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Investigation of the Release of Particles from a Nanocoated Product

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ABSTRACT

Manufactured products are being coated with nanoparticles in order to functionalize them with antibacterial or self-cleaning properties or to improve their durability etc. As the (eco-) toxicological effects of the nanoparticles are not well known yet, their use could lead to new potential risks for the workers, the consumers and the environment. This study focuses on the release of the nanoparticles during the operations related to the handling and processing of an automotive part. The part is made up of a metallic alloy and, in order to reduce friction, the part is nano-coated with inorganic fullerenes. The mechanical stresses appearing during these operations are reproduced in a nano-secured facility. The release of nanoparticles is found to be increasing with the wear energy applied on the surface.

Keywords: Nanoparticle; Coating; Emission; Wear; Friction; Release

1. Introduction

There is a growing interest in the development of nano-coatings in industrial applications. The finished products, once nano-coated, can be self-cleaning, antibacterial, UV-resistant etc. Due to these advantages, their usage is becoming frequent and thus, subjected to investigations regarding the release of nanoparticles (NP) in the air, which may pose toxicological effects on the human and the environment. These effects of NPs are not well known yet and represent thus a possible threat, especially, since a new era of this new technology is commencing hastily. During their whole life-cycle, the products are exposed to wear by their environment and utilization and thus degradation might occur, [1-3] which could lead to the release of NP. However, in view of the large variety of possible exposures, the present data are still not sufficient. Moreover, as workplace atmosphere inspection for manufactured nanoparticles is relatively complex and expensive to conduct, new comprehensive studies may be required to provide estimates of their release in the occupational exposure. The need for these studies will rapidly increase due to growing production and use of manufactured nanoparticles. Previous studies (**Table 1**) have shown the emission of particles from different types of materials when solicited mechanically or thermomechanically or by UV. For environmental degradation,

it has been shown that exposure to UV radiations can lead to the increase of the density of NPs on the surface of a nano-structured polymer composite [3]. Fewer works have studied the exposure risks during handling and operating nanoparticles [4]. But new heights have been achieved when the nanoparticles were started being characterized during their generation process through wear. Their characterization also got encouraged with the fact that in several cases of mechanical wear, very high particle emissions have been observed for nanostructured materials, e.g. when drilling, [5] grinding, [6] or sanding [7] (see **Table 1**). From comparisons of materials with and without nano-structure, it has been confirmed that emission of particles occurs in both cases [5,7]. The study presented here aims at characterizing a possible source of NP release to air, which may lead to their inhalation and thus the possible worker exposure to NP during the operations related to the handling and processing of an automotive nanocoated part. Analysis has been carried out to quantify and characterize the particle emission levels of the part during the subsequent wear processes. An attempt, simulating these wear processes in real life conditions experimentally, is presented.

2. Experiments

For the experimental investigation, an iron alloy shaft cap with a radius of 2 cm and height 5 cm was used. Its

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Table 1. Overview of references.

Material	Wear mechanism	Particle emission		Reference number
		Total number concentration and size range	Main observations and findings	
Coated surfaces with zinc oxide nanoparticles	Abrasion with a Taber	<20 particles/cm ³ [6 nm - 1 µm]	The total number of generated submicrometer or nanoparticles was extremely low and lay below the statistic significance threshold. The particles <100 nm remain embedded in the coarse wear particles.	[10]
Polycarbonate containing 3% wt Carbon Nanotube (CNT)	Abrasion with a Taber	50 - 1800 particles/cm ³ [7 nm - 10 µm]	The abrasion effect is enhanced when increasing the normal force. The particle emission is also increased with the speed (number of cycle per minute). No free CNTs were observed.	[8]
Polymethyl Methacrylate (PMMA) containing 10% wt Cu	Abrasion with a Taber	Up-to 10,000 particles/cm ³ [7 nm - 10 µm]	The type of abrasion tool has a high impact on particle emission intensity. No visible free Cu nanoparticles were observed even for one case they would be nearly released.	[9]
TiO ₂ nanopowder coated on different substrates including wood, polymer and tile.	UV + fan + rubber knife	~10 to 600 particles/cm ³ [15 - 661 nm]	These actions together can greatly reduce the binding force between the TiO ₂ nanopowder and the substrate surface, and, therefore produce particle emissions from the coating products.	[15]
Paints with or without engineered nanoparticles (NP). NP: carbon black and TiO ₂	Sanding	14,000 - 28,000 particles/cm ³ [5 nm - 20 µm]	The sander was the only source of particles smaller than 50 nm and they dominated the number concentration spectra. NPs released during the sanding do not occur in individual particles.	[7]
Non-nanodoted materials: Granite, Ceramic, Steel, Aluminum, PTFE, Hardwood	Grinding	10,000 - 800,000 particles/cm ³ [4 nm - 20 µm]	Ultrafine particles do have the potential to form a significant component of a grinding aerosol for a number of substrates. processes: 1) from within the grinding motor, 2) from the combustion of amenable grinding substrates and 3) from volatilization of amenable grinding materials at the grinding wheel/substrate interface.	[6]
Surfboard with or without carbon nanotubes (CNT)	Drilling and grinding	Upto 107 particles/cm ³ [7 nm - 10 µm]	When used, the CNT are not placed in the outside layer, but inside. Material with or without CNT: the particle emissions are comparable. The top layer (varnish) is the major source of particles.	[5]
Steel and fiber cement panels coated with the ZnO and Fe ₂ O ₃ nanoparticles	Sanding	Upto 109 particles/cm ³	For the given stress, the swarf mass, the particle size distribution of the released aerosol, and consequently the number of released particles depend primarily on the used surface coating.	[11]
Epoxy nanocomposites reinforced with CNTs	Weighing operation and sanding	With no local exhaust ventilation: 2.68 µg/m ³ Inside the fume hood: 21.4 µg/m ³	This study demonstrated that weighing bulk CNTs and sanding epoxy containing CNTs generate few airborne particles that are nano-sized. Furthermore, it was demonstrated that sanding epoxy containing CNTs may generate micrometer-sized particles with CNTs protruding from the main particle core.	[12]
PA6 nanocomposites	Drilling	20,000 particles/cm ³ [175 and 350 nm]	It is likely that the presence of nanoclay in some way retains the formation of high quantity of airborne particles and promotes particle deposition.	[13]
Concrete	Cutting	5.28 ± 1.26 mg/m ³	An increase in CS resulted in an increase in the four measured particle exposure levels resulting from its influence on the generation of coarse particles. For both Ctho and Cres which are known to have a higher content of fine particles, the increase in AF resulted in an increase in their concentrations. However, for the concentrations with a higher fractions of coarse particles (<i>i.e.</i> , Ctot and Cinh), the increase in AF led to an initial increase, followed by a decrease in measured particle exposure concentrations.	[14]

surface was coated homogeneously with inorganic fullerene, having a coating thickness of 0.1 µm. The coating ensures the lowering of its friction coefficient. An emission chamber *i.e.* a sealed glove box (model 830-ABC/EXP, Plas-Labs), has been used to ensure lim-

ited background particles concentration inside the chamber and to prevent the release of (nano) particles to the outside atmosphere, thus, providing a nano-secured facility. The details of all the instruments, used during the test, are given in **Table 2**. During the whole test, clean air

Table 2. Instrumentation.

Parameter	Technique	Instruments	Characteristics	Set sample time
Number concentration [4 nm - 1 μm]	Condensation Particle Counter (CPC)	TSI model 3785	Water-based	1 s
		Grimm model 5.400	Butanol-based	6 s
Size distribution [10 nm - 0.5 μm]	Scanning Mobility Particle Sizer (SMPS)	DMA TSI model 3081 + CPC TSI model 3010		235 s
Size distribution [0.3 - 20 μm]	Optical Counter (COP)	Grimm model 1.108		6 s

(Filtered Air Supply 3074B, TSI) is passed upwards through the chamber. All devices are fed from a unique sampling point, which is set above the testing part to collect the maximum of particles released into the air from the experiment (**Figure 1**). The clean air inlet flow rate is equal to 0.075 lps. It enhances to carry out the investigation in an atmosphere practically free of particles (2 particles/cm³) and thus to detect emitted particles even at the low emission levels. Two Condensation Particle Counters—water based CPC (model 3785, TSI) and a butanol-based CPC (model 5.400, Grimm)—have been used in parallel which enabled reliable measurements of even low particle number concentrations, in the range of particle sizes from 4 nm to 1 μm . The particle size distribution is measured by a Scanning Mobility Particle Sizer (SMPS) configured with a long Differential Mobility Analyzer (DMA, model 3081) for particles' size ranging from 10 nm to 500 nm and by an Optical Particle Counter (OPC, model 1.108, Grimm) for particles' size ranging from 300 nm to 20 μm . The total sample time for SMPS has been kept at 235 s (scan time 215 s, retrace time 20 s) whereas for other instruments, it has been kept at low values. For validating the scan by SMPS, it was operated only under stable particle concentration conditions. However, a rougher state of the particles concentration stability cannot be neglected and excluded from the scanning period in these types of experiments. The experimental investigations presented here are corresponding to the operations which occur when a part, under development, is handled e.g. for unpacking, roughness measurement, specimen preparation for microscopy observations and other tests. In order to simulate these real life operations at real scale, following experiments have been carried out sequentially on a given part with increasing levels of stresses: 1) Extraction from the safety bag; 2) Plastic bag agitation; 3) Part extraction from plastic bag; 4) Part moves in 3 dimensions; 5) Surface rubbing with glove; 6) Falling from a height of 20 cm; 7) Surface scratching in 3 directions; 8) Surface sawing; 9) Surface sanding. The sampling point to measure the particles number concentration has been kept in the vicinity of the source. All the above mentioned operations were carried out in a glove box measuring 300 cm $\times$ 240 cm $\times$ 360 cm. For test #9, the rotary tool's

motor (Dremel 400 DIGITAL with a rotating aluminum oxide abrasive wheel) has been placed outside the chamber to avoid the contamination from the particles released by the motor's brushes. Its velocity of rotation has been kept at 19,000 rpm. It is interesting to note that these activities are also arranged in ascending order of wear energy levels. The wear energy associated with a system in a process can be defined as the energy consumed during machining or erosion or deterioration of the system during that process. The precautions for nanosafety are achieved by placing the glove-box within a nanosecured venting hood and putting the sample parts into two plastic bags, when not in use. However, due to the lack of the information on homogenization of the inside air due to ventilation, inlet and exhaust valves, data available on the measure of the particles number concentration should be considered as semi-quantitative. In order to ensure the repeatability of data and results presented in the present work, the whole experimental procedure was repeated three times. The particle deposition on the surface walls is a critical issue in such experiments which are done in a confined area. This may lead to the carrying out of these particles from one experimental trial to another. An utmost care has been taken to avoid it by cleaning the surface walls using an ethanol absorbed paper after every test and trial.

3. Experimental Results

The number concentration of the released particles versus time, measured by means of COP Grimm, water based CPC and butanol based CPC, in a sequential order, is shown in **Figure 2**. Note that **Figure 2** depicts the number concentration during one of the three trials. The particle concentration level appearing before applying any stress (often referred as background noise level) is approximately 2 particles/cm³. During test #1, *i.e.* extraction of the part from its bag, a particle concentration of 4 particles/cm³ was measured. Tests #2 to #7 generate peaks of particle concentration of approximately 10 particles/cm³, while peaks for sawing (test #8) range from 50 to 1000 particles/cm³. Test #9 generates highest concentration level of 900 particles/cm³ which is measured for the particles lying in a size range of 10 nm to 0.5 μm , as shown in **Figure 3**. From **Figure 3**, it is evident that

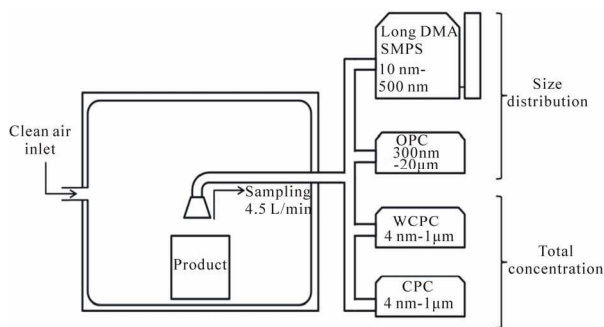


Figure 1. Experimental set-up.

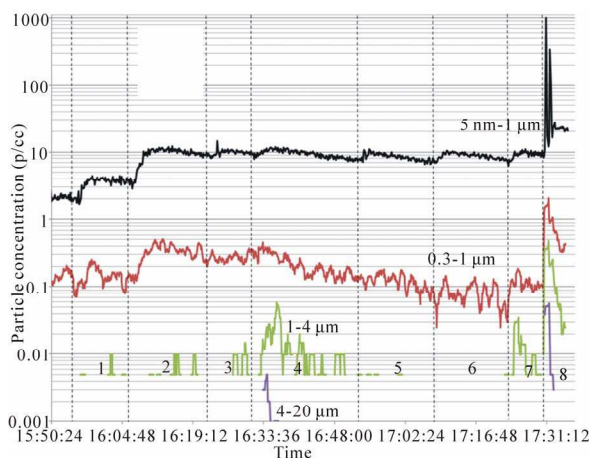


Figure 2. Particle concentration through time for different size ranges while testing a single nanocoated part (cf. Table 3, except sanding).

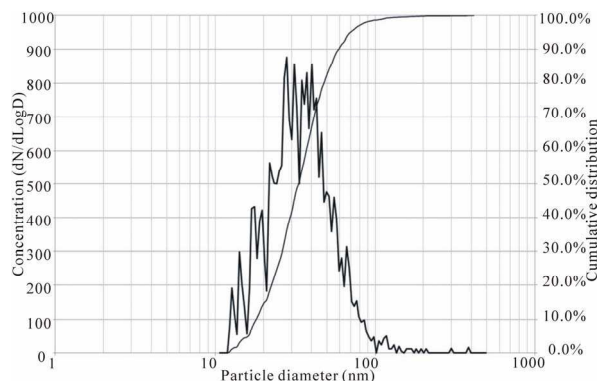


Figure 3. Particle size distribution while sanding measured by SMPS.

50% of the particles generated during sanding are smaller than 40 nm.

During tests #1 to #5, the wear energy levels are very low due to low level of stress application. For the particles lying in the size range of 5 nm to 1 μm, their concentrations were found to be in the range of 4 to 12 particles/cm³. For micron sized particles, concentration levels are also low. Therefore, from these observations, it has been found that for the present nanocoated shaft cap,

handling operations (tests #1 to #5) do not involve any particle emission or their exposure threats to the operator. However, for the tests #6 to #9, there may be possibility of a considerable particle emission because of the involvement of high levels of stress application. However, for tests #6 and #7, this hasn't been seen so far. The only phase where a significant emission has been seen is test #9. In this test, as soon as the sanding commences, there occurs a sharp peak of the number concentration due to the ejection of the particles from the material. To have a stable particle concentration during the particles size scanning by SMPS, the scan starts after the peak has arrived and it starts diminishing slowly with time, thus, providing a rough but stable particles presence.

4. The Energetic Approach

In the present approach, we have seen a series of operations involving increasing order of stresses acting on the nano-coated part. For the handling operations involving low stresses (#1 to #5), the particles emission is negligible. For the test #6, which involves accidental fall of the part, the same low emission continues. In order to get a better insight of this accidental scenario, the type of material in play should be noted. This is a light weight iron alloy which has got a high value of impact strength and ductility. The impact stress to which it is subjected at the end of the fall was calculated and found to be less than 100 Pa Equation (1) which is very less in magnitude as compared to its material impact strength. This means, all the impact applied on this nano-coated part gets sufficiently absorbed by rendering the part untouched. As a result, the wear energy applied on this part seemed to be lesser than required in order to have a higher particles emission. But if the same wear energy had been applied on a brittle material having very low impact strength (ex. Ceramics, concrete, glass etc.) the scenario of accidental fall would have produced high particles emission concentration. Therefore, it shows that the particles emission depends on both the wear energy level as well as the material to which it is being applied. For test #7, the low particle emission concentration during the peeling off of nanocoating signifies the sustainability of integrity of nanocoating during getting peeled off. However, it still requires a better view on structural changes occurring during this process. In **Table 4**, the values of wear energy are shown with their corresponding total particle number concentrations. For the calculation, following basic equations have been employed:

For accidental fall:

$$E_a = m * g * h \quad (1)$$

For scratching and sawing:

$$E_d = \mu * F_N * d \quad (2)$$

Table 3. Stress tests-description, aim and results.

#	Description	Aim	Wear Energy level	Total Particle emission concentration (#/cm ³)			
				5 nm - 1 µm	0.3 - 1 µm	1 - 4 µm	4 - 20 µm
1	Extraction from the safety bag		Very low	2	~0	~0	~0
2	Plastic bag agitation	Nanoparticles release while handling the part	Very low	10	<1	~0	~0
3	Part extraction from plastic bag		Very low	10	~0	~0	~0
4	Part moves in 3 dimensions + shock test		Very low	10	~0	10	+
5	Surface rubbing with glove	Accidental scenario #1	Very low	10	~0	~0	~0
6	Falling from a height of 20 cm (h)	Accidental scenario #2	Low	10	~0	~0	~0
7	Surface scratching (3 directions)	Nanoparticles release during roughness measurement	Low	12	<1	~0	~0
8	Surface sawing	Nanoparticles released during sample preparation for microscopy	Medium	300	2	<1	<1
9	Surface sanding	Extreme stress	High	100,000	X	X	X

For sanding:

$$E_d = \mu * F_N * r * \omega * t \quad (3)$$

with E_a is the energy absorbed (J), m is the mass (kg), g is the acceleration due to gravity (m/s²), h is the height of the fall, E_d is the energy loss (J), F_N is the normal load (N), d is the crossed distance (m), μ is the friction coefficient, r is the radius of the abrasive wheel (m), ω is the velocity of rotation (rad/s) and t is the stress duration (s). In **Figure 4**, the wear energy levels applied on the nano-coated part have been plotted on a log-log graph. The four points in the **Figure 4** correspond to the four energy levels in the tests #6 to #9. Clearly, with increasing wear energy level, particles emission concentration also increases, hence a direct proportionality between the two.

5. Discussions and Conclusions

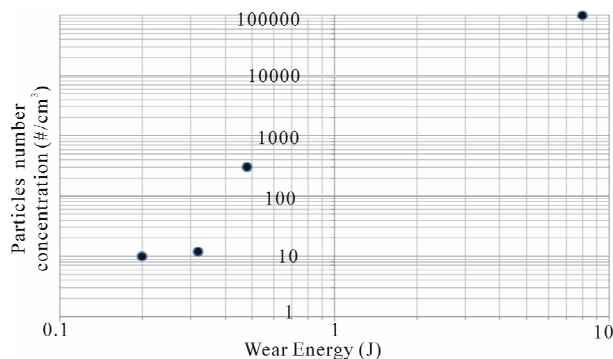
The study presented here investigates the release of nanoparticles during the operations related to the handling of and processing of a nano-coated shaft cap. For different types of wear, concentration levels of emitted particles in air have been measured. The whole experimental procedure was divided in nine carefully selected tests and done in a proper nano-secured facility. Using state of the art instrumentation, concentration and size levels of the emitted particles were quantified. There was no significant particle release during the handling operations (tests #1 to #6). There has been a presence of low concentration of micronic particles for the nano-coated part during tests #2 and #4. These results were also compared with the results obtained from a same part but uncoated. No considerable difference was observed between the two except for the test of surface scratching.

During the sanding test (#9), it has been found that the total number concentration of the released particles has been dominated by the nanoparticles with a mean diameter size of 35 nm. This conclusion agrees with the results of Koponen *et al.*, 2009 and Göhler *et al.*, 2010. This should be noted that because the aerosol was sampled directly in the vicinity of the particle generation source, the measured particle numbers represent maximum possible values in a practical case. The concept of wear energy has been introduced to the present study. It has proved to be a valuable indicator for forecasting the behavior of particles' emission. From the trend observed during experimental conditions, it may be suggested that with wear energy, particle emission increases. With the use of primary force-energy relationships, wear energy has been approximated and was found to be showing a positive correlation with particle emission. However, this hypothesis still needs more experimental trials to get verified.

Although the present study itself provides some of the valuable information on the particle emission behavior, it still demands both an improved energy model as well as refined experimental setup to understand the particle-energy interaction at microscale. The wear energy calculation must take into consideration that the frictional force is not a constant term. For the present study, there were limited number of test samples but authors will follow the work with more samples in order to provide a versatile asset to this study. As seen in the study, there has either been very weak release of the nanoparticles during handling operations or a strong release at higher energy levels. Therefore, in between these two extreme states, more sophisticated experimental procedures are

Table 4. Wear energy and particle emission associated with tests #6 to #9.

Test	Wear energy (J)	Total particle number concentration (#/cm ³)
Accidental fall	0.2	10
Scratching	0.32	12
Sawing	0.48	300
Sanding	8	100,000

**Figure 4. Total particle concentration versus dissipated energy.**

required to set and measure the applied wear energy. To verify the emissions and their characteristics, the scanning electron microscopy and chemical analysis of the fullerenes are required which will be followed in the subsequent work.

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