

# A TGA/FT-IR study for measuring OC and EC in aerosol samples

P. Fermo<sup>1</sup>, A. Piazzalunga<sup>1</sup>, R. Vecchi<sup>2</sup>, G. Valli<sup>2</sup>, and M. Ceriani<sup>2</sup>

<sup>1</sup>Dipartimento di Chimica Inorganica, Metallorganica e Analitica, University of Milan, Via Venezian 21, 20133 Milan, Italy

<sup>2</sup>Istituto di Fisica Generale Applicata, University of Milan, via Celoria 16, 20133 Milan, Italy

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**Abstract.** Carbon analysis consists in the evaluation of the carbonaceous content of the aerosol (TC) but, more importantly, of its distribution between the two components EC (Elemental Carbon) and OC (Organic Carbon) that are characterized by different physical-chemical properties. In spite of the numerous studies focused on this topic, nowadays a universal methodology for the determination of the two components EC and OC is not available. In fact OC and EC (also known as black carbon or soot) are operationally defined by the method of analysis and, as a consequence, different methods can produce different results.

In this paper we present results on the application of TGA/FT-IR (Thermo-gravimetric Analysis/Fourier Transformed Infrared Spectroscopy) to the characterization of carbonaceous aerosols. The analytical methodology was applied to PM<sub>10</sub> (particulate matter with aerodynamic diameter smaller than 10  $\mu\text{m}$ ) four-hour time resolution samples collected in Milan urban area. The method is a two-steps thermal one and it is based on the different thermal behaviour of OC and EC. It has been set up analyzing suitable standards containing both organic and elemental carbon. Carbon quantification is achieved by on-line, continuous monitoring of CO<sub>2</sub> infrared absorption at 2361  $\text{cm}^{-1}$ . A good separation between OC and EC on particulate matter (PM) samples has been obtained. Ranges and average values were 12–70  $\mu\text{g}/\text{m}^3$  and 20  $\mu\text{g}/\text{m}^3$  for OC and 0.2–6  $\mu\text{g}/\text{m}^3$  and 2  $\mu\text{g}/\text{m}^3$  for EC. On average OC and EC made up 29 ( $\pm 13$ )% and 2.5 ( $\pm 1.8$ )% of the PM<sub>10</sub> fraction, respectively. The method reliability has been verified by a preliminary comparison with TOT (Thermal Optical Transmission) technique. OC and EC values determined for ambient samples of PM<sub>10</sub> were correlated with meteorological parameters as well as with Radon concentrations.

## 1 Introduction

The overall composition of the particulate matter (PM) has become one of the fundamental topics in the field of environmental chemistry. One of the main topics is represented by carbonaceous fraction quantification since this component makes up about the 50% of the particulate mass concentration.

The total carbon content (TC) includes elemental carbon (EC) and organic carbon (OC) as well as a small percentage (less than 5%) of inorganic carbon mainly present as carbonate.

EC is essentially a primary pollutant (Seinfeld and Pandis, 1998) emitted during incomplete combustion of fossil and biomass carbonaceous fuels. In urban areas, diesel emissions are one of the major sources for black carbon, which is often used as a marker for urban pollution (Delumyea et al., 1980; Salma et al., 2004); furthermore, its temporal pattern could be related to traffic intensity (Ruellan and Cachier, 2001).

OC has both primary and secondary origin. Primary OC is mainly formed during combustion processes such as unleaded gasoline combustion in urban area or biomass and field agricultural burning (Duan et al., 2004). It is also directly emitted as plant spores, pollens and soil organic matter. Secondary OC can originate from different processes such as gas to particle conversion of low vapour pressure volatile organic compounds, condensation and physical and chemical adsorption. The presence of secondary organic aerosol (SOA) is suggested by an increase of the OC/EC ratio. Secondary organic aerosol can be easily estimated using EC as tracer of OC primary emission (Turpin and Huntzicker, 1995; Salma et al., 2004). As for OC chemical composition, this fraction contains a large amount of organic substances belonging to different classes (aliphatic or aromatic compounds, acids, etc.) the majority of which has not yet been identified.

Correspondence to: P. Fermo  
(paola.fermo@unimi.it)

EC is also indicated as black carbon (BC) because of its color. It has a graphitic-like structure with the presence of some functional groups containing elements such as oxygen, sulphur, hydrogen and nitrogen, which are able to enhance catalytic processes. EC is sometimes referred as *soot* when carbon to oxygen ratio during the combustion process is less than 1. Indeed soot, that represents the dark component of the carbonaceous aerosol, is a very complex mixture of both elemental carbon and highly polymerized organic substances.

The large concern on elemental carbon (EC) concentrations in PM samples is due to the adverse health effects (Summerhays, 1991; Oberdörster and Yu, 1990) and soiling of surfaces. At a global scale, EC might also play a role in radiative forcing effects, as it is the dominant light-absorbing component of atmospheric aerosols. As regards to the organic carbon (OC) the increasing awareness has to be attributed to the fact that it typically constitutes up to 70% of the total dry fine particle mass in atmosphere. Moreover, OC is an aggregate of hundreds of individual compounds, which can modify the thermodynamic and chemical properties of the atmosphere (Turpin et al., 2000).

The large number of studies carried out in the last years and devoted to OC and EC quantification, have demonstrated that, because of the carbonaceous aerosol complex nature, it is difficult to compare results produced by means of different techniques (Schmid et al., 2001; Chow et al., 2001, 2004). In fact, these methods gave results in fairly good agreement just with respect to TC concentration while difficulties have been encountered in the determination of the two fractions i.e. EC and OC.

It is worthy to note that up to now a reference method of analysis is not available and there is a significant disagreement in the experimental results (Schmid et al., 2001).

The methods most commonly applied are based on thermal evolution and reflectance/transmission measurements. Amongst them the IMPROVE (Chow et al., 1993) and the NIOSH (Birch and Cary, 1996) protocols are widely employed. Both methods require the sample (i.e. PM deposited on a filter support) first heating to 600°C in pure He and, after the temperature has decreased to 400°C, a second heating to 600°C in He containing a small percentage of O<sub>2</sub>. During the first step organic substances are removed since OC is less thermally resistant than EC. However, at the same time OC can partially be converted into EC through a pyrolysis. Because of this process, known as *charring*, OC would be underestimated. In order to overcome this problem, a pyrolysis correction is applied monitoring a laser beam transmitted or reflected by the filter. The IMPROVE protocol adopts the thermal-optical reflection (TOR) method while the NIOSH protocol uses the thermal-optical transmission (TOT) method. The comparison of NIOSH and IMPROVE carbon measurements has shown that they are equivalent as concerns TC while NIOSH EC is lower than IMPROVE EC (Chow et al., 2001). The discrepancy is mainly due to differences in the temperature steps and to the different correc-

tion system. Moreover, a recent comparison between TOR and TOT methods has shown that the EC concentration determined by TOT is often a factor two lower than the corresponding value determined by TOR (Chow et al., 2004).

Another well-established thermal method for OC/EC analysis has been proposed by Cachier et al. (1989). In this procedure inorganic carbon is preliminarily eliminated by exposure of the filter to HCl vapours. Afterwards, half a filter is heated at 340°C in oxygen in order to decompose the whole OC. The remaining carbon, i.e. EC, is then determined by coulometric titration. The remaining half of the filter is directly analyzed without the pre-treatment. OC concentration is determined by difference. This protocol is known as LSCE (Laboratoire des Sciences du Climat et de l'Environnement, Cachier et al., 1989).

A thermal method quite similar for some aspects to the LSCE protocol has been proposed in a preliminary work carried out by Fermo et al. (2003). In this study the carbonaceous component in the Milan urban particulate matter was estimated by means of a thermo-gravimetric analyzer combined with an infrared spectrophotometer (TGA/FT-IR). In the present paper the employed technique is the same but the protocol has been partially modified in order to overcome some drawbacks of the previous methodology such as a not satisfactory OC/EC separation and, in some cases, a not sufficient sensitivity.

The aims of this study are:

- to test the reliability of TGA/FT-IR technique for the detection of the carbonaceous component in particulate matter samples;
- to quantify OC and EC concentrations on PM samples collected in Milan (Italy) in a field campaign carried out during winter 2003.

The method set up has involved the analysis of both standard samples prepared in our laboratory and ambient samples of Milan urban particulate matter.

## 2 Experimental

### 2.1 Standards preparation

To optimize and calibrate the analytical technique applied in this study, some “standard samples” (carbon black or graphite mixed with SiO<sub>2</sub> and some suitable compounds) were generated. These standard mixtures have been prepared so that their carbon concentration, both for organic and elemental carbon, was similar to the one expected in the atmospheric aerosol samples.

For this reason, among the organic substances, some of those present in the particulate matter in higher concentration were chosen (Seinfeld and Pandis, 1998).

The following mixtures were prepared:

- *standard a* containing: SiO<sub>2</sub> (80%) and Na<sub>2</sub>C<sub>2</sub>O<sub>4</sub> (20%)
- *standard b* containing: SiO<sub>2</sub> (85%) – palmitic acid (5%) – stearic acid (5%) and benzoic acid (5%)
- *standard c* containing: SiO<sub>2</sub> (70%) – Na<sub>2</sub>C<sub>2</sub>O<sub>4</sub> (5%) – palmitic acid (5%) – stearic acid (5%) – benzoic acid (5%) – carbon black (10%)
- *standards d1–d3* containing: SiO<sub>2</sub> and graphite in different concentrations, i.e. 4.7%, 1730 ppm and 3300 ppm, respectively.

The substances employed for the preparation of the standard samples are commercial products. Graphite powder was obtained grinding E441 Shaped Carbon Rods (Assing, Italy); carbon black was a powder of Monarch (Cabot Corporation, France); silica, sodium oxalate, palmitic acid, stearic acid and benzoic acid were supplied by Aldrich (Italy).

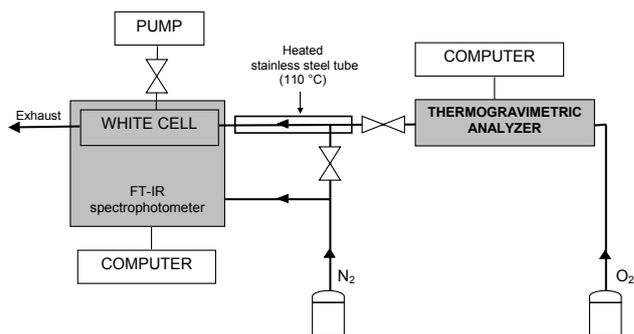
As for the mixtures preparation, the chosen materials were at first mixed in an agate mortar and then in a ball mill (Retsch model S1, operating with three balls, 3 mm diameter) in order to obtain a homogeneous sample of powder with small grain dimensions to be deposited on the filter supports.

The carbon concentration in our standards has been verified by CHN (Carbon/Hydrogen/Nitrogen Analyzer, Perkin Elmer instrument model 2400). In particular, as for standards d1–d3 used in the instrument calibration, the actual carbon concentrations were verified by CHN for standard d1 and by TOC (Total Carbon Analyzer, Dohrmann Apollo 9000 instrument equipped with a NDIR-detector and a solid sampler allowing to analyze small sample quantities) for standards d2 and d3. The use of TOC was necessary as CHN was not suitable to analyze samples d2 and d3 because of their low carbon content. It has been found that the measured carbon concentrations (i.e. those used for the calibration curve; see Sect. 3.2) were slightly lower than those calculated on the base of the weighed quantities, probably because of some losses occurred during mixture preparation.

The NIST (National Institute of Standards and Technology) Standard Reference Material 1649a (urban dust) and the NIES (National Institute for Environmental Studies) Certified Reference Material No. 8 (vehicular exhaust particulate) have been analyzed to test the method reliability.

In order to realize measurement conditions quite similar to those typical of particulate matter samples, standard d1 has been deposited on quartz fiber filters and these filters were used for the TGA/FT-IR calibration (see further on in the text).

The deposition was done by means of a home-made re-suspension system composed of two differently sized chambers in order to get a homogeneous dispersion of the powder mixture in the chambers volume. At the inlet of the air flow



**Fig. 1.** Diagram of the carbon analyzer obtained by coupling a thermo-gravimetric analyzer (TGA) with an infrared spectrophotometer (FT-IR).

there is a filter to prevent the entrance of ambient particulate matter in the re-suspension system.

The re-suspended powder was collected on the filter support (quartz fiber filters, Whatman QMA) after passing a size-selective inlet constituted by a cyclone, which allowed the sampling of particles with aerodynamic diameter smaller than 3.5  $\mu\text{m}$ . The grains size was checked by scanning electron microscopy (SEM) using a Hitachi 2400 instrument equipped with a Quantum Kevex energy dispersive X-ray microanalyzer (EDX).

Before and after the depositions the quartz fiber filters were exposed for 24–48 h on open but dust-protected sieve-trays in an air-conditioned weighing room ( $T=20\pm 1^\circ\text{C}$  and  $R.H.=50\pm 3\%$ ). The gravimetric determination of the mass was carried out using an analytical microbalance (precision 1  $\mu\text{g}$ ), which is installed and operates in the weighing room. In order to remove static electricity from filters the balance was equipped with a special kit consisting in a Faraday shield.

## 2.2 Analytical techniques – TGA/FT-IR

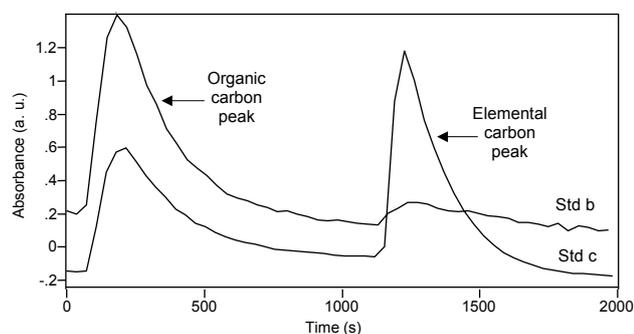
EC and OC determination has been performed by a TGA/FT-IR (Thermo-gravimetric Analysis coupled with Fourier Transformed Infrared Spectroscopy) home-made instrument, assembled using a JASCO-FTIR spectrophotometer Model 360 and a Dupont Thermo-gravimetric analyzer model 951. A schematic diagram of the carbon analyzer apparatus is reported in Fig. 1.

Basically, the method is a thermal evolved one where carbon quantification is achieved by on-line continuous monitoring of CO<sub>2</sub> infrared absorption. At the moment, the facility of the thermo-gravimetric analysis (weight loss evaluation) is not used for carbon quantification.

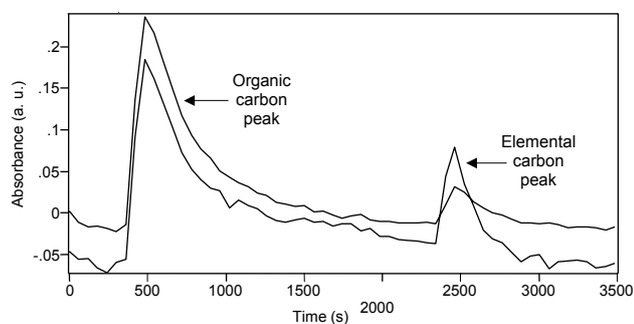
The TGA/FT-IR system is also advantageous because it allows the continuous monitoring of CO<sub>2</sub> as well as of other species (e.g. water vapour, SO<sub>2</sub>, NO<sub>x</sub> and NH<sub>3</sub>), which volatilize during the decomposition of sulphates and nitrates contained in the particulate matter. Indeed, from preliminary

**Table 1.** Experimental conditions used for TGA/FT-IR analyses; standards composition is reported in Sect. 2.1.

Experiment No.	Sample	First step T (°C)	First isotherm duration (min)	Second step T (°C)	Second isotherm duration (min)	Carbon % recovered in the first step
1	Std a	350	120	700	10	72
2	Std a	425	10	700	10	84
3	Std b	350	10	700	10	60
4	Std b	425	10	700	10	90
5	ambient sample	435	25	700	10	
6	Std b	435	25	700	10	90



(a)



(b)

**Fig. 2.** CECs (CO<sub>2</sub> Evolution Curves) obtained by TGA/FT-IR analyzing standard b and c (see Fig. 2a) and two particulate matter samples deposited on quartz fiber filters (Fig. 2b); the peaks due to organic carbon (OC) and elemental carbon (EC) are indicated with arrows.

measurements on a few PM samples both SO<sub>2</sub> and NO<sub>x</sub> have been detected from infrared spectra; nevertheless, the quantification of these species will require a suitable calibration and work is still in progress.

The system consists of a furnace (which is the core of the thermo-gravimetric analyzer) where the sample is heated under controlled conditions and the carbonaceous aerosol fraction is oxidized to CO<sub>2</sub> whose concentration is monitored by the FT-IR spectrophotometer. Two increasing tempera-

ture steps allow the separation of organic carbon from elemental carbon. CO<sub>2</sub> evolved during sample decomposition is transferred through a stainless steel tube heated at 110°C to a 15 cm long multipass White cell equipped with KBr windows (thus, the optical path reaches a length of 120 cm), which is located inside the infrared spectrophotometer. In this way, during each thermo-gravimetric analysis, infrared spectra series are collected automatically by the FT-IR with a spectrum collected every 60 s. By continuous monitoring of CO<sub>2</sub> absorbance at 2361 cm<sup>-1</sup> (which corresponds to the most intense signal in CO<sub>2</sub> infrared spectrum) vs. time, a CO<sub>2</sub> evolution curve (in the following called CEC) is obtained. The curve shows distinctive features and it is possible to distinguish, on the base of the different decomposition temperature, the two components OC and EC detectable as well separated peaks. CECs examples are shown in Fig. 2a–b. These curves were obtained analyzing both standards b and c, prepared as described in the experimental section (Fig. 2a), and a couple of four-hour time resolution particulate matter samples collected in Milan as described further on in the paper (Fig. 2b). Two fully separated peaks are detectable, the first one due to organic carbon and the second one due to elemental carbon. The area under each peak is proportional to OC and EC content in the sample. It is worthy to note that the small peak corresponding to EC in the CEC of standard b has to be assigned to some organic carbon *charring* since this standard mixture doesn't contain any elemental carbon. Furthermore, the EC peak falls at higher time values in the real samples since the experimental conditions were slightly different. In fact, CECs corresponding to standards b and c were acquired by using the heating program reported for experiment No. 4 (Table 1) while the ambient samples were analyzed as described in experiment No. 5 (Table 1).

All TGA/FT-IR analyses were carried out in O<sub>2</sub> atmosphere with a gas flow rate of 110 mL/min. Before each measurement the White cell was evacuated and purged by a nitrogen flux. Subsequently, oxygen was fluxed and the background spectrum acquired. CO<sub>2</sub> is carried through the stainless steel tube and the White cell by means of O<sub>2</sub> flux (Fig. 1).

A two-step heating procedure has been optimized in order to minimize both charring of organic carbon and pre-combustion of elemental carbon. The heating ramp, suitable to obtain the best separation between EC and OC, consists of a first jump at 435°C, an isotherm for 25 min, a jump at 700°C and an isotherm for 10 min. The heating program is displayed in Fig. 3. Each jump takes about 2 min with a heating preset rate of 160°C/min (corresponding to a rate of increase of about 200°C/min as shown in Fig. 3). The system takes about 5–10 min to equilibrate at the two desired temperatures of 435°C and 700°C. Each analysis takes altogether 55 min.

The analysis conditions, i.e. the two-step temperatures, the atmosphere and the isotherm duration, have been set-up analyzing standard samples (described in Sect. 2.1) and will be discussed later on in this paper.

In order to perform the TGA/FT-IR calibration, known quantities of standards d1-d3 were analyzed.

Thereafter, for each sample, the area of the peak due to elemental carbon was integrated and the obtained values plotted against the corresponding carbon concentrations. The instrument calibration has been carried out analyzing both powders of samples d1-d3 and standard d1 re-suspended on quartz fiber filter (Sect. 3.1). When analyzing real particulate matter samples (Fig. 2b), the quantification is achieved by single-peak area integration (since the two peaks corresponding to OC and EC are fully separated) taking into account the point where the baseline regains its initial value.

It is worthy to note that suitable standard carbon concentrations have been selected according to the carbon concentrations expected both for ambient samples analyzed in this study (see Sect. 3.3) and for SRM1649a. In this way the concentrations determined in the samples fall exactly within the calibration ranges.

For TGA/FT-IR analysis one half of the quartz filter (corresponding to a deposit area of 6 cm<sup>2</sup>) was accurately cut from the exposed filter area or from the standard sample. The filter was then divided into small pieces, which were put into a circular shaped and flat platinum sample pan (inner diameter=1 cm and height=2 mm) placed in the TGA furnace oven.

### 3 Results and discussion

#### 3.1 Operational conditions set-up

##### 3.1.1 Heating rate and carrier gas

A number of procedures have been recently proposed for OC and EC quantification on filter samples of ambient particulate (Schmid et al., 2001; Chow et al., 2001, 2004, 1993; Birch and Cary, 1996; Cachier et al., 1989; Fermo et al., 2003). Nevertheless, a unique and validated method doesn't exist because of some problems encountered in assigning correctly the carbon content to the organic or to the elemental fraction.

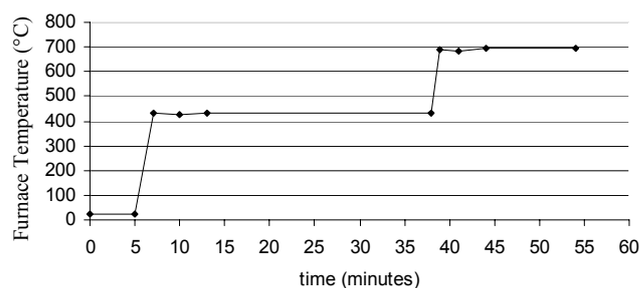


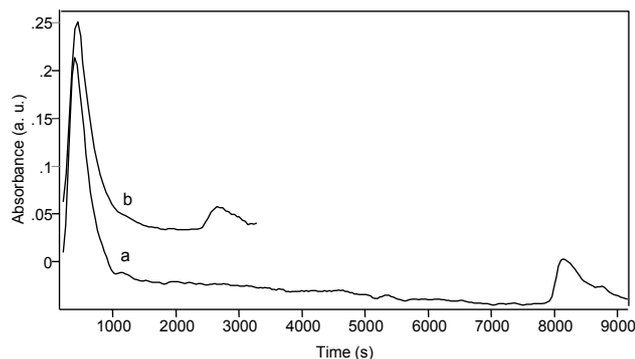
Fig. 3. Heating program employed for the analysis of particulate matter samples by TGA/FT-IR.

Our methodology for OC and EC analysis is a thermal one based on TGA/FT-IR measurements. It allows OC/EC separation thanks to the different thermal behaviour of the two carbon components. OC and EC are oxidized to carbon dioxide whose quantification is accomplished by a FT-IR spectrophotometer, which acquires infrared spectra simultaneously with the thermo-gravimetric analysis. While non-dispersive infrared detection (NDIR) of evolved CO<sub>2</sub> has already been used (Cadle et al., 1980; Tanner et al., 1982; Novakov, 1997), as far as we know, a FT-IR spectrophotometer as a detector is here applied for the first time (see the Experimental section for details on the apparatus).

Our approach shows some analogies with the earlier thermal methods presented in the literature taking into account the specific problems previously described.

Evolved gas analysis procedures proposed in the literature differ in the temperature programs and in the atmosphere conditions (oxidation or pyrolysis). As concerns the sample heating, up to now three different systems have been employed: (1) direct insertion of the filter into the heated zone (Cadle et al., 1980), (2) flash heating (Tanner et al., 1982; Cachier et al., 1989), i.e. a very rapid heating which allows to reach the final desired temperature within a few minutes, and (3) progressive heating (Chow et al., 1993; Birch and Cary, 1996; Novakov, 1997; Iwatsuki et al., 1998). As in our system the heating is realized by a thermo-gravimetric analyzer, we have tested both progressive and flash mode. In a previous work (Fermo et al., 2003) first attempts were made using a temperature rate of 5°C/min in the range going from ambient temperature up to 650°C. The method accuracy in total carbon determination was verified analyzing SRM 1649a. However, on examining particulate matter on filters, a good separation between OC and EC was not achievable. Therefore, in the present study the heating rate was increased at the maximum value allowed by the instrument (160°C/min), which means to realize a flash heating.

When starting the analysis, at first the system is left to equilibrate for some minutes at 30°C (Fig. 3) then a first jump at 425°C (at a rate of 160°C/min) takes place. At this point, after an isotherm step of 25 min, a second jump at 700°C is performed maintaining the same heating rate



**Fig. 4.** CECs obtained by TGA/FT-IR analyzing standard a and corresponding to experiment No. 1 – Table 1 (curve a) and experiment No. 2 – Table 1 (curve b).

(heating temperature values and isotherms duration are discussed in the next paragraph).

A very rapid heating allowed us to obtain two well distinguishable peaks in the CEC ( $\text{CO}_2$  Evolution Curve). Furthermore, in these conditions the determination of low quantity of organic and elemental carbon is possible and the organic carbon volatilization is favoured against carbonization. In fact it is well known that, while heating, organic carbon could char, i.e. a pyrolytic organic-to-elemental carbon conversion takes place. Previous works (Cadle et al., 1980; Cachier et al., 1989) have also estimated the percentage of OC to EC conversion, which is about 10–20% in the worst case.

In the literature, as carrier gas, He and a mixture of 10%  $\text{O}_2/\text{He}$  or 2%  $\text{O}_2/\text{He}$  are generally used for the OC and EC decomposition step, respectively (Chow et al., 1993; Birch and Cary, 1996; Cadle et al., 1980; Tanner et al., 1982). When an inert atmosphere is used during the first step just the organic fraction decomposition takes place and EC decomposition is prevented. However, some authors observed that in order to minimize *charring*, oxygen has to be chosen instead of an inert atmosphere (Cachier et al., 1989; Ohta and Okita, 1984). In particular Cachier et al. (1989) found that using both flash heating and oxygen as carrier gas, pyrolysis would affect 10% or less of the initial organic carbon. On the base of some experiments previously carried out by TGA/FT-IR (Fermo et al., 2003), it has been concluded that also in our case *charring* was more pronounced in non-oxidizing media such as helium. For this reason we selected pure oxygen as carrier gas.

### 3.1.2 Heating temperature and isotherm duration

Once flash-heating mode has been selected, some experiments were carried out to optimize the two heating step temperatures in order to find a split point between OC and EC, which is not arbitrary. Nevertheless, it is important to stress that up to now the OC and EC definition is an operative one, depending on the analytical method used for their quantifi-

cation and, as a consequence (see the introduction), this can produce different splitting between the two fractions.

Different temperatures have been proposed in the literature for the separation of organic carbon from elemental carbon such as 300°C–350°C (Cachier et al., 1989; Ohta and Okita, 1984; Ellis et al., 1884), 400°C–430°C (Tanner et al., 1982; Iwatsuki et al., 1998) and 650°C (Cadle et al., 1980).

Isotherm duration is also quite variable and goes from a few minutes (Tanner et al., 1982) to longer periods such as 30 min (Novakov, 1997), 60 min (Iwatsuki et al., 1998) or 120 min (Cachier et al., 1989).

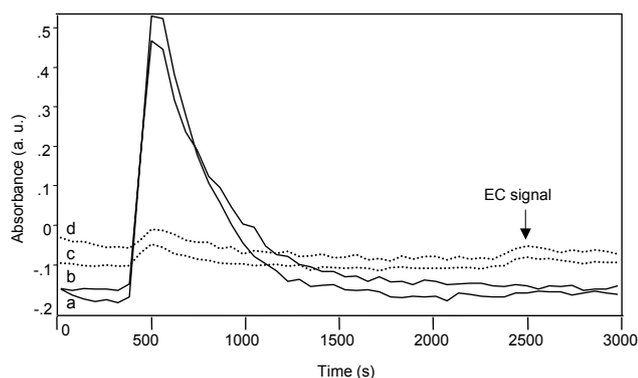
In setting the proper temperature it is mandatory taking into account some facts: temperature must be high enough to assure organic matter complete decomposition; *charring* must be minimized; elemental carbon pre-combustion must be avoided.

Some experiments were carried out on standard mixtures of known composition in order to establish the optimal temperature for the first step and the corresponding isotherm duration. We didn't check 650°C since the apparatus used in that case (Cadle et al., 1980) worked in an inert atmosphere, avoiding EC decomposition, while we had decided to work in oxygen (see Sect. 3.1.1).

The second temperature step has been in any case set at 700°C since, working in oxidizing conditions, it is enough to decompose elemental carbon (Cadle et al., 1980; Iwatsuki et al., 1998). This has been verified by our system analyzing both pure graphite and the standards d1-d3. No residual carbon was observed.

Results obtained in the optimization procedure are summarized in Table 1. Each experiment was replicated at least twice.

At first we reproduced the experimental conditions reported by Cachier et al. (1989), i.e. first step temperature of 350°C and a heating duration of 120 min in order to decompose all the OC fraction (experiment No. 1, Table 1). The CEC corresponding to this experiment is reported in Fig. 4, curve a. Only 72% of the total carbon, in the mixture as  $\text{Na}_2\text{C}_2\text{O}_4$ , is recovered in the first step (Table 1). From experimental results obtained in our laboratory by means of a standard thermo-gravimetric analysis, it has been noted that pure sodium oxalate shows two characteristic decomposition temperatures at about 500°C and 780°C respectively: at 500°C it decomposes and turns to carbonate which then decomposes at 780°C. This high temperature has been avoided in our analyses because of the possibility of carbonate decomposition during real PM samples measurements. The second peak present in the CEC may be due to some *charring* or to initial carbonate decomposition. Oxalate is the most abundant of the dicarboxylic acids in the particulate matter and can potentially act as cloud condensation nuclei. Its atmospheric concentration is about  $0.35 \mu\text{g}/\text{m}^3$  (Yao et al., 2002) and together with malonate and succinate represent on average 5.5% of sulphate concentration, one of the major PM component. It has also been demonstrated that mono and



**Fig. 5.** CECs obtained by TGA/FT-IR analyzing a 24-hour sample of particulate matter (on a filter). The filter was divided into two portions both heated at 425°C for 25 min (curves a and b); curve c was obtained with the heating program, reported in Fig. 3, on one half of the filter submitted to 24-h Soxhlet extraction; curve d was obtained by the same heating program on the second half of the filter not extracted.

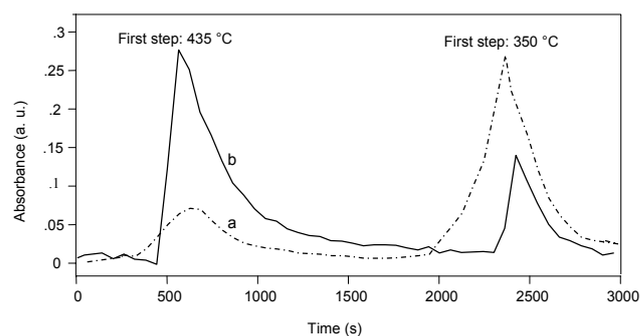
di-carboxylic acids represent one of the main classes in the so-called WSOC (water soluble organic carbon) fraction that accounts for about 50% of OC (Decesari et al., 2001). As a consequence, oxalate contribution to particulate matter mass is not negligible.

It is also important observing that the OC fraction could include high molecular weight organic substances having a thermal behaviour quite similar to the soot one. For example a mixture of amid, starch and sucrose has been chosen as a model for this fraction and its thermal behaviour studied. It has been found that in order to have a complete combustion of this class of substances, it is necessary to reach a threshold temperature of 485°C (Iwatsuki et al., 1998).

In our experiment, the first temperature step was then increased to 425°C in order to test an intermediate value between 350 and 485°C (experiment No. 2, Table 1). The duration step was drastically reduced to 10 min, which seems enough to regain the initial baseline signal and a recovery increase was observed (Fig. 4, curve b). Nevertheless, when thermal analysis is performed at temperatures lower than 780°C – as in our case – OC may be slightly underestimated if oxalate is present, since it does not completely decompose.

A temperature of 350°C is not enough to decompose standard b, a mixture containing organic acids (experiment No. 3, Table 1). Carbon recovery reaches 90% when heating at 425°C (experiment No. 4, Table 1). Taking into account the decomposition temperatures of the substances contained in standard b it is reasonable supposing that under these conditions a *charring* of 10% took place.

The temperature of 425°C was then checked for the analysis of samples of ambient particulate matter. For this purpose a 24-h filter was divided into two portions that were heated at 425°C with an isotherm duration of 25 min after which the



**Fig. 6.** CECs obtained by TGA/FT-IR analyzing SRM1649a using a first temperature step of 350°C (curve a) and 435°C (curve b) respectively.

baseline signal was regained (Fig. 5, curves a and b). After cooling, one half of the filter was undergone to a 24-h Soxhlet extraction in dichloromethane in order to dissolve the organic matter that was eventually still present. Soxhlet extraction is frequently used in the analysis of particulate matter organic compounds while dichloromethane has been chosen since it is one of the most widely used solvent (Pozzoli et al., 2004). The two halves were then submitted to the complete ramp program (heating at 425°C/isotherm/heating at 700°C/isotherm). A residual signal is detectable in both CECs in correspondence of the first step (Fig. 5, curves c and d) suggesting that also by means of the extraction it was not possible to eliminate all the organic substances (the second peak in both curves c and d is due to EC). Afterwards, the experiment was carried out working at a 435°C, which turned out to be high enough to decompose the whole OC fraction. As a consequence, we selected this value as the first step temperature and adopted the first isotherm duration of 25 min for ambient particulate samples analysis.

Some experiments were also performed in order to evaluate minimum EC combustion temperature. According to data reported by other authors (Iwatsuki et al., 1998) in oxidizing conditions the elemental carbon combustion starts beyond 485°C and it is complete under 650–660°C (Cadle et al., 1980; Iwatsuki et al., 1998). Loss of elemental carbon can be reduced by a short pre-combustion step (Cachier et al., 1989). Samples of pure graphite were analyzed by TGA/FT-IR using the following first step temperatures: 425°C, 450°C, 470°C and 500°C. Starting from 470°C, CEC shows a signal in correspondence of the first step indicating that in our operational conditions some EC pre-combustion can take place.

However, it is known that soot present in the atmospheric aerosol is not only composed by elemental carbon but contains a variety of highly condensed organic compounds (Cadle et al., 1980, Cachier et al., 1989). Standard reference material NIES No. 8, being a vehicle exhaust particulate, contains about 80% of carbon so that it was used to check pre-combustion conditions in real particulate samples. Four

**Table 2.** Standards used for the calibration curve shown in Fig. 7a; together with the standard type the mass of the standard, the carbon content present in each standard and the peak area observed in the CEC are reported.

Std type	Mass of Std (mg)	Carbon content (mg)	Peak area
No sample	0	0	11.4
d2	5.4	0.009	18.7
d2	5.8	0.010	24.0
d1	0.5	0.025	52.6
d2	44	0.074	114
d2	45	0.076	94.7
d3	23	0.077	109
d3	42	0.137	187
d3	43	0.142	167

standard samples, previously submitted to Soxhlet extraction in order to eliminate the soluble organic fraction, were analyzed by TGA/FT-IR testing the following first step temperatures: 425°C, 435°C, 450°C and 470°C. A signal attributed to pre-combustion already appeared at 450°C. We can therefore conclude that to avoid elemental carbon pre-combustion, in our operational condition the maximum temperature for the first step must be 435°C. Even if it cannot be excluded that some quite refractory organic substances could be not still decomposed at this temperature, 435°C is a good compromise between the need to decompose OC and to avoid EC pre-combustion.

As carbon loss is associated with a change in the filter color, in order to verify if some variation occurred, a particulate matter filter was extracted from the furnace at the end of first isotherm step and no change in the filter color was evident.

The final selected operational conditions for ambient samples analysis are summarized in Table 1.

We would stress that also with our method OC charring cannot be excluded when analyzing particulate matter samples. However, for a standard mixture of known composition (standard b, experiment No. 6, Table 1) where no elemental carbon was present, it was estimated to be 10% in accordance with literature data (Cachier et al., 1989). Pure substances with high molecular weights such as sucrose and starch have been analyzed and, from the ratio between areas of the first and the second peak, it was estimated a charring percentage of 10 and 14%, respectively.

SRM1649a has been also analyzed using the final selected operational conditions. Figure 6 shows the CECs obtained with a first temperature step of 350°C (curve a) and 435°C (curve b). As expected, 350°C wasn't high enough to allow complete OC evolution. On the contrary, using a first temperature step of 435°C a good agreement with literature values has been obtained (Currie et al., 2002). A total carbon con-

**Table 3.** Standards used for the calibration curve shown in Fig. 7b; together with the standard type the mass of  $\frac{1}{2}$  filter (area 6 cm<sup>2</sup>), the carbon content present in each standard and the peak area observed in the CEC are reported.

Std type	Mass of Std on $\frac{1}{2}$ filter (mg)	Carbon content (mg)	Peak area
$\frac{1}{2}$ blank filter	0	0	17.3
d1	0.172	0.008	23.5
d1	0.277	0.013	30.0
d1	0.523	0.025	37.5
d1	0.702	0.033	44.5
d1	1.299	0.061	65.6
d1	1.480	0.070	91.6
d1	1.791	0.084	96.5

tent of 17.0% has been obtained against a certified value of 17.68 ( $\pm 0.19$ )% and OC and EC concentrations were 81.4% and 18.6% of TC, both values within the range of variation determined by the other methods (Currie et al., 2002).

### 3.2 Calibration and method validation

Some reference samples containing pure graphite and SiO<sub>2</sub> in known quantities have been prepared in order to perform instrument calibration (standards d1-d3 whose composition is described in Sect. 2.1).

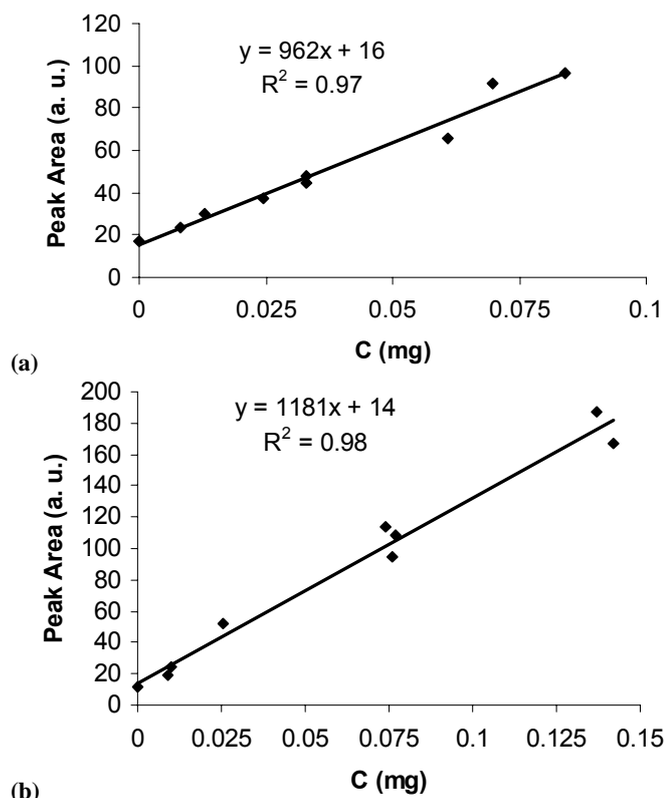
It is important to stress that carbon concentration chosen for standards d1–d3 are found in ambient samples such those collected in this study, i.e. samples with four-hour time resolution. These filters are characterized by low carbon concentrations and the total carbon content on  $\frac{1}{2}$  filter (that is the analyzed portion) is within the concentration range reported in Table 2 and 3.

The measurement conditions were the same employed for real samples analysis (experiment No. 5, Table 1) that allowed to obtain the best OC/EC separation. Two different calibration curves were constructed:

- the first curve (Fig. 7a) obtained analyzing different quantities of standard d1, d2 and d3 (Table 2) that can be used for OC/EC quantification when particulate matter powder samples have to be measured (e.g. SRM1649a);
- the second curve (Fig. 7b) obtained analyzing one half filter loaded with standard d1 in different quantities (Table 3) varying between about 0.17 mg and 1.8 mg; this curve is used for OC/EC quantification in real particulate matter samples.

The difference between the slopes of the two curves is within 20%.

As well known, TGA response depends on the physical status of the sample and this is the reason of our tests both



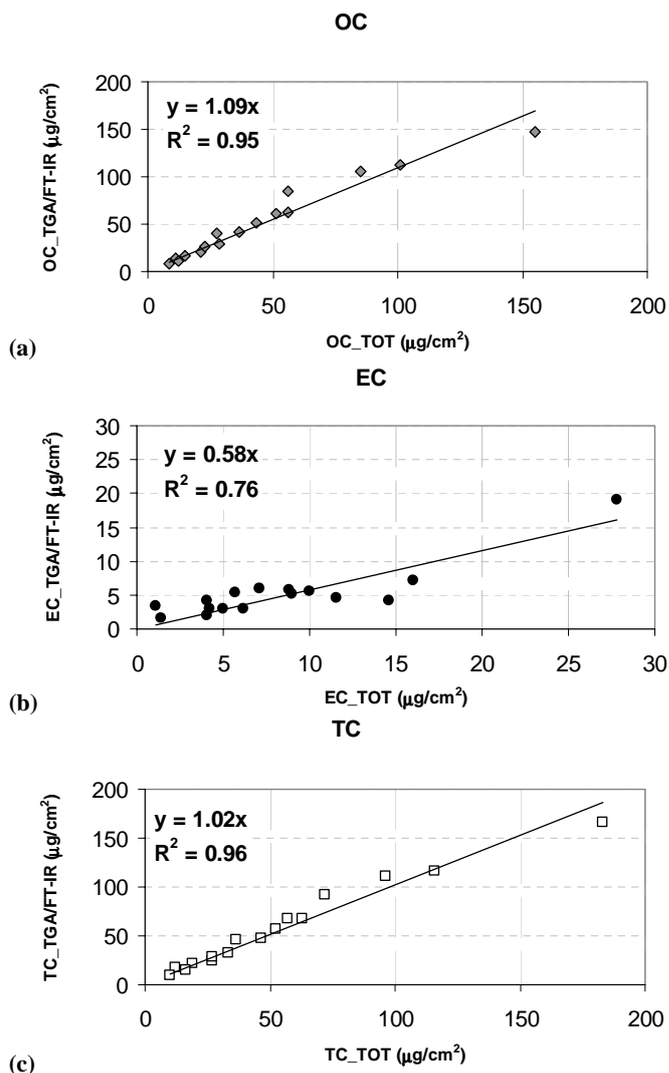
**Fig. 7.** Calibration curves obtained analyzing different quantities of standards d1, d2 and d3 (see Sect. 2.1 and Table 2) (Fig. 7a) and analyzing one half of the filter loaded with standard d1 in different quantities (see Table 3) (Fig. 7b).

on powder as is and on powder deposited on the filter. Working with two different matrices, slight different combustion efficiency can be hypothesized. However, the combustion efficiency at fixed analysis conditions shows a very good reproducibility.

In this work, all the carbon quantifications in PM samples have been made using the second curve (Fig. 7b).

It has been estimated that our technique has a detection limit of  $0.5 \mu\text{gC}/\text{cm}^2$  and the precision of the method is 10%. In comparison to NIOSH-TOT and LSCE our methodology has a lower precision while the detection limit is comparable to LSCE one and slightly higher than the NIOSH-TOT one ( $0.15 \mu\text{gC}/\text{cm}^2$ ).

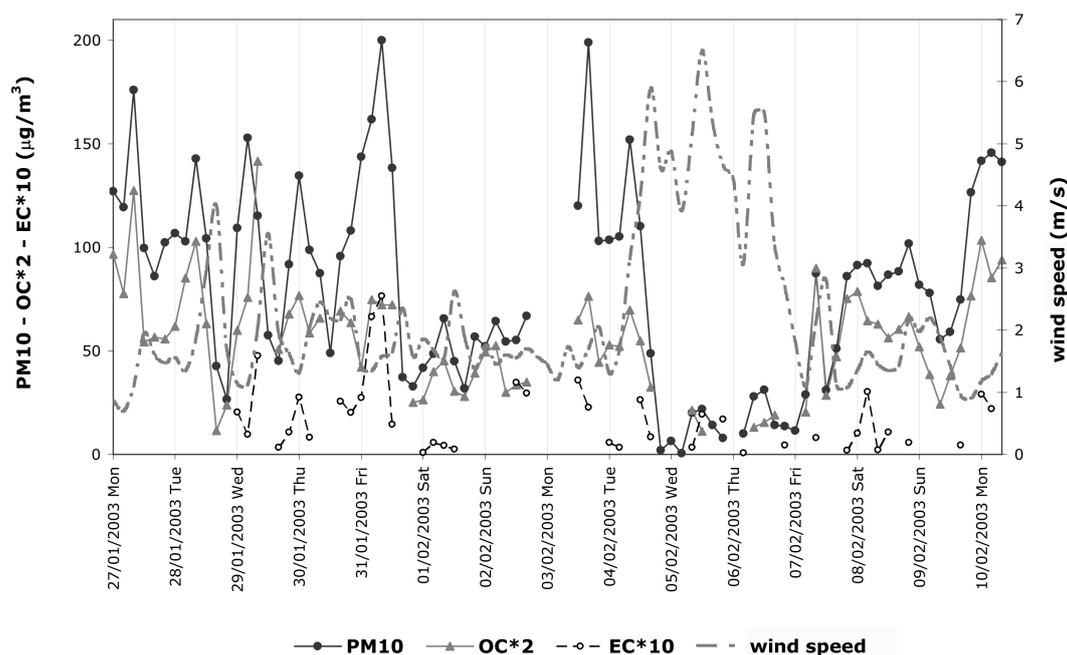
A few PM<sub>10</sub>, PM<sub>2.5</sub> and PM<sub>1</sub> samples (particulate matter with aerodynamic diameter smaller than 10, 2.5 and  $1 \mu\text{m}$  respectively) were also analyzed by a thermal-optical method TOT (Sunset Lab inc., USA) to perform a first comparison with TGA results on EC/OC/TC. The TOT instrument was designed to specifically address some of the problems observed in other methods in assigning carbon either to the organic or elemental fraction. By the continuous monitoring of the optical absorbance of the sample during the analysis,



**Fig. 8.** Comparison between TGA/FT-IR and TOT methodologies for the analysis of OC (Fig. 8a), EC (Fig. 8b), and TC (Fig. 8c) in particulate matter samples.

this method is able to both prevent any undesired oxidation of original elemental carbon and make corrections for the possible generation of carbon char produced by the pyrolytic conversion of organics into elemental carbon.

In Figs. 8a, b and c the results of the preliminary comparison between the TGA/FT-IR and TOT methodologies are shown. Generally, there is a good agreement on total carbon and OC while EC shows larger discrepancies. Differences can be due either to possible OC charring processes or to the EC pre-combustion with TGA/FT-IR method but also to the large uncertainty of TOT results in case of heavy loaded PM samples because of difficulties in the optical evaluation of the OC/EC splitting point. A more rigorous comparison on a larger number of samples is in progress at our laboratory.



**Fig. 9.** Temporal patterns of PM10, OC and EC concentrations together with wind speed (OC and EC values have been multiplied x2 and x10 respectively).

### 3.3 Analysis of carbon content in PM samples

The analytical methodology above described was applied to PM10 samples collected in the urban area of Milan to determine the EC/OC/TC content. Blank filters were also analyzed and no significant signals were observed.

The sampling site was a “background urban” location, according to the Italian Environmental Agency definitions, as it is not directly influenced by local emissions (i.e. traffic). The campaign was carried out in 2003, 27 January–10 February, and the PM10 was sampled daily with a four-hour time resolution. As a consequence during each day six four-hour time resolution filters were collected with a total number of 90 (=6×15) ambient samples during the whole period.

During the second week of the campaign the PM threshold values were exceeded so that the local authorities decided to introduce the condition of an odd/even plate’s traffic. However, this obligation was maintained only for a couple of days (starting from Monday, 3 February) because of the occurrence of a strong Föhn episode, which cleaned up the atmosphere on the whole Po valley (see Fig. 9). Thus, it was not possible to single out the effect of this action to prevent high pollution levels.

The samples were collected on quartz fiber filters (Whatman QMA) using low volume CEN-equivalent samplers (flow rate: 2.3 m<sup>3</sup>/h) equipped with PM10 (particulate matter with aerodynamic diameter smaller than 10 µm) inlets. The PM10 mass was gravimetrically determined using the same methodology described in Sect. 2.1.

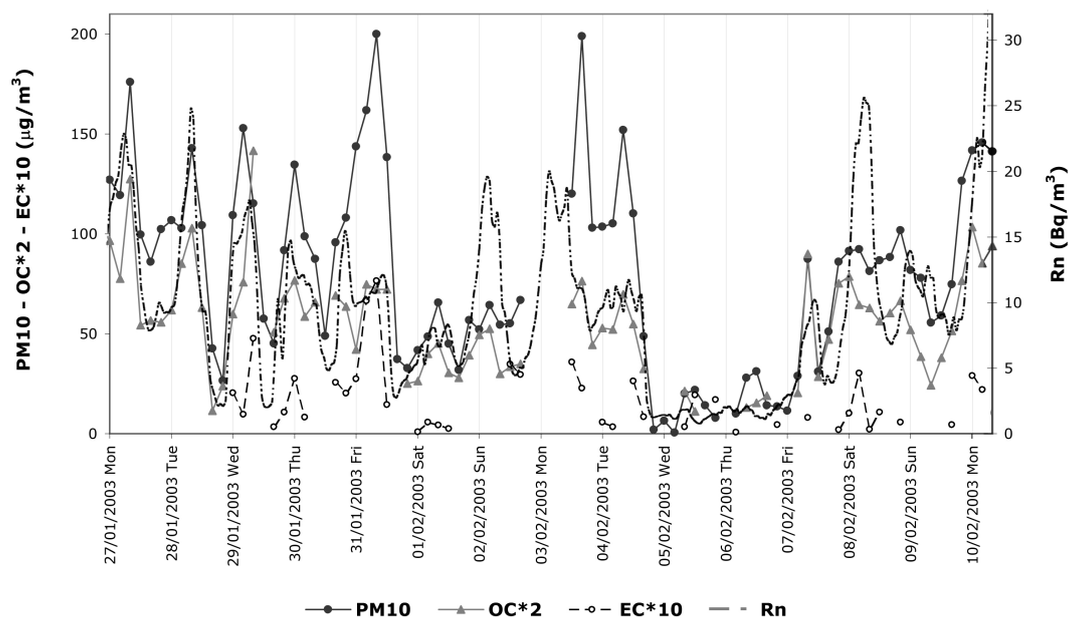
In Fig. 9 the temporal patterns of PM10, OC and EC concentrations are displayed together with the wind speed registered during the whole campaign. The good correlation between the PM10 mass concentration and its carbonaceous component is a first qualitative evidence of TGA/FT-IR reliability. As the examined filters are characterized by a sampling time of 4 h, the collected particulate matter amount is sometimes very low and EC is often below the detection limit (missing values in the figure).

During the campaign, PM mass concentration varied from 20.1 µg/m<sup>3</sup> to 200 µg/m<sup>3</sup> with an average value of 79 µg/m<sup>3</sup> and a standard deviation of 48 µg/m<sup>3</sup> (it is an expression of the observed variability in the data-set and not of the experimental error which is lower than 5%).

Ranges and average values were 12–70 µg/m<sup>3</sup> and 20 µg/m<sup>3</sup> for OC and 0.2–6 µg/m<sup>3</sup> and 2 µg/m<sup>3</sup> for EC. On average OC and EC made up 29.3 (±12.8)% and 2.5 (±1.8)% of PM10 mass, respectively.

Converting the organic carbon (OC) to organic matter (OM) concentrations using a mean molecular-to-carbon ratio of 1.6, as proposed by the recent literature (Salma et al., 2004; Turpin and Lim, 2001), an average contribution of about 53% to the PM10 mass has been found.

To study the observed temporal patterns the concentrations of the gaseous precursors of secondary aerosol as well as meteorological parameters (data available from the regional air quality network), have been also taken into account. Owing to the important role of the dispersion conditions of the atmosphere in determining PM10 concentration levels in the



**Fig. 10.** Temporal patterns of PM10, OC and EC concentrations together with Radon concentration (OC and EC values have been multiplied x2 and x10 respectively).

urban area of Milan, and in general in the Po valley, also Radon measurements were carried out during the campaign (Fig. 10). Radon can be considered a good natural tracer of vertical dispersion as it does not undergo chemical transformations in the air and it has a half-life of 3.8 days that is long enough to measure the atmospheric phenomena of interest here. It also has an approximately constant rate of exhalation from the ground over large, homogeneous areas and over time.

The Radon hourly concentration is routinely and continuously measured at ground level (6 m a.s.l.) in Milan by the group of the Institute of Physics. Radon activity concentration outdoor is detected with hourly resolution through the collection of its short-lived decay products attached to aerosol particles and the spectroscopic evaluation of their alpha activity (estimated detection limit =  $0.2 \text{ Bq m}^{-3}$ ). Details on the experimental methodology to measure  $^{222}\text{Rn}$  concentration are reported in Sesana et al. (2003).

The temporal patterns of PM10, OC and  $^{222}\text{Rn}$  show a fairly good correlation (Fig. 10); and when radon concentration reaches the maximum value (during the night and the early morning) the atmospheric dilution power is poorer and the pollutants accumulate (Vecchi et al., 2004).

#### 4 Conclusions

The method here presented appears suitable for the analysis of atmospheric aerosol carbonaceous fraction. It is based on evolved gas analysis and it consists of a simple home-made apparatus obtained by the coupling of a thermo-gravimetric

analyzer and a FT-IR spectrophotometer. In the framework of our study particular attention has been devoted to the method set-up. In particular, it is well known that the choice of both heating program and carrier gas are tricky points in the analysis of the two components OC and EC. The adopted analytical conditions led to the minimization of both charring of OC and pre-combustion of EC.

The method reliability for carbon quantification is supported by the fairly good agreement with TOT technique, which is one of the most widely used systems for TC/OC/EC analysis. Nevertheless, a clear separation between OC and EC is not easy to achieve either using more sophisticated systems such as TOT or TOR.

TGA/FT-IR was applied to PM10 samples collected in the urban area of Milan to determine the EC/OC/TC content. The technique reliability was also qualitatively demonstrated by the good correlation between the particulate matter mass concentration and its carbonaceous component. As for the method sensitivity, EC values were often below the detection limit but this is mainly due to the fact that the filters were sampled with four-hour time resolution.

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