

atems: Analysis tools for TEM images of carbonaceous particles

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DOI: 10.21105/joss.06416

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Submitted: 10 October 2023 Published: 25 July 2024

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Summary

The objective of atems is to provide a suite of open source analysis tools (largely in Matlab) for transmission electron microscopy (TEM) image analysis that are specifically designed for soot and related carbonaceous particles (e.g., tarballs). This codebase started as a manual analysis code by Dastanpour & Rogak (2014), with the first automated methods added by Dastanpour et al. (2016). The current, open source version has been streamlined and expanded to include a larger suite of automated analysis methods from the literature, as detailed in the following section. In this regard, a key contribution of this codebase is to provide open source implementations of multiple analysis methods spanning a range of laboratories. This codebase places these methods in the same framework, with the goal of enabling intercomparisons of analysis routines across a range of data.



Figure 1: Sample TEM image of soot demonstrating the aggregate structure, where a is an unlabeled image containing soot aggregates and b is that same image with the aggregates labeled.

Statement of need

Soot, carbon black, and other carbonaceous particles have important climate, health, and technological impacts that depend on their morphology. These particles have complex shapes composed of a collection of small, primary particles in fractal arrangements, as shown in Figure 1a. TEM images of these particles allow for detailed information about particle morphology that is unavailable in other characterization techniques. However, extracting this information requires image analysis across a statistically-significant number of particles,

Sipkens et al. (2024). atems: Analysis tools for TEM images of carbonaceous particles. *Journal of Open Source Software*, 9(99), 6416. 1 https://doi.org/10.21105/joss.06416.



with the quality of conclusions improving as the number of characterized particles increases. For instance, Kelesidis et al. (2020) suggested quantifying at least 400 primary particles per experimental condition in a premixed flame to get an accurate average primary particle diameter from manually drawing elipses (that study counted 800 primary particles). In the broader literature, a few hundred particles per condition seems to be standard, with other authors having employed between 150 and 400 particles per condition (Liati et al., 2014; Marhaba et al., 2019; Trivanovic et al., 2019, 2020), depending on the type of analysis. For multiple conditions, this can quickly expand to over 1000 particles. This characterization is often done manually, which at a minimum of several minutes per aggregate, is incredibly labour intensive. Unfortunately, the low contrast (carbonaceous particles on carbon films) and complex particle morphology of common carbonaceous particles makes automated analysis challenging, requiring unique analysis methods over those developed for traditional TEM image analysis of many engineered nanomaterials (Schneider et al., 2012). At the same time, existing automated methods across the literature are typically only applied to data from a single laboratory, with few exceptions (Anderson et al., 2017; Sipkens et al., 2021). This limits comparability between laboratories (Sipkens et al., 2023).

Methods

After loading images (with an automated method provided for doing so), analysis involves two major steps.

The first step is segmentation of the aggregates from their background. Available methods include the slider-based manual approach of Dastanpour & Rogak (2014); the common Otsu method; a modification of Otsu by Dastanpour et al. (2016) that employs morphological operations to improve segmentation; the *k*-means approach of Sipkens & Rogak (2021); and carboseg, which is the convolutional neural network (CNN) approach from Sipkens et al. (2021). Functionality is also available to prepare (e.g., read and crop image footers) and export images for external analysis, prior to reading the images in for subsequent analysis. This enables external extensions, such as the WEKA segmentation method of Altenhoff et al. (2020). Tools are then available to compute aggregate projected area, perimeter, and circularity, among other properties. A sampling of segmentations produced by these methods is presented in Figure 2.



Figure 2: Sample segmentations across a range of methods available in this code. The manual method corresponds to an updated version of the code development by Dastanpour et al. (2016). The Otsu segmentation is standard Otsu, without any adaptations. The *k*-means method is that described by Sipkens & Rogak (2021). TWS refers to trainable WEKA segmentation based on the method described by Altenhoff et al. (2020), which makes use of the code enabling external extensions. These first four panels correspond to images from Sipkens & Rogak (2021). The final panel corresponds to the convolutional neural network method described by Sipkens et al. (2021).



Second, this code works to identify primary particles, that is the small, roughly circular structures inside the aggregates. Available methods include a updated version of the Euclidean distance mapping–surface-based scale analysis (EDM-SBS) of Bescond et al. (2014), converted from SciLab to Matlab in association with Sipkens et al. (2021) (functionality between the two languages resulted in minor differences); the Euclidean distance mapping–watershed (EDM-WS) method of De Temmerman et al. (2014); the pair correlation method (PCM) of Dastanpour et al. (2016); the Hough transform method of Kook et al. (2016); and the Hough transform method of Altenhoff et al. (2020).

General plotting and other utilities (tools.*) are provided to enable further analysis and visualization (e.g., as in Figure 1b and Figure 2).

Use

This code has been used in a number of studies in the literature. This code was used by Sipkens et al. (2021) to compare multiple segmentation and primary particle analysis methods. The code was also used by Trivanovic et al. (2019), Kheirkhah et al. (2020), and Trivanovic et al. (2020) to perform image analysis of marine engine and flare soot. The *k*-means method in this code (Sipkens & Rogak, 2021) was also employed for soot by Li (2022).

Acknowledgements

We wish to acknowledge related code released in association with some of the cited work. These include Matlab code provided in Kook et al. (2016) and SciLab code released in association with Bescond et al. (2014). The code from Altenhoff et al. (2020) was provided by the authors and adapted to the present format.

We also wish to acknowledge funding by the Canadian Council of the Arts (Killam Fellowship), the Natural Sciences and Engineering Research Council of Canada (NSERC), and Transport Canada.

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