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Nitrogen in aramid-based activated carbon fibers
by TPD, XPS and XANES

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Abstract

Activated carbon fibers were prepared from Nomex® [poly(m-phenylene isophthalamide)] by either H3PO4 activation, H3PO4–CO2 activation, or simply CO2 or steam activation. These treatments converted amide groups from the polymer precursor into complex and heterogeneously distributed nitrogen functionalities. TPD, XPS and XANES were used to study the effects of these treatments on the local bonding environment around nitrogen in the resulting carbons. These analytical techniques showed that nitrogen atoms are present in the 6-membered rings located at the edges of condensed polyaromatic systems as pyridine-like sp2 nitrogen (N1 or N2) or in the interior, where nitrogen replaces one carbon atom and is bonded to three carbon neighbors (N3). The occurrence of a species (N2) hypothetically related to a pyridinic cycle bearing oxygen substituents or intracyclic oxygen atoms could be correlated with the degree of oxidation of the carbon surface. Assuming that a relative N3 increase is indicative of aromatization and that the reverse, correlated with a N2 increase, is indicative of surface oxidative denitrogenation, the ratio between these nitrogen species revealed that aromatization and oxidative denitrogenation processes occur sequentially or simultaneously to different extents according to the type of carbon activation and to the burn-off degree. Physical activation involves thermal aromatization reactions during the carbonization stage and the subsequent isothermal activation one. In this second activation stage, co-occurring thermal oxidation reactions lead to a less intense denitrogenation during CO2 activation than during steam activation. H3PO4 activation induces the largest nitrogen retention in the final product in a double process of aromatization and “auto-activation” producing a moderate oxidative attack of nitrogen. However, an increase of the H3PO4 ratio fostered the oxidation of the carbon surface and consequently enhanced nitrogen gasification during the subsequent activation.

Keywords: Activated carbon; Activation; Temperature programmed desorption; X-ray photoelectron spectroscopy; Functional groups

1. Introduction

In recent years, aramid fibers, more particularly Nomex® [poly(m-phenylene isophthalamide)] (henceforth designated as Nomex), have been proposed as precursors for activated carbon materials with very homogeneous micropore size, usable as adsorbents, molecular sieves, cata-

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desorption with detection of the nitrogen-containing gases by mass spectrometry (TPD-MS), X-ray photoelectron spectroscopy (XPS) and X-ray absorption near-edge structure (XANES). To date, TPD has often been used to study nitrogen in inorganic solids [3–6], but not so frequently in organic and carbon materials. Due to the expected interactions during thermal decomposition, nitrogen TPD can, at first sight, provide us with indirect information on the distribution and reactivity of nitrogen groups present in the sample [7,8]. XPS is element-specific, but is particularly sensitive to surface effects and suffers from a lack of resolution: peak separations from different chemical forms of nitrogen are comparable to or less than peak widths. Nitrogen K-edge XANES brings additional details allowing a more precise interpretation of N functionalities present in the sample [9–12]. The positions of the lowest energy $\nu$ resonances can generally be used as a characteristic energy for the identification and quantification of aromatic nitrogen functionalities present in carbon materials. However, problems exist in resolving the resonances and in estimating the baseline, which make it difficult to identify and quantify the various species. XANES is a technique sensitive to all the forms of nitrogen that are present in the sample, which can provide us only with an average structure as a result. In this context, one of the major aims of this work is to find correlations between spectral features in these techniques for attempting a structural characterization of nitrogen-containing species present in the studied carbons. These results should help to understand details of the changes in nitrogen content and nitrogen functionality distribution as a function of the activation mode and of the burn-off degree.

2. Experimental

2.1. Materials

The precursor material was crystalline Nomex provided by DuPont Asturias. Various series of carbon adsorbents were prepared from either pure Nomex (N series), or Nomex pre-impregnated with different amounts of $\text{H}_3\text{PO}_4$ (NP series). The impregnation process was carried out in a rotary evaporator, using each time 10 g Nomex and 200 ml of solutions of phosphoric acid with different concentrations in order to vary the impregnation ratio. The temperature was increased progressively while stirring until complete evaporation of water; this process lasted for 6 h. Then, the samples were dried overnight at 383 K in a vacuum furnace. The impregnation ratio ($X_p$), calculated as the weight gain after impregnation relative to the initial mass of Nomex, [i.e. $X_p = (\text{H}_3\text{PO}_4 \text{ weight/ Nomex weight}) \times 100$] amounted to 1, 7 and 75 wt.%, depending on the phosphoric acid concentration. The resulting samples were named NP 1, NP7 and NP75, respectively (cf. Table 1).

Pyrolysis/physical activation treatments with $\text{CO}_2$ of Nomex, fresh or impregnated with $X_p = 1$ or 7 wt.% $\text{H}_3\text{PO}_4$, was carried out in a vertical reactor made of quartz, using 4–5 g of sample, a flow of Ar (99.999% pure) of 50 cm$^3$ min$^{-1}$ and a heating rate of 10 K min$^{-1}$ until 1123 K; once this temperature was attained, the samples were either rapidly cooled down to room temperature (N sample, only pyrolyzed), or to 1073 K. In the latter case, the gas flow was then changed to 50 cm$^3$ min$^{-1}$ $\text{CO}_2$ (99.98% pure), and it was kept for different time intervals to attain different burn-offs. Pyrolysis/physical activation treatments with steam were conducted in the same reactor. In this case, batches of about 10 g of Nomex were pyrolyzed in Ar (99.999 vol.% pure, flow rate of 50 cm$^3$ min$^{-1}$) up to 1073 K (heating rate, 10 K min$^{-1}$) and then cooled down to 1053 K and activated with a steam/argon mixture (720/50 cm$^3$ min$^{-1}$) for various periods of time to attain different burn-offs.

Chemical activations were carried out in a U-shaped reactor made from quartz, using 3 g of impregnated Nomex (75 wt.% $\text{H}_3\text{PO}_4$), an Ar flow of 50 cm$^3$ min$^{-1}$ and a heating rate of 10 K min$^{-1}$ until the final temperature (873 or 1073 K). Once this temperature was attained, the sample was rapidly cooled down to room temperature. Then, the resulting materials were washed with water in a Soxhlet extractor to eliminate the excess of phosphoric acid and/or its decomposition products until the conductivity in the washing liquids was $<3$ gS cm$^{-1}$. The yield and the burn-off of samples prepared from pre-impregnated Nomex were calculated from the final mass (after washing and drying) relative to the initial mass of Nomex before impregnation.

The studied NACFs and their preparation conditions are listed in Table 1, whereas Table 2 gives their elemental analysis on a dry basis. The elemental analyses were carried out in a microanalysis apparatus, LECO CHNS-932, provided with an oxygen analyzer, LECO VTF-900. The mass loss of selected NACFs between room temperature and 1273 K was studied in a Stanton-Redcroft STA-1500.
thermobalance, using 15 mg sample, a heating rate of 10 or 30 K min\(^{-1}\) and an Ar flow of 50 cm\(^3\) min\(^{-1}\).

The porous texture of NACFs was analyzed from physical adsorption isotherms of N\(_2\) at 77 K, measured in an ASAP 2010 volumetric apparatus (Micromeritics) and CO\(_2\) at 273 K, measured in a NOVA 1200 volumetric apparatus (Quantachrome). In either case, the samples were previously degassed overnight at 523 K. Porous texture parameters obtained from the N\(_2\) and CO\(_2\) isotherms are presented in Table 3. The N\(_2\) isotherms were analyzed by means of the BET method (using data in the relative pressure range of 0.01–0.1) to calculate the specific surface area (S\(_{\text{BET}}\)), and the Dubinin–Radushkevich (DR) equation to calculate the micropore volume \(V_{\text{l,p}(N_2)}\); the total pore volume (V\(_{\text{pt}}\)) was calculated from nitrogen uptake at 0.975 relative pressure, assuming N\(_2\) to be in the liquid state (Gurvitsch rule). The CO\(_2\) isotherms were analyzed by means of the DR equation to calculate the corresponding micropore volume \(V_{\text{l,p}(CO_2)}\). Table 3 also includes data for the burn-off (mass loss during activation, relative to mass of char) and yield (mass loss during activation, relative to mass of Nomex precursor).

2.2. TPD

For TPD, the sample (ca. 5 mg) was placed in a He + 1% Ne flow of 10 cm\(^3\) min\(^{-1}\) under atmospheric pressure and heated at 30 K min\(^{-1}\). Evolved gases were continuously detected with a quadrupole mass spectrometer. N\(_2\) was detected after selective oxidation of CO as CO\(_2\) on MnO\(_2\) at 393 K. External calibration was done using mixtures of the gas to calibrate in pure helium with 1% neon. The signal amplitude of a selected mass was related to the m/z 20 amplitude of neon. In this work, physisorbed water occurs as a large peak, probably as a result of steam interactions with the inner wall of the transfer line between the oven and the detector. Calibration of the integrated signal of physisorbed water was used to assess the sample moisture and hence to express the TPD gas yield with respect to the dry matter.

2.3. XPS

XPS signals were recorded using a VG Scientific ESCA-LAB 250 system (based at ITODYS, Universite Paris 7, Paris, France) equipped with a micro-focused, monochromated Al K\(_\alpha\) X-ray source (1486.6 eV, 650 \(\mu\)m spot size, 200 W) and a magnetic immersion lens which permits to enhance the sensitivity by ca. one order of magnitude. The spectra were acquired with pass energies of 150 and 20 eV for the survey and the high resolution spectra, respectively. No N1s binding energy correction was done, with respect to the C1s one, because the sample, mounted onto a conducting double-sided adhesive tape, did not exhibit any charging effects. The Advantage software, version 1.85, was used for digital acquisition and data processing. The compositions (in at.%) were determined by considering the integrated peak areas of C1s, N1s, O1s and P2p from the survey spectra and the respective sensitivity factors.

2.4. XANES

XANES measurements were performed at the Laboratoire pour l’Utilisation du Rayonnement Electromagnétique (LURE, Orsay, France) on the VUV Super-Aco storage ring. They were carried out at a resolution of -0.2 eV at the nitrogen K edge. The samples were ground to about 20 \(\mu\)m particle size, dispersed in acetone, and applied drop-wise to high-purity tungsten plates, and the solvent was allowed to evaporate. All measurements were made at room temperature under ultra high vacuum.
Calibration of the photon energy scale at the N1s edge was made to match the N1s — n· of C=N in polyacrylonitrile (PAN, Aldrich) at 401.4 eV. The N K-edge XANES spectra of 1s → n transitions were deconvoluted with Gaussian–Lorentzian curves after subtraction of an error function step to the experimental XANES spectra.

3. Results and discussion

3.1. TPD

Temperature-programmed desorption profiles recorded at 30 K min⁻¹ from 300 to 1750 K are presented in Fig. 1. As shown in Fig. 2 and Table 4, total TPD gas yields are of the same order of magnitude as the total mass loss recorded by thermogravimetry at 30 K min⁻¹. TG curve tailing at temperatures above the peak maximum at 1075–1125 K (Fig. 2) could be attributed, in the case of samples NP75-873 and NP75-1073 (which underwent the largest mass losses), to vaporization of P2O5 (coming from decomposition of unreacted phosphoric acid and/or products from its reaction with Nomex); this process takes place above ca. 1000 K at lower heating rate [13]. For the rest of samples, mass losses above 1173 K (maximum temperature to which some of the samples were exposed during their preparation) are rather attributable to loss of very stable functional groups and/or other constituents. At the temperature of maximum mass loss rate, around 1100 K, the main gases released are carbon monoxide and water. Physisorbed water and carbon dioxide coming from postcooling exposure to the atmosphere evolve earlier, and molecular hydrogen together with several nitrogen containing gases occurs later. The thermal desorbed water peak was computed by subtraction of the whole water signal with the physisorbed water tailing curve. This background curve was obtained by extra- and interpolation of the physisorbed water tailing curve appearing before and after the thermal desorbed water peak that results from removal of chemisorbed oxygen. The background corrected water peak (desorbed water) occurs between ca. 750 and

![Fig. 1. TPD profiles of the Nomex char (N) and Nomex-based activated carbon fibers.](imageurl)
Fig. 2. Comparison between total evolution rates measured by thermogravimetry and by TPD.

1200 K with a symmetrical shape along an apparent pseudo-second-order kinetics. After correction, only NH42 and NP75-873 displayed some desorbed water, which occurs much above the main CO2 peak, so that it cannot result from H-bonded H2O associated with acidic oxygen complexes, but from the thermal condensation of hydroxyl groups initially present, or secondarily formed, in the solid sample [14–17]. Molecular hydrogen evolved from all the studied samples in two steps above 900 K, as a result of condensation between polyaromatic systems forming hydrogen radicals. The materials obtained by steam activation and by H3PO4 activation are the most hydrogenated ones (Table 2), and accordingly they liberate more hydrogen (Table 3) and this release occurs at lower temperatures, than for the rest of samples. In contrast, samples obtained by CO2 activation release much less hydrogen and this occurs at a higher temperature (>1450 K).

The quantification of the thermal desorption of CO2 or CO in the low temperature range is hindered by the simultaneous desorption of simply physisorbed species and the removal of gaseous compounds formed by the breakdown of fragile surface functional groups [18]. In the present work, the first peaks and shoulders at the beginning of programmed heating that are due to physisorbed CO2 or CO, were deleted from the whole profile (which was fitted by summation of a series of equations of first-order kinetics). The assignment of CO- and CO2-yielding groups was made on the basis on the current literature [19–25]. The proportion of CO2- and CO-generating complexes varies with the sample treatment: the samples activated with H3PO4 are the most acidic ones (highest CO2 evolution) and display the highest fraction of high-temperature CO2-generating complexes (lactone/anhydride type and/or adjacent acidic oxygen complexes formed at dangling carbon sites during post-activation ageing), whereas the samples activated physically display the highest fraction of high temperature CO-evolving complexes (chromene type). The lack of correlation between CO and CO2 yields and the porous texture (Table 3) suggests that CO- and CO2-evolving complexes would not have been formed over the entire surface area of the chars during the activation process [25]. Some of these complexes would have been formed after cooling by low temperature oxidation.

As shown in Fig. 1, the activation methods of the Nomex fibers produce several types of N-containing gas

Table 4
TPD and TG yields

<table>
<thead>
<tr>
<th>Sample</th>
<th>TPD yields, mmol g g-1</th>
<th>Total loss, wt.%</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>H2O</td>
<td>CO</td>
</tr>
<tr>
<td>Initial char</td>
<td></td>
<td></td>
</tr>
<tr>
<td>N</td>
<td>0.00</td>
<td>0.50</td>
</tr>
<tr>
<td>CO activation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>NC4</td>
<td>0.00</td>
<td>0.50</td>
</tr>
<tr>
<td>NC41</td>
<td>0.00</td>
<td>0.60</td>
</tr>
<tr>
<td>Steam activation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>NH10</td>
<td>0.00</td>
<td>0.50</td>
</tr>
<tr>
<td>NH42</td>
<td>0.10</td>
<td>0.50</td>
</tr>
<tr>
<td>H3PO4 (1%)/CO activation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>NP1</td>
<td>0.00</td>
<td>0.80</td>
</tr>
<tr>
<td>NP1C16</td>
<td>0.00</td>
<td>0.50</td>
</tr>
<tr>
<td>NP1C26</td>
<td>0.00</td>
<td>0.50</td>
</tr>
<tr>
<td>H3PO4 (7%)/CO activation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>NP7</td>
<td>0.00</td>
<td>0.60</td>
</tr>
<tr>
<td>NP7C14</td>
<td>0.00</td>
<td>0.70</td>
</tr>
<tr>
<td>NP7C32</td>
<td>0.00</td>
<td>0.60</td>
</tr>
<tr>
<td>H3PO4 (75%) activation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>NP75-873</td>
<td>1.20</td>
<td>0.90</td>
</tr>
<tr>
<td>NP75-1073</td>
<td>0.00</td>
<td>1.40</td>
</tr>
</tbody>
</table>

From 300 to 1723 K at 30 K minn.

From 300 to 1373 K at 10 or 30 K minn.
species distribution, which can be correlated with the H- and O-containing gas species distributions. The largest part of nitrogen evolves as molecular nitrogen in two steps: the first one below 1150 K (only in samples NP7, NP75-873 and NP75-1073), and the second one – the most important above this temperature – centered between the two \( \text{H}_2 \) peaks. HCN evolves in one peak in the same temperature range as \( \text{H}_2 \). In the most oxidized samples (those activated by \( \text{H}_3 \text{PO}_4 \)), HCN co-evolves with NO and its peak is almost as high as the \( \text{N}_2 \) one. The sample NP75-873, prepared at relatively low temperature, releases a small amount of \( \text{NH}_3 \), which co-evolves with the desorbed water just before the first \( \text{H}_2 \) peak. Positive correlations (not shown) of HCN/\( \text{N}_2 \) ratio with the carbonization/activation yield (Table 3) or with the atomic O/C ratio (which can be computed from Table 2) suggest that oxygen groups at the edge of condensed polyaromatic clusters would make the N polyaromatic skeleton more fragile and hence prone to earlier generation of N-containing radical species, which in the absence of a sufficient amount of hydrogen radicals are involved in secondary gas phase reactions leading first to hydrogen derivatives (HCN, \( \text{NH}_3 \)) and then giving \( \text{N}_2 \) in the absence of a sufficient amount of hydrogen radicals involved in the secondary gas phase reactions. A proof of that would be that hydrogen addition preserves the primary nitrogen fragments resulting from solid phase reactions (decomposition of functional groups in the carbon structure and interfacial reactions): the largest part of nitrogen evolves in the low temperature range, and consequently \( \text{NH}_3 \) and HCN evolution increases.

3.2. XPS and XANES

XPS survey spectra, together with N1s core level photoelectron spectra, the N K-edge XANES spectra and nitrogen TPD-MS profiles for selected NACFs, are compared in Fig. 3. In agreement with previous works [9,26], the XPS N1s spectra show, depending on the sample, three to five components: two dominant peaks plus some smaller contributions. In the \( \pi^* \) region of the XANES spectra, three peaks can be distinguished. To obtain acceptable curve fittings, the XPS spectra were decomposed into five peaks (labeled P1–P5) and the XANES spectra into three (labeled A1–A3) (Table 5). Note that the \( \pi^* \) resonances should fall at nearly the same energy than the photoemission peaks [10].

The components P1, P3, A1 and A3 appear as discernible peaks in the N1s envelop and can be structurally assigned as follows. P1 and A1 can be related to pyridinic nitrogen in non-functionalized heterocycles (i.e., where N atoms contribute to the \( n \) system with one \( n \)-electron bond to two sp\(^2\) carbons not linked to another heteroatom), and to a lesser extent to arylnitriles, which may be over-evaluated because the nitrile bond predominantly has a \( \pi \) system...

Fig. 3. N1s XPS, \( \pi \) region of the N K-edge XANES spectra.
Table 5
N1s XPS (P) and XANES (A) data

<table>
<thead>
<tr>
<th>Sample</th>
<th>XPS atomic ratios x 100</th>
<th>Relative area of N1s components, % of total area</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>O/C N/C P/C P1 P2 P3 P4 A1 A2 A3</td>
<td></td>
</tr>
<tr>
<td>N</td>
<td>3.19 5.09 – 36.1 11.5 40.5 11.8 42.0 21.0 36.0</td>
<td></td>
</tr>
<tr>
<td>NC41</td>
<td>4.54 3.42 – 33.6 15.8 37.1 12.8 40.8 21.5 37.7</td>
<td></td>
</tr>
<tr>
<td>NH42</td>
<td>3.69 3.08 – 30.1 11.2 41.6 13.3 35.5 17.9 46.6</td>
<td></td>
</tr>
<tr>
<td>NP1</td>
<td>5.47 5.34 – 37.0 16.9 40.2 11.0 40.2 24.0 35.8</td>
<td></td>
</tr>
<tr>
<td>NP1C26</td>
<td>5.19 4.61 0.16 33.9 14.4 41.7 10.0 34.5 24.0 41.5</td>
<td></td>
</tr>
<tr>
<td>NP7</td>
<td>6.38 4.44 0.56 29.6 19.6 38.6 15.2 40.4 24.9 34.4</td>
<td></td>
</tr>
<tr>
<td>NP7C32</td>
<td>6.66 3.62 0.60 34.0 16.6 38.8 11.7 35.2 22.4 42.3</td>
<td></td>
</tr>
<tr>
<td>NP7-873</td>
<td>9.66 5.56 1.84 33.6 28.8 27.0 10.6 47.6 19.6 30.6</td>
<td></td>
</tr>
<tr>
<td>NP7-1073</td>
<td>12.36 4.77 2.75 21.1 26.2 35.8 15.4 37.9 23.4 38.7</td>
<td></td>
</tr>
</tbody>
</table>

with large oscillator strength [12]. According to previous findings [8], P3 and A3 can be assigned to substitutional nitrogen in condensed polyaromatic systems where nitrogen substitutes carbon [10–12, 27–29]. They can be related to aromatization and microstructure ordering [8, 30]. When carbon surface bears hydroxyl groups, which generate water by thermal decomposition, as seen by water TPD for NH42 and NP7-873 (Fig. 1), the XPS peak P3 could also be assigned to quaternary nitrogen resulting from weak pyridine protonation by adjacent or nearby located hydroxyl groups forming hydrogen bridge with the pyridinic nitrogen [31–34].

The hidden P2 and A2 components show no clear relationship with each other (Fig. 4a) since P2 is assigned to some nitrogen species which would not have any $\pi^*$ resonance in arylnitrile or pyridine structures as found for A2 by Zhu et al. [9]. The formation of pyridone by pyridine oxidation, and/or pyrrole by elimination of CO from the pyridine [35] is not supported by our data: the XANES spectra of NACFs show neither any $\pi^*$ resonance at 401.9 eV (as expected for pyridone), nor around 403.5 eV (as expected from pyrrolic nitrogen). Despite the fact that nitrile group occurrence in the studied NACFs can be excluded on the basis of the DRIFTS examination (results not shown), the structural assignment of A2 mode at around 401.3 eV to nitrile group instead to pyridine centered at around 401.9 eV, [9], by reference to the N1s $\pi^*$ transition of cyanoanthracene, could be valid because arylnitriles are known to be thermally stable [36–38]. Li et al. [39] found that, during pyrolysis of a German bituminous coal, the nitrile group, not present in the raw coal and the tars produced at 873 and 973 K, appeared in the size exclusion chromatography fractions of the tar produced at 1073 K. The temperature at which the – C= N group is first observed coincided with the temperature at which nitrogen-containing model compounds were reported to begin to decompose. The XPS data indicated conversion of pyridinic nitrogen to nitrite nitrogen at temperatures in the range from 873–1073 K. However, in the case of the Nomex activated carbon fibers, the lack of any other satisfactory analytical evidence prevents us from regarding the A2/P2 component as resulting from any thermal resistant nitrile groups. In contrast, as supported both by previous studies [8, 40–43] and our own TPD, XPS and XANES observations, hydrogenated and/or oxygenated nitrogen species could be much better potential candidates.

![Fig. 4. Correlations between N1s XPS, XANES XPS hidden peak relative area (ra), atomic O/C ratio and HCN TPD yield.](image-url)
for A2/P2 components such as amide, pyridine, nitrile or nitroxy groups (these groups can thermally generate N₂, NO, ammonia and/or HCN at relatively low temperature instead of only N₂ at high temperature). Therefore, on the basis of reactional behaviour and correlations, we will assume that A2/P2 in activated carbons would be related to pyridinic cycle bearing oxygen substituents or intracyclic oxygen atoms making the cycle more prone to be open by thermal cracking during TPD and to release both HCN and N₂ at lower temperature rather than pure N₂ formed as a result of surface or gas phase radical recombination due to hydrogen starvation at higher temperature. The negative correlation between P3 or A3 and the oxidation degree (indirectly measured by P2 in NACF in Fig. 4b, itself well correlated with HCN TPD yields in Fig. 4c, as well as with atomic O/C ratio in Fig. 4d), suggests that, in agreement with previous works on nitrogen transformation during gasification or during wet oxidation [44,45], condensed polyaromatic systems bearing substitutional nitrogen are prone to be degraded by oxidation. These oxidation reactions would lead to a gradual denitrogenation and a transformation from the condensed pyridinic rings (center-N) to peri-located (top or valley-N) pyridone and pyridine.

The detailed observation of the nitrogen distribution given by N1s XPS and N1s XANES for the different modes of activation gives a more contrasted picture of the transformation of nitrogen functionalities during Nomex fiber activation. For instance, the relative abundance of substitutional nitrogen in condensed polyaromatic systems noticeably increases during NP75 carbonization from 873 to 1073 K (Fig. 5a), but decreases with increasing impregnation ratio at the same final carbonization temperature. These two opposite trends could be due to a double process of aromatization and “auto-activation” producing an oxidative attack of nitrogen, corresponding to the important release of gases above 873 K (Fig. 1), with an increase of the P2 nitrogen type – correlated with a systematic increase of the u/n ratio after activation. These reactions would occur more sequentially during physical activation, which involves a carbonization stage followed by an activation one with a co-occurrence of isothermal heating (aromatization) and thermal-oxidation. At equivalent burn-off, denitrogenation rate is lower during CO₂ activation than during steam activation (Fig. 4b), as the latter produces both hydrogen and hydroxyl radical attack of the nitrogen-containing polyaromatic systems. In this case, the P3 increase is not simply due to pyridine condensation but to pyridinic nitrogen protonation by bridge formation with adjacent or nearby located hydroxyl oxygen groups as evidenced by water TPD. As seen by elemental analysis (Table 2) and TPD (Table 4 and Fig. 1), CO₂ activation following H₃PO₄ impregnation and carbonization lead to a stronger denitrogenation than the one observed during unimpregnated Nomex char (N) activation. However, in this case, as shown in Fig. 5b, the apparent surface oxidation of nitrogen, measured by the relative change of lower than the one reached with char N activation – in spite of an equivalent development of the porosity measured by the wide micropore volumes (difference between the micropore volumes obtained by N₂ and CO₂ adsorption). In other words, increased denitrogenation during CO₂ activation, after Nomex impregnation with phosphoric acid and subsequent carbonization, would be due to the abundant oxygen substituents on N cycles introduced during H₃PO₄ activation.

### 3.3. Surface vs. bulk

Correlation plots between XPS atomic N/C or O/C ratios and elemental analysis atomic N/C ratios for NACFs are shown in Fig. 6a and b. The two correlation lines fall below the surface-bulk parity (indicated by the dashed line). The difference in N and O contents between the surface and the bulk of the carbon material could be attributed to an inaccurate calibration using inappropriate XPS sensitivity factors for the particular carbon samples under study. Though nitrogen and oxygen are expressed with respect to carbon, the oxidation propensity of the carbon surface exposed to atmospheric air could also explain the compositional deviation due to modification of the secondary electron travel through the oxidized surface layers of the carbon sample. As suggested by previous work [46], surface nitrogen depletion as seen by XPS could...
Fig. 6. Correlations between surface and bulk analyses for NACFs and reference samples [8,27,41,42,51]: (a) surface vs. bulk N/C ratio; (b) surface N/C and O/C ratios vs. elemental ratios; (c) TPD nitrogen yield vs. elemental nitrogen content; (d) nitrogen content or TPD nitrogen yield vs. activation yield.

be accounted for by a pure thermal preferential surface denitrogenation either of previously activated carbon material or during the activation process itself, which involves both purely thermal and thermo-oxidative reactions. This is suggested by Fig. 6c, where nitrogen appears to be more easily thermally desorbed from activated carbons with a larger surface area than from massic/non-activated ones.

Assuming that differential surface denitrogenation is the most important factor contributing to the deviation of the N correlation line below the surface-bulk parity, the smallest deviation for the samples containing the lowest nitrogen content could be explained by a more intense denitrogenation process that would have minimized the difference between surface and bulk. On the contrary, a better preservation of the initial nitrogen contained in the precursor polyaramid fibers would underline the nitrogen surface depletion observed by XPS. In this respect, the preferential surface denitrogenation and its specific effects on the nitrogen distribution (Fig. 5) are more obvious for the H3PO4 activated carbons, where the difference between surface and bulk N/C is linearly correlated with the carbonization/activation yield. Hence, the correlation between the nitrogen content and the carbonization/activation yield (Fig. 6d) illustrates the surface denitrogenation of the Nomex fibers by the activating agents. Freeman et al. [47] found that Nomex based chars contain much less nitrogen than comparable Kevlar-derived chars (as confirmed by Cuesta et al. [48]), possibly due to the accessible and more uniform nature of the char structure, thus allowing reaction of nitrogen groups with carbon dioxide or steam to occur more freely. Several authors have reported that, in comparison with CO2, steam specifically reacts with nitrogen groups, thereby reducing more the N/C ratio with increasing burn-off during physical activation of polyacrylonitrile chars [49,50].

4. Conclusions

In the framework of a work on the chemistry of porosity development during carbon activation and of the resulting carbon surface, we have conducted a combined study by TPD, XPS and XANES on the local bonding environment around nitrogen in Nomex-based activated carbon fibers, as a function of the activation mode and of the burn-off degree, in order to understand the changes in nitrogen content and nitrogen functionality distribution during carbonization/activation. The nitrogen bonding structure was shown to vary with the preparation treatments. Comparison of the data showed that in all cases thermal reactions occur preferentially at the sample surface and involve both aromatization reactions forming more condensed nitrogen species and degradative reactions of condensed pyridinic nitrogen with further oxidation of nitrogen atoms located at the edge of polyaromatic systems. These reactions occur more sequentially during physical activation, which involved a carbonization stage followed by an activation stage with a co-occurrence of isothermal heating (aromatization) and thermal-oxidation. At equivalent burn-off, the extent of nitrogen removal was lower during CO2 activation than during steam activation, which produced an attack of the nitrogen-containing polyaromatic systems with hydroxyl oxygen group formation by OH' or H' radical reaction with the carbon surface. Impregnation with
H:PO₄ led to a higher carbonization/activation yield and a greater preservation of N-containing groups due to early dehydration and crosslinking reactions forming the micropore network. In this activation mode, the nitrogen group distribution is also controlled by two simultaneous reactions: strong aromatization, and surface oxidative denitrogenation due to a secondary activation triggered by CO₂. This enhanced denitrogenation due to CO₂ activation in the combined cycles enhanced a stronger thermal char denitrogenation instead of pure N₂ formed by surface or gas phase radical interactions.

During subsequent CO₂ activation in the combined cycles oxygen substituents or intracyclic oxygen atoms making the cycle more prone to be open by thermal cracking. This opening would occur during TPD under isothermal 


