

- SESAMI APP: An Accessible Interface for Surface
- <sup>2</sup> Area Prediction of Materials from Adsorption
- 3 Isotherms
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#### **Software**

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### Statement of need

Surface area characterization is one of the most important material's characterization techniques for modern engineering application. The most widely used approach to calculate a material's gravimetric surface area, i.e. surface area per unit mass, is the Brunauer-Emmett-Teller (BET) method (Brunauer et al., 1938). The method computes the surface area given the adsorption isotherm of a probe gas (i.e., N<sub>2</sub> or Ar) in an adsorbent. Despite its importance, many researchers either obtain the BET area from commercial software that comes with equipment, or perform the analyses manually on a spreadsheet, which is time-consuming, and nearly impossible for some types of isotherms, which leads to large variability in BET-calculated areas (Osterrieth et al., 2022). These challenges have motivated the development of programs for the automated and standardized calculation of BET areas (Datar et al., 2020; lacomi & Llewellyn, 2019; Osterrieth et al., 2022; Sadeghi et al., 2020; Sinha et al., 2019). Furthermore, shortcomings of BET method as a tool for surface area characterization, such as its relatively poor performance in treating high surface area materials with multimodal pore sizes (Gómez-Gualdrón et al., 2016; Wang et al., 2015), have led to the development of alternate methods for surface area calculation from isotherms (Datar et al., 2020; Sinha et al., 2019).

## Theory background

- $_{\mbox{\tiny 28}}$  The surface area of a material is calculated using Equation 1. S is a material's surface area,
- $q_m$  is the molar amount of adsorbate forming a monolayer per unit mass of adsorbent, N is
- $_{\mbox{\tiny 30}}$  the Avogadro constant, and  $A_m$  is the area taken up by a single adsorbate molecule in the
- monolayer.

$$S = q_m N A_m \tag{1}$$

In order to attain  $q_m$ , as well as the BET constant, C, it is necessary to identify the monolayer loading region from the isotherm using the BET method (Fagerlund, 1973). This region is assigned to a section of the isotherm where  $\frac{p/p_0}{1-p/p_0} \cdot \frac{1}{q}$  is linear as a function of  $\frac{p}{p_0}$ , where p is the vapor pressure,  $p_0$  is the saturated vapor pressure, and q is the adsorbate loading. The linear region for BET analysis is usually chosen based on the consistency criteria proposed by Rouquerol et al. (F. Rouquerol et al., 2013; J. Rouquerol et al., 2007). The consistency criteria are as follows:



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- 1. The linear region should only be a range of  $p/p_0$  in which the value of  $q(1-p/p_0)$  monotonically increases with  $p/p_0$ .
- 2. The value of the C should be positive.
  - 3. The value of the monolayer loading capacity should correspond to a value of  $p/p_0$  which falls within the selected linear region.
- 4. The value of  $p/p_0$  calculated from BET theory,  $1/(\sqrt{C}+1)$ , and  $p/p_0$  calculated from the third consistency rule should be equal (with  $\pm 10\%$  tolerance).
- 5. The linear region should end at the knee of the isotherm.

Once a linear region is selected, the identified uptake value is multiplied by the molecular cross-sectional area of the adsorbate, typically derived from the bulk liquid density (16.2  $Å^2$ /molecule for  $N_2$ ; 14.2  $Å^2$ /molecule for Ar), to obtain the material's surface area, under the assumption that the adsorbate molecules only form a monolayer (Equation 1). The BET method is based on the BET equation (Equation 2).

$$\frac{p/p_0}{q(1-p/p_0)} = \frac{1}{q_m C} + \frac{C-1}{q_m C} \frac{p}{p_0}$$
 (2)

# Summary

The SESAMI web interface allows a user to make surface area predictions on their web browser simply by uploading isotherm data. The website facilitates access to the previously developed SESAMI models (SESAMI 1 and 2) for the evaluation of material's surface area (Datar et al., 2020; Sinha et al., 2019). The motivation for this interface is to lower the barrier of entry for research groups seeking to use SESAMI code, which was previously packaged in Python and Jupyter Notebook scripts.

SESAMI 1 applies computational routines to identify suitable linear regions of adsorption 63 isotherms for BET surface area predictions (Fagerlund, 1973). The automated workflow includes consideration of Rouquerol criteria and the use of coefficients of determination as a measure of linearity. Furthermore, SESAMI 1 supports a combined BET+ESW (excess sorption work) approach for linear region selection; this combined approach has been shown to outperform the BET method in some cases (Sinha et al., 2019). A user can specify a cutoff R<sup>2</sup> and a minimum R<sup>2</sup>, such that a candidate linear region is favored to be selected if it has an R<sup>2</sup> above the cutoff, and a candidate linear region is only considered if it has an R<sup>2</sup> above the minimum. On the other hand, SESAMI 2 applies a machine learning (specifically, regularized linear regression with LASSO) model for the accurate surface area prediction of high surface area materials, improving on BET performance for these materials (Datar et al., 2020). The LASSO model uses as input the average loading in seven isotherm pressure regions as well as pairwise products of these loadings. The SESAMI routines support isotherms with  $N_2$  and argon adsorbate at 77 K or 87 K, respectively. We note that a recent study shows that surface areas determined from  $N_2$  or Ar isotherms are similar, despite the latest 2015 IUPAC report's suggested use of Ar (Datar et al., 2022; Thommes et al., 2015). In addition, the SESAMI 1 code supports arbitrary user-specified adsorbates if their cross-section, saturation vapor pressure, and temperature are specified.

The SESAMI web interface has extensive error handling and clearly alerts users of issues with their adsorption isotherm data. For example, it alerts the user if no ESW minima is found by SESAMI 1 or if the data is incompatible with SESAMI 2 code due to data sparsity in certain pressure regions. As shown in Figure 1, the interface displays SESAMI 1 calculation results including information on the chosen linear region, namely the satisfied Rouquerol



criteria, the pressure range and number of data points in the region, and the coefficient of determination. The interface also displays intermediate SESAMI 1 values for surface area calculation, namely C and  $q_m$ . Furthermore, the SESAMI web interface allows the user to download figures generated by SESAMI 1 that indicate, among other things, the chosen linear monolayer loading regions by the BET and BET+ESW approaches as well as the excess sorption work plot (Figure 1). The user can convert output from commercial equipment to AIF format and upload the converted data to the interface for analysis. The SESAMI web interface is publicly available at <a href="https://sesami-web.org/">https://sesami-web.org/</a>, and source code is available at <a href="https://github.com/hjkgrp/SESAMI\_web">https://github.com/hjkgrp/SESAMI\_web</a>.





### a) SESAMI 1.0 (BET) results are:

BET area = 3827.6 m²/g C = 33.08 q<sub>m</sub> = 39.23 mol/kg Rouquerol consistency criteria 1 and 2: Yes Rouquerol consistency criterion 3: Yes Rouquerol consistency criterion 4: Yes Number of points in linear region: 4 Lowest pressure of linear region: 9000 Pa Highest pressure of linear region: 30000 Pa

R<sup>2</sup> of linear region: 0.9996

#### SESAMI 1.0 (BET+ESW) results are:

BET area = 2861.8 m²/g
C = 265.3
q<sub>m</sub> = 29.33 mol/kg
Rouquerol consistency criteria 1 and 2: Yes
Rouquerol consistency criterion 3: No
Rouquerol consistency criterion 4: No
Number of points in linear region: 10
Lowest pressure of linear region: 750 Pa
Highest pressure of linear region: 3500 Pa
R² of linear region: 0.9989

#### SESAMI 2.0 (LASSO) surface area prediction is:

2944.2 m<sup>2</sup>/g

### Figure type

b)

Isotherm BET fit 60 BET+ESW fit BET region BET+ESW region 50 ESW minimum Consistency 1 maximum BET data points 40 q / mol/kg 30 20 10 0 10-8 10-4 10<sup>-2</sup> 10<sup>0</sup>  $p/p_0$ **Download figure** 

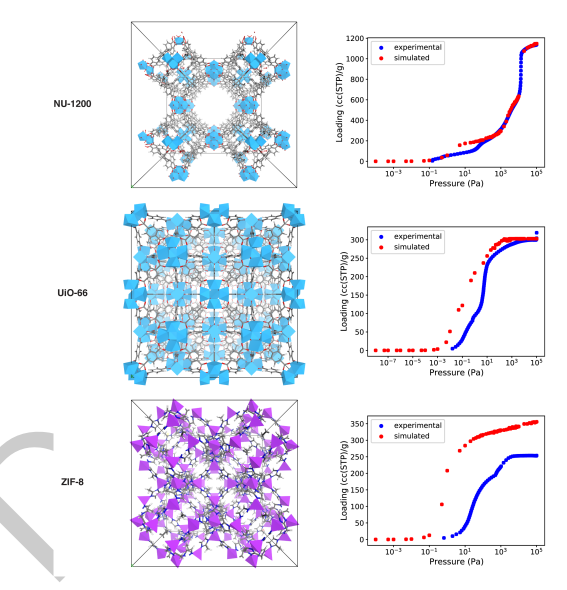
Figure 1: Information displayed by the SESAMI web interface after a calculation has been run, here for a GCMC isotherm of MIL-101. Apart from the inclusion of the LASSO prediction, default settings were used (e.g.  $N_2$  gas). a) Interface printout of information on the SESAMI 1 chosen linear regions, and SESAMI 1 and 2 calculation results. b) Figure download functionality for figures detailing the SESAMI 1 calculation.

## Benchmarking

- <sub>96</sub> To assess the performance of the SESAMI code in predicting surface areas from isotherms,
- $_{97}$  we benchmark the SESAMI routines against other similar programs for 13 simulated and 9
- $_{98}$  experimental  $N_2$  isotherms obtained at 77 K for 14 metal-organic frameworks (MOFs), some



of which are shown in Figure 2. Simulated isotherms are obtained from grand canonical Monte Carlo (GCMC) simulations using the open-source RASPA 2.0.47 software (Dubbeldam et al., 2016), and experimental adsorption isotherms are obtained from the experimental data reported by Islamogu and coworkers (Islamoglu et al., 2022). The data are then used to calculate the surface areas from the SESAMI website, BETSI (Osterrieth et al., 2022; Rampersad et al., 2020), pyGAPS (Iacomi, 2019; Iacomi & Llewellyn, 2019), and BEaTmap (Sadeghi et al., 2020).



**Figure 2:** The crystal structures and isotherms of 3 of the 14 MOFs used to benchmark different isotherm to surface area codes.

We find that over the set of 13 GCMC isotherms, the SESAMI machine learning model (run from the web interface) and BEaTmap have the best correlation with Zeo++ version 0.3 surface areas (Willems et al., 2012) calculated with a 1.67 Å probe  $N_2$  molecule (Tables 1 and 2). Nevertheless, all software are in generally good agreement, underscoring the benefit of a computational approach to surface area calculation. The agreement between software is also not surprising due to the similar approach taken by most of the codes of considering multiple subsets of consecutive data points and applying checks like the Rouquerol



criteria to select a linear region for BET analysis. Indeed, this agreement is also observed over the 9 experimental isotherms (Table 3). The benchmark isotherms, XLSX files of surface area predictions across different software tools for both GCMC and experimental isotherms, detailed settings used for each software, and analysis scripts employed are available at https://github.com/hjkgrp/SESAMI\_web.

**Table 1:** Calculated surface areas  $(m^2/g)$  for the 13 MOFs with GCMC isotherms. Cases where a software does not find a surface area are denoted by N/A. Zeo++ calculations are conducted with the same CIF files used to generate GCMC isotherms, and a 1.67 Å probe  $N_2$  molecule, the high accuracy flag, and 2,000 Monte Carlo samples per atom are used. All other software take as input the GCMC isotherms.

	SESAMI 1 (BET)	SESAMI 1 (BET+ESW)	SESAMI 2 (LASSO)	BETSI	py- GAPS	BEaTmap	Zeo++
HKUST-1	2001	1933	2089	1962	1902	1980	2397
IRMOF-1	3502	3543	3123	3519	3504	N/A	3722
MIL-100 (Cr)	2107	1853	2111	N/A	1852	2094	1957
MIĹ-100 (Fe)	2386	2438	2203	2426	82	2423	1933
MIĹ-101	3828	2862	2944	N/A	2939	3331	3164
MIL-53 (AI)	1221	1168	1405	1164	1183	1212	1510
MÓF-74 (Mg)	1828	1834	1902	N/A	1839	1791	1796
MOF-808	44	N/A	1275	N/A	N/A	1147	1690
NU-1000	2439	2181	2633	N/A	2144	2672	3050
NU-1200	2711	934	2601	N/A	1073	2930	3192
NU-1500	3543	3594	3111	N/A	3758	3492	3944
(Fe)							
UiO-66	1239	1239	1443	N/A	1242	1304	1289
ZIF-8	1429	1386	1575	1381	1390	1414	1588

**Table 2:** Comparison between surface area predictions from software for isotherm to surface area calculation and from Zeo++, over the 13 MOFs with GCMC isotherms. The mean absolute percent error and Pearson correlation coefficient are taken with respect to Zeo++ predictions for each software, over all successful surface area calculations for that software.

Software	Mean absolute percent error (MAPE)	Pearson correlation coefficient	Successful calculations (out of 13)
SESAMI 1 (BET)	19.4	0.85	13
SESAM1 1 (BET+ESW)	17.9	0.72	12
SESAMI 2 (LASSO)	12.4	0.95	13
BETSI	17.0	0.92	5
pyGAPS	23.0	0.75	12
BEaTmap	12.6	0.93	12



**Table 3:** Calculated surface areas  $(m^2/g)$  for the 9 MOFs with experimental isotherms. Cases where a software does not find a surface area are denoted by N/A. All other software take as input the experimental isotherms

	SESAMI 1 (BET)	SESAMI 1 (BET+ESW)	SESAMI 2 (LASSO)	BETSI	py- GAPS	BEaTmap
HKUST-1	1505	1466	1668	N/A	1495	1498
MOF-74 (Mg)	1580	1467	1692	N/A	1574	1565
MOF-808	1998	900	1727	N/A	2439	1752
NU-1000	2154	2090	2385	N/A	2654	2459
NU-1200	2893	2718	2781	2758	3917	3069
NU-1500 (Fe)	3305	3409	2809	N/A	3413	3227
SIFSIX-3 (Ni)	356	201	716	N/A	355	353
ÙiÓ-66	1251	1228	1413	1250	1249	1246
ZIF-8	1092	910	1214	N/A	1082	1047

**Table 4:** Settings used for software for isotherm to surface area calculation. All BET calculations by SESAMI 1 and pyGAPS reported in this work fulfill Rouquerol criteria 1 and 2.

Software	Mode of access	Settings		
SESAMI 1 (BET) SESAMI 1	Run from SESAMI web interface	Type of gas: Nitrogen Scope: BET and BET+ESW		
(BET+ESW)	Accessed February 2023	R <sup>2</sup> cutoff: 0.9995		
SESAMI 2 (LASSO)	Accessed 1 editary 2025	R <sup>2</sup> min: 0.998		
323/(WI 2 (E/1330)		Include ML prediction?: Yes		
BETSI	GUI started from the	Minimum number of points in the		
	command line	linear region: 10		
	GitHub version 1.0.20	Minimum R <sup>2</sup> : 0.998		
		Rouquerol criteria 1: Yes		
		Rouquerol criteria 2: Yes		
		Rouquerol criteria 3: No		
		Rouquerol criteria 4: No		
		Rouquerol criteria 5: No		
pyGAPS	Python package	Used function area_BET		
	Conda version 4.4.2	Default values for keyword arguments		
BEaTmap	Run from BEaTmap web	Adsorbate cross-sectional area: 16.2		
	interface	$ m \AA^2/molecule$		
	Accessed February 2023	Criteria 1: Yes		
		Criteria 2: Yes		
		Criteria 3: No		
		Criteria 4: No		
		Minimum number of data points: 5 BET calculation criteria: Maximum		
		data points		



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