

Application of small angle and wide angle X-ray scattering for the characterization of carbonaceous materials, aerosols, and particles

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Small angle X-ray scattering (SAXS) can provide quantitative information on internal surface areas, porosity, particle size, void size distributions, surface roughness and fractal dimension of surfaces, interfaces and particles and aggregate structures. Applied in-situ and ex-situ, SAXS even permits to derive kinetic parameters for chemical reactions and transformations. In particular homogeneous systems such as soot can be studied with SAXS, and carbonaceous materials have been widely used to develop this technique. Wide-angle X-ray scattering (WAXS) permits to measure crystallite sizes and to distinguish aromatic and aliphatic structures in carbon materials. WAXS was particularly applied for coal research. We present here studies that we made on diesel exhaust soot for combustion enginering and environmental science, as well as studies on model systems such as glassy carbon (pore size and connectivity evolution) and on aerogels (inclusion of third phases such as metal clusters).

Scattering techniques, including light scattering (LS), provide statistically very robust data, complementary and often even superior to microscopy data. Small angle X-ray scattering (SAXS), when done at synchrotron radiation sources, can provide data very fast. At best, a scattering curve can be obtained in a fraction of 1 second. High resolution data may need about 15 minutes to cover a length scale from several microns to 1 nanometer. A shortcoming of scattering techniques is that they do require some minimum amount of material, such as milligrams, which always exceeds the amount necessary for TEM studies, which ultimately needs one tiny particle only. Also, SAXS cannot be applied very well for chemically very inhomogeneous systems. However, for soot studies, SAXS is perfect. Wide angle X-ray scattering (WAXS), pretty mch like SAXs, is a diffuse scattering technique which basically uses profile and background scattering analysis. It has been extensively applied for coal research with much success.

A quantitative methodology has been developed for analysis of SAXS data obtained from diesel soots which is capable of distinguishing three characteristic size parameters that characterize the soot: a soot particle agglomerate size, a primary particulate size, and a particulate sub-unit size, depending on engine and fuel conditions. The most prominent feature is visible in the log-log plot of scattering curves between q = 0.001 °A-1 and 0.01, which is due to soot particle agglomerates. In addition to differences in the soot primary particle and aggregate size, the stiffness of the aggregates was probed by pressing soot powder into pellets under various pressures. The scattering pattern of soot experiences systematic changes upon pressurizing, in particular a systematic, pressure dependent shift of the aggregate size signature which can be used to characterize the stiffness of the aggregates. Diesel soot from idle and load engine condition was pressed into pellets at

pressures ranging up to approximately 8.5 GPa. Soot powder was also immersed in acetone in order to obtain soot aggregates without agglomeration. Small angle X-ray scattering was carried



out on the powder, the pellets, and on the acetone immersed soot. Powder and pellets show characteristic aggregate structure at small scattering vectors. Scattering curves of the pellets show a shift of the aggregate size related scattering feature towards larger scattering vectors for increasing pressure. For the highest pressures, this aggregate structure vanished, while the suspected primary particle scattering became visible as the asymptote of the aggregate scattering structure. Aggregate size of powder was about 290 nm for the idle soot and 240 nm for the load soot. Primary particle size was 14.3 and 10.2 nm, respectively. Idle soot showed a higher compressibility than the load soot. Pressing the soot into pellets eliminates scattering from aggregation of primary particles and provides a good route to reveal the otherwise inaccessible primary particle scattering. In addition, studying the aggregate structure as a function of pellet pressure permit to derive compacticity data of the soot. Without extracting volatiles, it is ultimately not possible to quantify the impact of the volatiles like lubricants and fuel on the compaction behaviour of the soot under pressure. This remains particularly unclear for idle soot, which contains more volatiles than load soot but yet strongly resists pressure. The USAXS technique combined with pressing pellets appears to be valuable for the study of materials which are built from aggregates of primary particles, such as soot, or carbon black, silica gel, and many other materials. The advantage of using scattering techniques is that a considerable amount of material can be studied with reasonable experimental effort, providing robust and statistically representative data compared to single aggregate analysis.

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Objective

Scattering techniques, including light scattering (LS), provide statistically very robust data, complementary and often even superior to microscopy data. Small angle X-ray scattering (SAXS), when done at synchrotron radiation sources, can provide data very fast. At best, a scattering curve can be obtained in a fraction of 1 second. High resolution data may need about 15 minutes to cover a length scale from several microns to 1 nanometer. A shortcoming of scattering techniques is that they do require some minimum amount of material, such as milligrams, which always exceeds the amount necessary for TEM studies, which ultimately needs one tiny particle only. Also, SAXS cannot be applied very well for chemically very inhomogeneous system However, for soot studies, SAXS is perfect. Wide angle X-ray scattering (WAXS), pretty mch like SAXs, is a diffuse scattering technique which basically uses profile and ckground scattering analysis. It has been extensively applied for coal research with much success



Left: Diesel exhaust from heavy duty truck. Right: Diesel soot sample generation at Univeritu of Utah.

Samples: Diesel PM from 50/50 Chevron/Phillips reference fuels T22/U15, oxygenated with DEC and ethanol, operated under idle/load. Oxygenated fuel is called "Mix A" and "Mix B". Addition of 1000 ppm ferrocene to some of the fuels was made to catalytically oxidize the soot and prevent graphitization.

Thermogravimetric Analysis



Left: TGA profiles of load (black) and idle (red) soot from oxygenated Diesel, including the temperature profile TGA was operated with N_{2^1} and at 750°C with O_{2^n} . Right: Summary of TGA results. Data in parentheses are based on Soxhlet extraction Idle soot contains more volatiles.

Particle and pore size and porosity in diesel soot and GC



Left: Log-log plot of small angle scattering curves reveal at least 5 size ranges in Diesel PM, with size $L=2\pi/q$. Curve with open symbols was obtained after subtraction of Porod- and constant background scattering. Exponent of decay allows determination of fractal dimension, and was close to -4 for high q range and thus indicates smooth surfaces of primary particles and sub-units. For low q, exponents of decay are close to -3. Right: Comparison of scattering curves for soot and glassy carbon, a standard SAXS material with many micropores.

Quantitative data for diesel soot

Seat	Elementary units Djanj	Sub anits D (nm)	Primary Particles D [am]	high q exponent	Fractal Encodes	low q exponent	Fractal dimension
Diesel, idle	1.5	17.4	49.16	1.99	2.01	3.28	2.72
Direct, load	1.6	14.5	41.50	3.86	2.14	3.12	2.88
Mix A, idle	1.9	21.1 (14.2	78.29	3.97	2.03	3.62	2.98
Mix A, load	1.4	13.8 (12)	36.78	1.96	2.04	3.09	2.91
Mix B, idle	2.0	143 (143	83.85	3.92	2.05	2.96	2.96
Mix B, load	14	22.0 (18.6	48.73	3.98	2.02	2.75	2.75

Elementary particles sizes 1-2 nm range. Form compact cluster to built subunits of 15-20 nm size. These build up larger structures (primary particles) of 40-80 nm, which form aggregates. Aggregates are found at q-values of 0.001 1/A, though harder to resolve in the SAXS curves. Idle soot has generally larger particles than load soot.

Pressure and compaction studies with USAXS



Left: SAXS scattering curves for diesel soot as poder, pressed pellet, and immersed in acetone. Sample environment modiefies soot aggregation slightly and has impact on scattering curve. Right: Shift of characteristic structures in scattering curves upon pressurizing of samples indicates aggregate rearrangement



Maxima in Kratky plots of scattering curves provide information about compactness of soot particles and size of agglomerates: L=n/q



systematic trend for aggregate size rearrangement, which can be used for elasticity estimation



Quantitative analysis of diffuse XRD diffractograms (WAXS) provides information on aromaticity (area under y-band neak vs. the entire neak area including (002) peak), ratio of crystalline/amorphous carbon (background scattering), and crystallite sizes (Scherrer Formula).



Left: X-ray diffractograms from load/idle soot, and reference Bragg peaks of graphite (2H Graphite PDF 26-1079). Center: Deconvolution of Peak into (002) and y-sideband for determination of aromaticity. Right: Comparison of load/idle soot XRD from Diesel and oxygenated Diesel Mix A, Mix B.

> 10.48 16.68

6.96

12.86 8.64 14.92

10.78 16.30

8.64 8.97 6.24

Aromaticiy of oxygenated and nonoxygenated diesel exhaust PM for idle and A B 0.34 0.31 load engine conditions. 0.22 0.15 0.05 Idle soot particles have smaller crystallites than load soot Aromaticity higher for idle soot from oxygenated Diesel (Mix A, Mix B). Idle soot contains more amorphous carbon than crystalline carbon. Adding oxygenates to Mx R. load fuel causes bigger differences in the structure between idle and load soot, in line with McA. Mr Mx A. load NEXAFS and TGA.

Soot structure depending on engine operation + fuel additives

Idle soot particles have smaller crystallites than load soot. Aromaticity is higher for idle soot from oxygenated Diesel (Mix A, B). Idle soot contains more amorphous carbon (upper part of column, grains) than crystalline Adding carbon (lower part, planes). Adding oxygenates to the fuel causes bigger differences in the structure between idle and oad soot, in line with observations form the NEXAES and TGA



Ferrocene is added to diesel fuel in order to prevent formation of graphitic soot structures. This facilitates subsequent soot oxidation in aftertreatment devices. WAXS data explicitly show that ferrocene soot lacks significantly in graphite-like structures. Instead, a aliphatic gamma side-band is observed. Results in line with NEXAFS spectroscopy data, as shown in the Figure below







Porosity and Chord Length distribution

Porosity evolution of activated (right side) and non-activated GC (left) was studied by SAXS. Based on first geometric principles (*Roiviaul linear integration principle*, storage and the side of the site of t













ples alters SAXS curves, too. Anomalous SAXS permits to ion. Metal clusters can be modelled by spheres (bottom, left). Ostwald ripening as coarsening process (upper, right)

References

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Inhomogeneity and Metal Inclusions