Carbonaceous aerosols and their potential role in atmospheric chemistry

D. J. Lary
Department of Geophysics and Planetary Sciences, Tel Aviv University, Tel Aviv, Israel

D. E. Shallcross
Centre for Atmospheric Science, Cambridge University, Cambridge, England, United Kingdom

R. Toumi
Department of Physics, Imperial College, London, United Kingdom

Abstract. This paper considers the nature of carbonaceous surfaces, the means by which they are activated, the nature of some functional groups that they support, and some reaction mechanisms that may be involved. Because of the strong affinity of carbonaceous surfaces for organic species and because of the ease with which compounds in a high oxidation state can oxidize the carbonaceous surface, it is highly likely that carbonaceous aerosols are interacting chemically with a range of organic species in ways that have, as yet, not been fully characterized but may significantly affect the oxidizing capacity of our atmosphere. If HONO is formed on the surface of carbonaceous aerosols then this could be a significant source of HOx as HONO is readily photolyzed to give OH, and it could explain the large values of HONO often observed in the troposphere. In general, the reduction of NOx on carbonaceous aerosols is an important consideration, and it is addressed here.

1. Introduction

Carbonaceous aerosols (soot, charcoal, elemental carbon, etc.) are one of the most ubiquitous materials in our atmosphere. All combustion processes lead to the formation of carbonaceous material. As a result, between 10 and 50% of tropospheric particulates are carbonaceous, with particularly high levels being found in the urban atmosphere. Because of air traffic, carbonaceous aerosols (CA) are also found in the stratosphere. In his excellent introduction to CA, Goldberg [1985] has pointed out that the universality of CA is related to their refractory nature (very high melting point) and to their origin in burning processes. They can even be formed at temperatures of 200°C or less. They are generally very impure forms of carbon. Environmental charcoals have H/C ratios of 0.25-0.69 and O/C ratios of 0.08-0.33 [Cope, 1979]. In addition, they can contain up to a percent or so of nitrogen and sulfur, as well as a host of metals and elements such as Al, Fe, Ca, Mg, K, and Si [Gay et al., 1983, 1984; Goss and Eisenreich, 1997; Medalia and Rivin, 1982a,b; Medalia et al., 1982, 1983].

Active carbon is extensively used in various industrial processes, often as a reducing agent. It is therefore not surprising that recent laboratory studies have shown that HNO3, NO2 and O3 are also all reduced on CA [e.g., Smith and Chughtai, 1997; Ammann et al., 1995, 1997a,b; Rogaski et al., 1997; Chughtai et al., 1994; Tabor et al., 1993, 1994; Thibi and Petit, 1994]. Modeling studies have shown that this could have a significant impact on atmospheric chemistry [Lary et al., 1997; Hauglustaine et al., 1996; Bekki, 1997]. Since the atmosphere is so strongly oxidizing, any process that can lead to the reduction of atmospheric constituents, such as CA, may have a significant impact on atmospheric chemistry as they oxidize the CA surface. The adsorptive property of black carbons to remove impurities from gases and solutions has been known for centuries. Such processes are therefore worthy of further study. This is particularly true if the reducing agent is also one of the most abundant particulates of the urban environment and is even present in the stratosphere. Examining the nature of these atmospheric processes is therefore important.

Some useful information on carbonaceous surfaces can be gained from the literature on activated carbon. However, considerable caution has to be applied when distinguishing atmospheric CA and activated carbon.
They may have similarities but also important differences. One should not necessarily assume that CA are 100% carbon, or that they are crystalline with a well-known structure, since they are often incompletely burned biomass with a complex and varying composition of organic phases.

Although carbonaceous materials can vary considerably in their chemical reactivity, they do have several features in common. They are all carbonaceous, they can all be oxidized, they have a wide range of surface functional groups, and they typically have large surface areas.

Sections 2-10 consider the nature of carbonaceous surfaces, the means by which they are activated, some functional groups they support, and some reaction mechanisms that may be involved.

2. Nature of the Surface

A surface of carbon, or carbon plus hydrogen, is strongly hydrophobic. The presence of oxygen makes the surface more hydrophilic. CA may be acidic or basic depending on the way in which they are formed. Exposure of black carbon to temperatures between 200° and 400°C yields an acidic surface whereas treatment at higher temperatures in CO₂ followed by exposure to oxygen produces a basic form [Goldberg, 1985].

CA found in the environment are usually formed at high temperatures and come from a variety of sources, such as the combustion of plants, woods, fossil fuels, and industrially produced substances. The properties of the CA, such as the surface morphology and size distribution, reflect their origin and their history since they left the place of formation. Small submicron particles have usually been formed from combustion in the vapor phase (e.g., the combustion of a gas), and the larger particles which are bigger than 10μm reflect the structure of the burnt material, the so called char. Both may come from the same source, such as the combustion of plants and other biomass [Goldberg, 1985]. A detailed review of the chemical and physical properties of many carbonaceous materials can be found in the work of Mantell, [1968].

In CA the elemental carbon may be considered to be a disordered form of graphite (whereas the organic carbon content can not be thought of in this way). The degree of ordering within the carbonaceous material is reflected in a number of its properties. e.g., the density of a single crystal of graphite is 2.25 g cm⁻³, the density of coke is 2.05 g cm⁻³, and the density of carbon lamp black is 1.90 g cm⁻³ [Mantell, 1968]. Another example is the heat of oxidation of graphite to CO₂ which is 32621 J g⁻¹, whereas the heat of oxidation of amorphous to CO₂ is more 33104 J g⁻¹, and the heat of oxidation of wood charcoal to CO₂ is greater still at 33807 J g⁻¹ [Mantell, 1968].

X-ray analysis of carbonaceous material shows that it has a graphite like structure with randomly oriented microcrystallites, with each of the platelets about 1 nm thick [de Voogt, 1983]. The disorder arises in the amorphous carbons from the smaller amount of carbon atoms in the layer planes, which have a mean diameter of the order of 2.5 nm [Goldberg, 1985]. Typically, only a percentage of the carbon surface is active, so poisoning is not very difficult. Usually, the smallest graphite crystallites are oxidized first thereby exposing more of the larger crystallites, which are less easily oxidized [van der Plas, 1970].

It is found that activated carbon absorbs hydrocarbon vapors in preference to water vapor [Mantell, 1968]. Heat is released when commercially available activated carbon samples are wetted. The heat release increases when the surfaces are highly oxidized. e.g., van Driel [1983] showed that when two different activated carbon samples were wetted by water between 34 and 35 J g⁻¹ of carbon was released. However, after oxidation this increased to almost double, 66 J g⁻¹ of carbon. This implies that part of the surface was of a more polar nature after oxidation. This explains why CA act as atmospheric cloud condensation nuclei, and it implies that atmospheric CA is oxidized.

In contrast, when the surface was wetted with toluene, instead of water, the heat released was between 129 and 131 J g⁻¹ of carbon, with only a slight increase to between 135 and 138 J g⁻¹ of carbon for the oxidized surface. The small increase in energy release was attributed to a possible increase in surface area. The heat release on wetting can provide energy for surface reactions.

The affinity of carbonaceous surfaces for hydrocarbons may have significant implications for atmospheric chemistry. If the effect of the HNO₃ reduction on CA can be observed in atmospheric observations, as it seems from the work of Lary et al. [1997] and Hauglustaine et al. [1996], then it is also likely that atmospheric interactions of organic compounds with CA should be observed, because they should be taken up preferentially on CA. Goss and Eisenreich [1997] measured the sorption of polar and nonpolar volatile organic compounds (VOCs) to particles from a combustion source. A decrease in the sorption with increasing relative humidity was observed for all VOCs. The calculated sorption enthalpies suggested stronger sorption for polar compounds compared with nonpolar compounds of comparable volatility. Existing field data indicate that soot may significantly affect the environmental speciation of polycyclic aromatic hydrocarbons (PAHs) [Gustafsson et al., 1997]. The interactions between organic species and CA are likely to be quite complex. e.g., McDow et al. [1996] have shown that 4 out of 10 of the major organic compound classes found in organic aerosols include compounds that accelerated the photodegradation of benz[a]anthracene.

The most reactive areas of the surface are likely to be where carbon is not exerting its full valency. These will often occur where the soot particles have
exposed aliphatic and aromatic chains. These chains are stripped off as the CA is oxidized leading to the formation of CO, CO₂, and water. The time taken to strip off these chains is prolonged by an increased H content in the soot [Smith and Chughtai, 1996].

Carbonaceous surfaces absorb hydrocarbon vapors in preference to water vapor [Mantell, 1968]. It is therefore very likely that many atmospheric interactions are occurring between hydrocarbons and CA. We suggest that an important class of reactions may be with organic peroxides which can oxidize the carbonaceous aerosols and which are reduced in the process.

3. Carbonaceous Aerosols as Cloud Condensation Nuclei

There have been several studies of the state of water adsorbed on carbonaceous surfaces. e.g., active carbons and soot were studied by Berezn et al. [1997] who found that the state of adsorbed water is similar to the state of a stretched liquid. CA can act as cloud condensation nuclei (CCN). e.g., Konopka and Vogelsberger [1997] studied possible water condensation processes during the formation of aircraft contrails for different kinds of condensation centers. They found that soot particles with or without sulfuric acid can be more easily activated than small H₂O/H₂SO₄ droplets for atmospheric situations.

Condensation properties of ultrafine carbon particles in the Aitken range (particle diameters between 20 and 100 nm) were investigated by Kotzick et al. [1997]. They studied condensation phenomena with dry monodisperse soot aerosols. After reaction with ozone, activation of soot particles occurred at lower supersaturations. Fourier Transform Infrared (FTIR) studies revealed that oxygen-containing functional groups are generated on the particle surface during oxidation, facilitating water uptake in the condensation process.

Lammel and Novakov [1995] found that the nucleation ability of chemically modified carbonaceous particles increased with increasing soluble mass fraction and was comparable to that of (NH₄)₂SO₄ when the soluble mass fraction exceeded about 10%. The hygroscopicity of particles generated by combustion of diesel fuel in a diffusion flame increased when a sulfur-containing compound was added to the fuel. The CCN characteristics of diesel soot appear to be comparable to that of wood smoke aerosol. Karcher et al. [1996] analysed the primary contrail particles (aqueous solution droplets nucleated in situ, emitted insoluble combustion aerosols, and entrained background aerosols) and found that soot must be involved as ice-forming nuclei if the visibility criterion is to be fulfilled.

4. Surface Activation

Industrially, the basic method of activation is the introduction of further amounts of oxygen into the surface of carbon. This is done by submitting the carbon surface to oxidizing agents. These are either gaseous oxidizing agents, such as oxygen, ozone, air, water vapor, carbon dioxide, and nitrogen oxides, or solutions of oxidizing agents, such as nitric acid, a mixture of nitric and sulfuric acid, hydrogen peroxide, acidic potassium permanganate, chlorine water, sodium hypochlorite, and ammonium persulphate. Our atmosphere is a strongly oxidizing atmosphere that contains both categories of oxidizing agents in significant quantities. Therefore it should be perfectly able to activate, and keep activated, any CA that are present. Jankowska et al. [1991] reports that even a carbonaceous surface activated with ariel oxygen has many oxygen-containing surface functional groups.

Carbon oxidation is a complex heterogeneous process. Though the exact mechanism for the overall process is not fully understood, the reaction of carbon with oxygen can be written stoichiometrically as the following two exothermic reactions:

\[
\begin{align*}
C + O_2 & \rightarrow CO_2 \quad \Delta H_R = -387 \text{ kJ mol}^{-1} \quad (1) \\
2C + O_2 & \rightarrow 2CO \quad \Delta H_R = -226 \text{ kJ mol}^{-1} \quad (2)
\end{align*}
\]

It is thought that both CO and CO₂ are primary products [Smisek and Cerny, 1970] and that the CO/CO₂ ratio increases with temperature.

5. Surface Functional Groups

All carbonaceous materials, even pure materials like diamond and graphite, contain surface functional groups [van Driel, 1983]. A range of independent investigations has shown that a large variety of surface compounds are formed on the carbon surface. A characterization of the functional groups produced by the combustion of hexane has led to a more detailed understanding of the overall surface structure [Keifer et al. 1981, Akhter et al., 1984, 1985abc, 1986, Chughtai et al. 1994, Smith and Chughtai 1995, 1996, 1997].

Those containing oxygen compounds are particularly important because of their universal occurrence [Jankowska et al., 1991, p.82, and references therein]. The origin of these functional groups can be the starting material from which the CA were formed (this is particularly true for CA produced by raw materials rich in oxygen), or they can be introduced later. All commercial activated carbons also contain mineral matter. It is very likely that this applies to CA too, and these may have a significant effect on the rates of heterogeneous reactions on CA.

The number of oxygen containing surface groups will depend on the oxidizing agent and the type of surface. Nonetheless, two points should be noted. First, oxygen-containing surface functional groups typically represent 90% of the total amount of bound oxygen [van der Plas, 1970]. Second, several types of groups are found in-
It has been found that there are various types of carbons: those produced at low temperatures that typically have a negative surface charge and acidic behavior, and those produced at high temperatures that typically have a positive surface charge and basic behavior. The surface groups are found mostly at the edges of graphite type basal planes, which supports the idea of electronic interaction between surface groups. The acidity of one group influences neighboring groups and an overlap is possible between acidic strengths.

On air oxidation the carbonaceous surface becomes more acidic and therefore more polar. This is probably because of the formation of monocarboxylic or dicarboxylic acid groups [van Driel, 1983]. The reducing properties of activated carbon are slightly decreased by oxidation, perhaps owing to the formation of hydroquinone groups [van Driel, 1983]. Carbon groups at the rim of the carbon plane are destroyed, and C=O groups are formed in a yet unknown configuration. Figure 1 shows the principal types of acidic surface groups formed on carbonaceous surfaces; Figures 2 and 3 show models of carbonaceous surfaces. It can be seen that the surface groups present on the activated carbon surface are identical to those found on carbonaceous aerosol surfaces. The carboxyl surface groups make the CA acidic, and so they can take up greater quantities of ammonia, something that may be relevant in the troposphere. Noh and Schwartz [1990] and Bastick et al. [1977] showed that there was a relation between ammonia adsorption and the presence of surface carboxyl groups.

Smith and Chugtai [1995] used FTIR spectroscopy to study the structure and reactivity of black carbon (in the form of n-hexane soot). They found that the soot structure, as produced by high-temperature incomplete combustion is predominantly aromatic with a surface coverage by oxygen-containing functional groups of about 50%. Sergides et al. [1987] also found that 50% of the CA surface was covered by oxygen containing groups and that the ratio of aromatic to aliphatic carbon in n-hexane soot is at least 9:1.

6. Mechanisms for Surface Reactions

The active sites on carbonaceous surfaces are thought to consist of carbon atoms that are not exerting their full valency. For example, the carbon-water reaction was explained by Long and Sykes [1948] by carbon atoms attached to the rest of the lattice by only three bonds. Such sites are likely to be found at the free edges of either the lamellar or the cross-linked aromatic rings, which are the basis of the carbonaceous structure.

Some processes occurring on the CA will be catalytic, and some will not. When considering the likely impact of noncatalytic processes, knowing the maximum number of carbon atoms that can be involved is important. Figure 4 shows the equivalent gas phase mixing ratio of carbon atoms if all the CA mass is assumed to be carbon for 1, 10 and 100 ng m$^{-3}$ as a function of the altitude. At the earth's surface (1000 mb) 1 ng m$^{-3}$ of carbon is equivalent to about 2 pptv of carbon. During the Mauna Loa Observatory Photochemistry Experiment (MLOPEX) II campaign, values up to 200 ng m$^{-3}$ of carbon can be observed with values of around 10 ng m$^{-3}$ being quite typical, equivalent to between 20 and 200 pptv of carbon. As pressure drops exponentially with height this has increased to 10 pptv at 15 km (100 mb). Blake and Kato [1995] report a typical Northern Hemisphere lower stratosphere value of 1.8 ng m$^{-3}$ that corresponds to about 20 pptv of carbon.

Smith and Chugtai [1995] have shown that FTIR methods are particularly well suited for following net changes in surface groups, and gas phase reactant/product concentrations. FTIR has been the key technique in determining the kinetics and mechanisms of some important heterogeneous reactions of black carbon with gas phase oxidant molecules. e.g., the reaction of NO$2$/N$2$O$4$ with soot follows a dual path mechanism, down to 2 ppm, which is reflected in the rate law, initial rate $R_1$:
The evidence suggests that dissociation of ozone yields a steadystate concentration of the excited oxygen atom, which is actually the oxidant. FTIR combined with chemical measurements has proven that a high solubility observed for carbon particles exposed to ozone has its origin in the hydrolysis of the surface carboxyls. Significant effects of simulated solar radiation on the reactions, especially in the soot/SO2/H2O/O2 system, has been revealed by FTIR. Infrared will continue its central role in the examination of increasingly complex systems containing black carbon, particularly through its interface with ancillary techniques.

7. Carbon and Ozone

Smith and Chughtai [1996] report that the reaction of ozone with a carbonaceous surface has three distinct stages. First, a rapid reaction with the surface produces functional groups such as hydroxyls and carboxylates. O3 begins the degradation of the soot particles by stripping them of aliphatic and aromatic chains with the for-
Figure 4. The equivalent gas phase mixing ratio of carbon atoms if all the CA mass is assumed to be carbon for 1, 10, and 100 ng m\(^{-3}\) as a function of altitude and atmospheric pressure. This is useful when considering whether or not chemical processes on CA will be significant even if they are not catalytic. v.m.r. means volume mixing ratio. Read 1E-12 as 1\(\times\)10\(^{-12}\).

mation of CO\(_2\) and water. This stage is prolonged by an increased H content in the soot.

Second, a sharp increase in the CO\(_2\) released, which increases to a maximum and then starts to decrease. This unusual increase undoubtedly means several reactions are occurring. After stripping off the H atoms in the first stage the O\(_3\) oxidizes the CO to CO\(_2\) besides its continued reaction with the most vulnerable parts of the soot surface. This is consistent with a reduction in CO release during this stage.

Thirdly, the CO\(_2\) release begins to decrease, probably because the more reactive portions of the soot have now been oxidized and the more organized parts of the soot react at a slower rate.

Compared to soot that has not been exposed to O\(_3\), after 300 hours the ozonated soot experienced a decrease in particle size of 40% and a decrease in surface area of 37% [Sergides et al., 1987]. Smith and Chughtai [1996] have summarized the individual reactions identified as shown in Figure 5. The catalytic portion of the reaction can be written stoichiometrically as the highly exothermic reaction

\[
2\text{O}_3 \rightarrow 3\text{O}_2 \quad \Delta H_R = -284 \text{ kJ mol}^{-1} \quad (4)
\]

This catalytic reaction could have a profound impact on the stratosphere [Lary et al., 1997; Bekki, 1997]. Current assessment of the impact is hampered by our lack of knowledge about the true "average" nature of the soot. Bearing in mind that the soot lifetime in the lower stratosphere is of the order of months, it seems likely that very aged soot may have lost most of its reactivity as the surface degrades or becomes poisoned. We certainly expect the available surface area to decrease with time. Coal and amorphous carbon are rapidly attacked by O\(_3\), producing primarily water soluble acids similar to benzene carboxylic acids. It appears that CO\(_2\) and oxalic acids result from the ozonation of aromatic structures [Sergides et al., 1987, and references therein]. There is an approximately thirty fold increase in carboxylic acid surface groups when soot is exposed to O\(_3\), the increase in the particle's weight parallels this increase [Sergides et al., 1987]. The proportion of black carbon likely to be transformed into polycarboxylic acids decreases as the attack proceeds deeper into the particle.

Smith and Chughtai [1997] have examined the soot-ozone reaction at low concentrations to determine any influence of solar radiation on its products and kinetics. The effect of simulated solar radiation is to change the product distribution toward CO\(_{2(g)}\), CO\(_{(g)}\), and H\(_2\)O\(_{(g)}\) at the expense of soot surface functional groups formation. They concluded that the rapid diminution of ozone in soot's presence is unaffected by solar radiation.

8. Carbon and NO\(_2\)

Chughtai et al. [1994] found that -ONO\(_2\), -NOO\(_2\), -NNO\(_2\), and oxygen surface groups are formed and that
the presence of water plays a significant role leading to the formation of NO and N2O [see also Pires et al., 1996]. Tabor et al. [1993, 1994], Thibidi and Petit [1994], and Rogaski et al. [1997] have found that the reduction of NO2 to NO is very fast and therefore important. The aerosol studies by Kalberer et al. [1996] showed that it is slow when proceeding via a chemisorbed species which has been characterized by Ammann et al. [1995]. The chemisorption releases 100 kJ mol-1 and reversible adsorption 30 kJ mol-1 [Ammann et al., 1995, 1997a,b]. The NO was produced only slowly with a first-order rate constant of around 10^-4 s^-1; yet even this small value could be significant when high levels of CA are present. 

Rogaski et al. [1997] studied the reactivity and hydration properties of amorphous carbon in a low-pressure Knudsen cell reactor at room temperature (298 K). The reaction with NO2 had an observed γ = 0.11±0.04. Treating the amorphous carbon with NO2 and O3 does not alter the H2O uptake, while treatment with SO2, HNO3, and H2SO4 increases significantly the H2O uptake.

HONO was first tentatively identified by Ammann et al. [1997a]. Then Ammann et al. [1998] and Gerecke et al. [1998] both showed that HONO is a major product of the soot/NO2 interaction. The formation of HONO involves several intermediate steps that are not easy to separate.

If, as reported, HONO is formed, then this could be a significant source of HOx as HONO is readily photolyzed to give OH. Even if this process is not catalytic and only a fraction of the nitrogen incident on the CA leads to HONO formation, this could be a significant HOx source as at the Earth’s surface 1 ng m^-3 of carbon is equivalent to about 2 pptv of carbon. During the MLOPEX II campaign, values up to 200 ng m^-3 of carbon were observed, equivalent to about 400 pptv of carbon. Tropospheric values of around 10 ng m^-3 of black carbon are quite typical and are equivalent to about 20 pptv of carbon. Since OH and HOx are present in the atmosphere on the pptv scale these interactions could be a significant source of HOx.

J. Kleffmann [personal communication, 1998] have recently observed that the reaction of NO2 on dry soot produces large quantities of HONO in the gas phase (50% yields) at low pressure (2 torr) with some NO and adsorbed HNO3. At atmospheric pressure, NO yields are high (50%) with only a small amount of HONO in the gas phase and 3-5% adsorbed on the soot. In the presence of water (50% humidity), significant increases in HONO formation are observed (740 torr). They conclude that a consecutive reaction is occurring:

\[
\text{NO}_2 \rightarrow \text{HONO} \quad (5) \\
\text{NO}_2 \rightarrow \text{NO} \quad (6)
\]

where the major channel is

\[
\text{NO}_2 + \text{soot/H}_2\text{O}_{(ads)} \rightarrow \text{HONO} + \text{soot}_{(ox.)} 
\quad (7)
\]

and the minor channel is

\[
2\text{NO}_2 + \text{H}_2\text{O}_{(ads)} \rightarrow \text{HONO} + \text{HNO}_3
\quad (8)
\]
followed by

$$\text{HONO} + \text{soot}_{(\text{reactive})} \rightarrow \text{NO} + \text{soot}_{(\text{ox.})} \quad (9)$$

where soot$_{(\text{ox.})}$ denotes oxidized soot.

9. Carbon and Nitric Acid

Rogaski et al. [1997] have studied the reduction of HNO$_3$ on a carbonaceous surface. They report that the main products are NO, NO$_2$, and H$_2$O. Constructing some overall stoichiometric reactions that are exothermic is possible, e.g.,

$$\text{HNO}_3 + \text{O}_3 \xrightarrow{\text{CA}} \text{HONO} + 2\text{O}_2 \quad (10)$$

$$2\text{HNO}_3 + 2\text{O}_3 \xrightarrow{\text{CA}} \text{NO} + \text{NO}_2 + \text{H}_2\text{O} + 4\text{O}_2 \quad (11)$$

which are both exothermic at 298 K. No detailed reaction mechanism is given by Rogaski, but on basis of the findings of Chughtai et al. [1994], who studied the interaction of soot with NO$_2$ and N$_2$O$_4$, it is likely that -ONO, -NOO, and -NNO$_2$ surface groups will be formed.

Rogaski et al. [1997] studied the reactivity and hydration properties of amorphous carbon in a low-pressure Knudsen cell reactor at room temperature (298 K). They found that the reaction of HNO$_3$ had an observed $\gamma = 0.038 \pm 0.008$.

As mentioned for NO$_2$ reacting on the surface of CA, if HONO is formed, then this could be a significant source of HO$_2$ as HONO is readily photolyzed to give OH. Even if this process is not catalytic and only a fraction of the nitrogen incident on the CA leads to HONO formation, Figures 6 and 7 show that this could be a significant source of OH and HONO. This is the case even if the CA surface area used is as low as 0.01 $\mu$m$^2$ cm$^{-3}$ and the reaction probability is as low as $10^{-3}$. This may explain why large values of HONO [e.g., Harrison and Kito, 1994] are often observed in the troposphere in regions where the numerical models do not predict such large amounts of HONO. If hydrogen is abstracted from HNO$_3$ then NO$_3$, a nighttime oxidizing agent, could be released.

Kinoshita [1988] and Mahajan et al. [1978] report that treatment with HNO$_3$ is one of the most effective ways to make the carbon surface acidic. The treatment of a carbonaceous surface with HNO$_3$ does not affect their physical morphology but rather alters their surface chemical properties.

During the Transport and Atmospheric Chemistry Near the Equatorial Atlantic (TRACE A) campaign, Smyth et al. [1996] observed high NO concentrations in the upper troposphere, over a region characterized by intense biomass burning. These workers have concluded that one possible explanation for these results is that HNO$_3$ is being rapidly recycled back to NO$_2$ in the upper troposphere. In addition, Jacob et al. [1996] have also come to the same conclusion from the TRACE A campaign. In the 8-12 km region these work-

**Figure 6.** The calculated effect at midlatitudes at equinox on the concentration of OH if the postulated reaction HNO$_3$ + O$_3$ $\rightarrow$ HONO + 2O$_2$ is included on carbonaceous aerosols with a reaction probability of only $10^{-3}$.
Figure 7. The calculated effect at mid-latitudes at equinox on the concentration of HONO if the postulated reaction $\text{HNO}_3 + \text{O}_3 \rightarrow \text{HONO} + 2\text{O}_2$ is included on carbonaceous aerosols with a reaction probability of only $10^{-3}$.

ers note that the median conversion of NO$_x$ to HNO$_3$ from a model is 57 ppt day$^{-1}$ whereas the median HNO$_3$ levels observed are 59 ppt. Therefore the lifetime of HNO$_3$ needs to be of the order of a day, whereas its lifetime with respect to reaction with OH and photolysis is of the order of 12 days. During biomass burning events, large quantities of black carbon are produced, and it is extremely likely that during deep convective events, elevated levels of black carbon are brought up to the upper troposphere. The conversion of HNO$_3$ to HONO/NO$_2$/NO and the conversion of NO$_2$ to HONO/NO on black carbon could well be the missing mechanism noted by Jacob et al. [1996] and Smyth et al. [1996] to explain the elevated levels of NO observed in this region.

The calculations shown in Figures 6 and 7 were made using the chemical model AUTOCHEM for midlatitude equinox conditions. No family or photochemical equilibrium assumptions are made. All species are time integrated separately with a 15 min time step. The time integration scheme used is an adaptive timestep Burlisch-Stoer [Stoer and Burlisch, 1980] scheme specifically designed for integration of stiff systems after Press et al. [1992]. The time integration package is as accurate as the often used Gear [1971] package but faster. Photolysis rates are calculated by using full spherical geometry and multiple scattering as described by Lary and Pyle [1992ab] after Meier et al. [1982], Nicolet et al. [1982], and Anderson [1983]. The average photolysis rate over a model time step is calculated using 10-point Gaussian quadrature as described by Press et al. [1992]. AUTOCHEM has also been used to perform for the first time four-dimensional variational analysis of chemical species [Fisher and Lary, 1995]. The reactions on the carbonaceous surface are treated in the same way as reactions on the surface of polar stratospheric clouds (PSCs) are, a surface area is specified, as well as a reaction probability. Because this is a preliminary study, no account is taken of surface ageing or poisoning, although both of these are likely to be important. For this to be treated accurately more kinetic data is required, which, as yet is unavailable. The model contains a total of 56 species with a total of 320 reactions. Fifty-one species are integrated, namely, O($^1$D), O($^3$P), O$_3$, N, NO, NO$_2$, NO$_3$, N$_2$O$_5$, HONO, HNO$_3$, HO$_2$NO$_2$, Cl, Cl$_2$, ClO, ClOO, OCIO, ClO$_2$, ClO$_3$, ClONO$_2$, HCl, HOCI, CH$_3$Cl, Br, Br$_2$, BrO, BrONO$_2$, BrONO, HBr, HOBr, BrCl, H$_2$, H, OH, HO$_2$, H$_2$O, CH$_3$, CH$_3$O, CH$_3$O$_2$, CH$_3$OH, CH$_3$OOH, CH$_3$ONO$_2$, CH$_3$O$_2$NO, HCO, HCHO, CH$_4$, CH$_3$Br, CF$_2$Cl$_2$, CO, N$_2$O, CO$_2$, and H$_2$O.

10. Entrainment Within Liquid Drops

Since oxidation of the CA makes them hydrophilic, it is very likely that CA will become incorporated into liquid drops, whether these are rain drops or sulphate aerosols. This is what is observed [e.g., Karcher et al., 1996]. It is also found that soot embedded in a liquid
coating containing H$_2$SO$_4$ and HNO$_3$ triggers heterogeneous freezing of water ice and leads to visible contrails \cite{Karcher, 1996}.

For a reaction to occur on a CA embedded within a droplet, several mass transfer steps must take place \cite{Seinfeld, 1986, Hanson et al., 1994}:

1. Gaseous species must diffuse from the bulk gas to the surface of the droplet. This has a timescale given by

\[ \tau_{dg} = \frac{r_p^2}{4D_g} \]  \hspace{1cm} (12)

where the radius of the drop is \( r_p \) and the typical molecular diffusivity \( D_g \) is 0.1 cm$^2$ s$^{-1}$. So for a droplet with a radius of 0.1 \( \mu m \) the timescale is just 25 ns.

2. Transfer across the gas-liquid interface must take place. This depends on the Henry's law coefficient and the accommodation coefficient via the relationship

\[ \tau_{p} = D_a \left( \frac{4RTH}{ac} \right)^2 \]  \hspace{1cm} (13)

where \( D_a \) is the aqueous phase diffusion coefficient of the species initially in the gas phase, typically around 10$^{-5}$ cm$^2$ s$^{-1}$. \( c \) is the average speed of gas molecules given by kinetic theory \( c = \sqrt{8RT/M} \). \( H \) is the Henry's law coefficient; as \( H \) increases, so does \( \tau_p \). If we take the accommodation coefficient \( \alpha \) to have its maximum value of 1, then for HNO$_3$, \( \tau_p \) is of the order of a second or so, whereas for O$_3$ and NO$_2$, \( \tau_p \) is approximately 10$^{-15}$ s.

3. If ionization of the species occurs, it will be rapid for strong acids such as HNO$_3$.

4. The dissolved species in the aqueous phase diffuse. The characteristic time for aqueous-phase diffusion is

\[ \tau_{da} = \frac{r_p^2}{\pi^2D_a} \]  \hspace{1cm} (14)

so for a radius of 1 \( \mu m \), \( \tau_{da} \) is just 10$^{-2}$ s.

5. If a chemical reaction occurs as these processes happen sequentially, the overall rate is the rate of the slowest step. In the case of CA for molecules such as HNO$_3$ with large Henry's law coefficients, the slowest step is likely to be the transfer across the gas-liquid interface or the reaction at the CA surface embedded in the drop. Even so, this is still quite rapid, occurring on the timescale of a few seconds. For molecules such as NO$_2$ and O$_3$ with small Henry's law coefficients, the slowest step is likely to be the reaction at the CA surface embedded in the drop. For droplets with a radius of less than a \( \mu m \), diffusion of the dissolved species within the aqueous phase is rapid.

Therefore the fact that CA become entrained within droplets does not necessarily mean that they will no longer be effective in chemically processing the surrounding air. However, a relevant question which needs further kinetic investigation is how reactive will the carbonaceous "core" be once entrained within a liquid aerosol?

### 11. Summary and Suggestions

Carbonaceous aerosols are readily oxidized in the atmosphere by both gas and aqueous phase oxidizing agents. In the process, the oxidizing agents are themselves reduced, something that in some cases does not rapidly occur in the atmosphere. e.g., HNO$_3$ is reduced to NO$_2$ on carbonaceous aerosols, something that will normally occur only slowly by reaction with OH or by photolysis.

The oxidation of the carbonaceous aerosols leads to the formation of a range of polar surface groups, such as carboxylates, which cause the carbonaceous aerosols to become more hydrophilic and acidic. As a result, they can act as effective cloud condensation nuclei. Newly formed carbonaceous aerosols are much more reactive as they contain more aromatic and aliphatic chains and sites where carbon is not exerting its full valency. This oxidation is slower when more H is present. Once the sites where carbon is not exerting its full valency are oxidized (with the associated release of CO, CO$_2$ and H$_2$O) the less reactive, graphite like, sheets become more exposed. As well as the mass loss. Because of the formation of CO and CO$_2$, there is a mass gain by the formation of groups such as surface carboxylates.

If HONO is formed on the surface of CA, then this could be a significant source of HO$_2$ as HONO is readily photolyzed to give OH. Even if this process is not catalytic and only a fraction of the nitrogen incident on the CA leads to HONO formation, this could be a significant source of OH and HONO. This is the case even if the CA surface area used is as low as 0.01 \( \mu m^2 \) and the reaction probability is as low as 10$^{-3}$. This may explain why large values of HONO are often observed in the troposphere in regions where the numerical models do not predict such large amounts of HONO. If the hydrogen is abstracted then NO$_3$, a nighttime oxidizing agent, could be released.

Carbonaceous surfaces absorb hydrocarbon vapors in preference to water vapor. It is therefore very likely that many atmospheric interactions are occurring between hydrocarbons and carbonaceous aerosols. An important class of reactions may be with organic peroxides which can oxidize the carbonaceous aerosols and which are reduced in the process. Also, if part, or all, of the reduction sequences, such as carboxylic acid \( \rightarrow \) aldehyde \( \rightarrow \) primary alcohol \( \rightarrow \) alkane, ketone \( \rightarrow \) secondary alcohol \( \rightarrow \) alkane, tertiary alcohol \( \rightarrow \) alkane, occur on carbonaceous surfaces, it could significantly affect the oxidizing capacity of the atmosphere. It appears that the carbonaceous aerosol studies to date have omitted considering the possibility of such reactions taking place. These reduction sequences clearly include the increasingly high concentrations of the many halogenated carbonyl compounds produced by the degradation of
hydrochlorofluorocarbons (HCFCs), hydrofluorocarbons (HFCs), halons, and chlorofluorocarbons (CFCs).

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