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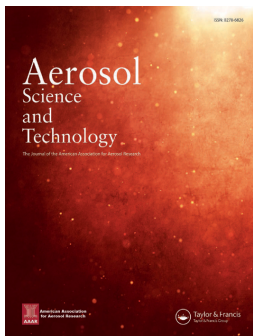
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In-service performance assessment of electrostatic precipitators serving a rubber vulcanization process

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ABSTRACT

We examined two emission abatement systems of some vulcanization ovens, serving a unit producing small rubber-based parts for automotive application. Each emission control unit treats the gases exhausted by three to five ovens. A heat exchanger cools down the fumes to a temperature suitable for the correct operation of a couple of two-stage electrostatic precipitators in series.

We performed quantitative analysis of concentrations and size distributions in these rubber fumes using aerosol technology instrumentation, namely optical particle spectrometers and electrical mobility particle sizers. The size of sampled particles was mainly between 100 nm and 1000 nm. We evaluated the performance of the exhaust fume abatement units, with focus on the electrostatic precipitator. Concerning batch ovens, the quantitative trend of the emissions

follows the thermal cycle of the post-curing process. Time interval since the last maintenance operation causes a gradual reduction of the removal efficiency.

The measured data demonstrate the reliability and the adequacy of aerosol instrumentation for the characterization of the emissions from rubber vulcanization ovens. The pair of electrostatic precipitators were shown to be effective in removing most of the particles detected in the fumes stream. The measurement protocol developed in this study allows assessing the influence of the maintenance schedule on the performance of the emission control units. New technologies for treating organic vapors can be evaluated in a reliable and effective way.

Introduction

Vulcanization irreversibly adds sulfur atoms to the elastomer molecules, improving their traction resistance and decreasing their vulnerability to oxidation (Callister 1999). Following the basic vulcanization process, molded elastomer parts frequently undergo "post-curing" treatments, such as exposure to UV radiation or further heating, which result in release of chemicals and formation of liquid or solid particles. However, phenomena of coagulation in fumes exhausted by ovens working at 220 °C yield mostly suspended compact particles having a size in the range of between a few nanometers up to ~1 µm (Kim et al. 2013).

Establishing effective control systems and emission abatement strategies minimizing, as well, the effects on the manufacturing chain is a crucial issue in production processes with high health and environmental hazards, as in the case of rubber vulcanization industry, which is characterized by carcinogenic exhausts (International Agency for Research on Cancer 1982).

An opinion from the Scientific Committee on Occupational Exposure Limits (SCOEL, Santonen *et al.*, 2016) defines rubber process fumes as “a variety of substances emitted from rubber compounds into a workplace atmosphere as a result of industrial processing, the composition of which depends on the formulation of the compounds concerned, the process technology in use, and the associated process parameters”. Thus, any discussion should consider all the possible physical forms in which they can be present in the workplace environment, i.e. gases, vapors, and aerosols (Willoughby 2003; Forrest 2015).

Due to lack of recent (post-2010) data on the variable composition of rubber dusts and fumes, it is not possible to provide a recommended threshold value for exposure of rubber dusts or fumes correlated to cancer risk (Santonen *et al.* 2016). However, some standards impose binding limit values to the emission of fumes exhausted by processes of rubber industry (Health and Safety Executive 2005b).

Detecting all the constituents of rubber fumes in an effective way is not a trivial task because of their complex chemical composition and their diverse forms. Nudel'man (2001) reported that aerosols from rubber goods production are formed both by mechanical stresses and local overheating at mixing stage and by condensation of emitted vapors of high-boiling-point substances during the vulcanization process. In both cases, the aerosol particles contain in addition a mix of substances captured by the liquid droplets forming the aerosol itself.

This implies that different sampling and analysis methods may affect the measurement outcome. In fact, concerning visible fume, i.e. the complex mix of aerosols derived from the condensation of hot vapors, the manner of monitoring is an integral part of the definition of rubber fume (Willoughby 2003). In particular, the development of a monitoring system for aerosols from rubber goods production should

carefully take into account the influence of plasticizer aerosols on the sampling process itself (Forrest 2015).

Giese (1999) followed the principles of applicable regulations, such as the German hazardous substances ordinance, to describe an efficient sampling and analysis method for monitoring air in the areas dedicated to vulcanization processes in rubber industry. Among the different substance categories present in field tests and analyzed, the aerosols were captured with glass-fiber filters, and vapors by XAD-2 polymer adsorption tubes. A 94.0-98.9% mass-based removal efficiency was measured.

Forrest mentions in his review (Forrest 2015) that Levin and Asplund captured aerosols and particulates released from a selection of rubber compounds employing quartz wool filters. Forrest also points out that Blanden and Isherwood employed an ESP and glass fiber filters in five British sites to collect the particulate matter in the fumes from different rubber production processes.

3M's Environmental Engineering and Pollution Control Laboratory (DyneonTM Fluoroelastomers, 2001) identified a two-stage dry or wet ESP as the most suitable abatement unit to reduce opacities and volatile matter in post-curing exhausts.

Kim et al. (2013) carried out a mapping analysis based on five 40-min particle number concentration measurements, conducted by counting particles with a diameter ranging from 0.02 to 1.0 μm at intervals of 30 minutes, using a CPC (Condensation Particle Counter). As a result, they obtained a concentration map of the rubber manufacturing plant, with a peak near the workbench.

Objectives

The experimental procedures described here mainly aimed at evaluating the efficacy of the electrostatic precipitators (ESPs) installed in a factory for the removal of particles generated by rubber post-curing.

Since the composition of volatile substances exhausted by rubber post-cured in the specific facility under examination is not known, it is not possible to state a priori whether electrostatic precipitation is the best available technology for the abatement of pollutants contained in the exhausts. Therefore, the first objective of the research project was the verification of the usefulness of ESPs in our specific case study, especially in lack of information from literature about the suitability of such mitigation measures in our specific case study.

Hence, a preliminary analysis was necessary to verify the presence of sufficient concentrations of particles and then correlate their size distribution to the performance of ESPs.

These elements led to the evaluation of the use of instruments like particle sizers for the quantification of the aerosol emissions from rubber post-curing ovens: the possibility of performing the sampling and the immediate analysis of data using the same instrument makes this solution very attractive. Some of the available aerosol instrumentation not only allows the analysis of the emissions with portable devices measuring with a time resolution on the order of seconds or minutes, but also providing estimates of the quantity of contaminants exhausted by the pieces undergoing vulcanization, even without knowing the chemical composition of these materials. That is particularly important when the chemical composition of vulcanized materials is patented and therefore undisclosed. Moreover, when one aims at evaluating the performance of devices that remove particulate contaminants, it is necessary to use the aerosol

instrumentation in order to combine and align the resulting efficiency with the data of concentrations. Indeed, sampling a chemical substance without knowing whether it is gaseous or liquid/solid does not allow evaluating the suitability of the technologies and devices used for controlling the emissions. We did not find any information about the use of aerosol instrumentation in the scientific literature for the purification of the exhaust fumes produced by this specific manufacturing process. In fact, the most common methods for the study of particles emitted by rubber curing are differential gravimetric analysis (DyneonTM Fluoroelastomers, 2001; Health and Safety Executive, 2005; de Vocht *et al.*, 2006) and Fourier transform infrared spectroscopy (Giese 1999). Hence, the aim of this paper is to provide scientific evidence about the possibility of using this kind of devices for the real-time detection of rubber fume emissions.

The paper provides a sound in-situ assessment of the performance of ESPs used for purifying vulcanization fumes: the results of the measurements, critically analyzed in this work, justify the adoption of ESPs as purifying devices for rubber fumes.

A further objective of the research work is to establish a correlation between the maintenance time interval and the removal efficiency of ESPs for typical rubber fumes. The performance of the abatement system progressively decreases, because of the build-up of particles on the collection plates, especially particles with high electric resistivity (Parker 2003).

Facility description

We studied two emission treatment units downstream of post-curing ovens, serving a plant producing small parts containing rubber for the automotive sector. We focused on the emissions exhausted by two different types of ovens, chosen by the technical staff of the factory. Five in-line ovens are connected to an emission control

unit, and three batch ovens are connected to another unit. The temperature of the studied post-curing ovens has a maximum of $220\pm 3^{\circ}\text{C}$.

The five in-line ovens (denoted as C_1 - C_5) work continuously, i.e. they steadily treat rubber parts that are loaded according to a uniform schedule with a quick opening and closing of the inlet shutter.

On the other hand, the three batch ovens (indicated as B_1 - B_3) follow an extended cyclic schedule. Each of the four daily cycles of oven B_2 lasts about six hours, consisting of a 25-min temperature ramp-up from 50 to 220°C , followed by a four-hour phase at a constant 220°C temperature, a final 50-min cool-down to 50°C and a 45-min pause at low temperature. The ovens B_1 and B_3 run three cycles per day lasting eight hours each: the difference from the schedule of oven B_2 consists in the additional two hours for the final low-T pause interval, as shown in Figure 1.

One of the two pollution control units (Figure 2) treats the fumes exhausted by the three batch ovens; the other one served the five in-line ovens. Each emission control unit removes particles from an exhausted flow rate of less than $1000\text{ m}^3\text{ h}^{-1}$ at a duct velocity of about 3.8 m s^{-1} . In each treatment unit, a heat exchanger cools the fumes down to a temperature value suitable for the correct operation of the downstream two-stage ESPs (around 30°C). Each stage of the precipitators has a corona-discharge ionization section and an array of parallel collection plates.

Protocol for efficiency measurements

We performed several measurements on particles in the exhaust fumes using the following instruments:

- TSI “Optical Particle Sizer (OPS) model 3330” (TSI 2011); the software used for the collection of data is the TSI Aerosol Instrument Manager (AIM).
- TSI “Nanoscan SMPS Nanoparticle Sizer model 3910” (TSI 2013); the instrument can be used in single mode by setting one particle size or scan mode (measuring particle concentration in 13 size channels between 10-420 nm). The data are provided by the TSI Nanoscan Manager (NM) software.

To know the size distribution of particles between 10 and 10,000 nm, we combined the data provided by those two instruments. We measured particle number concentrations with TSI 3910 operating in single mode between 10 and 400 nm. We set TSI 3910 in single mode, because size distributions in the exhaust air were fluctuating, resulting in “holes” at the lowest range of the spectrum (Figure 3). It should also be noted that although the TSI 3910 is specified to have a lower size limit at 10 nm, it may not be very accurate in its lower size range (Fonseca et al. 2016). Further, it cannot be excluded that particles are formed due to nucleation in the rubber curing process. These would be even smaller and not detectable with the TSI 3910.

We set the following particle size bounds for measurements with TSI 3330: 300, 400, 550, 700, 1000, 1300, 1600, 2200, 3000, and >10000 nm. In every test session, the dilution factor is set varying the bypass flow rate of sample fumes to the external diluter in order to avoid overloading the instrument measuring the particle concentrations upstream and downstream of the ESP.

Since the width of the size channels in the single size mode of the TSI 3910 is unknown, concentration data could not be normalized to $d\log(d_p)$. However, this is not essential here, as the main goal of using the single size mode was to monitor the variation of concentrations of particles of a known mobility size (TSI 2013).

The match when the scan mode was used by the TSI 3910 (Figure 3) shows good agreement between the two instruments at 350 nm.

We also performed validation tests and measurements of the removal efficiency approximately following ISO 16890:2016 (International Organization of Standardization 2016). Despite the differences among standards concerning contaminant conditions, scale and sampling (Tronville and Rivers 2005), we followed this International Standard for testing and rating air filters due to its minor variations for the equipment and procedure of particle capture efficiency measurement (Tronville and Rivers 2016). In general, we measured particle concentrations upstream and downstream of the ESP following the schedule in Table 1. Each of the seven sampling slots lasted three minutes (one for cleaning, two for the measurement of particle concentrations).

A sensitivity analysis using TSI OPS 3330 to measure particle concentrations optimized the schedule of Table 1. We found it desirable to perform a higher number of short-duration samples, getting a more accurate set of efficiency outcomes, rather than long-duration samples. The decision of sample duration is a trade-off aimed to reduce the time for measurements and enhance the goodness of results.

We obtained three different efficiency values $E_{i,j}$ for the j^{th} size channel in the i^{th} sampling period, then averaged to get a single representative efficiency result:

$$E_{i,j} = 1 - \frac{D_{i,j}}{\frac{U_{i,j} + U_{i+1,j}}{2}} \rightarrow E_j = \frac{\sum_{i=1}^3 E_{i,j}}{3} \quad (1)$$

Here U and D are the upstream and downstream particle concentrations, respectively.

Test rig and validation

In-situ validation measurement in controlled conditions

At first, we investigated the reliability of the results of measurements on the emission abatement units under examination. Therefore, we carried out a validation of the test rig by comparing the outcomes obtained in situ with those of a laboratory apparatus, operating according to international standards and in controlled conditions (Figure 4).

The larger circle in Figure 4 shows the location of the inlet for Di-Ethyl-Hexyl-Sebacate (DEHS) aerosol into the test rig, and the location of the sampling probe upstream of the ESP. A diagonal baffle plate cut short at its lower end is installed across the tube between these two locations: the small gap generates a turbulent flow at this point with high velocity which mixes the aerosol in the exhaust stream, hence ensuring a more constant concentration at the sampling point upstream of the ESP.

Three different efficiency values $E_{i,j}$ were obtained for the j^{th} size channel in the i^{th} sampling period, and then averaged to get a single efficiency result. We repeated the sampling procedure described in Table 1 four times, progressively modifying the value of concentration k , obtained regulating the pressure of the inlet flow of compressed air and consequently changing the dilution factor to avoid overloading the particle sizer with excessively high inlet concentrations. We replicated this measurement for four days (index d) in the same conditions and following the same method, in order to appreciate variations of fractional removal efficiency:

$$E_{d,i,j,k} = 1 - \frac{D_{d,i,j,k}}{\frac{U_{d,i,j,k} + U_{d,i+1,j,k}}{2}} \rightarrow E_{d,j,k} = \frac{\sum_{i=1}^3 E_{d,i,j,k}}{3} \rightarrow E_{d,j} = \frac{\sum_{k=1}^4 E_{d,j,k}}{4} \rightarrow E_j = \frac{\sum_{d=1}^4 E_{d,j}}{4} \quad (2)$$

Figure 5 compares the performance against DEHS aerosol of precipitators serving the batch ovens with the fractional efficiency curve of the ESPs downstream of the in-line ovens during usual post-curing operation (data come from preliminary tests assessing the effective operation of the ESP systems). The two curves of fractional removal efficiency vs. optical dimension of particles agree fairly well, although the particle concentrations vary in the two measurements due to different operating modes and sampling conditions of the two systems.

Validation with measurements in laboratory

The second step of the study aimed at evaluating whether the portable test apparatus used in situ provided data consistent with those measured by standard ISO 16890:2016.

As a first step, the size distribution of the DEHS aerosol produced with the portable test setup (Figure 4) used for in-situ measurements was compared with the one generated by a Laskin nozzle for standardized tests in the research laboratory on filtering media of the Department of Energy (DENERG) at Politecnico di Torino. As shown by the log-linear curves in Figure 6, the two size distributions agree quite well.

The layouts of the two setups studied present several differences that may lead to mismatching efficiency outcome:

- in the lab, an all-in-one program developed in-house is used instead of AIM and NM software by TSI: it allows at the same time to run the particle sizer, modify the speed of fans of the circuit to set the correct flow rate, and manage the sampling procedure;
- in the lab, a system of three different one-way valves automatically manages the opening and closing of upstream/downstream sampling line or cleaning HEPA unit following the schedule shown in Table 1. On the contrary, the

management of the sampling system for the in-situ measurements relies on a manual three-way valve.

We performed some further tests to verify the coherence between the portable test setup used in situ and the laboratory test apparatus. In the same test sessions, we also checked the consistency among different particle sampling instruments by measuring the removal efficiency of a filter with well-known performance in four different ways:

- TSI OPS 3330 with the lab test rig and in-house software;
- TSI OPS 3330 with the portable rig to be used in situ and AIM software;
- TSI SMPS 3910 with the portable rig to be used in situ and NM software;
- PMS LAS-X II with the lab test rig and in-house software.

We checked whether the efficiency data obtained with different devices are coherent. We set the SMPS 3910 spectrometer in single mode to the geometric mean size of the first sampling channel of the OPS 3330, i.e. at 346 nm. The uncertainty associated to particle concentration measurement performed in single mode with TSI 3910 does not affect appreciably the measured device efficiency because the efficiency is determined by the ratio of the upstream and downstream particle concentration. The results of a past inter-laboratory validation analysis shows good agreement of the efficiency data measured in single mode using the TSI 3910 and the data provided by more sophisticated instrumentation (Sachinidou et al. 2017).

Figure 7 shows the agreement of the results of the four different test setups specified before, confirming the goodness of the efficiency curve of the OPS 3330 and the coherence of the efficiencies obtained with SMPS 3910 and OPS 3330 in the overlapping sampling channel.

Measurements with ovens in service

Before proceeding with the analysis of efficiency measurements, we report some preliminary outcomes regarding the response of particle concentrations during the normal operation of batch ovens.

Influence of the operation of batch ovens on particle concentration

Unlike in-line ovens, batch ovens follow a cyclic, discontinuous schedule that we suppose affects the evolution in time of the particle concentration. In fact, this variation of the oven temperature determines a progressive evaporation of substances, starting from those characterized by a low boiling point.

We examined the variation of particle concentration across the cooler and the ESP in the emissions control unit serving the batch ovens. We employed both SMPS 3910 and OPS 3330, with sampling durations long enough to show the change in particle concentrations.

The results of both the tests (shown in Figure 8 and Figure 9) confirmed our expectations: apart from the changes due to the sampling uncertainty, the macroscopic evolution in time of particle concentrations matches fairly well the trend of the temperature inside the ovens. In fact, we can notice a quick increment during the startup, a plateau when the temperature stays constant, and a reduction in the cool-down phase. In some cases, the particle concentration exceeded the maximum value accepted by the instruments (especially when using OPS 3330, despite adopting the highest dilution factor possible).

Figure 10 shows the evolution in time of the particle number concentrations sampled with OPS 3330 and averaged on an hourly basis: it clearly highlights the correlation between the oven temperature (cf. Figure 9) and the particle concentrations

in the exhaust fumes. We have performed the measurements at the downstream sections of both the cooler and the ESP. One can also notice the progressive decrease of the measured particle concentrations as the optical diameter of particles increases.

Efficiency measurement of ESPs serving in-line ovens

We carried out a set of tests with OPS 3330 focused on the evolution in time of the removal efficiency of the ESPs placed in the emission control unit serving the in-line ovens. The analysis proved that the time elapsed from the last maintenance operation on the ESP is a key parameter influencing its performance. In fact, the degree of obstruction across the ESP due to the deposit of particles on the arrays of collection plates clearly depends on the maintenance schedule. This phenomenon is evident at the entrance of the capture section (Figure 11) and inevitably alters the uniformity of the electric field, with clear effects on the removal efficiency against particles.

To verify the progressive reduction of removal efficiency with the increasing clogging of particles on the collection plates, we carried out six efficiency tests throughout four days between two subsequent maintenance operations planned on Mondays and Thursdays around noon. The results in Figure 12 all refer to the ESP in the abatement system serving in-line ovens. Our expectation is confirmed by the evident proportionality linking the degradation of fractional removal efficiency in consecutive tests with the cumulative amount of post-cured rubber pieces, representing the activity of the ovens (numbers provided in Figure 12).

Discussion

All the experimental tests carried out on the emission abatement system serving the batch ovens highlighted the strict influence of the temperature of the fumes on

particle concentration. We also noted that the size of sampled particles is between 100 nm and 1,000 nm, with the mode around 150-200 nm.

Concerning batch ovens, the quantitative trend of the emissions follows the thermal cycle of the post-curing process. In particular, the peak occurs around 20 – 30 minutes after the startup: this is the approximate time for rubber parts to reach the maximum temperature after the evaporation of low-boiling substances. The peak first relates to fine particles that rapidly evaporate in the ovens; afterwards, complex compounds are post-cured giving rise to larger particles detected by the instruments.

ESPs can ensure a good removal of such pollutants, at least in the control unit serving the in-line ovens, if maintenance follows a suitable schedule and is effectively carried out. In fact, the time interval since the last maintenance operation is an insidious variable of the system since it causes a gradual reduction of the removal efficiency, particularly against low-diameter particles.

In this regard, we observed the reduction of removal efficiency in time, i.e. as a function of the cumulative number of processed pieces or the time passed since the last maintenance intervention. Figure 13 highlights that this performance loss against particles with optical diameter equal to 350 nm is linear. In the sampling period of 72 hours, the efficiency continuously dropped from around 99% to 85%, which means a 14-fold increase in the emission. The direct proportionality between the efficiency of the ESP and the mass of treated rubber pieces is valid also in the case of batch ovens: the decrease of the efficiency is much slower, probably because the ESP serving the batch ovens has been manufactured later than the one used for the in-line ovens.

The influence of maintenance on the removal performance is evident in Figure 14, where the concentration of 350-nm particles measured with Nanoscan 3910 in single mode upstream of the ESP is almost double with respect to the downstream value: this

implies a removal efficiency of about 50%. This value is very low with respect to the typical performance indicators of ESPs and may be correlated to an ineffective schedule for the maintenance of the batch control unit.

Conclusions

A preliminary sensitivity analysis regarding the measuring protocol of emissions from post-curing ovens points out that it is desirable to perform a high number of short-duration samples, getting a more accurate set of efficiency outcomes, rather than a single long-lasting sample unavoidably associated to uncertainty due to unpredictable trends of the particle concentrations.

We carried out several validation steps. The DEHS aerosol produced by the portable test setup for in-situ validation tests is characterized by a size distribution that agrees fairly well with that of the aerosol employed in laboratory tests. The portable test apparatus also gave coherent efficiency data as confirmed by the use of different aerosol measuring instruments. Thus, DEHS aerosol seems to mimic well the behavior of the particles contained in rubber fumes.

The results of the experimental tests carried out in this work confirmed our decision of using aerosol instrumentation for monitoring and measuring the fractional removal efficiency of the system intended to remove the pollutants in rubber post-curing fumes. Precipitators show a remarkable removal efficiency, usually over 90% for all particle sizes if maintenance follows an adequate schedule. In fact, time passed since the last maintenance operation causes a progressive degradation of the removal efficiency, particularly against low-diameter particles: a suitable maintenance strategy should limit this detrimental phenomenon, while limiting costs as much as possible. We can assume that continuous monitoring systems with low-cost sensors would be beneficial to optimize maintenance intervals. However, it is firstly necessary to check

the correlations with more expensive and accurate instruments, in order to be sure that low-cost devices would be able to sample exhausted particles without being overloaded. Indeed, the main limitation is that, at the moment, there is no low-cost diluter on the market.

In the plant under examination, the centralized ventilation strategy employed mixing of the contaminants with the air of the workplace. Instead, a control and abatement system, installed close to the source of emissions would be preferable. In fact, the removal efficiency of ESPs operating against highly concentrated pollutants in lower, undiluted flows of fumes would probably increase. This enhancement requires the precipitator maintenance schedule to be changed in order to avoid the clogging of the collection plates.

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Figure 1. Cyclic schedule of the batch post-curing ovens

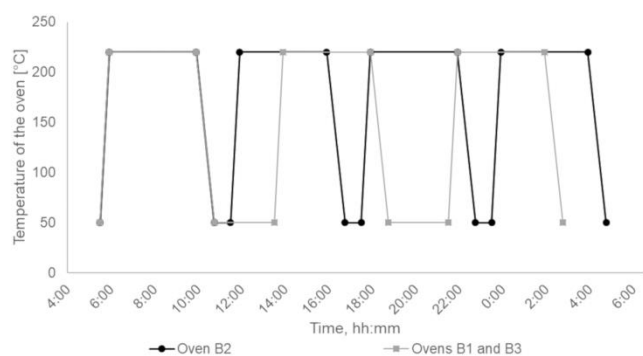


Figure 2. Emission abatement system of the facility studied

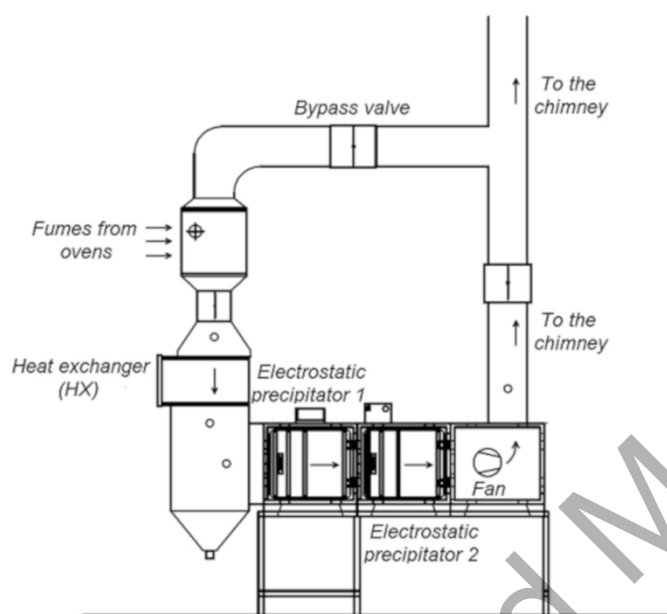


Figure 3. Typical normalized size distribution of the aerosol downstream of post-curing ovens

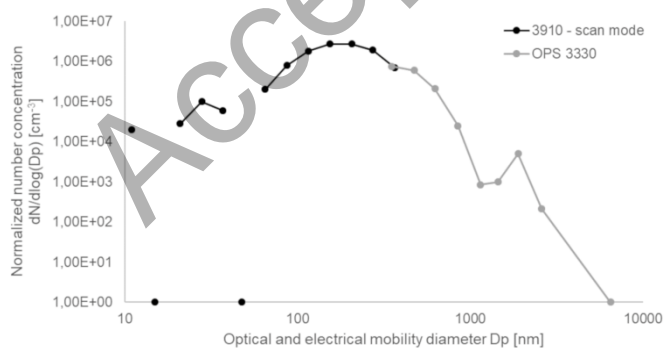


Figure 4. Portable test rig for the validation tests performed using DEHS

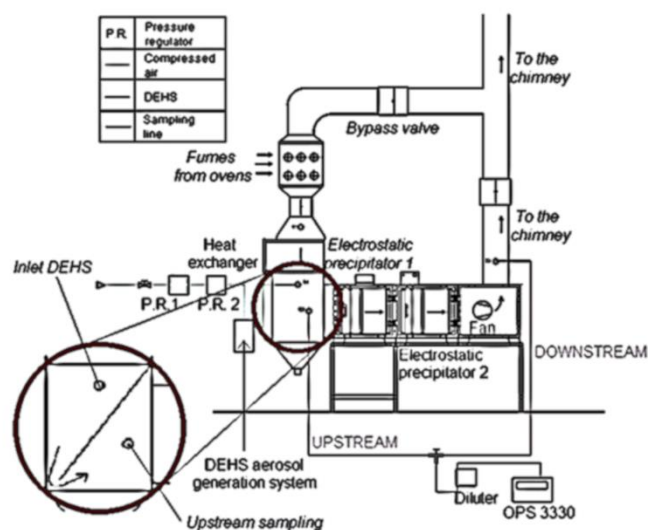


Figure 5. Validation of measurement with OPS 3330 through the comparison of fractional removal efficiency (data not shown up to 10 μm , due to few particles)

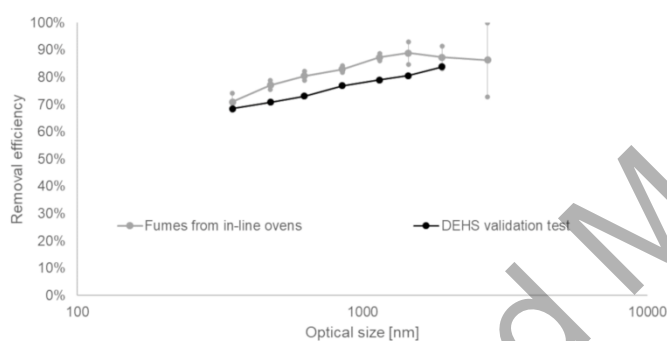


Figure 6. Size distributions of aerosols produced with different test setups

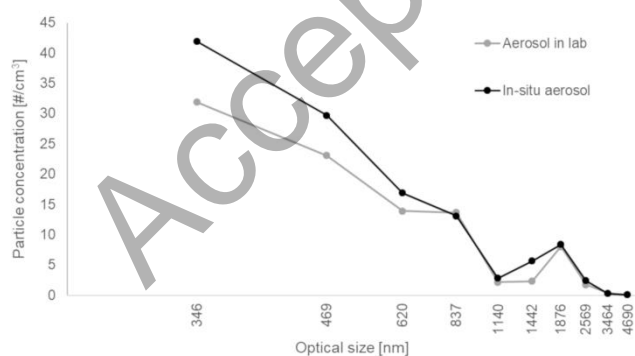


Figure 7. Comparison of fractional removal efficiency measured with different instrumentation and configurations

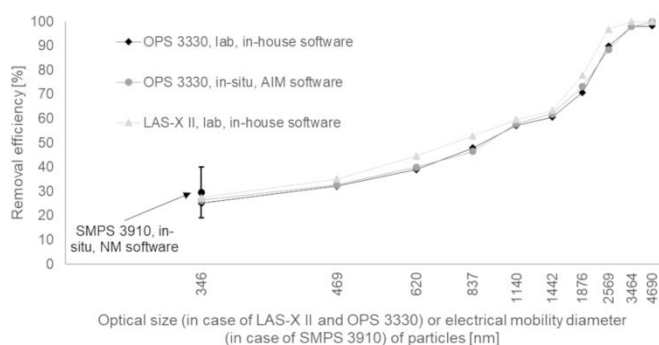


Figure 8. Particle concentration vs. time across the ESP downstream of batch ovens measured with SMPS 3910 – 100 nm – No dilution

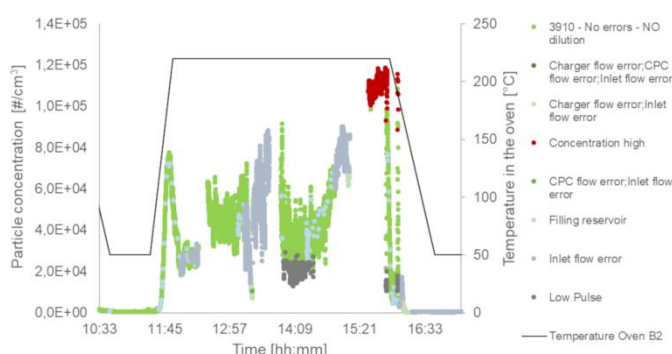


Figure 9. Particle concentration (downstream section of the cooler serving the batch ovens) vs. time measured with OPS 3330 - 350 nm - Dilution 1:15 before 11:16, 1:100 afterwards

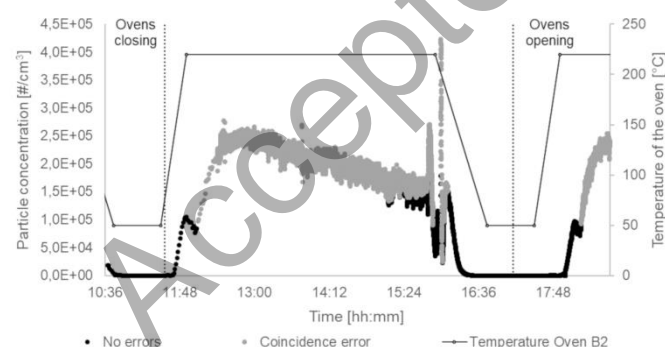


Figure 10. Hourly average particle concentrations downstream of batch ovens vs. time. Measured with OPS 3330 – Dilution 1:370 for cooler; for precipitator, 1:200 between 17:39 and 18:37, 1:300 afterwards

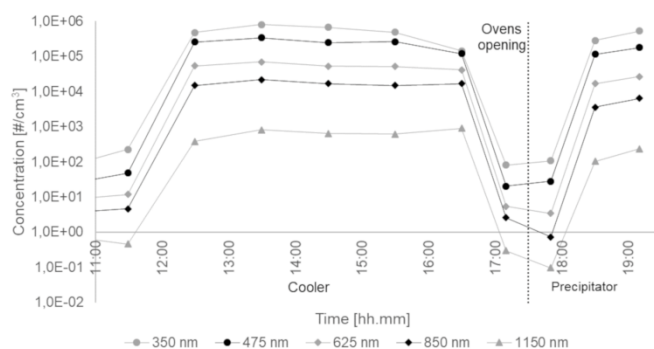


Figure 11. Top view of collection plates with progressive build-up of particles (fumes enter from left side)



Figure 12. Evolution in time of the fractional removal efficiency of ESPs serving in-line ovens; the number of pieces refers to the number of rubber pieces treated inside the oven since its last maintenance

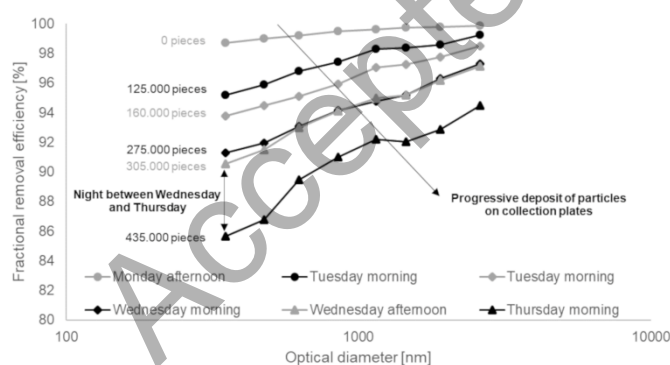


Figure 13. Dependence of the removal efficiency towards 350-nm particles on the amount of post-cured rubber parts

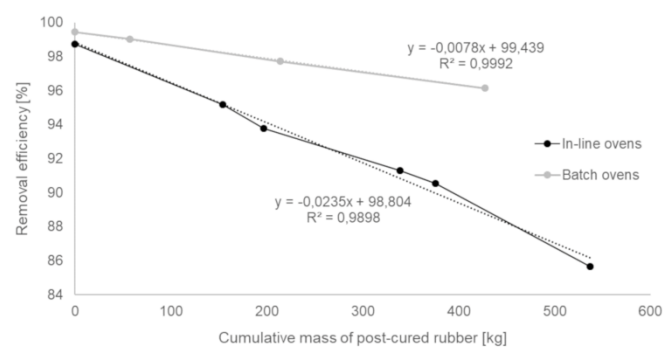


Figure 14. Particle concentration vs. time across the ESP downstream of batch ovens measured with SMPS 3910 – 350 nm – No dilution

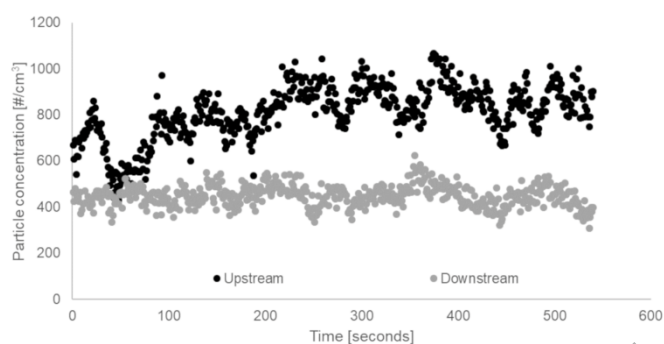


Table 1. Schedule of the samplings for one single measurement of efficiency

1		2		3		4		5		6		7	
Cleaning	U ₁	Cleaning		Cleaning	U ₂	Cleaning		Cleaning	U ₃	Cleaning		Cleaning	U ₄
			D ₁				D ₂				D ₃		
Efficiency N.1						Efficiency N.3							
				Efficiency N.2									