Review

Occupational Exposure to Diisocyanates in the European Union

Dag Rother* and Urs Schlüter

Federal Institute for Occupational Safety and Health (BAuA), Division 4 - Hazardous Substances and Biological Agents, Unit 4.1 - Exposure Scenarios, Friedrich-Henkel-Weg 1, D-44149 Dortmund, Germany

*Author to whom correspondence should be addressed. Tel: +49-231-9071-2749; e-mail: rother.dag@baua.bund.de

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Abstract

Objectives: Diisocyanates are a chemical group that are widely used at workplaces in many sectors. They are also potent skin- and respiratory sensitizers. Exposure to diisocyanates is a main cause of occupational asthma in the European Union. To reduce occupational exposure to diisocyanates and consequently the cases of diisocyanate-induced asthma, a restriction on diisocyanates was recently adopted under the REACH Regulation in the European Union.

Methods: A comprehensive evaluation of the data on occupational exposure to the most important diisocyanates at workplaces was made and is reported here. The diisocyanates considered are methylene diphenyl diisocyanate (MDI), toluene diisocyanate (TDI), and hexamethylene diisocyanate (HDI), accounting for more than 95% of the market volume in the EU. The exposure assessment is based on data from Chemical Safety Reports (CSRs) of REACH Registration Dossiers, workplace air monitoring data from Germany, from the UK Health and Safety Executive (HSE), and literature data relevant for the EU, and the USA.

Results: Occupational exposure to diisocyanates is particularly relevant in: (i) C.A.S.E. applications (Coatings, Adhesives, Sealants, Elastomers), (ii) production of polyurethanes (PUs) (e.g. slab-stock foam), (iii) handling of partly uncured PU products (e.g. cutting, demoulding, spray application of foam), and (iv) when diisocyanates/PUs are heated (e.g. hot lamination, foundry applications/casting forms). Ranking of the reported data on inhalation to diisocyanate exposure at workplaces (maximum values) leads to following order: (i) HDI and its oligomers in coatings, (ii) MDI in spray foam applications, (iii) TDI in manufacture of foam, (iv) TDI in manufacture of PUs and PU composite materials, (v) TDI in adhesives, (vi) MDI in adhesives, (vii) MDI in manufacture of PUs and PU composite materials, (viii) TDI in coatings, (ix) MDI in manufacture of foam, and (x) HDI in adhesives.

Keywords: asthma; diisocyanates; occupational exposure; workplace exposure
Introduction

Isocyanates are highly reactive compounds defined by the isocyanate group, R–N=C=O, where R can be an aliphatic, cycloaliphatic, or an aromatic group. Isocyanates undergo exothermic and usually very fast reactions with nucleophiles. The most common types of isocyanates in workplaces are diisocyanates and oligomers/polyisocyanates derived thereof, which act as cross-linking agents. Their predominant use (>90%) is in the manufacture of polyurethane plastic materials (PUs, also PURs) by reacting with polyols and/or other nucleophiles like polyamines. Depending on the choice of the diisocyanate species and the polynucleophiles a wide range of polymers with diverse properties can be realized (Engels et al., 2013). Consequently, the use of diisocyanates at the workplace is just as diverse and widespread.

Typical diisocyanate-based products include:

- flexible PUs
- rigid PUs
- PU foams (rigid and flexible foam systems, e.g. mattresses)
- assembly foams (e.g. insulation panels)
- foundry cores (casting)
- coating materials (paints, lacquers, varnishes)
- adhesives and glues
- elastomers
- sealants
- prepolymers in chemical synthesis
- engineering plastics
- PU fibres

Notwithstanding their versatile material properties isocyanates are also potent respiratory sensitizers, making exposure to diisocyanates the main cause of occupational asthma in Germany and in the European Union. The number of new cases of occupational asthma resulting from exposure to diisocyanates in the EU is estimated to be more than 5000 per year. As this number is considered to be unacceptably high, Germany has prepared a restriction dossier for diisocyanates under REACH and submitted it to the European Chemicals Agency (ECHA). The restriction proposes to limit the use of diisocyanates to those workplaces where appropriate technical and organizational measures have been implemented and workers received a standardized training package to prepare them for appropriate risk management measures in a most efficient way and thus effectively lower/minimize exposures to diisocyanates and the associated risks.

In the course of the restriction dossier preparation, also a comprehensive evaluation of the current data on toxicology, epidemiology, and exposure to diisocyanates from various sources was made. Additionally, a socioeconomic impact assessment of the cost and benefits of the proposed restriction in comparison with two other risk management options and a baseline scenario (i.e. no further action) has been made. As part of the dossier preparation, an overview of the exposure situation in the EU was compiled of which the results are presented in this paper. Updates to the initial assessment were also added in the course of drafting the paper.

Scope and method of the assessment

In the restriction dossier, overall 10 different types of diisocyanates are included in a non-exhaustive list of diisocyanates covered by the restriction proposal. However, there are large differences with regard to the market volumes and used quantities of different diisocyanate species. The most important diisocyanates in terms of quantities used are:

- methylene diphenyl diisocyanate (MDI),
- toluene diisocyanate (TDI), and
- hexamethylene diisocyanate (HDI),

which together account for more than 95% of the market volume (Falcke et al., 2017). Consequently, most of exposure monitoring data are on MDI, TDI, and HDI. For this reason, the exposure assessment in the restriction dossier was limited to the three diisocyanates with
the highest market volume rather than a comprehensive list of all diisocyanates, where only very limited information on the exposure was found. Since most of the published data are based on the measurement of the diisocyanate monomers, a focus is put on them.

MDI and TDI are aromatic diisocyanates, i.e. the NCO groups are attached to an aromatic ring, leading to particularly highly reactive diisocyanate species due to mesomeric interactions. They react extremely fast with polyols under catalysed conditions to PUs. Depending on the chemical nature of the chemical building blocks (type of isocyanates and polyols), a wide range of PUs with tailored material properties can be realized (soft elastic to hard).

Aliphatic diisocyanates as HDI, i.e. where the NCO group is not directly attached to an aromatic ring, have lower reactivity, but also form more inert urethane bonds than aromatic diisocyanates. PU materials made thereof are very durable and exhibit higher ultraviolet stability as well as chemical and mechanical resistance compared with PUs based on aromatic diisocyanates. Aliphatic diisocyanates, with HDI being the most commonly used species, are generally regarded as speciality materials and account for less than 5% of the total diisocyanate consumption. The main applications are as hardeners for high quality surface coatings, where high performance is required, and as adhesives.

For the exposure assessment in the restriction dossier the following sources of information were evaluated and the assessment basically is based on these:

- Chemical Safety Reports (CSRs) of the Registration Dossiers [prepared by the Netherlands Organisation for Applied Scientific Research (TNO), where the exposure estimates are based on measurement data published by ISOPA (2012a,b, 2014a,b,c)] and on modelled data [Advanced REACH Tool (ART) v 1.0];
- Workplace measurement data (gathered from 2000 to 2011) from Germany published by the Institute for Occupational Safety and Health of the German Social Accident Insurance (IFA, 2010, 2012, 2013);
- Workplace air monitoring data from the UK Health and Safety Executive (HSE); and
- Selected and evaluated literature data relevant for the EU (this includes data since 2000 mainly from Belgium, Finland, the Netherlands, Poland, Sweden, Spain, the UK and the USA).

The literature research was done using public databases (Web of Science, Google Scholar, and PubMed) and standard search engines (such as Ecosia and Google). Selection of the literature was made according to timeliness (since 2000) and relevance for the situation in the EU, but not limited to the Member States of the European Union. The search strategy included different terms for diisocyanates (‘isocyanate’ OR ‘diisocyanate’ OR ‘MDI’ OR ‘TDI’ OR ‘HDI’) and exposure. The research was then further narrowed down to studies with a focus on workplaces or occupational settings [‘exposure’ AND (‘occupational’ OR ‘worker’ OR ‘workplace’)]. The remaining titles and abstracts were screened and about 200 matches were obtained in full text out of which the 58 quoted papers were selected.

The data from the CSRs presented here are based on the information published by ISOPA (2012a,b, 2014a,b,c) and no data from CSRs not publically available are presented for confidentiality reasons in this publication. However, the full dataset, as available in the registration dossiers (including confidential data) was evaluated for the preparation of the restriction dossier.

In the beginning of the exposure assessment, a request was made to the Institute for Occupational Safety and Health of the German Social Accident Insurance (IFA) in addition to the other member states of the European Union via ECHA for data on occupational exposure to diisocyanates. The IFA made exhaustive evaluations of workplace measurement data from Germany for MDI, TDI, and HDI by IFA (2010, 2012, 2013). These data were representative for more than 6 h of time of workplace exposure and documented in accordance to the measurement system of the German Social Accident Insurance Institutions for exposure assessment (MGU) (Gabriel et al., 2010). Grouping of the data was done according to industry groups as well as work area groups. However, for drafting the restriction dossier, the available information on occupational exposure was grouped differently, i.e. according to the information provided in the REACH registration dossiers. Out of these, the following uses were considered to be particularly relevant for occupational exposure and therefore the focus of the assessment was laid on:

1. Manufacturing of diisocyanates,
2. Use in manufacture of PUs and PU composite materials,
3. Use in manufacture of foam,
4. Use in spray foam applications,
5. Use in coatings, and
6. Use in adhesives

Since the grouping of measurement data to industry groups and area groups as done by IFA is not based on use classifications as in the registration dossiers, a simple one-to-one translation/assignment of the
MEGA data to the registered uses was not possible. The same holds true for all data provided by HSE and the publicly available data identified during the literature research. However, to allow a meaningful comparison of all identified data it was necessary to assign all data in a consistent way to the uses above. This was based on expert judgement in a way deemed most plausible. More details on the selection and grouping of the original data can be found in the restriction report (German CA, 2017).

Unless stated otherwise, the exposure levels shown in this paper are given in the same metrics as in the original sources. For most of the data, exposure values refer to the measured masses (concentrations) of the respective monomers. However, since the NCO unit is assumed to be the toxicologically relevant functional group, it is difficult to compare exposure data for the different species (MDI, TDI, and HDI) and, if applicable, their polycyanates as such (Bello et al., 2004). For direct comparison of exposure values of different diisocyanates in the discussion part, the units were therefore converted into ‘total isocyanate group’ values (µg NCO m⁻³).

Dermal exposure
In addition to inhalation exposure, there is an obligation to assess dermal exposures in the worker exposure scenarios under REACH. The dermal route is of particular relevance when assessing the overall exposure to diisocyanates, as the role of skin exposure to diisocyanates for development of respiratory sensitization and occupational asthma is discussed (Fent et al., 2009; Tsui et al., 2020). Cowie et al. conducted a comprehensive study on where diisocyanates were used in the UK and found that the airborne diisocyanate exposure were minimal in almost all of the investigated workplaces, but ‘there was nearly always potential for skin contact’ (Cowie et al., 2005). Uncured or not fully cured PU products pose a source of skin exposure to diisocyanates (Bello et al., 2007). This aspect is of particular importance since it was demonstrated that diisocyanates (e.g. from uncured resins) can deposit on and penetrate into the skin (Liljelind et al., 2010). While airborne exposures could in principle be reduced by changing to less volatile diisocyanates (like MDI and/or prepolymer) for many uses, such substitutions do not necessarily reduce dermal exposure and there are ample opportunities for skin contact in workplaces. However, the assessment of dermal exposure at workplaces is often complicated by the irregular and random occurrence of skin exposure, such as spills, contact with contaminated surfaces or during clean-up (Bello et al., 2007; Heederik et al., 2012) and quantification of dermal exposure is particularly difficult. Measurement of dermal exposure, in general, is less established than air monitoring (Kasiotis et al., 2020) and data on dermal exposure to diisocyanates in workplaces are scarce (Liu et al., 2007). There are no standardized methods available for measuring dermal diisocyanate exposure (Lockey et al., 2015). Sampling of dermal exposure to diisocyanates is challenging as the analytical methods are adaptations of the methods for airborne diisocyanates and rely on presence of unreacted NCO groups. Because NCOs (and especially mixtures of diisocyanates with polyols or amines) are highly reactive and they also react with moisture on or proteins of the skin, timing of the sampling is particularly critical (Redlich, 2010). For these reasons, dermal exposure to diisocyanates is assessed indirectly by comparison of personal air samples with corresponding biomonitoring data (Cocker, 2011; Jones et al., 2017). However, as measurement data on dermal exposure to diisocyanates are very limited, they will not be further discussed in this paper.

Results
1. Manufacturing of diisocyanates
The main process to produce diisocyanates is the phosgenation of corresponding diamines (see Fig. 1):

![Figure 1. Phosgenation of a diamine for production of a diisocyanate.](image)

Owing to the dangerous properties of phosgene and isocyanates themselves, the production processes are carried out under containment in high integrity closed systems (Falcke et al., 2017). As long as the manufacturing processes run under normal operating conditions, occupational exposure to diisocyanates at this stage is generally considered to be low compared with the uses covering the application phases. This view is supported by occupational exposure data in the exposure scenarios for MDI and TDI published
by ISOPA (2012a, b, 2014a, b, c). The exposure estimates of the diisocyanate species for manufacture of the diisocyanates as taken from the CSRs are in following ranges:

- MDI: 5.6–29 µg m⁻³;
- TDI: 5–32 µg m⁻³;
- HDI: 3–23.5 µg m⁻³.

These ranges are based on 90th percentiles of occupational hygiene measurement data and cover all contributing scenarios within the manufacturing scenarios.

2. Use of diisocyanates in manufacture of PUs and PU composite materials

Production of PU materials is the predominant use of diisocyanates and has the by far highest volume. To produce PUs the diisocyanates are reacted with macropolyols and/or other polynucleophiles and usually optional additives like catalysts, surfactants, stabilizers, flame retardants, and the like. The polyaddition reaction of isocyanates with the nucleophiles is highly exothermic. Depending on the reaction quantities and conditions, the temperature can increase considerably during the process. The chemical equation below exemplifies the general mechanism of the reaction by the example of MDI and 1,5-pentanediol (see Fig. 2). It is based on a simple single-phase PU, which is just one species of reaction products between diisocyanates and diols:

Usually the reaction is largely completed within seconds up to 30 min, whereby the isocyanate groups form urethane bonds with the polyol in the polymer backbone. However, the final curing and post-curing of PUs, where exposure to unreacted isocyanates is still possible may take up to 72 h. Occupational exposure often takes place on a regular basis in the production of PU materials and can be expected to be frequent. On the other hand, exposure control measures by means of technical controls/measures are often applied at such workplaces, so that exposure levels can be expected to be moderate.

MDI is the most used diisocyanate species for production of PU materials. Compared with MDI, TDI plays a subordinate role in the production of PU materials, except for the production of block foams, which are considered separately later in this paper. The use of and data on HDI in the manufacture of PUs on the other hand is very limited and therefore not taken into account for this use.

Table 1 provides an overview of the inhalation exposure levels to MDI and TDI in the manufacture of PU materials. The data are based on information given in the CSRs, by the Institute for Occupational Safety and Health of the German Social Accident Insurance (IFA, 2010), the Health and Safety Executive of the United Kingdom (HSE UK) (due to the confidential nature of the data provided, the following statement has to be included to this section: ‘The data is not representative of any industry partly due to bias in selection of the sites where data has been collected and is determined by HSE interest in specific substance or process. Most of the data was collected between 1986 and 1993 after which the rate of data collection reduced significantly. It should be noted that NEDB itself has an inherent bias, in that HSE Specialist Occupational Hygiene Inspectors as part of their enforcement duties obtained approximately 90% of the samples. Consequently, a tendency towards high levels of exposure would be expected, as companies with no perceived problems were generally not sampled. Even so, NEDB still contains many samples indicating low exposure (<25% of the appropriate occupational exposure limit), so the actual bias is not as large as would be expected. Whether or not NEDB should be considered as containing worst case data is debatable, but it cannot be regarded as being truly representative of occupational exposure in Great Britain given that it does not come from a random selection of workplaces and circumstances.’), and literature data on measured exposure levels of diisocyanate at workplaces in the PU and PU composite materials industry.

3. Use of diisocyanates in manufacture of foams

PU foams are generally divided by their elasticity into flexible, semi-flexible, and rigid foams. Foams are also the largest market for PUs, with flexible foams being the larger part. Both, MDI and TDI are used in the production of foams. High molecular polyols with two to six hydroxy functionalities yield flexible foams. When combined with low molecular polyols and/or amines, semi-flexible foams can be realized, while rigid foams are made of highly branched polyols with a relatively low molecular mass (Adam et al., 2005). As very different

![Figure 2. Reaction of MDI and 1,5-pentanediol.](image-url)
manufacturing processes may be applied for different foams, generally a distinction is made between slab-stock foaming processes, foam moulding, and spray foaming, the latter of which is discussed separately (see below). Foam moulding can be seen as a special case in the manufacture of PU materials as described above, where the diisocyanate-containing component is mixed with the polyol and other components just before getting injected/transferred into moulds. As the reaction mostly takes place in the moulds this also allows closed systems. Slab-stock foaming on the other hand is a process where typically two (or more) polymer components are thoroughly combined in a mixing head and are then immediately dispensed onto conveyor belts. The polymerization takes place simultaneously to the foam formation of the slab-stock and usually just takes a few seconds but depending on the choice of the reacting agents can also last up to several minutes. While technical control measures to reduce exposures are usually applicable at the mixing unit and the following part of the conveyor belt (e.g. enclosure with enhanced exhaust ventilation) the end of the lines where curing of the slab-stock still takes place are open and exposure to residual diisocyanates is highly possible. After initial curing the slab-stock is cut and further processed. However, as the curing might have not completely finished, exposure to unreacted diisocyanates might be still possible at these stages (Cummings and Booth, 2002).

Due to the relatively low vapour pressure of MDI the ranges of inhalation exposure to MDI in manufacture of foam are usually low compared with TDI in the manufacture of foams. TDI is an important component in the production of flexible foams, mostly produced in slab-stock foaming processes. HDI is not considered to be relevant for the manufacture of foams.

Data from CSRs, IFA, HSE UK, and literature for inhalation exposures to MDI and TDI during the manufacture of foams are presented in Table 2. Some attention should be given to the values from the study by Tinnerberg and Mattsson (2008), where workplace measurement data from 13 Swedish industry plants were compared before and after installation of technical measures to improve the processes in order to reduce occupational exposures. After the modernization and improvements of the plants (mostly achieved through technical measures such as better enclosures, increased ventilation, and decreased reaction speed by reducing the amount of catalyst in the reaction mixture), the exposure levels were found to be around 80% lower compared with the levels before.

4. Use in spray foam applications

As spray foam applications are linked to particularly high exposure levels, compared with uses that take place in technically controlled environments (e.g. manufacture) or where only low mechanic energies are applied and therefore no or very low aerosol formation is to be expected (e.g. gluing), this use is considered as a special case.

Spray foams are typically two-component rigid foams, with one component being an isocyanate containing hardener (usually MDI based) and the other component being a polyol formulation (including catalysts, the blowing agent, and other additives such as flame retardants, surfactants, etc.). Depending on the formulation, open and closed cell foams can be

|                | CSR 90th perc. range | IFA 90th perc. range | HSE Range | Literature data Range |
|----------------|----------------------|----------------------|------------|-----------------------|
|                | Long term/short term | (Mean)               | (Mean)     | (Mean/median)         |
|                |                      | (LoQ)                | (Mean)     |                       |
|                |                      | (N = 559)            | (N = 13)   |                       |
| MDI            | 2–38/3–76            | <LoQ=18.0            | 0.09–32.8  | <0.03–3.3 (mean 0.7)  |
|                |                      | (Mean 2.3)           |            | (N = 131) [1]         |
|                |                      | (N = 293)            |            | 0.042–7.8 (med. 3.7)  |
|                |                      |                      |            | <1–7.2 (N = 70) [3]   |
|                |                      |                      |            | <0.6–3.3 (N = 46) [4] |
| TDI            | 1–32/1–64            | 4.0–67.3             | —          | 0.08–14.6 (med. 1.2–3.9) |
|                |                      | (Mean 1.3)           |            | (N = 14) [2]          |

*LoQ was not further specified in IFA (2010) report; [1] Kääriä et al. (2001b); [2] Sennbro et al. (2004); [3] Creely et al. (2006); [4] Brzeźnicki and Bonczarowska (2015).
realized. For application, the two components are pumped from separate containers into a spray gun, which also serves as the mixing unit. The reactivity of spray foam systems is usually very high so that the finished foam forms within seconds after spraying (Adam et al., 2005). Spray foams are mainly used for insulation of buildings or industrial installations but can also serve as a specialty packing material for fragile items. This use is particularly challenging in terms of exposure reduction and risk management as aerosol formation during spraying is inevitable. In addition, spray foam installation is often carried out in dynamic workplaces (e.g. at construction sites) in which cases all equipment, including the technical measures (e.g. mobile enclosures) has to be mobile as well, which makes technical exposure reduction measures more challenging. When spray foams are applied in confined spaces (e.g. insulation of crawl spaces under basement, etc.), some technical measures like enclosures or exhaust ventilation might be not feasible at all or to a

Table 2. Occupational inhalation exposure levels of diisocyanate species (µg m⁻³) for use of diisocyanates in manufacture of foams.

| CSRs | IFA | HSE | Literature data |
|------|-----|-----|-----------------|
|      | 90th perc. range | 90th perc. range | Range | Range |
|      | Long term/short term | Long term | (Mean) | (Mean/median) |
| MDI  | 6–29/12–58 | <LoQ⁻⁴.2 | 0.03–0.17 | • <0.6 (N = 26) [4] |
|      | (Mean 1.7) | (N = 1013) | (N = 3) | • <0.6 (N = 20) [5] |
| TDI  | 1–32/1–64 | <1.3–72.8 | 0.06–9.0 | • <0.2–230 (N = 96) [1b] |
|      | (Mean 4.7) | (N = 110) | [Short term: 1.37–45.0] | • 0.08–39.9 (med. 1.2–31.4) (N = 140) [2] |
|      |            |            | (N = 13) | • 0.2–58.8 (med. 4.0–9.8) (N = 26) [4] |
|      |            |            |            | • 0.2–58.9 (mean 3.6–26.3) (N = 20) [5] |
|      |            |            |            | • 46.5–73.6 (med. 62.9) b, 5.0–86.5 (med. 12.5) c [6] |
|      |            |            |            | • <7.2–17.4 (N = 26) [7] |
|      |            |            |            | • 4.2–142 (mean 31.1) (N = 21) [8] |
|      |            |            |            | • <71 (49 workers) [9] |
|      |            |            |            | • 0.03–3.1 (5 workers) [10] |

a LoQ was not further specified in IFA (2010) report.
b Before risk management measures (RMM) improvements.

Table 3. Occupational inhalation exposure levels (µg m⁻³) for use of MDI in spray foam applications.

| CSRs | IFA | HSE | Literature data |
|------|-----|-----|-----------------|
|      | 90th perc. range | 90th perc. range | Range | Range |
|      | Long term/short term | Long term | (Mean) | (Mean/median) |
| MDI  | 6–29/11–58 | <LoQ⁻¹ | 0.03–200 | • 0.07–2.47 (N = 36) [10] |
|      | (Mean 1.9) | (N = 33) | (N = 8) | • 10–570 (N = 61) [11] |
|      |            |            |            | • 70–2050 (N = 13) [12] |
|      |            |            |            | • 11–591 (med. 54.8) (N = 94) [13] |
|      |            |            |            | • <LoQ–770 [14] |
|      |            |            |            | • <4.6–410 [15] |
|      |            |            |            | • 30–90 (experimental set) [16] |
|      |            |            |            | • 0.9–123.0 (GM 13.8) (N = 62) [17] |

a LoQ was not further specified in IFA (2010) report; [10] Jones et al. (2017), levels given as total NCO (µg NCO m⁻³); [11] Crespo and Galán (1999); [12] Lesage et al. (2007); [13] Roberge et al. (2009); [14] RPS (2014); [15] Robert et al. (2014); [16] Puscasu et al. (2015); [17] Bello et al. (2019).
limited extend. Risk management and exposure reduction therefore largely depend on personal protective equipment (Allport et al., 2003).

According to information from literature and the registration dossiers, the only diisocyanate used for spray foam applications is MDI. Table 3 summarizes the inhalation exposure levels to MDI during spray foam applications based on data from CSRs, IFA, HSE UK, and selected literature.

5. Use in coatings

Coatings are often applied to surfaces by spreading or by spraying, and as during these applications often aerosols are generated and/or splashes occur, it is therefore often linked to particularly high exposures in comparison to uses with no (or minimal) aerosol/droplet formation.

PU coatings can be one-component (one-pack) or two-component systems. One-pack paints containing free isocyanates are usually high molecular prepolymer of polyols with excess isocyanate groups that undergo a cross link reaction with atmospheric moisture. Two-component systems form the 'conventional' PU coatings and paints and are by far the most important systems (Adam et al., 2005). The cross-linking constituents are polyisocyanates based on TDI, HDI, isophorone diisocyanate (IPDI), MDI, or 4,4'-methylenedicyclohexyl diisocyanate (HMDI). Solvent borne curing agent solutions have an isocyanate content of 5–16% (w/w), while solvent free may have up to 30% (w/w) isocyanate (Stoye et al., 2000). The other component of the paint contains polyols and/or polyamines as well as additives such as pigments, catalysts, and solvents. Both components are mixed immediately before application (preferably in an equimolar ratio). Due to their outstanding properties (especially high mechanical resistance, chemical resistance, and light and weather resistance) PU coatings are the systems of choice for protecting coatings like vehicle finishes and refinishes and in the building sector (floor coatings, anti-corrosion coatings, etc.). Aliphatic isocyanates (especially HDI) are important basic materials for protective and decorative coating systems, especially in vehicle body repair, where HDI-based spray paints are widely used.

All of the diisocyanate species covered in this assessment (MDI, TDI, and HDI) are used in coatings. Inhalation exposure to MDI as well as TDI during application of diisocyanates containing coatings is found to be relatively low compared with systems based on the more volatile HDI. Table 4 summarizes the inhalation exposure levels to MDI, TDI, and HDI for the use in coatings based on data from CSRs, IFA, HSE UK, and found literature relevant to the topic.

6. Use in adhesives

PU adhesives are used in a broad scope of applications and products ranging from extremely stable and weatherproof woodworking and construction glues to bonding automotive parts (e.g. windshields). The

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Table 4. Occupational inhalation exposure levels (µg m⁻³) for use of diisocyanates in coatings.

|     | CSRs 0th perc. range | IFA 90th perc. range | HSE Range | Literature data Range |
|-----|----------------------|----------------------|-----------|-----------------------|
|     | Long term/short term | Mean range | Long term | Range | (Mean/median) |
|     | Long term | (Mean) | (N = 685) | — | 0.06–8.1 \[10\] |
| MDI | 6–29/11–58 | <LoQ—18.8 | (Mean 2.4) | — | 0.35–208 \[10\] |
|     | (N = 809) | (Mean 1.3) | (N = 1221) | (Short term: 0.82–245 000 (N = 47)) | (Short term: 0.82–245 000 (N = 47)) |
| TDI | 1–35/1–70 | <1.3–6.0 | (Mean 1.3) | — | 421–423 \[10\] |
|     | (N = 809) | (Mean 1.3) | (N = 15) | 0.02–57.6 °(med. 0.08–7.4) (N = 95) | 0.02–946.7 °(GM 87.2) \[21\] |
| HDI | 110–430 | <2.3–12.0 | (Mean 2.3) | 0.35–208 | 421–423 \[10\] |
|     | (N = 15) | (N = 1221) | (N = 88) | 0.02–946.7 °(GM 87.2) \[21\] |

°LoQ was not further specified in IFA (2010) report.

°Data are presented for monomeric HDI; \[10\] Jones et al. (2017), levels given as total NCO (µg NCO m⁻³); \[18\] Sparer et al. (2004); \[19\] Pronk et al. (2006a); \[20\] Fent et al. (2009); \[21\] Bello et al. (2020).
Adhesives can be two-component or one-component systems, which themselves can be solvent-based, waterborne (aqueous dispersions), or solvent free (granulates, dry powders). They can be processed and/or cured at ambient temperatures or at elevated temperatures (from 50–80 to 180–200°C). With respect to potential exposure, it was shown that both the content of isocyanate monomers and the processing temperature have a significant impact on emissions (Cano et al., 2015).

Besides the broad spectrum of applications, there is only limited data available for the use of diisocyanates in adhesives. In fact, many of such systems, especially those that are used at room temperature, are often low emission glues.

Table 5 provides an overview of the inhalation exposure levels to MDI, TDI, and HDI in the use of adhesives as given in the CSRs, the MEGA evaluations by IFA, and as published by Brzeźnicki and Bonczarowska (2015).

### Biological monitoring

In addition to inhalation data some studies also provide biological monitoring data. Biological monitoring of diisocyanates is based on the analysis of isocyanate adducts with haemoglobin or albumin in the blood or the determination of corresponding diamines in urine or in plasma (Cocker, 2011; Świerczyńska-Machura et al., 2015). The most common way for biomonitoring is using urine samples and looking for corresponding diamines of the diisocyanates, i.e. methylene diphenyl diamine (MDA) for MDI, toluene diamine (TDA) for TDI, and hexamethylene diamine (HDA) for HDI. However, as the amines are not specific markers for diisocyanates, exposure to the corresponding diamines has to be ruled out since, otherwise, the results can be biased (Gries and Leng, 2013).

Table 6 provides summaries of the biological monitoring data for MDI in the manufacture of PUs and PU composite materials, for TDI in the manufacture of foam, and HDI in coatings.

As biological monitoring assesses the total burden of workers, it is not possible to distinguish between the exposure pathway and sources contributing to the burden. Diisocyanate metabolites can often be detected in biological monitoring samples even if the corresponding air monitoring measurements were below the limit of detection (Creely et al., 2006). On the other hand, biological monitoring can be used to assess also behavioural aspects regarding the effectiveness of risk management measures like proper use of PPE and efficacy of training interventions (Jones et al., 2013). However, recently the need for a harmonized approach for biological monitoring of isocyanates and to establish a baseline against which the effectiveness of the proposed restriction can be evaluated was highlighted (Scholten et al., 2020).

### Other uses

As stated before, diisocyanates are used in a wide range of sectors and products, not all of which are covered by this assessment. It is clear that other diisocyanates like 1,5-naphthylene diisocyanate or IPDI can also have an impact on the workplace exposure (e.g. Tinnerberg et al., 2014). Potentially high exposures can also arise in other uses than those described here (e.g. in foundry applications (Liljelind et al., 2010)). However, as information on those diisocyanates and/or uses was found

### Table 5. Occupational inhalation exposure levels (µg m⁻³) for use of diisocyanates in adhesives.

|     | CSRs 90th perc. range | IFA 90th perc. range (Mean) (N = 533) | HSE Range Long term | Literature data Range |
|-----|------------------------|----------------------------------------|---------------------|-----------------------|
| MDI | 5–43/9–87              | <LoQ=6.5 (Mean 2.8)                     | —                   | <0.6–5.2 (N = 20) [4] |
| TDI | 1–5/1–70               | <1.3–48.2 (Mean 1.9)                    | —                   | —                     |
| HDI | —/—                    | <2.3 (N = 294)                         | 0.8–1.0 (N = 20)    | [4]                   |

LoQ was not further specified in IFA (2010) report; [4] Brzeźnicki and Bonczarowska (2015).
to be too limited to allow further assessments of these exposure situations such aspects are not taken into the scope of this study.

**Discussion**

The potential for occupational exposure to diisocyanates is determined by intrinsic substance properties (e.g. volatility) or to the processes involved in their handling. Volatility is one key determinant for the potential for inhalation exposure and related to the molecular size. Diisocyanates with a low molecular weight, i.e. TDI and HDI, have significant vapour pressures already at room temperature, which can lead to relatively high concentrations at the workplace (i.e. the respective diisocyanates can be detected and the concentrations are measurable, which is not the case e.g. for MDI). This trend can be also seen in the evaluations of the MEGA database for diisocyanates made by IFA as shown in Fig. 3 (IFA, 2010, 2012, 2013). In addition, higher temperatures increase the vapour pressure thus the tendency of diisocyanates to

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**Table 6.** Overview of biological monitoring data of exposed workers.

| Biomonitoring metabolite (μmol mol⁻¹ creatinine) | Airmonitoring concentration (μg m⁻³) | Reference |
|-------------------------------------------------|-------------------------------------|----------|
| MDI in the manufacture of PUs and PU composite materials |                                      |          |
| 0.015–1.38 (MDA in urine) | <0.03–3.3 (64% <0.03) | Kääriä et al. (2001b) |
| • 90th perc. 6.29 nmol l⁻¹, median 1.34 (MDA in urine) | | Sabbioni et al. (2007) |
| • 90th perc. 0.177 pmol g⁻¹ Hb (haemoglobin adduct MDA) | | |
| <LOD–12.64 (MDA in urine) | <1–7.2 | Creely et al. (2006) |
| • 0.1–0.2 (MDA in urine, during working day) | | Henriks-Eckerman et al. (2015) |
| • <0.1 (MDA in urine, day off) | | |
| <LOD–12.64 (MDA in urine) | | Robert et al. (2007) |
| • <LOD–14.1 (MDA in urine) | | Gries and Leng (2013) |
| • <LOD–16.2 pmol g⁻¹ (ABP-Val-Hyd in blood) | | |
| • 0.5–8.4 μg l⁻¹ (MDA in urine) | 0.04–9.7 | Tinnerberg et al. (2014) |
| • 0.4–19.4 μg l⁻¹ (MDA in plasma) | | |
| TDI in the manufacture of foam |                                      |          |
| 0.05–39 (total TDA in urine) | <0.2–230 | Kääriä et al. (2001a) |
| <0.05–1.6 (total TDA in urine) | <3.5–8.4 | Austin (2007) |
| Before RMM improvements: | | Tinnerberg and Mattsson (2008) |
| • 2.9–27.2 μg l⁻¹, median 7.0 (2,4-TDA in plasma) | 46.5–73.6, median 62.9 | |
| • 8.2–62.1 μg l⁻¹, median 30.8 (2,6-TDA in plasma) | | |
| After RMM improvements: | | |
| • 0.5–2.0 μg l⁻¹, median 1.0 (2,4-TDA in plasma) | 5.0–86.5, median 12.5 | Święcistyńska-Machura et al. (2015) |
| • 2.0–11.8 μg l⁻¹, median 4.0 (2,6-TDA in plasma) | | |
| • <LOD–3.9 (total TDA in urine) | 0.2–58.9 | |
| • <LOD–5.4 (total TDA in urine) | 0.03–3.1* | Jones et al. (2017) |
| HDI in coatings |                                      |          |
| • 1.9–146.2 (HDA in urine) | 0.03–28.8 (HDI monomer) | Pronk et al. (2006b) |
| • <LOD–21.0 (HDA in urine) | 0.003–179 (HDI monomer) | Gaines et al. (2010) |
| Before SHAD*: | | Jones et al. (2013) |
| • 90th perc. 1.34 (HDA in urine) | | |
| After SHAD: | | |
| • 90th perc. 0.6 (HDA in urine) | | |
| • <LOD–1.0 (total HDA in urine) | 421–423* | Jones et al. (2017) |

*Exposure levels given as total NCO (μg NCO m⁻³).

*Safety and Health Awareness Day.
become airborne is correspondingly higher in hot processes. In addition, handling diisocyanate-containing products at high temperatures can lead to thermal degradation, which can release the original monomeric diisocyanate and other low molecular isocyanates or fragments during thermal decomposition processes.

Figure 3. 90th percentile values of air concentrations of MDI, TDI, and HDI for work area groups for selected uses from IFA reports (MEGA database; IFA, 2010, 2012, 2013), converted into total NCO units (µg NCO m⁻³).

Figure 4. Air concentrations of MDI, TDI, and HDI for uses from all sources presented before, converted into total NCO units (µg NCO m⁻³) (note the logarithmic display of the exposure levels).
(Simon et al., 1988; Delebecq et al., 2013; Wang et al., 2013). Very high exposures are found when a process is used where high levels of aerosols are formed (mostly spraying). Diisocyanate-based paints and varnishes are often used for spray painting; especially in vehicle body refinishing HDI-based spray paints are ubiquitously frequently used and lead to significant occupational exposures. Spray foaming, especially when applied to greater surfaces (e.g. insulation of ceilings/walls) also leads to high aerosol release (Christensen et al., 2014). Fig. 4 shows the found exposure levels for the different diisocyanates and sectors as presented before on a semi-logarithmic scale.

Many commercial products are not only based on diisocyanate monomers but can consist predominantly of oligomers and/or prepolymers (e.g. in HDI-based coating systems). Nevertheless, the vast majority of available exposure data is based on the measurement of the respective diisocyanate monomer, whereas measurement of ‘total isocyanate group’ values (Bello et al., 2004) has only recently become more common. It is understood that the overall risk of exposure to isocyanates is likely to be underestimated if these are not included. However, as written above, most of the available measurement data are for monomeric species, thus the focus in this paper is on these.

With regard to the dermal pathway, skin contact with products containing isocyanates (e.g. uncured PU foams, paint, or glue splashes) is reported to be a significant route of exposure (Austin, 2007) and there is almost always potential for dermal exposure when handling isocyanate containing formulations or reaction products thereof which are not fully cured (Cowie et al., 2005). For example, Creely et al. found that urinary levels of isocyanate metabolites of workers with observable dermal exposure were over two times that of workers who did not have evident skin contact (Creely et al., 2006). It was also observed that both dermal and inhalation exposures correlate significantly with urinary diisocyanate metabolite concentrations (Gaines et al., 2021).

Grouping of the exposure data in a ranking order according to the reported bandwidths of inhalation exposure levels results in the following order, inhalation exposure levels to:

- HDI and its oligomers in coatings—from 0.003 up to 5566.3 µg m⁻³ (90th percentile), total range: 0.003–245 000 µg m⁻³
- MDI in spray foam applications—from limit of quantification (LoQ) up to 2050 µg m⁻³
- TDI in manufacture of foam—from LoQ up to 203 µg m⁻³
- TDI in manufacture of PUs and PU composite materials—from LoQ up to 67.3 µg m⁻³
- TDI in adhesives—from LoQ up to 48.2 µg m⁻³
- MDI in adhesives—from LoQ up to 43 µg m⁻³
- MDI in manufacture of PUs and PU composite materials—from LoQ up to 32.8 µg m⁻³
- TDI in coatings—from LoQ up to From LoQ up to 35 µg m⁻³
- MDI in manufacture of foam—from LoQ up to 29 µg m⁻³
- HDI in adhesives—from LoQ up to 1.0 µg m⁻³

The uses found to give rise to the highest inhalation exposure levels are HDI (and its oligomers) in coatings and MDI in spray foam applications. In both uses the diisocyanate resins are applied by spraying, confirming that high exposures are to be expected when diisocyanates are applied in high energy processes and aerosols are formed. Relatively high inhalation exposure levels are also found for some uses of TDI such as in the manufacture of foam as well as in the manufacture of PUs and PU composite materials and, in parts, for the use in adhesives. The exposure levels of MDI on the other hand are for all of these uses significantly lower. These findings are in line with the expectation that use of less volatile diisocyanates leads to lower inhalation exposure levels.

While for the majority of the discussed uses, most of the measured data were quite low (near or below the LoQ), it has to be stressed that measurement of airborne diisocyanates is technically challenging. The target compounds are usually highly reactive and some measurement methods are less sensitive to this, hence resulting in systematic underestimation of the actual exposure levels at workplaces (Streicher et al., 2000, 2002; Bello et al., 2004; Brandt et al., 2013). Relatively high exposure levels can also occur in uses that appear to be well controlled at the first sight (e.g. TDI in adhesives). Such relatively high levels of inhalation exposure seem to occur in an unpredictable and unexplainable manner in all sectors and uses but could not be explained by analysis of the statistical data [see part B.9.9 of the Annex XV restriction dossier for diisocyanates (ECHA, 2017)]. The situation is further complicated by the fact, that different air sampling methods exist that measure differently (only monomers or total isocyanate mass concentrations, etc.), making comparison of measurement values between different studies more difficult (Bello et al., 2004).

It should also be highlighted that isocyanate adducts can often be detected in biological monitoring samples even if the corresponding air monitoring measurements were below the limit of detection (e.g. Creely et al., 2006). This might be explained by significant
uncertainties in (i) the contribution of the respective exposure pathway to the total burden (inhalation vs. dermal) and (ii) uncertainties in the air monitoring data themselves as measurement of airborne diisocyanates and particularly of peak exposures is technically challenging and may underestimate actual exposure levels.

**Conclusion**

The estimated number of annual incidences of diisocyanate-related occupational asthma in the EU is in the range from 2350 to 10150 cases. (ECHA, 2017) Annex XV restriction report—Diisocyanates—Part B. Helsinki, Fi: ECHA.) The result of the exposure assessment(s) for diisocyanates at European workplaces lead to the conclusion that risks are not sufficiently controlled for a proportion of situations as shown above. Occupational exposure to diisocyanates is particularly relevant in:

- C.A.S.E. applications (Coatings, Adhesives, Sealants, Elastomers),
- production of PUs (e.g. slab-stock foam),
- handling of partly uncured PU products (e.g. cutting, demoulding, spray application of foam),
- when isocyanates/PUs are heated (e.g. hot laminating, foundry applications/casting forms).

With regard to the data for inhalation exposure it has to be kept in mind that measurement of airborne diisocyanates is technically challenging and may underestimate actual exposure levels. In addition, peak exposures to diisocyanates are particularly difficult to detect, which can also lead to an underestimation of exposure.

To address the risks of occupational asthma caused by diisocyanates, the German competent authority for REACH has proposed a restriction of products containing more than 0.1% by weight of diisocyanates (individually and in combination) under the EU’s REACH Regulation in 2016. The restriction was recently published in the EU Official Journal (European Commission, 2020) and will apply after a transitional period of 3 years from 24 August 2023. Diisocyanates are defined therein as ‘O=C=NR=R=N=C=O, with R an aliphatic or aromatic hydrocarbon unit of unspecified length’. In the opinion of the authors this definition also applies to oligomers/prepolymers as long as they have two terminal NCO units. It is supposed that this will lead to increased substitution efforts to safer products, i.e. products containing less than 0.1% (w/w) of diisocyanates. However, a derogation from this ban can be made if industrial and professional users receive an obligatory standardized training on good working practices and risk management. Work training has been shown to be an effective measure to reduce occupational exposure to diisocyanates (Jones et al., 2013) and the training required by the restriction aims to improve compliance and make working with diisocyanates safer.

**Conflict of interest**

This work was part of the authors’ job routine. Their employer, the German Federal Institute for Occupational Safety and Health (BAuA), therefore funded it. As the data presented in the manuscript were part of the proposal for the restriction of diisocyanates under REACH, the authors were not only driven by scientific motivation, but also have a regulatory interest in the topic. Urs Schlüter, who is member of the Committee for Risk Assessment (RAC) at ECHA has declared his conflict of interest (CoI) in the topic when the restriction was discussed by RAC. The authors have sole responsibility for writing and content of the manuscript.

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