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Raman Spectroscopy Study of the Transformation of the Carbonaceous Skeleton of a Polymer-based Nanoporous Carbon Along the Thermal Annealing Pathway

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Abstract

We report a multi-wavelength Raman spectroscopy study of the structural changes along the thermal annealing pathway of a poly(furfuryl alcohol) (PFA) derived nanoporous carbon (NPC). The Raman spectra were deconvoluted utilizing G, D, D’, A and TPA bands. The appropriateness of these deconvolutions was confirmed via recovery of the correct dispersive behaviours of these bands. It is proposed that the \(I_D/I_G\) ratio is composed of two parts: one associated with the extent of graphitic crystallites (the Tuinstra-Koenig relationship), and a second related to the inter-defect distance. This model was used to successfully determine the variation of the in-plane size and intra-plane defect density along the annealing pathway. It is proposed that the NPC skeleton evolves along the annealing pathway in two stages: below 1600°C it was dominated by a reduction of in-plane defects with a minor crystallite growth, and above this temperature growth of the crystallites accelerates as the in-plane defect density approaches zero. A significant amount of transpolyacetylene (TPA)-like structures was found to be remaining even at 2400°C. These may be responsible for resistance to further graphitization of the PFA-based carbon at higher temperatures.

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1. Introduction
The non-crystalline nature of nanoporous carbons (NPCs) makes characterisation of their carbonaceous skeleton a challenge. It is generally accepted that the skeleton is largely composed of nanometre-sized \( sp^2 \)-dominated building blocks \([1-6]\). Careful diffraction \([5, 7-10]\) and transmission electron microscopy (TEM) \([2, 3, 11]\) studies suggest these building blocks could be characterised as defective graphene-like layers with limited stacking order. Apart from the abundance of inter-layer boundaries due to their nanoscale size, intra-layer defects have also been observed in recent atomic-resolution electron microscopy studies \([12-14]\), including in the form of non-hexagonal rings. Structural models, that captures microporosity as well as defective carbonaceous skeleton have been proposed of this material \([1, 9, 10, 12, 15, 16]\) although there is still debate over their validity \([1]\). A quantitative understanding of this defective structure is important for tailoring NPCs to have desired characteristics in the contexts of adsorption \([14]\), thermal \([17]\) and electrical \([18]\) conductivity, and Li-ion capacity \([19]\).

Although electron microscopy techniques have played a major role in the study of structural defects within NPCs, they are much less useful when a quantitative analysis representative of the bulk volume is required. Raman spectroscopy, on the other hand, is sensitive to structural defects in bulk-scale samples. It has, for example, long been used to determine the in-plane size of the \( sp^2 \) building blocks \( (L_a) \) from the ratio of the intensities of the D and G bands \( (I_D/I_G) \), the two most outstanding features in NPC Raman spectra. It is now known that the Tuinstra-Koenig relationship \([20]\) (and its recent modification \([21, 22]\)) used to achieve this has its origins in the link between the D band and edge sites of the graphenic building blocks \([23]\) on the one hand, and the dependence of the G band on the building blocks in their entirety on the other. This and the fact that the relationships were derived using data for highly graphitic materials that likely contain low levels of intra-layer defects suggests care should be exercised in their application to NPCs. One possibility to address this concern is to use the relationship between the \( I_D/I_G \) ratio and intra-plane point-like defects derived recently \([24-27]\) in conjunction with the Tuinstra-Koenig relationship. This has not, however, been done to date as far as the authors are aware.

Theoretical studies \([28, 29]\) have indicated that structural details of carbon materials are also carried in other parts of the Raman spectrum. Of interest here are those that have been experimentally observed for disordered carbons and are thought to be related to defects, namely the D’ \([22, 30-34]\), A \([35-41]\) and TPA \([40-42]\) bands. Similar to the D band, the D’ band is caused by defect-induced breaking of translational symmetry \([32]\). The origins of the A band, on the other hand, is less well understood. Early studies attributed it to the ‘amorphous phase’ \((i.e.\ non-crystalline\ sp^3\)-dominated carbon) \([35-37]\), whilst more recent theoretical and experimental studies \([38-41]\) suggested it may come from non-hexagonal ring defects in the \( sp^2 \) system. The TPA band has been linked to both
sp²-based transpolyacetylene-like chains (see Fig. 1) at layer edges [42] and, more recently, to zig-zag edges [40, 41], although this assignment is questionable given the contrast between the abundance of zig-zag edges in these materials and the relatively low intensity of TPA bands that appear in their Raman spectra.

![Fig. 1. Transpolyacetylene-like structure.](image)

Evidence [39] has been found for the coexistence of all five Raman bands discussed above in NPCs. A thorough interpretation of NPC Raman spectra should, therefore, in principle involve consideration of all these bands during deconvolution (this is not to say that all may ultimately be included in a deconvolution). The use of multi-wavelength laser sources can also help improve deconvolution reliability as the fitted results can be checked against the expected excitation energy dispersive behaviour [23, 31, 32, 40-43].

In this paper we report a multi-band/multi-wavelength Raman spectroscopy investigation of the change in the defects of a nominally non-graphitizing NPC along an annealing pathway. The laser sources used range from the UV (266 nm) to the near-IR (785 nm), and the first order spectra obtained using these sources were deconvoluted successfully by considering the five Raman bands discussed above. The success of these deconvolutions was validated against the expected energy dispersive behaviour as well as XRD-derived data for the carbons. The relationship between the \( I_D/I_G \) ratio and intra-plane point-like defects [24-27] was then used in conjunction with the Tuinstra-Koenig relationship to derive both the cluster size and intra-plane defect density. This relationship was tested by deliberately introducing point-like defects into selected samples using controlled Ar⁺ ion bombardment. The origins of the A band was also investigated.

2. Materials and methods

2.1. Sample preparation

In order to reduce as much as possible experimental uncertainties, the base carbon studied here, termed C45SCO₂ hereafter [44], was derived from a furfuryl alcohol (FA) precursor using a carefully controlled process. The full details of this process may be found elsewhere [44] but are briefly outlined here for convenience. The as-received FA was vacuum-distilled to remove any contaminants before being mixed under an inert atmosphere with oxalic acid dehydrate as a polymerization catalyst. Small batches of the mixture were then heated in a horizontal quartz tube-furnace to 150°C under continuous argon flow before being soaked for 1 hour to bring about polymerization. Carbonization was then done under argon by further increasing the temperature to 800°C before being soaked for 2 hours. The cooled char was then broken up into chunks using a clean zirconia press before being ball-milled and sieved.
to obtain a powder with a particle size distribution of 38-106 µm. The powdered material was then activated to 45% conversion using CO₂ at 805°C for 48 hours. It has been shown that this slow activation protocol leads to homogeneous particles [44]. The carbon was kept in glass vials under an argon atmosphere until used.

A range of NPCs were derived from the C45SCO₂ carbon by subjecting it to heating under an argon atmosphere at 1000, 1600, 2000 and 2400°C; these carbons are referred to henceforth as C1000, C1600, C2000 and C2400. The C1000 material was produced under an Ar (99.9%) atmosphere using a quartz tube-furnace (Lindberg, USA) with a heating rate of 10°C/min and soak time of 2 hours. All the remaining annealed products were produced under an Ar (99.999%) atmosphere using a graphite furnace (Xerion, Germany) with a heating rate of 15°C/min up to 1500°C and 10°C/min thereafter before being soaked for 1 hour.

Two final carbons, denoted hereafter by C45SCO₂-Ar and C2400-Ar, were obtained by Ar⁺ ion bombardment of the base C45SCO₂ and C2400 carbons respectively. This involved the particular carbon first being mounted onto a 1×1 cm silicon substrate using double-sided sticky tape with a copper interlayer before being inserted into an ultra-high vacuum system (SPECS, Germany) with a base pressure of a few 10⁻¹⁰ mbar. The sample was then uniformly bombarded with 1.6×10¹⁵ Ar⁺ ions produced at 1000 eV and 0.04 µA.

2.2. XRD analysis

The base and thermally annealed carbons were subject to XRD analysis using a Miniflex 600 bench-top diffractometer (Rigaku, Japan) equipped with 40 kV Cu Kα X-ray source. The diffraction patterns were collected over the 2θ range of 5° to 100° at a scanning rate of 10° per minute and 0.02° per step.

Although use of the graphitic crystallite model [45] as a basis for analysis of NPC XRD diffraction patterns is questionable [46], we report such an analysis here given the Tuinstra-Koenig relationship is based on high graphitic materials. We have used the 10 and 002 peaks in the diffraction patterns to obtain the average in-plane extent, \( L_a \), and height, \( L_c \), of the crystallites respectively via the Scherrer equation [45]

\[
L_i = \frac{k_i\lambda}{B_i\cos\theta_i}
\]

where \( \lambda \) is the X-ray wavelength (0.154 nm for the Cu Kα radiation used here), \( \theta_i \) and \( B_i \) are the scattering angle and full width at half maximum (FWHM) values for the relevant peak, respectively, and \( K_a = 1.84 \) and \( K_c = 0.89 \).

The average spacing between graphenic layers was also estimated using the Bragg equation

\[
d_{002} = \frac{\lambda}{2\sin\theta_{002}} = \frac{1}{k_{002}}
\]

where \( k_{002} \) is the wave vector magnitude associated with the 002 peak, which can be determined by first subtracting the small angle scattering intensity from the spectra and then fitting a Gaussian function to the peak.
2.3. Raman analysis

Raman spectroscopy was performed on a Labram HR Evolution spectrometer (Horiba Jobin-Ivon, France) equipped with Synapse charged coupled device detection system (Horiba Jobin-Ivon, France). Four laser sources were used: a near-IR (NIR) laser (785 nm, 1.58 eV), two visible lasers (633 nm, 1.96 eV; 532 nm, 2.33 eV) and an UV laser (266 nm, 4.66 eV). Objective lenses of 50× and 40× magnification were used for the visible/NIR and UV lasers respectively. Low laser powers (15 mW for visible/NIR and 10 mW for UV) were used to avoid sample damage due to over-heating. The materials were carefully examined after analysis and no visible damage was found.

In each Raman measurement, a thin layer of sample was spread onto a glass substrate and then mounted into a triple-axis sample stage. Spectra were collected in mapping mode using a step size of 6 µm in both X and Y directions over 30×30 µm areas. Spectra were collected using an exposure time of 3 seconds. A total of 36 spectra were collected from each mapping, and the sum of these was used for each carbon sample/laser source combination. No significant variation was found among the 36 spectra in one collection as well as scans on different particles of one sample for test purpose.

All Raman spectra were fitted with combinations of G, D, D’, A and TPA bands. The initial positions of the Raman bands were specified by adopting the values reported for graphenes and nanographites and also considering their excitation energy dispersion behaviours [23, 31, 32, 40-43]. The fitting was done using the Levenberg-Marquardt algorithm as implemented in Origin Pro 9.0 (OriginLab Corporation, US). The position, width and area of each band were varied simultaneously in the fitting. A total of 50000 iterations were done for each fit. At least 10 fits with reduced $\chi^2$ values between 1.1 and 1.5 were obtained for each spectrum analysed ($\chi^2$ values in the range of 1-3 indicate good fits [48]). The final band parameter values were the average of the ten best fits; standard deviations were also evaluated from these fits.

3. Results and Discussion

3.1. XRD analysis

Fig. 2 shows the XRD patterns of the base carbon (C45SCO₂) as well as the annealed products. The 002, 10 and 11 peaks, which are commonly found for NPCs [1, 46], exist for all samples. Although the 004 peak is absent from the base and C1000 carbons, it appears in the C1600 material and grows along the annealing pathway. A small high graphitic component (2$\theta = 26.5^\circ$, indicated by a red arrow in Fig. 2) also similarly appears and grows along the pathway.

Both the 10 and 11 peaks show increasing intensity and narrowing along the annealing pathway, indicating improving in-plane coherence length and atomic ordering. Similarly, the 002 peak narrows and grows in intensity between the base and C2400 material, where it is quite sharp and located at 2$\theta = 26.5^\circ$ in line with a high graphitic stacking phase [49]. These results
collectively suggest increasing graphitization of the carbon skeleton along the annealing.

**Fig. 2.** XRD patterns of base carbon and annealed products derived from it.

The average extents of the graphitic crystallites, $L_a$ and $L_c$, and average inter-layer spacing, $d_{002}$, evaluated from the XRD scattering patterns are plotted in Fig. 3 against the annealing temperature. The average inter-layer spacing of C45SCO$_2$ is about 0.44 nm, 32% higher than that of graphite. Together with the broadness of the 002 peak, it reflects the lack of inter-layer ordering in this material. The inter-layer spacing steadily decreases along the annealing pathway until it reaches 0.34 nm at 2400°C, just slightly greater than that of graphite. Fig. 3(b) and (c) show, on the other hand, that the crystallite extents steadily grow along the annealing pathway. The height of the crystallites in the C2400 material grows to around 4 times that in the base material whilst the in-plane extent is

**Fig. 3.** Change of the graphitic crystallite parameters along annealing pathway: (a) average inter-layer spacing; (b) in-plane extent; and (c) normal-to-plane extent. The values for the base carbon are plotted for reference at the temperature where activation took place. Error bars are one standard deviation from five repeat measurements.
nearly double. Despite the significant growth in size of the crystallites, they are still modestly sized as one may expect from a non-graphitizing carbon [50]. Interestingly, although the inter-layer spacing drops in a linear manner, the rate of growth in the crystallite extents is retarded up to 1600°C compared to above this temperature, suggesting some additional resistance to crystallite growth below this temperature.

3.2. Raman Band Analysis

3.2.1. Overview
Fig. 4 shows the Raman spectra of the base carbon (C45SCO2) collected using the four different lasers along with their deconvolution and associated residual; all the spectra are normalized to the maximum of the intensity of the G band located in every case at around 1600 cm⁻¹. The spectra for all but that obtained using the UV laser are similar in character. In addition to the G band, they possess a larger feature to its left that is the D band along with a clear small shoulder on its left. Comparison of Fig. 4(a)-(d) indicates that the area under the D band (henceforth termed the intensity) grows relative to that of the G band as the excitation energy decreases; this can be explained by the fact that the absolute cross section of the latter band is proportional to the fourth power of the energy (approximately, as we will see below) whilst that of the D band is independent of the energy [22]. The D band also upshifts with the excitation...
Fig. 5. Normalised Raman spectra of the carbons (black points) along the annealing pathway obtained using an excitation wavelength of 532 nm and their deconvolutions and associated residual (see Fig. 4 for colour scheme): (a) C1000; (b) C1600; (c) C2000; and (d) C2400.

energy, which is in line with the 50 cm⁻¹ per eV energy-dispersive nature of the D band [31, 32, 43, 51, 52]. Although the spectrum obtained using UV excitation differs from the others in that the G band is clearly the only dominate feature, with the D band being reduced to a small shoulder on its left at around 1430 cm⁻¹, this behaviour also has its origins in the differences in energy dispersive and absolute cross-section character of the two bands.

It was found that the best fits to the experimental spectra shown in Fig. 4 were achieved by using, in addition to the G and D bands, the TPA band [40-42], which is responsible for the shoulder on the left of the D band, the A band [35-41], which contributes to the intensity between G and D band, and the D' band [22, 30-34], which is located just to the right of the G band. Whilst the residual plots in Fig. 4 indicate satisfactory fits, this is confirmed below through the recovery of the correct excitation energy dependence of the TPA, D and D' band positions [22, 23, 31, 32, 40-43].

Fig. 5 shows the Raman spectra of the four carbons along the annealing pathway obtained using an excitation wavelength of 532 nm; once again, they have been normalised using the intensity of the G band. It is clear from this figure that the positions of the bands are invariant with temperature. Their intensities are, however, dependent upon temperature as one may expect given the XRD analysis suggests the carbonaceous skeleton is changing along the annealing pathway. The narrowing of the G and
D bands combined with diminishing TPA and A bands, with the latter disappearing in the C2400 material, is all suggestive of an increasing degree of graphitization along the annealing pathway. Greater detail of these changes will follow below when considering the individual bands more deeply.

3.2.2. G band

Fig. 6(a) shows the variation of the G band position along the annealing pathway for the four Raman excitation wavelengths; the differences between the variations for the different wavelengths is within the uncertainty of the data, in keeping with the non-dispersive nature of the G band. Whilst the trend in the figure is not unambiguous, it appears as if the G band position decreases along the annealing pathway towards graphite’s G band position of 1580 cm$^{-1}$. This suggests the decrease seen here reflects an increasing degree of graphitization [23, 53], in line with the XRD results presented in the previous section.

The upshift of the G peak position of the base material relative to graphite is in line with that seen in studies of other $sp^2$-dominated carbonaceous materials like pyrolytic chars [31, 43, 54, 55], carbon blacks [37, 56], activated carbons [36, 57], glassy carbons [43, 51] and carbon fibres [58, 59]. The upshift is, however, in keeping with that seen for carbon materials with significant $sp^3$ content, where shifts well beyond 1600 cm$^{-1}$ have been observed [52, 60-62]. Ferrari *et al* [52], for example, observed the G band position of a tetrahedral amorphous carbon to decrease from 1690 cm$^{-1}$ at 88% $sp^3$ content to 1585 cm$^{-1}$ at zero content. Comparing this with the G band positions of the carbons here suggests they contain little, if any, $sp^3$ carbon. This is also confirmed by the absence of the T band in the UV spectra [62] for the carbons studied here.

Fig. 6 shows the variation of G band characteristics with annealing temperature for the four laser excitation wavelengths: (a) position; and (b) FWHM. The values for the base carbon are plotted for reference at the temperature where activation took place. Lines are a guide to the eye only.

Fig. 6(b) shows the variation of the G band FWHM with annealing temperature; once again, within the uncertainty, no excitation energy-dependence is seen here. The FWHM does appear, however, to decrease along the annealing
pathway, with the decrease accelerating beyond 1600°C towards a value that is around double that of graphite, which is 15 cm$^{-1}$. As in-plane defects [24, 63] and small cluster size [22] are both known to broaden the G band, these trends suggest the level of defects is decreasing and/or the lateral dimensions of the clusters are increasing along the annealing pathway, once again in keeping with the XRD results.

### 3.2.3. D band

Fig. 7 shows the variation of the D band position of all the carbons studied here with the laser excitation energy. This shows that, within the uncertainty of the data, the position of the D band remains unchanged along the annealing pathway. It does, however, vary with the excitation energy at a rate of 50 cm$^{-1}$ per eV, entirely in line with prior observations for pyrolytic carbons [43] and a PFA char carbonized at 600°C [31]. This confirms the validity of the spectra deconvolutions.

![Fig. 7. Variation of D band position with excitation energy for the carbons.](image)

### 3.2.4. D’ band

The D’ band, which is a defect-induced feature that can be explained by intra-valley double resonance scattering process [32], is observed in all carbons along the annealing pathway for all excitation energies. This is in line with previous observations for multi-crystalline graphite [30], nanographites [40, 41], defective graphenes [24-26, 33, 34], soots [39], carbon blacks [31] and polymer derived carbons [31]. Fig. 8, which shows the variation of the D’ band position of all the carbons considered here with excitation energy, indicates that this position is dispersive for 10 cm$^{-1}$ per eV, which is once again consistent with that observed previously [23, 32]. This once again supports the validity of the spectra deconvolutions presented here.

![Fig. 8. Variation of D’ band position with excitation energy for the carbons.](image)

### 3.2.5. A band

The A band is observed for all the carbons excepting C2400 using the NIR and UV excitation. The presence of the A band here is consistent with previous observations for many types of disordered carbons, including carbon blacks [37], activated carbons [36], carbon onions [41], diamond like carbon films [35], ion-
bombarded graphites [63] and graphenes [24]. Fig. 9(a), which shows the A band position against excitation energy, indicates the position falls within a narrow range (±30 cm$^{-1}$) of 1530 cm$^{-1}$.

As the variations are largely within uncertainty, the band position appears to be non-dispersive. Whilst this possibility has not been commented upon in the literature before, previous work on Ar$^+$ ion bombarded graphites [63] suggests that the A band might be a density of states feature (in common with the G band), and thus a non-dispersive character is not unexpected.

![Fig. 9. Key characteristics of the A band: (a) A band position variation with excitation energy for the carbons; and (b) variation of $I_A/I_G$ ratio with annealing temperature for the four excitation wavelengths.](image)

Fig. 9(b) shows the variation of the $I_A/I_G$ ratio along the annealing pathway for the four excitation wavelengths. This figure shows that the intensity ratio decreases along the annealing pathway, with the ratio for the non-UV excitation wavelengths decreasing by an order of magnitude between the base and C2400 carbon. The decrease is far smaller for the UV excitation, reflecting the ratio’s clear excitation energy dependence. This suggests the A band cross section is energy dependent.

![Fig. 10. Raman spectra of C2400 and C2400-Ar obtained using 532 nm excitation.](image)

Two different types of defect have been posited as responsible for the A band. The first are $sp^3$ interstitial sites attached to the $sp^2$ network [35-37]. The absence of any evidence of $sp^3$ sites in the carbons here (see discussion of the G band) suggests, however, that this origin is unlikely here.

The second type of defect, which has been proposed based on a theoretical study, is $sp^2$ based point-like defects, in particular non-hexagonal rings [38], which are certainly known to exist in NPCs like those considered here [12-14]. To test this possibility, the Raman spectrum of the C2400 carbon is compared in Fig. 10 to its Ar$^+$ ion bombarded counterpart, C2400-Ar. This shows that the bombardment leads to the re-emergence of the A band when it had disappeared in the C2400 material. Based on the simulation of ion bombardment on graphene [64], it is thought
that of the defects introduced by the ion bombardment used here, around 50% will be single vacancies (SVs), with double vacancies (DVs) being less common, and Stone-Wales defects (SWs) very uncommon. Thus, it is proposed here that the A band is likely to be linked to SVs and DVs.

3.2.6. TPA band
The TPA band, which has been related to transpolyacetylene-like structures [42], is seen along the annealing pathway. Fig. 11(a), which shows the TPA band position variation with excitation energy for all the carbon samples, indicates it is energy dispersive. Whilst it is not unambiguous, the results in this figure suggest there may be two types of energy dependence. For C45SCO2 and C1000, TPA band position shows a positive shift with excitation energy at a rate of \( \sim 35 \text{ cm}^{-1} \) per eV. For the remaining samples, however, the TPA band position shifts in the opposite direction at the same rate. Such positive and negative shifts of the TPA band have been reported elsewhere for nanographites [40, 41] and diamond-like carbons [42], and hence the results here may have some basis in fact, but clearly more detailed study is required.

Fig. 11(b) shows the variation of the \( I_{TPA}/I_G \) ratio along the annealing pathway. This figure shows that the intensity ratio decreases along the pathway for all the excitation wavelengths. However, unlike the A band, the TPA band remains present in all spectra right up to 2400°C. This suggests that the transpolyacetylene-like structures in the carbon considered here are relatively resistant to annealing. The figure also shows that the \( I_{TPA}/I_G \) ratio is influenced by the excitation energy. This is in part due to the substantial reduction in the G band cross section at low energies, but an energy dependence of the TPA cross section may also play a role.

Fig. 11. Key characteristics of the TPA band: (a) TPA band position variation with excitation energy for the carbons; and (b) variation of \( I_{TPA}/I_G \) ratio with annealing temperature for the four excitation wavelengths.

3.3. \( L_a \) and defect density from the \( I_D/I_G \)
The pioneering work of Tuinstra and Koenig [20] showed that \( I_D/I_G \) is inversely proportional to \( L_a \) in carbons of high graphitic content. More recent work on nanographites of controlled sizes ranging from 20 to 65 nm [22] suggested that the Tuinstra-Koenig relationship is in fact excitation-energy, \( E_t \), dependent as per
\[(I_D/I_G)_\alpha = \frac{cE_{l}^{-4}}{L_a} \]  \hspace{1cm} (3)

where \(c\) is a constant, and we have assigned the \(\alpha\)-subscript to the intensity ratio for a reason that will become clearer below. This excitation energy dependence, which is often termed the \(\omega^4\) (= \(2\pi E_l/h\)) dependence [65], originates from the energy dependence of the G band scattering cross section [22]. Fig. 12, which shows the variation of \(I_D/I_G\) against the inverse of the excitation energy for the carbons, indicates that the \(\omega^4\) dependence indeed holds for the carbons obtained at the two higher annealing temperatures, C2000 and C2400. This dependence does not, however, appear to hold for the other carbons, suggesting that \(\omega^4\) dependence is not applicable to materials whose structures are some way from being graphitic in nature. It is, therefore, proposed that the following more general form of Eq. (3) be used for NPCs

\[(I_D/I_G)_\alpha = \frac{cE_{l}^{-B}}{L_a} \]  \hspace{1cm} (4)

This equation suggests that plotting \((I_D/I_G)E_{l}^{B}\) against \((L_a)^{-1}\) should collapse all the data for the carbons onto a single line. As Fig. 13(a) shows, this is indeed the case for the two most annealed carbons here, C2000 and C2400. The others, however, fall above the line defined by these two carbons. This suggests there is another factor at play. It is now known that the Raman D band is not only affected by the in-plane extent, \(L_a\), but also in-plane defects [53]. The relationship between the \(I_D/I_G\) ratio and point-like defects

![Fig. 12. Variation of the \(I_D/I_G\) ratio with inverse excitation energy. The parameters obtained by fitting Eq. (4) are shown in the inset table.](image1)

![Fig. 13. Contribution to \(I_D/I_G\) from \(L_a\) and \(L_D\). (a) plot of A values \((E_{l}^{B}(I_D/I_G))\) against \(L_a^{-1}\), (b) the locations of C45SCO2, C1000, C1600, C45SCO2-Ar and C2400-Ar on the two-stage defect trajectory.](image2)
have been studied in recent years by considering graphenes modified by controlled ion bombardment [24-26, 33]. This work showed that \( \frac{I_D}{I_G} \) ratio is related to the average distance between nearest defects, \( L_D \) in a so called two-stage trajectory. At low in-plane defects densities (stage one, i.e. large \( L_D \)), this dependency was found to be inversely proportional to the square of the inter-defect distance

\[
\frac{I_D}{I_G} = \left( \frac{4.3 \times 10^3}{I_D} \right)^{0.5} E_t^{-B}
\]

(5)

where the \( \omega^4 \) dependence observed in this prior work [25] has been replaced by the more general energy dependence observed here, and the \( D \)-subscript indicates the intensity ratio is related to the in-plane defects. At \( L_D \) of around 3 nm, this Stage One dependency was found to change to Stage Two [27, 60] (the more general power dependency has once again been used) to

\[
\frac{I_D}{I_G} = 18.5 E_t^{-B} L_D^2
\]

(6)

Returning to Fig. 13(a), on the basis that the three less-annealed carbons appear to fall on a roughly straight line parallel to the line that passes through the C2000 and C2400 carbons, it is proposed here that the ratio of the intensities of the D and G bands be composed of two parts as per

\[
\frac{I_D}{I_G} = \left( \frac{I_D}{I_G} \right)_a + \left( \frac{I_D}{I_G} \right)_D
\]

(7)

Using this equation with Eq. (4) for \( \left( \frac{I_D}{I_G} \right)_a \) allows evaluation of the defect component, \( \left( \frac{I_D}{I_G} \right)_D \). Comparing it with the two-stage trajectory as expressed in Eq. (5) and (6) yields Fig. 13(b) This is further supported by the results obtained from the Ar-ion bombarded versions of the C45SCO_2 and C2400 carbon, which are also shown in Fig. 13.

The defect density, \( N_D \), can be evaluated given \( L_D \) using

\[
N_D (cm^{-2}) = \frac{10^{14}}{\pi L_D^2}
\]

(8)

Using this equation yields the defect densities in the inset table of Fig. 13(b). This table suggests that the in-plane defect density in the C1600 carbon is approximately half that of the base carbon, and the Ar-ion bombardment that the base carbon was subject to roughly doubled the defect density.

3.4. Defect evolution along the annealing pathway

The structural evolution along the annealing pathway studied here can be broken up into two separate stages. As the analysis of the \( \frac{I_D}{I_G} \) ratio in the previous section showed, annealing up to somewhere below 1600°C sees a rapid removal of defects accompanied by a modest increase of in-plane extent. Above 1600°C, however, this analysis suggests that growth of the in-plane extent accelerates. This analysis is further supported by the observed variation of the FWHM of the G band and \( I_A/I_G \) ratio along the annealing pathway (see Fig. 6 and Fig. 10 respectively), as well as the XRD analysis (see Fig. 3). Support for this proposed mechanism can also be found from an annealing study of a phenolic resin based carbon [6], which is also non-graphitizing, where they observed a rapid growth of \( L_a \) using XRD between 1600 and 2500°C.
Whilst Raman analysis clearly does not elucidate the atomistic-level annealing process, processes can be conjectured that are consistent with the analysis and other results, including that based on molecular modelling. The two stage process observed here suggests the presence of at least two groups of defects. The first group are particularly mobile at lower temperatures, and are thus likely to be constituted by, for example, point defects such as Stones-Wales defects, single point vacancies, and out-of-plane atoms (e.g. those cross-linking between adjacent stacked planes) [66-68]. These will migrate at the lower annealing temperatures (between the 800°C at which the material is formed and 1600°C) to form more stable structures, including higher order defects such as double vacancies [68]. The second group only become more mobile at higher temperatures, and are thus constituted by more complex defects. An example that may be relevant here are dislocation-like defects which, once removed, bring an increase of the in-plane extent.

4. Conclusion
The structural evolution of a PFA derived NPC along a thermal annealing pathway spanning from 1000 to 2400°C was studied using multi-wavelength Raman spectroscopy. The spectra obtained were deconvoluted using G, D, D', A and TPA bands. The validity of the deconvolutions was verified through inspection of the residuals and, more importantly, the recovery of the expected dispersive character of the various bands. Analysis of the A band suggests that it is associated with single- and double-point vacancies, such as 5- and 7-membered ring structures. There is clear support for this finding in the form of recent high resolution TEM images.

The $\omega^4$ dependence of the $I_D/I_G$ ratio was found to be valid only for the upper end of the annealing pathway. Below this, the power of the excitation energy dependency decreased with decreasing annealing temperature.

The analysis here strongly suggests that the $I_D/I_G$ ratio is the sum of two parts: one inversely proportional to the in-plane extent of the graphitic crystallites, $L_a$, (i.e. the Tuinstra-Koenig type relationship), and a second that is related to the inter-defect distance, $L_D$ [24-27].

Finally, drawing on the Raman results collectively, it is proposed that the carbonaceous skeleton transforms during annealing via a two stage process. For the annealing temperature range up to somewhere below 1600°C, removal of structural defects dominates, especially point-like defects, whilst the crystallite size undergoes a mild increase. At temperatures beyond 1600 °C, the rate of growth of the crystallites accelerates substantially. This mechanism is supported by XRD analysis along the annealing pathway.

There is room for further study of the transformations observed here. For example, there is a clear need to consider the transformations below 1600 °C with a higher degree of resolution (e.g. every 100 °C) and with further techniques such as X-ray and UV photoelectron spectroscopy in the hope to better...
understand the in-plane defect evolution in carbons prepared below 1600 °C.

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