mask to 0.1 mg/m<sup>3</sup> (MMAD 0.5 µm) resulted in an accelerated bronchial mucociliary clearance of Fe<sub>2</sub>O<sub>3</sub> (MMAD 7.5) compared to sham exposure, whereas exposures to 0.3 mg/m<sup>3</sup> gave rise to a variable bronchial clearance, and 1 mg/m<sup>3</sup> caused a transient slowing of clearance. Tracheal mucociliary transport rates following the acid exposures did not differ from those obtained after sham exposures. The 7.5-µm particles used for assessment of clearance were calculated to be primarily deposited in the larger bronchial airways, where submicrometer H<sub>2</sub>SO<sub>4</sub> has little deposition (151). Another study was therefore designed to determine the effect of submicrometer H<sub>2</sub>SO<sub>4</sub> on clearance from the distal ciliated airways (152) (both studies also reported by

Lippmann et al 1981) (158). Clearance of 4.2-µm Fe<sub>2</sub>O<sub>3</sub>-particles inhaled before the aerosol exposure was measured in 8 healthy volunteers (4 of each sex) exposed via nasal mask to 0, 0.1, 0.3, or 1 mg/m<sup>3</sup> (MMAD 0.5  $\mu$ m) H<sub>2</sub>SO<sub>4</sub>. In contrast to results obtained with larger radiolabelled particles, bronchial mucociliary clearance of the smaller particles was slower after all H<sub>2</sub>SO<sub>4</sub> inhalations than after sham exposure. Respiratory mechanics and tracheal mucociliary transport rates were not affected by any aerosol exposure. The authors' explanation to the differences between results obtained with larger and smaller tagged particles, respectively, is the deposition patterns of the Fe<sub>2</sub>O<sub>3</sub> and acid aerosols. Submicrometer  $H_2SO_4$  is primarily deposited in the distal small airways in a pattern more like that of the  $4.2 - \mu m$  than the  $7.5 - \mu m$  particles. In the larger bronchial airways, the 0.1 mg/m<sup>3</sup> acid dose acted as a small stimulatory dose to mucociliary transport, whereas in the distal ciliated airways, the acid dose was sufficient to depress mucociliary transport (152). The ability of an irritant to stimulate mucociliary clearance at a low dose while slowing it at higher doses has also been shown by others; reviewed by Wanner (241).

In a parallel study, 10 asthmatics (6 males and 4 females) were exposed according to essentially the same protocol (inhalation of 3.9-µm tagged Fe<sub>2</sub>O<sub>3</sub>-particles preceding H<sub>2</sub>SO<sub>4</sub>-exposure at 0, 0.1, 0.3 and 1 mg/m<sup>3</sup> for 1 hour). For the 6 asymptomatic mild asthmatics, bronchial mucociliary clearance was delayed in a dose-dependent manner and significantly so at the highest exposure level. In addition, exposure to 1 mg/m<sup>3</sup> produced decrements in respiratory function. The remaining 4 asthmatics on daily medication exhibited variable and inconsistent clearance patterns. Mean tracheal mucociliary transport rates were not altered for either group following any of the acid exposures, although their variability was increased compared with sham exposure. Sham exposure clearance rates were slower among both the medicated and non-medicated asthmatics in this study (219) than among the healthy volunteers in the study by Leikauf *et al* (152).

In another study, tracheobronchial mucociliary clearance of 5.2-µm Fe<sub>2</sub>O<sub>3</sub>-particles was reduced following exposure of 10 healthy male volunteers to 0.1 mg/m<sup>3</sup> of H<sub>2</sub>SO<sub>4</sub> aerosol (MMAD 0.5) for 1 or 2 hours compared with exposure to a distilled water aerosol. Respiratory mechanics were not affected (220).

Exposure to larger droplets of  $H_2SO_4$  at 0.47 mg/m<sup>3</sup> (MMAD 10.3 µm) for 40 minutes at rest followed by 20 minutes during exercise accelerated *tracheal* clearance and small airway mucociliary clearance of a radioaerosol (MMAD 3.4 µm) in 7 healthy male subjects compared with saline fog. No effects were observed on symptom ratings (headache, irritation of eyes, upper or lower airways) ventilatory function or airway reactivity (149).

Exposure of 10 healthy volunteers (9 males, 1 female) to 1 mg/m<sup>3</sup> (MMAD 0.5  $\mu$ m) for 2.5 hours increased bronchial mucociliary clearance of a radiolabelled aerosol (MMD 3  $\mu$ m) compared with exposure to a distilled water mist. Tracheal clearance was not studied (177).

In a comparative study in rabbits and humans (10 males and 2 females) of a single exposure by inhalation, similarities in effects on pulmonary immuno-

competence were observed in response to  $H_2SO_4$  exposure (1 mg/m<sup>3</sup>  $H_2SO_4$  for 3 hours, MMAD 0.9 µm) (Section 10.2). The ability of recovered human pulmonary macrophages to attach to a solid substrate *in vitro* and the capacity to produce superoxide anion was reduced. Cell viability and phagocytosis of latex particles were not significantly affected (252).

# Sulphuric acid: Conclusion

A reduced bronchial mucociliary clearance has been reported in healthy volunteers after exposure to 0.1 mg/m<sup>3</sup> for 1 or 2 hours. An accelerated tracheal clearance was observed in healthy volunteers exposed for 1 hour to larger droplets at 0.47 mg/m<sup>3</sup>. A 1-hour exposure to larger droplets at 0.5 mg/m<sup>3</sup> was associated with a small increase in upper and lower respiratory symptom ratings. One occupational study indicates that symptoms of respiratory irritation occur at exposures in the range 0.1-0.5 mg/m<sup>3</sup> (presented as conference proceedings). Pulmonary function changes (bronchial hyperreactivity) were demonstrated in healthy volunteers exposed 16 minutes to 1 mg/m<sup>3</sup>.

In adolescent asthmatics, one research team reported small pulmonary function changes at  $0.1 \text{ mg/m}^3$  or even lower. In adult asthmatics, the LOAEL for pulmonary function effects is 0.35 (low oral ammonia levels) or 0.45 mg/m<sup>3</sup>. However, there are other studies showing no effects at or even above 1 mg/m<sup>3</sup>.

## Hydrochloric acid

Human inhalation exposure studies with HCl are few. In a relatively recent study, 5 male and 5 female adult asthmatics were exposed via half-face mask to air containing 0, 1.12, and 2.52 mg/m<sup>3</sup> HCl for 45 minutes. The 45-minute exposure was divided into three equal periods: exercise, rest, exercise. Pulmonary function was measured and self-reported symptoms registered. The test subjects did not show any adverse respiratory health effects and did not report any symptoms (upper and lower respiratory, and non-respiratory symptoms) from the HCl exposure (228). The authors also listed earlier human studies and case reports of HCl exposure. It was estimated that the odour threshold was in the range 1.5-7.5 mg/m<sup>3</sup>, intolerable at 75-150 mg/m<sup>3</sup>, and that 1 950-3 000 mg/m<sup>3</sup> is a lethal concentration (228). Other sources assign odour thresholds between 1 and 50 mg/m<sup>3</sup> (0.7-35 ppm) (Table 2).

In a report compiling HCl air levels and subjective irritant effects among workers (number not given) in steel pickling facilities, it was concluded that no irritation of the mucous membranes was observed at 3-4.5 mg/m<sup>3</sup>, initial mild irritation of the airway mucosa, which regressed rapidly, occurred at 5.2 mg/m<sup>3</sup>, slight irritation at 7-11 mg/m<sup>3</sup>, and breathing difficulties at 26-34 mg/m<sup>3</sup> (165). The study is based on many years of observations but does not comply with current standards (Table 15).

		Exposure		No. of volunteers <sup>a</sup>	Effect	Reference
Level, mg/m <sup>3</sup>	Droplet size, MMAD, µm	Duration (at rest if not otherwise stated)	System			
0.035	0.6	45 or 90 min during intermittent moderate exercise	Mouthpiece	14 asthmatic adolescents	<ul> <li>45 min exposure: 6 % fall in FEV<sub>1</sub> immediately after exposure compared to the before level. The changes were no longer significant 20 minutes after the exposure.</li> <li>90 min exposure: No significant change in FEV<sub>1</sub>. No change in FVC or total respiratory resistance after any exposure.</li> </ul>	(136)
0.068	0.6	40 min (30 min rest and 10 min intermittent moderate exercise)	Mouthpiece prestudy	9 asthmatic adolescents	3 % fall in $\text{FEV}_1$ after exposure compared to the before level (non-significant).	(135)
0.07	0.6	45 or 90 min during inter- mittent moderate exercise	Mouthpiece	14 asthmatic adolescents	<i>45 min exposure:</i> Decreasing trend in FEV <sub>1</sub> (-3 %), but not statistically significant. No other lung function changes. <i>90 min exposure:</i> No significant change in FEV <sub>1</sub> , FVC or total respiratory resistance.	(136)
0.07+0.13	0.72	40 or 45 min during inter- mittent moderate exercise	Mouthpiece	22 asthmatics adolescents	Small changes in FEV <sub>1</sub> and FVC. The changes were no longer significant 20 minutes after exposure.	(103)
0.1	0.5	2 h during intermittent exercise	Chamber	6 healthy, 6 asthmatics	No effects on symptoms (upper and lower respiratory symptoms and non-respiratory symptoms) or lung function.	s (20)
0.1	0.6-1	16 min during exercise	Mouthpiece	17 asthmatics	No effects on SGaw or FEV <sub>1</sub> .	(236)
0.1	0.6-1	16 min	Chamber	14 healthy, 17 asthmatics	No effects on SGaw or FEV <sub>1</sub> .	(237)
0.1	1.0	1 h with alternate 10 min periods of heavy exercise and rest	Chamber	15 healthy, 15 asthmatics	No effect on the incidence of symptoms of irritancy, bronchial reactivity to methacholine, or lung function.	(14)

		Exposure		– No. of volunteers <sup>a</sup>	Effect	Reference
Level, mg/m <sup>3</sup>	Droplet size, MMAD, µm	Duration (at rest if not otherwise stated)	System			
0.1	0.64	3 h	Chamber	30 healthy, 30 asthmatics	No effects on symptoms or pulmonary function in healthy or asthmatic subjects.	(83)
0.1	0.5	6.5 h with 50 min exercise periods and 10 min rest	Chamber	<ol> <li>15 healthy,</li> <li>30 asthmatics</li> </ol>	No changes in symptoms of irritancy, bronchial reactivity, or lung function.	(156)
0.1	0.5	1 h	Nasal mask	6 asthmatics	No significant effect on bronchial mucociliary clearance of 3.9- $\mu$ m Fe <sub>2</sub> O <sub>3</sub> -particles (clearance reduced in a dose-dependent manner, non-significantly at 0.1 and 0.3 mg/m <sup>3</sup> and significantly at 1.0 mg/m <sup>3</sup> ). No effect on mean tracheal mucociliary transpor- rates or lung function.	y
0.1	0.6	30 min rest and 10 min moderate exercise	Mouthpiece	10 asthmatic adolescents	Small but significant reductions in $FEV_1$ , total respiratory resistance and $FEF_{50}$ , all reversible. No changes in symptom scores (nasal, respiratory and non-respiratory effects).	(134)
0.1	0.5	1 h	Nasal mask	8 healthy	Reduced bronchial mucociliary clearance of $Fe_2O_3$ -particles (MMAD 4.2 µm). Average half-time increased from 80 to 110 min. Same effect seen at 0.3 and 1 mg/m <sup>3</sup> . No effect on mucociliary tracheal transport, FEV <sub>1</sub> , FVC, FEF <sub>50</sub> .	(152)
0.1	0.5	1 h and 2 h on separate occasions	Nasal mask	10 healthy	Reduced bronchial mucociliary clearance of $Fe_2O_3$ -particles (MMAD 5.2 µm), persistent 2 h after exposure. Average clearance half-time increased from 42 to 84 and 110 min respectively (for volunteers exposed 1 and 2 h). No effects on FEV <sub>1</sub> , FVC, FEF <sub>50</sub> , PEPR, or airway resistance.	(220)
0.1	0.5	1 h	Nasal mask	10 healthy	Markedly increased bronchial clearance of $7.5 - \mu m Fe_2O_3$ - particles in 6/10 exposed volunteers. No change in tracheal mucociliary transport rate or in pulmonary mechanics.	(151)

	Exposure				Effect	Reference	
Level, mg/m <sup>3</sup>	Droplet size, MMAD, µm	Duration (at rest if not otherwise stated)	System	volunteers <sup>a</sup>			
0.2	0.3	1 h	Head-only, cast acrylic head dome	12 healthy, 12 asthmatics	No changes in symptom scores, ventilation, lung function (FEV <sub>1</sub> , FEF <sub>50</sub> , FVC), antioxidant levels in nasal lavage or exhaled nitric oxide.	(232)	
0.3	0.5	1 h	Nasal mask	10 healthy	Variable bronchial clearance of $7.5-\mu m Fe_2O_3$ -particles (both faster and slower than the controls). No change in tracheal mucociliary transport rate or in pulmonary mechanics.	(151)	
0.3	0.5	1 h	Nasal mask	8 healthy	Reduced bronchial mucociliary clearance of $4.2 - \mu m Fe_2O_3$ - particles. Tracheal transport was not significantly affected. Same results after exposure to 1 mg/m <sup>3</sup> .	(152)	
0.3	0.5	1 h	Nasal mask	6 asthmatics	No significant effect on bronchial mucociliary clearance $(3.9-\mu m Fe_2O_3)$ . Clearance was reduced (dose-dependently), non-significantly at 0.1 and 0 and significantly at 1.0 mg/m <sup>3</sup> ). No effect on mean tracheal mucociliary transport rates or lung function.	(219)	
0.35	0.8	20 min at rest and 10 min of exercise	Mouthpiece	15 asthmatics	Following exercise, low oral $NH_3$ caused significantly greater re- ductions in $FEV_1$ and maximum expiratory flow at 60 % of total lung capacity compared to saline or acid + high oral $NH_3$ levels.	(238)	
0.38	0.9	1 h including three 10 min periods of heavy exercise	Chamber	21 healthy, 21 asthmatics	No change in pulmonary function, airway reactivity or reporting of upper, lower respiratory or non-respiratory symptoms.	(21)	
0.45	0.6-1	16 min	Chamber	17 asthmatics	Bronchial hyperreactivity after provocation with carbachol.	(237)	
0.45	0.6-1	16 min during exercise	Mouthpiece	17 asthmatics	Bronchoconstriction (reduced SG <sub>aw</sub> ).	(236)	
0.47	10.3	40 min at rest and 20 min of exercise	Chamber	7 healthy	Accelerated tracheal and small airways mucociliary clearance of a sulphur colloid (MMAD 3.4 $\mu$ m). No effects on symptom ratings (headache, irritation of eyes, upper or lower airways) ventilatory function or airway reactivity.	(149)	

 Table 13. Volunteer studies with exposure to sulphuric acid aerosol.

	Exposure			No. of	Effect R	Reference
Level, mg/m <sup>3</sup>	Droplet size, MMAD, µm	Duration (at rest if not otherwise stated)	System	volunteers <sup>a</sup>		
0.5	10 (VMD)	1 h including three 10 min periods of heavy exercise	Chamber	<ul><li>22 healthy,</li><li>22 asthmatics</li></ul>	Small increase in upper and lower respiratory symptoms among healthy. Scores were successively increased at 1 and 2 mg/m <sup>3</sup> . No increase beyond that from sham exposure among asthmatics.	(22)
1	10 (VMD)	1 h including three 10 min periods of heavy exercise	Chamber	<ul><li>22 healthy,</li><li>22 asthmatics</li></ul>	Dose-dependent increase in upper and lower respiratory symptoms in healthy and asthmatics. Also seen at 2 mg/m <sup>3</sup> . No pulmonary function effects apart from a trend to dose-dependent decrease in PEFR in asthmatics at this and higher level (2 mg/m <sup>3</sup> ).	(22)
1	0.9	1 h including three 10 min periods of heavy exercise	Chamber	<ul><li>21 healthy,</li><li>21 asthmatics</li></ul>	In healthy, no change in pulmonary function, airway reactivity or reporting of upper or lower respiratory or non-respiratory symptoms.	(21)
					In asthmatics, increases in lower respiratory and non-respiratory symptoms and decrements in pulmonary function (FVC and $FEV_1$ ) but no changes in airway reactivity.	
1	0.9	3 hours, with intermittent exercise	Chamber	12 healthy	Reduced ability of recovered pulmonary macrophages to attach to a solid substrate <i>in vitro</i> and production of superoxide anion by stimulated pulmonary macrophages. Viability or phagocytosis of latex particles was not significantly affected.	(252)
1	0.1	10 min	Mouthpiece	11 healthy, 11 asthmatics	No effects on ventilation, lung function, or cardiopulmonary function. Same results obtained at 0.01 and 0.1 $mg/m^3$ .	(195)
1	0.6-1	16 min during exercise	Mouthpiece	17 asthmatics	Bronchoconstriction (reduced $SG_{aw}$ and $FEV_1$ ).	(236)
1	0.6-1	16 min	Chamber	14 healthy, 17 asthmatics	In healthy, no effect on $SG_{aw}$ , $FEV_1$ or maximum flow rates at 40 or 60 % of total lung capacity. Bronchial hyperreactivity after provocation with carbachol in asthmatics and healthy subjects.	(237)

	Exposure			No. of	Effect	Reference
Level, mg/m <sup>3</sup>	Droplet size, MMAD, µm	Duration (at rest if not otherwise stated)	System	volunteers <sup>a</sup>		
1	0.5	2.5 h, intermittent exercise during the 1st hour	Chamber	10 healthy	Increased bronchial mucociliary clearance of tagged albumen saline aerosol (MMAD 3.5 µm).	(177)
1	0.5	1 h	Nasal mask	Healthy 8 (small tracers) 10 (large tracers)	Reduced bronchial mucociliary clearance of both small (MMAD 4 $\mu$ m) and large (MMAD 7.5 $\mu$ m) Fe <sub>2</sub> O <sub>3</sub> -particles. No effect on tracheal transport.	(151, 152)
1	0.5	1 h	Nasal mask	6 asthmatics	Transient slowing of tracheobronchial mucociliary clearance of $Fe_2O_3$ -particles (MMAD 3.9 µm) and decrements in respiratory function (SGaw, FEV <sub>1</sub> , FEV <sub>1</sub> /FVC, FEF <sub>25</sub> , FEF <sub>50</sub> ). No effect on mean tracheal mucociliary transport rates.	(219)
1	0.5	1 h	Nasal mask	10 healthy	Decreased bronchial clearance of $Fe_2O_3$ -particles (MMAD 7.5 $\mu$ m). No effect on tracheal transport.	(151)
1	0.9	2 h with intermittent exercise	Chamber	12 healthy	No effects in total protein, cell recovery, cell differential counts or influx of inflammatory cells into the alveolar space, nor in alveolar macrophage function (superoxide anion release or ability to inactivate influenza virus).	(82)
1	0.9	2 h	Chamber	12 healthy	No change in airway mucin glycoproteins.	(61)
1.5	0.9	1 h including three 10 min periods of heavy exercise	Chamber	21 healthy, 21 asthmatics	In healthy, no change in pulmonary function and airway reac- tivity, and no upper or lower respiratory and non-respiratory symptoms. In asthmatics, increases in lower respiratory and non-respiratory symptoms and decrements in pulmonary function (FVC and $FEV_1$ ) but no changes in airway reactivity.	(21)
2.0	0.3	1 h	Head-only, cast acrylic head dome	12 healthy, 12 asthmatics	No differences in symptom scores, ventilation, lung function $(FEV_1, FEF_{50}, FVC)$ , antioxidant levels in nasal lavage or exhaled nitric oxide.	(232)

<b>I</b> and <b>a</b>				No. of	Effect	Reference
Level, mg/m <sup>3</sup>	Droplet size, MMAD, µm	Duration (at rest if not otherwise stated)	System	volunteers <sup>a</sup>		
3.0	0.4 and 6 (VMD)	16 min rest and exercise, respectively, with mouthpiece, later 1 h intermittent moderate exercise in chamber	Mouthpiece, chamber	18 asthmatics	No effect on specific airway resistance, cough, upper and lower respiratory, or non-respiratory symptom scores, apart from in- creased throat irritation at low relative humidity.	. ,

<sup>a</sup> Adult unless otherwise stated.

FEF<sub>x</sub>: forced expiratory flow at x % of FVC, FEV<sub>1</sub>: forced expiratory volume in one second, FVC: forced vital capacity, MMAD: mass median aerodynamic diameter, PEFR: peak expiratory flow rate, SG<sub>aw</sub>: specific airway conductance, VMD: volume median diameter.

### Nitric acid

Ten healthy volunteers (6 males, 4 females) exercised for 2 hours in an atmosphere containing 0.4 mg/m<sup>3</sup> HNO<sub>3</sub> fog (VMD 6  $\mu$ m, relative humidity 100 %). No significant differences in pulmonary function (FEV<sub>1</sub>, FVC, specific airway resistance, respiratory rate, tidal volume) or symptom scores (lower and upper respiratory symptoms or non-respiratory symptoms) were reported as compared to exposures to water fog or clean-filtered air (16).

Ten healthy volunteers (8 males, 2 females) exposed to 0 and 0.5 mg/m<sup>3</sup> of HNO<sub>3</sub> vapour for 4 hours during moderate exercise did not report any exposure-related symptoms. Further, no proximal airway or distal lung injury could be detected with lung function tests (FEV<sub>1</sub>, FVC or specific airway resistance), proximal airway lavage, bronchoalveolar lavage or bronchial biopsies (17).

Exposure of 1 female and 8 male healthy subjects to 0 and 0.2 mg/m<sup>3</sup> of HNO<sub>3</sub> vapour for 2 hours, of which 100 minutes during moderate intermittent exercise, did not affect pulmonary function (spirometry and airway resistance) and subjective symptoms. Nor were there any significant increases in indicators of airway injury or inflammation assessed by bronchoalveolar lavage 18 hours post-exposure. However, there was an effect from HNO<sub>3</sub> on alveolar macrophage function (significant increases in the phagocytic activity and the resistance to infection with respiratory syncytial virus, and decreased superoxide anion production) (25).

Adolescent asthmatics (19 males, 9 females, 12-19 years old), inhaled a combination of 0.05 ppm ( $0.125 \text{ mg/m}^3$ ) HNO<sub>3</sub>, 0.12 ppm O<sub>3</sub> and 0.30 ppm NO<sub>2</sub> for 90 minutes via a mouthpiece. Pulmonary function parameters were not affected when compared with changes after clean air exposure. The study protocol also included exposure to oxidants alone and oxidants combined with H<sub>2</sub>SO<sub>4</sub>. Six subjects left the study before completion because of unpleasant symptoms (137).

In an old study published 1907, the author and a colleague were exposed to concentrations of 11-12 ppm (27-30 mg/m<sup>3</sup>). Symptoms described included sneezing, pressure in the chest, pains in the trachea and larynx, coughing, secretion from the nose and salivary glands, moderate burning in the eyes, and lachrymation, burning and itching of the facial skin. The author considered exposure at those levels for more than 1 hour intolerable and dangerous to human health. Exposure to 84 ppm (210 mg/m<sup>3</sup>) was tolerated by the author for only 2-3 minutes (Diem<sup>1</sup>, cited in (66)).

Other reports on observations in man are from accidental exposure to HNO<sub>3</sub> (Section 11.1).

# Phosphoric acid

No studies were found. A few case reports are included in Section 11.1.

<sup>&</sup>lt;sup>1</sup> Diem L. Untersuchungen über die Einatmung von Saltpetersäure-dämpfen (thesis). Würzburg, 1907.

# 11.4 Effects of long-term exposure

Only a few studies have demonstrated effects of long-term exposure to the acids in humans, most of them concerning dental effects (Table 15). Exposure to inorganic acids in the working environment can lead to dental etching and erosion. Systems for grading dental etching and erosion are exemplified by the one presented by Ten Bruggen Cate in 1968 (230) (Table 14). In a recent review of occupational risk factors for dental erosion (including the studies presented below), it was concluded that battery, galvanising and associated workers exposed to  $H_2SO_4$  or HCl and to a lesser degree to  $H_3PO_4$ , HNO<sub>3</sub> and hydrofluoric acid were at higher risk of dental erosion based on prevalence studies including a control group. However, prevalence data in both acid-exposed workers and controls exhibited a great variation, amounting to 26-100 % for battery and galvanising workers and to 0-80 % for controls (244).

The prevalence of dental erosion varies greatly with age, time period, and dietary factors. Relatively little is known about the prevalence of erosion in the general population. In a Swiss study, a prevalence of grade 2 erosion was found to be 62 % in the age group 46-50 years (161). In a review, prevalences were reported to range from 4 % to 82 % in adults aged 18-88 years (121).

# Sulphuric acid

There was no association between exposure to  $H_2SO_4$  (average 0.15 mg/m<sup>3</sup>) and respiratory symptoms (cough, phlegm, dyspnoea and wheezing) among 248 workers (243 males, 5 females) in 5 battery plants in a cross-sectional study. Differences were analysed between workers having a cumulative exposure above  $15 \text{ mg/m}^3 \times \text{months}$  (average exposure level 0.21 mg/m<sup>3</sup>) and workers with cumulative exposure below 7 mg/m<sup>3</sup> × months (average exposure level 0.10)  $mg/m^3$ ). FVC was impaired in the highly exposed workers compared to the less exposed workers. Tooth etching and erosion was strongly associated with  $H_2SO_4$ exposure and was observed in 15-33 % of the workers in 4 of the 5 plants investigated (and in no workers in the remaining plant). Dental etching was observed in 38 % of the highly exposed workers, and in 8 % in the low exposure group workers (p < 0.0005 adjusted for age and smoking). The earliest cases of etching and erosion, respectively, occurred after 4 and 30 months of exposure to an estimated average exposure of 0.23  $mg/m^3$  (see Table 15 for details) (87). The exposure concentrations were estimated for the various workplaces after personal sampling on presumably one occasion, and can therefore only be taken as

**Table 14.** Grades of etching and erosion according to Ten Bruggen Cate (230).

Grade	Description
Etching	Dull, ground-glass appearance of the enamel surface without loss of contour.
Grade 1 erosion (G1)	Loss of enamel only.
Grade 2 erosion (G2)	Loss of enamel with involvement of dentine.
Grade 3 erosion (G3)	Loss of enamel and dentine with exposure of secondary dentine.
Grade 4 erosion (G4)	Loss of enamel and dentine resulting in pulpal exposure.

approximate. The authors also reviewed previous studies<sup>1</sup> of  $H_2SO_4$ -exposed workers. Most of these studies reported on the prevalence of respiratory diseases such as bronchitis but showed no effect on the respiratory system clearly attributable to low levels of  $H_2SO_4$  in battery plants and  $H_2SO_4$  plants. However, the potential effect on pulmonary function was not examined in these studies. Tooth etching and erosion was common.

The effects of  $H_2SO_4$  exposure on the teeth have been documented in other studies. Usually, only current and no historical exposure levels were given. In a cross-sectional study, Petersen and Gormsen (1991) reported dental erosion and attrition in relation to exposure to  $H_2SO_4$  (0.4-4.1 mg/m<sup>3</sup> at the time of the investigation) in a modern battery factory in 61 male dentate workers, in the processes known as forming and charging. Dental erosion and attrition was seen in 42 % of the workers exposed for less than 10 years and in 56 % of the workers exposed for more than 10 years (186).

In a Finnish study on acid exposed workers from battery and galvanising factories exposed predominantly to  $H_2SO_4$ , blind dental examinations were used. Referents were from acid-free departments of the same companies. Of the acid exposed workers, 18 % had one or more teeth with erosion compared to 9 % of the referents (p=0.075). The number of teeth with erosion (all three grades according to Eccles' classification, which is similar to that of Ten Bruggen Cate) was significantly higher among the acid exposed workers than among the referents. Concentrations of  $H_2SO_4$  fumes varied from 0.06 to 2.0 mg/m<sup>3</sup> (233). In another study, the prevalence of tooth surface loss was 63 % among acid exposed workers in Tanzania, compared to 38 % among controls. Reported  $H_2SO_4$  levels ranged from below 1 to above 5 mg/m<sup>3</sup>. The occurrence of tooth surface loss was significantly higher among workers than among controls already after 1-5 years of employment (234).

Among workers at an electro-winning facility in South Africa, the odds ratio (OR) for dental erosion was 5.5 for workers exposed to  $0.3-1 \text{ mg/m}^3$  compared to those exposed to  $0.1-0.3 \text{ mg/m}^3$  (referents). There was also a significant difference in the severity of tooth surface loss between the two groups. The authors expressed some uncertainty regarding the reliability of the stated concentrations as represent-tative of the exposure (51).

In battery workers (males), prevalences of erosion and etching were 87 % (exposure 3-17 mg/m<sup>3</sup>), and 47 % (exposure < 0.8-2.5 mg/m<sup>3</sup>). The average exposure of those showing etching was 5 years (163).

<sup>&</sup>lt;sup>1</sup> Anfield BD, Warner CG. A study of industrial mists containing sulphuric acid. *Ann Occup Hyg* 1968;11:185-194.

El-Sadik YM, Osman HA, El-Gazzar RM. Exposure to sulphuric acid manufacture of storage batteries. *J Occup Med* 1972;14:224-226.

Morando A. Experimental and clinical contribution to human pathology due to sulphuric acid fumes. *Med Lav* 1956;47:55-61.

Pelnar T. The influence of work in sulfuric acid production in employee health. *Prac Lek* 1951;3:287-294.

Williams MK. Sickness absence and ventilatory capacity of workers exposed to sulphuric mist. *Br J Ind Med* 1970;27:61-66.

El-Sadik *et al* (1972) reported the prevalence of tooth erosion to be 39 % in plants with very high exposure (>  $12.6 \text{ mg/m}^3$ ) (71). Ten Bruggen Cate (1968) also found a high prevalence and incidence of tooth etching and erosion among workers in battery plants, but no air concentrations were reported (230).

Nasal symptoms and alterations of the nasal mucosa were studied in 52 workers exposed to H<sub>2</sub>SO<sub>4</sub> mists in 5 anodising plants. The workers included were not exposed to other vapours or metals. Work-related nasal symptoms (e.g. itching, bleeding and discharge) were reported by 21 workers (40 %). The major clinical findings in the exposed were hyperaemia (30 %), pale mucosal patches (29 %) and ulcerations (12 %). Stationary measurements during a 5-day working week revealed average geometric mean air levels between 0.035 and 2.1 mg/m<sup>3</sup> in the 5 plants. Pale mucosal patches or ulcerations were only observed in workers (15/37) from the three plants with the highest exposure levels (above  $0.2 \text{ mg/m}^3$ ) and in none of 15 workers exposed at lower levels. Histopathological evaluation (nasal biopsy) showed that squamous metaplasia, squamous atypia and mild dysplasia were more frequent among the exposed (20 selected workers) than among unexposed (n=11). According to the authors, the risk of squamous atypia or dysplasia increased with increasing exposure to H<sub>2</sub>SO<sub>4</sub> but did not correlate with exposure duration (mean 6 years, range: 4 months to 16 years) (95). The study has weaknesses in design and reporting, e.g. regarding control groups, data handling and statistics.

## Hydrochloric acid

In a report compiling HCl air levels and subjective irritant effects among workers in steel pickling facilities, chronic bronchitis was reported after years of exposure at approximately  $30 \text{ mg/m}^3$ . It was also stated (no further details given) that no damage to the teeth occurred at average concentrations of 4.5-7.7 mg/m<sup>3</sup> (165). The study is based on many years of observations but does not comply with current standards.

In a hot dip zinc galvanising plant, 90 % of 38 workers had some dental erosion (grade 1 or 2) of the incisor teeth. Exposure levels (geometric means) ranged from 1.8 to 12.4 mg/m<sup>3</sup> HCl at different sites. The dental effect could not be causally linked to the HCl exposure from the pickling process (where a 15 % HCl solution was used to remove corrosion) as the number of workers was small and there was no control group (191).

A high prevalence of dental erosion was observed among industrial workers exposed to HCl but no air concentrations were reported (230).

### Nitric acid

Dental erosion from occupational exposure to  $HNO_3$  fumes was suggested by some authors but no corresponding levels of exposure were presented (3, 230).

*Phosphoric acid* No studies were found.

### Mix of acids

Dental erosion, especially among battery formation workers (exposed to  $H_2SO_4$ ), and black staining of the teeth among iron picklers (who were exposed to various acids including H<sub>2</sub>SO<sub>4</sub>, HCl, HNO<sub>3</sub>) were reported to be due to acid exposure in the working environment. Etching or erosion (grade 1-3) was observed in more than 30 % of the workers. No grade 4 cases were observed. HCl and H<sub>2</sub>SO<sub>4</sub> accounted for far more erosion cases than HNO<sub>3</sub> or chromic acids, but it is not possible to conclude whether this was due to differences in erosive potential of the acids or in exposure levels. Dental etching increased to a peak in workers exposed for 2-5 years and then diminished, while dental erosion grade 1 cases started to appear in workers exposed from 4-6 months and increased steadily. Dental erosion grade 2 cases first appeared after 2-5 years and increased while the dental erosion grade 3 cases were seen after 6-10 years. The exposure levels were not given, and other effects were not described (230). Exposure to  $H_2SO_4$  at the time when the study was published (pre 1970s) has been characterised as high for workers in metal pickling (>  $1 \text{ mg/m}^3$  8-hour time weighted average) and low for workers in lead battery production ( $< 0.1 \text{ mg/m}^3$ ) (197).

Long-term (up to 5 years) exposure to low levels of SO<sub>2</sub>, HCl and SO<sub>4</sub><sup>2-</sup> (7-hour TWA maximum 0.30, 2.1 and 0.5 mg/m<sup>3</sup>, respectively) was not associated with any increase in the airway hyperresponsiveness to histamine among male workers in synthetic fibre plants. The source of SO<sub>4</sub><sup>2-</sup> was H<sub>2</sub>SO<sub>4</sub> (141). In previous reports on the same study population, the authors found an association between exposure to SO<sub>2</sub>, HCl and SO<sub>4</sub><sup>2-</sup> and a higher prevalence of work-related cough and nasal symptoms compared with the reference group (139, 140).

#### Conclusion

Only a few studies have demonstrated effects of long-term exposure to the acids in humans (none on H<sub>3</sub>PO<sub>4</sub>). Dental etching or erosion is a concern after exposure to acids and has been reported after exposure to approximately 0.2 mg/m<sup>3</sup> of H<sub>2</sub>SO<sub>4</sub>. Histopathological changes of the nasal mucosa were reported at about the same levels of H<sub>2</sub>SO<sub>4</sub> exposure. Respiratory symptoms and effects on pulmonary function have been investigated or reported to a small degree. In one study, FVC was impaired in workers with cumulative H<sub>2</sub>SO<sub>4</sub> exposure above 15 mg/m<sup>3</sup> × months (average 0.21 mg/m<sup>3</sup>) compared to less exposed workers (below 7 mg/m<sup>3</sup> × months (average 0.10 mg/m<sup>3</sup>). Chronic bronchitis is reported after exposure to approximately 30 mg/m<sup>3</sup> of HCl.

#### **11.5 Genotoxic effects**

The number of chromosomal aberrations, micronuclei and sister chromatid exchanges were increased in lymphocytes from workers of a phosphate fertiliser factory. The workers had been exposed to phosphoric acid, but also to other chemicals and radioactivity (91). No studies on the other acids were found.

In vitro studies of human cells are presented in Section 10.4.

Exposure level, mg/m <sup>3</sup>	Droplet size, MD, µm	Exposure duration	Industry	Study size	Study design/con- founders adjusted for	Effect	Reference
Sulphuric acid							
Effects of short-term	exposure						
< 0.15 (low) 0.15-0.50 (high)	NG	$\geq$ 6 months	13 workplaces representing several industries	82 workers 75 controls	Cross-sectional/ smoking	Highly significant trends between individual symptom reporting and exposure level. The most commonly reported symptoms were sneezing, irritated nose and cough. Runny nose increased most markedly with exposure.	
< 0.07 mg/m <sup>3</sup> (low) > 0.3 mg/m <sup>3</sup> (high)	3-10	10 years (average for all plants as given by Gamble <i>et al</i> (87).	5 battery plants	225 mostly male workers <sup>a</sup> 116 (low) 41 (high)	Cross-sectional/ age, smoking	Workers with the higher exposure (n=41) did not have an increased rate of acute work-related symptoms as compared to the low-exposure group (n=116). Changes in pulmonary function over the shift were not related to levels of airborne acid.	(86)
Effects of long-term	exposure						
0.06-2.0	NG	1-39 years	2 battery and 2 galvanising factories	76 male workers, 81 controls	Cross-sectional/ age, smoking, food and drink intake	Number of teeth with erosion significantly higher among acid exposed than among controls.	(233)
0.15 (average) 0.10 (average, low <sup>b</sup> ) 0.21 (average, high <sup>c</sup> )		10 years (average for all plants)	5 battery plants	243 male/5 female workers no referents 99 (low) 100 (high)	Cross-sectional/ age, smoking	No increase in respiratory symptoms. FVC reduced in the high-exposure group (4.83 litres) compared to the low-exposure group (5.11 litres). Tooth etching and erosion associated with exposure.	(87)

**Table 15.** Effects of occupational exposure to the inorganic acids.

Exposure level, mg/m <sup>3</sup>	Droplet size, MD, µm	Exposure duration	Industry	Study size	Study design/con- founders adjusted for	Effect	Reference
0.035-2.1	NG	6 years	5 anodising plants	52 workers (histopathology on 20) 11 controls	Cross-sectional/ age, smoking	Nasal symptoms reported by 21 workers. Pale mucosal patches and ulcerations only in workers exposed to > $0.2 \text{ mg/m}^3$ . Squamous metaplasia, squamous atypia and mild dysplasia of the nasal mucosa more frequent among a subset of exposed than among unexposed. The risk of squamous atypia or dysplasia increased with increasing concentration of H <sub>2</sub> SO <sub>4</sub> but did not correlate with exposure duration.	(95) 1
0.1-0.3 (referents) 0.3-1	NG	1 month-24 years, average 4.2 years	Electro-winning facility	103 workers 102 referents	Cross-sectional/ age, length of service	Significantly increased risk for dental erosion (OR 5.5). Significantly increased severity of tooth surface loss. No relation- ship between exposure time and erosion.	(51)
0.4-4.1 (range) (average NG)	NG	< 10 years > 10 years	Battery factory	61 workers no referents	Cross-sectional/NG	Dental erosion and attrition in 42 % of the workers exposed for < 10 years and in 56 % of the workers exposed for > 10 years.	(186)
< 0.8-2.5 (charging) 3-17 (forming)	NG	Years	Battery plant	60 workers 117 controls	Cross-sectional/ socio- economical status, age	Significantly higher incidence of erosion in men in forming than in charging or controls.	(163)
< 1 - > 5	NG	1-19 years	Fertiliser factory	68 workers 61 controls	Cross-sectional/ health, dietary habits	Significantly increased prevalence of tooth surface loss (63 % vs. 38 %). Prevalence increased among workers already after 1-5 years of exposure.	(234)
12-35	NG	NG	Battery factories	<ul><li>33 workers</li><li>20 controls</li></ul>	Cross-sectional/age	Prevalence of erosion 39 % versus 0 % among controls.	(71)

**Table 15.** Effects of occupational exposure to the inorganic acids.

Exposure level, mg/m <sup>3</sup>	Droplet size, MD, µm	Exposure duration	Industry	Study size	Study design/con- founders adjusted for	Effect	Reference
Hydrochloric acid							
1.8-12.4 (average)	NG	NG	Hot dip zinc galvanising plant	38 workers no controls	Cross-sectional	90 % had dental erosion (grade 1 or 2) of the incisor teeth.	(191)
3-4.5 5.2 4.5-7.7 7-11 26-34 ~30	Vapour	Years	Steel pickling	NG	Cross-sectional	No irritation of the mucous membranes. Initial mild irritation. No dental damage. Slight irritation. Breathing difficulties. Chronic bronchitis.	(165)
Mix of irritants							
2.1 HCl 0.5 H <sub>2</sub> SO <sub>4</sub> 0.30 SO <sub>2</sub>	NG	< 5 years	Synthetic fibre plants	119 workers 180 referents	Cross-sectional/ smoking, age, allergy	Higher prevalence of work-related cough and nasal symptoms. No increase in chronic respiratory symptoms or airway hyper- responsiveness.	(139- 141)
H <sub>2</sub> SO <sub>4</sub> , HCl, HNO <sub>3</sub> and others. Levels not given.	NG	< 10 years	Battery factory and iron pickling facility	555 workers 293 referents	Prospective/age	Dental etching and erosion (grade 1-3), especially from exposure to HCl and $H_2SO_4$ .	(230)

Table 15. Effects of occupational exposure to the inorganic acids.

<sup>a</sup> Basically the same population as in Gamble *et al* (87).
<sup>b</sup> Cumulative exposure < 7 mg/m<sup>3</sup>×months.
<sup>c</sup> Cumulative exposure > 15 mg/m<sup>3</sup>×months.
FVC: forced vital capacity, MD: median diameter, NG: not given, OR: odds ratio.

# **11.6 Carcinogenic effects**

An overview of epidemiological studies is given in Table 16.

### Sulphuric acid

The exposure is poorly characterised in most studies, but for the older studies it is assumed that the pre 1970s exposure levels presented for various types of industries in Chapter 6 can be applied.

Occupational exposure to strong inorganic acid mists containing H<sub>2</sub>SO<sub>4</sub> was classified as carcinogenic to humans (Group 1) by the International Agency for Research on Cancer (IARC) in 1992 (116). Several studies were evaluated and the classification was primarily based on the human studies by Ahlborg et al (7), Beaumont et al (24), Siemiatycki (214), Steenland and Beaumont (226), Steenland et al (225), Soskolne et al (217, 218) and Weil et al (243). In all studies, H<sub>2</sub>SO<sub>4</sub> mists were regarded to be the commonest exposure. Additional support was provided by Forastiere et al (80) and Hagmar et al (100). In several of these studies, exposure to acid mists was associated with laryngeal cancer. In the years following the publication of the IARC monograph, a follow-up of the Steenland et al study (225) has been published, showing a positive association between exposure to acid mist and laryngeal cancer (227). In this study, the incidence of laryngeal cancer was investigated in a cohort of male steel workers exposed primarily to H<sub>2</sub>SO<sub>4</sub> mists 1940-65. Exposure data from 1975-79 from the two plants indicated that personal exposure levels of H<sub>2</sub>SO<sub>4</sub> had averaged 0.19 mg/m<sup>3</sup> and area samples  $0.29 \text{ mg/m}^3$ , and that the exposure duration was on average 9.2 years. The average first year of exposure was 1949. The rate ratio for laryngeal cancer among the 1 031 workers was 2.2 (95 % confidence interval (CI) 1.2-3.7) as compared to national rates (the expected number of laryngeal cancers in the general population) after adjustment for tobacco and alcohol consumption (227).

In a cohort of 1 409 workers employed in  $H_2SO_4$  production, with or without previous exposure in mines, no excess mortality in laryngeal (4 cases versus 3.1 expected) or lung cancer (27/32.8) was observed. An increased mortality from myeloid leukaemia could not be attributed to any of the exposures (184).

In a recent nested case-control study in a cohort of nickel refinery workers exposed to low levels of  $H_2SO_4$  (generally less than 0.5 mg/m<sup>3</sup>) with 213 cases and 525 age-matched controls, no excess risk of lung cancer from  $H_2SO_4$  mist was found. The resulting OR (CI) for the different exposure groups (mg/m<sup>3</sup> × years) were: low: 1.0 (0.4-2.3), medium: 1.0 (0.4-2.4), high 0.8 (0.3-2.0) (97).

In a recent case-control study, no association was found between exposure to environmental pollution of  $H_2SO_4$  (historic levels exceeding 0.5 mg/m<sup>3</sup>) and lung cancer in Lithuanian men. After control for smoking and age, the OR for lung cancer was 1.03 (95 % CI 0.76-1.39) (187) (NEG noted that historic levels, if correct, were extremely high).

To test the hypothesis of a role of occupational risk factors in the aethiology of gastric cancer, Cocco *et al* conducted a case-control study based on death certificates concerning stomach cancer in the United States. The data base

included 41 957 deaths from gastric cancer. No excess risk was associated with  $H_2SO_4$  exposure at the workplace (OR 0.99, 95 % CI 0.95-1.03) after adjustment for other exposures, ethnic origin, marital and socio-economic status, and residence (56). The same team evaluated the risk of gastric cardia cancer by occupation and industry in a case-control study (n=1 056) based on death certificates. Exposure intensity was classified in three categories without quantitative dose measures. Among white males, a significantly increased risk with increasing exposure was observed (55).

One case of laryngeal cancer and three cases of naso-pharyngeal carcinoma were reported.  $H_2SO_4$  was suggested to be the causative agent (111, 112).

Histopathological effects of the nasal mucosa were reported in a study by Grasel *et al* (95) detailed in Section 11.3.

There is one population-based case-control study suggesting an excess risk of oesophageal cancer with exposure to  $H_2SO_4$  (214) (also published by Parent *et al* (183)). The exposure was coded by chemists and hygienists based on the confidence that the exposure had occurred, the frequency of exposure during a normal working week and the relative concentration of  $H_2SO_4$  and other agents. Exposure was classified as none, non-substantial or substantial.

In conclusion, one follow-up study published after the IARC monograph supports the conclusion about laryngeal cancer whereas the evidence for an association with lung cancer is weak. These conclusions are in accordance with other recent evaluations (27, 106, 197).

### Hydrochloric acid

There is inadequate evidence for the carcinogenicity in humans of HCl according to IARC (115). No association was found between exposure to HCl and risk of lung cancer (129 exposed cases) in a nested case-control study among chemical factory workers. The exposure was up to  $3 \text{ mg/m}^3$  for several years. A latency period of 15 years was applied in the analysis (31, 32). An excess of lung cancer was observed in a study of steel-pickling workers in a subset of 189 workers who had been exposed to mists of acids other than H<sub>2</sub>SO<sub>4</sub>, primarily HCl (24).

In the population-based case-control study by Siemiatycki (described above), an increased risk of non-Hodgkin's lymphoma (n=18), rectum cancer (n=18) and lung oat-cell carcinoma (n=19) was suggested in workers exposed to HCl (p < 0.10). No risk was observed for other histological types of lung cancer (214).

### Nitric acid

No association was found between exposure to nitrogen products (HNO<sub>3</sub> and urea) and mortality from bladder cancer in exposed chemical plant workers (166). The study is however, inconclusive with regard to risk for bladder cancer and exposure to HNO<sub>3</sub> due to a small number of cases (n=4) and the relatively good prognosis of bladder cancer making mortality an inappropriate outcome to study.

In a study by Hilt *et al*, the possibility of co-exposure to asbestos limits the interpretability with respect to HNO<sub>3</sub> exposure and lung cancer (109).

In the population-based case-control study described above, an excess risk for pancreas (n=5), prostate (n=9) and kidney (n=4) cancer was suggested in workers exposed to HNO<sub>3</sub> (p < 0.10) (214).

# Phosphoric acid

The population-based case-control study by Siemiatycki indicated an excess risk of kidney (n=6) and lung (n=14) cancer among  $H_3PO_4$  exposed (p <0.10) (214).

### Mixed acids

In a cohort study among former and present battery and steel work employees exposed to acid mists, the standard mortality ratio for upper aerodigestive tract tumours was 0.92 (95 % CI 0.85-0.98). A case-control study in the same cohort showed that the risk was moderately but non-significantly increased among workers exposed for at least five years to H<sub>2</sub>SO<sub>4</sub> or HCl (H<sub>2</sub>SO<sub>4</sub> was substituted with HCl ) in excess of 1 mg/m<sup>3</sup> (OR 2.0, 95 % CI 0.4-10). This included cases of lip cancer. The authors distinguished between three levels of exposure to acids: zero, low (< 1 mg/m<sup>3</sup> H<sub>2</sub>SO<sub>4</sub> or HCl) and high ( $\geq$  1 mg/m<sup>3</sup>), taking into account the time periods during which jobs had been held (57). Hathaway later commented on the data analysis, e.g. that exclusion of the lip cancer cases would presumably result in an OR near unity, and that the study produced no evidence that acid aerosols may cause upper aerodigestive cancer (104)

In cohort mortality studies from the phosphate industry, the workers were exposed to  $H_3PO_4$ , as well as to  $H_2SO_4$ . The standard mortality ratios for lung cancer (SMR 1.22 and 1.24) was slightly increased among white as well as non-white workers in a study (44), which was later analysed anew and described as non-causal in relation to occupational exposure (45).

In another study, a pronounced increase of lung cancer among African American workers was reported (SMR 4.11). The levels of  $H_2SO_4$  and  $H_3PO_4$  were 0.013-0.22 mg/m<sup>3</sup> and 0.03-0.52 mg/m<sup>3</sup>, respectively. Exposure to other substances known as carcinogens during vessel cleaning had occurred (224). There were no adjustments for smoking in the above-mentioned studies.

A multicentre case-control study was conducted to evaluate the role of occupational exposures in risk of laryngeal/hypopharyngeal cancer. No overall excess risk was found linked to inorganic acid mists, but a significantly higher risk for hypopharyngeal cancer (4 exposed cases) among those exposed for at least 15 years. No association, however, was observed with cumulative exposure. Exposure was mainly to HCl but included also HNO<sub>3</sub>, H<sub>3</sub>PO<sub>4</sub>, H<sub>2</sub>SO<sub>4</sub>, hydrofluoric, and chromic acid (213).

### 11.7 Reproductive and developmental effects

No studies were located regarding reproductive or developmental effects in humans for any of the four acids but effects at relevant exposure levels are unlikely (see Section 10.7).

Acid exposure level (co-exposure)	Design	Industry/product, Country	Study size	Adjustment in addition to age <sup>a</sup>	Outcome and risk estimate (95 % CI), no. of cases (cohort) no. of exposed cases (case-control)	Reference
Sulphuric acid						
NG	Cohort, morbidity	Isopropanol manufacture, USA	182 (71 exposed > 5 years)	No	Sinonasal cancer: 4 cases represented an apparent, large excess compared with the national rate.	(243)
NG	Cohort, mortality	Steel workers - sheet and tin mills (incl. pickling), USA	2 763	No	Lung cancer: SMR 1.10 (0.73-1.60), 27.	(167)
NG (diethyl sulphate)	Cohort, mortality, morbidity	Alcohol manu- facturing plant, USA	335	No	Laryngeal cancer: SIR 5.04 (1.36-12.90), 4. (Assumed to be due to diethyl sulphate).	(162)
NG	Cohort, mortality	Isopropyl alcohol plant, United Kingdom	262	No	Non-significant excess of deaths from neoplasms. 1 man died from nasal cancer (0.02 expected), and 2 each from kidney cancer ( $p = 0.039$ ) and brain cancer ( $p = 0.007$ ).	(11)
NG (125 also exposed to epichlorohydrin)	Cohort, mortality	Isopropanol production, USA	433	No	Lung cancer: RR 2.48 (0.67-6.36), 4; for workers co- exposed to epichlorohydrin, RR 0.69 (0.14-2.02), 3; for workers not exposed to epichlorohydrin.	(73)
"Moderate and high exposure"	Nested case- control, morbidity	Refinery and chemical plant, USA	50 cases of upper respiratory cancer, each matched to at least 3 controls, 30 cases of laryngeal cancer	Smoking, alcohol, history of ear, nose or throat disease	Dose-response for laryngeal cancer risk: OR 4.6 (0.83-25.4) and OR 13.4 (2.08-86.0), for moderate and high exposure, respectively. Number of cases in each group not given.	(217)
NG (lead)	Cohort, mortality	Lead battery plant, USA	4 519 (comparison with national rates)	No	Laryngeal cancer: RR 1.28 (0.47-2.80), 6. Lung cancer RR: 1.24 (1.02-1.50), 109.	(60)

Table 16. Carcinogenicity studies in workers (predominantly males) exposed to the inorganic acids (in chronological order).

Acid exposure level (co-exposure)	Design	Industry/product, Country	Study size	Adjustment in addition to age <sup>a</sup>	Outcome and risk estimate (95 % CI), no. of cases (cohort) no. of exposed cases (case-control)	Reference
0.64-1.12 mg/m <sup>3</sup> (nickel)	Cohort, mortality and morbidity	Soap production, Italy	361	No	Lung cancer: SMR 1.69 (0.55-3.86), 5. Laryngeal cancer: SMR 2.3 (0.09-11.4), 1. Laryngeal cancer: SIR 6.94 (2.25-16.2) or 3.47 (1.13-8.10), 5; depending on reference population.	(80, 116)
NG (SO <sub>2</sub> , fluorides, silica dust, radiation from radon decay products)	Cohort, mortality	Phosphate company, USA	2 607 Caucasians 840 African Americans	No	Lung cancer: SMR 1.62 (1.14-2.23), 37. Laryngeal cancer: SMR 1.91 (0.23-6.9), 2. No excess risk among African Americans.	(28)
0.1-3.1 mg/m <sup>3</sup>	Cohort, mortality and morbidity	Sulphuric acid plant, Sweden	400	No	Respiratory tract cancer: SIR 2.0, p=0.11. Urinary bladder cancer: SIR 3.77, p=0.01.	(72)
NG	Case-control, morbidity	Population-based, USA	183 cases 250 controls	Smoking, alcohol	Laryngeal cancer: RR 0.76 (0.42-1.35), 22.	(38)
NG	Case-control, morbidity	Population-based, Canada	Lung cancer: 857 cases 1 360 controls Oesophageal cancer:	No	Lung cancer: OR 1.2 (0.8-1.9) <sup>b</sup> , 60. Lung (oat-cell) cancer: OR 1.7 (1.0-2.9) <sup>b</sup> , 16. Lung (squamous-cell) cancer: OR 1.5 (1.0-2.4) <sup>b</sup> , 38.	(214)
			99 cases 2 546 controls		Oesophageal cancer: OR 2.2 (1.3-3.6) <sup>b</sup> , 15 (also published by Parent <i>et al</i> (183)).	

Table 16. Carcinogenicity studies in workers (predominantly males) exposed to the inorganic acids (in chronological order).

Acid exposure level (co-exposure)	Design	Industry/product, Country	Study size	Adjustment in addition to age <sup>a</sup>	Outcome and risk estimate (95 % CI), no. of cases (cohort) no. of exposed cases (case-control)	Reference
NG	Case-control, morbidity	Population-based, Canada	183 cases 183 controls	Smoking, alcohol	Laryngeal cancer: OR 1.97 (0.63-6.13), $10; \le 10$ yrs low exposure. OR 3.57 (1.19-10.73), $10; \le 10$ yrs high exposure. OR 4.30 (1.69-10.91), 50; >10 yrs low exposure. OR 5.57 (2.00-15.50), 63; >10 yrs high exposure.	(218)
NG	Cohort, mortality	Two chemical plants: ethanol and isopropanol production, USA	1 031	No	<i>Strong acid process:</i> Deaths from laryngeal cancer from both plants combined: 2 observed, 1 expected. <i>Weak acid process</i> (currently used method): No evidence for carcinogenicity related to the exposure.	(231)
NG	Case-control, morbidity	Population-based, Canada	183 cases 183 controls	Smoking, alcohol	Laryngeal cancer: OR 1.97 (0.63-6.13), $10; \le 10$ yrs low exposure. OR 3.57 (1.19-10.73), $10; \le 10$ yrs high exposure. OR 4.30 (1.69-10.91), 50; >10 yrs low exposure. OR 5.57 (2.00-15.50), 63; >10 yrs high exposure.	(218)
Estimated from detailed job histories	Case-control, morbidity	Population based, Canada	99 cases of which 63 squamous cell carcinomas 533 population	Respondent status birthplace, education,	Oesophageal cancer (OC): OR 2.2 (1.2-4.3), 15; any exposure. OR 2.0 (1.0-4.0), 12; non-substantial exposure. OR 4.1 (1.0-17.2), 3; substantial exposure.	(183)
			controls + 533 patient controls	alcohol, smoking, carotene intake	OC/Squamous cell carcinoma: OR 2.8 (1.2-6.1), 10; any exposure. OR 2.2 (1.2-6.3), 9; non-substantial exposure. OR 3.1 (0.3-28.1), 1; substantial exposure. (Data originally published by Siemiatycki 1991 (214))	
$> 0.5 \text{ mg/m}^3$	Case-control, mortality	Population-based, Lithuania	277 cases 1 108 controls	Smoking	Lung cancer: OR 1.03 (0.76-1.39), 96.	(187) <sup>c</sup>

Table 16. Carcinogenicity studies in workers (predominantly males) exposed to the inorganic acids (in chronological order).

Acid exposure level (co-exposure)	Design	Industry/product, Country	Study size	Adjustment in addition to age <sup>a</sup>	Outcome and risk estimate (95 % CI), no. of cases R (cohort) no. of exposed cases (case-control)	eference
$< 0.5 \text{ mg/m}^3$	Nested case-control, morbidity	Nickel refinery workers, Norway	213 cases 525 controls	Smoking	No excess risk of lung cancer relating to $H_2SO_4$ exposure ORs 0.8-1 (0.3-2.0), 9-14 depending on exposure level.	(97) <sup>c</sup>
Hydrochloric acid						
NG (Cl <sub>2</sub> , SO <sub>2</sub> , CCl <sub>4</sub> )	Case-control, mortality	Chemical plant, USA	28 cases. Two matched control groups from same company.	No	Brain tumours: OR 1.40 (0.70-2.80), 13. OR 1.02 (0.81-1.29), 13 (ORs calculated for comparisons with both control groups).	(29)
NG	Case-control, mortality	Chemical plant, USA	26 cases. Two matched control groups from same company (n = 92 +98).	No	Renal cancer: OR 0.90 (0.44-1.83), 12 <sup>b</sup> OR 0.86 (0.40-1.86), 12 <sup>b</sup> (ORs calculated for comparisons with both control groups).	(30)
Up to $\geq 2.8 \text{ mg/m}^3$ , (also e.g. Cl <sub>2</sub> , SO <sub>2</sub> , CCl <sub>4</sub> , rarely e.g. butadiene)	Nested case- control, mortality	Same as (29), USA	308 cases 2×308 controls	No	Lung cancer: OR 1.02 (0.77-1.35), 129; pooled controls.	(31)
NG (acrylamide)	Cohort, mortality	Four chemical plants, USA and the Netherlands	161 lung cancer deaths	Smoking	Excess of lung cancer (SMR 1.32) (no further details given). The excess was confined to 2 groups in one facility, including the muriatic acid department (11 deaths).	(58)
< 2.8 mg/m <sup>3</sup> $\ge$ 2.8 mg/m <sup>3</sup>	Nested case- control, mortality (follow up with focus on HCl)	Same as (29), USA	See (31) 308 cases 2×308 controls	Race, smoking	Exposure for several years gave no excess lung cancer risk when either duration, cumulative or highest average exposure was used as indices of HCl exposure. Highest average exposure: $< 2.8 \text{ mg/m}^3$ : RR 0.8 (0.5-1.2), 41. $\ge 2.8 \text{ mg/m}^3$ : RR 1.2 (0.8-1.6), 88.	(32)

Table 16. Carcinogenicity studies in workers (predominantly males) exposed to the inorganic acids (in chronological order).

Acid exposure level (co-exposure)	Design	Industry/product, Country	Study size	Adjustment in addition to age <sup>a</sup>	Outcome and risk estimate (95 % CI), no. of cases (cohort) no. of exposed cases (case-control)	Reference
NG	Case-control, morbidity	Population-based, Canada	Rectum cancer: 257 cases 1 299 controls Lung (oat cell) cancer: 159 cases 1 360 controls Non-Hodgkin's lymphoma: 215 cases 2 357 controls		Rectum cancer: OR 1.9 (1.1-3.4) <sup>b</sup> , 18. Lung (oat-cell) cancer: OR 1.6 (1.0-2.6) <sup>b</sup> , 19. Non-Hodgkin's lymphoma: OR 1.6 (1.0-2.5) <sup>b</sup> , 18.	(214)
Nitric acid						
NG (asbestos)	Cohort, incidence	HNO <sub>3</sub> production plant, Norway	287	Smoking	Lung cancer: RR 1.6-5 in different groups, but only statistically significant for maintenance workers.	(109)
NG	Case-control, morbidity	Population-based, Canada	Pancreas cancer: 116 cases 2 454 controls Prostate cancer: 449 cases 1 550 controls Kidney cancer: 177 cases 2 481 controls	No	Pancreas cancer: OR 4.6 (1.9-11-1) <sup>b</sup> , 5. Prostate cancer: OR 3.9 (1.3-11.7) <sup>b</sup> , 9. Kidney cancer: OR 3.1 (1.2-7.8) <sup>b</sup> , 4.	(214)
Nitrogen products, esp. HNO <sub>3</sub> and urea. Exposure measured by duration in years.	Cohort, mortality Nested case- control, mortality	Chemical plant, Lima Ohio, USA	995	Smoking	Bladder cancer: SMR 3.31 (0.90-8.47), 4.	(166)

Table 16. Carcinogenicity studies in workers (predominantly males) exposed to the inorganic acids (in chronological order).

Acid exposure level (co-exposure)	Design	Industry/product, Country	Study size	Adjustment in addition to age <sup>a</sup>	Outcome and risk estimate (95 % CI), no. of cases (cohort) no. of exposed cases (case-control)	Reference
Phosphoric acid						
NG	Case-control, morbidity	Population-based, Canada	Lung cancer: 857 cases 1 360 controls <i>Kidney cancer:</i> 177 cases 2 481 controls	No	Lung cancer: OR 1.9 (1.0-3.8) <sup>b</sup> , 14. Kidney cancer: OR 3.7 (1.7-8.1) <sup>b</sup> , 6.	(214)
Mixed acids						
H <sub>2</sub> SO <sub>4</sub> , HNO <sub>3</sub> (1950s), oxalic acid, ammonium bifluoride and soap (1960s-70s)	Cohort, morbidity	Steel pickling, Sweden	110	No	Laryngeal cancer: 3 observed cases, 0.06 expected (>10 years' induction- latency time).	(7)
H <sub>2</sub> SO <sub>4</sub> H <sub>3</sub> PO <sub>4</sub> (ionising radiation)	Cohort, mortality	Phosphate industry, USA	17 601 white and 4 722 non-white male workers	No	Lung cancer: SMR 1.22 (p<0.05), 117; compared to national rates. SMR 1.03; compared to local rates (Florida).	(44)
H <sub>2</sub> SO <sub>4</sub> H <sub>3</sub> PO <sub>4</sub> ionising radiation	Cohort, mortality	Phosphate industry, USA	Sub-cohorts from previous study (44)	No	RR 0.87 for length of employment > 10 years and work area with exposure to $H_2SO_4$ and $H_3PO_4$ .	(45)
H <sub>2</sub> SO <sub>4</sub> : 0.013-0.22 mg/m <sup>3</sup> H <sub>3</sub> PO <sub>4</sub> : 0.03-0.52 mg/m <sup>3</sup>	Cohort, mortality	Phosphate fertiliser production facility, USA	3 199	No	Tracheal, bronchial and lung cancer: SMR 1.13 (0.61-1.92), 10 (entire cohort). SMR 4.11 (p<0.05) (African American workers with > 10 years employment and follow-up).	(224)
H <sub>2</sub> SO <sub>4</sub> , HNO <sub>3</sub> , HCl and others (self-reported)	Case-control, morbidity	Population based, UK	<ul><li>698 cases of which</li><li>477 self respondent,</li><li>1 683 controls</li></ul>	Sex, race	Multiple myeloma: OR 1.0 (0.6-1.9), 20 (all cases). OR 1.5 (0.8-2.8), 19 (self-respondent cases).	(173)

Table 16. Carcinogenicity studies in workers (predominantly males) exposed to the inorganic acids (in chronological order).

Acid exposure level (co-exposure)	Design	Industry/product, Country	Study size	Adjustment in addition to age <sup>a</sup>	Outcome and risk estimate (95 % CI), no. of cases I (cohort) no. of exposed cases (case-control)	Reference
$H_2SO_4$ (and other acids mainly HCl, 0.2 mg/m <sup>3</sup> ). MMAD probably ~ 5 µm	Cohort, mortality	Steel industry, pickling process, USA	1 165	Smoking	Lung, tracheal and bronchial cancer: SMR 1.64 (1.14-2.28), 35; any acid. Laryngeal cancer: SMR 1.93 (0.23-6.99), 2; any acid. Lung cancer: SMR 1.92 (1.10-3.13), 14; $\geq$ 20 years latency, daily exposure to H <sub>2</sub> SO <sub>4</sub> only. SMR 2.24 (1.02-4.25), 9; subset of 189 workers exposed to acids other than H <sub>2</sub> SO <sub>4</sub> , primarily HCl.	(24)
$H_2SO_4$ : 0.2 mg/m <sup>3</sup> and other acid mists	Cohort, incidence	Steel industry, pickling, USA	879	Smoking, alcohol	Laryngeal cancer: SIR 2.30 (1.1-4.4), 9.	(225)
$H_2SO_4$ mist: 0.2 mg/m <sup>3</sup> and other acid mists	Cohort, mortality,	Steel industry, pickling operations, USA	1 165 follow-up of (24)	Smoking	Lung cancer: SMR 1.36 (0.97-1.84), 41. SMR 1.50 (1.05-2.07), NG; >20 yr since 1st exposure	(226)
	Cohort, mortality	Phosphate fertiliser manufacture, Sweden	1 148	No	Respiratory cancer: SMR 1.51 (1.03-2.2), 29. Laryngeal cancer: SMR 1.60 (0.44-4.10), 4.	(100)
Inorganic acids	Case-control, morbidity	Population-based, Canada	Lung cancer: 857 cases 1 360 controls <i>Kidney cancer:</i> 177 cases 2 481 controls	No	Lung cancer: OR 1.2 (1.0-1.6) <sup>b</sup> , 129; mixed acids. Kidney cancer: OR 1.7 (1.2-2.4) <sup>b</sup> , 32; mixed acids.	(214)

Table 16. Carcinogenicity studies in workers (predominantly males) exposed to the inorganic acids (in chronological order).

Acid exposure level (co-exposure)	Design	Industry/product, Country	Study size	Adjustment in addition to age <sup>a</sup>	Outcome and risk estimate (95 % CI), no. of cases (cohort) no. of exposed cases (case-control)	Reference
$H_2SO_4$ (later substituted with HCl) >1 mg/m <sup>3</sup>	Cohort, mortality, and case-control	Battery manu- facturers and steel workers, United Kingdom	4 401 total cohort, 15 cases 73 controls	No	Laryngeal cancer (cohort): SMR 0.48 (0.01-2.70), 1. Lung cancer (cohort): 0.98 (0.78-1.22), 83. Upper aerodigestive cancer (case-control): OR 2.0 (0.4-10), 12.	(57)
$H_2SO_4$ mist (0.19+0.29 mg/m <sup>3</sup> ) and other acids	Cohort, mortality, morbidity	Steel industry, pickling operations, USA, follow-up of (225).	1 031	Smoking, alcohol	Laryngeal cancer: RR 2.2 (1.2-3.7), 14 RR 2.5 (1.7-4.7), NG; daily contact with $H_2SO_4$ mist.	(227) <sup>c</sup>
Mainly HCl. Also HF, HNO <sub>3</sub> , H <sub>3</sub> PO <sub>4</sub> , H <sub>2</sub> SO <sub>4</sub> and chromic acid	Multi-centre, case-control	Population-based	316 laryngeal cancer + 34 hypopharyngeal cases, 728 controls	Country, smoking, alcohol	Laryngeal cancer: OR 0.94 (0.60-1.49), 37. Hypopharyngeal cancer: OR 3.72 (1.08-12.81), 5; exposure for >15 years.	(213)

Table 16. Carcinogenicity studies in workers (predominantly males) exposed to the inorganic acids (in chronological order).

 $^{\rm a}$  Smoking and alcohol are the strongest known confounders for laryngeal cancer .  $^{\rm b}$  90 % CI.

<sup>c</sup> Published following the IARC monograph (116). CI: confidence interval, NG: not given, SMR: standardised mortality rate, SIR: standardised incidence ratio, RR: relative risk (risk ratio), OR: odds ratio.

# 12. Dose-effect and dose-response relationships

Comprehensive compilations of toxicity data have been presented in Chapters 10-11 (Tables 6-16). The  $LC_{50}$ s suggest that  $H_2SO_4$  and  $HNO_3$  have a higher acute toxicity than HCl and probably also than  $H_3PO_4$ .

## 12.1 Sulphuric acid

Toxicity data from low-level exposures to  $H_2SO_4$  are summarised in Table 18.

# 12.1.1 Single/short-term exposure

### Animal studies

Details have been presented in Chapter 10 (Tables 6, 7 and 9).

Single and repeated, short-term inhalation of low concentrations of  $H_2SO_4$  has resulted in non-specific bronchial hyperresponsiveness *in vitro* and effects on the defences of the rabbit lung (accelerated respiratory clearance, depressed phagocytic capacity, reduced TNF $\alpha$  release and superoxide anion production in macrophages). Effects have appeared at 0.05-0.075 mg/m<sup>3</sup> (70, 201, 205, 250). A lowered intracellular pH in pulmonary macrophages was demonstrated after exposure of rabbits at 0.125 mg/m<sup>3</sup> (49).

A dose-related increase in pulmonary flow resistance in guinea pigs was reported during exposure to 0.1-1 mg/m<sup>3</sup> for 1 hour (MMD 0.3 and 1  $\mu$ m). The smaller droplets also reduced pulmonary compliance in a dose-related manner during and after exposure (13).

Non-specific airway hyperresponsiveness *in vivo* was induced in male guinea pigs exposed for 1 hour to ultrafine  $H_2SO_4$ -particles at 0.2 mg/m<sup>3</sup> (mean diameter 0.06 µm) (48).

Rabbits were exposed to  $H_2SO_4$  at 0.25, 0.5 or 1 mg/m<sup>3</sup> (MMD 0.3 µm), 1 hour/day for 5 days. A dose-dependent decrease in eicosanoid concentration in the lung lavage fluid was reported (203) suggesting modulation of the inflammatory response.

At 0.3 mg/m<sup>3</sup>, the phagocytic capacity of macrophages was depressed by ultrafine acid particles (MMD 0.04  $\mu$ m) and enhanced by larger, sub-micrometer acid particles (MMD 0.3  $\mu$ m) in guinea pigs after both single and repeated exposure. Further, an increased release of TNF $\alpha$  by macrophages was reported after some exposures (47, 48). A 1-hour exposure at the same level (MMD 0.08  $\mu$ m) decreased single breath diffusion capacity for carbon monoxide (46).

Minimal squamous metaplasia in ciliated epithelium in the larynx in rats was observed after a 28-day inhalation exposure to 0.3 mg/m<sup>3</sup> (6 hours/day, 5 days/ week). At higher concentrations (1.38 and 5.52 mg/m<sup>3</sup>), changes were more severe (some cases with parakeratosis) and were accompanied by cell proliferation in the larynx. No effects were observed in the nose or lungs (127). The changes observed at 0.3 mg/m<sup>3</sup> can be regarded as an adaptive response, but suggest a risk of respiratory tract epithelial changes following exposures of longer duration.

Increased *in vitro* tracheal reactivity in rabbit was reported after single exposure for 3 hours to 0.5 mg/m<sup>3</sup> (MMD 0.3  $\mu$ m). At the same level, depressed tracheal mucociliary clearance in dogs was reported after 1-hour exposures to aerosols with larger particles (MMAD 0.9  $\mu$ m), whereas no such effect was reported after exposures to aerosols with MMAD 0.3  $\mu$ m even at 5 mg/m<sup>3</sup> (70, 246). Depressed ciliary beating in trachea was observed in hamster at single exposures to 0.9 (98) and 1.1 mg/m<sup>3</sup> (98, 199). At the higher level, also the respiratory epithelium of the trachea was damaged (98, 199).

Clear dose-effect relationships regarding airway irritation and changes in pulmonary function were reported in three whole-body plethysmograph guinea pig studies with exposures in the range 10-55 mg/m<sup>3</sup> (198), 14-43 mg/m<sup>3</sup> (192), and 24-73 mg/m<sup>3</sup> (249).

### Human studies

Acute effects of  $H_2SO_4$  aerosols have been studied in a number of controlled exposure experiments with healthy and asthmatic volunteers (Table 13), with exposure duration ranging from 10 minutes to 6.5 hours, droplet sizes ranging from 0.1-10 µm (MMAD), relative humidities varying between below 10 to 100 % and exposures carried out at rest, during exercise or combinations thereof.

In three studies on healthy volunteers, bronchial mucociliary clearance was affected after exposures for 1 hour to  $0.1 \text{ mg/m}^3$  (MMAD 0.5 µm) (151, 152, 220). In those and other studies, tracheal transport rates were not affected after exposures to 0.1-1 mg/m<sup>3</sup>. However, an effect on tracheal mucociliary clearance was observed in a study with exposure to larger, fog droplets (MMAD 10.3 µm) at 0.47 mg/m<sup>3</sup> (149).

Respiratory symptoms increased among healthy volunteers exposed to large particles (VMD 10  $\mu$ m) at 0.5 and 1 mg/m<sup>3</sup> for 1 hour with intermittent heavy exercise. Exposure to submicrometer particles (MMAD 0.9  $\mu$ m) at 1 mg/m<sup>3</sup> did not increase symptom ratings compared with air exposure (21, 22). One study (presented as conference proceedings) indicated that workers suffered symptoms of respiratory irritation at exposures in the range 0.1-0.5 mg/m<sup>3</sup> (81).

Bronchial hyperreactivity (after provocation with carbachol) was demonstrated in healthy subjects after a 16-minute exposure to 1 mg/m<sup>3</sup> (237). Generally, no effects on lung function are reported among healthy volunteers at exposures up to 1 mg/m<sup>3</sup> (14, 20-22, 83, 149, 151, 152, 156, 195, 219, 220, 232, 237). However, a few studies indicate that asthmatics respond to lower concentrations. One research team exposed adolescent asthmatics to acid and air or saline in a slightly humid atmosphere (65-75 %). FEV<sub>1</sub> was reduced in subjects exposed to H<sub>2</sub>SO<sub>4</sub> for 45 minutes to as little as 0.035 and 0.07 mg/m<sup>3</sup> (MMAD 0.6  $\mu$ m), significantly so at the lower exposure level. The effect did not remain after 90 minutes of exposure. Exposure to 0.1 mg/m<sup>3</sup> (MMAD 0.6  $\mu$ m) produced reversible significant changes in lung function in 10 adolescent asthmatics following exposure during exercise (134, 136). In adult asthmatics, enhanced bronchoconstriction as well as increased reactivity to carbachol was demonstrated after exposure for 16 minutes to 0.45 mg/m<sup>3</sup> (236, 237). The same research team also reported that prevention of oral ammonia-induced neutralisation of inhaled acid resulted in decrements in pulmonary function following exercise in asthmatics exposed 30 minutes to 0.35 mg/m<sup>3</sup> (238). Other studies showed unaffected pulmonary function in adult asthmatics exposed to up to 1 mg/m<sup>3</sup> or more (195, 232).

Exposure of workers to high but unknown concentrations has been associated with severe effects such as burns, ARDS, pulmonary oedema and fibrosis, bronchioectasis, emphysema and even death (26, 93, 132, 208).

# 12.1.2 Long-term exposure

# Animal studies

Details have been presented in Chapter 10 (Tables 11-12).

The LOAELs after long-term exposure are 0.1 mg/m<sup>3</sup> (1 hour/day, 5 days/week for 6 months) at which impaired mucociliary clearance in donkeys was reported (200) and 0.125 mg/m<sup>3</sup> (2 hours/day, 5 days/week for up to 12 months) at which an accelerated tracheobronchial mucociliary clearance and bronchial epithelial secretory cell hyperplasia were observed in rabbits (204). At 0.25 mg/m<sup>3</sup> (4-12 months, same daily dose as in the previous study) tracheobronchial mucociliary clearance was retarded and a narrowing of the airways, airway hyperresponsiveness, and secretory cell hyperplasia were observed in rabbits (88-90).

In monkeys exposed to larger particles (MMD 2.15  $\mu$ m) at 0.38 mg/m<sup>3</sup> for 78 weeks, an increased respiratory rate, slight hyperplasia of bronchiolar epithelium and thickening of the walls of the respiratory bronchioles were observed. No such changes were reported after exposure to submicrometer particles (MMD 0.54  $\mu$ m) at 0.48 mg/m<sup>3</sup> using an otherwise similar protocol (8).

Dogs exposed to  $0.9 \text{ mg/m}^3$  for 620 days exhibited impaired lung function and reduced heart weights (154).

Monkeys exposed to  $0.88-0.99 \text{ mg/m}^3$  combined with sulphur dioxide and/or fly ash for 78 weeks developed a variety of lung lesions such as hypertrophy, hyperplasia and metaplasia of the bronchiolar epithelium. The effects were attributed to H<sub>2</sub>SO<sub>4</sub>. These effects were not seen in monkeys exposed to  $0.1 \text{ mg/m}^3$  (9).

The few cancer studies in animals have weaknesses in design and reporting. Thus, no conclusion can be drawn regarding the carcinogenic potential of  $H_2SO_4$  in animals.

### Human studies

A number of studies on workers exposed to  $H_2SO_4$  for several years show tooth etching and erosion (Table 15). This has been reported after estimated average exposure levels of 0.2 mg/m<sup>3</sup> (87) and above (186, 230) and after exposure in the range 0.06-2 mg/m<sup>3</sup> (233). In the study by Gamble *et al*, the earliest cases of etching and erosion, respectively, occurred after 4 and 30 months of exposure (87).

In the same study, reduced FVC was observed in the high (average 0.21 mg/m<sup>3</sup>, cumulative exposure > 15 mg/m<sup>3</sup> × months) but not in the low (average 0.10 mg/m<sup>3</sup>, cumulative exposure < 7 mg/m<sup>3</sup> × months) exposure group (87).

In another field study, exposure averaged 0.035-2.1 mg/m<sup>3</sup>. Pale mucosal patches and ulcerations were found only in the workers exposed to more than 0.2 mg/m<sup>3</sup>. Squamous metaplasia, squamous atypia and mild dysplasia of the nasal mucosa were more frequent among exposed than among unexposed workers. The risk of squamous atypia or dysplasia increased with increasing concentration of H<sub>2</sub>SO<sub>4</sub>, but did not correlate with exposure duration. Work-related nasal symptoms were also reported (95).

Human carcinogenicity studies have previously been presented in Table 16, Chapter 11. The IARC classification of strong inorganic acid mist containing  $H_2SO_4$  as carcinogenic to humans (Group 1) was based on several studies (7, 24, 214, 217, 218, 225, 226, 243) with additional support provided by yet another few (80, 100). The risk for laryngeal cancer after exposure to  $H_2SO_4$  was increased in several cohort and case-control studies, but the exposure levels were poorly characterised. Later, one follow-up (227) supporting an association with laryngeal cancer has been published. The evidence for an association with lung cancer is weak (Table 17).

The only study of oesophageal cancer suggests an excess risk with exposure to  $H_2SO_4$  (183, 214).

It should be stated that the validity regarding measurements of the acids in both older and in some more recent field studies is uncertain.

Cancer type/ Industry	Risk estimate	95 % CI	Reference
Laryngeal cancer			
Soap production	SIR 6.94 or SIR 3.47 $^{\rm a}$	2.25-16.2 or 1.13-8.10	(80)
Steel-pickling	SIR 2.30	1.1-4.4	(225)
Steel-pickling	RR 2.2	1.2-3.7	(227) <sup>b</sup>
Steel-pickling	3 cases vs. 0.06 expected		(7)
Refinery and chemical plant	OR 4.6 OR 13.4	0.83-25.4 (moderate exposure) 2.08-86.0 (high exposure)	(217)
Population-based	OR 1.97 OR 3.57 OR 4.30 OR 5.57	0.63-6.13 (≤ 10 yrs low exposure) 1.19-10.73 (≤ 10 yrs high exposure) 1.69-10.91 (> 10 yrs low exposure) 2.00-15.50 (> 10 yrs high exposure)	(218)
Lung cancer			
Steel-pickling	SMR 1.92	1.10-3.13 ( $\geq$ 20 yrs latency, daily exposure to H <sub>2</sub> SO <sub>4</sub> )	(24)
	SMR 1.64 (incl trachea/ bronchus)	1.14-2.28 (any acid)	
Steel-pickling	SMR 1.36 SMR 1.50	0.97-1.84 1.05-2.07 (> 20 yrs since 1st exposure	(226) e)
Phosphate fertiliser manufacture	SIR 1.51	1.03-2.20 (respiratory tract)	(100)
Population-based	OR 1.2 OR 1.7 (oat-cell) OR 1.5 (squamous-cell)	1.0-1.6 <sup>°</sup> (mixed acids) 1.0-2.9 <sup>°</sup> (H <sub>2</sub> SO <sub>4</sub> ) 1.0-2.4 <sup>°</sup> (H <sub>2</sub> SO <sub>4</sub> )	(214)

**Table 17.** Cohort and case-control studies of laryngeal and lung cancer with significantly elevated risk estimates related to exposure to strong inorganic acid mist containing sulphuric acid.

<sup>a</sup> Depending on reference population. <sup>b</sup> 10-year extension of the 1988-study (225).

° 90 % CI.

CI: confidence interval, OR: odds ratio, RR: relative risk (risk ratio), SIR: standardised incidence ratio, SMR: standard mortality ratio.

NOAEL (mg/m <sup>3</sup> )	LOAEL (mg/m <sup>3</sup> )	Species/ subjects <sup>a</sup>	Exposure duration	Effects	Reference
Animal studies					
-	0.05	Rabbit	4 h/d, 14 d	Accelerated respiratory clearance.	(201)
0.05	0.075	Rabbit	2-3 h	Depressed phagocytic capacity, superoxide anion production and TNF $\alpha$ activity by pulmonary macrophages. Increased bronchial hyperresponsiveness ( <i>in vitro</i> ).	(70, 205, 250)
-	0.1	Guinea pig	1 h	Increased pulmonary resistance and decreased compliance.	(13)
-	0.1	Donkey	1 h/d, 5 d/wk, 6 mo	Impaired bronchial mucociliary clearance.	(200)
-	0.125	Rabbit	2 h/d, 5 d/wk, up to 1 yr	Accelerated tracheobronchial mucociliary clearance. Increased secretory cell density in the small airways and possible focal epithelial hyperplasia.	(204)
-	0.2	Guinea pig	1 h	Non-specific airway hyperresponsiveness in vivo.	(48)
-	0.25	Rabbit	1h/d, 5d	Decreased concentration of the eicosanoid $TxB_2$ in broncho- pulmonary lavage fluid.	(203)
-	0.25	Rabbit	1 h/d, 5 d/wk, up to 1 yr	Airway hyperresponsiveness. Decreased tracheobronchial mucociliary clearance, increased airway secretory cell density, and narrowing of the bronchial airways.	(88-90)
-	0.3	Guinea pig	1 h	Decreased single breath diffusion capacity for carbon monoxide.	(46)
-	0.3	Guinea pig	3 h and 3 h/d, 4 d	Increased release of TNF $\alpha$ from pulmonary macrophages. Phagocytic capacity depressed by 0.04-µm particles and enhanced by 0.3-µm particles.	(47)

Table 18. No observed adverse effect levels (NOAELs) and lowest observed adverse effect levels (LOAELs) for low-level inhalation exposures to sulphuric acid.

**Table 18.** No observed adverse effect levels (NOAELs) and lowest observed adverse effect levels (LOAELs) for low-level inhalation exposures to sulphuric acid.

NOAEL (mg/m <sup>3</sup> )	LOAEL (mg/m <sup>3</sup> )	Species/ subjects <sup>a</sup>	Exposure duration	Effects	Reference
0.3 (5 days)	0.3 (28 days)	Rat	5 d and 28 d	Minimal squamous metaplasia in ciliated epithelium of the larynx, not accompanied by cell proliferation. The severity increased in a dose-dependent manner at 1.4 and 5.5 mg/m <sup>3</sup> and was accompanied by cell proliferation.	(127)
0.48 (MMD 0.54 μm)	0.38 (MMD 2.15 μm)	Monkey	23 h/d, 78 weeks	Slight hyperplasia of bronchiolar epithelium and thickening of walls of respiratory bronchioles. Increased respiratory rate. More severe effects at 2.4 mg/m <sup>3</sup> and 4.8 mg/m <sup>3</sup> .	(8)
0.25	0.5	Rabbit	3 h	Increased tracheal reactivity.	(70)
-	0.9	Hamster	2 h	Tracheal mucociliary impairment (depressed ciliary beating).	(98)
-	0.9	Dog	21 h/d, 620 d	Impaired pulmonary function and reduced heart weights.	(154)
0.09-0.11 (with $SO_2$ and/or fly ash)	0.88-0.99	Monkey	23 h/d, 78 weeks	Lung lesions: Focal epithelial and goblet cell hypertrophy and hyperplasia, erosion, thinning and squamous metaplasia of bronchiolar epithelium.	(9)
0.48 (MMD 0.54 μm)	2.4 (MMD 3.6 μm)	Monkey	23 h/d, 78 weeks	Impaired lung function (increased respiratory rate, decreased oxygen tension, deterioration of the distribution of ventilation) and histopathological pulmonary changes.	(8)
Human studies					
-	0.035 and 0.07	Adolescent asthmatics	45, 90 min	Marginal fall in $\text{FEV}_1$ immediately after exposure for 45 but not for 90 min. Values had returned to normal 20 min post-exposure.	(136)
-	0.1	Adolescent asthmatics	40 min incl exercise	Impaired pulmonary function (reversible reductions in $FEF_{50}$ , $FEV_1$ and total respiratory resistance).	(134, 136)
-	0.1	Healthy	1-2 h	Increased bronchial mucociliary clearance of large tracers (MMAD 7.5 $\mu$ m). Reduced bronchial mucociliary clearance of smaller tracers (MMAD 4-5 $\mu$ m).	(151, 152, 220)

NOAEL (mg/m <sup>3</sup> )	LOAEL (mg/m <sup>3</sup> )	Species/ subjects <sup>a</sup>	Exposure duration	Effects	Reference
-	~ 0.1	Workers	> 6 months	Symptoms of eye and respiratory irritation.	(81)
-	~ 0.2	Workers	10 yr average	Tooth etching and erosion.	(87)
0.1	~ 0.2	Workers	10 yr average	Reduced FVC.	(87)
-	> 0.2	Workers		Cellular changes of the nasal mucosa.	(95)
0.35 (high oral NH <sub>3</sub> )	0.35 (low oral NH <sub>3</sub> )	Asthmatics	20 min rest + 10 min exercise	Impaired pulmonary function (reduced $\text{FEV}_1$ and maximum expiratory flow rates at 60 % total lung capacity).	(238)
0.1	0.45	Asthmatics	16 min	Enhanced bronchoconstriction.	(236, 237)
-	0.47	Healthy	1 h incl exercise	Accelerated tracheal clearance and accelerated small airway mucociliary clearance.	(149)
0.47 (MMAD 10 µm)	0.5 (VMD 10 µm)	Healthy	1 h incl exercise	Small increase in upper and lower respiratory symptoms.	(22, 149)
0.38 (MMAD 0.9 µm)	1 (MMAD 0.9 µm)	Asthmatics	1 h incl exercise	Lower respiratory and non-respiratory symptoms increase.	(21)
0.5 (VMD 10 µm)	1 (VMD 10 µm)	Asthmatics	1 h incl exercise	Upper and lower respiratory symptoms increase.	(22)
0.3	1.0	Asthmatics	1 h	Reduced bronchial mucociliary clearance.	(219)
0.5 (VMD 10 µm)	1 (MMAD 0.6-1)	Healthy	16 min	Enhanced bronchoconstriction.	(22, 237)

Table 18. No observed adverse effect levels (NOAELs) and lowest observed adverse effect levels (LOAELs) for low-level inhalation exposures to sulphuric acid.

<sup>a</sup> Adult; unless stated otherwise.

FEV<sub>1</sub>: forced expiratory volume in one second, FEF<sub>50</sub>: forced expiratory flow at 50 % of forced vital capacity, FVC: forced vital capacity, MMAD: mass median aerodynamic diameter, MMD: mass median diameter, TNF: tumour necrosis factor, Tx: thromboxan, VMD: volume median diameter.

# 12.2 Hydrochloric acid

Toxicological effects after low-level exposures are compiled in Table 19.

## 12.2.1 Single/short-term exposure

### Animal studies

Details are presented in Chapter 10 (Tables 6 and 8). In male guinea pigs exposed 2 hours/day, 5 days/week for 7 weeks, a NOAEL of 15 mg/m<sup>3</sup> was identified for histological changes in the lungs and airways and for effects on pulmonary function (181).

In mice, the respiratory rate decreased by approximately 10 % at 56 mg/m<sup>3</sup> and RD<sub>50</sub> was calculated to be 432 mg/m<sup>3</sup> (23). Male mice exposed for 6 hours/day to the same level were all dead or moribund after 3 exposures. Severe exfoliation, erosion, ulceration and necrosis, and mild inflammation of the respiratory epithelium were observed, as well as mild ulceration and necrosis of the olfactory epithelium, and serous exudate. No lesions were induced in the lower respiratory tract (41). At approximately the same level (448 mg/m<sup>3</sup>), sensory and pulmonary irritation in guinea pigs is reported (42).

In mice and guinea pigs, 700 mg/m<sup>3</sup> (lowest dose tested) reduced the respiratory frequency. The exposure resulted in the death of 4/6 mice and of 1/9 guinea pigs (124). In baboons, there was no significant effect in pulmonary function at the same level of exposure (123). In both studies, the exposure duration was 15 minutes (head-only). Mice, guinea pigs, and rats exposed to higher levels (3 500, 5 880 and 5 880 mg/m<sup>3</sup>, respectively) experienced damaged respiratory tract epithelium and lung injuries. In baboons, higher level (7 000-14 000 mg/m<sup>3</sup>) of exposure increased respiratory frequency and minute volume registered by plethysmography.

Exposure of rats to 1 820 mg/m<sup>3</sup> HCl in plethysmographs for 30 minutes caused marked toxicity in the nasal region in nose breathers. Even higher toxicity was seen after forced mouth breathing, such as major tissue disruption in the trachea, accumulation of inflammatory cells and exudates in the airways and more peripheral lung damage. Breathing frequency, pulmonary ventilation, and tidal volume was reduced by 4-7 % in animals breathing through the nose compared to the controls. In the mouth breathing group, pulmonary ventilation increased during exposure (223).

### Human studies

The toxicological database for humans is very limited.

No pulmonary function impairment and no lower or upper respiratory symptoms were observed in asthmatic adults exposed to 1.12 and 2.52 mg/m<sup>3</sup> HCl for 45 minutes (228). The authors also listed earlier human studies and case reports of HCl exposure in which it was estimated that the odour threshold was in the range 1.5-7.5 mg/m<sup>3</sup>, that work can be carried out undisturbed at 15 mg/m<sup>3</sup>, is difficult at 15-75 mg/m<sup>3</sup>, intolerable at 75-150 mg/m<sup>3</sup>, and that 1 950-3 000 mg/m<sup>3</sup> is a lethal concentration (228).

NOAEL	LOAEL	Species/	Exposure	Effects	Reference		
$(mg/m^3)$	$(mg/m^3)$	subjects	duration				
Animal studies							
2.8 <sup>a</sup>	14	Rat	6 h/d, 5 d/wk, for life	Increased incidences of hyper- plasia in larynx and trachea.	(212)		
-	15	Rat	6 h/d, 5 d/wk, 90 d	Nasal irritation.	<sup>b</sup> cited in (68)		
-	56	Mouse	10 min	Respiratory rate reduced by 10 % $(RD_{50} = 432 \text{ mg/m}^3).$	(23)		
15	-	Guinea pig	2 h/d, 5 d/wk, 7 wk	No histological changes in the lungs and airways or effects on pulmonary function.	(181)		
Human stu	ıdies						
4.5-7.7 <sup>°</sup>	1.8-12.4	Workers	Occupational	Tooth erosion indicated (lack of control group).	(165, 191)		
2.5	-	Asthmatics	45 min	No irritation of the respiratory tract and no pulmonary function impairment.	(228)		
3-4.5 °	5.2 °	Workers	Occupational	Initial mild irritation of the airway mucosa, which regressed rapidly.	(165)		
-	26-34 °	Workers	Occupational	Breathing difficulties, chronic bronchitis (after years of exposure).	(165)		

**Table 19.** No observed adverse effect levels (NOAELs) and lowest observed adverse effect levels (LOAELs) for low-level inhalation exposures to hydrochloric acid.

<sup>a</sup> Calculated from the LOAEL by the German MAK-committee by linear interpolation between the incidences at 0 and 14 mg/m<sup>3</sup>.

<sup>b</sup> Chemical Industry Institute of Toxicology. 90-day inhalation study of hydrogen chloride gas in B6C3F1 mice, Sprague-Dawley rats, and Fischer-344 rats. ToxiGenics, Inc for CIIT Research Triangle Park, NC, USA, CIIT Docket No. 20915.

<sup>c</sup> No further details given (see below).

No irritation of the mucous membranes was observed among workers in steel pickling facilities exposed to 3-4.5 mg/m<sup>3</sup> whereas initial mild irritation of the airway mucosa, which regressed rapidly was observed at 5.2 mg/m<sup>3</sup>, slight irritation at 7-11 mg/m<sup>3</sup> and breathing difficulties at 26-34 mg/m<sup>3</sup> (165). The study is based on many years of observations but is not conducted and reported according to current standards.

Accidental exposure to high air levels of HCl has produced severe symptoms, and respiratory impairments (35, 126). Exposure of the skin to a high concentration of HCl gas or to a concentrated solution of HCl will cause burns; repeated or prolonged exposures to dilute solutions may cause dermatitis (5).

## 12.2.2 Long-term exposure

### Animal studies

In male and female mice and rats exposed to 0, 10, 20, or 50 ppm (0, 15, 30, or  $75 \text{ mg/m}^3$ ) for 6 hours/day, 5 days/week for 90 days, local irritative effects on the

nose were observed in both species. The systemic NOAEL was found to be 30 mg/m<sup>3</sup>, and the LOAEL for irritation 15 mg/m<sup>3</sup> (industry report<sup>1</sup> cited in (68)).

Animal carcinogenicity studies are listed in Table 12. Male rats exposed to air containing 14 mg/m<sup>3</sup> HCl for life developed hyperplasia in larynx and trachea, but no cancer. The incidences of laryngeal and tracheal hyperplasia were 22 % and 26 % compared with 2 % and 6 % among the controls. No other level of exposure was tested (212). In a previous study with equal exposure level and duration, no nasal cancer was observed (10).

# Human studies

Effects of occupational exposure are presented in Tables 15-16.

In a hot dip zinc galvanising plant, 90 % of 38 examined workers in the pickling process in which a 15 % HCl solution was used had grade 1-2 dental erosion of the incisor teeth. Exposure levels (geometric mean) ranged from 1.8 to 12.4 mg/m<sup>3</sup>. The dental effect could not be causally linked to the HCl exposure due to the lack of control group and the small number of workers, which made dose-response analysis impossible (191).

In a report from steel pickling facilities, chronic bronchitis was reported after exposure to approximately  $30 \text{ mg/m}^3$  for years. It was also stated (no further information given) that no damage to the teeth occurred at average concentrations of 4.5-7.7 mg/m<sup>3</sup> (165). The study does not comply with current standards.

IARC concluded in 1992 that HCl was not classifiable as to its carcinogenicity to humans.

### 12.3 Nitric acid

Toxicity data from low-level exposures to nitric acid are presented in Table 20.

### 12.3.1 Single/short-term exposure

### Animal studies

Details are presented in Chapter 10 (Tables 6 and 10).

The lowest effect levels reported concern the rabbit lung defence. A reduced production of superoxide anions by alveolar macrophages was observed after 4 weeks of exposure to  $\geq 0.05 \text{ mg/m}^3$ . At  $\geq 0.15 \text{ mg/m}^3$ , reduced bronchial responsivity to smooth muscle constrictor challenge and reduced TNF $\alpha$  activity were seen (207). Decreased superoxide anion production in rat pulmonary macrophages was also observed after a 4-day repeated exposure to 0.25 mg/m<sup>3</sup>, the only dose tested, as well as increased elastase inhibitory capacity of lung lavage fluid (175).

No bronchoconstriction was seen in normal sheep exposed for 4 hours to  $4.1 \text{ mg/m}^3$ . However, in allergic sheep this concentration caused increased airway hyperreactivity to carbachol (2).

<sup>&</sup>lt;sup>1</sup> Chemical Industry Institute of Toxicology (1984). 90-Day inhalation study of hydrogen chloride gas in B6C3F1 mice, Sprague-Dawley rats, and Fischer-344 rats. ToxiGenics, Inc for CIIT Research Triangle Park, NC, USA, CIIT Docket No. 20915.

NOAEL	LOAEL	Species/	Exposure	Effects	Reference
$(mg/m^3)$	$(mg/m^3)$	subjects	duration		
Animal stu					
-	0.05	Rat	40 wk	Elevated stress-inducible heat-shock protein 70 in the lungs.	(164, 215, 248)
-	0.05	Rabbit	4 wk	Decreased superoxide anion production in alveolar macrophages.	(207)
0.05	0.15	Rabbit	4 wk	Reduced <i>in vitro</i> bronchial respons- ivity to smooth muscle constrictor challenge and TNF $\alpha$ activity in alveolar macrophages.	(207)
-	0.25	Rat	4 h/d, 4d	Decreased superoxide anion production in isolated pulmonary macrophages. Increased elastase inhibitory capacity of lung lavage fluid.	(175)
4.1 (n, a)		Sheep	4 h	No bronchoconstriction.	(2)
4.1 (n)	4.1 (a)	Sheep	4 h	Hyperreactivity (carbachol).	(2)
Human sti	udies				
0.125		Adolescent asthmatics	40 min	No effects on pulmonary function.	(137)
-	0.2	Healthy	2 h	Alveolar macrophage function effected (increased phagocytic activity and resistance to infection, and decreased superoxide anion production.	(25)
0.5		Healthy	4 h	No effect on symptom scores, no proximal airway and distal lung injuries assessed by pulmonary function measurements, lavage and biopsies.	(17)

**Table 20.** No observed adverse effect levels (NOAELs) and lowest observed adverse effect levels (LOAELs) for low-level inhalation exposures to nitric acid.

a: allergic, n: normal, TNF: tumour necrosis factor.

At repeated exposure to much higher levels (190-320 mg/m<sup>3</sup>), impaired pulmonary function, histopathological findings such as chronic inflammation of smaller airways, and hyperresponsiveness to histamine were observed in dogs (85, 185).

### Human studies

Pulmonary function parameters were not affected in adolescent asthmatics (n=28), 12-19 years old, after inhalation of a combination of 0.05 ppm ( $0.125 \text{ mg/m}^3$ ) HNO<sub>3</sub>, 0.12 ppm O<sub>3</sub> and 0.30 ppm NO<sub>2</sub> for 90 minutes via a mouthpiece (137).

Exposure of 9 healthy subjects to 0 and 0.2 mg/m<sup>3</sup> of HNO<sub>3</sub> vapour for 2 hours, did not affect pulmonary function, subjective symptoms or indicators of airway injury and inflammation. However, there was an effect of HNO<sub>3</sub> on alveolar macrophage function (increased phagocytic activity and resistance to infection, and decreased superoxide anion production) (25).

In 10 healthy volunteers, no effects on symptoms or pulmonary function were seen after exposure to  $0.5 \text{ mg/m}^3 \text{ HNO}_3$  for 4 hours (17) nor after exposure for 2 hours to 0.4 mg/m<sup>3</sup> (16), the only levels tested.

In an old study, exposure to 11-12 ppm (27-30 mg/m<sup>3</sup>) for more than 1 hour was considered intolerable and dangerous to human health. Exposure to 84 ppm (210 mg/m<sup>3</sup>) was tolerated for only 2-3 minutes. The results were based on only 1-2 individuals (Diem<sup>1</sup>, cited in (66)).

Inhalation of high, but unknown, concentrations of HNO<sub>3</sub> fumes has caused numerous deaths; death can be delayed several days. Contact with concentrated acid causes severe skin burns and corneal injury leading to blindness.

# 12.3.2 Long-term exposure

# Animal studies

Elevated stress-inducible heat shock protein 70 was observed in lungs from rats exposed to  $HNO_3$  vapour at 0.05 mg/m<sup>3</sup> for 40 weeks. Body weight, lung polyamine contents and lung clearance was not affected (164, 215, 248). No other concentrations were tested.

# Human studies

No quantitative human data on effects from long-term exposure to HNO<sub>3</sub> have been found in the peer-reviewed literature.

# 12.4 Phosphoric acid

Apart from an  $LC_{50}$  of > 850 mg/m<sup>3</sup> (Table 6) no quantitative data on effects from exposure to  $H_3PO_4$  have been found in the peer-reviewed literature.

 $H_3PO_4$  is a mild irritant of the eyes, upper respiratory tract and the skin, and the dust is especially irritating to skin in the presence of moisture. Severe reactions including death have been described after excessive exposure.

# 13. Previous evaluations by national and international bodies

# 13.1 Sulphuric acid

The Nordic Expert Group for Documentation of Occupational Exposure Limits (previous name for NEG) concluded in 1992 that the critical effect of acid aerosols is acute and chronic irritation of the airways. NEG stated further that some human volunteer studies (not replicated by others, NEG claims in 1992) indicated that  $H_2SO_4$  aerosols in the one-micron size range may induce moderate increase of airway resistance in asthmatics at concentrations around 0.1 mg/m<sup>3</sup> and even lower, and that submicron long-term exposures to  $H_2SO_4$  aerosols in the 0.1-0.5 mg/m<sup>3</sup> range induced changes in airways and lungs in animal models.

<sup>&</sup>lt;sup>1</sup> Diem L. Untersuchungen über die Einatmung von Saltpetersäure-dämpfen (thesis). Würzburg, 1907.

Long-term acid aerosol exposures in doses found in several industries should be suspected as a causative factor for laryngeal cancer (142).

In 1992, *IARC* classified occupational exposure to strong inorganic acid mists containing H<sub>2</sub>SO<sub>4</sub> as carcinogenic to humans (Group 1) (116).

The Agency for Toxic Substances and Disease Registry (ATSDR) (1998) was not able to derive a minimal risk level (MRL) for inhalation exposure in humans to  $H_2SO_4$ . ATSDR argued that physiological factors and conditions are just as important as the exposure concentration, and that the response depends on individual factors. Because the occupational studies identified NOAELs higher than LOAELs in acute-duration studies in animals, they were not considered appropriate for MRL derivation (19).

The German Research Foundation (DFG) evaluation of  $H_2SO_4$  for MAKvalues concluded in 2001 that the most sensitive endpoint for the local effects in man has proved to be the alterations in mucociliary clearance seen after exposure to concentrations of 0.3 mg/m<sup>3</sup> or more (67).

*The US National Toxicology Program* (*NTP*) published a Report on Carcinogens (RoC) in 2002 titled "Strong inorganic acid mists containing sulfuric acid". In the report,  $H_2SO_4$  was identified as a carcinogen (180).

The Dutch Expert Committee on Occupational Standards (DECOS) conducted an evaluation of carcinogenicity and genotoxicity of strong inorganic acid mists containing  $H_2SO_4$  in 2003, and concluded it to be carcinogenic to humans (comparable with EU category 1), acting as non-stochastic genotoxic carcinogens (meaning that an OEL can be derived using a threshold model) (106).

The American Conference of Governmental Industrial Hygienists (ACGIH) (2004) has recommended an 8-hour threshold limit value (TLV) of 0.2 mg/m<sup>3</sup> for  $H_2SO_4$  (thoracic particulate fraction) to avoid mucociliary clearance and pulmonary function changes, and also other effects (6).

The Scientific Committee on Occupational Exposure Limits (SCOEL) in the European Union concluded in 2007 that experimental studies in a range of animals suggest respiratory tract effects from repeated exposure to concentrations around 0.3 mg/m<sup>3</sup>, with the possibility of effects of some health significance even at concentrations as low as 0.1 mg/m<sup>3</sup>. SCOEL emphasised the human carcinogenicity data and the larynx as a site of particular concern. SCOEL stated that the presumed mechanism by which laryngeal cancer arose in workers is chronic inflammation of the epithelium in this region, caused by the acidity of H<sub>2</sub>SO<sub>4</sub> aerosols, a hypothesis that links with the findings of the rat inhalation study of Kilgour *et al* (2000). A threshold would apply to this presumed carcinogenic mechanism, that being the dose at which the buffering capacity of the epithelial cells is overwhelmed and a significant fall in cellular pH occurs. SCOEL concluded that long-term exposure should be maintained below 0.1 mg/m<sup>3</sup> and hence recommended an 8-hour TWA limit of 0.05 mg/m<sup>3</sup> (210).

### 13.2 Hydrochloric acid

*The International Programme on Chemical Safety (IPCS)* considered in 1982 sensory irritation and objective changes in pulmonary function to be likely critical effects from exposure to chlorine and HCl, but was unable to establish a value for the protection from effects from HCl (117).

*IARC* stated in 1992 that HCl is not classifiable as to its carcinogenicity to humans (Group 3) (115).

ACGIH recommended in 2003 that a TLV-Ceiling of 2 ppm  $(2.8 \text{ mg/m}^3)$  for HCl should be based on the NOAEL from Stevens *et al* (228) of 1.8 ppm (2.52 mg/m<sup>3</sup>) in human volunteers exposed for 45 minutes with intermittent exercise. The asthmatic subjects in this study showed no adverse respiratory health effects of inhalation of this low exposure. The recommended TLV should prevent acute irritation (5).

The *DFG* evaluation of HCl for MAK-values (Germany) concluded in 1982 that it could not be excluded that long-term exposure of workers to the current MAK-value of 5 ppm (dated 1958) could cause changes in respiration mechanics and possible effects on lung function should be investigated (65). Later, the MAK-value was reduced from 5 ppm to 2 ppm (2004) based on the 2-year studies in rats exposed to 10 ppm HCl by inhalation, and the finding of hyperplasia in larynx and trachea (10, 212). Linear interpolation was carried out between the incidence at 10 ppm and that of controls. This resulted in 2 ppm HCl as an estimated concentration at which the incidence of hyperplasia was not significantly different from the controls (68).

# 13.3 Nitric acid

The *DFG* evaluation of HNO<sub>3</sub> (Germany) concluded in 1992 that the MAK-value should be lowered to  $2 \text{ ml/m}^3$  (2 ppm) to protect against irritation of the airways and lungs and be regarded as a provisional value. HNO<sub>3</sub> was classified in category I for the limitation of exposure peaks because it is a local irritant (66).

The *European commission* recommended in 1996 a short-term exposure limit (STEL) of 1 ppm to protect against acute irritation. The limit value was based on the study by Sackner and Ford in which inhalation of 1.6 ppm  $(4.2 \text{ mg/m}^3)$  for 10 minutes was without effects on the pulmonary function in healthy volunteers and an old study from 1907 indicating that humans are more sensitive than cats and rabbits (75).

*IPCS* summarised the studies of pulmonary response to  $HNO_3$  vapour in sheep (Section 10.2) and in rats (Section 10.3) (118) in 1997. Sheep exposed head-only to 4.12 mg/m<sup>3</sup> for 4 hours showed decreased specific pulmonary flow resistance "indicating the absence of any bronchoconstriction". Effects on alveolar macrophages and elastase inhibitory capacity of bronchoalveolar lavage were seen in rats exposed for 4 hours to 1 mg/m<sup>3</sup>, or for 4 hours/day for 4 days to 0.25 mg/m<sup>3</sup>.

*ACGIH* recommended in 2001 a TLV-TWA of 2 ppm and a TLV-STEL of 4 ppm for HNO<sub>3</sub>. The values were considered sufficiently low to prevent ocular and upper respiratory tract irritation and also dental corrosion (3).

*SCOEL* in the European Union recommended an 8-hour TWA of 1 ppm (2.6 mg/m<sup>3</sup>) and a STEL of 2 ppm (5.2 mg/m<sup>3</sup>) in 2001 (209), based on the ACGIH-document and an abstract from Sackner and Ford (1981, not fully published) (3). Sackner and Ford briefly described a study on healthy volunteers exposed to 1.6 ppm ( $4.2 \text{ mg/m}^3$ ) for 10 minutes with no effect on pulmonary function.

# 13.4 Phosphoric acid

Based on data presented as brief communication to ACGIH, *the European Commission* concluded in 1992 that fumes of phosphorous pentoxide are unlikely to produce significant irritation at concentrations  $0.8-5.4 \text{ mg/m}^3$ . It was also noted that phosphorous pentoxide is a powerful dehydrating agent combining with atmospheric moisture or in the respiratory tract to produce H<sub>3</sub>PO<sub>4</sub>. Since this reaction generates heat and desiccates tissues it is likely to cause more tissue damage than pre-formed H<sub>3</sub>PO<sub>4</sub>. Applying the results of studies on phosphorous pentoxide would therefore supply an adequate margin of safety (74).

Based on analogy from comparable experience and data for  $H_2SO_4$  (they recommended 1 mg/m<sup>3</sup> at the time), *ACGIH* recommended in 2001 a TLV-TWA of 1 mg/m<sup>3</sup> and a TLV-STEL of 3 mg/m<sup>3</sup> for  $H_3PO_4$ . The TLV-TWA was said to be below the concentration that causes throat irritation among unacclimated workers (4).

# 14. Evaluation of human health risks

## 14.1 Assessment of health risks

Sulphuric, hydrochloric and nitric acids ( $H_2SO_4$ , HCl, HNO\_3) are strong mineral acids, whereas phosphoric acid ( $H_3PO_4$ ) is weaker. HNO\_3 and  $H_2SO_4$  are also oxidants of which HNO\_3 is the stronger. Acute toxicity data (Table 6) suggests that these two acids have a higher acute toxicity than HCl and possibly also than  $H_3PO_4$ . All acids are hygroscopic. The more volatile HCl and HNO\_3 will appear in air as vapours or aerosols, whereas the less volatile  $H_2SO_4$  and  $H_3PO_4$  will be present in air primarily as aerosols. Droplet size is usually the critical factor governing the site of deposition and hence the respiratory response to the acid aerosols. Industrial aerosols can have MMADs as large as 14 µm (160).

All four acids are direct irritants causing adverse effects at the site of contact. Severe reactions including corrosion and destruction of body tissues from burns leading to ulcers, as well as blindness and death have been described after contact with skin, eyes, and mucous membranes. Repeated or prolonged skin exposure to dilute solutions may cause dermatitis. There are no data describing dermal absorption but the polarity of the acids suggests little absorption via an intact skin barrier.

Following absorption, the toxic effects of the acids will mainly be from their protolysis, yielding  $H^+$  dissolved in the mucosa. The protons will lower the local pH and induce cell membrane injuries and ulcerations. The anions are essential and enter the body pool. No systemic effects are expected at relevant exposure levels. However, effects from both protons and anions such as acidosis, cyanosis, hyperphosphataemia and hypocalcaemia may occur after excessive exposure.

No sensitising effects have been reported after exposure to any of the acids.

The risk for dental erosion has been shown to increase with increasing concentration of acids or with exposure time and duration of employment.

Low pH has produced positive responses such as clastogenic effects and DNAdamage in some *in vitro* mutagenicity assays. There are no *in vivo* data.

An excess risk of laryngeal cancer has been found among workers exposed to strong inorganic acid mists containing  $H_2SO_4$  and this exposure has also been classified by IARC as carcinogenic to humans (Group 1). It is not possible to conclude as to the carcinogenicity of the other acids.

NOAELs and LOAELs for low-level inhalation exposures are presented in Tables 18-20.

Except for  $H_2SO_4$ , the number of studies is limited, and many studies are old. The validity regarding exposure measurements of the acids in both older and in some more recent field studies is uncertain.

### Sulphuric acid

 $H_2SO_4$  is hygroscopic enough to char the skin. The reaction with water is rapid and liberates sufficient heat to produce burns similar to thermal burns. The potent dehydrating ability combined with the heat generation means that  $H_2SO_4$  causes more tissue damage than expected from acidity alone.

The effect of  $H_2SO_4$  aerosols on pulmonary defence mechanisms has been investigated in both animals and humans. In some animal studies, tracheobronchial mucociliary or alveolar clearance of inert particles has been affected and in humans, an effect on bronchial mucociliary clearance has been reported in several studies. Both decreases and increases in clearance have been observed. The outcome seems to depend on acid dose and the sizes of the  $H_2SO_4$  aerosol droplets (hence of deposition sites) and of the inert test particles, respectively.

Alterations in bronchial mucociliary clearance after single exposures in healthy volunteers were reported at  $0.1 \text{ mg/m}^3$  in a few studies (lowest dose tested) and in experimental animals after short- and long-term exposure at the same or even lower levels. Changes in the phagocytic capacity and in biological modifiers critical for maintaining pulmonary immunocompetence (such as reduced super-oxide anion production and TNF $\alpha$  release in pulmonary macrophages) as well as increased hyperresponsiveness have also been observed in animals at these low levels.

In humans, tracheal clearance or tracheal transport rates were affected only in one study in which exposure was to larger droplets (MMAD 10.3  $\mu$ m).

Cellular changes in the respiratory tract have been shown after  $H_2SO_4$  exposure. Alterations in the nasal mucosa have been observed in workers after long-term exposure at > 0.2 mg/m<sup>3</sup>. Epithelial secretory cell hyperplasia in the small airways of rabbits has been reported after long-term exposure to 0.125 and 0.25 mg/m<sup>3</sup> (same daily doses). Minimal squamous metaplasia in ciliated epithelium of the larynx not accompanied by cell proliferation occurred in rats after a 28-day exposure at 0.3 mg/m<sup>3</sup>. The severity of the lesions increased after exposure at higher air levels. The changes observed at 0.3 mg/m<sup>3</sup> were regarded as an adaptive response and suggest a risk of respiratory tract epithelial changes following exposures of longer duration. In monkeys, long-term exposure to submicrometer droplets (MMD 0.54 µm) at 0.48 mg/m<sup>3</sup> produced no alterations of pulmonary structures whereas droplets with MMD 2.15 µm at 0.38 mg/m<sup>3</sup> produced slight histological changes (bronchiolar epithelial hyperplasia and thickening of walls of respiratory bronchioles and alveoli) and an increased respiratory frequency, indicating that droplet size was the important factor.

Overall, the above-mentioned results correspond well with data reviewed by Wanner 1996, who concluded that short-term exposure to cigarette smoke, atmospheric pollutants (including  $H_2SO_4$ ) and oxygen at low to intermediate concentrations/doses may cause transient ciliostimulation, mucous secretion, and stimulation of mucociliary clearance. In contrast, long-term exposure to low to intermediate concentrations or short-term exposure to high concentrations can produce changes of the airway mucosa with disruption of the ciliated epithelium, mucous cell hyperplasia and metaplasia, hypersecretion and impaired clearance.

In healthy volunteers, the LOAEL for upper and lower respiratory symptoms was 0.5 mg/m<sup>3</sup> after a single exposure to large droplets (VMD 10  $\mu$ m) in a humid atmosphere. Smaller particles (MMAD 0.9  $\mu$ m) did not induce respiratory symptoms in healthy subjects even at 1 mg/m<sup>3</sup>. In occupationally exposed, respiratory and eye irritation were reported at 0.1-0.5 mg/m<sup>3</sup> in one study.

Bronchial hyperreactivity in healthy subjects were demonstrated in a study after a 16-minute exposure to 1 mg/m<sup>3</sup>. However, pulmonary function has in general not been affected after single exposures up to 1 mg/m<sup>3</sup> although some studies indicate that asthmatics, especially adolescent asthmatics, may be more vulnerable. Thus, FEV<sub>1</sub> was reduced in asthmatics (13-18 years) exposed for 45 minutes to 0.035 and 0.07 mg/m<sup>3</sup>, significantly so at the lower dose. No effect on lung function was seen when the same subjects were exposed for 90 minutes. Exposure to 0.1 mg/m<sup>3</sup> produced significant changes in lung function in 10 adolescent asthmatics after exercise. Thus, the LOAEL for impaired pulmonary function in adolescent asthmatics may be at or even below 0.1 mg/m<sup>3</sup>. At the same level (0.1 mg/m<sup>3</sup>) impaired pulmonary function was reported in guinea pigs exposed for 1 hour. Enhanced bronchoconstriction was observed in adult asthmatics exposed to 0.45 mg/m<sup>3</sup> at rest, and, following exercise at 0.35 mg/m<sup>3</sup> in combination with low oral ammonia levels. Other studies have shown no pulmonary function effects in adult asthmatics exposed to up to  $1 \text{ mg/m}^3$  or above.

In workers, a reduced FVC was observed at an average exposure to  $0.21 \text{ mg/m}^3$  (cumulative exposure above 15 mg/m<sup>3</sup> × months) whereas no effect was seen at an average exposure of  $0.10 \text{ mg/m}^3$  (cumulative exposure < 7 mg/m<sup>3</sup> × months).

Tooth etching and erosion in workers has been shown after exposure to approximately  $0.2 \text{ mg/m}^3$  and above. At an estimated average exposure of  $0.23 \text{ mg/m}^3$ , the earliest cases of etching and erosion, respectively, occurred after 4 and 30 months.

An excess risk of laryngeal cancer has been found among workers exposed to strong inorganic acid mists containing  $H_2SO_4$ . Although  $H_2SO_4$  was the commonest exposure, the levels of  $H_2SO_4$  were poorly characterised in the studies and the validity regarding past measurements of the acid is uncertain. It is therefore not possible to associate an increased cancer incidence with a particular exposure level.

The development of cancer seems to be secondary to the tissue damage caused by the acid. Thus a threshold is likely, i.e. induction of laryngeal cancer will only occur at exposure levels surpassing the buffering capacity of the epithelial cells.

### Hydrochloric acid

The toxicological database is small.

In asthmatics, no airway irritation or pulmonary function changes were reported after exposure to up to 2.5 mg/m<sup>3</sup> for 45 minutes.

In a report not complying with current standards regarding performance and scientific documentation, it was stated that no irritation of the mucous membranes was observed among workers in steel pickling facilities exposed to 3-4.5 mg/m<sup>3</sup> but initial mild irritation of the airway mucosa, which regressed rapidly, at 5.2 mg/m<sup>3</sup>. Slight irritation was observed at 7-11 mg/m<sup>3</sup> and breathing difficulties at 26-34 mg/m<sup>3</sup>. In the same report, it was also stated that no damage to the teeth occurred at average concentrations of 4.5-7.7 mg/m<sup>3</sup>. Dental erosion was indicated in a study of workers (prevalence 90 %) where average exposure levels ranged from 1.8 to 12.4 mg/m<sup>3</sup>. The lack of control group, however, weakens the result.

Hyperplasia in larynx and trachea, but no cancer, was reported after life-time exposure of male rats at  $14 \text{ mg/m}^3$  HCl, the only level tested. Incidences of laryngeal and tracheal hyperplasia were 22 % and 26 %, respectively, compared with 2 % and 6 % among the controls.

### Nitric acid

The documentation of health effects from exposure to HNO<sub>3</sub> is scarce.

Pulmonary function parameters were not affected in adolescent asthmatics inhaling a combination of  $0.125 \text{ mg/m}^3 \text{ HNO}_3$ ,  $0.12 \text{ ppm O}_3$  and  $0.30 \text{ ppm NO}_2$  for 90 minutes via a mouthpiece.

In healthy volunteers, NOAELs for effects on pulmonary function and inflammatory response was reported after exposure to  $0.5 \text{ mg/m}^3 \text{ HNO}_3$  for 4 hours, the only level tested, and after a similar exposure for 2 hours at 0.4 mg/m<sup>3</sup>. Following a 2-hour exposure to  $0.2 \text{ mg/m}^3 \text{ HNO}_3$ , defence functions of alveolar macrophages were affected (increased phagocytic activity and resistance to infection, decreased superoxide anion production) (25).

Like for  $H_2SO_4$ , the lowest reported effect levels after short- or long-term exposure in animals relate to pulmonary defence mechanisms. At 0.05-0.25 mg/m<sup>3</sup>, reduced level/production of superoxide anion and TNF $\alpha$  activity in alveolar macrophages, increased elastase inhibitory capacity of lung lavage fluid, reduced bronchial responsivity to smooth muscle constrictor challenge and elevated stress-inducible heat shock protein 70 were observed. Heat-shock proteins are highly active within the immune system (168) but the relevance of this latter finding is unclear.

Irritation of the airways seen as bronchoconstriction in sheep exposed to  $4.1 \text{ mg/m}^3$  for 4 hours is reported.

In an old study based on only 2 individuals, exposure to  $27-30 \text{ mg/m}^3$  for more than 1 hour was considered intolerable and dangerous to human health.

### Phosphoric acid

No relevant studies were identified. Severe effects including death have been reported after excessive exposure.

# 14.2 Groups at extra risk

Data indicate that asthmatics, especially adolescent asthmatics are more susceptible to respiratory effects from  $H_2SO_4$  exposure.

Incidents with extremely polluted air containing H<sub>2</sub>SO<sub>4</sub> aerosols have caused increased mortality, e.g. among persons with chronic respiratory tract disease.

Subjects with chronic bronchitis and smokers may be at extra risk, but little is known about their susceptibility to the four acids.

# 14.3 Scientific basis for an occupational exposure limit

#### Sulphuric acid

The toxicological databases from short- and long-term animal and human exposures are quite consistent although a clear-cut overall NOAEL is difficult to identify from the available data.

Based on short- and long-term exposure data in both animals and humans, the critical effects are effects on bronchial mucociliary clearance, pulmonary function, and airway and eye irritation. The effects appear at approximately  $0.1 \text{ mg/m}^3$ . The impairment of pulmonary function at this level has been observed in adolescent asthmatics. At slightly higher levels, dental erosion, and pathological changes in the nasal mucosa have been reported in humans. In animals, cellular changes of the respiratory tract epithelium have occurred in rats, rabbits and monkeys at repeated exposure to concentrations in the range  $0.125-0.38 \text{ mg/m}^3$ . An excess risk of laryngeal cancer has been found among workers exposed to strong inorganic acid mists containing  $H_2SO_4$ . The development of laryngeal cancer from

 $H_2SO_4$  exposure is likely to have a threshold and thought to be secondary to local irritation and damage of the respiratory epithelium. Damage to the respiratory tract epithelium resulting in cancer development is unlikely at exposure levels below those affecting mucociliary clearance.

The reliable measurement of exposure around the lowest effect levels is nonproblematic for 8-hour samplings when only  $H_2SO_4$  aerosol is present, but challenging when sampling times are short.

# Hydrochloric acid

The critical effect is airway irritation. Initial mild irritation of the airway mucosa, which regressed rapidly, was reported at 5 mg/m<sup>3</sup> (LOAEL) in workers. No airway irritation was observed at 2.5 mg/m<sup>3</sup> (NOAEL) in asthmatics exposed 45 minutes.

Tracheal and laryngeal hyperplasia observed after chronic exposure of rats at 14 mg/m<sup>3</sup> is regarded secondary to airway irritation in analogy with  $H_2SO_4$ .

A higher prevalence of dental erosion among battery and galvanising workers exposed to  $H_2SO_4$  and HCl has been found but the HCl level at which it appears is not known.

### Nitric acid

There is a general lack of data that could serve as the scientific basis for an occupational exposure limit for HNO<sub>3</sub>.

In healthy volunteers, a NOAEL for effects on pulmonary function and inflammatory response was reported after exposure to 0.5 mg/m<sup>3</sup> HNO<sub>3</sub> for 4 hours, the only level tested. Following a 2-hour exposure to 0.2 mg/m<sup>3</sup>, results indicating a stimulatory as well as an inhibitory effect on the defence functions of alveolar macrophages were obtained.

Based on animal acute toxicity data, the potency of  $HNO_3$  seems to be similar to that of  $H_2SO_4$ .

### Phosphoric acid

No relevant studies that could serve as the basis for an occupational exposure limit have been identified.

The assessment must be based on the conclusion made by the European commission in 1992 that fumes of phosphorous pentoxide at concentrations  $0.8-5.4 \text{ mg/m}^3$  are unlikely to produce significant irritation. It was also noted that phosphorous pentoxide is a powerful dehydrating agent combining with atmospheric moisture or in the respiratory tract to produce H<sub>3</sub>PO<sub>4</sub> in a reaction generating heat and is likely to cause more tissue damage than H<sub>3</sub>PO<sub>4</sub> itself. Thus, applying the NOAEL for irritation for phosphorous pentoxide would supply an adequate margin of safety.

# 15. Research needs

- Repeated inhalation exposure studies on HCl, HNO<sub>3</sub> and H<sub>3</sub>PO<sub>4</sub> in animals and human volunteers focusing on pulmonary function, sensory irritation, respiratory effects and, in addition, animal studies on cytotoxicity and laryngeal cancer.
- Epidemiological studies on dental etching and erosion, pulmonary effects and laryngeal cancer with better exposure data, or the assessment of past exposure by developing job-exposure matrices in relevant industries.
- Studies of the mechanism behind the carcinogenic effect of strong inorganic mists containing H<sub>2</sub>SO<sub>4</sub>.

## 16. Summary

van der Hagen M, Järnberg J. *The Nordic Expert Group for Criteria* Documentation of Health Risks from Chemicals. 140. Sulphuric, hydrochloric, nitric and phosphoric acids. Arbete och Hälsa 2009;43(7):1-122.

Sulphuric, hydrochloric, nitric and phosphoric acids are common inorganic or mineral acids. The first three acids are strong, whereas phosphoric acid is weaker. They are all important industrial chemicals used in a variety of applications, e.g. in the manufacture of chemicals and metal or food products. The relatively nonvolatile sulphuric and phosphoric acids will occur in air primarily as aerosols and the more volatile hydrochloric and nitric acids as vapours or aerosols. Following absorption, the toxic effects of the acids will be mainly from protolysis yielding protons in the mucosa. The reaction between sulphuric acid and water generates heat.

Except for sulphuric acid, the toxicological database is poor or very poor.

The acids are corrosive and will cause chemical burns when in contact with eyes, skin and mucous membranes. Acid vapours and aerosols are respiratory tract irritants and may cause pulmonary impairment, as well as dental erosion, and laryngeal cancer.

Sulphuric acid: The critical effects are alterations in bronchial mucociliary clearance, lung function effects and airway and eye irritation. The effects begin to appear at approximately  $0.1 \text{ mg/m}^3$  in humans. At slightly higher levels, dental erosion and pathological changes of the nasal mucosa have been reported. Cellular changes of the respiratory tract epithelium have been observed in animals after repeated exposures to concentrations in the range  $0.125-0.38 \text{ mg/m}^3$ . An excess risk of laryngeal cancer has been found among workers exposed to strong inorganic acid mists containing H<sub>2</sub>SO<sub>4</sub>. The mechanism of laryngeal cancer from acid mist exposure seems to be secondary to the local airway irritation caused by the acid.

*Hydrochloric acid:* The critical effect is airway irritation. No airway irritation at 2.5 mg/m<sup>3</sup> was reported in asthmatics but mild irritation, which regressed rapidly, at 5 mg/m<sup>3</sup> in workers. Tracheal and laryngeal hyperplasia observed in animals after chronic exposure to 14 mg/m<sup>3</sup> is regarded secondary to airway irritation in analogy with sulphuric acid.

*Nitric acid:* There is a general lack of data. In healthy volunteers, no effects on pulmonary function and inflammatory response were noted after a single exposure to  $0.5 \text{ mg/m}^3$  but defence functions of alveolar macrophages were affected at  $0.2 \text{ mg/m}^3$ . The potency of nitric acid seems to be similar to that of sulphuric acid.

*Phosphoric acid:* As data are lacking, the assessment has to be based on comparison with the stronger irritant phosphorous pentoxide, which is converted to the acid in the airways.

*Keywords:* aerosol, hydrochloric acid, hyperplasia, irritation, laryngeal cancer, nitric acid, occupational exposure limit, phosphoric acid, respiratory tract, review, risk assessment, sulphuric acid, toxicity

## 17. Summary in Norwegian

van der Hagen M, Järnberg J. *The Nordic Expert Group for Criteria* Documentation of Health Risks from Chemicals. 140. Sulphuric, hydrochloric, nitric and phosphoric acids. Arbete och Hälsa 2009;43(7):1-122.

Svovelsyre, saltsyre, salpetersyre og fosforsyre er vanlige uorganiske syrer (mineralsyrer). De er alle viktige industrikjemikalier i bruk til ulike formål som produksjon av industrikjemikalier, metaller og matvarer. Alle unntatt fosforsyre er sterke syrer. Svovelsyre og fosforsyre er mindre flyktige og finnes primært som aerosoler. Saltsyre og salpetersyre er mer flyktige og kan forekomme som damp eller aerosol. Etter at stoffene er absorbert er den toksiske virkningen knyttet til proteolyse og frigjøring av protoner i slimhinnen. Reaksjonen mellom svovelsyre og vann frigjør varme.

Databasen har tilstrekkelig informasjon om svovelsyre, mens det er lite informasjon om de andre syrene. De er etsende og kan føre til kjemiske brannskader når de kommer i kontakt med øyne, slimhinner og hud. Syredamper og aerosoler kan føre til irritasjon i luftveiene og nedsatt lungefunksjon. Det er også dokumentert økt risiko for tannerosjon og strupekreft.

*Svovelsyre:* Kritiske effekter er endringer i slimheisfunksjonen, lungefunksjon, samt irritasjon i luftveier og øyne. Disse fremkommer ved eksponeringer omkring 0,1 mg/m<sup>3</sup>. Ved litt høyere konsentrasjoner oppstår tannerosjoner og vevsforandringer i neseslimhinnen, Celleskade i respiratorisk epitel oppstår hos dyr eksponert for 0,125-0,38 mg/m<sup>3</sup>. Det er påvist økt risiko for kreft i strupen blant arbeidere eksponert for sterk uorganiske tåke som inneholder H<sub>2</sub>SO<sub>4</sub>. Mekanismen for kreftutviklingen synes å være sekundær til luftveisirritasjonen forårsaket av syren.

*Saltsyre:* Kritisk effekt er luftveisirritasjon, rapportert som mild og raskt forbigående hos arbeidstakere ved en konsentrasjon på 5 mg/m<sup>3</sup>. Ingen irritasjon ble vist hos astmapasienter ved 2,5 mg/m<sup>3</sup>. Fortykkelse av slimhinnen i hovedluftrør og strupe ble påvist ved kronisk eksponering for 14 mg/m<sup>3</sup>, bedømt som sekundært til luftveisirritasjon.

*Salpetersyre:* Det er få data i litteraturen. Hos en gruppe friske frivillige ble det ikke påvist utfall i lungefunksjon eller betennelsesrespons etter en engangseksponering på 0,5 mg/m<sup>3</sup>. Det ble også påvist endringer i alveolare makrofagers forsvarsevne ved 0,2 mg/m<sup>3</sup>. Evnen til å forårsake effekt synes sammenliknbar med svovelsyre.

*Fosforsyre:* Det finnes ikke data som gir grunnlag for å vurdere kritisk effect. I dette dokumentet er det foreslått å vurdere den i forhold til den sterkere irritanten fosforpentoksid som omdannes til syre i luftveiene.

*Nøkkelord:* administrative norm, aerosol, fosforsyre, hyperplasi, irritasjon, luftveier, review, risikovurdering, salpetersyre, saltsyre, strupekreft, svovelsyre, toksisitet

## 18. References

- Aas W, Solberg S, Berg T, Manö S, Yttri KE. Overvåkning av langtransportert forurenset luft og nedbör. Atmosfærisk tilförsel, 2005. NILU 955/2006. Norway, Oslo: Statens forurensningstilsyn, Norskt institutt for luftforskning: Statlig program for forurensningsovervåkning, 2006 (in Norwegian).
- 2. Abraham WM, Kim CS, King MM, Oliver W, Jr., Yerger L. Effects of nitric acid on carbachol reactivity of the airways in normal and allergic sheep. *Arch Environ Health* 1982;37:36-40.
- ACGIH. Nitric acid. Documentation of the threshold limit values and biological exposure indices. 7th ed. Cincinnati, Ohio: American Conference of Governmental Industrial Hygienists, 2001.
- 4. ACGIH. Phosphoric acid. *Documentation of the threshold limit values and biological exposure indices.* 7th ed. Cincinnati, Ohio: American Conference of Governmental Industrial Hygienists, 2001.
- 5. ACGIH. Hydrogen chloride. *Documentation of the threshold limit values and biological exposure indices.* 7th ed. Cincinnati, Ohio: American Conference of Governmental Industrial Hygienists, 2003.
- 6. ACGIH. Sulfuric acid. *Documentation of the threshold limit values and biological exposure indices.* 7th ed. Cincinnati, Ohio: American Conference of Governmental Industrial Hygienists, 2004.
- 7. Ahlborg G, Hogstedt C, Sundell L, Åman C. Laryngeal cancer and pickling house vapors [Letter]. *Scand J Work Environ Health* 1981;7:239-240.
- 8. Alarie Y, Busey WM, Krumm AA, Ulrich CE. Long-term continuous exposure to sulfuric acid mist in cynomolgus monkeys and guinea pigs. *Arch Environ Health* 1973;27:16-24.
- Alarie YC, Krumm AA, Busey WM, Urich CE, Kantz RJ. Long-term exposure to sulfur dioxide, sulfuric acid mist, fly ash, and their mixtures. Results of studies in monkeys and guinea pigs. *Arch Environ Health* 1975;30:254-262.
- Albert RE, Sellakumar AR, Laskin S, Kuschner M, Nelson N, Snyder CA. Gaseous formaldehyde and hydrogen chloride induction of nasal cancer in the rat. *J Natl Cancer Inst* 1982;68:597-603.
- 11. Alderson MR, Rattan NS. Mortality of workers on an isopropyl alcohol plant and two MEK dewaxing plants. *Br J Ind Med* 1980;37:85-89.
- 12. Aldrich Chemical Co., Inc. *Aldrich Catalog/Handbook of fine chemicals (1996-1997)*. Milwaukee, Wisconsin: Aldrich Chemical Co., 1996.
- 13. Amdur MO, Dubriel M, Creasia DA. Respiratory response of guinea pigs to low levels of sulfuric acid. *Environ Res* 1978;15:418-423.
- Anderson KR, Avol EL, Edwards SA, Shamoo DA, Peng RC, Linn WS, Hackney JD. Controlled exposures of volunteers to respirable carbon and sulfuric acid aerosols. *J Air Waste Manage Assoc* 1992;42:770-776.
- 15. Aris R, Christian D, Sheppard D, Balmes JR. Lack of bronchoconstrictor response to sulfuric acid aerosols and fogs. *Am Rev Respir Dis* 1991;143:744-750.
- 16. Aris R, Christian D, Sheppard D, Balmes JR. The effects of sequential exposure to acidic fog and ozone on pulmonary function in exercising subjects. *Am Rev Respir Dis* 1991;143:85-91.
- Aris R, Christian D, Tager I, Ngo L, Finkbeiner WE, Balmes JR. Effects of nitric acid gas alone or in combination with ozone on healthy volunteers. *Am Rev Respir Dis* 1993;148:965-973.
- 18. Arts JH, de Heer C, Woutersen RA. Local effects in the respiratory tract: relevance of subjectively measured irritation for setting occupational exposure limits. *Int Arch Occup Environ Health* 2006;79:283-298.

- ATSDR. *Toxicological profile for sulfur trioxide and sulfuric acid.* Atlanta, Georgia: U.S. Department of Health and Human Services, Public Health Service, Agency for Toxic Substances and Disease Registry, 1998.
- Avol EL, Jones MP, Bailey RM, Chang NM, Kleinman MT, Linn WS, Bell KA, Hackney JD. Controlled exposures of human volunteers to sulfate aerosols. Health effects and aerosol characterization. *Am Rev Respir Dis* 1979;120:319-327.
- Avol EL, Linn WS, Whynot JD, Anderson KR, Shamoo DA, Valencia LM, Little DE, Hackney JD. Respiratory dose-response study of normal and asthmatic volunteers exposed to sulfuric acid aerosol in the sub-micrometer size range. *Toxicol Ind Health* 1988;4:173-184.
- 22. Avol EL, Linn WS, Wightman LH, Whynot JD, Anderson KR, Hackney JD. Short-term respiratory effects of sulfuric acid in fog: a laboratory study of healthy and asthmatic volunteers. *JAPCA* 1988;38:258-263.
- Barrow CS, Alarie Y, Warrick JC, Stock MF. Comparison of the sensory irritation response in mice to chlorine and hydrogen chloride. *Arch Environ Health* 1977;32:68-76.
- Beaumont JJ, Leveton J, Knox K, Bloom T, McQuiston T, Young M, Goldsmith R, Steenland NK, Brown DP, Halperin WE. Lung cancer mortality in workers exposed to sulfuric acid mist and other acid mists. J Natl Cancer Inst 1987;79:911-921.
- Becker S, Roger LJ, Devlin RB, Horstman DH, Koren HS. Exposure to nitric acid stimulates human alveolar macrophage function but does not cause inflammation or changes in lung function. *Inhal Toxicol* 1996;8:185-200.
- 26. Benomran FA, Hassan AI, Masood SS. Accidental fatal inhalation of sulfuric acid fumes. *J Forensic Leg Med* 2008;15:56-58.
- Blair A, Kazerouni N. Reactive chemicals and cancer. *Cancer Causes Control* 1997;8:473-490.
- Block G, Matanoski GM, Seltser R, Mitchell T. Cancer morbidity and mortality in phosphate workers. *Cancer Res* 1988;48:7298-7303.
- 29. Bond GG, Cook RR, Wight PC, Flores GH. A case-control study of brain tumor mortality at a Texas Chemical plant. *J Occup Med* 1983;25:377-386.
- Bond GG, Shellenberger RJ, Flores GH, Cook RR, Fishbeck WA. A case-control study of renal cancer mortality at a Texas chemical plant. *Am J Ind Med* 1985;7:123-139.
- 31. Bond GG, Flores GH, Shellenberger RJ, Cartmill JB, Fishbeck WA, Cook RR. Nested casecontrol study of lung cancer among chemical workers. *Am J Epidemiol* 1986;124:53-66.
- 32. Bond GG, Flores GH, Stafford BA, Olsen GW. Lung cancer and hydrogen chloride exposure: results from a nested case-control study of chemical workers. *J Occup Med* 1991;33:958-961.
- Boulet LP. Increases in airway responsiveness following acute exposure to respiratory irritants. Reactive airway dysfunction syndrome or occupational asthma? *Chest* 1988;94:476-481.
- 34. Boutoux M, Bernard R, Leroyer C, Dewitte JD. Reactive airways dysfunction syndrome après exposition aux vapeurs d'acide phosphorique. *Arch Mal Prof* 1995;56:45-47.
- 35. Boyce SH, Simpson KA. Hydrochloric acid inhalation: who needs admission? *J Accid Emerg Med* 1996;13:422-424.
- Branday J, Arscott GD, Smoot EC, Williams GD, Fletcher PR. Chemical burns as assault injuries in Jamaica. *Burns* 1996;22:154-155.
- 37. Breuer D, Howe A. Performance of methods for measurement of exposure to inorganic acids in workplace air. *J Environ Monit* 2006;8:120-126.
- Brown LM, Mason TJ, Pickle LW, Stewart PA, Buffler PA, Burau K, Ziegler RG, Fraumeni JF, Jr. Occupational risk factors for laryngeal cancer on the Texas Gulf Coast. *Cancer Res* 1988;48:1960-1964.
- 39. Brusick D. Genotoxic effects in cultured mammalian cells produced by low pH treatment conditions and increased ion concentrations. *Environ Mutagen* 1986;8:879-886.

- 40. Bråtveit M, Haaland IM, Moen BE, Målsnes A. Exposure to sulfuric acid in zinc production. Ann Occup Hyg 2004;48:159-170.
- Buckley LA, Jiang XZ, James RA, Morgan KT, Barrow CS. Respiratory tract lesions induced by sensory irritants at the RD50 concentration. *Toxicol Appl Pharmacol* 1984;74:417-429.
- Burleigh-Flayer H, Wong KL, Alarie Y. Evaluation of the pulmonary effects of HCl using CO<sub>2</sub> challenges in guinea pigs. *Fundam Appl Toxicol* 1985;5:978-985.
- 43. Caravati EM. Metabolic abnormalities associated with phosphoric acid ingestion. *Ann Emerg Med* 1987;16:904-906.
- Checkoway H, Mathew RM, Hickey JL, Shy CM, Harris RL, Jr., Hunt EW, Waldman GT. Mortality among workers in the Florida phosphate industry. I. Industry-wide cause-specific mortality patterns. J Occup Med 1985;27:885-892.
- 45. Checkoway H, Mathew RM, Hickey JL, Shy CM, Harris RL, Jr., Hunt EW, Waldman GT. Mortality among workers in the Florida phosphate industry. II. Cause-specific mortality relationships with work areas and exposures. *J Occup Med* 1985;27:893-896.
- 46. Chen LC, Miller PD, Lam HF, Guty J, Amdur MO. Sulfuric acid-layered ultrafine particles potentiate ozone-induced airway injury. *J Toxicol Environ Health* 1991;34:337-352.
- Chen LC, Fine JM, Qu QS, Amdur MO, Gordon T. Effects of fine and ultrafine sulfuric acid aerosols in guinea pigs: alterations in alveolar macrophage function and intracellular pH. *Toxicol Appl Pharmacol* 1992;113:109-117.
- 48. Chen LC, Miller PD, Amdur MO, Gordon T. Airway hyperresponsiveness in guinea pigs exposed to acid-coated ultrafine particles. *J Toxicol Environ Health* 1992;35:165-174.
- 49. Chen LC, Qu Q, Amdur MO, Schlesinger RB. Alteration of pulmonary macrophage intracellular pH following inhalation exposure to sulfuric acid/ozone mixtures. *Exp Lung Res* 1995;21:113-128.
- 50. Chen LC, Schlesinger RB. Considerations for the respiratory tract dosimetry of inhaled nitric acid vapor. *Inhal Toxicol* 1996;8:639-654.
- 51. Chikte UM, Josie-Perez AM. Industrial dental erosion: a cross-sectional, comparative study. *SADJ* 1999;54:531-536.
- 52. Christensen TG, Lucey EC, Breuer R, Snider GL. Acid-induced secretory cell metaplasia in hamster bronchi. *Environ Res* 1988;45:78-90.
- Cipollaro M, Corsale G, Esposito A, Ragucci E, Staiano N, Giordano GG, Pagano G. Sublethal pH decrease may cause genetic damage to eukaryotic cell: a study on sea urchins and Salmonella typhimurium. *Teratog Carcinog Mutagen* 1986;6:275-287.
- 54. Coalson JJ, Collins JF. Nitric acid-induced injury in the hamster lung. *Br J Exp Pathol* 1985;66:205-215.
- 55. Cocco P, Ward MH, Dosemeci M. Occupational risk factors for cancer of the gastric cardia. Analysis of death certificates from 24 US states. *J Occup Environ Med* 1998;40:855-861.
- 56. Cocco P, Ward MH, Dosemeci M. Risk of stomach cancer associated with 12 workplace hazards: analysis of death certificates from 24 states of the United States with the aid of job exposure matrices. *Occup Environ Med* 1999;56:781-787.
- 57. Coggon D, Pannett B, Wield G. Upper aerodigestive cancer in battery manufacturers and steel workers exposed to mineral acid mists. *Occup Environ Med* 1996;53:445-449.
- 58. Collins JJ, Swaen GM, Marsh GM, Utidjian HM, Caporossi JC, Lucas LJ. Mortality patterns among workers exposed to acrylamide. *J Occup Med* 1989;31:614-617.
- 59. Columbia University. *The Columbia Encyclopedia*. 6 th ed. Columbia, New York: Columbia University Press, 2004.
- 60. Cooper WC, Wong O, Kheifets L. Mortality among employees of lead battery plants and lead-producing plants, 1947-1980. *Scand J Work Environ Health* 1985;11:331-345.

- 61. Culp DJ, Latchney LR, Frampton MW, Jahnke MR, Morrow PE, Utell MJ. Composition of human airway mucins and effects after inhalation of acid aerosol. *Am J Physiol* 1995;269:L358-370.
- 62. Dalton P. Evaluating the human response to sensory irritation: implications for setting occupational exposure limits. *AIHAJ* 2001;62:723-729.
- 63. Darke CS, Warrack AJ. Bronchiolitis from nitrous fumes. *Thorax* 1958;13:327-333.
- 64. Darmer KI, Jr, Kinkead ER, DiPasquale LC. Acute toxicity in rats and mice exposed to hydrogen chloride gas and aerosols. *Am Ind Hyg Assoc J* 1974;35:623-631.
- 65. DFG. Hydrogen chloride. In: Greim H ed. Occupational toxicants: critical data evaluation for MAK values and classification of carcinogens. Vol VI. Deutsche Forschungsgemeinschaft. Weinheim: Wiley-VCH: 1982:231-238.
- DFG. Nitric acid. In: Greim H ed. Occupational toxicants: critical data evaluation for MAK values and classification of carcinogens. Vol 3. Deutsche Forschungsgemeinschaft. Weinheim: Wiley-VCH: 1992:233-240.
- DFG. Sulfuric acid. In: Greim H, ed. Occupational toxicants: critical data evaluation for MAK values and classification of carcinogens. Vol 15. Deutsche Forschungsgemeinschaft. Weinheim: Wiley-VCH: 2001:165-222.
- DFG. Hydrogen chloride. In: *The MAK-Collection Part 1: MAK value documentation*. Vol 24. Deutsche Forschungsgemeinschaft. Weinheim: Wiley-VCH: 2007:133-147.
- 69. Dilawari JB, Singh S, Rao PN, Anand BS. Corrosive acid ingestion in man a clinical and endoscopic study. *Gut* 1984;25:183-187.
- El-Fawal HA, Schlesinger RB. Nonspecific airway hyperresponsiveness induced by inhalation exposure to sulfuric acid aerosol: an in vitro assessment. *Toxicol Appl Pharmacol* 1994;125:70-76.
- 71. El-Sadik YM, Osman HA, el-Gazzar RM. Exposure to sulfuric acid in manufacture of storage batteries. *J Occup Med* 1972;14:224-226.
- 72. Englander V, Sjöberg A, Hagmar L, Attewell R, Schutz A, Möller T, Skerfving S. Mortality and cancer morbidity in workers exposed to sulphur dioxide in a sulphuric acid plant. *Int Arch Occup Environ Health* 1988;61:157-162.
- 73. Enterline PE. Importance of sequential exposure in the production of epichlorohydrin and isopropanol. *Ann N Y Acad Sci* 1982;381:344-349.
- 74. European Commission. Occupational exposure limits: Criteria document for phosphoric acid. Health and safety. EUR 14178 EN. Commission of the European Communities, Directorate-General, Telecommunications, Information Industries and Innovation. Luxemburg: Office for Official Publications of the European Communities, 1992.
- European Commission. Occupational exposure limits: Criteria document for nitric acid. EUR 16668 EN. Directorate-General for employment, industrial relations and social affairs. Luxemburg: Office for Official Publications of the European Communities, 1996.
- European Commission. *ClassLab Database*. Joint Research Centre. Institute for Health and Consumer Protection. http://ecb.jrc.ec.europa.eu/classification-labelling/search-classlab/. Accessed October 2008.
- 77. European Committee for Standardization (CEN). *Workplace atmospheres Size fraction definition for measurement of airborne particles.* EN 481:1993.
- Faraj Z, Khattabi A, Soulaymani R. [Reactive airways dysfunction syndrome (Brooks syndrome)]. *Presse Med* 2002;31:1410-1413.
- 79. Flammiger A, Maibach H. Sulfuric acid burns (corrosion and acute irritation): evidence-based overview to management. *Cutan Ocul Toxicol* 2006;25:55-61.
- 80. Forastiere F, Valesini S, Salimei E, Magliola ME, Perucci CA. Respiratory cancer among soap production workers. *Scand J Work Environ Health* 1987;13:258-260.

- Foster G, Murdoch C, Apthorpe L, Mandryk J. Sulphuric acid mist: Exposures, controls and respiratory symptoms. Perth, WA: Australian Institute of Occupational Health, 1996:171-177 (conference proceedings).
- Frampton MW, Voter KZ, Morrow PE, Roberts NJ, Jr., Culp DJ, Cox C, Utell MJ. Sulfuric acid aerosol exposure in humans assessed by bronchoalveolar lavage. *Am Rev Respir Dis* 1992;146:626-632.
- 83. Frampton MW, Morrow PE, Cox C, Levy PC, Condemi JJ, Speers D, Gibb FR, Utell MJ. Sulfuric acid aerosol followed by ozone exposure in healthy and asthmatic subjects. *Environ Res* 1995;69:1-14.
- Fujimaki H, Katayama N, Wakamori K. Enhanced histamine release from lung mast cells of guinea pigs exposed to sulfuric acid aerosols. *Environ Res* 1992;58:117-123.
- Fujita M, Schroeder MA, Hyatt RE. Canine model of chronic bronchial injury. Lung mechanics and pathologic changes. *Am Rev Respir Dis* 1988;137:429-434.
- Gamble J, Jones W, Hancock J. Epidemiological-environmental study of lead acid battery workers. II. Acute effects of sulfuric acid on the respiratory system. *Environ Res* 1984;35:11-29.
- Gamble J, Jones W, Hancock J, Meckstroth RL. Epidemiological-environmental study of lead acid battery workers. III. Chronic effects of sulfuric acid on the respiratory system and teeth. *Environ Res* 1984;35:30-52.
- 88. Gearhart JM, Schlesinger RB. Sulfuric acid-induced airway hyperresponsiveness. *Fundam Appl Toxicol* 1986;7:681-689.
- 89. Gearhart JM, Schlesinger RB. Response of the tracheobronchial mucociliary clearance system to repeated irritant exposure: effect of sulfuric acid mist on function and structure. *Exp Lung Res* 1988;14:587-605.
- 90. Gearhart JM, Schlesinger RB. Sulfuric acid-induced changes in the physiology and structure of the tracheobronchial airways. *Environ Health Perspect* 1989;79:127-136.
- 91. Ghiassi-Nejad M, Varzegar R, Zakeri F, Rasouli-Nejad S. Analysis of chromosomal aberrations, micronuclei, and sister chromatid exchanges in lymphocytes of workers of a phosphate fertilizer factory. *Central European Journal of Occupational and Environmental Medicine* 2002;8:277-282.
- Giuriati C, Cristofori MC, Gorni A, Abballe F. Ion chromatography applications in the determination of HF, HCl, NOx, SOx on stationary emissions. *Ann Ist Super Sanita* 2003;39:223-228.
- 93. Goldman A, Hill WT. Chronic bronchopulmonary disease due to inhalation of sulfuric acid fumes. A M A Arch Ind Hyg Occup Med 1953;8:205-211.
- 94. Granata M, Ammendolea C, Nicoletti M, Martelletti P, Giacovazzo M. Headache induced by accidental nitric acid inhalation. *Cephalalgia* 2004;24:238.
- Grasel SS, Alves VA, da Silva CS, Cruz OL, Almeida ER, de Oliveira E. Clinical and histopathological changes of the nasal mucosa induced by occupational exposure to sulphuric acid mists. *Occup Environ Med* 2003;60:395-402.
- 96. Greenwood NN, Earnshaw A. Chemistry of the elements. Oxford: Pergamon Press Ltd, 1986.
- Grimsrud TK, Berge SR, Haldorsen T, Andersen A. Can lung cancer risk among nickel refinery workers be explained by occupational exposures other than nickel? *Epidemiology* 2005;16:146-154.
- Grose EC, Gardner DE, Miller FJ. Response of ciliated epithelium to ozone and sulfuric acid. *Environ Res* 1980;22:377-385.
- 99. Hackney JD, Linn WS, Avol EL. Acid fog: effects on respiratory function and symptoms in healthy and asthmatic volunteers. *Environ Health Perspect* 1989;79:159-162.
- Hagmar L, Bellander T, Andersson C, Linden K, Attewell R, Moller T. Cancer morbidity in nitrate fertilizer workers. *Int Arch Occup Environ Health* 1991;63:63-67.

- 101. Hajela R, Janigan DT, Landrigan PL, Boudreau SF, Sebastian S. Fatal pulmonary edema due to nitric acid fume inhalation in three pulp-mill workers. *Chest* 1990;97:487-489.
- 102. Hall JN, Cooper CE. The effects of the inhalation of the fumes of nitric acid. *JAMA* 1905;5:396-398.
- 103. Hanley QS, Koenig JQ, Larson TV, Anderson TL, van Belle G, Rebolledo V, Covert DS, Pierson WE. Response of young asthmatic patients to inhaled sulfuric acid. *Am Rev Respir Dis* 1992;145:326-331.
- 104. Hathaway JA. Upper aerodigestive cancer in battery manufacturers and steel workers exposed to mineral acid mists. *Occup Environ Med* 1997;54:141-142.
- 105. Hawkins DB, Demeter MJ, Barnett TE. Caustic ingestion: controversies in management. A review of 214 cases. *Laryngoscope* 1980;90:98-109.
- 106. Health Council of the Netherlands. *Strong inorganic acid mists containing sulphuric acid; Evaluation of the carcinogenicity and genotoxicity.* The Hague: Health Council of the Netherlands, 2003; publication no. 2003/07OSH.
- 107. Heidelberger C, Freeman AE, Pienta RJ, Sivak A, Bertram JS, Casto BC, Dunkel VC, Francis MW, Kakunaga T, Little JB, Schechtman LM. Cell transformation by chemical agents--a review and analysis of the literature. A report of the U.S. Environmental Protection Agency Gene-Tox Program. *Mutat Res* 1983;114:283-385.
- 108. Hery M, Hecht G, Gerber JM, Bemer D, Gorner P. Exposure to sulphuric acid and sulphur dioxide in the manufacturing of titanium dioxide. *Ann Occup Hyg* 1992;36:653-661.
- 109. Hilt B, Langard S, Andersen A, Rosenberg J. Asbestos exposure, smoking habits, and cancer incidence among production and maintenance workers in an electrochemical plant. *Am J Ind Med* 1985;8:565-577.
- 110. Hinkamp DL. Acids, Inorganic. In: Stellman JM, ed. *Encyclopaedia of occupational health and safety*. 4th ed. Vol IV. Geneva: International Labour Office, 1998:104.5-104.7.
- 111. Ho CK, Lo WC, Huang PH, Wu MT, Christiani DC, Lin CT. Suspected nasopharyngeal carcinoma in three workers with long-term exposure to sulphuric acid vapour. Occup Environ Med 1999;56:426-428.
- 112. Houghton DJ, White PS. The carcinogenic risk of exposure to sulphuric acid fumes from lead acid batteries. *J Laryngol Otol* 1994;108:881-882.
- 113. Hsu YM, Wu CY, Lundgren DA, Birky BK. Size-resolved sulfuric acid mist concentrations at phosphate fertilizer manufacturing facilities in Florida. *Ann Occup Hyg* 2007;51:81-89.
- Hsu YM, Wu CY, Lundgren DA, Nall JW, Birky BK. Chemical characteristics of aerosol mists in phosphate fertilizer manufacturing facilities. J Occup Environ Hyg 2007;4:17-25.
- 115. IARC. Hydrochloric acid. In: Occupational exposures to mists and vapours from strong inorganic acids; and other industrial chemicals. IARC monographs on the evaluation of carcinogenic risks to humans. Vol 54, pp 189-211. Lyon: International Agency for Research on Cancer, World Health Organization, 1992.
- 116. IARC. Occupational exposures to mists and vapours from sulfuric acid and other strong inorganic acids. In: Occupational exposures to mists and vapours from strong inorganic acids; and other industrial chemicals. IARC monographs on the evaluation of carcinogenic risks to humans. Vol 54, pp 41-119. Lyon: International Agency for Research on Cancer, World Health Organization, 1992.
- 117. IPCS. *Environmental health criteria 21: Chlorine and hydrogen chlorine*. Geneva: International Programme on Chemical Safety, World Health Organization, 1982:1-95.
- 118. IPCS. *Environmental health criteria 188: Nitrogen oxides.* 2nd ed. Geneva: International Programme on Chemical Safety, World Health Organization, 1997:1-507.
- 119. IUPAC goldbook. *The IUPAC compendium of chemical terminology*. Electronic version: http://goldbook.iupac.org. Union of Pure and Applied Chemistry, 2007.
- 120. Jacobs GA. OECD eye irritation tests on two strong acids. J Am Coll Toxicol 1992;11:734.

- Jaeggi T, Lussi A. Prevalence, incidence and distribution of erosion. *Monogr Oral Sci* 2006;20:44-65.
- 122. Jones W, Gamble J. Epidemiological-environmental study of lead acid battery workers. I. Environmental study of five lead acid battery plants. *Environ Res* 1984;35:1-10.
- 123. Kaplan HL, Anzueto A, Switzer WG, Hinderer RK. Effects of hydrogen chloride on respiratory response and pulmonary function of the baboon. *J Toxicol Environ Health* 1988;23:473-493.
- 124. Kaplan HL, Switzer WG, Hinderer RK, Anzueto A. Studies of the effects of hydrogen chloride and polyvinylchloride (PVC) smoke in rodents. *Journal of Fire Sciences* 1993;11:512-552.
- 125. Kauppinen T, Toikkanen J, Pedersen D, Young R, Ahrens W, Boffetta P, Hansen J, Kromhout H, Maqueda Blasco J, Mirabelli D, de la Orden-Rivera V, Pannett B, Plato N, Savela A, Vincent R, Kogevinas M. Occupational exposure to carcinogens in the European Union. *Occup Environ Med* 2000;57:10-18.
- 126. Kilburn KH. Effects of a hydrochloric acid spill on neurobehavioral and pulmonary function. *J Occup Environ Med* 1996;38:1018-1025.
- 127. Kilgour JD, Foster J, Soames A, Farrar DG, Hext PM. Responses in the respiratory tract of rats following exposure to sulphuric acid aerosols for 5 or 28 days. *J Appl Toxicol* 2002;22:387-395.
- 128. Kimmel TA, Chen LC, Bosland MC, Nadziejko C. Influence of acid aerosol droplet size on structural changes in the rat lung caused by acute exposure to sulfuric acid and ozone. *Toxicol Appl Pharmacol* 1997;144:348-355.
- Kirk RE, Othmer, DF, Mark HF eds. *Encyclopedia of chemical technology. Vol 17.* 3rd ed. New York: John Wiley and Sons, 1982.
- 130. Kjaergaard SK, Hodgson M. The assessment of irritation using clinical methods and questionnaires. *AIHAJ* 2001;62:711-716.
- Kleinman MT, Phalen RF. Toxicological interactions in the respiratory system after inhalation of ozone and sulfuric acid aerosol mixtures. *Inhalation Toxicology* 2006;18:295-303.
- 132. Knapp MJ, Bunn WB, Stave GM. Adult respiratory distress syndrome from sulfuric acid fume inhalation. *South Med J* 1991;84:1031-1033.
- 133. Kobayashi T, Shinozaki Y. Effects of exposure to sulfuric acid-aerosol on airway responsiveness in guinea pigs: concentration and time dependency. *J Toxicol Environ Health* 1993;39:261-272.
- 134. Koenig JQ, Pierson WE, Horike M. The effects of inhaled sulfuric acid on pulmonary function in adolescent asthmatics. *Am Rev Respir Dis* 1983;128:221-225.
- Koenig JQ, Covert DS, Pierson WE. Effects of inhalation of acidic compounds on pulmonary function in allergic adolescent subjects. *Environ Health Perspect* 1989;79:173-178.
- 136. Koenig JQ, Covert DS, Larson TV, Pierson WE. The effect of duration of exposure on sulfuric acid-induced pulmonary function changes in asthmatic adolescent subjects: a dose-response study. *Toxicol Ind Health* 1992;8:285-296.
- 137. Koenig JQ, Covert DS, Pierson WE, Hanley QS, Rebolledo V, Dumler K, McKinney SE. Oxidant and acid aerosol exposure in healthy subjects and subjects with asthma. Part I: Effects of oxidants, combined with sulfuric or nitric acid, on the pulmonary function of adolescents with asthma. *Res Rep Health Eff Inst* 1994:1-36.
- 138. Korhonen K, Liukkonen T, Ahrens W, Astrakianakis G, Boffetta P, Burdorf A, Heederik D, Kauppinen T, Kogevinas M, Osvoll P, Rix BA, Saalo A, Sunyer J, Szadkowska-Stanczyk I, Teschke K, Westberg H, Widerkiewicz K. Occupational exposure to chemical agents in the paper industry. *Int Arch Occup Environ Health* 2004;77:451-460.

- 139. Kremer AM, Pal TM, Boleij JS, Schouten JP, Rijcken B. Airway hyper-responsiveness and the prevalence of work-related symptoms in workers exposed to irritants. *Am J Ind Med* 1994;26:655-669.
- 140. Kremer AM, Pal TM, Boleij JS, Schouten JP, Rijcken B. Airway hyperresponsiveness, prevalence of chronic respiratory symptoms, and lung function in workers exposed to irritants. *Occup Environ Med* 1994;51:3-13.
- 141. Kremer AM, Pal TM, Schouten JP, Rijcken B. Airway hyperresponsiveness in workers exposed to low levels of irritants. *Eur Respir J* 1995;8:53-61.
- 142. Kristensen P. Inorganic acid aerosols. In: *Criteria documents form the Nordic Expert Group* 1992. Solna: National Institute of Occupational Health, Arbete och Hälsa 1993;1:7-54.
- 143. Kroschwitz JI, Howe-Grant M eds. *Kirk-Othmer encyclopedia of chemical technology*. 4th ed. New York: John Wiley and Sons, 1995.
- 144. Krämer W, Bender HF, Leuppert G, Fischer P, Gusbeth K, Breuer D. Messung von Schwefelsäure in verschiedenen Arbeitsbereichen. *Gefahrstoffe - Reinhaltung der luft* 2002;62:45-51.
- 145. Kudoh I, Miyazaki H, Ohara M, Fukushima J, Tazawa T, Yamada H. Activation of alveolar macrophages in acid-injured lung in rats: different effects of pentoxifylline on tumor necrosis factor-alpha and nitric oxide production. *Crit Care Med* 2001;29:1621-1625.
- 146. Larson TV. The influence of chemical and physical forms of ambient air acids on airway doses. *Environ Health Perspect* 1989;79:7-13.
- 147. Laskin S. and Sellakumar A. Comparison of pulmonary carcinogenicity of known carcinogens with and without added H<sub>2</sub>SO<sub>4</sub> mists, airborne respirable particles, and gases. Final report of progress to the Environmental Protection Agency, 1978 (project no. 68-02-1750).
- 148. Last JA, Pinkerton KE. Chronic exposure of rats to ozone and sulfuric acid aerosol: biochemical and structural responses. *Toxicology* 1997;116:133-146.
- 149. Laube BL, Bowes SM, 3rd, Links JM, Thomas KK, Frank R. Acute exposure to acid fog. Effects on mucociliary clearance. *Am Rev Respir Dis* 1993;147:1105-1111.
- 150. Lee MM, Green FH, Roth SH, Karkhanis A, Bjarnason SG, Schurch S. Sulfuric acid aerosol induces changes in alveolar surface tension in the guinea pig but not in the rat. *Exp Lung Res* 1999;25:229-244.
- 151. Leikauf G, Yeates DB, Wales KA, Spektor D, Albert RE, Lippmann M. Effects of sulfuric acid aerosol on respiratory mechanics and mucociliary particle clearance in healthy nonsmoking adults. *Am Ind Hyg Assoc J* 1981;42:273-282.
- 152. Leikauf GD, Spektor DM, Albert RE, Lippmann M. Dose-dependent effects of submicrometer sulfuric acid aerosol on particle clearance from ciliated human lung airways. *Am Ind Hyg Assoc J* 1984;45:285-292.
- 153. Levin J-O ed. *Principer och metoder för provtagning och analys av ämnen på listan över hygieniska gränsvärden*. Stockholm: National Institute for Working Life, Arbete och Hälsa 2000;23:1-73 (in Swedish).
- 154. Lewis TR, Moorman WJ, Ludmann WF, Campbell KI. Toxicity of long-term exposure to oxides of sulfur. *Arch Environ Health* 1973;26:16-21.
- 155. Lidén G, Melin B, Lidblom A, Lindberg K, Norén JO. Personal sampling in parallel with open-face filter cassettes and IOM samplers for inhalable dust--implications for occupational exposure limits. *Appl Occup Environ Hyg* 2000;15:263-276.
- 156. Linn WS, Shamoo DA, Anderson KR, Peng RC, Avol EL, Hackney JD. Effects of prolonged, repeated exposure to ozone, sulfuric acid, and their combination in healthy and asthmatic volunteers. *Am J Respir Crit Care Med* 1994;150:431-440.
- 157. Lippmann M, Yeates DB, Albert RE. Deposition, retention, and clearance of inhaled particles. *Br J Ind Med* 1980;37:337-362.

- Lippmann M, Leikauf G, Spektor D, Schlesinger RB, Albert RE. The effects of irritant aerosols on mucus clearance from large and small conductive airways. *Chest* 1981;80:873-877.
- 159. Lippmann M, Schlesinger RB. Interspecies comparisons of particle deposition and mucociliary clearance in tracheobronchial airways. *J Toxicol Environ Health* 1984;13:441-469.
- Lippmann M, Gearhart JM, Schlesinger RB. Basis for a particle size-selective TLV for sulfuric acid aerosols. *Appl Ind Hyg* 1987;2:188-199.
- 161. Lussi A, Schaffner M, Hotz P, Suter P. Dental erosion in a population of Swiss adults. *Community Dent Oral Epidemiol* 1991;19:286-290.
- 162. Lynch J, Hanis NM, Bird MG, Murray KJ, Walsh JP. An association of upper respiratory cancer with exposure to diethyl sulfate. *J Occup Med* 1979;21:333-341.
- 163. Malcolm D, Paul E. Erosion of the teeth due to sulphuric acid in the battery industry. *Br J Ind Med* 1961;18:63-69.
- 164. Mannix RC, Phalen RF, Oldham MJ, Mautz WJ, Kleinman MT. Effects of repeated exposure to nitric acid vapor and ozone on respiratory tract clearance in the rat. *Inhalation Toxicology* 1996;8:595-605.
- 165. Mappes R. [MAC value of hydrochloric acid in the steel industry]. Zentralbl Arbeitsmed Arbeitsschutz Prophyl Ergonomie 1980;30:172-173.
- Marsh GM, Gula MJ, Youk AO, Cassidy LD. Bladder cancer among chemical workers exposed to nitrogen products and other substances. *Am J Ind Med* 2002;42:286-295.
- Mazumdar S, Lerer T, Redmond CK. Long-term mortality study of steelworkers. IX. Mortality patterns among sheet and tin mill workers. J Occup Med 1975;17:751-755.
- 168. Meyer NJ, Garcia JG. Wading into the genomic pool to unravel acute lung injury genetics. *Proc Am Thorac Soc* 2007;4:69-76.
- 169. Morita T, Watanabe Y, Takeda K, Okumura K. Effects of pH in the in vitro chromosomal aberration test. *Mutat Res* 1989;225:55-60.
- 170. Morita T, Nagaki T, Fukuda I, Okumura K. Effect of pH on the activity and stability of clastogens in the in vitro chromosomal aberration test with Chinese hamster ovary K1 cells. *Mutat Res* 1991;262:159-166.
- 171. Morita T, Nagaki T, Fukuda I, Okumura K. Clastogenicity of low pH to various cultured mammalian cells. *Mutat Res* 1992;268:297-305.
- 172. Morita T. Low pH leads to sister-chromatid exchanges and chromosomal aberrations, and its clastogenicity is S-dependent. *Mutat Res* 1995;334:301-308.
- 173. Morris PD, Koepsell TD, Daling JR, Taylor JW, Lyon JL, Swanson GM, Child M, Weiss NS. Toxic substance exposure and multiple myeloma: a case-control study. *J Natl Cancer Inst* 1986;76:987-994.
- 174. Murray FJ, Schwetz BA, Nitschke KD, Crawford AA, Quast JF, Staples RE. Embryotoxicity of inhaled sulfuric acid aerosol in mice and rabbits. *J Environ Sci Health C* 1979;13:251-266.
- 175. Nadziejko CE, Nansen L, Mannix RC, Kleinman MT, Phalen RF. Effect of nitric acid vapor on the response to inhaled ozone. *Inhalation Toxicology* 1992;4:343-358.
- 176. Naumann BD, Schlesinger RB. Assessment of early alveolar particle clearance and macrophage function following an acute inhalation of sulfuric acid mist. *Exp Lung Res* 1986;11:13-33.
- 177. Newhouse MT, Wolff RK, Dolovich M, Obminski G. Effect of TLV levels of SO<sub>2</sub> and H<sub>2</sub> SO<sub>4</sub> on bronchial clearance in exercising man. *Arch Environ Health* 1978;33:24-32.
- 178. Nielsen GD, Wolkoff P, Alarie Y. Sensory irritation: risk assessment approaches. *Regul Toxicol Pharmacol* 2007;48:6-18.
- 179. NIOSH. Acids, inorganic: method 7903, issue 2, 6 p. In: *Manual of analytical methods* (*NMAM*), 4th ed, 8/15/94. U.S. National Institute for Occupational Safety and Health, Centers for Disease Control & Prevention. Department of Health and Human Services, 1994.

- 180. NTP. Strong inorganic acid mists containing sulfuric acid CAS no. 7664-93-9 (sulfuric acid). In: *Report on carcinogens, eleventh edition*. U.S. Department of Health and Human Services, Public Health Service, National Toxicology Program, 2002.
- 181. Oddoy A, Drabke P, Felgner U, Kirsch H, Lachmann B, Merker G, Robertson B, Vogel J. [The effect of intermittent hydrogen chloride exposure on the lung function of the guinea pig]. Z Erkr Atmungsorgane 1982;158:285-290.
- 182. OSHA. Sulfuric acid in workplace atmosphere (ID-113). Occupational Safety & Health Administration. U. S Department of Labor. http://www.osha.gov/dts/sltc/methods/inorganic/id113/id113.html. Accessed September 2008.
- 183. Parent ME, Siemiatycki J, Fritschi L. Workplace exposures and oesophageal cancer. *Occup Environ Med* 2000;57:325-334.
- 184. Pesatori AC, Consonni D, Rubagotti M, Bonzini M, Catalano P, Bertazzi PA. [Mortality study in a cohort of workers employed in a plant producing sulphuric acid ]. *Med Lav* 2006;97:735-748.
- 185. Peters SG, Hyatt RE. A canine model of bronchial injury induced by nitric acid. Lung mechanics and morphologic features. *Am Rev Respir Dis* 1986;133:1049-1054.
- Petersen PE, Gormsen C. Oral conditions among German battery factory workers. *Community Dent Oral Epidemiol* 1991;19:104-106.
- 187. Petrauskaite R, Pershagen G, Gurevicius R. Lung cancer near an industrial site in Lithuania with major emissions of airway irritants. *Int J Cancer* 2002;99:106-111.
- 188. Prevent. *Chemical substances database.* http://www.prevent.se/verktyg/kemiska\_amnen/default.asp. Accessed October 2007.
- 189. Rajan B, Alesbury R, Carton B, Gerin M, Litske H, Marquart H, Olsen E, Scheffers T, Stamm R, Woldbaek T. European proposal for core information for the storage and exchange of workplace exposure measurements on chemical agents. *Appl Occup Environ Hyg* 1997;12:31-39.
- 190. Reist PC. Basic aerosol science. In: Harris E, ed. *Patty's Industrial Hygiene*. 5th ed. Vol 1. New York: John Wiley and Sons, 2000:355-410.
- 191. Remijn B, Koster P, Houthuijs D, Boleij J, Willems H, Brunekreef B, Biersteker K, van Loveren C. Zinc chloride, zinc oxide, hydrochloric acid exposure and dental erosion in a zinc galvanizing plant in the Netherlands. *Ann Occup Hyg* 1982;25:299-307.
- 192. Roth SH, Bjarnason SG, De Sanctis GT, Feroah T, Jiang X, Karkhanis A, Green FH. Ventilatory responses in awake guinea pigs exposed to acid aerosols. *J Toxicol Environ Health A* 1998;54:261-283.
- 193. RTECS. Registry of Toxic Effects of Chemical Substances. Compiled by the Canadian Centre for Occupational Health and Safety. Licensed through MDL Information Services, Inc. San Leandro, CA: MDL Information Services, Inc., 2006.
- 194. Rubin AE, Wang KP, Liu MC. Tracheobronchial stenosis from acid aspiration presenting as asthma. *Chest* 2003;123:643-646.
- 195. Sackner MA, Ford D, Fernandez R, Cipley J, Perez D, Kwoka M, Reinhart M, Michaelson ED, Schreck R, Wanner A. Effects of sulfuric acid aerosol on cardiopulmonary function of dogs, sheep, and humans. *Am Rev Respir Dis* 1978;118:497-510.
- 196. Sarangapani R, Wexler AS. Growth and neutralization of sulfate aerosols in human airways. *J Appl Physiol* 1996;81:480-490.
- 197. Sathiakumar N, Delzell E, Amoateng-Adjepong Y, Larson R, Cole P. Epidemiologic evidence on the relationship between mists containing sulfuric acid and respiratory tract cancer. *Crit Rev Toxicol* 1997;27:233-251.
- 198. Schaper M, Kegerize J, Alarie Y. Evaluation of concentration-response relationships for histamine and sulfuric acid aerosols in unanesthetized guinea pigs for their effects on ventilatory response to CO2. *Toxicol Appl Pharmacol* 1984;73:533-542.

- 199. Schiff LJ, Bryne MM, Fenters JD, Graham JA, Gardner DE. Cytotoxic effects of sulfuric acid mist, carbon particulates, and their mixtures on hamster tracheal epithelium. *Environ Res* 1979;19:339-354.
- Schlesinger RB, Halpern M, Albert RE, Lippmann M. Effect of chronic inhalation of sulfuric acid mist upon mucociliary clearance from the lungs of donkeys. *J Environ Pathol Toxicol* 1979;2:1351-1367.
- 201. Schlesinger RB. Exposure-response pattern for sulfuric acid-induced effects on particle clearance from the respiratory region of rabbit lungs. *Inhalation Toxicology* 1990;2:21-27.
- 202. Schlesinger RB, Chen LC, Finkelstein I, Zelikoff JT. Comparative potency of inhaled acidic sulfates: speciation and the role of hydrogen ion. *Environ Res* 1990;52:210-224.
- Schlesinger RB, Gunnison AF, Zelikoff JT. Modulation of pulmonary eicosanoid metabolism following exposure to sulfuric acid. *Fundam Appl Toxicol* 1990;15:151-162.
- 204. Schlesinger RB, Gorczynski JE, Dennison J, Richards L, Kinney PL, Bosland MC. Long-term intermittent exposure to sulfuric acid aerosol, ozone, and their combination: alterations in tracheobronchial mucociliary clearance and epithelial secretory cells. *Exp Lung Res* 1992;18:505-534.
- 205. Schlesinger RB, Zelikoff JT, Chen LC, Kinney PL. Assessment of toxicologic interactions resulting from acute inhalation exposure to sulfuric acid and ozone mixtures. *Toxicol Appl Pharmacol* 1992;115:183-190.
- 206. Schlesinger RB, Chen LC. Comparative biological potency of acidic sulfate aerosols: implications for the interpretation of laboratory and field studies. *Environ Res* 1994;65:69-85.
- 207. Schlesinger RB, El-Fawal HA, Zelikoff JT, Gorczynski JE, McGovern T, Nadziejko CE, Chen LC. Pulmonary effects of repeated episodic exposures to nitric acid vapor alone and in combination with ozone. *Inhalation Toxicology* 1994;6:21-41.
- 208. Schultz G, Henkind P, Gross EM. Acid burns of the eye. Am J Ophthalmol 1968;66:654-657.
- 209. SCOEL. Recommendation of the Scientific Committee on Occupational Exposure Limits for nitric acid. SCOEL/SUM/61, 2001.
- 210. SCOEL. Recommendation of the Scientific Committee on Occupational Exposure Limits for sulphuric acid. SCOEL/SUM/105, 2007.
- 211. Scott D, Galloway SM, Marshall RR, Ishidate M, Jr., Brusick D, Ashby J, Myhr BC. International Commission for Protection Against Environmental Mutagens and Carcinogens. Genotoxicity under extreme culture conditions. A report from ICPEMC Task Group 9. *Mutat Res* 1991;257:147-205.
- 212. Sellakumar AR, Snyder CA, Solomon JJ, Albert RE. Carcinogenicity of formaldehyde and hydrogen chloride in rats. *Toxicol Appl Pharmacol* 1985;81:401-406.
- 213. Shangina O, Brennan P, Szeszenia-Dabrowska N, Mates D, Fabianova E, Fletcher T, t'Mannetje A, Boffetta P, Zaridze D. Occupational exposure and laryngeal and hypopharyngeal cancer risk in central and eastern Europe. *Am J Epidemiol* 2006;164:367-375.
- 214. Siemiatycki J ed. *Risk factors for cancer in the workplace*. 325 pp. Boca Raton: CRC press, 1991.
- Sindhu RK, Mautz WJ, Kikkawa Y. Chronic exposure to ozone and nitric acid vapor results in increased levels of rat pulmonary putrescine. *Arch Toxicol* 1998;72:445-449.
- 216. Smeets MA, Kroeze JH, Dalton PH. Setting occupational exposure limits in humans: contributions from the field of experimental psychology. *Int Arch Occup Environ Health* 2006;79:299-307.
- Soskolne CL, Zeighami EA, Hanis NM, Kupper LL, Herrmann N, Amsel J, Mausner JS, Stellman JM. Laryngeal cancer and occupational exposure to sulfuric acid. *Am J Epidemiol* 1984;120:358-369.
- 218. Soskolne CL, Jhangri GS, Siemiatycki J, Lakhani R, Dewar R, Burch JD, Howe GR, Miller AB. Occupational exposure to sulfuric acid in southern Ontario, Canada, in association with laryngeal cancer. *Scand J Work Environ Health* 1992;18:225-232.

- 219. Spektor DM, Leikauf GD, Albert RE, Lippmann M. Effects of submicrometer sulfuric acid aerosols on mucociliary transport and respiratory mechanics in asymptomatic asthmatics. *Environ Res* 1985;37:174-191.
- 220. Spektor DM, Yen BM, Lippmann M. Effect of concentration and cumulative exposure of inhaled sulfuric acid on tracheobronchial particle clearance in healthy humans. *Environ Health Perspect* 1989;79:167-172.
- 221. Spengler JD, Keeler GJ, Koutrakis P, Ryan PB, Raizenne M, Franklin CA. Exposures to acidic aerosols. *Environ Health Perspect* 1989;79:43-51.
- SPIN database. Substances in Preparations in Nordic Countries. http://www.spin2000.net. Accessed August 2008.
- 223. Stavert DM, Archuleta DC, Behr MJ, Lehnert BE. Relative acute toxicities of hydrogen fluoride, hydrogen chloride, and hydrogen bromide in nose- and pseudo-mouth-breathing rats. *Fundam Appl Toxicol* 1991;16:636-655.
- 224. Stayner LT, Meinhardt T, Lemen R, Bayliss D, Herrick R, Reeve GR, Smith AB, Halperin W. A retrospective cohort mortality study of a phosphate fertilizer production facility. *Arch Environ Health* 1985;40:133-138.
- Steenland K, Schnorr T, Beaumont J, Halperin W, Bloom T. Incidence of laryngeal cancer and exposure to acid mists. *Br J Ind Med* 1988;45:766-776.
- 226. Steenland K, Beaumont J. Further follow-up and adjustment for smoking in a study of lung cancer and acid mists. *Am J Ind Med* 1989;16:347-354.
- 227. Steenland K. Laryngeal cancer incidence among workers exposed to acid mists (United States). *Cancer Causes Control* 1997;8:34-38.
- Stevens B, Koenig JQ, Rebolledo V, Hanley QS, Covert DS. Respiratory effects from the inhalation of hydrogen chloride in young adult asthmatics. J Occup Med 1992;34:923-929.
- 229. Swenberg JA, Beauchamp RO, Jr. A review of the chronic toxicity, carcinogenicity, and possible mechanisms of action of inorganic acid mists in animals. *Crit Rev Toxicol* 1997;27:253-259.
- 230. Ten Bruggen Cate HJ. Dental erosion in industry. Br J Ind Med 1968;25:249-266.
- 231. Teta MJ, Perlman GD, Ott MG. Mortality study of ethanol and isopropanol production workers at two facilities. *Scand J Work Environ Health* 1992;18:90-96.
- 232. Tunnicliffe WS, Harrison RM, Kelly FJ, Dunster C, Ayres JG. The effect of sulphurous air pollutant exposures on symptoms, lung function, exhaled nitric oxide, and nasal epithelial lining fluid antioxidant concentrations in normal and asthmatic adults. *Occup Environ Med* 2003;60:e15.
- Tuominen M, Tuominen R, Ranta K, Ranta H. Association between acid fumes in the work environment and dental erosion. *Scand J Work Environ Health* 1989;15:335-338.
- Tuominen ML, Tuominen RJ, Fubusa F, Mgalula N. Tooth surface loss and exposure to organic and inorganic acid fumes in workplace air. *Community Dent Oral Epidemiol* 1991;19:217-220.
- Uleckiene S, Griciute L. Carcinogenicity of sulfuric acid in rats and mice. *Pathol Oncol Res* 1997;3:38-43.
- Utell MJ, Morrow PE, Speers DM, Darling J, Hyde RW. Airway responses to sulfate and sulfuric acid aerosols in asthmatics. An exposure-response relationship. *Am Rev Respir Dis* 1983;128:444-450.
- 237. Utell MJ, Morrow PE, Hyde RW. Airway reactivity to sulfate and sulfuric acid aerosols in normal and asthmatic subjects. *J Air Pollut Control Assoc* 1984;34:931-935.
- 238. Utell MJ, Mariglio JA, Morrow PE, Gibb FR, Speers DM. Effects of inhaled acid aerosols on respiratory function: the role of endogenous ammonia. *J Aerosol Med* 1989;2:141-147.
- Utell MJ, Frampton MW. Sulfur dioxide and sulfuric acid aerosols. In: Rom N ed. *Environmental and occupational medicine*. 2nd ed. Boston, MA: Little, Brown and Company,1992:519-527.

- 240. van Thriel C, Schaper M, Kiesswetter E, Kleinbeck S, Juran S, Blaszkewicz M, Fricke HH, Altmann L, Berresheim H, Bruning T. From chemosensory thresholds to whole body exposures-experimental approaches evaluating chemosensory effects of chemicals. *Int Arch Occup Environ Health* 2006;79:308-321.
- 241. Wanner A, Salathe M, O'Riordan TG. Mucociliary clearance in the airways. *Am J Respir Crit Care Med* 1996;154:1868-1902.
- 242. Weast, RC, ed. *CRC Handbook of chemistry and physics*. 52nd ed. The Chemical Rubber Co. Cleveland, Ohio, United States, 1971.
- 243. Weil CS, Smyth HF, Jr., Nale TW. Quest for a suspected industrial carcinogen. A M A Arch Ind Hyg Occup Med 1952;5:535-547.
- 244. Wiegand A, Attin T. Occupational dental erosion from exposure to acids: a review. *Occup Med* (*Lond*) 2007;57:169-176.
- 245. Witschi HP, Last JA. Toxic responses of the respiratory system. In: Klaassen CD ed. Casarett and Doull's. Toxicology: The basic science of poisons. New York: McGraw-Hill, 2001:515-534.
- 246. Wolff RK, Muggenburg BA, Silbaugh SA. Effect of 0.3 and 0.9 micron sulfuric acid aerosols on tracheal mucous clearance in beagle dogs. *Am Rev Respir Dis* 1981;123:291-294.
- 247. Vollmuth TA, Schlesinger RB. Measurement of respiratory tract ammonia in the rabbit and implications to sulfuric acid inhalation studies. *Fundam Appl Toxicol* 1984;4:455-464.
- Wong CG, Bonakdar M, Mautz WJ, Kleinman MT. Chronic inhalation exposure to ozone and nitric acid elevates stress-inducible heat shock protein 70 in the rat lung. *Toxicology* 1996;107:111-119.
- 249. Wong KL, Alarie Y. A method for repeated evaluation of pulmonary performance in unanesthetized, unrestrained guinea pigs and its application to detect effects of sulfuric acid mist inhalation. *Toxicol Appl Pharmacol* 1982;63:72-90.
- Zelikoff JT, Schlesinger RB. Modulation of pulmonary immune defense mechanisms by sulfuric acid: effects on macrophage-derived tumor necrosis factor and superoxide. *Toxicology* 1992;76:271-281.
- 251. Zelikoff JT, Sisco MP, Yang Z, Cohen MD, Schlesinger RB. Immunotoxicity of sulfuric acid aerosol: effects on pulmonary macrophage effector and functional activities critical for maintaining host resistance against infectious diseases. *Toxicology* 1994;92:269-286.
- 252. Zelikoff JT, Frampton MW, Cohen MD, Morrow PE, Sisco M, Tsai Y, Utell MJ, Schlesinger RB. Effects of inhaled sulfuric acid aerosols on pulmonary immunocompetence: A comparative study in humans and animals. *Inhalation Toxicology* 1997;9:731-752.

## 19. Data bases used in the literature search

The major literature searches were performed in January 2006. The following databases were used:

Arbline Cheminfo CISDOC HSELINE MHIDAS NIOSHTIC2 OSHLINE PubMed RILOSH RTECS Toxline

A final search in PubMed and Toxline was performed on October 29, 2008.

Submitted for publication June 12, 2009.

## Appendix 1. Occupational exposure limits

Country	Sulphuric		Hydrochloric		Nitric		Phosphoric		Reference
	acid		acid		acid		acid		
	8-h	STEL	8-h	STEL	8-h	STEL	8-h	STEL	
Denmark	1	2	7 <sup>a</sup>	-	1.3	2.6	1	2	(1)
Finland	0.2	1	-	7.6	1.3	2.6	1	2	(2)
Germany (DFG)	$0.1^{b}$	0.1 <sup>b</sup>	3	6	-	-	2 <sup>b</sup>	4 <sup>b</sup>	(3)
• • •		$0.2^{a, b}$							
The Netherlands	-	-	8	15	-	1.3	1	2	(4)
Norway	0.1	0.3	7 <sup>a</sup>	-	5	10	1	3	(5)
Sweden	1	3	8 <sup>a</sup>	-	5	13	1	3	(6)
United Kingdom	-	-	2	8	-	2.6	1	2	(7)
US (ACGIH)	0.2 °	-	-	2.8 <sup>a</sup>	5	10	1	3	(8)
US (NIOSH)	1	-	7 <sup>a</sup>	-	5	10	1	3	(9)
US (OSHA)	1	-	7 <sup>a</sup>	-	5	-	1	-	(9)
EU	-	-	8	15	-	2.6	1	-	(10, 11)

Occupational exposure limits  $(mg/m^3)$  for the inorganic acids in different countries as time-weighted averages.

<sup>a</sup> Ceiling value.

<sup>b</sup> Inhalable fraction.

<sup>c</sup> Thoracic fraction.

STEL: short-term exposure limit, OSHA: Occupational Safety and Health Administration.

### References

- 1. *Grænsværdier for stoffer og materialer*. At-vejledning. Stoffer og materialer-C.0.1. Køpenhavn: Arbejdstilsynet, August 2007.
- 2. *HTP-värden 2007*. Koncentrationer som befunnits skadliga. Helsingfors: Social- och hälsovårdsministeriets publikationer, 2007:20.
- 3. *MAK- und BAT-Werte-liste* 2007. Senatskommission zur Prüfung gesundheitsschädlicher Arbeitsstoffe. Mitteilung 43. Bonn: Deutsche Forschungsgemeinschaft (DFG), 2007.
- 4. *Wijziging Arbeidsomstandighedenregeling*. Staatscourant 28 december 2006, nr. 252 / pag. 23. Den Haag: Ministerie van Sociale Zaken en Werkgelegenheid, 2007.
- 5. Administrative normer for forurensning i arbeidsatmosfære. Veiledning til arbeidsmiljøloven, Oslo: Arbeidstilsynet, 2009.

http://www.arbeidstilsynet.no/c28864/artikkel/vis.html?tid=28880.

- Hygieniska gränsvärden och åtgärder mot luftföroreningar. Arbetsmiljöverkets författningssamling, AFS 2005:17. Solna: Arbetsmiljöverket, 2005.
- 7. *EH40/2005 Workplace exposure limits*. Table 1: List of approved workplace exposure limits (as consolidated with amendments October 2007). London: Health and Safety Executive, 2007.
- 8. 2007 TLVs and BEIs. Based on the documentation of the "Threshold limit values for chemical substances and physical agents and biological exposure indices". Cincinnati, Ohio: The American Conference of Governmental Industrial Hygienists (ACGIH), 2007.
- 9. *NIOSH pocket guide to chemical hazards*. Cincinnati, Ohio: National Institute for Occupational Safety and Health, 2005.
- Indicative occupational exposure limit values. Commission directives: 2000/39/EC of 8 June 2000. Official Journal of the European Communities: L 142/47-50. Brussels: The commission of the European communities, 2000.
- 11. *Indicative occupational exposure limit values*. Commission directives: 2006/15/EC of 7 February 2006. Official Journal of the European Communities: 9.2.2006. Brussels: The commission of the European communities, 2006.

## Appendix 2. Previous NEG criteria documents

Substance/Agent	Arbete och Hälsa issue
Acetonitrile	1989:22, 1989:37*
Acid aerosols, inorganic	1992:33, 1993:1*
Acrylonitrile	1985:4
Allyl alcohol	1986:8
Aluminium	1992:45, 1993:1*
Ammonia	1986:31, 2005:13*
Antimony	1998:11*
Arsenic, inorganic	1981:22, 1991:9, 1991:50*
Arsine	1986:41
Asbestos	1982:29
Benomyl	1984:28
Benzene	1981:11
1,2,3-Benzotriazole	2000:24*D
Boric acid, Borax	1980:13
1,3-Butadiene	1994:36*, 1994:42
1-Butanol	1980:20
γ-Butyrolactone	2004:7*D
Cadmium	1981:29, 1992:26, 1993:1*
7/8 Carbon chain aliphatic monoketones	1990:2*D
Carbon monoxide	1980:8
Ceramic Fibres, Refractory	1996:30*, 1998:20
Chlorine, Chlorine dioxide	1980:6
Chloromequat chloride	1984:36
4-Chloro-2-methylphenoxy acetic acid	1981:14
Chlorophenols	1984:46
Chlorotrimethylsilane	2002:2
Chromium	1979:33
Cobalt	1982:16, 1994:39*, 1994:42
Copper	1980:21
Creosote	1988:13, 1988:33*
Cyanoacrylates	1995:25*, 1995:27
Cyclic acid anhydrides	2004:15*D
Cyclohexanone, Cyclopentanone	1985:42
n-Decane	1987:25, 1987:40*
Deodorized kerosene	1985:24
Diacetone alcohol	1989:4, 1989:37*
Dichlorobenzenes	1998:4*, 1998:20
Diesel exhaust	1993:34, 1993:35*
Diethylamine	1994:23*, 1994:42
2-Diethylaminoethanol	1994:25*N
Diethylenetriamine	1994:23*, 1994:42
Diisocyanates	1979:34, 1985:19
Dimethylamine	1994:23*, 1994:42
Dimethyldithiocarbamates	1990:26, 1991:2*
Dimethylethylamine	1990:20, 1991:20
Dimethylformamide	1983:28
Dimethylsulfoxide	1991:37, 1991:50*
Dioxane	1982:6
Enzymes, industrial	1994:28*, 1994:42
Epichlorohydrin	1994.20, 1994.42
Ethyl acetate	1990:35*
Empractiale	1770.33

NEG criteria documents published in the scientific serial Arbete and Hälsa (Work and Health):

Substance/Agent	Arbete och Hälsa issue
Ethylbenzene	1986:19
Ethylenediamine	1994:23*, 1994:42
Ethylenebisdithiocarbamates and Ethylenethiourea	1993:24, 1993:35*
Ethylene glycol	1980:14
Ethylene glycol monoalkyl ethers	1985:34
Ethylene oxide	1982:7
Ethyl ether	1992:30* N
2-Ethylhexanoic acid	1994:31*, 1994:42
Flour dust	1996:27*, 1998:20
Formaldehyde	1978:21, 1982:27, 2003:11*D
Fungal spores	2006:21*
0 1	
Furfuryl alcohol	1984:24
Gasoline	1984:7 1997 20*P 1998 20
Glutaraldehyde	1997:20*D, 1998:20
Glyoxal	1995:2*, 1995:27
Halothane	1984:17
n-Hexane	1980:19, 1986:20
Hydrazine, Hydrazine salts	1985:6
Hydrogen fluoride	1983:7
Hydrogen sulphide	1982:31, 2001:14*D
Hydroquinone	1989:15, 1989:37*
Industrial enzymes	1994:28*
Isophorone	1991:14, 1991:50*
Isopropanol	1980:18
Lead, inorganic	1979:24, 1992:43, 1993:1*
Limonene	1993:14, 1993:35*
Lithium and lithium compounds	2002:16*
Manganese	1982:10
Mercury, inorganic	1985:20
Methacrylates	1983:21
Methanol	1984:41
Methalor Methyl bromide	1987:18, 1987:40*
Methyl chloride	1992:27*D
Methyl chloroform	1981:12
Methylcyclopentadienyl manganese tricarbonyl	1982:10
Methylene chloride	1979:15, 1987:29, 1987:40*
Methyl ethyl ketone	1983:25
Methyl formate	1989:29, 1989:37*
Methyl isobutyl ketone	1988:20, 1988:33*
Methyl methacrylate	1991:36*D
N-Methyl-2-pyrrolidone	1994:40*, 1994:42
Methyl-tert-butyl ether	1994:22*D
Microbial volatile organic compounds (MVOCs)	2006:13*
Microorganisms	1991:44, 1991:50*
Mineral fibers	1981:26
Nickel	1981:28, 1995:26*, 1995:27
Nitrilotriacetic acid	1989:16, 1989:37*
Nitroalkanes	1988:29, 1988:33*
Nitrogen oxides	1983:28
N-Nitroso compounds	1990:33, 1991:2*
Nitrous oxide	1990.33, 1991.2
Oil mist	1982.20
Organic acid anhydrides	1990:48, 1991:2*
Ozone	1986:28
Paper dust	1989:30, 1989:37*
Penicillins	2004:6*
Permethrin	1982:22
Petrol	1984:7

Substance/Agent	Arbete och Hälsa issue
Phenol	1984:33
Phthalate esters	1982:12
Platinum	1997:14*D, 1998:20
Polyethylene,	1998:12*
Polypropylene, Thermal degradation products in the	1998:12*
processing of plastics	
Polystyrene, Thermal degradation products in the	1998:12*
processing of plastics	
Polyvinylchloride, Thermal degradation products in the	1998:12*
processing of plastics	
Polytetrafluoroethylene, Thermal degradation products	1998:12*
in the processing of plastics	
Propene	1995:7*, 1995:27
Propylene glycol	1983:27
Propylene glycol ethers and their acetates	1990:32*N
Propylene oxide	1985:23
Refined petroleum solvents	1982:21
Refractory Ceramic Fibres	1996:30*
Selenium	1992:35, 1993:1*
Silica, crystalline	1993:2, 1993:35*
Styrene	1979:14, 1990:49*, 1991:2
Sulphur dioxide	1984:18
Synthetic pyretroids	1982:22
Tetrachloroethane	1996:28*D
Tetrachloroethylene	1979:25, 2003:14*D
Thermal degradation products of plastics	1998:12*
Thiurams	1990:26, 1991:2*
Tin and inorganic tin compounds	2002:10*D
Toluene	1979:5, 1989:3, 1989:37*, 2000:19 <sup>3</sup>
1,1,1-Trichloroethane	1981:12
Trichloroethylene	1979:13, 1991:43, 1991:50*
Triglycidyl isocyanurate	2001:18*
n-Undecane	1987:25, 1987:40*
Vanadium	1982:18
Vinyl acetate	1988:26, 1988:33*
Vinyl chloride	1986:17
Welding gases and fumes	1990:28, 1991:2*
White spirit	1986:1
Wood dust	1987:36
Xylene	1979:35
Zinc	1981:13
* in English remaining documents are in a Scandinavian	

\* in English, remaining documents are in a Scandinavian language.

D = collaboration with the Dutch Expert Committee on Occupational Standards (DECOS).

N = collaboration with US NIOSH.

To order further copies in this series, please contact: Arbets- och miljömedicin, Göteborgs universitet Att: Cina Holmer, Box 414, SE-405 30 Göteborg, Sweden E-mail: arbeteochhalsa@amm.gu.se. The NEG documents are also available on the web at www.nordicexpertgroup.org www.amm.se/aoh

# Latest issues in the scientific serial WORK AND HEALTH

**2006:10. M C Nelson (Ed)** Occupational Health and Public Health. Lessons from the Past – Challenges for the Future.

**2006:11. J Montelius (Ed)** Scientific Basis for Swedish Occupational Standards. XXVII

### 2006:12. M Barnekow Bergkvist.

Arbetslivsinstitutets expertgrupp för ergonomisk dokumentation - Dokument 5. Kan fysisk träning i arbetslivet förbättra muskuloskeletal hälsa? En kunskapsöversikt.

**2006:13.** A Korpi, J Järnberg and A-L Pasanen. The Nordic Expert Group for Criteria Documentation of Health Risks from Chemicals 138. Microbial volatile organic compounds (MVOCs).

**2006:14. M Oudhuis och A Olsson.** Spelar värderingar någon roll för arbetsmiljön? En studie om konsekvenser vid övergång till utländskt ägande och vid generationsskiften i företag.

**2006:15. A Hedlund och B Pontén.** Införande av systematiskt arbetsmiljöarbete på träföretag – utvärdering av en metod, dess resultat och påverkan på arbetets attraktivitet.

### 2006:16 K Håkansson och T Isidorsson.

Arbetsmiljöarbete och långsiktigt hållbara arbetsorganisationer. Ett delprojekt inom Arbetslivsinstitutets tema Strategier, metoder och arbetssätt för fungerande arbetsmiljöarbete SMARTA.

2006:17. J Eklund, B Hansson, L Karlqvist, L Lindbeck och W P Neumann. Arbetsmiljöarbete och effekter – en kunskapsöversikt.

### 2006:18. L Rose och U Orrenius.

Arbetslivsinstitutets expertgrupp för ergonomisk dokumentation - Dokument 6. Beräkning av arbetsmiljöns ekonomiska effekter på företag och organisationer. En översikt av ett urval modeller och metoder.

**2006:19. C Stenlund och M Torgén**. Arbetsledare i processindustrin. Arbetsuppgifter, förutsättningar, psykosocial arbetsmiljö och självskattad hälsa.

2006:20. I-M Andersson, J Laring, M Åteg och G Rosén. Arbetsmiljöfrågans väg. Samverkan mellan kundföretag och företagshälsovård.

**2006:21. W Eduard**. Fungal spores. The Nordic Expert Group for Criteria Documentation of Health Risks from Chemicals139.

### 2006:22. M Björkman, I Carlsson.

Känsla av sammanhang på arbetet. Vilka faktorer på arbetsplatsen och hos individen främjar arbetsterapeuters upplevelse av ett meningsfullt arbete? **2007;41:1. A Lindegård Andersson.** Working technique during computerwork. Associations with biomechanical and psychological strain, neck and upper extremity musculoskeletal symptoms.

2008;42:1. P Westerholm (red.) Psykisk arbetsskada

**2008;42:2. G Johanson, M Rauma.** Basis for skin notation. Part 1. Dermal penetration data for substances on the Swedish OEL list.

**2008;42:3. J Montelius (Ed.)** Vetenskapligt Underlag för Hygieniska Gränsvärden 28. Kriteriegruppen för hygieniska gränsvärden.

**2008;42:4. P Wiebert.**The impact of airway-irritating exposure and wet work on subjects with allergy or other sensitivity - epidemiology and mechanisms

**2008;42:5. E Månsson**. Att skapa en känsla av sammanhang -om resultatet av hälsofrämjande strategier bland lärare.

**2008;42:6. J Montelius (Ed.)** Scientific Basis for Swedish Occupational Standards. XXVIII

2008;42:7. B Melin Experimentell och epidemiologisk forskning –relationen psykosocial exponering, stress, psykisk belastning, muskelaktivitet och värk i nacke-skuldra.

**2009;43(1) J Montelius (Ed.)** Vetenskapligt Underlag för Hygieniska Gränsvärden 29. Kriteriegruppen för hygieniska gränsvärden.

**2009;43(2) J Weiner.** Könsskillnader i ersättning vid arbetsskador? – en 10- årsuppföljning av arbetsskador 1994.

2009;43(3) G Aronsson, K Gustafsson och C Mellner. Samband mellan sjuknärvaro, sjukfrånvaro och självskattad hälsa i den

yrkesaktiva befolkningen.

**2009;43(4) J Montelius (Ed.)** Scientific Basic for Swedish Occupational Standards XXIX. Swedish criteria Group for Occupational Standards

2009;43(5) K Kruse och W Eduard. Prøvetaking av inhalerbart melstøv.

**2009;43(6) E Gustafsson** Physical exposure, musculoskeletal symptoms and attitudes related to ICT use.

**2009;43(7) M van der Hagen and J Järnberg.** The Nordic Expert Group for Criteria Documentation of Health Risks from Chemicals140. Sulphuric, hydrochloric,nitric and phosphoric acids.