# Reactivity of polycyclic aromatic hydrocarbon radicals: implications for soot formation

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#### Abstract

This paper presents a systematic study of the reactivity of polycyclic aromatic hydrocarbons (PAH), identifying crosslinks that permit the combination of physical  $\pi$ -stacking interactions and covalent bonding. Dispersion corrected hybrid density functional theory was used to identify the location of reactive sites on PAHs using the average local ionisation potential. The bond energies formed between these various reactive sites were then computed.  $σ$ -radicals were found to be the most reactive, forming bonds with other radicals and some reactive closed shell edge types. Partially saturated rim-based pentagonal rings were found to form localised  $\pi$ -radicals with high reactivity. This site, in addition to resonantly stabilised  $\pi$ -radicals, was found to be capable of bonding and stacking, which is explored for a variety of larger species. Localised  $\pi$ -radicals, in particular, were found to form strongly bound stacked complexes indicating a potentially important role in soot formation.



#### **Highlights**

- The reactivity of aromatic soot precursors are studied.
- Covalent bond energies between aromatic soot precursors are calculated.
- Molecular structures combining physical interactions and covalent bonds are shown.

## **Contents**



### <span id="page-4-0"></span>1 Introduction

Soot emitted from human activities leads to respiratory disease, contributes to warming the atmosphere and increasing ice loss  $[3, 27, 34]$  $[3, 27, 34]$  $[3, 27, 34]$  $[3, 27, 34]$  $[3, 27, 34]$ . Significant research efforts to understand the mechanisms responsible for the formation of soot are ongoing. The critical stage of soot formation (and the least well understood) is the inception process, in which gas phase aromatics form condensed clusters, resulting in carbonaceous nanoparticles [\[58\]](#page-24-0).

Reactive aromatic species have long been thought to contribute to soot inception [\[18,](#page-20-0) [42\]](#page-22-1). These species can be grouped into either open shell radical species or closed shell aromatic species. The former are significantly more reactive but are present in lower concentrations, and the latter less reactive and are present in higher concentrations. Therefore, reactions between radical species and closed shell species are often the focus of soot mechanisms. Beginning with the radicals, the most well studied reactive species, σ-radicals, are generated by hydrogen being abstracted from the rim of aromatics by collision with gas phase radicals [\[19\]](#page-20-1). This provides the reactive site for acetylene addition and the extension of the aromatic network through the well known hydrogen abstraction acetylene addition (HACA) growth mechanism [\[12,](#page-20-2) [13\]](#page-20-3). Radicals can also arise in the  $\pi$ -bonding network if there is an odd number of  $\pi$ -electrons. These  $\pi$ -radicals can be stabilised due to delocalisation, providing long-lived resonantly stabilised radicals (RSR), although these have reduced reactivity. They are critical for the formation of the first aromatic ring [\[16\]](#page-20-4) and have long been suggested to be present in aromatic species, indicated by the odd-numbered carbon species measured using flame mass spectrometry [\[22\]](#page-21-1). High-resolution atomic force microscopy (HR-AFM) and photoionisation mass spectrometry (PIMS) have also recently confirmed the presence of RSR [\[21,](#page-21-2) [44\]](#page-23-0). It was suggested that these react with PAH  $\sigma$ -radicals leading to a chemical polymerisation that did not require subsequent hydrogen abstraction [\[21\]](#page-21-2). However, Keller et al. [\[22\]](#page-21-1) found that for PAH with molecular mass greater than 400 Da odd-numbered carbon fragments in mass spectrometry were of similar concentration to even-numbered carbon fragments (more likely to be closed shell), suggesting that the growth mechanism does not favour reactions with RSR. Evidence for another  $\pi$ -radical has recently been found that arises from a partially protonated rim-based pentagonal ring, which has unknown reactivity [\[44\]](#page-23-0).

The aromatic  $\sigma$ - and  $\pi$ -radicals have also been suggested to react with closed shell species that possess significantly electrophilic sites such as double or triple bonds. This is well illustrated by the reaction between the aromatic  $\sigma$ -radical with the acetylene triple bond, which is critical for the HACA mechanism, but is not rapid enough to explain soot formation [\[12\]](#page-20-2). Benzene and high symmetry aromatic species are strongly aromatically stabilised leading to aromatic bonds with low reactivity compared with double bonded carbon. However, in many low symmetry polycyclic aromatic hydrocarbons, some rings possess a reduced aromaticity. Well known examples are the 9,10-free edge of phenanthrene and pyrene, which is more vulnerable to electrophilic or free radical attack [\[6\]](#page-19-2). Rim-based pentagonal rings have also been observed and shown to be thermally stable [\[20,](#page-20-5) [49\]](#page-23-1). Five-membered rings are not aromatically stabilised and thus provide a free edge with double-bond character and significant reactivity. These low aromaticity free edges have been suggested to react with carbon  $\sigma$ -radicals, forming nanoparticles in the flame through the aromatic aliphatically linked hydrocarbon (AALH) mechanism [\[10,](#page-19-3) [56\]](#page-24-1).

Evidence for this growth mode has been shown by mass spectrometry of benzene-oxygen flames where high radical concentrations allow for significant carbon  $\sigma$ -radical concentrations [\[56\]](#page-24-1). Partial integration of the pentagonal ring within the hexagonal network has been directly imaged [\[7\]](#page-19-4) and provides edges with greater reactivity, with HACA growth on the five-membered bay site found to proceed rapidly [\[40\]](#page-22-2). Fully integrated pentagonal rings curve the aromatic network and have been experimentally observed [\[26,](#page-21-3) [32\]](#page-22-3). This curvature leads to an electric polarisation of the  $\pi$ -electrons [\[31\]](#page-21-4) and reduced aromatic-ity [\[11\]](#page-20-6) due to reduced  $\pi$ -overlap on pyramidalised carbon atoms. This has been shown to increase the edge oxidation reactivity [\[41,](#page-22-4) [50\]](#page-23-2) and speed up HACA growth on an armchair edge of hexagonal rings [\[40\]](#page-22-2). This curvature integrates a permanent dipole moment into the molecules, suggesting strong dipole-ion interactions are possible, which have been proposed as a possible explanation of the electrical aspects of soot formation [\[32\]](#page-22-3). The presence of these reactive aromatics leads to a number of questions: how does the reactivity of different sites compare? How thermally stable are the crosslinks formed between these sites?

The proposed chemical inception mechanisms involving reactive aromatics are not fast enough to explain the experimental observations [\[58\]](#page-24-0). Physical condensation, however, could rapid enough to explain soot formation if sufficient intermolecular interactions existed between the fragments [\[12\]](#page-20-2). Direct evidence for physical dimerisation has been found using photoionisation mass spectrometry of species sampled from the flame, revealing a series of peaked distributions separated by approximately 500 Da (without any change in the C/H ratio, which would imply a chemical reaction) [\[5,](#page-19-5) [17\]](#page-20-7). Laser induced fluorescence (LIF) experiments have provided evidence for  $\pi$ -stacked aromatics that are able to form an excited eximer state [\[38\]](#page-22-5). Time resolved LIF has shown this signal to be long lived, providing further evidence for the stacked eximer hypothesis [\[36,](#page-22-6) [45\]](#page-23-3). However, we have shown that van der Waals or physical dispersion interactions are not sufficient to allow the 500 Da aromatic species (as commonly found in flames) to cluster at temperatures in the flame where soot forms, 1500–2000 K [\[53\]](#page-23-4). Previous authors have suggested a combination of covalent bonding and physical interactions to explain the rapid condensation [\[18,](#page-20-0) [24,](#page-21-5) [37\]](#page-22-7). Experimentally, fluorescence within the flame suggested excimer states from stacked aromatic molecule [\[38\]](#page-22-5). Also recent electron spin resonance study found a significant reduction in concentration and change in the radical character when soot growth begins indicting a significant number of radical termination reactions occurring at soot inception [\[57\]](#page-24-2). The authors and others suggested a combination of physical stacking interactions and covalent bonding between  $\pi$ -radicals such as pancake bonds (multicentre  $\pi$ -bonds) or biradicals to provide the significant intermolecular interactions required to hold PAH within clusters at flame temperatures [\[9\]](#page-19-6). This poses a further question: can a combination of physical interaction and covalent bonding explain the rapid growth of soot?

In this paper, we seek to locate reactive sites and compare their reactivity using the average local ionisation potential. The energies of the bonds formed between the different site types is then systematically calculated revealing possible covalent bonding options. Finally, physically and covalently bonded structures are computed revealing the possibility of  $\pi$ -radical pancake- and rim-bonded complexes.

### <span id="page-6-0"></span>2 Methodology

For determining the minima and transition state geometries we employed a range of electronic structure methods computed in the Gaussian 09 package [\[14\]](#page-20-8). For all geometries, frequency calculations were performed to confirm the minima or transition state and also to compute the zero-point energy correction to the electronic structures. We have previously benchmarked a range of hybrid density functional theory methods against experimental and wavefunction methods [\[19\]](#page-20-1). We found geometry optimisations at the B3LYP/6-31G(d,p) level of theory and single point calculations at the M06-2X/cc-pVTZ level of theory provide energetics within chemical accuracy for hydrogen abstractions (<1 kcal/mol) [\[19\]](#page-20-1).

For the aromatic complexes we have previously found that dispersion corrected B97 hybrid density functional theory performs well for geometries and electrostatics describing the dipole moment of curved arenes [\[15,](#page-20-9) [30\]](#page-21-6). A variety of geometry optimisations were performed for each edge couple to determine the lowest energy isomer. The B97D/ccpVTZ level of theory was used in order to ensure minimal basis set superposition errors (<2 kcal/mol) as the bonded and stacked geometries were unable to be corrected for this error (counterpoise corrected). The bond dissociation energies were calculated from a homolytic bond cleavage using single point energies calculated using the dispersion corrected hybrid density functional M06-2X-D3/cc-pVTZ, which uses the same dispersion correction that was added to the B97D functional but provides significantly better bond energies, as will be detailed in the next section. We compared the van der Waals complexes with our reference SAPT(DFT) calculations [\[33,](#page-22-8) [53\]](#page-23-4) and found the DFT calculations to overestimate the binding energy by −6±1 kcal/mol, which will be further discussed in the results section.

The average local ionisation potential is used to locate and compare reactive sites on aromatic species to electrophilic attack by, for example, carbon radicals [\[46\]](#page-23-5) and is computed as,

$$
\bar{I}(\mathbf{r}) = \sum_{i}^{N_{occ}} \frac{\rho_i(\mathbf{r}) |\varepsilon_i|}{\rho_{tot}(\mathbf{r})},
$$
\n(1)

where  $\rho_i$  is the electron density of the *i*-th molecular orbital at the point **r**,  $\varepsilon_i$  is the orbital energy and  $\rho_{tot}(\mathbf{r})$  is the total electron density at the point **r** where the sum is over the  $N_{\text{occ}}$  occupied molecule orbitals.  $\bar{I}(\mathbf{r})$  is then a local measure of the energy required to ionise an electron from a certain point in space and therefore a lower value indicates a higher reactivity to electrophiles such as radicals. The Fukui function  $f^-(\mathbf{r})$  has also been used to compute the reactivity of sites towards hydrocarbon radicals in the context of soot [\[59\]](#page-24-3).  $f^-(\mathbf{r})$  requires that the electron density of the ionised molecule be computed and the difference taken with the molecule in its standard state. This is often approximated with the highest occupied molecular orbital (HOMO), but this is not possible for larger aromatics with many low lying occupied molecular orbitals that could also contribute, e.g. HOMO–1, HOMO–2 etc. Therefore, two calculations would be required to compute *f*<sup>−</sup>(**r**). The average local ionisation potential and the Fukui functions are formally related in the local density approximation as  $\bar{I}(\mathbf{r}) = \sum_i |\varepsilon_i| f_i^-(\mathbf{r})$ , where the index is over the *i*th occupied molecular orbitals, indicating they will both provide the most electrophilic site [\[51\]](#page-23-6). The advantage of using  $\bar{I}(\mathbf{r})$  is that only a single electronic structure is required as it considers ionisation from all of the occupied MO while also providing a convenient comparison across our molecular series. Multiwfn 3.3.9 was used to compute the average local ionisation potential on the isosurface 0.002 atomic units (a.u.) of the electron density (near the van der Waals surface) computed from the electronic structure calculations [\[29\]](#page-21-7). We also prepared spin population isosurfaces for the resonantly stabilised radicals using Multiwfn.

In the context of aromatic species,  $\bar{I}(\mathbf{r})$  has been found to accurately predict the sites most reactive to electrophilic attack of hydrogen radicals [\[4\]](#page-19-7). To further confirm this we demonstrated that the enthalpy of hydrogenation at edges of closed shell species correlates well with the minima near the edge carbon atom that was protonated,  $min_r[\bar{I}(r)]$  (see Figure [S1](#page-18-0)) in the Appendix). We can then suggest that  $\min_r[\bar{I}(r)]$  is appropriate for comparison between the different radical and closed shell species' reactivity. For ease of reading, the minimum in the average local ionisation of the site min<sub>*r*</sub> [ $\bar{I}$ (**r**)] will be referred to as  $\bar{I}_S$ .

### <span id="page-7-0"></span>3 Results and discussion

#### <span id="page-7-1"></span>3.1 Reactivity of aromatic edges

Figure [1](#page-8-0) shows the average local ionisation potential calculated on the molecular surface for aromatic species ordered by their  $\bar{I}_s$ . This serves two main purposes: classification and ordering of the different radicals and closed shell species, as well as an understanding of the location of the potential reactive sites that are capable of crosslinking reactions, which will be explored in the next section.

We will begin with closed shell species as they will introduce the concepts of reactivity and aromaticity using the  $\bar{I}_S$  metric. Partially embedded cyclopentaphenathrene-type pentagonal rings D) possess the lowest  $\bar{I}_s$  site with values  $\bar{I}_s = 7.28 - 7.41$  eV. The  $\bar{I}_s$ are concentrated on the aromatic face on the pentagonal's edge, with the pentagonal ring being partially embedded with three bond sharing hexagonal rings. This non-aromatic site has been observed in HR-AFM and is an intermediate towards completely curved species [\[44\]](#page-23-0). We also explored other partially embedded pentagonal rings that had also been observed, such as fluoranthene or benzo[ghi]pyrene type, but these were found to have  $\bar{I}_S$  matching that of low-aromaticity free edges shown in the scheme below.



This indicates that aromaticity is lowest for the cyclopentaphenathrene type edges providing the most reactive configuration of a partially embedded pentagonal ring. This could explain why this site type is most often seen to be methylated, fluorene-like, having two hydrogens bonded to the edge [\[7\]](#page-19-4).

<span id="page-8-0"></span>

Figure 1: *The average localised ionisation potential is plotted on the molecular surface produced from the electron density at the isovalue*  $\rho = 0.002$  *a.u.* (B97D/6-*311G(d,p)) for a variety of aromatic species. Overlayed on these molecular surfaces are one the Kekulé structures.*

Rim-based pentagonal rings E) possess values of  $\bar{I}_s = 7.45 - 7.64$  eV. The minima is concentrated on the edge carbon atoms of the pentagonal ring. In this case only two neighbouring bonds are shared with the hexagonal aromatic rings and due to the antiaromatic pentagonal ring the  $\bar{I}$  indicates a free edge with a double-bond character. There is little effect due to system size and many of these sites can be present on a single aromatic. Experimentally this was found to be a common edge type [\[7\]](#page-19-4).

Low aromaticity free edges F) possess values of  $\bar{I}_s = 7.91 - 8.08$  eV. These arise in hexagonal aromatic networks due to the topology of the network. Clar provided a framework for describing these low-aromaticity edges [\[2,](#page-19-8) [6\]](#page-19-2). The maximum number of sextets are placed within the network with double bonds placed elsewhere. The symmetrically equivalent arrangements of sextets (denoted by a circle within the ring) and double bonds are then considered to be in resonance in more modern formulations [\[48\]](#page-23-7). The edge sites where double bonds are prevalent in the resonance structures are found to be reactive. These Clar descriptions match with the sites with low  $\bar{I}_S$  and with the reactivity trends for these species  $[6]$  and has been used to describe PAH growth  $[28]$ . These Clar descriptions match with the sites with low  $\bar{I}_s$  and with the reactivity trends for these species [\[6\]](#page-19-2) and has been used to describe PAH growth [\[28\]](#page-21-8). Some examples are shown in the scheme below.



Naphthalene contains a single sextet that is in resonance (otherwise known as a travelling sextet). This allows for the free edges to possess a double-bond character with an increased reactivity. Phenanthrene is an example of a structure with a single Clar structure with two sextets leaving the 9,10-free edge with a significant double bond character and higher reactivity than the rings with the sextets [\[6\]](#page-19-2). Corannulene can have two sextets and five resonance structures [\[2\]](#page-19-8). This provides low-aromaticity free edges with double-bond character. Corannulene has an added reduction in aromaticity due to the pyramidalisation of the carbon network, as shown by others [\[11\]](#page-20-6). This might explain the lower  $\bar{I}_S$  compared with phenanthrene or naphthalene found for corannulene. Another cause of low aromaticity edges was methylation of hexagonal aromatic edges site. In the Clar framework this is due to the reduction in the number of sextets that can be placed. Finally, high symmetry PAH G) such as benzene or coronene are found to possess the highest  $\bar{I}_S$  value of all of the closed shell edges with values  $\bar{I}_s = 8.12 - 8.21$  eV. The symmetry of these species, being close to circular, provides highly pericondensed networks with high aromaticity and no low-aromaticity free edges.

Turning now to the radical, the site of lowest  $\bar{I}_S$  was found to be the  $\sigma$ -radical A) with minimum values from 7.00–7.17 eV. The reactive site minima lies on the edge of the aromatic, parallel to the carbon–hydrogen bonds. Little difference was found between the rim-based, pentagon-based  $\sigma$ -radical compared with the hexagonal ring-based one. This reactive site is known to react with acetylene readily and drives the growth of the aromatic network [\[12\]](#page-20-2). The next lowest  $\bar{I}_s$  is the localised  $\pi$ -radical **B**) with minimum values from 7.51–7.61 eV. In both cases the minima lies on the rim-based pentagonal carbon atom with a single hydrogen attached. Unlike the  $\sigma$ -radical, the  $\bar{I}_S$  is on the face of the aromatic plane. There are also regions of low  $\bar{I}$  around the rim, which have similar values to the low aromaticity free edges, indicating this site can modify the other edge sites on the same molecule. This will be further discussed later in the text.

In order to determine the localisation of this radical, two partially saturated pentagonal rings were added to the seven-ring aromatic coronene (Figure [2a\)](#page-10-1). The triplet with two free electrons (a biradical) was found to be significantly more stable (−24.3 kcal/mol) compared to the singlet. If the radical was able to delocalise, the diradical would provide a stable singlet spin configuration across the aromatic network. However, as the triplet dominates we can confirm that the radical is indeed localised to the pentagonal ring and that multiple localised  $\pi$ -radicals could be formed on a single aromatic with the low  $\bar{I}_S$  = 7.54 *eV* preserved as shown in Figure [2a.](#page-10-1)

A Clar analysis provides insight into why the radical is localised (Figure [2b\)](#page-10-1). Considering the average local ionisation potential of 1,2-dihydroacenaphthylen-1-yl, four major Clar

<span id="page-10-1"></span>

(a) *Mechanism for forming two B) edge types showing the two independent localised*  $\pi$ *-radicals.* 



(b) *Clar analysis of 1,2-dihydroacenaphthylen-1-yl.*

Figure 2: ¯*I*(r) *surface plotted as in Figure [1](#page-8-0) for the partially hydrogenated edges of type B).*

structures can be drawn. The first two resonance structures have a sextet on the hexagonal rings with the radical localised to the pentagonal ring. The third and fourth resonance structures have the radical on the aromatic ring, *i.e.* the  $\beta$ - and  $\gamma$ -position from the pentagonal site. Similar patterns in the  $\bar{I}(\mathbf{r})$  are also found for the larger species in Figure [2a,](#page-10-1) where the radical site only appears to be delocalised to the  $\beta$ -position relative to the pentagonal site. It can therefore be suggested that the radical is localised in order to keep the high aromaticity of the six-membered ring network.

The radical with the highest  $\bar{I}_s$  was found to be the resonantly stabilised  $\pi$ -radical RSR C) with  $\bar{I}_s = 7.88 - 8.12$  eV. These arise from an odd number of  $\pi$ -electrons in an aromatic network providing a radical stabilised by delocalisation [\[23\]](#page-21-9). The smallest species, indenyl, was found to have the lowest  $\bar{I}_s$  value (7.88 eV) concentrated on the surface of the pentagonal ring, while for increasing size the value drops to 8.12 eV and is concentrated on the face of the aromatic on the zig zag edge sites. We will explore later whether this reduction impacts the reactivity of the RSR as they enlarge and the radical becomes more resonantly stabilised.

The reactive edges have been described and grouped. Importantly, the location of reactive sites has been found allowing for potential crosslinks between these reactive sites to be explored on a smaller subset of edges in the following section.

#### <span id="page-10-0"></span>3.2 Crosslinking reactions between reactive aromatics

In order to compare the reactivity of a variety of edge types, we have chosen to consider only the first bond formed between a series of small species. There are a number of reasons for this. Firstly, the formation of the first bond will provide insight into the edge site reactivity and provide further evidence for the reactivity ordering predicted from  $\bar{I}_s$ .

Secondly, the first bond is important for the crosslinking reaction as the reaction is not entropically favoured due to the reduction in number of species, and therefore the energy of the first bond directly corresponds to the likelihood of the complex fragmenting. Thirdly, molecular beam mass spectrometry shows that the clustering process, in low temperature flames, does not involve crosslinking reactions with dehydrogenation but a constant C/H ratio [\[5,](#page-19-5) [17\]](#page-20-7). Finally, we did not include any aliphatic bridges with methyl or longer aliphatics that have recently been observed in specific inverse diffusion flames [\[1\]](#page-19-9). These would modify the C/H ratio and therefore would represent a different growth mode than what we are focusing on in this study. Furthermore, in premixed flames these aliphatic bridges would be prone to radical-induced fragmentation, just as the fuel is. Therefore, we restricted our search to crosslinks that do not modify the C/H ratio and have been seen in aromatics directly imaged using HR-AFM [\[7\]](#page-19-4).

Figure [3](#page-12-0) shows the matrix of bond energies between the species containing different edge types ordered by their bond energy with the  $\sigma$ -radical. A general trend was found where edge sites with low  $\bar{I}_s$  produced more stable bonds compared to those with high  $\bar{I}_s$  (as seen in the inset graph). However, steric effects also contribute to these bond energies leading to slightly different orderings compared with  $\bar{I}_S$  values. We did find that the site with the lowest  $\bar{I}_s$ , or its neighbour, was always found to be the most reactive. Bond energies can be compared to the bond enthalpy benchmark values (NIST thermodynamic database) as the thermal correction is minimal at  $\langle 1 \text{ kcal/mol}$ . The biphenyl C–C bond has a benchmark value of −117.6 kcal/mol [\[55\]](#page-24-4); using our methodology provided a value of −119.4 kcal/mol, showing a slight overestimation of the bond energy (+1.5%). Therefore, these values should be viewed within this error to provide a comparative look at the reactivity of the different edge sites. To consider the flame stability of bonded species, a comparison with other species is helpful. The thermal energy necessary for homolytic bond cleavage can be approximated as 6*RT* (3/2RT for the translational and 3/2RT for rotational degrees of freedom for each species), which is 17 kcal/mol at 1500 K and 24 kcal/mol at 2000 K. However, to be persistently stable at flame temperature, larger bond energies have been found to be required. The primary fragmentation pathway for fuels is radical induced fragmentation, such as  $\beta$ -scission, where the  $\beta$  site can have bond energies of 30 – 40 kcal/mol that readily break at flame temperatures. For the physical dimerisation of flat PAH, a full statistical approach revealed that a binding energy of  $<-40$  kcal/mol was required for clustering above 1500 K [\[58\]](#page-24-0). Single bonds C–H and C–C in benzene and aliphatics have bond dissociation energies of −113 and −90 kcal/mol respectively, and require high temperatures for unimolecular decomposition (>2000 K). We therefore consider anything below  $-40$  kcal/mol to be of interest for soot formation and anything above −10 kcal/mol is not considered for forming any thermally stable bond.

The most significant bond energies are formed are with  $\sigma$ -radicals A), as shown by the first and second row of the grid in Figure [3.](#page-12-0) The bond energies follow the ordering suggested from  $\bar{I}(\mathbf{r})$  for the radicals. The strongest bonds are formed with other sigma radicals with  $\langle -120 \text{ kcal/mol}$ . These bond energies are lower than single bond energies (−90 kcal/mol), indicating that these species are significantly conjugated and stabilised by delocalisation. We have previously demonstrated that crosslinking between two aromatics leads to a small drop in the band gap supporting such a limited conjugation [\[35\]](#page-22-9). The bond energy between two pentagonal ring  $\sigma$ -radicals Ai) is particularly interesting as the minimum energy isomer is planar providing the greatest degree of delocalisation and

<span id="page-12-0"></span>

Figure 3: *The bond energies (kcal/mol) between different species representing edge types shown in Figure [1.](#page-8-0) Inset graph shows the correlation between the bond energies and the*  $\bar{I}_S$  *and a selection of geometries.* 

the highest bond energy. This crosslink being planar would also allow for more effective van der Waals clustering and have been experimentally observed [\[44\]](#page-23-0). These σ-radical crosslinks would be long-lived in the flame based on the calculated bond energies. However, the low concentration of radicals makes this mechanism possible only in systems with many aromatic radicals, such as in the pyrolysis of PAHs [\[39\]](#page-22-10). The next strongest crosslink with A) is the localised  $\pi$ -radicals B) with bond energies indicating single C–C bond formation between these edge types. Subsequent hydrogen abstraction could provide an  $Ai$ ) +  $Ai$ ) bond type, which as mentioned is favourable for clustering. The partially embedded pentagonal ring **D**) also reacts readily with this site to form bonds of strong energy ( $\lt -73$  kcal/mol). The concentration of this site in flame aromatics is unknown and will likely be considerable given its possible equilibrium with hydrogen radicals and the methylated version as shown in Figure [2a.](#page-10-1) Resonantly stabilised radicals follow with bond energies of  $< -61$  kcal/mol. One question surrounding this crosslink is how the bond energy varies with the size of the RSR, as mentioned by Homann [\[22\]](#page-21-1). Computing the bond energies of site Aii) with the small RSR indenyl Cii) through to a 10-ring RSR species (shown in Figure [5\)](#page-16-0) showed no decrease in the bond energy with system size with a mean value of  $-65.9 \pm 2.8$  kcal/mol. This indicates that while the bond energies with RSR are lower than a single bond, they do not vary greatly and that upon reaction with a  $\sigma$ -radical,

the  $\pi$ -radical is easily localised to form a bond. Calculations have shown that hydrogen can be easily lost from the RSR, reforming the radical and leading to potential chain reactions [\[21\]](#page-21-2). However, the concentration of RSR in the flame needs to be determined to see how common these reactions could be. The presence of odd-numbered carbon species is not sufficient to established a RSR due to the hydrogenation of the edge, as has been recently demonstrated in HR-AFM; i.e., species with an odd number of carbon atoms that are protonated can become closed shell species [\[7\]](#page-19-4).

The reaction between a rim-based pentagonal ring E) and a  $\sigma$ -radical A) is the first step detailed in the AALH mechanism [\[10,](#page-19-3) [56\]](#page-24-1) and provides bond energies  $\sim$  −50 kcal/mol. For curved aromatics with low-aromaticity free edges Fi) we found that bonds formed with A) species were at the threshold of stability at flame temperatures, while crosslinks with phenanthrene Fii) and naphthalene Fiii) did not provide stable bonds at flame temperatures. The mechanisms including  $\sigma$ -radicals often require crosslinking followed by a hydrogen abstraction or loss to transform a weakened bonded structure into a  $A$ ) +  $A$ ) type crosslink. These mechanisms require a large number of  $\sigma$ -radicals and hydrogen radicals to allow for chemical polymerisations [\[58\]](#page-24-0). However, in most atmospheric flames, clustering occurs with a constant C/H ratio indicating a lack of chemical polymerisation [\[17\]](#page-20-7). While this is the case, crosslinking of  $\sigma$ -radicals providing strong bonds with energies  $<-40$  kcal/mol is found to be possible with edge types Fi), E), C), D) and B).

After the  $\sigma$ -radicals, the  $\pi$ -radicals are found to also form strong bonds with themselves and some high reactivity closed shell species. Localised  $\pi$ -radicals on rim-based pentagons B), row three of the table in Figure [3,](#page-12-0) show the next highest reactivity, forming bonds with itself of energy −62 kcal/mol. Bonds with D) type edges are found to provide bonds below −40 kcal/mol while bonds with C) and E) type edges are above this threshold. Of particular interest are the bonds formed with the rim-based pentagonal rings E) as these closed shell species are prevalent in the flame. The RSR radicals are found to form weak bonds with themselves and with the partially embedded pentagonal rings and no stable bonds with the rim-based pentagonal rings E) were found to be possible. The exception is the small indenyl species that forms a strong bond with itself,  $Cii$ ) +  $Cii$ ), showing that reactivity between RSR decreases with the size of the species as indicated from the  $\bar{I}_s$ .

Finally, rim-based pentagonal rings E) were found to form a weak bond with themselves, which is unlikely to be of interest. This [2+2] pericyclic reaction is not thermally allowed and requires optical excitation to form due to spin restrictions [\[43\]](#page-23-8). We have recently measured and calculated the band gap to be between  $3.6-3.7$  eV ( $\approx$ 350 nm deep UV) [\[35\]](#page-22-9). These high energy photons are not prevalent in flames, indicating this bond is unlikely to form in a flame. The bond energy is also very low. Santos *et al.* have previously experimentally and computationally shown that the energetics of acenaphthylene dimerisation are not possible above 550 K [\[43\]](#page-23-8), further supporting the weak nature of these bonds.

From this systematic study, only  $\sigma$ -radicals A) and localised  $\pi$ -radicals B) are found to be capable of forming strong enough bonds to be long-lived in the flame. However, the  $\pi$ -radicals being able to bond on the aromatic face hold the possibility of combined physically held and covalently bound configurations, which is explored next.

#### <span id="page-14-0"></span>3.3 Combining  $\pi$ -stacking and covalent bonding

As mentioned, physical dispersion forces are insufficient to stably bind clusters of small aromatic species at flame temperatures [\[53,](#page-23-4) [58\]](#page-24-0) and so covalently stabilised  $\pi$ -stacked structures are sought. The  $\sigma$ -radicals are unable to accomplish this feat as the reactive radical site points out from the aromatic ring, making bonding only achievable parallel to the aromatic planes, which will not allow for  $\pi$ -stacking. The thick-bordered entries in the matrix of Figure [3](#page-12-0) show the reactive edges were able to covalently bond as well as  $\pi$ -stack. These binding energies will be compared with van der Waals interactions and the unstrained covalent bonds, i.e. the energies of the covalent bond when there is no stacking from Figure [3.](#page-12-0)

Figure [4](#page-15-0) shows the binding energy and molecular geometries of the reactive dimerisations for their average monomer mass. For comparison, physical dimerisation due to dispersion or van der Waals interactions have also been computed (black dashed lines). We have also shown our benchmark SAPT(DFT) calculation from previous works, which can accurately predict the virial coefficient of benzene and the exfoliation energy of graphite (solid black lines) [\[52\]](#page-23-9). From this we can see an overbinding of nonbonded planar aromatics by the hybrid density functional method M06-2X-D3, which we have previously seen for these empirically corrected DFT methods [\[30\]](#page-21-6). Given that this overbinding is linear in molecular mass, we can compare the enhancement  $\Delta E = E_{VdW} - E_{C+VdW}$  due to any covalent interactions  $E_{C+VdW}$  compared to the van der Waals dimers  $E_{VdW}$  calculated with the dispersion corrected DFT for an equivalent mass monomer (blue and green arrows in Figure [4\)](#page-15-0).

Figure [4a](#page-15-0) insets shows the single bonds formed between the species and, in the case of the localised  $\pi$ -bonds **B**) as referred to in Figure [1,](#page-8-0) how rotation about the single bond does not impact the binding energy of these species. Figure [4b](#page-15-0) shows how as the molecular mass increases this binding is enhanced by van der Waals interactions, either by maintaining a rim-based bond across the fragments, in the case of B) type edges, or through multicentre  $\pi$ -bonds, in the case of RSR C).

The localised  $\pi$ -radicals **B**) are found to most strongly bind with themselves **B**) having an average enhancement of some  $\Delta E = 50 \pm 4$  kcal/mol. This is allowed for by the minimal strain in going from the tetrahedrally oriented species to the  $\pi$ -stacked configuration due to the ability of the saturated pentagonal ring hydrogens to be placed far from each other. These edges also form strong bonds with partially embedded pentagonal rings D) with an enhancement of  $\Delta E = 35 \pm 6$  kcal/mol. For the reaction of **B**) with the rim-based pentagonal ring **E**) we found a minor enhancement of  $\Delta E = 8 \pm 4$  kcal/mol. This small enhancement might be explained by the low initial bond energy in the unstacked configuration. When stacked the added strain provides reduced bond energies.

Resonantly stabilised  $\pi$ -radicals are able to form multicentre  $\pi$ -bonds – a so-called "pancake bond" – and have been suggested to stabilise clustering aromatics [\[47,](#page-23-10) [57\]](#page-24-2). We found enhancements of  $\Delta E = 6 \pm 3$  kcal/mol compared with non-bonded configurations indicating a minor increase in binding energy. This is comparable to binding energies previously computed [\[23\]](#page-21-9). We found that for the most stable isomers the multicentre bond was not formed across the entire molecule but was partially localised to the most triangulene-like part of the fragment. This is illustrated in Figure [5](#page-16-0) where we have plotted the orientation,

<span id="page-15-0"></span>

Figure 4: *a) Binding energy (kcal/mol) as a function of molecular mass of the monomers (Da) is shown for different covalently bonded structures from Figure [3.](#page-12-0) b) Bonded and stacked geometries are found for enlarged monomers. c) Binding* energy solely from van der Waals interactions  $E_{VdW}$  is also shown for compar*ison with the method chosen compared with our reference calculations from Totton et al. [\[54\]](#page-23-11) d) The bonding enhancement compared with the non-bonded case* ∆*E is also shown.*

spin population and HOMO of four of the RSR molecules. From the pancake bonding molecular orbital we can see that the multicentre bond is concentrated on one end of the dimer. This is geometrically manifested in the optimised structures where the region of bonding possesses a lower interlayer distance of 3.38 Å compared with the opposing edge of 3.7 Å. This can be understood from examining the spin population in Figure [5,](#page-16-0) indicating that the  $\pi$ -radical is concentrated on the most triangulene-like region of the molecule from where it forms the multicentre bond.

<span id="page-16-0"></span>

Figure 5: *a-d) Dimer geometries, spin population at iso* = 0.003 *a.u. and HOMO at iso* = 0.025 *a.u. for a selection of RSRs showing the localisation of the radical on the most triangulene-like edge.*

Considering the implications for soot formation. we can discuss whether these covalently bound complexes could allow for an explanation for the fluorescence signal interpreted as stacked aromatic eximer state [ $38$ ]. Recent calculations by Krueger and Blanquart of  $sp<sup>3</sup>$ bonded aromatics through aliphatics that are stacked showed that these long lived eximer states can indeed exist in stacked and bonded complexes [\[25\]](#page-21-10). Therefore, we anticipate that the localised  $\pi$ -radical **B**) bonded complexes will possess a fluorescence signal. For the pancake bonded complexes, however, it is unclear whether an eximer state would be bound given that the  $\pi \to \pi^*$  transition would be expected to lead to a breaking of the pancake bond. Pancake bonds are also highly phase dependent, breaking if they are rotated by 30° [\[8\]](#page-19-10), further questioning whether they could be stable at flame temperatures.

These considerations provide some possible mechanisms for the covalently stabilised soot nucleation mechanism with high sticking coefficients that have been found to be computationally required for modelling soot nucleation [\[18,](#page-20-0) [24,](#page-21-5) [37\]](#page-22-7). In particular, the ability of multiple localised  $\pi$ -radicals to be formed on a single aromatic suggests that the rim-based

pentagonal ring, in partial equilibrium with hydrogen radicals in the flame (as shown in Figure [2a\)](#page-10-1), could provide a constant supply of reactive sites to polymerise soot formation.

### <span id="page-17-0"></span>4 Conclusions

The average local ionisation potential was used to explore the reactivity of a variety of edge types that have been recently directly imaged in aromatic soot precursors. The radical sites and closed shell edge types were characterised with seven reactive edge types, which were detailed and compared. The location of reactive sites also allowed for a systematic study of the bond energies between different covalently crosslinked reactive edge types. σ-radicals were found to form strong bonds with curved PAH low aromaticity free edges Fi, rim-based pentagonal ring free edges E), resonantly stabilised  $\pi$ -radicals C), partially embedded pentagonal ring edges D) and localised  $\pi$ -radical on a partially saturated rim based pentagonal ring B), indicating the importance of such crosslinks in certain flame types.  $\pi$ -radicals possess lower covalent bond energies, however, they were found to allow for stacked and bonded configurations significantly enhancing the interaction energy as the monomers mass increased. Localised  $\pi$ -radicals **B**) are suggested to be important due to their significant binding energy and the potential to rapidly generate them via protonation of abundant rim-based pentagonal rings. More work is required to determine the thermal stability of these covalently stabilised stacked complexes and to establish the concentration of each edge type within sooting flames. However, we have demonstrated that covalent bonds and stacked configurations are not necessarily mutually exclusive and that strongly bound complexes of interest to soot formation can be formed.

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## <span id="page-17-1"></span>A Appendix

#### <span id="page-17-2"></span>A.1 Hydrogenation

Figure [S1](#page-18-0) plots the reaction enthalpy of hydrogen abstraction and addition to a collection of aromatics that will be explored in this work. For the hydrogen abstraction we do not find any correlation with the carbon rim  $\min[\bar{I}]$  value, which is consistent with what we have found previously [\[19\]](#page-20-1). The enthalpy of hydrogenation, however, strongly depends on

<span id="page-18-0"></span>the carbon rim min $[\bar{I}]$  value to which it is being added to. This indicates that  $\bar{I}$  is accurate in predicting the reactivity of aromatic edges to radical attack over a range of different aromatic species.



Figure S1: *Enthalpy of reaction for i) hydrogen addition (open circles) and ii) hydrogen abstraction (filled circles).*

### <span id="page-19-0"></span>References

- <span id="page-19-9"></span>[1] B. D. Adamson, S. A. Skeen, M. Ahmed, and N. Hansen. Detection of aliphatically bridged multi-core polycyclic aromatic hydrocarbons in sooting flames with atmospheric-sampling high-resolution tandem mass spectrometry. *The Journal of Physical Chemistry A*, 122(48):9338–9349, 2018. [doi:10.1021/acs.jpca.8b08947.](http://dx.doi.org/10.1021/acs.jpca.8b08947)
- <span id="page-19-8"></span>[2] A. T. Balaban and D. J. Klein. Claromatic carbon nanostructures. *The Journal of Physical Chemistry C*, 113(44):19123–19133, 2009.
- <span id="page-19-1"></span>[3] T. C. Bond, S. J. Doherty, D. W. Fahey, P. M. Forster, T. Berntsen, B. J. Deangelo, M. G. Flanner, S. Ghan, B. Kärcher, D. Koch, S. Kinne, Y. Kondo, P. K. Quinn, M. C. Sarofim, M. G. Schultz, M. Schulz, C. Venkataraman, H. Zhang, S. Zhang, N. Bellouin, S. K. Guttikunda, P. K. Hopke, M. Z. Jacobson, J. W. Kaiser, Z. Klimont, U. Lohmann, J. P. Schwarz, D. Shindell, T. Storelvmo, S. G. Warren, and C. S. Zender. Bounding the role of black carbon in the climate system: A scientific assessment. *Journal of Geophysical Research Atmospheres*, 118(11):5380–5552, 2013. [doi:10.1002/jgrd.50171.](http://dx.doi.org/10.1002/jgrd.50171)
- <span id="page-19-7"></span>[4] F. A. Bulat, J. S. Burgess, B. R. Matis, J. W. Baldwin, L. Macaveiu, J. S. Murray, and P. Politzer. Hydrogenation and fluorination of graphene models: Analysis via the average local ionization energy. *The Journal of Physical Chemistry A*, 116(33): 8644–8652, 2012. [doi:10.1021/jp3053604.](http://dx.doi.org/10.1021/jp3053604)
- <span id="page-19-5"></span>[5] F. Carbone, M. R. Canagaratna, A. T. Lambe, J. T. Jayne, D. R. Worsnop, and A. Gomez. Exploratory analysis of a sooting premixed flame via on-line high resolution (APi-TOF) mass spectrometry. *Proceedings of the Combustion Institute*, 37: 1–8, 2018. [doi:10.1016/j.proci.2018.08.020.](http://dx.doi.org/10.1016/j.proci.2018.08.020)
- <span id="page-19-2"></span>[6] E. Clar, B. McAndrew, and M. Zander. The establishment of double bond character in methyl derivatives of phenanthrene, pyrene, chrysene and coronene by nmr. *Tetrahedron*, 23(2):985–993, 1967.
- <span id="page-19-4"></span>[7] M. Commodo, K. Kaiser, G. De Falco, P. Minutolo, F. Schulz, A. D'Anna, and L. Gross. On the early stages of soot formation: Molecular structure elucidation by high-resolution atomic force microscopy. *Combustion and Flame*, 205:154–164, 2019.
- <span id="page-19-10"></span>[8] Z.-H. Cui, H. Lischka, H. Z. Beneberu, and M. Kertesz. Rotational barrier in phenalenyl neutral radical dimer: Separating pancake and van der Waals interactions. *Journal of the American Chemical Society*, 136(15):5539–5542, 2014.
- <span id="page-19-6"></span>[9] A. D'Anna and H. Wang. Personal communication at the 37th International Symposium on Combustion, 2018.
- <span id="page-19-3"></span>[10] A. D'Anna, A. Violi, A. D'Alessio, and A. F. Sarofim. A reaction pathway for nanoparticle formation in rich premixed flames. *Combustion and Flame*, 127(1): 1995–2003, 2001. [doi:10.1016/S0010-2180\(01\)00303-0.](http://dx.doi.org/10.1016/S0010-2180(01)00303-0)
- <span id="page-20-6"></span>[11] M. A. Dobrowolski, A. Ciesielski, and M. K. Cyrański. On the aromatic stabilization of corannulene and coronene. *Phys. Chem. Chem. Phys.*, 13:20557–20563, 2011. [doi:10.1039/C1CP21994D.](http://dx.doi.org/10.1039/C1CP21994D)
- <span id="page-20-2"></span>[12] M. Frenklach. Reaction mechanism of soot formation in flames. *Physical Chemistry Chemical Physics*, 4(11):2028–2037, 2002. [doi:10.1039/b110045a.](http://dx.doi.org/10.1039/b110045a)
- <span id="page-20-3"></span>[13] M. Frenklach, M. Ramachandra, and R. Matula. Soot formation in shock-tube oxidation of hydrocarbons. *Symposium (International) on Combustion*, 20(1):871–878, 1985. [doi:10.1016/S0082-0784\(85\)80576-2.](http://dx.doi.org/10.1016/S0082-0784(85)80576-2) Twentieth Symposium (International) on Combustion.
- <span id="page-20-8"></span>[14] M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. A. Robb, J. R. Cheeseman, G. Scalmani, V. Barone, G. A. Petersson, H. Nakatsuji, X. Li, M. Caricato, A. Marenich, J. Bloino, B. G. Janesko, R. Gomperts, B. Mennucci, H. P. Hratchian, J. V. Ortiz, A. F. Izmaylov, J. L. Sonnenberg, D. Williams-Young, F. Ding, F. Lipparini, F. Egidi, J. Goings, B. Peng, A. Petrone, T. Henderson, D. Ranasinghe, V. G. Zakrzewski, J. Gao, N. Rega, G. Zheng, W. Liang, M. Hada, M. Ehara, K. Toyota, R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda, O. Kitao, H. Nakai, T. Vreven, K. Throssell, J. A. Montgomery, J. E. Peralta, F. Ogliaro, M. Bearpark, J. J. Heyd, E. Brothers, K. N. Kudin, V. N. Staroverov, T. Keith, R. Kobayashi, J. Normand, K. Raghavachari, A. Rendell, J. C. Burant, S. S. Iyengar, J. Tomasi, M. Cossi, J. M. Millam, M. Klene, C. Adamo, R. Cammi, J. W. Ochterski, R. L. Martin, K. Morokuma, O. Farkas, J. B. Foresman, and D. J. Fox. Gaussian 09, Revision A 02, 2009.
- <span id="page-20-9"></span>[15] S. Grimme. Semiempirical GGA-type density functional constructed with a longrange dispersion correction. *Journal of Computational Chemistry*, 27(15):1787– 1799, 2006. [doi:10.1002/jcc.20495.](http://dx.doi.org/10.1002/jcc.20495)
- <span id="page-20-4"></span>[16] N. Hansen, T. A. Cool, P. R. Westmoreland, and K. Kohse-Höinghaus. Recent contributions of flame-sampling molecular-beam mass spectrometry to a fundamental understanding of combustion chemistry. *Progress in Energy and Combustion Science*, 35(2):168–191, 2009. [doi:10.1016/j.pecs.2008.10.001.](http://dx.doi.org/10.1016/j.pecs.2008.10.001)
- <span id="page-20-7"></span>[17] J. Happold, H.-H. Grotheer, and M. Aigner. Soot precursors consisting of stacked pericondensed pahs. In H. Bockhorn, editor, *Combustion Generated Fine Carbonaceous Particles*, chapter 18, pages 277 – 288. KIT Scientific Publishing, Karlsruhe, Germany, 2009.
- <span id="page-20-0"></span>[18] S. J. Harris and A. M. Weiner. A picture of soot particle inception. *Symposium (International) on Combustion*, 22(1):333 – 342, 1989. [doi:10.1016/S0082-](http://dx.doi.org/10.1016/S0082-0784(89)80039-6) [0784\(89\)80039-6.](http://dx.doi.org/10.1016/S0082-0784(89)80039-6)
- <span id="page-20-1"></span>[19] D. Hou and X. You. Reaction kinetics of hydrogen abstraction from polycyclic aromatic hydrocarbons by H atoms. *Physical Chemistry Chemical Physics*, 19(45): 30772–30780, 2017. [doi:10.1039/c7cp04964a.](http://dx.doi.org/10.1039/c7cp04964a)
- <span id="page-20-5"></span>[20] J. B. Howard. Carbon addition and oxidation reactions in heterogeneous combustion and soot formation. 23(1):1107–1127, 1991.
- <span id="page-21-2"></span>[21] K. Johansson, M. Head-Gordon, P. Schrader, K. Wilson, and H. Michelsen. Resonance-stabilized hydrocarbon-radical chain reactions may explain soot inception and growth. *Science*, 361(6406):997–1000, 2018.
- <span id="page-21-1"></span>[22] A. Keller, R. Kovacs, and K.-H. Homann. Large molecules, ions, radicals and small soot particles in fuel-rich hydrocarbon flames. part iv. large polycyclic aromatic hydrocarbons and their radicals in a fuel-rich benzene–oxygen flame. *Physical Chemistry Chemical Physics*, 2(8):1667–1675, 2000.
- <span id="page-21-9"></span>[23] M. Kertesz. Pancake bonding: An unusual pi-stacking interaction. *Chemistry–A European Journal*, 25(2):400–416, 2019.
- <span id="page-21-5"></span>[24] M. R. Kholghy, G. A. Kelesidis, and S. E. Pratsinis. Reactive polycyclic aromatic hydrocarbon dimerization drives soot nucleation. *Phys. Chem. Chem. Phys.*, 20: 10926–10938, 2018. [doi:10.1039/C7CP07803J.](http://dx.doi.org/10.1039/C7CP07803J)
- <span id="page-21-10"></span>[25] R. A. Krueger and G. Blanquart. Predicting aromatic exciplex fluorescence emission energies. *Phys. Chem. Chem. Phys.*, pages 10325–10335, 2019. [doi:10.1039/C9CP02027F.](http://dx.doi.org/10.1039/C9CP02027F)
- <span id="page-21-3"></span>[26] A. L. Lafleur, J. B. Howard, K. Taghizadeh, E. F. Plummer, L. T. Scott, A. Necula, and K. C. Swallow. Identification of  $C_{20}H_{10}$  Dicyclopentapyrenes in Flames: Correlation with Corannulene and Fullerene Formation. *The Journal of Physical Chemistry*, 100(43):17421–17428, 1996. [doi:10.1021/jp9605313.](http://dx.doi.org/10.1021/jp9605313)
- <span id="page-21-0"></span>[27] P. J. Landrigan, R. Fuller, N. J. R. Acosta, O. Adeyi, R. Arnold, N. N. Basu, A. B. Baldé, R. Bertollini, S. Bose-O'Reilly, J. I. Boufford, P. N. Breysse, T. Chiles, C. Mahidol, A. M. Coll-Seck, M. L. Cropper, J. Fobil, V. Fuster, M. Greenstone, A. Haines, D. Hanrahan, D. Hunter, M. Khare, A. Krupnick, B. Lanphear, B. Lohani, K. Martin, K. V. Mathiasen, M. A. McTeer, C. J. L. Murray, J. D. Ndahimananjara, F. Perera, J. Potočnik, A. S. Preker, J. Ramesh, J. Rockström, C. Salinas, L. D. Samson, K. Sandilya, P. D. Sly, K. R. Smith, A. Steiner, R. B. Stewart, W. A. Suk, O. C. P. van Schayck, G. N. Yadama, K. Yumkella, and M. Zhong. The Lancet Commission on pollution and health. *The Lancet*, 2017. [doi:10.1016/S0140-6736\(17\)32345-0.](http://dx.doi.org/10.1016/S0140-6736(17)32345-0)
- <span id="page-21-8"></span>[28] M. Liu and W. H. Green. Capturing aromaticity in automatic mechanism generation software. *Proceedings of the Combustion Institute*, 37(1):575 – 581, 2019. [doi:10.1016/j.proci.2018.06.006.](http://dx.doi.org/10.1016/j.proci.2018.06.006)
- <span id="page-21-7"></span>[29] T. Lu and F. Chen. Multiwfn: a multifunctional wavefunction analyzer. *Journal of computational chemistry*, 33(5):580–592, 2012.
- <span id="page-21-6"></span>[30] J. W. Martin, G. J. McIntosh, R. Arul, R. N. Oosterbeek, M. Kraft, and T. Söhnel. Giant fullerene formation through thermal treatment of fullerene soot. *Carbon*, 125: 132–138, 2017. [doi:10.1016/j.carbon.2017.09.045.](http://dx.doi.org/10.1016/j.carbon.2017.09.045)
- <span id="page-21-4"></span>[31] J. W. Martin, R. I. Slavchov, E. K. Y. Yapp, J. Akroyd, S. Mosbach, and M. Kraft. The polarization of polycyclic aromatic hydrocarbons curved by pentagon incorporation: the role of the flexoelectric dipole. *The Journal of Physical Chemistry C*, 121 (48):27154–27163, 2017. [doi:10.1021/acs.jpcc.7b09044.](http://dx.doi.org/10.1021/acs.jpcc.7b09044)
- <span id="page-22-3"></span>[32] J. W. Martin, M. Botero, R. I. Slavchov, K. Bowal, J. Akroyd, S. Mosbach, and M. Kraft. Flexoelectricity and the formation of carbon nanoparticles in flames. *The Journal of Physical Chemistry C*, 122(38):22210–22215, 2018.
- <span id="page-22-8"></span>[33] J. W. Martin, K. L. Bowal, A. Menon, R. I. Slavchov, J. Akroyd, S. Mosbach, and M. Kraft. Polar curved polycyclic aromatic hydrocarbons in soot formation. *Proceedings of the Combustion Institute*, 37(1):1117–1123, 2019. [doi:10.1016/j.proci.2018.05.046.](http://dx.doi.org/10.1016/j.proci.2018.05.046)
- <span id="page-22-0"></span>[34] J. R. McConnell, R. Edwards, G. L. Kok, M. G. Flanner, C. S. Zender, E. S. Saltzman, J. R. Banta, D. R. Pasteris, M. M. Carter, and J. D. W. Kahl. 20th-Century Industrial Black Carbon Emissions Altered Arctic Climate Forcing. *Science*, 317 (5843):1381–1384, 2007. [doi:10.1126/science.1144856.](http://dx.doi.org/10.1126/science.1144856)
- <span id="page-22-9"></span>[35] A. Menon, J. Dreyer, J. W. Martin, J. Akroyd, J. Robertson, and M. Kraft. Optical band gap of cross-linked, curved, and radical polyaromatic hydrocarbons. *Phys. Chem. Chem. Phys.*, 2019.
- <span id="page-22-6"></span>[36] X. Mercier, O. Carrivain, C. Irimiea, A. Faccinetto, and E. Therssen. Dimers of polycyclic aromatic hydrocarbons: the missing pieces in the soot formation process. *Physical Chemistry Chemical Physics*, 21(16):8282–8294, 2019.
- <span id="page-22-7"></span>[37] J. H. Miller. The kinetics of polynuclear aromatic hydrocarbon agglomeration in flames. *Symposium (International) on Combustion*, 23(1):91 – 98, 1991. [doi:10.1016/S0082-0784\(06\)80246-8.](http://dx.doi.org/10.1016/S0082-0784(06)80246-8) Twenty-Third Symposium (International) on Combustion.
- <span id="page-22-5"></span>[38] J. H. Miller. Aromatic excimers: evidence for polynuclear aromatic hydrocarbon condensation in flames. *Proceedings of the Combustion Institute*, 30(1):1381–1388, 2005.
- <span id="page-22-10"></span>[39] J. Mukherjee, A. F. Sarofim, and J. P. Longwell. Polycyclic aromatic hydrocarbons from the high-temperature pyrolysis of pyrene. *Combustion and Flame*, 96(3):191– 200, 1994.
- <span id="page-22-2"></span>[40] A. Raj. Structural effects on the growth of large polycyclic aromatic hydrocarbons by C2H2. *Combustion and Flame*, 204:331 – 340, 2019. [doi:10.1016/j.combustflame.2019.03.027.](http://dx.doi.org/10.1016/j.combustflame.2019.03.027)
- <span id="page-22-4"></span>[41] A. Raj, S. Y. Yang, D. Cha, R. Tayouo, and S. H. Chung. Structural effects on the oxidation of soot particles by  $O_2$ : Experimental and theoretical study. *Combustion and Flame*, 160(9):1812 – 1826, 2013. [doi:10.1016/j.combustflame.2013.03.010.](http://dx.doi.org/10.1016/j.combustflame.2013.03.010)
- <span id="page-22-1"></span>[42] K. Rummel and P.-O. Veh. Die Strahlung Leuchtender Flammen. Erster Teil: Schrifttumsgrundlagen, Arbeitshypothesen und Vorversuche (Translation: The radiance of glowing flames. Part 1: Foundations, working hypotheses and preliminary experiments). *Archiv für das Eisenhüttenwesen*, 14(10):489–499, 1941.
- <span id="page-23-8"></span>[43] R. C. Santos, C. E. Bernardes, H. P. Diogo, M. F. M. Piedade, J. N. Canongia Lopes, and M. E. Minas da Piedade. Energetics of the thermal dimerization of acenaphthylene to heptacyclene. *The Journal of Physical Chemistry A*, 110(6):2299–2307, 2006.
- <span id="page-23-0"></span>[44] F. Schulz, M. Commodo, K. Kaiser, G. D. Falco, P. Minutolo, G. Meyer, A. D'Anna, and L. Gross. Insights into incipient soot formation by atomic force microscopy. *Proceedings of the Combustion Institute*, 37(1):885 – 892, 2019. [doi:10.1016/j.proci.2018.06.100.](http://dx.doi.org/10.1016/j.proci.2018.06.100)
- <span id="page-23-3"></span>[45] M. Sirignano, A. Collina, M. Commodo, P. Minutolo, and A. D'Anna. Detection of aromatic hydrocarbons and incipient particles in an opposed-flow flame of ethylene by spectral and time-resolved laser induced emission spectroscopy. *Combustion and Flame*, 159(4):1663 – 1669, 2012. [doi:10.1016/j.combustflame.2011.11.005.](http://dx.doi.org/10.1016/j.combustflame.2011.11.005)
- <span id="page-23-5"></span>[46] P. Sjoberg, J. S. Murray, T. Brinck, and P. Politzer. Average local ionization energies on the molecular surfaces of aromatic systems as guides to chemical reactivity. *Canadian Journal of Chemistry*, 68(8):1440–1443, 1990.
- <span id="page-23-10"></span>[47] D. Small, S. V. Rosokha, J. K. Kochi, and M. Head-Gordon. Characterizing the dimerizations of phenalenyl radicals by ab initio calculations and spectroscopy:  $\sigma$ bond formation versus resonance π-stabilization. *The Journal of Physical Chemistry A*, 109(49):11261–11267, 2005. [doi:10.1021/jp054244n.](http://dx.doi.org/10.1021/jp054244n)
- <span id="page-23-7"></span>[48] M. Solà. Forty years of clar's aromatic π-sextet rule. *Frontiers in chemistry*, 1:22, 2013.
- <span id="page-23-1"></span>[49] S. E. Stein and A. Fahr. High-temperature stabilities of hydrocarbons. *The Journal of Physical Chemistry*, 89(17):3714–3725, 1985. [doi:10.1021/j100263a027.](http://dx.doi.org/10.1021/j100263a027)
- <span id="page-23-2"></span>[50] D. Su, R. Jentoft, J.-O. Müller, D. Rothe, E. Jacob, C. Simpson, Z. Tomovic,´ K. Müllen, A. Messerer, U. Pöschl, R. Niessner, and R. Schlögl. Microstructure and oxidation behaviour of Euro IV diesel engine soot: a comparative study with synthetic model soot substances. *Catalysis Today*, 90(1):127 –132, 2004. [doi:10.1016/j.cattod.2004.04.017.](http://dx.doi.org/10.1016/j.cattod.2004.04.017) Air Pollution Abatement Symposium.
- <span id="page-23-6"></span>[51] A. Toro-Labbé, P. Jaque, J. S. Murray, and P. Politzer. Connection between the average local ionization energy and the Fukui function. *Chemical Physics Letters*, 407(1):143 – 146, 2005. [doi:10.1016/j.cplett.2005.03.041.](http://dx.doi.org/10.1016/j.cplett.2005.03.041)
- <span id="page-23-9"></span>[52] T. S. Totton, A. J. Misquitta, and M. Kraft. Assessing the polycyclic aromatic hydrocarbon anisotropic potential with application to the exfoliation energy of graphite. *The Journal of Physical Chemistry A*, 115(46):13684–13693, 2011.
- <span id="page-23-4"></span>[53] T. S. Totton, A. J. Misquitta, and M. Kraft. A quantitative study of the clustering of polycyclic aromatic hydrocarbons at high temperatures. *Physical chemistry chemical physics*, 14(12):4081–94, 2012. [doi:10.1039/c2cp23008a.](http://dx.doi.org/10.1039/c2cp23008a)
- <span id="page-23-11"></span>[54] T. S. Totton, A. J. Misquitta, and M. Kraft. A quantitative study of the clustering of polycyclic aromatic hydrocarbons at high temperatures. *Physical Chemistry Chemical Physics*, 14(12):4081–4094, 2012. [doi:10.1039/C2CP23008A.](http://dx.doi.org/10.1039/C2CP23008A)
- <span id="page-24-4"></span>[55] R. S. Tranter, S. J. Klippenstein, L. B. Harding, B. R. Giri, X. Yang, and J. H. Kiefer. Experimental and theoretical investigation of the self-reaction of phenyl radicals. *The Journal of Physical Chemistry A*, 114(32):8240–8261, 2010.
- <span id="page-24-1"></span>[56] A. Violi, A. Kubota, T. Truong, W. Pitz, C. Westbrook, and A. Sarofim. A fully integrated kinetic monte carlo/molecular dynamics approach for the simulation of soot precursor growth. *Proceedings of the Combustion Institute*, 29(2):2343–2349, 2002. [doi:10.1016/S1540-7489\(02\)80285-1.](http://dx.doi.org/10.1016/S1540-7489(02)80285-1)
- <span id="page-24-2"></span>[57] G. Vitiello, G. De Falco, F. Picca, M. Commodo, G. D'Errico, P. Minutolo, and A. D'Anna. Role of radicals in carbon clustering and soot inception: A combined epr and raman spectroscopic study. *Combustion and Flame*, 205:286–294, 2019.
- <span id="page-24-0"></span>[58] H. Wang. Formation of nascent soot and other condensed-phase materials in flames. *Proceedings of the Combustion Institute*, 33(1):41 – 67, 2011. [doi:10.1016/j.proci.2010.09.009.](http://dx.doi.org/10.1016/j.proci.2010.09.009)
- <span id="page-24-3"></span>[59] H. Yuan, W. Kong, F. Liu, and D. Chen. Study on soot nucleation and growth from PAHs and some reactive species at flame temperatures by reaxff molecular dynamics. *Chemical Engineering Science*, 195:748 – 757, 2019. [doi:10.1016/j.ces.2018.10.020.](http://dx.doi.org/10.1016/j.ces.2018.10.020)