Mechanical and thermal properties of graphene random nanofoams via Molecular Dynamics simulations

Andrea Pedrielli a, b, Simone Taioli b, c, Giovanni Garberoglio b, **, Nicola Maria Pugno a, d, e, *

a Laboratory of Bio-Inspired and Graphene Nanomechanics, Department of Civil, Environmental and Mechanical Engineering, University of Trento, Italy
b European Centre for Theoretical Studies in Nuclear Physics and Related Areas (ECT*-FBK), and Trento Institute for Fundamental Physics and Applications (TIFPA-INFN), Trento, Italy
c Faculty of Mathematics and Physics, Charles University, Prague, Czech Republic
d School of Engineering and Materials Science, Queen Mary University of London, UK
e Ket Lab, Edoardi Amaldi Foundation, Italian Space Agency, Italy

Article history:
Received 9 January 2018
Received in revised form 21 February 2018
Accepted 22 February 2018
Available online 27 February 2018

Abstract

Graphene foams have recently attracted a great deal of interest for their possible use in technological applications, such as electrochemical storage devices, wearable electronics, and chemical sensing. In this work, we present computational investigations, performed by using molecular dynamics with reactive potentials, of the mechanical and thermal properties of graphene random nanofoams. In particular, we assess the mechanical and thermal performances of four families of random foams characterized by increasing mass density and decreasing average pore size. We find that the foams’ mechanical performances under tension cannot be rationalized in terms of mass density, while they are principally related to their topology. Under compression, higher-density foams show the typical slope change in the stress–strain curve at 5 – 10% strain, moving from linear elasticity to a buckling region. At variance, lower density foams display a quasi-linear behavior up to 35% strain. Furthermore, we assess the thermal conductivity of these random foams using the Green–Kubo approach. While foam thermal conductivity is affected by both connectivity and defects, nevertheless we obtain similar values for all the investigated families, which means that topology is the critical factor affecting thermal transport in these structures.

1. Introduction

Recently, an increasing interest has been paid to nanoporous materials. Porosity, indeed, can strongly increase the surface-to-volume ratio and enhance the specific mechanical properties, such as the specific modulus and strength, with respect to the bulk material. For example, a high surface-to-volume ratio is desirable for gas adsorption and separation [1], while improving specific mechanical properties using carbon-based porous materials is of interest for building lightweight structural components [2].

Moreover, after the discovery of novel bi-dimensional materials [3], such as the hexagonal allotrope of boron nitride (h-BN) and graphene, several investigations have been focused onto the search of unconventional 3D structures that inherit the outstanding electrical [4,5], thermal and mechanical properties of their 2D counterpart in order to achieve specific requirements.

In particular, graphene shows excellent tensile properties, such as fracture strength (σ ≈ 130 GPa) and Young’s modulus (E ≈ 1 TPa) coupled with relatively low density due to its bidimensionality, and thus it is the best candidate material to be used in the synthesis of foam assemblies with superior properties. Graphene-based nanofoams can be synthesized by using Chemical Vapor Deposition (CVD) on metallic scaffold as well as nanoparticles assemblies [6–8] or chemically derived by reducing graphene oxide [9]. In mechanical and thermal applications, critical parameters are the concentration of defects, the topology as well as the inter-flake contacts. Despite this technological interest, only a few
computational investigations have been performed to characterize their electronic, thermal and mechanical properties [10–12]. In particular, mechanical properties of porous materials at microscale can be studied by the Ashby-Gibson theory, in which a unit cell approach is combined with dimensional analysis [13]. While this approach can be useful to derive scaling laws of mechanical properties with respect to density, the effective properties of porous materials are not often a simple function of porosity. At odds, they usually strongly depend on features at the nanoscale, where local atomic interactions start to play a crucial role, or on the presence of struts and of random pores with very special shapes that destroy structural periodicity. Furthermore, deformation mechanisms at the mesoscale can be very different at the nanoscale, where the fine details of graphene topology come into play, and a multiscale approach should be devised [2]. Additionally, it turns out that carbon-based nanoporous materials with random porosity distribution exhibit poor scaling of the mechanical properties with decreasing density, even more pronounced than that of metal and polymeric foams [14]. However, nanoporous graphene foams easily outperform polymeric foams at high density and can compete with their high-performance rivals, such as the metal foams. Thus, the interest in studying these random porous structures for energy storages and damping devices remains high.

Moreover, the high porosity of random foams suggests a concurrent application of these materials as thermal insulators. In particular, our goal is to assess the dependence of the thermophysical properties on pore density and size, and to compare thermal insulation performances of graphene-based 3D structures with other widely used carbon-based foams, such as polyurethane and metal foams.

This work is thus aimed at shedding some light on the mechanical and thermal properties of random graphene nanofoams. In particular, we present molecular dynamics (MD) simulations of random-pore foams under tension and compression by modeling atomic interactions via reactive potentials. Several random-pore foams, characterized by different density and porosity, are produced (called A, B, C, and D, see Fig. 1a, which are provided as xyz coordinate files with this submission). Each family is characterized by a different initial number of randomly-positioned support particles while, within each family, the only difference is the initial random distribution of the support points (Fig. 1a). During the whole procedure we impose periodic boundary conditions.

The LJ parameters used for the support (S) and foam (F) particles, respectively, are the following: \( \epsilon_{SS} = 100.0 \text{ eV}, \sigma_{SS} = 0.3 \text{ nm}, \epsilon_{FF} = 0.1 \text{ eV}, \sigma_{FF} = 0.32 \text{ nm}, \epsilon_{SF} = 0.23 \text{ nm}, \sigma_{SF} = 1.0 \text{ nm} \). The starting cell side length is 6.0 nm, expanded up to a length of 12.5 nm from step a) to step b) of Fig. 1. These parameters were chosen in such a way that the typical distance between the support particles was smaller than the equilibrium length between the foam and the support particles. In this way, the support particle surface is smooth, being obtained by several atoms lying nearby.

The topology of the nanofoams studied here is inspired by the graphene foams grown on nickel scaffolds (Fig. 2). However, we note that the pore size in the experimentally synthesized samples is larger than in our computational models. The way in which our graphene random foams are built is substantially different from that one presented in Ref. [14], where three-dimensional graphene assemblies were synthesized by starting from randomly distributed and oriented rectangular graphene flakes and spherical inclusions, and by repeating NPT-NVT cycles to obtain condensed graphene foams.

3. Characterization of graphene foams

Five different samples for each of the four families were prepared by varying the initial distribution of the support particles. In Fig. 3 we report representative models (3.0 nm slices) for each of these foam families. From A to D the foams present a decreasing average pore size, and an increasing mass density.

The geometrical analysis of the graphene porous foams and of their voids was carried out using the simulation code Zeo++ [16]. In particular, we characterize our prepared structures using the Pore Size Distribution (PSD) function, which can be experimentally obtained by adsorption/desorption measurements. PSD analysis delivers a quantitative description of the range of pore sizes present in a given sample.

Moreover, we perform a coordination analysis to find possible signature of under- or over-coordination of the carbon atoms usually forming a network of sp\(^2\) hybrid bonds. The computed quantities are reported in Table 1.

The averaged PSDs for all our graphene foam families are reported in Fig. 4 (continuous lines), showing the standard deviation within each group as a colored shaded area. By comparing these PSDs with those obtained in the case of regular pore foams [12, reported in Fig. 5, we notice that the random foams present similar average pore dimension and similar mass density. Indeed, the random foam PSDs are characterized by a maximum, representing the most likely pore size in each case, decreasing from 2.3 nm to 1.7 nm from A to D foam type. The average values of the pore size are reported in Table 1. These values vary reasonably well with
those reported for regular foams, where PSD peaks at about 2 nm (see Fig. 5). Finally, while regular foams present mass densities in the range 0.6 – 0.7 g cm$^{-3}$, our families of random foams have mass densities in the range 0.5 – 0.8 g cm$^{-3}$ (see Table 1). We notice that an almost linear relation (coefficient of determination $R^2 = 0.98$) with negative slope relates the mass density and the average pore dimension of the four random foam families, as reported in Fig. 6.

### 4. Computational methods

To perform MD simulations, carbon-carbon interatomic forces were modeled using the AIREBO potential [17]. To find the minimum energy structures with respect to defect positions, the samples were annealed at 3500 K, equilibrated at this temperature for 100 ps, and eventually cooled down to 700 K in 100 ps using a viscous damping force. The annealing was performed using the standard value for the cutoff parameter for the REBO part of the potential and performed within the microcanonical ensemble (NVE). For the simulations of compressive and tensile regimes, all samples were equilibrated at zero pressure and at the temperature of 1 K using the Nosé–Hoover barostat and thermostat. Furthermore, the adaptive cut-off parameter of the potential was set to 0.2 nm to better describe the near-fracture regime [18]. The equations of motion were integrated via the velocity-Verlet algorithm with time step of 1 fs. Mechanical properties were assessed in the isobaric-isothermal ensemble (NPT), adding a drag term to smooth out the pressure oscillations.

A uni-axial tensile strain was applied up to sample fracture in each case. The strain parallel to the direction of deformation is...
defined as

\[ \epsilon = \frac{L - L_0}{L_0} = \frac{\Delta L}{L} \tag{1} \]

where \( L_0 \) and \( L \) are the starting and current length of the sample in the direction of loading. To determine the stress, the pressure stress tensor components in response to the external deformation are computed as \[19\]:

\[ P_{ij} = P_{N_k m_k v_k v_i} + \sum_k \frac{r_{kj}}{V} \tag{2} \]

where \( i \) and \( j \) label the coordinates \( x, y, z \); \( k \) runs over the atoms; \( m_k \) and \( v_k \) are the mass and velocity of \( k \)-th atom; \( r_{kj} \) is the position of \( k \)-th atom; \( f_{kj} \) is the \( j \)-th component of the total force on the \( k \)-th atom due to the other atoms; and, finally, \( V \) is the volume of the simulation box.

The pressure in Eq. (2) includes both kinetic energy (temperature) and virial terms. Notice that the forces appearing in Eq. (2) are the sum of the pairwise, angle, dihedral, improper and long-range contributions. The computed stress is the true stress because the pressure is measured with respect to the instantaneous section area of the samples. The uni-axial compressive strain was applied up to reaching 35% total strain. The applied strain rate is chosen equal to \( 0.001 \text{ ps}^{-1} \), that we tested appropriately for ensuing convergence in the case of regular nanofoams \[12\]. Stress and strain were saved every 1000 time steps.

The stress–strain curve was computed at 1 K, since MD is usually computationally faster than minimization procedures. The same approach has been previously adopted by other groups dealing with similar problems (see e.g. Ref. \[11\]). While thermal fluctuations of the order of a few K, thus higher than the absolute value of the thermostat temperature, are found during MD runs,
they do not significantly affect numerical MD simulations. Indeed, we demonstrated in a previous work [1] that the contribution of the kinetic energy to the pressure tensor at a few K is approximately 2% of the total. Thus, a small kinetic contribution due to using low temperature MD does not prevent our simulations from reaching and overpassing local minima. The use of low-temperature MD simulations was chosen instead of standard minimization procedures because stress, for example due to compressive load, can be more effectively and more continuously redistributed within the entire structure during the dynamics by applying a deformation rate (providing this rate delivers converged results with respect to its value) instead of using a sequence of deformation-minimization steps. Indeed, under compressive strain the temperature is likely to increase: coupling the system with a thermostat leaks away this excess of thermal energy and allows for a minimization of the structures by using MD.

The observables that we calculate to characterize the mechanical properties of the nanofoams are the Young modulus, fracture stress and fracture strain. The toughness is also evaluated as the area under the stress-strain curve up to the fracture stress. Indeed, the samples have no plastic deformation but several sequential fractures. Stress-strain characteristics of carbon random nanofoams present a linear behavior at low strain. Thus, the Young modulus is obtained as the tangent at zero strain.

We also performed the calculation of the Poisson ratio $\nu$, defined as the negative ratio between the transverse $\Delta_T$ and the longitudinal deformations $\Delta_L$:

$$\nu = \frac{\Delta_T}{\Delta_L}$$

Here we extend the concept of Poisson ratio to deformations beyond the linear regime, and use it to quantify the lateral deformation of the material. A similar extension is done for the Young modulus.

Phonon thermal conductivity was assessed using the equilibrium Green-Kubo approach [20,21] since it is less sensitive to the simulation box dimension than non-equilibrium MD methods [22]. To this aim, first the atomic positions were relaxed and equilibrated at 300 K using the Berendsen thermostat method (NVT ensemble).

Then, in the NVE ensemble, the equilibrium thermal conductivity $k$ according to the Green-Kubo formalism, can be calculated as follows:

$$k = \frac{V}{3k_B T^2} \int_0^\infty \langle \vec{J}(0) \cdot \vec{J}(t) \rangle dt$$

where $V$ is the volume of the simulation cell, $t$ is the correlation time, and $k_B$ is the Boltzmann constant. The heat current $\vec{J}$, appearing in Eq. (4), is defined by:

$$\vec{J} = \frac{1}{V} \left( \sum_i E_i \vec{v}_i + \frac{1}{2} \sum_{i<j} \left( \vec{F}_{ij} \cdot (\vec{v}_i + \vec{v}_j) \right) \right)$$

where $\vec{v}$ is the velocity of a particle, $\vec{r}_{ij}$ and $\vec{F}_{ij}$ are the distance and force between the particles $i$ and $j$, and $E_i$ is the total energy per atom. The first term in the right hand side corresponds to convection, while the second term to conduction. The integrand in the expression for thermal conductivity is the heat current auto-correlation function (HCACF). To get a proper sampling of the phase space multiple runs are required with different initial conditions. Simulations to obtain MD trajectories to perform accurate ensemble averages were performed over a time span of 500 ps, using a step of 0.5 fs. HCACF has been computed by dividing the total time of computation into 250 fs beads and by performing the integral in Eq. (4) by sampling every 5 fs. Finally, we average over all the beads.

The thermal conductivity was calculated by using a version of the Tersoff potential [23] optimized to reproduce accurately the experimental phonon dispersion curves and the thermal properties of carbon structures, such as graphene and graphite. MD simulations were carried out using LAMMPS [24]. Atomic configurations were visualized by using the OVITO package [25] or VMD [26].

5. Results and discussion

5.1. Tension

In Fig. 7, we report the stress–strain characteristics for the four foam families investigated in this work, while in Fig. 8 the stress–strain curves are normalized with respect to the mass density.

The stress–strain curves show a typical elastic behavior for small deformations up to the tensile strength, followed by a

![Fig. 7. Stress–strain curves of the four graphene foam families under uni-axial tension along with the standard deviation for each family, reproduced by a colored shaded area on the top of the relevant curve. (A colour version of this figure can be viewed online.)](image-url)
curves should nearly overlap. Indeed, if the mass density were the factor most critically affecting density, such as the pore size distribution or the connectivity. Four foam families are due basically to features other than mass density: 0.366 g/cm³, Young modulus: 2.8 GPa), which compares to 3D graphene assemblies reported in Ref. [14]. Graphene assemblies have a specific Young modulus of 7.65 MNm kg⁻¹ (mass density: 0.366 g/cm³, Young modulus: 2.8 GPa), which compares with the lowest mass density foam family studied here (see Table 2). At variance, the specific strength of 7.4 MNm kg⁻¹ found in graphene assemblies is only slightly higher than in our random foams, mainly due to the higher connectivity of graphene sheets composing the assemblies. For completeness, we notice that the mechanical tests reported in Ref. [14] have been performed at a temperature of 300 K, while our simulations are performed at 1 K. Movies of our foams under tensile load are provided with this submission.

5.2. Compression

In this section we present the results obtained for our samples under compressive load. In Fig. 9 we report the stress–strain curves for the four foam families. The maximum deformation reaches 35% strain for the largest compression load. Beyond 35% strain the foams are mechanically unstable. From Fig. 9, we observe that at small strain foams are in the elastic regime, and the material is characterized by a full recovery to the original shape when the load is removed. Subsequently, we find a region with a slope similar for all our foam families, which models the structural collapse at a nearly constant stress by buckling or fracture of the building blocks.

Finally, at higher strain (not shown) one finds a steep ramp in the stress–strain curve, representing the complete collapse of the structures. The random foam family A, characterized by the lowest density, presents this ramp at 70% compressive strain. At variance, higher density random foams are not stable under compression before their respective final ramps, and present a structural transition from graphene ordered layers to amorphous carbon, with a strong stress decrease followed by an increase.

The most visible mechanical characteristics of our families of random foams (see the stress–strain curves in Fig. 9) is that, with increasing foam density and decreasing pore size average dimension, the elastic part presents an increasing slope, while beyond 5–10% strain the slopes are very similar. This behavior suggests a change in the compression mechanism: below 5–10% strain the stress is mainly due to the connectivity among graphene layers and this regime is characterized by structural stability, while beyond that the structures start collapsing with a relatively small increase of the stress, due to the bending of the graphene sheets inside the foams.

The foam family with the lowest mass density presents an almost linear stress–strain characteristic. This suggests that the collapse is dominated by bending. In higher density foams the slope change is more marked, showing that bending of graphene sheets occurs at higher strain.

We notice that the similarity of the slope of the stress–strain curves between 10% and 30% strain is due to a similar mechanism for collapse. This similarity can be rationalized by observing Fig. 10, where a 2.0 nm slice of a sample from the foam families A and D under 12% compressive strain is reported. As by Fig. 9 this strain value sets the transition between the elastic regime and the collapsing region. This transition for foams of higher porosity, such as those belonging to the type A, is related to the closure of the interstitial space when graphene sheets touch upon, as those highlighted by red circles.

Table 2

<table>
<thead>
<tr>
<th>Foam type</th>
<th>Young modulus (GPa)</th>
<th>Standard deviation (GPa)</th>
<th>Tensile strength (GPa)</th>
<th>Standard deviation (GPa)</th>
<th>Fracture strain (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>3.9</td>
<td>2.1</td>
<td>0.5</td>
<td>0.3</td>
<td>19</td>
</tr>
<tr>
<td>B</td>
<td>14.0</td>
<td>3.1</td>
<td>1.5</td>
<td>0.5</td>
<td>15</td>
</tr>
<tr>
<td>C</td>
<td>27.3</td>
<td>3.7</td>
<td>3.1</td>
<td>0.4</td>
<td>12</td>
</tr>
<tr>
<td>D</td>
<td>36.6</td>
<td>1.9</td>
<td>4.6</td>
<td>0.3</td>
<td>13</td>
</tr>
</tbody>
</table>
In Fig. 11 we report (as performed in the case of tensile load) the specific stress–strain curves, which are normalized per mass density. Similarly to the tensile case, the stress–strain characteristics are almost unaffected by this normalization, once more suggesting that the mechanical performances of random foams are related to the connectivity and the topology of the foams rather than to the mass density.

Moreover, under compression the behavior of the random foam families C and D, characterized by higher density, is very similar to that of regular nanofoams presented in Ref. [12] with comparable specific stress. At variance, lower density random foams present an almost linear behavior under compression that differs from that of regular nanofoams. Young modulus and collapsing region slope for the four random foam families under compressive load are reported in Table 4.

Finally, in Fig. 12 we report the stress–strain curves of the four random foam families, initially loading the samples up to 15% strain and subsequently unloading them. At odds with regular foams that can fully recover their initial shape when unloaded after reaching high deformations (up to 25%) [12], the higher density of defects in random foams and the local concentration of stress cause an incomplete elastic behavior even for relatively small strain (15%). Movies of our foams under compressive load are provided with this submission.

5.3. Poisson ratio

To better characterize the four random foam families, we computed the Poisson ratio of these structures. The Poisson ratio for each sample is computed as the average in the two directions transverse to the loading. The Poisson ratios under uni-axial compression of the four foam families are reported in Table 3.

### Table 3

<table>
<thead>
<tr>
<th>Foam type</th>
<th>Specific modulus (MN m kg⁻¹)</th>
<th>Standard deviation (MN m kg⁻¹)</th>
<th>Specific strength (MN m kg⁻¹)</th>
<th>Standard deviation (MN m kg⁻¹)</th>
<th>Specific toughness (MJ kg⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>7.1</td>
<td>3.8</td>
<td>0.9</td>
<td>0.5</td>
<td>0.1</td>
</tr>
<tr>
<td>B</td>
<td>20.6</td>
<td>4.6</td>
<td>2.2</td>
<td>1.5</td>
<td>0.2</td>
</tr>
<tr>
<td>C</td>
<td>35.0</td>
<td>4.7</td>
<td>3.9</td>
<td>3.1</td>
<td>0.3</td>
</tr>
<tr>
<td>D</td>
<td>44.1</td>
<td>2.3</td>
<td>5.4</td>
<td>4.6</td>
<td>0.4</td>
</tr>
</tbody>
</table>

In Fig. 10 we report (as performed in the case of tensile load) the specific stress–strain curves, which are normalized per mass density. Similarly to the tensile case, the stress–strain characteristics are almost unaffected by this normalization, once more suggesting that the mechanical performances of random foams are related to the connectivity and the topology of the foams rather than to the mass density.

### Table 4

<table>
<thead>
<tr>
<th>Foam type</th>
<th>Young modulus (GPa)</th>
<th>Collapsing region slope (GPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>3.5</td>
<td>1.6</td>
</tr>
<tr>
<td>B</td>
<td>12.3</td>
<td>3.0</td>
</tr>
<tr>
<td>C</td>
<td>24.4</td>
<td>3.7</td>
</tr>
<tr>
<td>D</td>
<td>31.8</td>
<td>3.1</td>
</tr>
</tbody>
</table>
tension and compression regimes are plotted in Figs. 13 and 14, respectively. Notably, the Poisson ratios are positive over the whole deformation range for all our random foam families.

In particular, for near zero strain under tension, the Poisson ratio is in the range $0.1 - 0.25$ with values increasing with mass density and foam connectivity. Furthermore, up to in 15% strain there is a small increment ($0.2 - 0.05$) in the Poisson ratio for all the foams families. Finally, the Poisson ratio at higher tensile strain goes to zero. This behavior is explained as mainly due to the fracture of the samples in this deformation regime, which prevents the sample from further contraction.

At variance, under compression the decrease of the Poisson ratio is due to the internal rearrangement of the graphene layers. At higher strain (about 30%) the Poisson ratio ranges between $(0.05 - 0.1)$. It can be worth noting that in Figs. 13 and 14, the standard deviation, reported as a shaded area across the relevant curve, is significantly smaller for lower density random foam families than that for higher densities.

5.4. Thermal conductivity

The thermal conductivity was assessed for all the samples using the averaging procedure of the HCACF explained before. In general, HCACF dies off within 100 ps and subsequently oscillates. This makes possible the time bead division, discussed in sec. 4.

In Fig. 15 we plot the average of the integral of HCACF as a function of simulation time for each foam family. The thermal conductivity is given by the asymptotic values of the time-integrated HCACFs. These values for the four foam families, obtained by averaging in the range 400 – 500 ps, are reported in Table 5.

We notice that two major factors affect the thermal conductivity, that are the foam connectivity and the presence of defects. In the foam families studied here, we devise that the low connectivity found in lower density foams is counterbalanced by the higher number of three-coordinated atoms, while the opposite trends occurs in higher density foams. Indeed, foams with different connectivity (see Fig. 10 of the paper) and different number of defects, show comparable values of the specific thermal conductivity (thermal conductivity per density mass unit). This could be due to a number of reasons. Our most likely explanation of this finding is as follows: smaller pore sizes generally means higher density as reported in Fig. 6. Nevertheless, a smaller size of the pores implies a larger number of defects, due to a larger local curvature, that is a
lower number of three-coordinated carbon atoms. We remind that graphene, due its particular topology, has a very large thermal conductivity, so that it is clear that a high level of carbon three-coordination is connected to large thermal conductivity. The presence of a larger number of defects in high-density foams with small size pores should then decrease the thermal conductivity. Thus we can argue that mass density (or pore size) and number of defects counteract in the determination of the thermal conductivity. In particular, in Fig. 16 we plot the thermal conductivity as a function of the mass density for the families of four different foams, finding an almost linear relation between these two observables (coefficient of determination $R^2 = 0.98$). The thermal conductivity of random foams is similar to that of glass (1 Wm$^{-1}$ · K$^{-1}$) for lower density foams, with a small increase (1.5 Wm$^{-1}$ · K$^{-1}$) for higher density foams.

### 6. Conclusions

In this work, we investigated by means of MD simulations with reactive potentials the mechanical and thermal properties of graphene random foams with a topology experimentally achievable by growing graphene on stacked nickel nanoparticles. In particular, we tested the mechanical performances under tension and compression of four random foam families, characterized by different mass density and pore size distribution. The samples were prepared using a multi-step approach based on the Voronoi partitioning of a triangulated surface, obtained by tessellation of the simulation cell using a LJ potential forcing the carbon atoms towards a rigid support.

Under compression, we found the typical elastic deformation regime with a Young modulus significantly increasing with a decreasing average pore size dimension. A behavior, common to all the random foam families here studied, was found for compressive strain in the collapsing region, with a positive slope of the stress–strain curve similar for all the four foam families. For the lowest density random foam family the stress–strain characteristic is almost linear.

Finally, we calculated the Poisson ratio, a quantity used to assess the transverse response of materials to deformation, of these random foams. Under tension, the Poisson ratio is positive for all the random foam families, indicating a transverse contraction under tensile load. The values of the Poisson ratio under compression are again positive for all the considered strain and tend to stabilize as the strain increases.

As a major outcome of our computational analysis, we find that mechanical properties under tension are characterized by an overall decrease of Young modulus with respect to regular nanofoams, while a tensile strength of the same order of that found for regular foams was obtained for higher density random foams.

Due to the interest of using foams as a mean for achieving thermal resistance, we computed the thermal conductivity of random foams using the Green-Kubo approach with a Tersoff potential optimized for these simulations. The thermal conductivity is comparable to that of glass, thus higher than materials typically used as thermal insulators, such as polyurethane rigid foams. Still, random foams do not display good thermal conductive properties, which can be related to the low connectivity in case of high porosity foams, and to the presence of defects in low porosity foams. Nevertheless, combining outstanding mechanical performances with light weight, low density and good thermal insulating properties, carbon random foams could be promising candidates as reinforcing fillers in nanocomposites or elastomers to tailor their properties or to replace polymer materials in applications where thermal stability and mechanical strength are needed.

### Acknowledgements

N.M.P. is supported by the European Commission H2020 under the Graphene Flagship Core 1 No. 666656 (WP14 “Polymer composites”) and under the Fet Proactive “Neurofibres” No.732344. S.T and G.G. acknowledge funding from previous WP14 “Polymer composites” grant. Access to computing and storage facilities owned by parties and projects contributing to the Czech National Grid Infrastructure MetaCentrum provided under the programme “Projects of Large Research, Development, and Innovations Infrastructures” (CESNET LM20150402), is greatly appreciated (https://www.metacentrum.cz/en/). The authors thanks Dr. V. Morandi and Dr. M. Christian for graphene foam SEM image and useful discussions.

### Appendix A. Supplementary data

Supplementary data related to this article can be found at https://doi.org/10.1016/j.carbon.2018.02.081.

### References


A. Pedrielli, S. Taïoli, G. Garberoglio, N.M. Pugno, Designing graphene based nanofoams with nonlinear auxetic and anisotropic mechanical properties under tension or compression, Carbon 111 (2017) 796—806.


