# Nanostructure of Gasification Charcoal (Biochar)

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#### Abstract

In this work, we investigate the molecular composition and nanostructure of gasification charcoal (biochar) by comparing it with heat-treated fullerene arc-soot. Using ultrahigh resolution Fourier transform ion-cyclotron resonance and laser desorption ionisation time of flight mass spectrometry, Raman spectroscopy and high resolution transmission electron microscopy we analysed charcoal of low tar content obtained from gasification. Mass spectrometry revealed no magic number fullerenes such as  $C_{60}$  or  $C_{70}$  in the charcoal. The positive molecular ion m/z 701, previously considered a graphitic part of the nanostructure, was found to be a breakdown product of pyrolysis and not part of the nanostructure. A higher mass distribution of ions similar to that found in thermally treated fullerene soot indicates that they share a nanostructure. Recent insights into the formation of all carbon fullerenes reveals that conditions in charcoal formation are not optimal for fullerenes to form, but instead curved carbon structures coalesce into *fulleroid-like* structures. Microscopy and spectroscopy support such a *stacked*, *fulleroid-like* nanostructure, which was explored using reactive molecular dynamics simulations.



#### Highlights

- Gasification charcoal (biochar) was analysed using high resolution mass spectrometry revealing no magic number fullerenes i.e. C<sub>60</sub> or C<sub>70</sub>.
- Heated fullerene arc-carbon also lacked magic number fullerenes but contained oxygenated fragments matching that of charcoal indicating a shared *fulleroid-like* nanostructure.
- Raman spectroscopy and high resolution transmission electron microscopy supported a *stacked*, *fulleroid-like* nanostructure.

# Contents

1	Introduction					
2	Materials and methods Results					
3						
	3.1	High resolution mass spectrometry	8			
	3.2	Raman spectroscopy	11			
	3.3	High resolution transmission electron microscopy	12			
4	4 Discussion					
5	Con	clusions	16			
	Refe	erences	17			

## **1** Introduction

Charcoal is formed from the thermal transformation of a carbon-rich substance, often woody biomass, into a solid carbon product. It has been produced in fires and kilns for millennia and has been used for creating art, smelting ores to produce metals and as a smokeless fuel [3]. Recently, advanced applications have been developed for charcoal such as electrode materials for lithium-ion batteries [36], activated carbons [6] and supercapacitors [55]. Environmental concerns have also prompted research into charcoal as a carbon capture technology. Storing atmospheric carbon is now possible considering that biomass, as it grows, photosynthetically captures carbon dioxide. Subsequent thermal treatment traps some of that carbon in a stable charcoal that will not break down. As this charcoal is also known to enhance crop yields (allowing costs to be offset), many researchers are suggesting addition of charcoal in order to improve soils while storing carbon (charcoal produced for application to the soil is often called biochar) [34, 44].

One means of offsetting biochar's cost is by producing energy through a process called gasification [10]. The gasification process injects a restricted supply of oxygen into a sealed vessel that oxidises some of the carbon in the charcoal to produce high temperatures (1000 °C). The surface of the hot carbon is then able to perform the gasification reactions, reducing carbon dioxide and water to carbon monoxide and hydrogen. The hot carbon is also able to crack unburnt hydrocarbons (tar) [4, 43] leading to a clean syngas fuel that can power a turbine or internal combustion engine. The high temperatures reached inside a gasifier produces a high quality charcoal with low tar content (low hydrogen content), which is necessary for the long-term stability of the material [34, 51]. The nanostructural understanding of gasification charcoal is therefore important for determining the reactivity of the char for producing syngas and also for considering the long-term stability of such a material for carbon capture and storage. Understanding of the material properties could also stimulate interest into new applications for carbon nanostructured materials produced from natural biomass precursors.

Concentrating on woody biomass and considering how it thermally transforms into charcoal in the absence of air (pyrolysis) a dynamic molecular structure has previously been proposed [31]. In this scheme, initially the crystalline and non-crystalline saccharide precursors dehydrate, decarboxylate and depolymerise forming pyrogenic amorphous carbon [35]. Analysis of hydrothermally treated saccharide suggests a pyrogenic structure containing six-membered aromatic rings and species such as hydroxymethylfurfural [5, 53]. From 400–700 °C ring condensation leads to the formation of stacked aromatic domains [29]. These small layered turbostratic (graphitic without ideal ABA stacking) disordered regions are completely developed by 700 °C and have been recently imaged in charcoal [54]. At the temperature the material is nearly entirely composed of carbon (95 wt%). Thermal treatment to 1000 °C further decreases the amount of oxygen to 1-3%, removes any remaining hydrogen (<1 wt%) and increases the graphitic turbostratic domains. The gasification process also includes oxidation and activation of the charcoal during reduction, which reduces the density and integrates porosity [23].

Significant experimental evidence has emerged suggesting a *fullerene-like* nanostructure for charcoal [21] (where a fullerene is a molecule with carbon atoms arranged in a closed network of pentagonal and hexagonal rings [48]). Pentagonal and heptagonal rings have

been resolved by use of high resolution transmission electron microscopy (HRTEM) in microporous carbons and charcoal [19, 23, 54] as well as closed cages, and curved fragments in carbon heated to higher temperatures (>1500 °C) such as glassy carbon [20]. A recent review of the nanostructure of non-graphitising carbon, focusing on charcoal in particular, has also highlighted the role of oxygen in inhibiting planarisation of the structure [42] and experimental results have suggested the mechanism for pentagon integration is the loss of oxygenated fragments along the zig-zag edge of aromatic species [1].

Laser desorption time of flight mass spectrometry (LDI-TOF MS) is one of the many techniques that has been used to study the structure of these non-graphitising carbons. In a study of graphite, diamond, glassy carbon, carbon nanotubes and *diamond-like* thin film, Sedo et al. observed only in glassy carbon some high intensity, positive ions in the range m/z 400-800, the highest being m/z 701 [49]. The ions were discarded as an artefact by the authors but were later observed by Bourke and co-workers in biomass and sugar-derived charcoals prepared with flash pyrolysis at 950 °C [7]. Sedo et al. also found that by increasing the laser power a distribution of more massive ions was produced. However, the absence of the m/z 720 ion (C<sub>60</sub> fullerene) in charcoal and also of large even-numbered carbon cages in glassy carbon called into question a *fullerene-like* nanostructure.

In this article, we aim to understand how the mass spectrometry of charcoal, indicating no fullerenes, can be understood in light of the microscopy and spectroscopy, suggesting curved fullerene-like fragments, by comparing a gasification charcoal with thermally treated fullerene arc-carbon, which we have previously studied [39].

### 2 Materials and methods

#### **Production of charcoal**

Untreated pinus radiata wood sourced from Carter Holt Harvey Wood Products, New Zealand (NZ), was cut into  $(30 \times 30 \times 10 \text{ mm})$  blocks that were then carbonised with the use of a custom built downdraft gasifier based on the microlab class gasifier with Fluidyne Gasification Ltd, NZ. Figure 1 shows the reaction segment of the gasifier. Wood flowed under gravity and heated from below was dehydrated and then thermally pyrolysed into charcoal. Nozzles were then used to inject air and the charcoal was oxidised. Once the oxygen had been consumed, reduction of CO<sub>2</sub> and H<sub>2</sub>O occurred on the surface of the hot carbon. Charcoal then dropped through the grate and was collected in the lower chamber.

Positive pressure was supplied to the air inlet connected to the nozzles and the gasifier was ignited and run for 20 minutes. The reactor was then starved of oxygen and cooled for a period of two hours. Charcoal was taken from the interface between the pyrolysis and oxidation zones, which had yet to undergo significant oxidation and activation but had attained high temperatures. These zones are well separated in the downdraft gasifier we have employed. Charcoal from this region is not significantly oxidised but does not have soot present on its surface, which was determined by a visible change in the charcoal's appearance. Before analysis of the charcoal by LDI-TOF MS the carbon was stored in evacuated glassware in order to prevent contamination from plasticisers and other envi-



**Figure 1:** Photograph of the Microlab class gasifier (left). Schematic of the gasifier throat showing the different reaction zones (right). Used with permission from Fluidyne Gasification Ltd, NZ.

ronmental species.

Ultimate analysis of the C, H, N, S and ash percentage weights (%wt) was determined by use of the ASTM D 3176 standard (Table 1). Oxygen %wt was determined through difference and does not include oxygen contained in the ash oxides. The Campbell Micro Analytical Laboratory at the University of Otago conducted the analysis of the oven dried charcoal.

**Table 1:** Ultimate analysis of gasification charcoal (%wt)

C	Н	0	Ν	Ash
90.80	< 0.1	3.34	< 0.1	5.8

The charcoal prepared in this study was found to contain 3.34% oxygen. This is consistent with the flash pyrolysis process which produced charcoals with an oxygen content of between 2–5% at 950 °C [7]. A low percentage of hydrogen was found in the structure, indicating tar is not present in the charcoal, i.e. a highly crosslinked nanostructure.

#### LDI-TOF mass spectrometry of charcoal

A Bruker UltrafleXtreme Matrix-Assisted Laser Desorption Ionisation Time of Flight/Time of Flight (MALDI-TOF/TOF) mass spectrometer was used for characterising the positive ions corresponding to the mass distribution of carbon fragments. Ions were produced by use of a nanosecond pulsed 355 nm frequency-tripled Nd:YAG laser (1 kHz repetition rate, 5 ns pulse duration, maximum power of 2.3  $\mu$ J, spot size of 100  $\mu$ m with a maximum laser fluence of 29.3 mJ/cm<sup>2</sup>). The Bruker Peptide calibration standard was used to calibrate the instrument.

#### ESI FT 9.4 T ICR mass spectrometry

For analysis the charcoal sample was ground into a fine powder with a mortar and pestle. 700 mg of powder was suspended in 30 mL of toluene and sonicated for three hours to extract any soluble sample components. 2 mL aliquots were placed in four centrifuge tubes and centrifuged for 30 minutes and the supernatant (stock solution) pulled. For (+) Electrospray Ionisation (ESI) analysis, the stock solution was diluted with methanol to a final composition of 50:50 MeOH:Stock with 1% HCOOH modifier and analysed by ESI Fourier Transform Ion Cyclotron Resonance Mass Spectrometry (FT ICR MS) at the National High Magnetic Field Laboratory, Florida State University.

#### **Raman spectroscopy**

The Raman spectrometer (Horiba Jobin Yvon LabRAM HR) was equipped with an Olympus BX41 microscope (×10 objective). The instrument was calibrated based on the 520.5 cm<sup>-1</sup> line of silicon. Raman spectra were excited by a Nd:YAG laser (532.8 nm, 16 mW) at a nominal resolution of 4–6 cm<sup>-1</sup> in the range 1000–1800 cm<sup>-1</sup>. The acquisition was repeated four times with an exposure time of 8 s to improve the signal to-noise ratio. A linear baseline was applied to the graph which was optimised as the peaks were fitted (ax + b;  $a = 0.405 \pm 0.003$ ,  $b = 985 \pm 5$ ).

The fitting procedure was based on the work by Smith et al. who applied electronic structure calculations to a variety of aromatic molecules with defects and non-hexagonal rings [50]. Each species was found to have Raman excitations at many different frequencies. Due to the variety of carbon structures and defective carbon in charcoal it would seem inappropriate to fit a single Lorentzian but instead to fit regions of the spectrum with Gaussian functions. This was justified by supposing that very disordered materials will have a random distribution of Lorentzian functions about each major resonance which would have a Gaussian shape. In chars prepared at lower temperatures the ring vibrations indeed display a more Gaussian profile [50]. Structures with more homogeneous aromatic molecules such as in soot show more of a Lorentzian profile and the fitting of a single Lorentzian to each is more justified. The goal then of fitting regions with Gaussian functions is to quantify the amount of certain types of defects determined from electronic structure calculations. It should be noted, however, that the conclusions made are also found if Lorentzian functions are used instead of Gaussian functions for the D and G bands as there is still substantial intensity between the D and G band which is usually fit using a smaller Gaussian function.

#### High resolution transmission electron microscopy

Images were captured by a FEI<sup>TM</sup> Tecnai F20 which operated at an accelerating voltage of 200 kV. This instrument is equipped with a SuperTwin objective lens with a  $C_S$  of 1.2 mm. Images were captured with a Gatan Imaging Filter (GIF) 200, fitted with a 1k by 1k CCD digital camera. Phase contrast in high resolution TEM images with 0.2 nm resolution were obtained from graphitic regions oriented so that their basal plane is at

Bragg angles with the electron beam. Charcoal was gently ground in a mortar and pestle. This did not alter the nanostructure of the carbon significantly, as noted by Harris et al. [22]. The carbon was then suspended in a solution of iso-propyl alcohol and was placed onto a C-Flat<sup>TM</sup> Holey carbon support containing holes one micron in diameter and only regions extending into the vacuum were imaged.

#### **Model construction**

We made use of a molecular geometry from the work of de Tomas et al. [12] (made accessible on the website carbonpotentials.org). This geometry was produced from a quenched molecular dynamic simulation by use of the REBO-II reactive potential in a periodic 7.58 nm wide cube [8] with periodic boundary conditions. In order to describe the surface, the geometry was sliced along the diagonal and an NVT molecular dynamics simulation was performed in the LAMMPS software package [45] with the same reactive forcefield. The Nose-Hoover chain thermostat was used to maintain a system temperature of 3000 K. A timestep of 0.5 fs was used with a total simulation time of 3.6 ns. In order to close the defective graphene that would normally be attached to the periodic image the simulation box was extended past the carbon structure to allow the surfaces to anneal. Figure 2 shows the molecular geometry. The non-hexagonal rings were determined by use of the algorithm developed by Franzblau [18]. Polypy, a python program by Jaap Kroes (https://github.com/jaapkroe/polypy), was used to render the geometry [28].

HRTEM images of this carbon model were generated using the multislice method. The software detailed in the book *Advanced computing in electron microscopy* by Kirkland [32] was used to simulate the propagation and scattering of electrons through the molecular structure. Parameters used for the simulation were an accelerating voltage of 200 kV, objective aperture of 20 mrad and a defocus of 20 nm.



Figure 2: Protocol for preparing the molecular geometry. A three dimensional periodic model prepared using the REBO forcefield [8] was used from de Tomas et al. [12]. The corner was cut and the surface was annealed using the same forcefield for 3.6 ns at 3000 K to anneal the surface. The structure is shown coloured by the coordination number (2 - red, 3 - white, 4 - blue). The different rings were then determined and are coloured according to ring number (5 - blue, 6 - white, 7 - red, 8 - yellow). Dashed box around the region is shown expanded in Figure 7

### **3** Results

### **3.1** High resolution mass spectrometry

Charcoal was taken from the interface of the pyrolysis and oxidation zones where temperatures of 1000 °C were attained (as measured via a K-type thermocouple) and soot was not present on the char in this region, which allowed the solid state charcoal to be resolved without interference from soot produced in the gas phase. Figure 3 shows the LDI-TOF mass spectrum of charcoal with a low laser fluence (10.3 mJ/cm<sup>2</sup>) providing ions with the prominent peak at m/z 701 and low abundances of other ions compared to flash pyrolysis, indicating low amounts of extractable tars and no magic number fullerenes  $C_{60}$  or  $C_{70}$  [7]. Elemental analysis also revealed low amounts of hydrogen, further indicating low tar, highly crosslinked and well developed nanostructure.



**Figure 3:** LDI-MS low mass positive ions from m/z 150–850 for gasification charcoal showing the prominent m/z 701. Inset right: a mass-scale expanded segment with mass-to-charge (m/z) in the range 700–705 following positive-ion ESI 9.4 T FT-ICR mass spectrometry analysis of a toluene charcoal extract. Represented by an asterisk is the calculated isotopic distribution for the species with elemental formula  $C_{36}H_{61}O_{13}$  corresponding to the peak assignment.

To resolve the elemental structure of the m/z 701 ion and determine if the ion is a constituent of the nanoscale structure, high resolution mass spectrometry was used. Toluene was used to extract the species. The extract was then injected into the electrospray ionisation ion source and analysed by a 9.4 tesla Fourier Transform Ion Cyclotron Resonance mass spectrometer (ESI 9.4 T FT-ICR MS). Figure 3 inset shows the experimental mass spectrum with the various isotopic peaks for the ion, with a measured accurate mass of 701.4092 Da. While wood contains a variety of elements, the presence of the m/z 701 ion in feedstocks containing only C, H and O, studied previously by the authors and others, [7, 49] led us to propose  $[C_{36}H_{60}O_{13}+H]^+$  as the candidate ion of which the difference between measured and calculated mass is 1.1 ppm. The candidate ion's elemental formula contains a significant amount of hydrogen and oxygen, indicating the ion is not aromatic but an early breakdown product of pyrolysis. The elemental composition determined from the ultimate analysis indicated significantly less hydrogen and oxygen in the material that would suggest a highly aromatic nanostructure (Table 1). Further analysis of the LDI-TOF mass spectrum of charcoal after passing through the entire gasification reactor showed the loss of the m/z 701 ion, indicating that it was broken down during the gasification process. Charcoals containing the m/z 701 ion were also sensitive to oxygen with exposure to air leading to the loss of the ion after two to three months. These results support the conclusion that the m/z 701 ion is an unstable trace product of pyrolysis and

is not part of the nanostructure. The FT ICR MS analysis also confirmed that there were no detectable amounts of magic number fullerenes such as  $C_{60}$  or  $C_{70}$  in this gasification charcoal.

Considering the LDI-TOF MS experiment again we found that by increasing the laser fluence (12.8 mJ/cm<sup>2</sup>) a distribution of higher m/z ions could be observed that is similar to that found for thermally treated fullerene arc-carbon [39]. Figure 4 a) shows the skewed distribution of peaks. Figure 4 b) shows an enlarged region of the mass spectrum showing that the distribution is made up of a collections of peaks that are separated by m/z 24 (C<sub>2</sub>). The isotopic peaks are indicative of predominately carbon species with one, two and three oxygen atoms integrated and the lack of intensity between these collection of peaks indicates firstly that no all-carbon fullerenes cages ions are present and secondly that these species are not dimers, trimers etc. of PAH.



Figure 4: a) LDI-TOF-MS mass spectrum of charcoal at high laser fluence (12.8 mJ/cm<sup>2</sup>). A distribution of peaks is observed separated by m/z 24. b) shows a region between m/z 1245–1265 with three species isotope-simulated responses overlayed (coloured). c) similar m/z region of the mass spectrum for heated fullerene arc-carbon showing similar oxygenated fragments from Martin et al. [39].

We have previously found a similar distribution of oxygenated fragments separated by m/z 24 in thermally treated fullerene arc-carbon where oxygen present in the structure, from exposure to air, became chemically integrated into a crosslinked curved nanostructure upon heating [39]. Figure 4 b) shows an enlarged region of the mass spectrum showing triplicate peaks of oxygenated ions. For comparison the same region is shown in Figure 4 c) for thermally treated fullerene arc-carbon we have previously collected [39] which show a similar triplicate separated by m/z 24.

Care must be taken, however, when analysing such a mass distribution in gasification charcoal due to the possibility of modifying the structure as a result of laser ablation [16]. We ensured that no fragmentation could be observed under m/z 200 for charcoal, which we have previously found to modify the distribution of higher ions found in fullerene arc-carbon [39]. By ablating graphite at the maximum laser fluence we produced these small fragments and did not observe fullerene ions, indicating the laser desorption process and vacuum chamber were not conducive to fullerene formation [39]. Another important consideration is that laser ablation could produce hot fragment that could close curved fragments in the gas phase [9]. With these caveats the similarities between the mass spectra of charcoal and thermally treated fullerene arc-carbon indicates a shared, curved nanostructure.

#### **3.2 Raman spectroscopy**

Raman scattering was also used to analyse the structure. The Horiba LabRam Raman microscope, equipped with a 532 nm laser, provided excitation of the sample and spectra were collected near the ring breathing modes  $(1000-1800 \text{ cm}^{-1})$  of the graphitic carbon. Recent electronic structure calculations of the vibrational frequencies for various defective polycyclic aromatic hydrocarbons have provided insights into fitting this highly structured region of the Raman spectrum for charcoal [2, 41, 50]. In particular, the region from 1400–1460 cm<sup>-1</sup> has been suggested to arise from the breathing mode of 5-membered rings [2, 50]. Figure 5 shows the Raman spectrum after baseline correction with the peaks from least-squared fitting (Values and errors in the fitted values can be seen in Table 2). We did not observe as much structure in the gasification charcoal Raman spectrum due to the higher temperature treatment as compared with the Raman spectrum collected for cellulose char heated from 400–700 °C by Smith et al. [50]. In particular, the SL and D<sub>S</sub> bands could not be observed. We also found the S band appeared to contain a strongly Lorentzian tail on the lower wavenumber shoulder leading us to fit the S peak with a Lorentzian. We were unable to fit two A peaks as has previously been fitted for Raman spectra of wood charcoal prepared via pyrolysis by Smith et al. [50]. We fit the A<sub>1</sub> peak, which corresponds to the breathing mode of the 5-membered carbon structures. Fitting a single Gaussian peak between the D and G band provided a peak position of 1457 cm<sup>-1</sup> which was closer to the A<sub>1</sub> peak (1400 - 1460 cm<sup>-1</sup>) than the A<sub>2</sub> peak (1480 - 1550 cm<sup>-1</sup>). This led us to assign this peak to the  $A_1$  breathing mode of 5-membered rings. We were unable to fit a second peak  $(A_2)$  in this region, previously attributed to sp<sup>2</sup> defects [50], which we consider another indicator of the high crosslinking in the sample. We also did not observe the same shoulder on the lower wavenumber side of the G peak as seen in low temperature charcoal [50], further indicating the lack of the A2 peak. The high temperature treatment appears to have significantly reduced or removed the peaks associated with disordered carbon such as the  $A_2$  and  $S_L$ .



**Figure 5:** Raman spectrum (532 nm excitation) for gasification charcoal (black points) with the peaks used to fit the spectrum shown (dotted line). The fit is shown (red line). For peak assignments see Table 2.

A strong C peak was also observed at  $1702 \text{ cm}^{-1}$  indicative of carbonyl species. Due to this peak the Lorentzian tail on the higher wavenumber shoulder could not be fitted with a second G<sub>L</sub> Lorentzian that describes the E<sub>2g</sub> mode for larger PAHs. Compared with the low temperature charcoal this strong C band indicates a larger amount of carbonyl oxygen species. This would align with our observation for thermal treatment of fullerene arc-carbon with oxygen converted to carbonyl species [39].

Position	Width	Peak	Shape	Assignment
1209±3	138±7	S	Lorentzian	Breathing mode for 7-membered rings or small PAH
				or breathing modes adjacent to heteroatom defects.
$1329 \pm 4$	$138{\pm}11$	D	Gaussian	Combined breathing modes for PAHs
$1457\pm2$	$115 \pm 1$	$A_1$	Gaussian	Breathing mode for 5-membered rings
1577±1	$141\pm3$	$G_G$	Gaussian	Asymmetric vibrations for small PAHs
$1702 \pm 1$	67±2	С	Gaussian	Carbonyl stretching mode

**Table 2:** *Peak assignments and fitted parameters (values in*  $cm^{-1}$ *)* 

### 3.3 High resolution transmission electron microscopy

Electron microscopy also indicated such a curved structure. We found that the lack of uncrosslinked extractable fraction (liquid tar) in the gasification charcoal improved the stability of the structure under the electron beam. Figure 6 shows transmission electron micrographs along the edges of different regions of the charcoal protruding into the

vacuum where observation of the geometry was least obscured. Figure 6a) shows a low magnification image of the edge of a charcoal particle. Curved regions can be seen around the edge. Figure 6b,c) shows higher magnification images. An *onion-like* structure can be observed in b). At the highest magnification, curved nanostructures with radii of curvature up to 1 nm observed. Alongside the curved structures, regions of layered graphitic planes can also be observed.



Figure 6: HRTEM images of gasification charcoal taken at 200 kV accelerating voltage.

Curved structures have been previously observed in HRTEM of charcoal [13, 36], multiwalled curved *onion-like* structures also seen in HRTEM [24] and scanning probe microscopy [33]. This curved nanostructure was similar to that found for sucrose char heated to 1000 °C in an inert environment with curved isotropic arrangement of fringes and dissimilar to that found for anthracene carbonised under similar conditions that produced anisotropic arrangement of fringes with low degree of curvature [21].

### **4** Discussion

We firstly discuss the reason for the absence of magic number fullerenes, which previously has led to doubt around a fullerene-like nanostructure for charcoal [7]. Recent advances in understanding magic number fullerene formation might help to explain why such species are unlikely to be found in charcoal [11, 15, 30, 39, 47]. Synthesis conditions required for magic number fullerenes are high-energy, non-equilibrium, dissipative environments such as in carbon arcs or low pressure oxygen-benzene combustion [26]. These environments allow curved aromatic structures to form and eventually close into disordered fullerenes [30]. The high-energies allows cages to eject and gain  $C_2$  [11, 15, 27, 47]. During this shrinking ( $C_2$  ejection) [30] and closed network growth ( $C_2$  gain), [15] the magic number fullerenes can be produced in high abundance due to the larger energy barriers for  $C_2$  loss or gain from these symmetric cages compared with their neighbours i.e. residing in *local* energy minima as showed by Curl et al. [11].

In contrast to the gas phase synthesis we have previously found that, by heating arcproduced magic number fullerenes in the solid state, magic number fullerenes are consumed and coalesce into giant fullerenes [39]. The ability of curved carbon nanostructures to fuse and coalescence is well known, for example the coalescence of fullerenes inside nanotubes leads to double walled nanotubes [25] and the fusing of single walled nanotubes into multiwalled nanotubes has been shown [37]. This reactivity has been studied via electronic structure methods and found to be due to the pyramidalisation of the carbon atoms which reduces the aromaticity around the defect and allows sp<sup>3</sup> bonds to be formed fusing the curved regions [14, 39, 55]. Importantly temperatures in most charcoal production do not reach those necessary to evaporate  $C_2$ , which is needed for evaporation and gain of  $C_2$  to form small magic number fullerenes. Therefore, if curved *fullerenelike* fragments are present in gasification charcoal the synthesis conditions would not be conducive to magic number fullerene formation.

Returning to the LDI TOF MS higher mass distribution of ions we found charcoal with an increase laser intensity, we can consider potential interpretations of what this distribution indicates about the nanostructure. One option could be that the distribution is due to dimers, trimers, tetramers etc. of predominantly planar stacked aromatic layers with oxygen functionalisation. We think this is unlikely for a few reasons. Firstly, the monomodal distribution is indicative of a continuous solid. Crosslinking of small PAHs is known to yield a multimodal distribution of peaks with a periodicity relating to the average molecular mass of the subunits [17]. Physical clustering in the case of aromatic molecules in soot formation is also known to lead to a multimodal distribution [46]. Secondly, the separation of the peaks by m/z 24 and the amount of oxygen per carbon peak is indicative of a similar curved nanostructure between heat treated fullerene arc-carbon and charcoal, which are both able to form fragments that close into oxygenated cages upon ablation. Thirdly, our previous study showed that ablation of the microcrystalline graphite containing planar carbon did not lead to formation of a distribution of peaks separated by m/z 24 as well as a lack of C<sub>60</sub> and C<sub>70</sub>, indicating that the laser power and vacuum inside the mass spectrometer is unable to produce fragmentation products and fullerenes when ablated [39]. However, from the HRTEM we do see evidence of stacking of aromatic layers interacting via van der Waals forces. One suggestion for why this stacking is not reflected in the mass spectrum could be due to an effect that Marsh commented on when discussing activated carbons [38] - many experimental techniques that probe carbon's nanostructure, such as HRTEM and x-ray diffraction, are most sensitive to regions where aromatic sheets are stacked, however, these sheets should extend past the stacked regions. This would lead to an underestimation of the aromatic network from considering only the stacked regions and not the single layers. The mass spectrum and HRTEM is consistent with a continuous curved nanostructure with regions of layering.

Figure 7 shows a molecular nanostructure which aims to consider the possible structures present in gasification charcoal generated from a modified quenched molecular dynamics study [12]. We have chosen not to consider oxygen in the model as it is only a minor component and therefore does not correspond to the long range graphitic nanostructure. The low amount of hydrogen in the elemental composition led us to consider a model only containing carbon. Figure 7 a) shows the surface of the nanostructure with the non-hexagonal rings that form due to the closure of the surface shown coloured. Both pentagonal ring providing bowl shaped surfaces and rings with greater than six rings providing saddle-shape geometries were observed, which allow for a large proportion of non-reactive sp<sup>2</sup> carbon atoms to be assembled at the surface. The effectiveness of the nanostructure closure was seen in Figure 2 where atoms have been coloured by coordination, revealing low numbers of reactive edges. We are currently exploring how this could confer oxidation

resistance to biochar.

Figure 7 b) and c) shows the simulated HRTEM images of the model produced using the multislice method, for two different orientations. Fringes with similar degrees of curvature (radii of curvature of  $\approx 1$  nm) were found. Some stacking was also observed for Figure 7 c). Significant improvements could be made to such a model to ensure quantitative agreement between the model and the experimental HRTEM, however, we have shown here a preliminary model which is able to demonstrate the main features of the HRTEM of significantly curved fringes, with some stacked fringes and with low amounts of unsaturated sp and sp<sup>3</sup> hybridised carbon.



Figure 7: a) Proposed stacked, fulleroid-like molecular nanostructure model for charcoal with rings coloured blue, white, red and yellow for 5-, 6-, 7- and 8-membered rings respectively (the dashed box in Figure 2 shows the location of this expanded section). c) and d) show simulated HRTEM images of the model structure with 2 nm scale bars shown.

These non-hexagonal rings have been directly imaged in wood charcoal using aberration corrected HRTEM as mentioned in the introduction [19]. We consider these topological defects to be present on a three dimensional graphene network. When considering what to call such a structure we decided not to refer to the structure as *fullerene-like* (fullerenes are defined as closed triply connected carbon networks containing pentagonal and hexagonal rings [48]) instead using the term *fulleroid-like* (fulleroid refers to a closed carbon network containing rings with any number of carbon atoms [48, 52]). A *stacked, fulleroid-*

*like* nanostructure could form an extended porous, curved, *foam-like* nanostructure with enclosed fulleroids and regions of layering.

We have previously shown that large electric fields are present around regions of positive Gaussian curvature in pericondensed aromatic [40]. These regions would be present in such a *stacked*, *fulleroid-like* structure. Understanding how the flexoelectric effect impacts adsorption properties of charcoal and activated carbon could provide considerable insight into their ability to adsorb ions and polar species. Further work is also needed to determine how oxygen is integrated into charcoal's nanostructure but its presence in all higher ions desorbed from charcoal indicate it is a part of the *stacked*, *fulleroid-like* nanostructure.

### 5 Conclusions

A highly crosslinked gasification charcoal was prepared allowing for the analysis of the nanostructure using a variety of techniques. High resolution mass spectrometry using LDI-TOF and ESI 9.4 T FT-ICR MS was able to resolve the m/z 701 ion near the m/z region of buckminsterfullerene as a breakdown product of pyrolysis, which was removed during reduction in the gasifier. The lack of C<sub>60</sub> and C<sub>70</sub> was explained based on the gas phase formation mechanism for fullerenes where small magic number fullerenes are only produced under non-equilibrium high-energy conditions. In a material originally containing magic number fullerenes, fullerene arc-carbon, heating in the solid-state led to coalescence of reactive highly curved structures into polymerised foam-like nanostructures. Using higher laser fluences in LDI-TOF MS a large distribution of ions separated by m/z 24 (C<sub>2</sub>) was observed for charcoal with integrated oxygen, which matched that found for thermally treated fullerene arc-carbon with oxygen becoming intercalated. We have previously found ablation of microcrystalline graphite under the same conditions produced no such distribution suggesting the distribution arises from a shared curved nanostructure. High resolution transmission electron microscopy indicates significantly curved fringes. Raman spectroscopy revealed ring breathing modes up shifted from the D band, which indicate strained enclosed pentagonal ring breathing modes. Using reactive molecular dynamics simulations possible carbon nanostructures were discussed with a stacked, fulleroid-like model being proposed.

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