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Dermal exposure assessment in occupational epidemiologic research

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Recognition of the importance of skin exposure in industrial settings has steadily increased over the last few decades. Whereas the 1980s saw widespread recognition of the importance of skin exposure in the field of industrial hygiene, the 1990s saw the beginnings of a systematic characterization of the field (1, 2). Unfortunately, the growing attention to dermal exposure has rarely been reflected in the field of occupational epidemiology. Although it is widely recognized that exposure to hazardous substances may occur by inhalation, ingestion, and dermal contact, the focus of exposure assessment in occupational epidemiology traditionally, explicitly, or implicitly has been on inhalation exposures. This situation is even true for contaminants, for example, pesticides, polycyclic aromatic hydrocarbons, and polychlorinated biphenyls (PCB), and solvents and diseases such as dermatitis and skin cancer, for which dermal exposure is known to contribute significantly to internal dose or the disease.

Valid and reliable exposure assessments are crucial in occupational epidemiology because new risks are likely to be lower than those seen historically and they are therefore more difficult to detect. They are also crucial because the emphasis in epidemiology has shifted from qualitative risk identification to quantitative risk assessment, which incorporates exposure-response relationships (3). Thus it is important that exposures through multiple routes (ie, oral, dermal, and inhalation) and from different sources (ie, occupational, environmental, and dietary) be accurately assessed in epidemiologic research. Inaccurate and imprecise exposure estimates may lead to a loss of power, precision, and attenuation in health risk estimates, depending on the type of error structure (4, 5).

Proper exposure assessment strategies for estimating dermal exposure in epidemiologic research depend on the chosen study design, for example, prospective, cross-sectional, retrospective (cohort, case-referent), and the

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health outcome of interest (eg, chronic, acute, systemic, local). In prospective and cross-sectional studies, one has the opportunity to collect current dermal exposure data and information on possible determinants that could subsequently be used to estimate group or individual dermal exposure levels. However, in retrospective studies one depends on historical exposure information or expert judgment. The health outcome under consideration determines the locations where measurements should be made. In the case of local effects like hand dermatitis, dermal exposure is important at the body location of interest, while for systemic effects the total dermal body exposure is the key for estimating internal dose.

We conducted a literature survey to identify dermal exposure assessment methods that have been applied in epidemiologic research. Using the identified studies and recognized determinants of dermal exposure and uptake as a basis, we describe factors that should be considered and postulate methodologies that could be applied for dermal exposure assessment in occupational epidemiologic research. Attention is given to estimating the intensity and duration of exposure, the exposed surface area, personal, temporal and spatial variability in dermal exposure, and uptake. The focus of this paper is primarily on historical dermal exposure assessment in relation to systemic effects, but many of the considerations described apply to other study designs and local effects as well.

Exposure metrics and analyses

The dose surrogates commonly used in occupational epidemiology are exposure intensity, exposure duration, cumulative exposure, and average exposure over the work history. For chronic health outcomes cumulative exposure is generally thought to be the most appropriate measure of exposure (6). The dose surrogate most preferred, however, would be an estimate of the cumulative dose of the active substance or metabolite at the target organ. Biomarkers of exposure potentially reflect internal dose and have the advantage that they integrate exposure from all sources through all routes of exposure, including the dermal route. However, for most chemical exposures, no biomarkers of the (historical) internal dose are readily available, and therefore their current use is limited in (retrospective) exposure assessment. As a result, substances absorbed dermally will require, at least for the time being, the development of separate exposure estimates for both inhalation and dermal exposure.

Separate estimates of inhalation and dermal exposure in an epidemiologic study can be used in several ways. One could be to combine them into a single estimate of internal dose. This procedure would require, however, quantitative estimates of the two exposure routes in the same measurement units and information on the respiratory and dermal absorption rates. For most substances, such information is not readily available. Moreover, it is at least questionable whether accurate quantitative estimates can be derived for the dermal route with current measurement methods. [For an overview of dermal exposure assessment techniques see Brouwer et al (7), Cherrie et al (8), and Soutar et al (9).] Therefore, an integrated estimate of the internal dose cannot, in many cases, be derived easily.

The alternative is to develop separate estimates for inhalation and dermal exposure and use them as independent estimates of exposure or use them in a stratified analysis. This approach allows the simultaneous use of quantitative inhalation estimates and semiquantitative or qualitative dermal estimates in the same analysis. The stratified approach does, however, have consequences for the statistical power of the study, and its applicability is, in many studies, limited to exposures with a moderate-to-high prevalence.

Assessment of dermal exposure in occupational epidemiologic studies

Historically, several different exposure indices have been used for inhalation exposure in occupational epidemiologic research, varying from simple surrogates of exposure to sophisticated measures of internal dose (3, 10) and from qualitative (exposed, unexposed) and semiquantitative (low, medium, high) estimates to quantitative measures. Because the latency of most chronic diseases is now within the time frame in which historical airborne measurements are available, deterministic and stochastic modeling of historical inhalation exposure data has become possible and therefore enables an evaluation of quantitative exposure-response relationships (11–13).

Unfortunately, the evaluation of dermal exposure is less developed. This lack of development is, in part, due to the scarceness of quantitative exposure data, and standardized methods for quantifying dermal exposure to many substances are still lacking (14). Only a limited number of epidemiologic studies can be found in the peer-reviewed literature that has assessed dermal exposure in any way. It is, however, difficult to present a complete overview of the methods and indices used because much of the published literature does not explicitly indicate which exposure routes were considered in the exposure assessment. In fact, even when dermal
exposure is identified as being considered, it is usually not clear how it was evaluated.

**Qualitative estimates**

Several qualitative estimates have been applied for dermal exposure in epidemiologic research. These estimates have almost exclusively been based on assumed determinants (subjective or objective) of dermal exposure, which were subsequently used as qualitative exposure proxies (yes;no). For example, in an epidemiologic study on acute pyrethroid poisoning among cotton farmers, dermal exposure was measured, and exposure determinants were identified (15). These determinants, contamination of clothing (yes;no), concentration of pyrethroid in the application solution (%), and the occurrence of leakage or blockage of the sprayer (yes;no), were used in the epidemiologic analyses, and each was found to be related to neurological symptoms. A similar approach was used in a study of signs and symptoms of pesticide toxicity among Indonesian farmers (16). In this study each spray operation was observed, and variables thought to influence exposure were recorded. The following four types of variables were consequently considered: the frequency and mechanics of the spray operations (applications/week, mixing with bare hands, leaky equipment, spraying against the wind), contact of the clothing and body with the spray solution (percentage of spray operations, hands or feet becoming wet during mixing or pouring, splashed body during application), the type of clothing worn (footgear, gloves, eyeglasses, long pants, short-sleeved shirt, headgear), and the kind of chemicals sprayed and their management (pesticides, hazard grade of the World Health Organization, other chemicals). The number of applications per week, the use of hazardous pesticides (grade 1B and II, established by the World Health Organization), and skin and clothing becoming wet with the spray solution were found to be associated with neurobehavioral signs and symptoms.

**Semi-quantitative estimates**

Semi-quantitative estimates of dermal exposure were used in a study of adverse health effects among tannery workers (17, 18). The assessment of skin exposure was based on a three-point scale of no contact with the agent of interest, moderate contact (infrequent skin contact with the agent, eg, contact occurred during specific activities that were not part of the daily work routine), and frequent contact (frequent skin contact, eg, regular contact was unavoidable due to the activities performed daily). In a study of workers at a coal liquefaction plant, a similar ranking system was used, based primarily on the likelihood of skin contact with various products as assessed by expert judgment (minimal, low, medium, high) (19). Self-reported frequency of direct dermal contact with PCB was used in a study of clinical and metabolic abnormalities to divide PCB-exposed workers into four exposure groups (eg, never, rarely, occasionally, and frequent contact) (20). In a study of cancer mortality among workers exposed to acrylonitrile, a dermal score was developed by multiplying the percentage concentration of acrylonitrile in the liquid with the estimated frequency of dermal contact (21, 22).

**Quantitative estimates**

Only a handful of examples of studies developing quantitative estimates of dermal exposure were found in the literature. Brouwer et al (23) developed an algorithm for calculating a cumulative exposure index (milligrams/lifetime) in a study of the health effects of pesticides in the flower-bulb industry in The Netherlands. The algorithm included the application method (tractor- or boom-, backpack-, bike-spraying), the method of mixing and loading (direct tank filling, pouring, or scooping), the method of bulb disinfection (manual or mechanical dipping), the application rate (milligrams per hectare), the bulb acreage (hectares), the number of applications, the number of bulbs disinfected per year (number of containers), and a protection factor for personal protective equipment (range: 0 = no protection to 1 = complete protection) (24). In a study of Pliofilm™ workers, exposure to benzene was calculated on the basis of the concentration of benzene in the cement being used, the number of skin contacts with the cement per day, the surface area of the contacted skin (square centimeters), and the contact time (hours per day) (25–27). Dermal uptake (milligrams per kilogram per day) was subsequently calculated using these dermal exposure estimates, the dermal absorption rate (milligram per square centimeter per hour), and body weight of the individual (kilograms). The absorbed dose was then converted to an estimate of the corresponding airborne concentration in parts per million based on the respiratory absorption rate of benzene.

**Basic parameters of dermal exposure**

The qualitative and quantitative parameters used in the occupational exposure algorithms across studies are remarkably similar and rely on or reflect the following three basic estimates: (i) the concentration or mass of the contaminant (eg, intensity), (ii) the exposed surface area, and (iii) the duration or frequency of the exposure. Not surprisingly, these basic parameters have also been
described in several conceptual models for the assessment of dermal exposure (1, 28). In the following paragraphs these parameters are discussed in somewhat more detail.

**Concentration or mass of the contaminant**

The passage of hazardous substances through the skin is governed by a diffusion process that is driven by the concentration of the contaminant on the skin (29). It has been suggested, therefore, that the concentration of the contaminant on the skin, rather than the mass, determines the internal exposure, and, therefore, the assessment of this quantity should provide a more reliable indicator of exposure than mass (30). Although the mass of the contaminant may not be directly important for dermal uptake, it is a measure of the reservoir of contamination on the skin that is potentially available for dermal uptake. The distinction between the concentration and mass of the contaminant on the skin may, therefore, be trivial when the concentration of the material handled is constant (eg, single source with constant concentration) or when there is not much material (ie, mass) available for uptake. However, in many occupational settings, these requirements will not be met, and, in these circumstances, it can be argued that the metric that should be assessed is the actual concentration of the contaminant on the skin, rather than the mass of the contaminant on the skin.

However, current monitoring techniques only measure mass (eg, skin loading). Therefore, an estimate of the concentration of the contaminant on the skin can only be assessed subjectively at present, using the concentration of the contaminant in the source or on contaminated surfaces as an estimate of the concentration of the hazardous substance on the skin. Information about the concentration of the contaminant in the source can be retrieved historically, at least in certain occupational settings, as was shown in two epidemiologic studies (21, 25). Retrospective exposure assessment studies should therefore attempt to collect as much information as possible about the concentration of the active ingredient in the source(s) or on contact surface(s) over time, together with information about the actual mass of the contaminant on the skin.

**Surface area exposed**

The surface area exposed, together with the concentration or mass of the contaminant, determines the total amount available for uptake at a certain time point. The first consideration in an evaluation of the surface area exposed is the identification of the parts of the body exposed. The identification depends largely on inherent process characteristics (eg, aerosol generation, ejection of particles, and splashes), which describe the interaction between the source, process, and equipment with the work environment and the dermal exposure pathway (eg, deposition, direct contact, immersion), which describes the interaction of the workers’ skin with the work environment.

Several studies using visualization techniques have examined the surface area exposed under certain exposure scenarios. These studies were almost exclusively restricted to the agricultural setting and to spray painting activities (31–33), and therefore the possibility to generalize the results are limited. Nonetheless, these studies clearly showed that the distribution of the dermal contaminant is, in most cases (if not all), inhomogeneous across the body. The notion of an inhomogeneous distribution of contaminants across the body was recognized long before it was actually visualized properly (34). To overcome this problem, it became general practice to divide the body into a number of specific body locations to meet the assumption of a homogeneous exposure distribution within the defined body locations (35, 36). Although, there is no limit to the number of body locations that one could define, for practical reasons, the body is often divided into nine (35) or ten (36) distinctive regions, namely, the head, trunk (chest and back), upper arms, forearms, hands, legs, thighs, lower legs, and feet. It is assumed that each location is homogeneously exposed and that the area exposed is equivalent to the skin surface area of that particular location. However, Fenske (32) demonstrated, among pesticide applicators, that the actual proportion of the skin surface of specific body regions receiving exposure is relatively small (4–22%) and highly variable. Estimation of the actual surface area exposed within an a priori defined body location is, therefore, imperative for an accurate estimate of dermal exposure. The estimation is, however, difficult if no visualization data are available or can be collected (because the tracer cannot be added to the source of exposure) or the occurrence of dermal exposure is random like, for instance, a spill. In this case, it may only be possible to derive an estimate of the exposed surface area by carefully considering the involved exposure pathways and work practices. In general, it can be assumed that dermal exposure originating from the immersion or deposition of airborne particulates on the skin will result in a more uniform exposure distribution for a specific body location than exposure originating from splashes and direct skin contact with contaminated surfaces.

The exposed surface area not only depends on external factors, however, but also on subject-specific parameters, including the use of (protective) clothing, individual work practices, and the actual skin surface area of the person in question. The use of protective clothing [the term protective clothing is used here, but
it also includes gloves and regular clothing, which can provide protection from dermal exposure as well (37–38).) is an important determinant of the actual skin area exposed. The most basic evaluation of this parameter is covered and uncovered skin. For the skin covered by protective clothing, the area exposed and mass of the contaminant on the skin depends on the protection factor provided by the clothing. However, little is known about the field effectiveness of protective clothing, which depends both on the selection (eg, type of material, degree of permeation, fit) and use (eg, replacement, wear and tear, cleanliness). A complicating factor is that clothing can potentially increase the uptake of the chemical through the skin by occlusion (39). Occlusion describes the process of increased hydration and temperature of the skin due to the coverage of the contaminated skin by clothing (or any other material), which thereby increases absorption secondarily. In addition, clothing may actually pull contaminants inside the clothing as a result of a “pumping effect” caused by normal body movements during work (40). Furthermore, studies have demonstrated that hands cannot be considered protected from exposure even if “appropriate” gloves are worn. This lack of protection may be due to contamination on the interior surface of the gloves, removal of gloves when fine hand movements is required, and handling of the outside of the gloves when putting them on or taking them off (37, 41). It could well be, therefore, that, although clothing can substantially reduce dermal contact with chemicals by reducing both the area exposed and the mass of the dermal contaminant, it may not decrease uptake by as much as one would expect on the basis of a laboratory-evaluated protection factor (42). Conservative estimates of the protection provided by protective clothing should be applied, as proper use or proper functioning often cannot be assumed.

The total skin surface area obviously differs from person to person. These individual differences are seldom taken into consideration, except in a study by van Rooij et al (43). In this study each body location was assigned a fixed percentage, based on the area distribution of the body surface of a “standard man” (36), of the person’s total body surface area estimated on the basis of the subject’s weight and height (44). Although this approach may seem unnecessarily precise, inclusion of subject-specific estimates for the total body surface area of a person may, in some circumstances, be important because it has been estimated that up to a threefold difference in surface area can be found among adult men (45). However, it is unclear whether differences in a person’s total body surface area automatically results in differences in the actual surface area exposed. In some environmental settings this assumption probably holds true (eg, for showering or swimming in contaminated waters). In these situations it seems reasonable to assume a constant relationship between a person’s total skin surface area and the exposed surface area. However, for many scenarios within industrial settings, such a direct relationship cannot be assumed, particularly if the surface area of a contaminated tool or product limits the area of contact with, for example, the hands. Extrapolation of dermal exposure from the estimated skin surface area may be necessary only when a homogeneous exposure distribution across the body or within a body location is a justifiable assumption and a direct association between the total individual skin surface and the exposed surface area can be assumed (eg, immersion).

Thus retrospective exposure assessment should focus on identifying likely exposed body locations on the basis of visualization studies or the involved exposure pathways and use of protective clothing. In a second step, the actual skin surface area exposed within exposed body locations needs to be assessed. Incorporating differences in the individual skin surface area seems less important, given the underlying assumptions and limited effect.

Frequency and duration of exposure

In addition to estimating the concentration or mass of the contaminant on the skin and the surface area of the exposed skin, two components of time (eg, frequency and duration of dermal contact) have to be considered. Depending on the mass transport process involved and the penetration rate of the contaminant, either the frequency or duration of the exposure will be more important. In the case of an event-based exposure process (eg, intermittent exposures or sporadic contact transfer from surface to skin through accidental contact) in combination with a high percutaneous penetration rate of the contaminant (ie, quick absorption), the exposure will be driven by the frequency of contacts with the contaminant (figure 1). In other words, if the percutaneous penetration rate is not rate limiting, the component of time can be described by the frequency of dermal contacts with the contaminant. In contrast, if the mass transport process is continuous (eg, deposition of airborne particulates, condensation of fumes on the skin, solvent vapors) or the uptake of the contaminant through the skin is rate limiting or uptake is limited by the rate of dissolution of the contaminant in the sweat layer (eg, particulates), the time component is best described by the duration of exposure. In this case, it is important to realize that, in contrast to inhalation exposure, dermal exposure does not necessarily stop after the workshift has ended, but continues until the contaminant is removed from the skin either by absorption, evaporation, mechanical loss, or deliberate removal from the skin. Recently, Kissel & Fenske (46) proposed a model for estimating dermal absorption for re-entry workers.

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The model can easily be applied to other occupations (figure 2). It considers retention of the contaminant on the skin during (i) a net accumulation phase (ie, the workshift) with continuous exposure until a washing event and (ii) a postworkshift period (until the washing event), over which the external load declines as the material is absorbed (neglecting evaporation and mechanical losses from the skin). This model assumes a 100% removal efficiency of the contaminant from the skin by a single washing event. However, even if the skin is washed, it may still be contaminated. For example, two studies found pesticide residues on the skin of pesticide workers 1 to 3 days after exposure (47, 48). In addition, some reports have shown incomplete removal of PCB from the skin by washing (49) and actual enhancement of dermal uptake for hydrocortisone due to washing of the hands (50). The duration of the postworkshift period should therefore carefully be considered for the exposure of interest and, if necessary, adjusted according to knowledge of removal efficiencies for the particular contaminant.

Furthermore, it should also be recognized that, although hands may be washed directly after the workshift, areas such as the arms and face are less likely to be cleaned immediately. Wearing contaminated clothing beyond the workshift can also prolong dermal exposure, as it serves as a continuous reservoir of contaminant and may continue to be a source even after the clothing has been laundered (51). The fact that dermal exposure (and consequently dermal uptake) can continue through residual contamination on the skin or on clothing after the actual activity or work has stopped is unique to dermal exposure and should be carefully evaluated in the assessment of dermal exposure.

Figure 1. Concept for using frequency or duration as the components of time. The dotted line and consequent gray area indicates the point where the penetration rate of the contaminant is not rate limiting relative to the exposure scenario (ie, intermittent to continuous).

Figure 2. Illustrative exposure scenarios with explicit treatment of the time dependence of absorption after the workshift. Exposure is accumulated over the workshift (shown as 8 hours). Scenario A: effective washing event immediately after finishing work; scenario B: effective washing event some time after the workshift (shown here as 2 hours); scenario C: partial removal of contaminants (75% effective) by washing event some time after the workshift (shown here as 3 hours); scenario D: no washing event after the workshift. For the last three scenarios (B–D) the chemical remaining on the skin at the end of the shift or after partial decontamination of the skin continues to be absorbed. [Figure modified based on an example developed by Kissel & Fenske (46)]
For retrospective dermal exposure assessment, it is thus important to collect information about the duration and frequency of exposure through careful consideration of work practices. After the percutaneous penetration rate is considered, an evaluation can be made of whether duration or frequency of exposure is the best component of time. In addition, information on skin and clothing decontamination procedures can be used to estimate the contribution of postworkshift exposure.

**Variability issues**

**Exposure**

Recognition of the personal, temporal, and spatial variability in exposure concentrations is imperative for the design of efficient and effective sampling strategies, for the interpretation of existing exposure data, and for assigning exposure estimates in an epidemiologic survey (52). The distribution of exposure over the body (ie, spatial variability) has been studied extensively, at least in the relative sense. In contrast, a formal evaluation of temporal and personal variability in dermal exposure measurements has been almost entirely absent. Recently, a database of dermal exposure measurements (DERMDAT) was constructed to analyze these two types of dermal exposure variability (14). These analyses showed median values of the total and within- and between-worker geometric standard deviations of 2.55, 1.98 and 1.47, respectively, which are strikingly similar to those published for respiratory exposure (53).

From this database, several factors were identified that increased the between-worker variance, for example, working in an indoor environment, localized sources, and a random measurement strategy (14). Interestingly, these factors only influenced the between-worker variance and not the day-to-day variance. That none of the studied factors explained the day-to-day variance suggests that dermal exposure is event-based in many occupational settings, with a certain probability that a contaminated surface is touched by a worker or a splash or spill lands on the skin. Additional justification for the event-based scenario of dermal exposure is that the sampling method partly determined the temporal exposure variability of the hand (14). Patches, which measured exposure at only a small part of the hand or wrist, exhibited more day-to-day variability than the hand wash method, which measured the total hand and wrist exposure. This finding suggests that the occurrence of dermal exposure at a specific spot on the body depends, at least partly, on chance.

The third variance component for dermal exposure is the between-body location component (ie, among skin locations, spatial variability). This source of variability is generally considered to be the most prominent component. Interestingly, in the DERMDAT database, the day-to-day variance component almost disappeared when the variability of the between-body location was taken into account (14). This finding suggests that the day-to-day variability in dermal exposure is important for a specific body location, but not for the overall average of total body exposure. A valid assessment of total dermal exposure thus depends on an accurate assessment of both the exposure distribution across the body and the intensity of the exposure at different body locations. Dermal exposure estimates based on measurements on a single day or a single body location should, therefore, be interpreted with caution.

In epidemiology, it is often assumed, for practical reasons, that workers employed in the same job at a given location are uniformly exposed. This assumption has led to observational schemes for classifying workers into homogeneous exposure groups on the basis of job title, location, and other identifiable features of the work environment (54). In figure 3 the cumulative distribution of the between- and within-worker values of the $R_{0.95}$'s (the ratio of 97.5th and 2.5th percentiles of the log-normal distribution of the between ($R_{0.95}$) and within ($R_{0.95}$) worker distribution), respectively, are shown for both inhalation and dermal exposure, using the WAUNC (53) and the DERMDAT (14) databases. The exposure groups were defined by job title and factory (eg, homogeneously exposed groups) and, in case of dermal exposure, also by body location. It can be seen that, in general, the range in the day-to-day variability exceeds the range in the between-worker variability for both inhalation and dermal exposure. According to Rappaport (55) an observational group is uniformly exposed if $R_{0.95}$ is less than two, while the Health and Safety Executive uses a criterion for uniformity that translates to $R_{0.95}$ being less than four (56). For inhalation exposure, 25% of the groups had 95% of the individual mean exposures within a factor of two, and 50% had a $R_{0.95}$ of less than four. For dermal exposure 40% of the groups had a $R_{0.95}$ within a factor of two, and 49% had a $R_{0.95}$ of less than four. Although the DERMDAT database has very limited power for generalization, it can be deduced that considerable variation in dermal exposure levels among persons within “homogeneous exposure groups” (as defined by job title and body location) exists. Thus, as is the case for inhalation exposure, exposure assessors should not rely blindly on observational schemes to guarantee that groups of workers are uniformly exposed by either dermal contact or by inhalation. Retrospective exposure assessment strategies should therefore adopt methods to address the variance components, if possible, by carrying out pilot studies to estimate the contribution from these sources of variance.
and identifying their determinants to optimize exposure assessment methods, strategies, and grouping schemes. These pilot studies should perform dermal exposure measurements at various body locations of several persons on multiple days.

Uptake

Dermal absorption depends not only on the time-dependent concentration of the contaminant on the skin surface, but also on compound-specific factors (eg, polarity, chemical structure, volatility), the presence of absorption enhancing concomitant exposures, and skin-specific factors. Compound-specific factors are inherent to the chemical under consideration and therefore are unlikely to lead to differences in percutaneous penetration between workers or groups of workers. On the other hand, concomitant exposures to ethanol and other short-chain alkanols, polyethylene glycols, acetone, and other solvents have been found to enhance the percutaneous penetration of several compounds (40, 57) and could result in substantial differences in uptake between (groups of) workers.

Skin-specific factors include the condition of the skin, skin temperature, skin thickness, skin perfusion, and lipid-protein makeup. The latter three differ substantially by anatomic region and thus affect dermal absorption rates. The effect of the anatomic region on percutaneous absorption has revealed reasonably similar results for a variety of chemicals, the head, neck, and scrotum being indicated as the more permeable body areas (58). To determine what parts of the body are most important in occupational exposure assessment, we calculated mean relative absorption rates per body location relative to the forearm by taking the ratio between the percutaneous penetration rate of each body location to that of the forearm (table 1). This value, multiplied by the actual surface area, results in a permeability-corrected skin surface area, which indicates the importance of each location as a function of the surface area and the percutaneous penetration rate. The scrotal area was not included in the calculation because it is not a standard measurement site in occupational dermal exposure assessment (65). The corrected surface areas indicate the importance of the upper body region (eg, the head, neck, and upper arms), which accounts for more than 40% of the permeability-corrected surface area. It should be noted that the estimated average percutaneous absorption coefficients are based on a very limited dataset and that differences in the percutaneous absorption rates among anatomic regions cannot be assumed to be independent of the physicochemical properties of the tested chemicals (eg, mostly lipophilic compounds) (38). Although the results of these studies seem reasonably consistent in their relative ranking of the different body locations, the exact estimates are imprecise. Nevertheless, they could be used when no better data are available. This finding suggests that, if a limited number of body locations is to be measured, they should be selected on the basis of the anticipated exposure distribution, taking into account the permeability of the specific body locations. For many industrial settings, the basic set of dermal measurements should therefore include the hands, head, neck, and shoulders.

Dermal uptake can be increased by several factors, of which occlusion and adverse skin conditions are the most important (66). The effect of occlusion has been shown to increase in vivo percutaneous absorption for several pesticides by a median factor of 3.5 (range 2.3–9.2) (67). Occlusion could therefore change substantially the absorption rates of the various body locations covered by clothing or gloves (39). This effect could well
counterbalance most of the differences in percutaneous absorption rates between the different body areas indicated in Table 1. However, in general, it is difficult to take occlusion into account due to the limited knowledge on the occurrence of occlusion and its actual effect on percutaneous absorption under field circumstances.

Adverse skin conditions (eg, hydration, dryness, skin diseases) have been shown to affect percutaneous absorption under both laboratory and field circumstances (67–71). Of the adverse skin conditions, skin abrasions (eg, dermatitis, cuts, burns) have been shown to have a potentially dramatic effect on percutaneous absorption. The median effect of skin abrasions on in vivo percutaneous absorption for several pesticides was estimated to be 6.3 (range 2.6–9.5) (67). However, for some exposures, like parquat, this effect could even be more dramatic, with almost 0% absorption through intact skin but 100% absorption through damaged skin (72). It seems, therefore, essential to take the presence of adverse skin conditions into account when systemic uptake through the skin is considered, especially since skin diseases are estimated to account for 9–35% of all occupational diseases (73–74).

In prospective and case-referent studies, questions should be asked about the occurrence of adverse skin conditions (ie, dermatitis, cuts). However, several studies have shown that standard questionnaires for identifying skin diseases or symptoms have limited use in epidemiologic studies because the overall validity is moderate with generally a moderate-to-high specificity (range 71–99%) but lower sensitivity (range 31–76%) (75–80). Recently, questionnaires using pictures of visual symptoms of skin diseases have been developed, and it is anticipated that their validity, and thus their applicability for epidemiologic research, will improve. In retrospective cohort studies, one could rely on medical records to assess skin diseases. However, relatively few people with skin complaints seek medical assistance (81), and, therefore, this approach seems futile. In circumstances in which no direct information on skin disorders can be obtained, the overall risk for the occurrence of skin diseases in a particular occupational group, if known based on the peer review literature, could be used as an exposure modifier at the group level. However, because the magnitude of the effect seems to be related to the physicochemical properties of the substance and because the relationship between exposure and adverse skin conditions (linear, nonlinear) is unknown, it would be preferable to include the occurrence of adverse skin conditions as an interaction term with the dermal exposure estimates in epidemiologic analyses.

### Table 1. Mean relative percutaneous penetration rates and permeability-corrected surface areas by body region. (AM = arithmetic mean)

| Body location | N | Compounds | Percutaneous penetration ratios | Surface area (cm²) | Percentage of total surface area | Permeability-corrected percentage of total surface area |
|---------------|---|-----------|--------------------------------|-------------------|---------------------------------|--------------------------------------------------------|
| Head          | 8 | a to g    | AM 2.55 SD 1.34 Range 0.95–4.75 | 1300              | 62                              | 11.8                                                   |
| Neck          | 2 | a, c      | AM 8.45 SD 6.43 Range 3.9–13.0 | 1210              | 58                              | 7.8                                                   |
| Forearms      | 11| a to k    | AM 1.00 SD 0.55 Range 0.8–1.9  | 820               | 3.9                             | 4.0                                                   |
| Hands         | 3 | a, b, c   | AM 2.04 SD .          | 2910              | 14.0                            | 21.1                                                  |
| Upper arm     | 1 | k         | AM 1.22 SD 0.47 Range 0.62–2.10 | 6840              | 32.8                            | 29.7                                                  |
| Trunk         | 10| a to j    | AM 1.00 SD 0.23 Range 0.75–1.21 | 3820              | 18.3                            | 13.6                                                  |
| Upper leg     | 3 | h, i, k   | AM 0.48 SD 0.08 Range 0.42–0.53 | 2380              | 11.4                            | 4.0                                                   |
| Lower leg     | 2 | c, k      | AM 0.83 SD 0.56 Range 0.14–1.60 | 1310              | 6.3                             | 3.9                                                   |
| Feet          | 5 | a, b, c, h, i | AM 26.90 SD 21.4 Range 11.8–42.0 | - | - | - |
| Scrotum       | 2 | a, c      | AM 26.90 SD 21.4 Range 11.8–42.0 | - | - | - |

* Number of compounds for which percutaneous penetration data were available for the particular body location (58, 60–64).
*b Chemicals studied: (a) parathion, (b) malathion, (c) hydrocortisone, (d) benzoic acid sodium, (e) caffeine, (f) benzoic acid, (g) acetyl salicyl acid, (h) fentanyl, (i) sufentanil, (j) ketoprofen, (k) polycyclic aromatic hydrocarbons.
* The percutaneous penetration ratio of the particular body location relative to that of the forearm calculated for each study and then averaged per body location.
* Adapted from reference 59.
* The permeability-corrected percentage of the total surface area was calculated by multiplying the surface area (cm²) of a particular body location by the mean percutaneous absorption ratio for that location divided by the total permeability-corrected surface area (cm²). The total permeability-corrected surface area was calculated as the sum of all individual permeability-corrected body locations and equaled 28181 cm². For example, head: [(1300 × 2.55)/28181] × 100 = 11.8.

### Applicability to exposure assessment for epidemiologic studies

#### Variability issues

Proper assignment of dermal exposure estimates in epidemiologic studies requires knowledge about the level...
of exposure (eg, intensity, exposed surface areas, duration), exposure variability (personal, temporal, spatial), and variability in uptake. The importance of accounting for exposure variability in inhalation exposures is widely recognized, and methods to account for these sources of variability in epidemiologic exposure assessment strategies have been well described in the literature (54, 82–85). Approaches have been developed to address the variability between groups, the variability among people within groups, and the variability over time. Careful consideration of these sources of variability increases the efficiency and sensitivity of the assessment and reduces bias.

These approaches also can be used in dermal exposure assessments, and many of the conclusions for inhalation assessment apply to dermal exposure assessment. The between-body location variance is, however, unique to dermal exposure, and methodologies have not yet been established to incorporate this source of variability into the design of exposure assessment strategies. Nevertheless, in principle, the between-body location variance can be considered a component of the day-to-day variance. When the spatial variability component is low, relatively few body areas would have to be measured to arrive at a good estimate of total body exposure.

In a study by Vermeulen et al (86) the relation between the dermal exposure of individual skin regions and total body contamination was investigated among rubber workers. A strong correlation (Pearson correlation \( r=0.87 \) and \( r=0.95 \) for production workers and technical engineers, respectively) was found between exposure at the wrist and total body exposure; this finding suggests that, for this particular occupational setting and epidemiologic purposes, dermal exposure could be estimated solely from the exposure of the hands. This assumption allowed the investigators to reduce the number of body locations that needed to be measured to estimate dermal exposure levels and facilitated the collection of repeated measurements from more persons to address personal and temporal variability issues. Careful consideration of the minimum number of measurements necessary to arrive at an accurate estimate of total body exposure, based on spatial autocorrelation analyses, could significantly increase the effectiveness and efficiency of the exposure assessment strategy.

Variability in the percutaneous penetration rates among body areas is generally small, only the head, neck, and scrotal areas having substantially different absorption rates than other locations. When these body locations are likely to be significantly exposed, the differences in their absorption rates should be taken into account. An example of such a situation can be found in a study by de Cock et al (87), who studied fruit growers applying pesticides. In this study an association was found between tetrahydrophthalimide in urine and captan exposure on the neck, but not with the total dermal exposure. In exposure situations in which only one area predominates exclusively or no significant exposure occurs to the head, neck, or scrotal area, it seems reasonable to assume a single percutaneous absorption rate for all body areas, as is common in models used for the dermal uptake of environmental contaminants (88–90) or merely the use of the external dermal exposure level as the estimate of exposure.

**Exposure assessment**

Methodologies that can be used to estimate dermal exposure levels depend strongly on the available exposure data. If comprehensive measurement data are available, (retrospective) exposures can be estimated using stochastic modeling of the dermal measurement data. Unfortunately, historical measurements are scant, except perhaps for pesticides, for which over 100 studies have been published that provide quantitative exposure data. It has to be noted, however, that probably many more dermal measurement data exist that have never been published in peer-reviewed scientific journals. However, even with the over 100 pesticide studies, historical assessment of the pesticide data is problematic due to the range of pesticides measured, the use of different measurement techniques (patches versus whole body), the measurement of different body locations, exposure measurements outside (potential exposure) and inside (actual exposure) clothing, and differences in extrapolation procedures. In addition, these studies often lack a comprehensive evaluation of the determinants of dermal exposure, and this lack limits their value in assessments of historical dermal exposure levels. Another source of information that could be used to develop pesticide exposure scores is the Pesticide Handlers Exposure Database (PHED) (91) and the European Predictive Operator Exposure Model (EUROPOEM) (92). These pesticide exposure databases reflect exposures to operators (mixers or loaders and applicators) under “representative” field circumstances, by which is meant highly standardized use scenarios. There is, therefore, some concern about its relevance to actual exposure situations because of the controlled, almost experimental, conditions under which the application occurs (eg, best case monitoring). However, relative comparisons between different application methods and various types of protective equipment can be inferred from these databases.

Statistical modeling of current dermal exposure measurements to identify and quantify exposure determinants may be a more fruitful approach. Although issues concerning differences in measurement techniques and strategies apply as well, “successful” examples of this approach can be found in the literature. These studies applied a single measurement method and collected...
expert judgment relies heavily on the familiarity of studies seems, at least for the time being, limited. Given the limitations of the model, its applicability to combine the expert opinions, as differences within expert panels can be considerable (107). One approach to minimize this variability would be to develop a formal structure for evaluating dermal exposures based on a priori rigorous evaluation criteria.

Concluding remarks

The assessment of dermal exposure is still in its infancy, and many problems have been discussed with regard to the measurement and estimation of dermal exposure. The inclusion of quantitative dermal exposure estimates in retrospective cohorts will therefore be very limited for many years to come. However, this limitation should not automatically lead to the conclusion that the dermal route should not be included in current studies, because potentially severe misclassification could occur by the omission of a significant exposure route (108). For example, in the case of pesticides, 90–95% of the absorbed dose has been found to be from dermal exposure (34). The extent and the structure of the misclassification will depend greatly on the relation between inhalation and
dermal exposure estimates and the way the exposure estimates are used (continuous, categorical). Several investigations have studied the correlation between dermal and airborne exposure estimates and found correlation coefficients ranging from as low as 0.06 to as high as 0.99, with a median correlation coefficient of $r=0.4$ (37, 43, 57, 86, 94, 107, 109, 110). As these statistics indicate, assuming that the optimal grouping strategy for inhalation and dermal exposure is the same is often not justified, because exposure sources and pathways for the two exposure routes may have distinct characteristics.

We have given an overview of the methods applied in epidemiologic studies, described the basic parameters to be considered when dermal exposure is assessed, and described procedures that can be applied in current epidemiologic research. The relative importance of these parameters has been indicated in Table 2. It goes without saying that the importance of these parameters depends heavily on the specific scenario under consideration, but the presented ratings can be used as a starting point in the exposure assessment procedure. In addition, we have indicated the current level of knowledge about these parameters. Overall, it can be anticipated that, in the coming years, retrospective dermal exposure assessment for epidemiologic research will generally be based on expert judgment and, to some degree, on process-specific exposure models. It should be stressed that, to increase the comparability across studies that rely on expert judgment, an attempt should be made to document the derivation of quantitative dermal exposures, even if only basic and limited information is available.

Field studies collecting quantitative dermal exposure data and statistical modeling to identify exposure determinants will, however, be imperative for progress in this field. In addition, the identification of the structure and magnitude of exposure variability (personal, temporal, and spatial) will provide information for more efficient and more effective measurement and grouping strategies. Some degree of standardization may be beneficial for current measurement methods, and it would increase comparability between studies. We urge that special attention be given to studies comparing different measurement methods, identifying dermal exposure determinants, evaluating exposure variability, and developing grouping strategies.

**Table 2** Overview of considerations for the estimation of (historical) dermal exposure levels in occupational epidemiologic research.

| Parameters to be considered                                           | Relative importance | Current level of knowledge |
|-----------------------------------------------------------------------|---------------------|----------------------------|
| Intensity of the contaminant                                          | ++++                | –                          |
| Concentration of contaminant on skin                                 | ++                  | +                          |
| Mass of contaminant on skin                                           | ++                  | ++/++                      |
| Exposure determinants to develop process-specific algorithms          | ++                  | +                          |
| Location and surface area exposed                                    | ++                  | +                          |
| Identification of exposed areas by visualization techniques or multiple sample sites | ++++                | +/-                        |
| Observation of exposure pathways (immersion, airborne deposition, direct contact) | ++++                | +/-                        |
| Observation of process characteristics (aerosol generation, ejection of particles, splashes) | ++++                | +/-                        |
| Determinants of exposure distribution and surface area exposed (eg, clothing) | ++                  | –                          |
| Individual’s total body surface                                      | +                   | +++                        |
| Duration and frequency of exposure                                    | ++++                | –/+                        |
| Frequency and duration of exposure                                    | ++++                | –/+                        |
| Skin decontamination behavior or procedures                           | ++++                | –/+                        |
| Clothing decontamination or procedures                                | +/-                 | –/+                        |
| Exposure variability                                                  | ++++                | –/+                        |
| Personal, temporal and spatial variability                            | ++++                | –/+                        |
| Uniform exposure groups                                               | +++                  | –/+                       |
| Uptake modifiers                                                      | ++                   | –/+                       |
| Adverse skin conditions                                               | ++                   | –/+                       |
| Occlusion (clothing, work climate)                                    | +/-                  | –/+                       |
| Body location, especially head, neck and scrotal area                 | ++                   | –/+                       |

* = mildly, ++ = moderately, +++ = very, ++++ = extremely important.  
* – = poor; + = limited; ++ = fair; +++ = good.

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