Health-based Reassessment of Administrative Occupational Exposure Limits

Gezondheidsraad

Voorzitter

Health Council of the Netherlands

Aan de Staatssecretaris van Sociale Zaken en Werkgelegenheid

Onderwerp : Aanbieding adviezen herevaluatie bestuurlijke MAC-waarden

Uw kenmerk : ARBO/AMIL/97/00648 Ons kenmerk : U 343/HS/mj/563-J5

Bijlagen : 13

Datum : 7 maart 2002

Mijnheer de staatssecretaris,

Op verzoek van uw ambtsvoorganger bied ik u hierbij 13 adviezen aan van een reeks over de gezondheidskundige basis van uit het buitenland overgenomen grenswaarden voor beroepsmatige blootstelling aan stoffen. Het verzoek om deze adviezen is in algemene zin vervat in brief nr ARBO/AMIL/97/00648 en in latere stadia door uw departement toegespitst op afzonderlijke stoffen. De adviezen zijn opgesteld door een daartoe door mij geformeerde internationale commissie van de Gezondheidsraad en beoordeeld door de Beraadsgroep Gezondheid en Omgeving.

De beoogde reeks van in het Engels gestelde adviezen zal losbladig worden gepubliceerd onder ons publicatienummer 2000/15OSH en, conform de aan de Gezondheidsraad voorgelegde toespitsingen van de adviesaanvraag, betrekking hebben op 168 stoffen. Het u thans aangeboden derde pakket bestaat uit de adviezen genummerd 2000/15OSH/030 tot en met 2000/15OSH/042, respectievelijk betrekking hebbend op: *p-tert-butyltolueen, cellulose, 2-chloor-6-(trichloormethyl)pyridine, ethylformaat, 4-ethylmorfoline, indeen, perchloormethylmercaptaan, o-, m-, p-terfenyl (mengsel), tetramethylsuccinonitril, tricarbonyl(eta-cyclopentadiënyl)mangaan, ijzerpentacarbonyl, zetmeel en zwaveltetrafluoride*.

De u thans aangeboden adviezen heb ik vandaag ter informatie doen toekomen aan de Minister van Volkshuisvesting, Ruimtelijke Ordening en Milieubeheer.

Hoogachtend,

prof. dr JA Knottnerus

Health-based Reassessment of Administrative Occupational Exposure Limits

Committee on Updating of Occupational Exposure Limits, a committee of the Health Council of the Netherlands

No. 2000/15OSH, The Hague, 14 December 2000

The Health Council of the Netherlands, established in 1902, is an independent scientific advisory body. Its remit is "to advise the government and Parliament on the current level of knowledge with respect to public health issues..." (Section 21, Health Act).

The Health Council receives most requests for advice from the Ministers of Health, Welfare & Sport, Housing, Spatial Planning & the Environment, Social Affairs & Employment, and Agriculture, Nature Preservation & Fisheries. The Council can publish advisory reports on its own initiative. It usually does this in order to ask attention for developments or trends that are thought to be relevant to government policy.

Most Health Council reports are prepared by multidisciplinary committees of Dutch or, sometimes, foreign experts, appointed in a personal capacity. The reports are available to the public.

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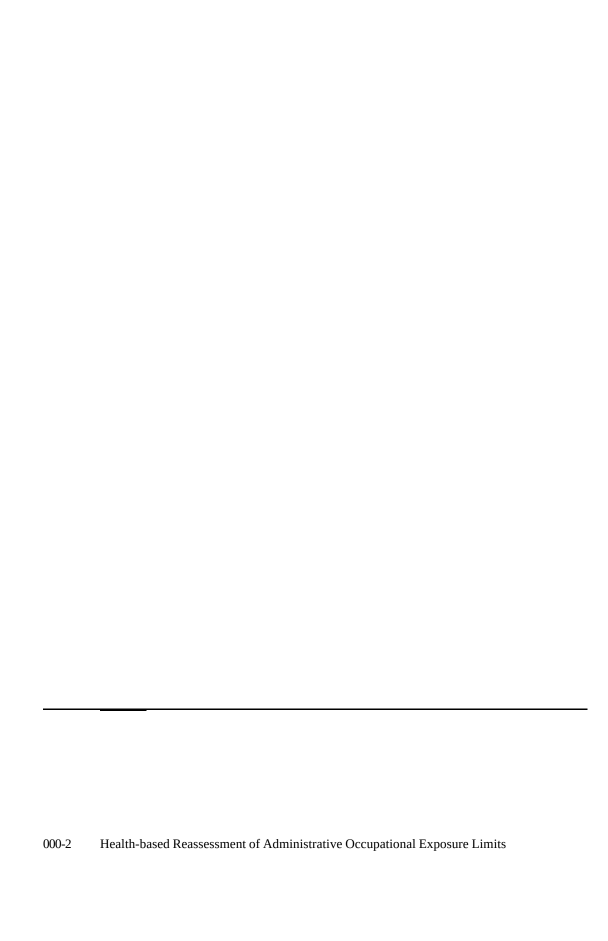
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General remarks

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General introduction

1.1 Background of the project

In September 1995, TNO Nutrition and Food Research Institute (Zeist, the Netherlands) reported the Minister of Social Affairs and Employment about the degree of health protection of the administrative occupational exposure limits (OELs) in the list of Maximal Accepted Concentrations (MAC) of 1994. The major part of these OELs were adopted from the American Conference of Governmental Industrial Hygienists' list of Threshold Limit Values in the 1970s. For the purpose of the evaluation concise toxicity profiles were prepared based on the documentation of the ACGIH and European criteria documents on almost 300 substances. TNO concluded that for 109 substances the current OEL was suspected to be too high from a health protection point of view, with deviations ranging from a factor of 2 to 250. For another 106 substances the toxicological data base was judged to be too poor to recommend a health-based OEL.

On September 24, 1996 the State Secretary of Social Affairs and Employment presented a plan to re-evaluate the administrative OELs to the Social and Economic Council. After the Social and Economic Council had given a positive reaction the State Secretary requested the Health Council on April 16, 1997, to re-evaluate these substances in a condensed procedure and to recommend health-based OELs. Given the international character of the request and in view

of the European harmonization, the State Secretary requested the Health Council to invite scientists from outside the Netherlands to participate in the project.

1.2 Setting OELs in the Netherlands

In the Netherlands the legally-binding occupational exposure limits for chemical substances are set using a three-step procedure. In the first step a scientific evaluation of the data on the toxicity of the substance is made by the Dutch Expert Committee on Occupational Standards (DECOS), a committee of the Health Council, on request of the State Secretary of Social Affairs and Employment. This evaluation should lead to a health-based recommended occupational exposure limit for the concentration of the substance in workroom air.*

In the next phase of the three-step procedure the Social and Economic Council advises the State Secretary on the feasibility of using the health-based value as a legally-binding OEL, or recommends a different OEL. In the final step of the procedure the State Secretary sets the regulatory OEL.

The three-step procedure will also be followed for health-based OELs derived in the present re-evaluation project.

1.3 Committee

As a first step the Health Council requested several regulatory authorities of European countries to nominate internationally acknowledged experts in toxicology, epidemiology or occupational medicine with experience in setting OELs. All nominees were requested to send a brief curriculum vitae and a list of publications. The committee was selected from the list of nominees using the following criteria: one member from each foreign country, the expertise within the committee should cover all aspects of hazard assessment of chemical substances, and good spread in (former) affiliation of the members (academia, national institutes and industry). All members were invited on a personal title, except for the corresponding member representing ACGIH. The members of the Committee on Updating of occupational exposure limits are listed in Chapter 2.

^{*} For genotoxic carcinogens DECOS does not derive an OEL but presents an exposure-response relationship.

1.4 Procedure

Under the authority of the Ministry of Social Affairs and Employment, for each substance a short document is prepared by a toxicologist at a research institute in the Netherlands. The requirements for the contents of these short documents were established by the committee during a trial phase*. The documents should be based on a full literature search including at least the data bases Medline, Toxline and Chemical Abstracts. From the published literature a key study is identified serving as the basis for deriving a health-based OEL. For extrapolation of the data (from the key study) to the occupational exposure situation an overall assessment factor is applied covering the aspects inter- and intraspecies variation, differences in duration and pattern of exposure between the key study and the situation of the worker, type of the critical effect, dose-effect relationship and quality of the total data base. In deriving the overall assessment factor the committee uses a check list of the different aspects, adopted from a TNO-report, to discuss and thoroughly weigh all available data. In each case the committee considers the appropriateness of applying default values, see Chapter 3. In case the key study refers to an oral animal experiment, scaling to humans is based on caloric demand (body weight to the power 0.75) rather than on body weight. For the recommendation of a health-based OEL the committee decided to make use of the preferred value system** given the inherent uncertainty of any OEL.

To derive a health-based OEL for a substance at least data on acute toxicity, including irritation (and sensitization), and on repeated-dose toxicity are required. The committee considers as a minimum the availability of a multi-dose study in which a relevant animal species was exposed via a relevant exposure route for a relevant exposure time and in which relevant toxicological endpoints were studied (preferably including body weight gain, haematology, clinical biochemistry, gross and microscopic pathology). Preferably the study provides information about the target organ and critical effect, and produces a no observed adverse effect level.

^{*} The aims of this phase were to determine the requirements for the documents, to develop and test the procedure, and to judge the feasibility of the project. The fourteen substances under consideration during this phase were selected by the Ministry of Social Affairs and Employment in consultation with regulatory authorities in the participating countries.

The preferred value system implies that OELs are rounded up or down to 'preferred values', e.g. 0.1, 0.2, 0.5, 1, 2, 5, 10 mg/m³ etc.

In case the data base is insufficient to recommend a health-based OEL the committee will strive to judge in a semi-quantitative way whether or not the current administrative limit is health-protective.

Carcinogenic substances are not taken into consideration by the committee. When the data base of a substance indicates carcinogenic and/or genotoxic potential, the committee recommends an evaluation and classification of the substance by DECOS. For substances tested for and proven to be toxic to fertility or development, the committee ascertains that the recommended health-based OEL protects against these effects as well. For quite a number of substances, however, these data were not available.

A draft of each document was released for public review during a period of six weeks and comments received were taken into account in the final version of the document.

The Hague, 14 December 2000, for the committee

dr CA Bouwman, scientific secretary

prof. dr J Noordhoek chairman

The committee

- J Noordhoek, chairman professor of toxicology; University of Nijmegen, Nijmegen, The Netherlands
- A Aitio senior scientist; International Programme on Chemical Safety, World Health Organization, Switzerland
- PL Chambers † co-ordinator toxicology studies; University of Dublin, Ireland
- VJ Feron professor of toxicology; TNO Nutrition and Food Research Institute, Zeist, The Netherlands
- H Greim professor of toxicology; GSF National Research Center for Environment and Health, Oberschleissheim, Germany
- U Hass senior researcher in toxicology; Institute of Food Safety and Toxicology; Søborg, Denmark
- CJ Högberg professor of toxicology; National Institute for Working Life and Karolinska Institutet, Stockholm, Sweden

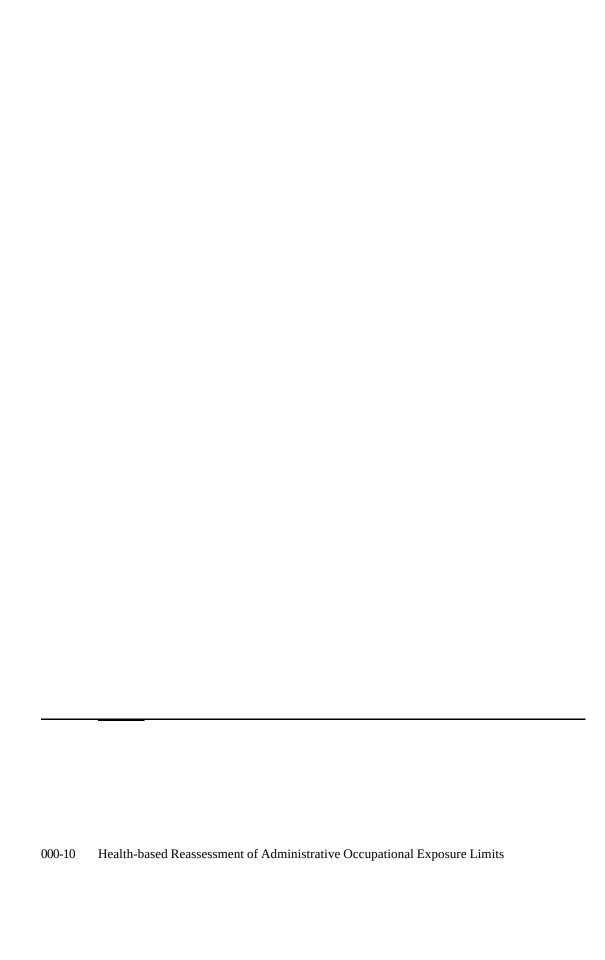
- G De Mik
 - toxicologist; National Institute of Public Health and the Environment, Bilthoven, The Netherlands
- A Moses consultant toxicologist; Cheshire, United Kingdom
- W Seinen professor of toxicology; Utrecht University, Utrecht, The Netherlands
- GMH Swaen epidemiologist; University of Maastricht, Maastricht, The Netherlands
- WMD Wagner, corresponding member
 American Conference of Governmental Industrial Hygienists,
 Cincinnati, Ohio, USA
- RD Zumwalde senior scientist; National Institute for Occupational Safety and Health, Cincinnati, Ohio, USA
- LCMP Hontelez, advisor
 Ministry of Social Affairs and Employment, The Hague, The Netherlands
- WF Passchier, observer
 Health Council of the Netherlands, The Hague, The Netherlands
- CA Bouwman, *scientific secretary*Health Council of the Netherlands, The Hague, The Netherlands

3

Default values

Adopted from the report 'Methods for establishment of Health-based Recommended Occupational Exposure Limits for existing substances', V96.463, 4 July 1996, by TNO Nutrition and Food Research Institute, Zeist, the Netherlands (see also De Raat *et al*, Reg. Toxiol. Pharmacol. 25, 1997: 204-210).

aspects	default value	
interspecies differences	3	
intraspecies differences	3	
differences between experimental conditions and exposure pattern of the worker	1-10	
type of critical effect	1	
dose-response curve	1	
confidence of the data base	1	



Abbreviations

Organisations and occupational exposure limits

ACGIH American Conference of Governmental Industrial Hygienists

DECOS Dutch Expert Committee on Occupational Standards

DFG Deutsche Forschungsgemeinschaft

EPA Environmental Protection Agency (USA) FDA Food and Drug Administration (USA)

HBR-OEL health based recommended occupational exposure limit

HSE Health and Safety Executive (UK)

IARC International Agency for Research on Cancer (WHO)

IPCS International Programme for Chemical Safety

MAC maximaal aanvaarde concentratie (maximal accepted concentration)

MAK Maximale Arbeitsplatz Konzentration

MEL maximum exposure limit

NIOSH National Institute for Occupational Safety and Health (USA)

NTP National Toxicology Programme (USA)

OECD Organisation for Economic Cooperation and Development

OEL occupational exposure limit OES occupational exposure standard

OSHA Occupational Safety and Health Association (USA)

PEL permissible exposure limit recommended exposure limit

RTECS Registry of Toxic Effects of Chemical Substances
SCOEL Scientific Committee for Occupational Exposure Limits
SER Social and Economic Council (Sociaal-Economische Raad NL)

STEL short term exposure limit
TLV threshold limit value
TWA time weighted average
WHO World Health Organisation

Toxicological terms

BALF bronchio-alveolar lavage fluid

bw body weight

CNS central nervous system

 EC_{50} concentration at which a described effect is found in 50% of the exposed animals or at which the effect is decreased up to 50% of the control value

Freunds Complete Adjuvans
forced expiratory volume

FEV forced expiratory volume FVC forced vital capacity GD gestation day(s)

GPMT guinea pig maximisation test

GSH glutathione

h hour

FCA

 IC_{50} concentration at which inhibition of a certain function is found up to 50% of

the control value

im intramuscular

ip intraperitoneal

it intratracheal

iv intravenous

 LC_{50} lethal concentration for 50% of the exposed animals

 LC_{lo} lowest lethal concentration

 LD_{50} lethal dose for 50% of the exposed animals

 LD_{lo} lowest lethal dose LDH lactate dehydrogenase

LOAEL lowest observed adverse effect level minimal alveolar concentration

MFO mixed function oxidase

MMAD mass median aerodynamic diameter

MOAEL minimal observed adverse effect level

NOAEL no observed adverse effect level

n number

ppb parts per billion (v/v)10⁻⁹ ppm parts per million (v/v)10⁻⁶ PNS peripheral nervous system

po per os (= oral) RBC red blood cells

RD₅₀ concentration at which a 50% decrease of respiratory rate is observed

SCE sister chromatid exchange

sc subcutaneous

UDS unscheduled DNA-synthesis

 V_{max} maximal reaction velocity of an enzyme

w week

Statistical terms

CL confidence limits
 GM geometric mean
 OR odds ratio
 RR relative risk
 SD standard deviation
 SEM standard error of mean
 SMR standard mortality ratio

Analytical methods
BE1 biological exposure index
GC gas chromatography
HPLC high performance liquid chromatography
MS mass spectrometry
NMR nuclear magnetic resonance
PAS personal air sampling
TLC thin layer chromatography

