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A roadmap towards a globally harmonized approach for occupational health surveillance and epidemiology in nanomaterial workers

Running Title: Roadmap for harmonizing nanomaterial worker studies

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1 Abstract

Objective: Few epidemiological studies have addressed the health of workers exposed to novel manufactured nanomaterials. The small current workforces will necessitate pooling international cohorts. Method: A roadmap was defined for a globally harmonized framework for the careful choice of materials, exposure characterisation, identification of study populations, definition of health endpoints, evaluation of appropriateness of study designs, data collection and analysis, and interpretation of the results. Results: We propose a roadmap to reach global consensus on these issues. The proposed strategy should ensure that the costs of action are not disproportionate to the potential benefits, and that the approach is pragmatic and practical. Conclusions: We should aim to go beyond the collection of health complaints, illness statistics or even counts of deaths: the manifestation of such clear endpoints would indicate a failure of preventive measures.

2 Clinical significance

Epidemiological research is needed to confirm that proposed protective measures are effective. This roadmap proposes a strategy to assess materials, exposure, health endpoints, populations, ethical-societal aspects, study designs, data collection and analysis, and interpretation of the results while ensuring that the approach is cost-efficient, pragmatic and practical.
3 Why harmonize world-wide?

Nanomaterials, due to their novel physico-chemical properties, are making their way into businesses and products. After a phase of "nano-hype" with few concrete nanomaterials applications, development of real-life products and processes including nanomaterials is now progressing steadily. Already today, about 1% of all companies are involved with nanomaterial production or development (1, 2), although each of them only has a few workers actually dealing with nanomaterials. However, with the growing number of existing and novel nanomaterials, and their applications and incorporation in products, an increasing number of workers is expected to become exposed to these materials throughout the products' life cycles. In anticipation of this growing market, and in response to debates about the potential health risks of nanomaterials, research is being carried out worldwide in an effort to ensure that there are no adverse health effects from working with nanomaterials (3). The early identification of potential health and safety issues indeed allows us to redirect investments for safer future steps in technology development.

Most current research projects in this field focus on exposure measurement techniques and toxicological research that identifies mechanisms of effect and no adverse effect levels in test systems (4). However, occupational health studies of exposed nanomaterial workers are needed to confirm that the derived levels are safe and that the applied safety and preventive health measures are effective. Epidemiological studies of the health of workers producing and using the classic carbon black and amorphous silica nanomaterials were carried out in the late 1980s/mid 1990s (5, 6). However, the workforce in individual countries and companies (1, 7) is still small, and there is a large diversity of nanomaterials, which poses challenges for the recruitment of sufficiently large cohorts. This leads to the necessity for pooling cohorts internationally for consideration of novel nanomaterials. The implementation of such studies is currently hampered by: i) few standardised, accurate and reliable approaches for estimating exposure; ii) large variability in nanomaterial metrics and co-exposures; iii) the lack of information on health effects and biomarkers; iv) lack of large production forces in this emerging industry, rather a considerable proportion in research and development; v) questionable statistical power related to currently small workforce sizes and short latency for disease occurrence; and v) a changing regulatory framework without harmonized registration systems for workers employed in these industries (8).

Numerous different approaches could be taken for monitoring the health effects of nanomaterials, including occupational health reporting schemes, health surveillance, health
risk appraisal surveys and self-reporting questionnaires (3, 9, 10). Acceptability of different approaches will be strongly influenced by the social contexts and regulatory backgrounds in different countries. These determine where responsibilities lie, and who ensures that the required steps in both exposure assessment and health effect monitoring are implemented.

In this report, the most pressing needs in terms of global harmonization were identified as follows:

(a) Outline the range of information necessary for epidemiological studies in nanomaterial workers;

(b) Evaluate exposure data and models useful for pooled exposure assessments and provide task-based exposure profiles for specific nanomaterials;

(c) Link different worker activities and task profiles to job titles, job descriptions, and industries on the one hand and to production processes on the other;

(d) Evaluate health effects and biomarkers for use in future occupational health reporting schemes and epidemiological studies, based on knowledge derived from toxicology studies of the subject nanomaterial and inference from epidemiological studies of other nanomaterials;

(e) Define the requirements for epidemiological studies and identify suitable cohorts;

(f) Provide recommendations for small and medium enterprises (SMEs) and associated organisations regarding early assessment and management of possible risks for nanomaterial workers and the setup of exposure and health effect registries.

Consequently, the rationale for creating this is that before embarking on any large-scale, human-health monitoring studies, it is of paramount importance to determine their feasibility and whether they will ensure useful, reliable results. Basic preparatory work is thus needed to evaluate and analyse existing knowledge, data, and practices for exposure and health effect assessment.

### 4 The roadmap development procedure

This roadmap is a product of expert discussions that started in 2008 with the definition of the NanolImpactNet project plan (FP7-Grant 218539, www.nanoimpactnet.eu). In 2009, during a NanolImpactNet workshop in Lausanne, Switzerland, participants were asked to answer a series of questions (Table 1) related to the design of occupational health surveillance studies
for both short- and long-term monitoring of the health of workers exposed to or handing nanomaterials.

Table 1 Initial questions about issues that a workers' health surveillance program needs to take into account:

1. Assessing and recording exposure
   a. How to gather qualitative and quantitative information on exposure (nature of worker tasks, areas of workplaces, materials being handled, control measures in place, field studies etc)
   b. How to record and access this information

2. Assessing and monitoring health / health surveillance
   a. How to define a harmonized approach for data recording and access
   b. How to identify "effects" to be monitored, preferably early markers of effect or indicators of biological response as opposed to final outcome (disease, death)

3. Information dissemination
   a. How to raise awareness amongst relevant health professionals
   b. How to disseminate the results of the monitoring to workers and management
   c. How to provide support to policy makers and other decision takers

The insight gained from the workshop was condensed into a report (11), which was shared with the audience of a NIOSH-organized meeting in Keystone, Colorado, in 2010 (12). Realizing the potential need for world-wide harmonisation, a global group of WHO Collaborating Centres and other partners was formed to extend that report's ideas to create this roadmap. The roadmap presented here will also serve as a guide for nanomaterial-related activities of the co-authoring researchers when preparing their contributions towards the implementation of the WHO-Action plan 2012-2017.

5 The roadmap

The present roadmap proposes a strategy to gain global acceptability by many different actors for occupational health studies in the field and to harmonize the collection and storage of data to yield maximum benefits for all the parties involved. It covers three main domains: exposure, health, and framework conditions related to risk management and study design (Figure 1).
Global agreement is needed to outline the scope of information that should be collected for epidemiological studies in nanomaterial workers to address the following issues:

- Exposure data measurements and models need to be evaluated and validated so that they are linkable to worker activities, task profiles, job titles, job descriptions, and industries on the one hand, and to production processes on the other.
- Potential health concerns need to be identified and biomarkers of effects need to be assessed for detecting short- and long-term effects.
- To facilitate global harmonisation, differences in risk assessment and management, but also data protection philosophies are important when designing epidemiological studies, identifying suitable cohorts and setting up exposure and health effect registries.

Feedback loops are needed in each domain of activity. Furthermore, the groups working on harmonising the exposure, health, and risk management and study design domains need to be tightly linked and considerable cross-talk will be needed to ensure a meaningful and efficient harmonisation of methods and approaches.

The roadmap focuses on occupational health studies in nanomaterial workforces. However, data collected following such a harmonized scheme will also be helpful to identify adequate risk management strategies, and to evaluate whether existing protective measures are efficient. The data will thus be interesting to industries using or producing nanomaterials, safety and health experts consulting for these companies, trade unions and politicians who need to respond to concerned constituents, and governmental and international bodies dealing with occupational health and safety issues surrounding nanomaterials.

### 5.1 Exposure to nanomaterials

Analysis and recording of exposure to nanoparticles is an important pre-requisite for the assessment of their health effects in workers. However, in most instances, this still represents one of the weakest stages in the risk management process.

A key requirement for all studies is good quality data on exposed workers. Measurements need to be feasible and based on direct factual observations, taking into account the use of protective equipment and ventilation of facilities and processes that may limit exposures.
Ideally these exposure studies would be carried out by an occupational hygienist able to link data to activities and work processes, but in reality, this may not always be possible. A good nanomaterial exposure assessment should follow the steps below:

1. Quantitative and qualitative assessment of potential exposures, including recording the number of companies and workers handling nanomaterials, quantities of materials used, handling procedures and whether nanoparticles are dispersed or in powder form during handling, for example.

2. Identification of potential sources of nanoparticle emission and exposure, noting that “emission sources” are different from “exposure sources”, i.e. depending on the risk management measures in place, working with a source of nanoparticles does not necessarily correlate with worker exposure to the engineered nanoparticle of interest. This process should include an awareness of work practices.

3. Agreement on and measurement of possible exposure parameters according to a harmonized protocol for all sites. This includes particle metrics and co-pollutants, and contextual information such as ventilation parameters, room size, protection strategies in use, duration of tasks, etc.

4. Identification of descriptors for job titles, activities, processes and industry sectors, so as to build a multidimensional job-activity-exposure matrix for the estimation of personal exposure with statistical models.

5. Incorporation of the results into industrial, national or international exposure registries.

Before exposure registries can start receiving data, a consensus needs to be reached on: the nature and minimal quantity of exposure data and contextual information required; how the data should be collected and managed; where it should be stored (e.g. in exposure registries, which may need to be country-specific); and on who is allowed access and under which conditions (13). In addition, cross-talk with health specialists and epidemiologists is necessary to ensure that the parameters allow for the examination of potential links between exposure and effect.

5.1.1 Qualitative assessment of exposure

Exposure to nanomaterials is plausible at all stages of their life-cycle from formulation and production to application in products, use and disposal. The levels and duration of exposure, as well as the number of individuals affected will vary at each stage (14). Ideally studies would be based on sufficiently large populations, exposed to specific, relevant, well-characterised nanomaterials at different levels and for different durations. However, this is a real challenge and not easily achievable in today’s nanotechnology industries. This is due to
a range of complex issues including costs, identification of workers handling nanomaterials, isolation of effects from specific manufactured nanomaterials versus those from other nanomaterials or chemicals involved in the process, and the fact that identified “high” exposures may lead to immediate corrective actions under the current precautionary stance being recommended by regulators and many other stakeholders (15).

In reality, although the nanotechnology industries are growing, the number of nanomaterial workers is still quite small, the materials handled are heterogeneous, and the exposures diverse and continually changing. Information to identify where workers may be exposed can be derived from investigations of the types of materials companies are using, the types of workplaces and protective measures in place (16, 17), as well as larger, representative surveys to identify the sectors of industry that are using nanomaterials and that can provide information about the quantities involved (1). Currently, research facilities are one of the places where many people seem to be handling novel nanomaterials (18, 19). However, universities may not be ideal places for initial monitoring studies, because of highly varied work activities and the turnover of personnel. Manufacturing sites with fewer short-term changes may be preferable; where there is potential for different processes to give rise to different potential exposures (high versus low); where best practice in minimizing exposure to chemicals in general is already in place and a culture of safety exists. Exposure levels will also be industry-specific. An initial pragmatic solution may be to build on existing field studies, such as those being conducted by national occupational safety and health institutes (e.g. the US National Institute for Occupational Safety and Health (NIOSH)) (7) or by larger research projects (e.g. EU Framework 7 studies investigating exposure and protection strategies in the workplace, of which an overview is given in the NanoSafety Cluster Compendium (4)).

5.1.2 Agreement on parameters and measurement of exposure

There is ongoing debate about which nanomaterial exposure parameters should be recorded (20). Good quality data are required, but it is not feasible to take large amounts of measurement equipment into busy workplaces on a routine basis; measurements need to be ongoing and practical. Furthermore, few validated data on which physico-chemical characteristics make nanomaterials hazardous or dangerous are available. As this debate continues, pragmatic approaches to exposure measurements are needed. For research purposes, a very detailed analysis of a wide range of parameters may be required, including mass, particle number, surface charge, surface reactivity, chemical composition, and characterization by electron microscopy (21, 22). However, in many other instances (e.g. for checking the efficiency of protective measures), measurement of mass, number, surface or charge of particles may be sufficient, regardless of the size distribution of the nanomaterials.
Ideally, any strategy should rely on practical personal and/or real-time monitoring of exposure to nanoparticles in areas where exposure has occurred or is very likely to occur at high concentrations. The best measurement methods are those that are specific to or highly correlated with the manufactured nanomaterial of interest.

Around the world, leading research groups are using slightly different instruments and methods, and thus measuring different aspects of exposure to particles in the ultrafine or nanoscale range (23). Whilst the methodological details differ slightly (e.g., demonstrated by (20, 22)) approaches such as those followed in NEAT (24, 25) and NANOSH (22) can be integrated to produce a standardized protocol for general nanoscale materials, but they may not indicate exposure to specific nanomaterials. More investigation is needed for these generalized emission-based approaches to be shown to correlate with more traditional, filter-based industrial hygiene measurements for specific engineered nanomaterials (26, 27).

A series of workshops has been initiated to foster the process of harmonization and the integration of strategies for exposure measurement, analysis and storage of data (e.g. by NanolImpactNet and TNO in Europe, and by NIOSH in the US). Supplementing these measurement approaches, exposure can also be assessed using mathematical models involving key elements, such as the air dispersion characteristics, the manufactured nanoparticle emission rate, the worker’s distance from the emission source, and other factors. Mechanistic models to estimate exposure to nanoparticles have yet to be developed; however, concepts and tools for such models have been proposed (28) and can be used to structure the exposure assessment. For some specific activities, i.e. powder handling, a more detailed model has been developed (29). It is likely that in the near future further detailed models will be proposed although their calibration will remain a challenge. Currently, some initiatives are being carried out to harmonize strategies to analyse and report measurement data to facilitate future pooling and storage of data (23). Such retrievable data bases will play an important role in the process of model calibration and validation.

5.1.3 Job-activity exposure models

For the use of exposure models in epidemiological studies, it is essential that the models are activity- or task-based and provide exposure distributions to account for within- and between individual variation of exposure. Activity- or task-based models can then be used as building blocks for multidimensional job-activity exposure matrices for estimating personal exposure. An important condition for their use is that the description of tasks and activities is unambiguous. The same holds for the description of other input parameters, and contextual information. Another challenge will be to create a comprehensive overview of job titles and activities related to exposure scenarios, involving varying levels of exposure. Recent studies
suggest that the types of industries where manufactured nanoparticles are used and the types of exposure scenarios vary considerably (1, 30). Because the formation of cohorts for an epidemiological study may involve combining workers from different companies and possibly different countries, it is essential that job-activity exposure linkages are consistent among populations so that they can be included in pooled epidemiological studies.

An important problem, however, is measuring small numbers of manufactured nanoparticles against a significant background of ambient ultrafine particles, where currently only costly off-site analyses allow distinction of particle types (21). Contextual information about work processes, such as the kinetic, thermal and other energies involved, ventilation systems, and protective equipment, can help in modelling the contributions of task-specific exposure sources and existing hygiene measures. Here, a set of contextual information may replace such measurements for the purpose of large epidemiological studies where detailed particle assessments at every workplace would not be feasible.

5.1.4 Incorporation into exposure registries

For either short or longer-term assessment of worker health, good quality exposure data are required from workplaces where nanomaterials are routinely used. Thus, studies and exposure characterization campaigns should be promoted and the information gathered then needs to be shared broadly and made available to researchers investigating occupational health. Companies should be encouraged to keep records of work activities that are as detailed as feasible. Awareness of the need to collect exposure data and establish exposure registries should be raised as widely as possible and at a high level in businesses, with senior managers, safety officers and occupational health professionals, potentially via continuing professional development and other continuous education programs and professional societies (3). These results should then be incorporated into industrial, national or international exposure registries. An approach to creating such registries is discussed below.

5.2 Occupational health effect assessment

Workers’ health should be monitored to ensure that their occupational interaction with nanomaterials does not result in any temporary or permanent harm. To monitor health effects, it is necessary to identify potential biomarkers and assess their relationship with exposure, differentiating between sub-clinical biomarker effects and health endpoints. Diversity of particle types is a major challenge. Effect markers may provide a useful strategy to address this issue. One proposed approach to addressing the diversity of particle types is
to de-emphasize exposure assessment in favor of grouping materials that produce similar pathways to disease (8).

The following steps should be taken for collecting data for (large scale) occupational health studies:

1. Identify the pathophysiological mechanisms potentially involved. Identification can be inspired from the mode of action of traditional particles and by deducing the likely consequences of novel physicochemical properties and nano-bio interactions.

2. Once potential pathophysiological mechanisms are identified, it must be determined which bodily responses and diseases might be expected and which markers can be used to evaluate and validate their presence. As potential exposure to manufactured nanomaterials is very new the initial focus should be on short-term effects, i.e. effects that can be observed after a relatively short exposure duration.

3. Some of the long-term effects that need to be assessed will correspond to the accumulation of low level effects over time, whilst the onset of others will be delayed such that a relatively short duration of exposure leads only much later to manifestations of effects. Markers are needed that are able to detect both types of long-term effects, and early in the disease’s progression when corrective actions are still possible.

4. Health data will need to be collected and fed into health surveillance databases. It may be necessary to set up multiple interlinked databases to account for national differences in health systems and legislation.

Before any (global) health database can be built, a consensus needs to be reached about the health endpoints, the markers and the methods to be included. The steps towards effective occupational health data collection, as described above, require feedback loops and also regular revision of the parameters needed and which exposure data can be made available.

The implementation of sound risk management practice should not be delayed until long-term health studies are completed. While long-term effects can be expected and confirmed in research studies only after nanomaterials have been present for some time, there is an immediate need to take precautionary steps to prevent long-term effects. For example, the recognition of the potential for diseases with delayed manifestation, such as mesothelioma from inhalation of some types of carbon nanotubes (31, 32), implies the need for immediate and strict precautionary measures. Research will have to confirm at a later stage that the recommended measures were indeed sufficient.
In parallel to health effect studies in workers, full assessment of nanomaterial hazards is required. This will require analysis of all the bodily systems potentially involved and represents one of the main toxicological challenges posed by nanomaterials. Much discussion has taken place on whether the current suite of regulatory toxicity tests is suitable for nanoparticles. NanolImpactNet, other European projects and the WHO Collaborating Centres are putting emphasis on the development and validation of *in vitro* approaches to test the health effects of nanoparticles as these reduce the use of experimental animals. Nevertheless it must be kept in mind that the unique aspects of nanomaterials may influence both the experimental design and outcomes.

5.2.1 Identify potential health concerns and mechanisms

When reducing the size of materials down to the nanoscale, physicochemical properties and interactions with biological systems change. There is considerable uncertainty with regard to the health consequences of exposure to these materials. This is partly due to lack of toxicological data, but also a consequence of the complexity of the nanomaterial-biotic system interaction (33). The testing and sponsorship programme of the Organisation for Economic Cooperation and Development (34) lists over 30 material properties as being potentially interesting for study. However, combining the number of physico-chemical properties with the number of health endpoints and testing methods results in an almost unsolvable task. Thus, to save time and efficiently use limited resources (not just financial, but also manpower and the number of institutions able to do such extensive testing) prioritisation is required.

Prioritisation can be based on those characteristics believed to be most relevant and on effect mechanisms. The particle characteristics scientists most frequently cite as important to defining the interaction between nanomaterials and biological systems are size, surface properties, biopersistence (or solubility in biological media) and morphology (35). Size is a key aspect because small individual particles may translocate through membranes and tissue barriers allowing them to travel through the body and reach distal target organs such as the brain and liver (36). The aggregation/agglomeration state of nanoparticles may affect their potential for translocation while smaller primary particle size is associated with a larger surface area per unit of mass and thus a greater surface for biological interactions. Other relevant mechanisms via which nanomaterials are believed to cause effects are particles acting as transport vehicles (37-39), as leading to the generation of toxic substances such as free radicals and oxidative stress (40-42), and by the fibre paradigm, according to which long nanotubes are believed to act similar as long asbestos fibres (32, 43-45).
There is already a wealth of information in the epidemiological literature about exposure to “traditional” or “classic” particles and other (non-engineered) materials that fall within the EC’s definition of nanomaterials. These workplace fibres, traditional nanomaterials and environmental airborne particulate matter give indications about the types of effects that should be considered - although as yet unknown effects should not be excluded. Studies on ambient particles suggest that particle exposure causes oxidative stress and inflammation resulting in the release of chemical messengers such as cytokines, and vasomotor factors with subsequent generalised inflammation, thrombosis, atherosclerosis, and potentially chronic obstructive lung disease and pulmonary fibrosis if the target organ is the lungs (46-48). Studies of workplace fibres suggest that long, biopersistent fibres can lead over time to serious health outcomes such as mesothelioma, and consequently experts and some national authorities recommend precautionary measures similar to those for asbestos (15, 49, 50).

Experience from the occupational health monitoring of metal welders and flame cutters provides the most applicable example of an available data-set – although these workers are not exposed to one single nano-sized material, but to heterogeneous mixtures of highly reactive metal particles (some in the nanoscale size range) and gases. Reviews of the health effects of these fumes have, however, revealed few effects beyond modest decreases in lung function, airway irritation and pulmonary siderosis (51), with effects in the reproductive systems and central nervous system being inconsistent. The International Agency for Research on Cancer (IARC) has classified welding fumes and gases as category 2B, possibly carcinogenic with limited evidence in humans and inadequate evidence in experimental animals.

In contrast to traditional particles, only limited data are available about novel nanomaterials. A group in Taiwan (52) has shown promising early results for the use of biomarkers of small airway damage and inflammation, as well as biomarkers of injuries to endothelium and sympathetic nerve activation among workers exposed to nanoparticles. Another study reported on seven Chinese workers handling a range of chemicals and nanoparticles who became ill, with subsequent pathological examinations of lung tissue revealing evidence of pulmonary inflammation, fibrosis and foreign-body granulomas, and nanoparticles were observed in pulmonary epithelial and mesothelial cells (53). However, poor occupational hygiene and the workers’ complex exposures preclude definitive conclusions about the contribution of nanoparticles to the effects observed.
5.2.2 Strategy to identify short and long term health effect markers

Once potential endpoints or mechanisms of action have been identified, markers of effect need to be identified and validated. Biochemical tests or functional parameters to be assessed should be supported by consistent pathophysiological mechanisms. Attention could be focussed on exposure via inhalation and the skin, since these routes of exposure are better understood for non-nanoscale chemicals. Pulmonary and cardiovascular diseases (leading to increased morbidity and mortality amongst vulnerable groups in particular) have been linked to pollution and levels of ultra-fine particles, but their use as health endpoints for workers exposed to manufactured nanomaterials have limitations: they are non-specific (and certainly not nano-specific), have a high prevalence in the general population and share multiple non-occupational risk factors. Thus, information on exposure to these other risk factors would also have to be acquired to allow attribution of effects (and it would be only on the level of worker populations and not individuals). Although these factors make long-term studies more challenging to conduct and interpret, they have been used in studies of many occupational diseases.

A variety of other potential short or long-term effect parameters have been proposed for a targeted assessment of personnel exposed to nanomaterials. These include heart rate variability, blood-clotting parameters, pro-inflammatory cytokines, up-regulation of adhesion molecules or antioxidant capacity, and biomarkers of pulmonary fibrosis (46, 54-56). These biomarkers are increasingly used to assess cardiovascular effects of fine particulate matter, quasi-ultrafine (i.e. PM < 0.25 µm) and primary carbon aerosols derived from traffic-related sources (57-60). These biochemical parameters have consistent pathophysiological mechanisms that have been investigated for combustion-derived ultrafine particles and diesel exhaust particles generated in laboratory settings (61). Although promising in epidemiological research as putative biomarkers of effect, these parameters are still not assessable for their predictive value of health risk at an individual level; they are not routinely applicable and need to be further validated.

On the basis that the potentially relevant health endpoints that have been tentatively ascribed to manufactured nanoparticles are cardiovascular, pulmonary and inflammatory effects, possible health monitoring endpoints include:

- Assessment of markers of exposure (e.g. presence of chemicals in the blood or the urine; this can readily be done for chemicals such as metals - strictly speaking not an endpoint).
- Chemical changes in exhaled air or exhaled breath condensate suggested to reflect abnormalities of the airway lining fluid and lung inflammation (62-64) but also potentially exposure (65)
• Local effects: inflammatory changes, short-term respiratory changes, respiratory, eye or skin irritation, depending on the route of exposure/site of uptake (with special tests to study biopersistent long fibres such as some forms of carbon nanotubes).

• Systemic effects to confirm cardiovascular changes and inflammatory mechanisms: heart-rate variability, platelet aggregation and other pro-thrombotic effects as well as cytokines and differential blood cell counts (54, 66).

• Medical tests for early detection of health effects at a preclinical stage (e.g. clinically validated biological markers of cardiovascular, hepatic, renal, haematological or respiratory dysfunctions).

We do not yet know whether or how such health effects may differ between chemically or structurally different particles at the nanoscale. To date, biomarkers of exposure cannot be adequately developed owing to the lack of consistent toxicokinetic studies of nanomaterials, which may be partly related to the enormous variability of surface properties for each type of nanomaterial and even between batches of the same material. However, it is likely that traditional biomarkers of exposure (e.g., mass quantification in serum or urine) will be more feasible for nanoscale metals than for carbonaceous nanomaterials. It is at present unknown whether manufactured nanoparticles in general, and carbon nanotubes in particular, can exacerbate pre-existing medical conditions or increase the susceptibility to certain diseases.

5.2.3 Occupational health databases

Health data collection should be tightly linked to exposure assessment. At a minimum, health data collection should be accompanied by a general exposure assessment at work places with identified risk potential. An occupational health database will need to allow the identification of exposed individuals and/or worker populations. Ideally, such a database would be held at a centralized data collection site with extended health screening at specific nanomaterial facilities. In addition to this basic data collection, targeted research studies will be needed with detailed exposure assessment and extended health effects monitoring, including mechanistic studies of the effects of nanomaterials. As for exposure, companies should be encouraged to keep records, awareness about the need for data collection should be raised, and the data should be incorporated into industry, national or international health registries. It would be useful if the databases can be linked to existing databases such as the national or international registries of death and cancer.
5.3 **Harmonized study designs and data collection strategies**

The success of epidemiological studies for quantitative risk assessment depends on the quality of available exposure and health response data. To achieve a coherent approach that leads to valid conclusions, data collection needs to be defined in anticipation of future (ideally prospective) studies that will pool and compare different situations world-wide. Over the coming years, the following points need to be addressed:

- Risk assessment and management cultures in different industrial sectors and countries need to be identified.
- Data collection strategies and data protection philosophies and the associated legal systems of different countries need to be assessed, and strategies for dealing with them defined.
- The most suitable epidemiological designs for different purposes need to be identified.
- Exposure and health registries need to be established that can link to health surveillance systems or epidemiological research projects.

It is unlikely that a single, global project on this scale could be set up to follow workers exposed to nanomaterials. However, a more modest approach with harmonized exposure and health registries could be established, recording details of workers’ activities, available information on exposure levels, the nanomaterials handled and health condition. Ideally, workers’ samples (e.g. blood, urine) would be stored for later analysis as more potential biomarkers of effect and exposure become available.

5.3.1 **Identify ethical, cultural and regional differences**

Any global data collection strategy needs to account for regional differences in existing technologies, exposure protection strategies, safety culture, data protection philosophies, and ethical aspects (67). These challenges are superimposed on the challenges and data needs for the conceptual models described above. For each of the recognized data collection needs, potential challenges for collecting (in the sense of being able to obtain) and defining (in the sense of using terminology that is free of sensitive connotations) such data needs to be identified. Differences in data protection laws might pose challenges to the collection (68), but also the pooling of data across frontiers. Usually, data can only be used for a pre-defined purpose and subjects’ identities must not be revealed. In some countries individuals can request their data be withdrawn at any moment during the study, often there is an expiration date relating how long data can be stored, and data may only be shared with researchers located in countries that have at least the same regulation of data protection.
All of these challenges need to be described and addressed in a strategy on how to overcome the differences and gaps when defining the data needed for collection. Thus, information about cultural and regional differences should be an integral part of the recommendations for a globally harmonized data collection strategy. WHO Collaborating Centres and the OECD may have an important role to play in the collection of such data about cultural and regional differences.

5.3.2 Define epidemiological designs
Well conducted epidemiological studies are an important source of information to risk assessment as they provide directly relevant human data. Epidemiological studies face challenges due to uncertainties in exposure measurements, dose to the target organ, and health effects resulting from the interaction of cells with nanomaterials. Nevertheless, examples exist [e.g., radon-exposed miners (69) asbestos-exposed textile workers (70)] in which prospective cohort studies have produced data that form the basis of quantitative risk assessments.

Currently, the number of workers facing a potential exposure to manufactured nanomaterials is not known. Major challenges in conducting prospective cohort studies in nanomaterial workers include developing a large cohort size and the long time periods required to draw firm conclusions regarding chronic health effects. In the interim, small-scale studies of 50 to 100 workers could be conducted within the next 5 years to assess biomarkers of exposure or of early effect (9).

Whilst there is currently no firm basis for the recommendation of targeted nano-specific occupational health surveillance for most manufactured nanomaterials, this should not be a reason for paralysis or inactivity. There are clear knowledge gaps and achievable recommendations can be made:

- General health surveillance should target those working with nanomaterials where exposure is likely (e.g. processes that are not contained), and the systems for recording the processes and types of nanomaterials used by the workers should be improved.

- Workplaces should apply measures to control exposure, containing particle emissions and deploying personal protective equipment for workers when appropriate if potential exposure to nanomaterials cannot be excluded. The measure should be recorded, ideally in a standardized reporting format, during occupational health studies.

- Simple questionnaires could be used for self-reporting symptoms so that focussed studies can be undertaken for different types of nanomaterials and industries.
- Individual cases of ill-health in those working with nanomaterials should be scrutinised to assess whether nanomaterials are a likely attributable source of the ill-health.

- Improved knowledge-sharing should be encouraged at a national and international level, through occupational health reporting networks.

- Simple nanomaterial measurement techniques are required that can be applied under most occupational circumstances without advanced technical knowledge.

- Use should be made of biological monitoring (where applicable).

Overall, there is a clear need to gather experimental, clinical and epidemiological data in order to characterize the relationship between exposure and health outcomes, and to provide a basis upon which to build and explore the effectiveness of preventative measures.

5.3.3 Health surveillance and setting up registries
The rapid development of exposure registries will help to provide data for the epidemiological studies of the future (3, 71). However, there are currently no validated methods for health monitoring and reporting that are specific to nanoparticles, but existing approaches for non-nanoscale substances could be adapted. In the UK, occupational health reporting systems have been developed, but they rely on reporting (sometimes self-reporting) of adverse outcomes and they rarely record information on exposure levels. Furthermore, if effective preventive approaches are adopted in workplaces handling nanomaterials, the number of expected adverse outcomes could be very small or even non-existent. In France basic information on worker health status is already being collected via mandatory occupational health surveillance, and this will be available in the future for retrospective studies (10). The approach recommended by NIOSH in the US is to consider the hazard and exposure levels of the nanomaterial when making a decision about whether to employ routine medical surveillance (72).

Historically, in occupational medicine, the practice of health surveillance has represented the final step in a process based on the integration of both experimental and epidemiological studies that identify the hazards and are supported by the implementation of occupational exposure limits (OELs). Medical surveillance requires at least a qualitative risk assessment and can be implemented when a residual risk exists and when the target population has been clearly identified (72). One major issue is the correct classification of exposure situations and linking them to the medical records. Currently, validated or even calibrated exposure models are lacking, however, and the application of such models for estimating the exposure of individuals will be hampered by the lack of indications of within/ between individual (worker) variances.
While there are no specifically validated methods for the risk assessment of nanomaterials in the workplace, the occupational safety community may be under-utilising its existing knowledge in hazard and exposure control of ionising radiation, biological agents, pharmaceuticals, nuisance dusts and pollution (73). With technology developing so rapidly, it will be necessary to make continual re-assessments and re-evaluations of the risks to workers' health and safety from nanotechnologies and nanomaterials in particular. It is suggested that existing regulations can address the emerging issues and potential hazards presented by nanoparticles in the workplace, and although some disagree with this view, guidance and codes of conduct have been produced by different organisations (e.g. European Commission, UK’s Health and Safety Executive (HSE) (15), and NIOSH (74)). Although incomplete, the body of toxicological evidence on manufactured nanomaterials suggests that occupational hygienists and health and safety professionals should recommend precautionary management measures in the workplace, including in research laboratories handling nanoparticles.

Risk management of nanomaterials in the workplace would involve recognition of potential worker exposure and the implementation of measures to reduce or minimise it (75). Occupational health surveillance and epidemiological research can support the risk management process by assessing whether health effects can be found at levels believed to be safe, but also by identifying the risks to workers where the exposure is insufficiently controlled (3). Thus, health surveillance and epidemiology are important tools to ensure workers' health. Medical screening constitutes just one part of a complete health and safety management program. NIOSH has published interim guidance on medical screening for workers handling nanomaterials (72), which discusses the value of medical screening for asymptomatic workers. A second NIOSH guidance document (50) describes the need for medical surveillance for carbon nanotubes, which demonstrate a specific hazard in toxicological studies.

### 5.3.4 Harmonisation of studies

To enable the harmonisation of studies, preparatory step such as the establishment of exposure and health effect registries may be needed. Occupational exposure and health registries have been used in public health for over 50 years, and are especially useful when the risks to workers are not well-defined. Recently, the Dutch Health Council and the Social and Economic Council of the Netherlands strongly recommended the establishment of an early warning system and exposure registries for nanomaterial workers (http://www.gezondheidsraad.nl/sites/default/files/Nanotechnologie2.pdf, http://www.gezondheidsraad.nl/sites/default/files/200818.pdf, http://www.ser.nl/~media/DB_Adviezen/2000_2009/2009/b27741.ashx).
We propose generation of an international framework under which such registries could be developed with a harmonized format and interlinked across Europe and globally. The proposed actions and related objectives are shown in Table 2.

**Table 2 Central issues for study of workers' exposure and health**

<table>
<thead>
<tr>
<th>Actions</th>
<th>Objectives</th>
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</thead>
<tbody>
<tr>
<td>Collect, collate, analyze and evaluate existing accessible data on exposure and contextual information, including activity-based exposure profiles with regard to their potential use in cohort studies and protection strategies</td>
<td>Review of existing exposure data to identify which risk management measures are most effective in reducing potential exposure, including during non-routine operations/accidents</td>
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<tr>
<td>Identify existing descriptors for job titles, activities, processes, and industrial sectors, and subsequently build a multidimensional job-activity-exposure matrix for the estimation of personal exposure</td>
<td>Develop models to link job activities to potential exposure to allow to evaluate the effect of current risk management practices on exposure levels. Test the feasibility of using the proposed matrices in epidemiology studies.</td>
</tr>
<tr>
<td>Identify and critically evaluate candidate biomarkers of effect and exposure to nanomaterials</td>
<td>Evaluate and select candidate biomarkers to contribute to improved methods for health impact assessment, particularly for monitoring early health effects in exposed populations.</td>
</tr>
<tr>
<td>Evaluate practical issues for use of biomarkers in epidemiological studies and/or health registries of nanomaterial workers</td>
<td>Evaluate practical issues for use of biomarkers in epidemiological studies and/or health registries of nanomaterial workers.</td>
</tr>
<tr>
<td>Outline the needs and possibilities for building occupational epidemiological studies for workers dealing with nanomaterials.</td>
<td>Test feasibility of the proposed study designs, using existing data formats.</td>
</tr>
<tr>
<td>Review and evaluate existing and emerging tools for early assessment and management of risks among workers exposed to nanoparticles, and provide guidance for their harmonization and improvement</td>
<td>To obtain harmonized exposure and health effect monitoring that allows a better evaluation of the various existing risk management tools. It also may eventually help identifying potential risky materials and strategies for managing these risks in a conservative manner.</td>
</tr>
<tr>
<td>Develop guidance for exposure and health registration to be used by SMEs</td>
<td>Development of such registries would ideally occur on an international scale to increase the numbers of workers recruited, although national views and regulations on data gathering and security may complicate this. These problems could be avoided by establishing national registries that collaborate internationally.</td>
</tr>
</tbody>
</table>

Such a system for international surveillance can be set up in analogy to initiatives done for assessment of background exposure in the normal population. Indeed, within the frame of the FP7 EU project COPHES and the Life+ supported project DEMOCOPHES a strategy for harmonized human biomonitoring in 17 EU countries is currently worked out. In parallel with which was done there, also for nanomaterials occupational exposure, a pilot monitoring program can be worked out to start organizing registries and/or measurements starting on a
limited scale. The study should be coordinated and steered by a limited group of a few partners who prepare the pilot study. In this context, it would be good to include partners of industry and occupational associations in this steering group. If there are enough resources, other participants can join and should be fully supported by providing all (technical) advice and/or training needed.

To obtain wide acceptance and support for such approaches, communication will be a very important element. This communication needs to reach workers, managers and also policy makers and research funding agencies. It was recommended that communication about nanomaterial risks should adopt established concepts of risk perception and risk communication strategies (76).

5.3.5 Check appropriateness of screening strategies

When hazard data are absent (as with many nanomaterials), the question is whether it is possible or informative to initiate specific health surveillance for nanotechnology workers (71). Unnecessary health surveillance can be associated with risks itself if it leads to incorrect diagnoses, uncertain interpretation, and the perception that work may not be “safe” for workers. Criteria of appropriateness should therefore be assessed; i.e. any screening strategy for a worker health should be justified on the basis that either: (i) finding a medical condition at an earlier stage significantly improves the potential outcome compared to a situation with no screening; (ii) especially in the case of chronic diseases, it helps reduce the population’s exposure to a level that prevents the development of these early stages of disease; or (iii) the screening program is used as part of a wider strategy to ensure effective exposure control.

A range of factors needs to be considered to determine whether implementing health surveillance in the workplace is appropriate. Among the existing criteria are assessment of the burden of suffering (which the precautionary approach aims to prevent), the accuracy and reliability of current test methods, the effectiveness of early detection in the absence of clear correlations to health endpoints from manufactured nanomaterials, and an assessment of the benefits versus any harm resulting from the screening itself. On the basis of these criteria, the known risks associated with some screening methods (e.g. chest x-rays or computed tomography) would have to be outweighed by the benefits, such as implementation of workplace risk management measures or preventative health interventions, before such a recommendation could be made. Routine screening has to be justified on the basis of a sufficient likelihood of the expected exposure to manufactured nanomaterials causing the condition in question; the absence of any indication of a risk attributable to nanoparticle exposure precludes conclusions on this point at present.
6 Conclusions / Recommendations

To best understand and control their health and exposure risks of workers dealing with manufactured nanomaterials, studies of groups and cohorts of those workers are necessary. At the moment, the necessary framework conditions to conduct such studies are not in place. To provide a coherent approach and make future epidemiological research a reality, a well defined framework is needed for the careful choice of materials, exposure characterisation, identification of study populations, definition of health endpoints, and evaluation of the appropriateness of study designs. Particularly needed are:

- A basis for prioritizing which engineered nanomaterials merit investigation (e.g., based on toxicological or inferential studies)
- A consistent, evidence-based set of job titles and task-based exposure profiles for epidemiological studies (informed by actual individual workplace assessments)
- A method for linking industry, company and job descriptors to exposure and consistent exposure metrics; this will need to provide recommendations on the type and format of data to be collected and how it can be interlinked so that future studies can use the data for developing exposure estimates for large cohorts of workers.
- Criteria for potentially useful biomarkers and (pre)clinical parameters for epidemiological studies on workers in small and medium enterprises and transnational companies. Recommendations on the feasibility of human population studies based on these biomarkers.
- Recommendations on the requirements for harmonized approaches for human biomonitoring and health effect studies tailored to nanomaterial workers.
- Recommendations for harmonization and the improvement of tools for early assessment and management of risks, and for exposure registration. These should be made available to relevant stakeholders, including small and medium enterprises, along the global value chain.

For either short or longer-term assessment of worker health, good quality exposure data are required from workplaces where nanomaterials are routinely used, and methods for collection of exposure data should be tested, validated and agreed internationally with some urgency. Concomitantly, field studies and exposure characterization campaigns should be promoted and information shared broadly. Companies should be encouraged to keep records of work activities that are as detailed as feasible, and if possible, full use should be made of simple sensors such as particle counters (recognizing that these are not at all
specific for the manufactured nanomaterial of interest). Awareness of potential hazards and the need to collect exposure data and establish exposure registries should be raised as widely as possible, potentially at a high level in businesses, with senior managers, safety officers and occupational health professionals, potentially via continuing professional development and other programs of continuing education.

Although at present no nano-specific screening procedure exists for the medical surveillance of workers potentially exposed to nanoparticles, periodic general medical examinations are to be recommended, preferably based on non-invasive procedures. Increasing general health surveillance for these workers could lead to an earlier recognition of adverse effects and information about which symptoms should be followed in other similarly exposed workers. This might also lead to informative individual index cases being identified.

There is an urgent need for further toxicological evaluation and physico-chemical characterization of all the types of nanomaterials currently handled in workplaces, and a determination of the relationship between particle characteristics and health effects so as to facilitate prediction of the effects of new materials, and to identify the most important exposure characteristics to be assessed. Biomarkers of exposure and health effects are also required, and novel, ideally point-of-care, detection techniques need to be explored for assessing those exposures, e.g. via determination of particle numbers or pro-inflammatory cytokines in exhaled breath condensate.

In spite of the advances in establishing a conceptual framework leading to a higher degree of worker protection, epidemiological studies of workers potentially exposed to manufactured nanomaterials will be difficult to conduct, for both ethical and practical reasons. These include: the heterogeneity of nanomaterials used in occupational settings, the overlap in exposure between combustion- or pyrolysis-derived ultrafine particles and manufactured nanoparticles, the lack of standardized exposure metrics, the long time-frame required to develop informative exposure histories, and the international cultural differences related to surveillance systems, data recording and storage, and data protection. Further issues include how to distinguish health effects that may arise from exposure to nanomaterials from those due to exposure to other workplace hazards and potential toxicants, and the effects of multiple exposures and confounding factors, such as smoking and underlying health conditions.

The proposed roadmap addresses a joint strategy and the flow of actions needed to achieve these goals. Such a joint strategy will ensure that the costs of action are not disproportionate to the potential benefits and, very importantly, that the strategy will be pragmatic and practical. The discussion has only just begun about the benefits and challenges associated
with the different potential approaches for occupational health surveillance and epidemiological studies for workers exposed to nanomaterials. However, results from such studies are needed for the assessment of nanomaterial workers’ risk and for evaluation of exposure controls.
7 References


34. OECD. List of manufactured nanomaterials and list of endpoints for phase one of the sponsorship programme for the testing of manufactured nanomaterials: revision Paris: Organisation for Economic Co-operation and Development (OECD); 2010.


