Response to Reviewer 1:

Review of “A Study on the Key Factors Determining the Hygroscopic property of Black Carbon” by Su et al.

In this study, the authors investigate the hygroscopicity of five types of black carbon (BC) samples in the laboratory by measuring the change in mass of each sample as a function of relative humidity. They also try to interpret the hygroscopicity data by analyzing BC samples with the OC/EC analyzer, GC-MS, IC, and Raman spectroscopy. Because the hygroscopicity of BC-containing particles is an important parameter affecting the atmospheric behavior and climatic impacts of BC, the data presented here is valuable. However, many clarifications are required so that the readers can understand their experimental procedures and the validity of their results and interpretation. The reviewer believes major revisions are necessary before its publication in ACP.

Reply: Thank you very much for the time and effort spent in reviewing the manuscript. Your guidance is greatly appreciated, and we believe it will significantly improve our manuscript. The following is a point-by-point response to all the comments, and the manuscript has been revised substantially according to your comments.

Major comments

Q1. Due to a lack of information and explanation, the reviewer could not understand the details of the experiments. For example, were the BC particles generated by the combustion of n-hexane, decane, and toluene collected on filters for further experiments? Is the U black carbon (UBC) sample originally suspension or a dry powder? No details of the aging experimental setup were also provided. Are there any previous studies using the same setup? What are the diesel engine operating conditions for collecting diesel BC (DBC) samples? It is difficult for the reader to reproduce their experiments with only the information provided in this manuscript.

A1. Thanks for your reminding. Here, we show the detail information about these experimental setups and added in the revised manuscript.

1. For prepared BC particles, they were obtained by burning n-hexane, decane and
toluene (AR, Sinopharm Chemical Reagent Lo., Ltd) in a co-flow system as described in our previous studies (Han et al., 2012; Zhao et al., 2017). Soot was collected on a quartz disc (7 cm in diameter) over a diffusion flame, eventually stored in a brown bottle (Agilent).

2. UBC (Printex U from Degussa, CAS No.: 1333-86-4) samples are dry powders and used as purchased. The aging experiments were performed in a quartz flow tube reactor (12 cm in length, 10 mm in diameter), as seen below. UBC was placed into a quartz flow tube reactor. SO$_2$ with certain concentration was introduced with carrier air. The RH in the flow could be established by adjusting the ratios of dry and humid air. A Xenon was used as light source of photo oxidation. This method has been used in our previous study (Zhang et al., 2022).

3. Diesel black carbon (DBC) was collected from the diesel particle filter (DPF) of a China VI heavy-duty diesel engine (ISUZU from China). A diesel engine bench test was run under the conditions of World Harmonized Transient Cycle (WHTC). China VI fuels were used in the study, meeting the GB T32859-2016 standard. The mechanism of SO$_2$ photooxidation on these DBC was studied in our previous study (Zhang et al., 2022).

This detailed information has been added in the revised manuscript (Line 88-102)

Q2. The definition of the term “BC” should be clearly stated. In this study, the authors seem to refer to a BC particle as an entire soot particle that can contain other organic or inorganic compounds. However, this treatment can make various descriptions unclear. For example, an expression like "OC in BC" (Line 112) would be confusing for many
researchers in the community. As discussed in previous studies (e.g., Petzold et al., 2013), it may be clear if mixed particles containing a BC fraction are termed “BC-containing particles” instead of “BC particles.” Please consider the terminology used in this study and state it clearly.

A2: Thanks for your suggestion. Previous study (Petzold et al., 2013) have mentioned that soot denotes the ensemble of the particles emitted during incomplete combustion, i.e., the sum of black carbon and organic carbon. The term soot generally refers to the source mechanism of incomplete combustion of hydrocarbon fuels rather than to a material property. In this study, five types of BC particles were analyzed. Three of the black carbon were produced by the co-flow burner and fueled with n-hexane, decane, and toluene. Diesel black carbon was produced in a diesel engine with China VI fuels. All of them are products of incomplete combustion of hydrocarbon fuels, and Printex U powder was used as a model soot. Therefore, we think that “soot” is more appropriate than black carbon for samples used in this study.

Names of samples have been changed in the revised manuscript:

n-hexane / decane / toluene flame BC → n-hexane / decane / toluene flame soot

Diesel BC → diesel soot (DS)

Printex U BC → Printex U powder (U-soot).

All “black carbon” and “BC” was changed to “soot” in the revised manuscript.

Q3. In the reviewer's understanding, this study analyses bulk samples, which means that the information on the mixing states of the particles collected on filters (e.g., mixing states of the particles collected from diesel exhaust) is not obtained. Because this study discusses the hygroscopicity based on the increase in the mass of each sample, does it implicitly assume that all non-BC components are internally mixed with BC? In other words, is the measured hygroscopicity considered an upper limit of the hygroscopicity of BC-containing particles? (If some non-BC components are externally mixed with BC, then the hygroscopic growth (mass increase) of the non-BC component should not contribute to the hygroscopic growth of BC-containing particles.) Explanations should
According to the previous comment (Q2), we have replaced black carbon to soot to describe the particles which contain EC, OC and other components. It is known that soot can exist in external mixing state (distinct from other components) and internal mixing state (incorporated within other components) (Jacobson, 2001). In this study, all soot samples are products of incomplete combustion of hydrocarbon fuels. During combustion processes, organic carbon (OC) is cogenerated with elemental carbon (EC) and mixed with EC through adsorption (Ogren and Charlson, 1983). Thus, OC and EC are usually assumed in an internal mixing state in soot particles.

As for inorganic components in soot, previous study demonstrated that diesel soot produced from diesel fuel contains up to 3.5% water-soluble substances, with the major fraction being comprised of inorganic ions such as sulfate (SO$_4^{2-}$) and chloride (Cl$^-$) (Lammel and Novakov, 1995). In our study, the IC results revealed the significant presence of sulfate (SO$_4^{2-}$) and nitrate (NO$_3^-$) ions within Diesel Soot particles. In order to clarify the mixing state of these water-soluble inorganic salts with DS, we performed TEM characterization with different dispersible liquids (ethanol and ultrapure water).

![Figure s1. TEM images of diesel soot, (a) sample were ultrasonically dispersed in ethanol, (b) sample were ultrasonically dispersed in ultrapure water.](image)

As seen in Figure s1a, the images of diesel soot aggregates dispersed in ethanol show a typical chain-like agglomerated structure composed of small spherical primary particles. In contrast, the samples dispersed in ultrapure water show many regular rectangle images due to salt crystallization (Figure s1b). Thus, it can be inferred that diesel soot and inorganic salts exist in a state of internal mixing.

As for SO$_2$ aged samples, soot particles and sulfate species could also exist in a
state of internal mixing, since our previous study demonstrated that SO\textsubscript{2} is oxidized by photo induced OH radicals on soot surface to form sulfuric acid (Zhang et al., 2022). Therefore, in our study, the mixing states of all soot particles could be considered in internal mixing state.

**Q4.** A clearer description of this study's new findings should be provided. The authors point out “a lack of comprehensive understanding regarding the factors that determine the hygroscopic properties of fresh BC” in the abstract and introduction (Lines 15 and 75). From that perspective, however, aging experiments with SO\textsubscript{2} are not intended to gain insight into fresh BC and are, therefore, not directly relevant. Also, it is well known that coatings of inorganic substances increase the hygroscopicity of BC-containing particles.

**A4.** Thanks for your comments and suggestions. The new findings in this study include quantitative measurement of vapor sorption capacity of various soot particles, the relationship between the hygroscopicity and the microstructure of soot, the effect of typical mixing components on the hygroscopicity of soot. As for fresh soot, we originally meant soot particles produced by fuel combustion, which is always considered to be hydrophobic. Because the hygroscopicity of soot plays an important role in its aging process, exploring the critical factors affecting the hygroscopicity of fresh produced soot is of significance for understanding their aging process. We intended to reveal the influence of microphysics and the content and composition of OC on the hygroscopicity of fresh soot particles. Nevertheless, we think that the use of the term "fresh" in the abstract and introduction sections is not necessary and could lead to misunderstandings. Therefore, we have deleted the word "fresh" from these sections. (Line 15 and 75).

We agree that it is well known that coatings of inorganic substances increase the hygroscopicity of BC-containing particles. Field studies showed that coating nitrate and sulfate on BC particles can increase the growth factor of these BC particles up to ~1.4-1.6 (Li et al., 2018; Liu et al., 2013). In laboratory investigations exploring the impact of sulfate on the hygroscopicity of BC, condensation of gaseous sulfuric acid has been
commonly used to simulate the mixing process of sulfate and BC in the atmosphere. Previous study showed that the hygroscopic growth factor of BC particles exposed to sulfuric acid vapor can reach 1.52 at 90% RH which is lower than pure sulfuric acid particles (2.03) (Zhang et al., 2008). Khalizov et al. found similar results with the hygroscopic growth factor of sulfuric acid coated soot particles in the range from 1.32 to 1.49 at 90% RH (Khalizov et al., 2009). These results represent the case where soot particles are suspended in the ambient atmospheric with high H$_2$SO$_4$ level, which may not suitable for most real atmosphere. Our previous study found that the photo oxidation of SO$_2$ on soot surface could be potential formation pathway to form internal mixture of sulfuric acid and soot (Zhang et al., 2022). Thus, in this study, we prepared SO$_2$ aged soot particles by the direct heterogeneous photochemical reaction between SO$_2$ and soot, which will promote the understanding of the hygroscopicity of soot particles containing sulfuric acid in the atmosphere.

**Q5.** The title is too broad and should be more specific. English proofreading throughout the manuscript is also desired.

**A5.** The title was changed to “A study on the influence of inorganic ions, organic carbon, and microstructure on the hygroscopic property of soot”. We have also had native English speaker polish our English writing.

**Specific comments:**

**Q1.** L34, There have been many studies on the radiative effects of BC after 2001, and their understanding has been much updated. Please consider adding or updating the references.

**A1.** We have updated the reference in the revised manuscript (Line 35):

Soot aerosol can influence climate by directly absorbing solar radiation and affecting cloud formation and surface albedo through deposition on snow and ice (Liao et al., 2015; Peng et al., 2016), which results in the contribution of soot to anthropogenic radiative forcing second only to that of CO$_2$ (Bond et al., 2013; Cappa et al., 2012; Liu et al., 2017)
Q2. L51, Is "commercial BC" not "fresh BC"? Is the UBC a commercial BC in this study?
A2. In this sentence, “fresh BC” means soot prepared by fuel combustion in the laboratory, which was used for comparison with aged samples. UBC is a kind of commercial BC (Printex U from Degussa, CAS No.: 1333-86-4). The ion chromatography analysis results indicate that UBC contains a small amount of sulfate, which may be due to aging during transportation and storage. Therefore, we think UBC could not be considered fresh BC. The corrections have been made in the manuscript (Line 49-51):
The hygroscopic behavior of soot has been widely studied. It was found that soot prepared in the laboratory or commercial soot appears to be hydrophobic as there is no noticeable uptake of water at unsaturated humidity.

**Q3.** L111, EC = EC1+EC2 + EC3?

**A3.** When the content of OC and EC was measured using a thermal-optical transmittance OC/EC analyzer, OC is usually defined as OC1+OC2+OC3+OC4+OP and EC is defined as EC1+EC2+EC3-OP (Chow et al., 1993; Li et al., 2016). The reason is that the EC fraction formed by OC conversion during pyrolysis is referred to as pyrolyzed carbon (OP) (Boparai et al., 2008). Thus, OP should be subtracted from the EC fraction and added to the OC fraction.

**Q4.** L166: P has already been defined in L165.

**A4.** The sentence “P is the equilibrium pressure of the adsorbing gas” has been deleted from the revised manuscript.

**Q5.** L183: v and v_m are lowercase letters in Eq. (3) but uppercase letters in Eq. (1).

**A5.** Thank you for your careful check. The corrections have been made in the manuscript (Line 201):

\[ V = \frac{V_m c P}{(P_0 - P)(1 + (c - 1)(P/P_0))} \]

**Q6.** L195: Table 2 lists the concentrations of \( \text{SO}_4^{2-} \) and \( \text{NO}_3^- \). Can the mass of EC and OC also be listed in the same Table?

**A6.** The ratio of OC/EC of soot samples has been added in the Table 2. (Line 221):

<table>
<thead>
<tr>
<th>Soot</th>
<th>Mass concentration of ( \text{SO}_4^{2-} ) (µg mg(^{-1}))</th>
<th>Mass concentration of ( \text{NO}_3^- ) (µg mg(^{-1}))</th>
<th>OC/EC</th>
</tr>
</thead>
<tbody>
<tr>
<td>n-hexane flame soot</td>
<td>0.00</td>
<td>0.19</td>
<td>0.41±0.02</td>
</tr>
</tbody>
</table>
Q7. L213: Five different samples were used in this study, but the results from the OC/EC, GC-MS, and Raman analyses are only shown for the three samples. Why is that?

A7. In section 3.1, the inorganic ions present in five different black carbons were analyzed using IC. The findings revealed that DBC and UBC contained significant amounts of SO$_4^{2-}$ and NO$_3^-$ ions. On the other hand, the three prepared black carbons exhibited minimal levels of inorganic ions. Thus, the three prepared black carbons were chosen to investigate the impact of factors other than inorganic ions on the hygroscopic properties of black carbon. Because inorganic components affect the hygroscopicity more significantly than OC and microstructure, the information of OC and microstructure of DBC and UBC was not shown.

Q8. L273: If I_D is the sum of I_D1 and I_D4, why are some values of I_D1/I_G larger than I_D/I_G in Table 3?

A8. Thank you for your careful check. We realized that there were some errors in the data handling process. As mentioned above, the first-order Raman spectra of three prepared soots has good curve-fitting results, four Lorentzian-shaped bands (D1, D2, D4, and G, centered at about 1360, 1620, 1180, and 1580 cm$^{-1}$, respectively) and one Gaussian-shaped band (D3, centered at around 1500 cm$^{-1}$) were used in the curve-fitting process. The intensities of G bands (IG) have been widely determined using the sum of D2 and G bands. The value of ID1/IG is the area of the D1 band divided by the sum of

<table>
<thead>
<tr>
<th>Sample</th>
<th>I_D1</th>
<th>I_D4</th>
<th>I_D1/I_G</th>
</tr>
</thead>
<tbody>
<tr>
<td>Toluene flame soot</td>
<td>0.00</td>
<td>0.18</td>
<td>0.24±0.04</td>
</tr>
<tr>
<td>Decane flame soot</td>
<td>0.00</td>
<td>0.22</td>
<td>0.16±0.06</td>
</tr>
<tr>
<td>DS</td>
<td>11.46</td>
<td>1.44</td>
<td>0.14±0.02</td>
</tr>
<tr>
<td>U-soot</td>
<td>2.55</td>
<td>0.24</td>
<td>0.12±0.03</td>
</tr>
<tr>
<td>U-soot aged 2h</td>
<td>4.83</td>
<td>0.20</td>
<td>---</td>
</tr>
<tr>
<td>U-soot aged 6h</td>
<td>7.14</td>
<td>0.19</td>
<td>---</td>
</tr>
<tr>
<td>U-soot aged 10h</td>
<td>9.61</td>
<td>0.20</td>
<td>---</td>
</tr>
</tbody>
</table>
the areas of the D2 and G bands. However, I incorrectly calculated the area of the D1 band divided by the area of the G band. The same error exists in the calculation of the value of ID4/IG and ID2/IG. Considering that the value of ID2/IG has no meaning in this paper, I have deleted it. I fixed the error and put the new result in the table 3 (Line 290):

Table 3. Parameters ID1/IG, ID4/IG, ID2/IG and Lₐ of n-hexane, decane and toluene flame soot.

<table>
<thead>
<tr>
<th>Fuels</th>
<th>ID1/IG</th>
<th>ID4/IG</th>
<th>ID2/IG</th>
<th>Lₐ (Å)</th>
</tr>
</thead>
<tbody>
<tr>
<td>n-hexane</td>
<td>2.59±0.13</td>
<td>0.33±0.01</td>
<td>2.87±0.13</td>
<td>15.34±0.69</td>
</tr>
<tr>
<td>toluene</td>
<td>2.87±0.03</td>
<td>0.42±0.09</td>
<td>3.23±0.09</td>
<td>13.62±0.38</td>
</tr>
<tr>
<td>decane</td>
<td>2.69±0.08</td>
<td>0.75±0.01</td>
<td>3.49±0.15</td>
<td>12.63±0.53</td>
</tr>
</tbody>
</table>

Q9. L295: Under the experimental RH conditions lower than the deliquescence RH of ammonium sulfate, ammonium sulfate would not contribute to hygroscopic growth even if present in abundance?

A9. The IC experiment results of UBC show that it does not contain NH₄⁺ cations. Our previous study showed that heterogeneous reaction between SO₂ and BC leads to the formation of sulfuric acid coating on the surface of BC (Zhang et al., 2022). The presence of sulfuric acid would not contribute to hygroscopic growth under low RH condition.

Q10. L306: Since this study examines not the ice nucleating property but the hygroscopicity of BC-containing particles, the comparison with previous studies focusing on the ice nucleating properties of BC may not be appropriate here. I suggest adding more explanation or eliminating the description of the ice nucleation. In addition, since it is now common to refer to ice nucleating particles (INPs) rather than IN, I suggest modifying or eliminating the use of the term IN in the abstract.

A10. Thanks for your suggestion. Due to its lack of relevance in the article, the mention of black carbon's ice nucleation property was deemed inappropriate and subsequently removed.
REFERENCE


