Machine Learning Prediction of the Energy Gap of Graphene Nanoflakes Using Topological Autocorrelation Vectors

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ABSTRACT: The possibility of band gap engineering in graphene opens countless new opportunities for application in nanoelectronics. In this work, the energy gaps of 622 computationally optimized graphene nanoflakes were mapped to topological autocorrelation vectors using machine learning techniques. Machine learning modeling revealed that the most relevant correlations appear at topological distances in the range of 1 to 42 with prediction accuracy higher than 80%. The data-driven model can statistically discriminate between graphene nanoflakes with different energy gaps on the basis of their molecular topology.

KEYWORDS: graphene, band gap engineering, machine learning, molecular topology, topological autocorrelation vectors

The unique electric,1 magnetic, and optical properties2 of graphene, single atomic layers from graphite, have attracted considerable attention in recent years. By tuning of graphene properties, which are intrinsically related to its shape, size, and edge conformation, graphene sheets could be incorporated into a wide variety of electronics, optoelectronics, and electromagnetic devices. However, controlling the precise structure of individual graphenes remains challenging.3 To circumvent the need for exquisite control at the atomic level,4 computer simulations can elucidate how the nanostructure variability affects the functional properties.

The fact that graphene lacks a band gap around the Fermi level, which is the defining feature of semiconductor materials, limits the control of the conductivity of graphene by electronic means, hampering its revolutionary application in microelectronics. Theoretically, the band gap can be tuned by periodic modulations of the graphene lattice5 and control of the zigzag and armchair edges.6 Meanwhile, experimental evidence has been reported for band gap opening by patterned adsorption of atomic hydrogen onto the Moiré superlattice positions7 and manipulation of the width of graphene nanoribbons and hydrogen passivation on the edges.8 Alternatively, assembly of two-dimensional atomic crystals into stacks, i.e., graphene/boron nitride heterostructures, has emerged as a very promising system for band engineering of graphene.9

Experimental and theoretical evidence shows that differences in the electronic structures and associated properties of nanomaterials can be linked to discrepancies in physical structure, which can be measured using structural fingerprints or molecular descriptors. These fingerprints encoding topological, geometrical, or electronic features10 can be conveniently combined with machine learning techniques to unravel complex structure–property patterns and build accurate predictive models.10 Recently, we found that the energy gap \((E_G)\) between the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO) of graphene nanoflakes can be predicted from the interatomic distance distribution10 and geometrical features11 with accuracies of 70% and 90%, respectively. However, there is still room for further exploration of how the topology of the molecular graph can impact the energy gap of graphene nanoflakes.

In this Letter, we demonstrate that the topology of the molecular graph can accurately predict the energy gap of graphene nanoflakes regardless of atom position information, which could thereby accelerate the efficient rational design of functional graphene nanomaterials. More specifically, machine learning models have been developed to correlate graph topological fingerprints to the energy gap of graphene nanoflakes in quantitative terms. A virtual data set of 417 structures was used to calibrate correlation models with topological autocorrelation scores derived from graphene molecular graphs to predict the \(E_G\) of another set of 205 nanoflakes as a parametric function of the molecular graph topology. In brief, the remainder of this Letter is organized as follows: first, a description of the data set of graphene nanoflake structures and the topological autocorrelation descriptors used.
to calibrate the machine learning models is given, and then an account of how the predictive models were trained and validated is provided along with concluding remarks.

The influence of the graphene nanoflake topology on the size of the energy gap was explored using a data set of 622 virtual nanographene samples with a large range of sizes (16 to 2176 carbon atoms) that includes the ranges observed experimentally, which were simulated using the density functional tight binding (DFTB) method described elsewhere. The set of structures is available free of charge at the CSIRO Data Access Portal (DAP). The in

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The set of topological autocorrelation scores (ATS) described below is a very convenient way to describe the topology of graphene nanoflakes, as they are independent of the original atom numbering, so they are canonical and independent of the size of the molecular graph. Therefore, a substantial reduction in data by limiting the topological distance can be achieved, which allows the analysis of a large data set with a wide range of molecular size and chemical diversity. ATS and related descriptors were first proposed to relate the molecular graphs of organic compounds to their biological activities14 and later successfully extended to elucidate how the distribution of physicochemical features along protein sequences affects their conformational stability15 and ligand affinities.16

In Figure 1, the graphene nanoflake is represented as a molecular graph, which can be transformed into a numerical ATS code that is independent of the size of the structure. To differentiate between unpassivated and passivated edges, ATS values are weighted using bond order values of 2 and 3, respectively, while internal fully connected carbon atoms are assigned bond order values of 3. The ATS vectors can be interpreted as correlations of the bond order at different topological distances $L$ in the molecular graph according to eq 1:

$$\text{ATS}^L_i = \sum_{ij} N_{ij} \times P_i \times P_j$$

In order to simplify the autocorrelation indices, the summation in eq 1 runs over only the $N$ pairs of carbon atoms that are not fully connected in the molecular graph, i.e., edges and defects, while $d_{ij}$ is the topological distance or shortest path between carbon atoms $i$ and $j$, in this case the bond order of the carbon atoms in graphene, and $\delta_{ij}^L$ is a delta function defined as follows:

$$\delta_{ij}^L = \begin{cases} 1 & \text{if } L = d_{ij} \\ 0 & \text{otherwise} \end{cases}$$

In the case of graphene nanoflakes, the use of topological indices to characterize the molecular structure is particularly convenient considering that the edge topology and overall shape have been shown to influence the energy gap to a large extent.6 In this respect, connectivity information in graphene can be derived from experimental structure analysis of graphene films using high-resolution transmission electron microscopy (HRTEM),17 as has been successfully reported by McNerny et al.18

The ATS fingerprints averaged over hexagonal, rectangular, and trigonal graphene nanoflakes in Figure 2a illustrate differences among the graph representations for three shapes, which mainly differ in height, with rectangular nanoflakes showing the higher values, followed by trigonal and hexagonal structures in that order. Rectangular and trigonal ATS vectors show a maximum around topological distance of 20 steps, while hexagonal structures exhibit a maximum at around 30

Figure 1. Molecular graph representation of a graphene nanoflake with edges connected by solid lines and all other interior nodes connected by dashed lines.

Figure 2. (a) ATS averaged over hexagonal, rectangular, and trigonal graphene nanoflakes. (b) Histograms of the ATS in the optimum SVM models of the energy gap of graphene nanoflakes in 200 independent GA runs.
topological steps in the molecular graphs. However, the complexity of the ATS profiles suggests that sophisticated pattern recognition techniques rather than simple correlation
methods can correlate the $E_G$ values with a selected number of relevant ATS.

Simple regression models, namely, multiple linear regression (MLR) and binary decision tree (DT) calibrated on a training set of 70% of the data set, yielded cross-validation correlation coefficients of $\sim 0.73$ for the entire set of autocorrelation vectors (see the Supporting Information for details). Meanwhile, more sophisticated nonlinear mapping techniques, namely, support vector machines (SVM) and artificial neural networks (ANN), yielded cross-validation correlation coefficients of $\sim 0.77$ (see the Supporting Information for details on the machine learning models).

We further improved the machine learning models by exploring optimum combinations of ATS vectors simultaneously that evolve for different generations according to a genetic algorithm (GA) previously described elsewhere. The analysis of 100 independent GA runs in Figure 2b depicts that the most informative ATS for the energy gap appear at a topological distance range of 1 to 42.

Details of the “best” topological SVM model to predict the $E_G$ of graphene appear in Table 1, while the rest of the GA-optimized machine learning predictors and further details of the optimum SVM model appear in the Supporting Information. It is worth noting that the “best” model in Table 1 has 17 topological distance inputs in a topological distance input range from 1 to 87 and yields a cross-validation accuracy of $\sim 86%$. This fact suggests that experimental characterization of graphene nanoflakes and probably other nanomaterials can be focused on a reduced set of topological distances across the topological structure; characterization of entire structures with atomic-level precision is not necessary.

The ability of the SVM model to predict the $E_G$ of the test set of 30% of the data set is illustrated in the scatter plot in Figure 3, and the squared correlation coefficient ($R^2_{\text{cross}}$) and root-mean-square error ($\text{RMSE}_{\text{cross}}$) of test set predictions appear in Table 1. Remarkably, the $R^2_{\text{cross}}$ value matched the cross-validation accuracy in Table 1 with a value higher than 0.81 and a low $\text{RMSE}_{\text{cross}}$ of $\sim 0.46$ eV, which demonstrates that the machine learning model not only “learned” the topological pattern relevant for opening a gap but, more importantly, successfully generalized to new graphene structures.

The predictions of the highest values of $E_G$ in Figure 3 are less accurate, but the model correctly ranks the majority of their values. However, it is worth noticing that the topological approach exhibits test set accuracy that is $\sim 10$ percent units higher than that of a partial least squares model of the interatomic distance distributions. More importantly, with more than 80% accuracy, the topological predictions compare favorably to our recently reported machine learning model trained on a sophisticated combination of geometrical features, namely, aspect ratio, average carbon coordination number, edge conformation, and average bond distances and angles of graphene nanoflakes.

Considering that experimental comparison is not possible at the moment, our results for over 600 regular graphene nanoflakes with trigonal, hexagonal, and rectangular shapes and different degrees of passivation and edge conformation demonstrates that the size of the energy gap can be controlled to a large extent via the connectivity of the carbon atoms. This result suggests that different nanoflakes can be statistically discriminated into gap energy groups according to their topologies, which could guide engineering of nanoflake structures with specific topologies using, for example, scanning tunneling microscope lithography, which allows patterning of nanoribbons with well-defined widths and predetermined crystallographic orientations at nanometric precision.

In summary, a machine learning model calibrated with topological information successfully predicted $E_G$ values for graphene nanoflakes with a squared correlation coefficient higher than 0.8 and an absolute error lower than 0.5 eV. The topological model can be useful to rapidly estimate the energy gap while providing some rationale for tuning the topology of graphene nanoflakes. To the best of our knowledge, this is the first machine learning prediction of the energy gap of graphene nanoflakes solely from molecular graph information. Furthermore, this approach could accelerate the development of graphene nanotechnologies by guiding the synthesis and/or lithographic preparation of graphene nanoflakes with moderated control over molecular connectivity and edge characteristics, and in general it can help experimentalists to identify rational strategies for the optimization of not only graphene but also other two-dimensional materials. In general, data-driven models can be integrated into in silico HT platforms for more efficient computational screening and analysis of the relationship between functional properties and structural features and imperfections of large nanomaterials libraries.
ASSOCIATED CONTENT

Supporting Information
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acscombsci.6b00094.

Details of the DFTB calculations, machine learning implementation, calibration and testing, and outlier removal details (PDF)

ATS descriptors and $E_G$ values for the graphene data set (ZIP)

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Notes
The authors declare no competing financial interest.

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