Material properties and structure of natural graphite sheet

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Natural graphite sheet (NGS) is compressible, porous, electrically and thermally conductive material that shows a potential to be used in fuel cells, flow batteries, electronics cooling systems, supercapacitors, adsorption air conditioning, and heat exchangers. We report the results of an extensive material characterization study that focuses on thermal conductivity, thermal diffusivity, electrical conductivity, coefficient of thermal expansion (CTE), compression strain, and emissivity. All the properties are density-dependent and highly anisotropic. Increasing the compression from 100 to 1080 kPa causes the through-plane thermal and electrical conductivities to increase by up to 116% and 263%, respectively. The properties are independent of the sheet thickness. Thermal and electrical contact resistance between stacked NGS is negligible at pressures 100 to 1080 kPa. In the in-plane direction, NGS follows the Wiedemann-Franz law with Lorenz number $6.6 \times 10^{-8} \frac{W m K^{-2}}{W m K^{-1}}$. The in-plane CTE is low and negative (shrinkage with increasing temperature), while the through-plane CTE is high, increases with density, and reaches $33 \times 10^{-6} K^{-1}$. Microscope images are used to study the structure and relate it to material properties. An easy-to-use graphical summary of the forming process and NGS properties are provided in Appendices A and B.

Natural graphite sheet (NGS) is a compressible, porous, electrically and thermally conductive material that has been used primarily to make sealing gaskets because of its ability to conform to rough surfaces, withstand high temperatures, and resist corrosive fluids. Presently, it is used—or considered to be used—in numerous applications, including bipolar plates in fuel cells and flow batteries for its electrical conductivity and chemical stability, supercapacitors for its corrosion resistance, adsorption cooling systems for its high thermal diffusivity, heat spreaders for its high in-plane thermal conductivity, heat exchangers for its thermal conductivity and corrosion resistance, and heat sinks for its thermal conductivity, low weight, and ease of forming into complex shapes. Multiple other names are used in the literature to refer to NGS, such as compressed exfoliated natural graphite (CENG), graphite foil, or flexible graphite. However, the term flexible graphite can be misleading because flexing thicker sheets results in a brittle fracture.

NGS is manufactured from natural graphite flakes in a process that involves soaking the flakes in an intercalation compound that penetrates between the graphite layers. During subsequent rapid heating, the vaporized compound forces the graphite layers apart, forming exfoliated natural graphite (ENG) particles also known as worms. NGS at densities 0.5 to 1.73 g cm$^{-3}$ is formed by compressing ENG particles at pressures ranging from 1 to 31 MPa.

In this study, we measure an extensive set of material properties and relate them to the structure observed by an SEM microscope. The sheet density is the primary parameter with the strongest influence on the properties. We found the through–plane thermal and electrical conductivity to increase with through–plane compression, and observed that the increase is density-dependent. The thickness of the sheet did not affect the measured properties.

The through–plane conductivities are low (2 W m$^{-1}$ K$^{-1}$ to 6 W m$^{-1}$ K$^{-1}$ and 3 Scm$^{-1}$ to 25 Scm$^{-1}$) while the in–plane ones are high (100 W m$^{-1}$ K$^{-1}$ to 350 W m$^{-1}$ K$^{-1}$ and 50 Scm$^{-1}$ to 1700 Scm$^{-1}$). In the in–plane direction, both thermal and electrical conductivities increase linearly with density. At low densities, our results agree with the previous studies, but at high densities we observed values 20 % to 25 % lower than those reported in the literature. In the through–plane direction, the conductivities decrease with density. The data from previous studies show high scatter and incomplete coverage of the density spectrum, which prevented us from forming a clear comparison with our results. The thermal and electrical contact resistance between stacked sheets

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of NGS is negligible at pressures higher than 100 kPa. We found the emissivity at wavelengths from 2 to 26 μm to decrease with density in the range 0.39 to 0.52.

When heated from room temperature to 100°C, NGS shows a dimensional stability superior to conventional materials such as aluminum or copper; however, the expansion in the through-plane direction is high and strongly dependent on the density. Our measurements show a very low and negative coefficient of thermal expansion (CTE) in the in–plane direction (−0.6 × 10⁻⁶ K⁻¹ to −1.4 × 10⁻⁶ K⁻¹) while in the through-plane direction it is positive and ranges from 10 × 10⁻⁶ K⁻¹ to 33 × 10⁻⁶ K⁻¹. Coupling the results with our study of the compression behaviour (outlined here and detailed elsewhere) allows for a quantitative analysis of thermomechanical stresses in devices such as fuel cells.

The structure of NGS is composed of tens-of-nanometers thin, highly anisotropic graphitic features whose orientation, spacing, and contact quality dictate the overall macroscopic properties of NGS. Compressing NGS causes reversible and irreversible structural changes that reflect in the overall properties. The irreversible changes happen during the forming process, and the reversible ones during the low-pressure loading of a finished sheet.

In summary, the goal of the present study is to compile a comprehensive resource describing the NGS properties and their relationship with the material structure. While the authors believe that this is the most comprehensive resource to date, we clearly state the need for further research. The most pressing issue is the lack of understanding of how the parameters of exfoliation methods and raw graphite flakes affect the structure and material properties of NGS.

Results

We first describe the structure of NGS, and then address each of the measured material properties. The forming process is not covered in this publication, however, a brief summary is provided in Appendix A. The reader is referred to other publication for details. Alongside the detailed description of the measurements and results further in this section, a simplified graphical summary of the measured material properties with explicit formulas of best fits is provided in Appendix B.

Structure. The structure of NGS is a result of compaction of ENG particles that are composed of highly crystalline cell walls as can be seen in Appendix A. The cell walls are 15 nm to 60 nm thick and their dimensions in the a and b crystal directions are equal to or less than the size of raw flakes. Upon compression, the cell walls within a single ENG particle deform into a platelet shape but the particles remain cohesive.

The cross-section structure at macro scale (hundreds of micrometers, Fig. 1a) changes significantly with the sheet density. At low densities, it is heterogeneous with large slit shape pores, and becomes homogeneous at higher densities. Stitched images in Supplementary Figure S1 do not show the through-plane density variation predicted by Bonnissel et al. At the scale of tens of micrometers (Fig. 1b), the microscope images show layers of flattened ENG particles whose preferred orientation increases with increasing density. Previous X-ray diffraction studies stated that the mean angle of graphite basal planes with respect to the sheet plane varies from 9° to 20° and that its change with sheet density is low at densities higher than 0.5 g cm⁻³. Assuming that the the linear features in Fig. 1b correlate with the graphite basal planes, the observed angles are qualitatively consistent with the previous work. The highest-magnification images at a micrometer scale shown in Fig. 1c reveal thin features whose thickness and spacing decrease with increasing density. The decrease of thickness with increasing density suggests that the features are not crystalline because graphite crystals are not expected to undergo permanent thickness reduction under compression at pressures relevant to NGS. The achieved level of magnification prevents forming definitive conclusions, however, we interpret the thin features as clusters of cell walls whose porosity allows for their thickness to be reduced by the pressure during the forming process.

The images of sheet faces shown in Fig. 1d,e indicate that the surface structure changes from rough to smooth with increasing density. The boundaries of the ENG particles are visible in the low magnification image of the 0.55 g cm⁻³ and 1.05 g cm⁻³ sheets. At higher densities, the surface appears uniform at low magnifications, and signs of overlapping cell walls are apparent in the high magnification images.

For relating the structure of NGS to its material properties, it should be noted that crystalline graphite is highly anisotropic. The orientation of crystalline cell walls dictate the anisotropy of NGS. At high NGS densities, the cell walls are aligned parallel to the sheet plane, which leads to the properties closer to those of a graphite crystal. However, due to porosity, imperfect contact between the cell walls, crystal defects, and other imperfections, the properties never reach those of a graphite crystal.

Material properties. The measured in-plane thermal and electrical conductivities (kth and σth) increase linearly with density as is shown in Fig. 2a,c. In the through-plane direction, the thermal and electrical conductivities (kth and σth) decrease with density as can be seen in Fig. 2b,d. Both kth and σth increase with through-plane compression pressure. For kth, the increase is not significant at low densities, but becomes progressively higher with increasing density. For kth, the increase is approximately constant over the entire measured density range. The experimental methods employed in this study are not suitable for measuring kth and σth at pressures below 100 kPa due to the high and unpredictably variable contact resistance between sensors and samples. Attempts of low-pressure measurements indicated a steep decrease in kth and σth, but future measurements using novel methods are necessary to confirm or reject the observed trend. Measurement of kth at varying pressures did not reveal a pressure dependence, and the in-plane electrical conductivity σth was measured in uncompressed state only.
Within the measured range of densities, pressures, and thicknesses, the thermal and electrical conductivities do not vary significantly with the sheet thickness, which can be supported by the overlap of the triangle and circular symbols in Fig. 2. The different sample thickness was achieved by using NGS at three surface densities ($d_s$) of 70, 140 and 210 mg cm$^{-2}$. Since stacks of NGS were measured in the through-plane direction, the insignificant variation with sheet thickness also suggests that the electric and thermal contact resistance at the sheet-to-sheet interfaces is negligible within the measured pressure range 100 kPa to 1080 kPa.

The low density of NGS results in high thermal diffusivity ranging from 230 to 270 mm$^2$s$^{-1}$ in the in-plane direction ($\alpha_{in}$) and from 1.5 to 12 mm$^2$s$^{-1}$ in the through-plane direction ($\alpha_{th}$). The plots of thermal diffusivity versus density are given in Appendix B.

The magnitude of thermal and electrical conductivity is dictated by the conductivity and orientation of the cell walls and the contact resistance between them. In the in-plane direction, the increase of $k_{in}$ and $\sigma_{in}$ with density arises from the decrease of cell wall misalignment and the decreased contact resistance due to improved contact quality or formation of new contact points. In the through–plane direction, the observed decrease of $k_{th}$ and $\sigma_{th}$ is contrary to the structure features seen in Fig. 1a,c. Closing of the slit–shaped pores and the decrease of the cell wall spacing with increasing density is expected to increase the through-plane conductivities. However, since the measured trend is opposite, the decrease in cell wall misalignment must be the dominating factor.

Besides the density and through-plane compression pressure, the material properties of NGS have been shown to be affected by the apparent density of ENG particles $d_{ENG}^{17,22}$, flake size$^{44}$, impurity content$^{14,45}$, crystal...
defects, rolling direction, or heat treatment. A comparison of the present thermal and electrical conductivities with the literature values is given in Supplementary Figures S2 and S3, but since the relevant parameters are varying or have not been reported, it is not possible to reliably explain the reasons for observed differences. Moreover, different measurement methods were used across the literature sources, which further broadens the list of possible sources of the discrepancies. Extensive future work is required to evaluate the sensitivity of NGS properties to the parameters listed above.

The detailed compression behavior of $k_{th}$ and $\sigma_{th}$ together with the through-plane stress-strain compression curves for three selected densities is shown in Fig. 3a–c. The change of $k_{th}$ with pressure is qualitatively proportional to the through-plane compression strain $S_{th}$. Both $k_{th}$ and $S_{th}$ of the 0.55 g cm$^{-3}$ sheet show a linear increase with pressure. At higher densities, $k_{th}$ and $S_{th}$ become progressively nonlinear with a higher rate of change at low pressures. The through-plane electrical conductivity $\sigma_{th}$ does not show the same proportionality to $S_{th}$ as $k_{th}$. The relative increase of $k_{th}$ and $\sigma_{th}$ at 1080 kPa with respect to the values at 100 kPa was plotted in Fig. 3d. The pressure sensitivity increases with increasing density, and $\sigma_{th}$ is more sensitive to pressure than $k_{th}$. The increase of $k_{th}$ and $\sigma_{th}$ with pressure is likely caused by the elastic deformation of cell walls and the improved contact quality between them, as is shown in Fig. 3e. Contrary to the forming process, no significant change in the misalignment of cell walls is expected.

To verify the claim that NGS follows the Wiedemann–Franz law, Lorenz number was calculated and plotted in Fig. 4 where it is seen to remain constant at $6.6 \times 10^{-6}$ W$\Omega^{-1}$K$^{-2}$ in the in-plane direction, but increase in the...
through-plane direction. The Wiedemann–Franz law is therefore concluded to be valid in the in-plane direction, but not in the through-plane direction.

The in–plane CTE of NGS (CTE\textsubscript{gr}) shown in Fig. 5a is low, negative, and its magnitude increases with density. CTE in the through-plane direction (CTE\textsubscript{th}) is large, positive, and increases with density up to \(33 \times 10^{-6} \text{ K}^{-1}\) as is shown in Fig. 5b. The anisotropy and density dependence arises from the NGS structure, which was illustrated in Fig. 5c. At room temperature, the CTE of cell walls is \(-1.8 \times 10^{-6} \text{ K}^{-1}\) in the \(ab\) crystal directions (CTE\textsubscript{ab,gr}) and \(26 \times 10^{-6} \text{ K}^{-1}\) in the \(c\) direction (CTE\textsubscript{c,gr})\textsuperscript{41,42,43}. The deformation of a single cell wall with increasing temperature was outlined in Fig. 5d where the arrows symbolize the direction and magnitude of the dimension change. The CTE of NGS is dictated by the orientation and interaction of all cell walls. At high densities, CTE\textsubscript{th} approaches CTE\textsubscript{ab,gr} as expected based on the highly-oriented structure. At densities above 1.2 g cm\(^{-3}\), CTE\textsubscript{th} appears to be higher than CTE\textsubscript{c,gr}, which is contrary to the expectations. However, the measurement uncertainty

Figure 3. The pressure dependence of (a) through-plane electrical conductivity and (b) through-plane thermal conductivity with (c) through-plane compression strain. The relative increase of the through-plane thermal and electrical conductivities at 1080 kPa relative to 100 kPa is shown in (d). Only the 140 mg cm\(^{-2}\) samples are shown to improve plot clarity. A schematic of the change of NGS structure with through-plane compression is shown in (e).
Figure 4. Lorenz number of NGS in the in-plane and through-plane directions as a function of density. Only the results for 140 mg cm\(^{-2}\) samples are shown to improve plot clarity.

Figure 5. Results of CTE measurements in (a) in-plane and (b) through-plane directions. The relevant graphite crystal values \(CTE_{ab,gr}\) and \(CTE_{c,gr}\)\(^{41,42,43}\) are demarcated by the dotted lines. In (c) a schematic structure of NGS is shown, and (d) illustrates the deformation of a single cell wall with increasing temperature.
prevents a definitive conclusion. It is likely that the expansion of air in the pores or the residual intercalation agent between graphite layers can cause an increase in CTE beyond CTE_{C}. The total emissivity at faces of NGS for wavelengths 2 \mu m to 26 \mu m decreases with increasing density, from 0.52 at 0.55 g cm^{-3} to 0.39 at 1.7 g cm^{-3} as shown in Fig. 6a. The emissivity of graphitic materials has been previously related to the in-plane electrical conductivity \sigma_{in} and surface roughness. As shown in the simplified illustration in Fig. 6b, \sigma_{in} increases with increasing density, while the surface structure changes from rough to smooth. Both of the factors contribute to the decrease in emissivity, the former via the Hagen–Rubens relation, and the latter via an increased surface area and the small pores that act as radiation black bodies.

**Experimental methods**

In-plane and through-plane thermal conductivities were measured using the transient plane source method (Hot Disk TPS 2500S) in the slab and one-dimensional modes, respectively. Electrical conductivity was measured using a four probe method with the Raytech Micro Junior 2 micro ohm meter. The compression behavior was measured in a mechanical tester with circular compression platens 50 mm in diameter and 3 kN load cell (Bose Electroforce 3300 Series II). CTE was measured using a thermomechanical analyzer (TA Instruments TMA Q400). Emissivity was determined from reflectivity measurements, which were performed in accordance with the Method C of ASTM E408-13 using an infrared reflectometer (Surface Optics Corporation 400T). All measurements were performed at room temperature except for CTE, which was determined in the range 30 \degree C to 100 \degree C.

The samples were prepared by calendering a low-density 0.2 g cm^{-3} NGS into the desired density and thickness. NGS was supplied by Nano Carbon Technology CO. and its properties were fixed carbon content 99.27 \%, d_{ENG} 4 mg cm^{-3}, flake size composition 81 \% larger than 300 \mu m (50 mesh, before exfoliation), intercalation agent H_{2}SO_{4}+H_{2}O_{2}+KMnO_{4}, exfoliation temperature 950 \degree C, and exfoliation time 5 min. A number of samples was tested to capture the variability of the properties. For example, for the through-plane thermal conductivity, 20 samples were prepared for each of the measured densities, resulting in the total of 240 samples. A random combination of 20 samples was measured at least three times at each of the compression pressures, which resulted in the total of 222 measurements. The density of the samples was determined by measuring the dimensions and weight using an OHAUS AX124 scale and a Darson Instruments 0-1” analog micrometer. The samples for cross-section SEM images were prepared by impregnating NGS in epoxy resin, polishing, and removing the smeared flakes by plasma etching. FEI Nova NanoSEM microscope was used for imaging. The compression behavior measurements were reported elsewhere. Details about the measurement methods, sample preparation, and uncertainty analysis were addressed at length in the supplementary document. The raw data, data processing scripts, microscope images in high resolution, and results in a tabular form are available in the supplementary dataset.
Appendix A

Forming of natural graphite sheet

1) Natural graphite flakes

2) Exfoliated natural graphite (ENG) particles

3) Compression forming

Recommended design and manufacturing process:
1) Determine desired free-standing density \( \rho \) based on considered application and NGS properties in Appendix B
2) Determine desired free-standing thickness \( t \) [mm]
3) Calculate (or estimate graphically) surface density: \( \sigma = d \frac{t}{10} \)
4) Calculate the mass of ENG particles: \( m_{ENG} = \sigma A \)
5) Compress ENG particles at forming pressure: \( p_f = 0.35e^{2.6d} \)

4) Natural graphite sheet (NGS)
Appendix B

Summary of natural graphite sheet properties

Through-plane direction

In-plane direction

All properties are at room temperature
Specific heat capacity at 25°C $c_p=729.3$ J kg$^{-1}$ K$^{-1}$
Summary of natural graphite sheet properties

\[ d \text{ - free-standing density [g cm}^{-3}\text{]} \]

\[ CTE_{th} \]

\[ CTE_{in} \]

\[ S_{th} \]

\[ S_{in} \]

\[ k_{th} = (34.63 - 34.44d + 29.61d^2)p^{0.1015 - 0.000842d} \\
- 30.59 + 34.99d - 31.75d^2 \]

\[ k_{in} = 212d - 22 \]

\[ \sigma_{th} = 2667 - 3806d + 4.1113p + 2945d^2 - 1.795dp - 0.004019p^2 - 992d^3 \\
+ 0.9967d^2p - 0.0003251dp^2 + 1.877 \cdot 10^6p^3 \]

\[ \sigma_{in} = 1.167 \cdot 10^5d - 1.934 \cdot 10^4 \]

\[ \epsilon = -0.11d + 0.58 \]

\[ CTE_{th} = -119d^{-0.151} + 142 \]

\[ CTE_{in} = -0.132d^{0.45} - 0.6 \]

\[ S_{th} = (0.0026 - 0.014d + 0.0166d^2)(1 - e^{-30})^{0.03d^{14}} \\
+ (0.00012 - 0.000147d + 4.8 \cdot 10^{-5}d^2)(p - 30) \]

\[ \pm 0.4 \text{ [W m}^{-1}\text{K}^{-1}] \]

\[ \pm 15 \text{ [W m}^{-1}\text{K}^{-1}] \]

\[ \pm 150 \text{ [S m}^{-1}] \]

\[ \pm 10\% \]

\[ \pm 5\% \]

\[ \pm 18\% \text{ [K}^{-1}] \]

\[ \pm 0.2 \text{ [K}^{-1}] \]

\[ \pm 0.004 \text{ [J]} \]

\[ d \text{ - density [g cm}^{-3}\text{]}, p \text{ - pressure [kPa]}, \text{relationships valid for } 0.5 \leq d \leq 1.7 \text{ g cm}^{-3} \text{ and } 100 \leq p \leq 1000 \text{ kPa} \]

\[ \text{Through-plane direction} \]

\[ \text{In-plane direction} \]

All properties are at room temperature, CTE for the range 35 to 95 °C

Specific heat capacity at 25°C \( c_p = 729.3 \text{ Jkg}^{-1}\text{K}^{-1} \)
References

1. Qian, P. et al. A novel electrode–bipolar plate assembly for vanadium redox flow battery applications. Journal of Power Sources 175, 613 – 620 (2008). http://www.sciencedirect.com/science/article/pii/S0378775307017806.

2. Lourenssé, K., Williams, J., Ahmadpour, F., Clemmer, R. & Tasnim, S. Vanadium redox flow batteries: A comprehensive review. Journal of Energy Storage 25, 100844 (2019). http://www.sciencedirect.com/science/article/pii/S2352152X19302798.

3. Kopasz, J. P., Benjamin, T. G. & Schenck, D. Bipolar plate workshop summary report. Tech. Rep., Argonne National Lab (ANL), Argonne, IL (United States) (2017).

4. Blomquist, N. et al. Metal-free supercapacitor with aqueous electrolyte and low-cost carbon materials. Scientific reports 7, 39836 (2017).

5. Blomquist, N., Koppolu, R., Dahlström, C., Toivakka, M. & Olin, H. Influence of substrate in roll-to-roll coated nanographite electrodes for metal-free supercapacitors. Scientific Reports 10, 1–9 (2020).

6. Wang, L., Metcalf, S., Crripe, R., Thorpe, R. & Tamanait-Jeoh, Z. Thermal conductivity and permeability of consolidated expanded natural graphite treated with sulpheric acid. Carbon 49, 4812 – 4819 (2011). http://www.sciencedirect.com/science/article/pii/S0008622310015452.

7. Chung, D. & Takizawa, Y. Performance of isotropic and anisotropic heat spreaders. Journal of Electronic Materials 41, 2580–2587 (2012). https://doi.org/10.1007/s11664-012-2177-4.

8. Thermal Performance of Natural Graphite Heat Spreaders, vol. Advances in Electronic Packaging, Parts A, B, and C of International Electronic Packaging Technical Conference and Exhibition. 10.1115/IPACK2005-75073. https://asmedigitalcollection.asme.org/InterPACK-proceedings-pdf/InterPACK2005/4200579/453138679_1.pdf.

9. Jamzad, P., Kenna, J. & Bahrami, M. Development of novel plate heat exchanger using natural graphite sheet. International Journal of Heat and Mass Transfer 131, 1205 – 1210 (2019). http://www.sciencedirect.com/science/article/pii/S0017931018337098.

10. Norley, J., Tseng, J. J., Getz, G., Klug, J. & Fedor, B. The development of a natural graphite heat-spreader. In Seventeenth Annual IEEE Semiconductor Thermal Measurement and Management Symposium (Cat. No.1CH37189), 107–110 (2001).

11. Cermak, M., Faure, X., Sætæn, M. A., Bahrami, M. & Ordonez, M. Natural graphite sheet heat sinks with embedded heat pipes. IEEE Access 8, 80827–80835 (2020).

12. Kaburagi, Y., Aoki, H. & Yoshida, A. Thermal and electrical conductivity and magnetoresistance of graphite films prepared from aromatic polyniflode. Tansu 5–9 (2012).

13. Chung, D. D. L. A review of exfoliated graphite. Journal of Materials Science 51, 554–568 (2016). https://doi.org/10.1007/s10853-015-9284-6.

14. Dowell, M. & Howard, R. Tensile and compressive properties of flexible graphite foils. Carbon 24, 311 – 323 (1986). http://www.sciencedirect.com/science/article/pii/0008622386902320.

15. Cermak, M. & Bahrami, M. Compression behavior of natural graphite sheet. SN Applied Sciences 2, 357 (2020).

16. Chung, D. D. L. Exfoliation of graphite. Journal of Materials Science 22, 4199–4190. https://doi.org/10.1007/BF01132908 (1987).

17. Wei, X. H., Liu, L., Zhang, J. X., Shi, J. L. & Guo, Q. G. Mechanical, electrical, thermal performances and structure characteristics of flexible graphite sheets. Journal of Materials Science 45, 2449–2455. https://doi.org/10.1085/jmse-101-4126-y (2010).

18. Liu, R. et al. Anisotropic high thermal conductivity of flexible graphite sheets used for advanced thermal management materials. In 2013 International Conference on Materials for Sustainable Energy and Environment, vol. 1, 107–111 (2013).

19. Bonnissel, M., Luo, L. & Tondeur, D. Compacted exfoliated natural graphite as heat conduction medium. Carbon 39, 2151 – 2161 (2001). http://www.sciencedirect.com/science/article/pii/S000862230100032X.

20. Luo, X. & Chung, D. Electromagnetic interference shielding reaching 130 db using flexible graphite. MRS Proceedings 445, 235 (1996).

21. Chen, P. H. & Chung, D. Thermal and electrical conduction in the compaction direction of exfoliated graphite and their relation to the structure. Carbon 37, 538 – 550 (2014). http://www.sciencedirect.com/science/article/pii/S0008622310051324.

22. Afanasov, I. M. et al. Thermal conductivity and mechanical properties of expanded graphite. Inorganic Materials 45, 486–490. https://doi.org/10.1134/S0020168509050057 (2009).

23. Gu, J. L., Leng, Y., Kang, E. & Shen, W. C. Microstructure effect on mechanical properties of flexible graphite sheet. Engineering Fracture Mechanics (1985). http://citeeseerx.ist.psu.edu/viewdoc/summary?doi=10.1.1.417.1336.

24. Gu, J. et al. Fracture mechanism of flexible graphite sheets. Carbon 40, 2169 – 2176 (2002). http://www.sciencedirect.com/science/article/pii/S0008622300007571.

25. Ionov, S. G., Avdeev, V. V., Kuvshinnikov, S. V. & Pavlova, E. P. Physical and chemical properties of flexible graphite foils. Molecular Crystals and Liquid Crystals Science and Technology. Section A. Molecular Crystals and Liquid Crystals 340, 349–354 (2000). https://doi.org/10.1080/10587250008025491.

26. Luo, X. & Chung, D. Flexible graphite under repeated compression studied by electrical resistance measurements. Carbon 39, 985 – 990 (2001). http://www.sciencedirect.com/science/article/pii/S000862230000213X.

27. Chen, P.-H. & Chung, D. Viscoelastic behavior of the cell wall of exfoliated graphite. Carbon 61, 305 – 312 (2013). http://www.sciencedirect.com/science/article/pii/S0008622310004223.

28. Luo, X., Chung, R., Biller, B. C., Hoi, Y. & Chung, D. D. L. Electronic applications of flexible graphite. Journal of Electronic Materials 31, 535–544. https://doi.org/10.1007/s11664-002-0111-x (2002).

29. Chung, D. D. L. Flexible graphite for gasketing, adsorption, electromagnetic interference shielding, vibration damping, electrochemical applications, and stress sensing. Journal of Materials Engineering and Performance 9, 161–163. https://doi.org/10.1361/1059940077346105 (2000).

30. Gu, J. L., Cao, W. Q., Cao, N. Z. & Shen, W. C. Effect of microstructure on compressive properties of flexible graphite sheet. In The European Carbon Conference “Carbon 96”, 346–347 (1996).

31. Cézard, A., Schneider, S. & Maréché, J. Densification of expanded graphite. Carbon 40, 2185 – 2191 (2002). http://www.sciencedirect.com/science/article/pii/S0008622300007775.

32. Toda, H. et al. Compression and recovery micro-mechanisms in flexible graphite. Carbon 59, 184 – 191 (2013). http://www.sciencedirect.com/science/article/pii/S0008622313002273.

33. Spencer, H. Empirical heat capacity equations of gases and graphite. Ind. and Eng. Chem. 40, (1948).

34. Cermak, M., Collins, M. & Bahrami, M. Emissivity of natural graphite sheet. International Journal of Heat and Mass Transfer (2020). Under review.

35. Cermak, M. Natural graphite sheet heat sinks for power electronics. Ph.D. thesis, Simon Fraser University (2020).

36. Cermak, M., Perez, N. & Bahrami, M. Material properties and structure of natural graphite sheet - dataset. https://doi.org/10.20383/101.00216 (2020).

37. Cézard, A., Maréché, J. & Ferdin, G. Surface area of compressed expanded graphite. Carbon 40, 2713 – 2718 (2002). http://www.sciencedirect.com/science/article/pii/S000862230018135.

38. Cézard, A., Maréché, J. F. & Ferdin, G. Modelling of exfoliated graphite. Progress in Materials Science 50, 93–179 (2005).
39. Celzard, A., Mareche, J., Furdin, G. & Puricelli, S. Electrical conductivity of anisotropic expanded graphite-based monoliths. Journal of Physics D: Applied Physics 33, 3094 (2000).
40. Inagaki, M., Kaburagi, Y. & Hishiyama, Y. Thermal management material: Graphite. Advanced Engineering Materials 16, 494–506 (2014). https://onlinelibrary.wiley.com/doi/abs/10.1002/adem.201300418.
41. Pierson, H. O. 3 - graphite structure and properties. In Pierson, H. O. (ed.) Handbook of Carbon, Graphite, Diamonds and Fullerenes, 43 – 69 (William Andrew Publishing, Oxford, 1993). http://www.sciencedirect.com/science/article/pii/B9780815513391500086.
42. Kelly, B. T. B. Physics of graphite / B.T. Kelly. (Applied Science, London ; Englewood, N.J., 1981).
43. Tsang, D., Marsden, B., Fok, S. & Hall, G. Graphite thermal expansion relationship for different temperature ranges. Carbon 43, 2902 – 2906 (2005). http://www.sciencedirect.com/science/article/pii/S0008622305003374.
44. Leng, Y., Gu, J., Cao, W. & Zhang, T.-Y. Influences of density and flake size on the mechanical properties of flexible graphite. Carbon 36, 875 – 881 (1998). http://www.sciencedirect.com/science/article/pii/S0008622397001966.
45. Smalc, M., Norley, J., Reynolds, R. A., Pachuta, R. & Krassowski, D. W. Advanced Thermal Interface Materials Using Natural Graphite. In 2003 International Electronic Packaging Technical Conference and Exhibition, Volume 2, 253–261 (2003). http://proceedings.asmedigitalcollection.asme.org/proceeding.aspx?doi=10.1115/IPACK2003-35113.
46. Hooker, C. N., Ubbelohde, A. R. J. P. & Young, D. A. Thermal conductance of graphite in relation to its defect structure. Proceedings of the Royal Society of London. Series A. Mathematical and Physical Sciences 276, 83–95 (1963). 10.1098/rspa.1963.0194.
47. Kitaoka, S., Wada, M., Nagai, T., Osa, N. & Konno, T. Increasing the thermal diffusivity of flexible graphite sheets by superheated steam treatment. Journal of Materials Science 46, 1132–1135. https://doi.org/10.1007/s10853-010-4991-5 (2011).
48. Uher, C., Hockey, R. L. & Ben-Jacob, E. Pressure dependence of the c-axis resistivity of graphite. Phys. Rev. B 35, 4483–4488. https://doi.org/10.1103/PhysRevB.35.4483 (1987).
49. Autio, G. & Scala, E. The normal spectral emissivity of isotropic and anisotropic materials. Carbon 4, 13 – 28 (1966). http://www.sciencedirect.com/science/article/pii/0008622366900054.

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Author contributions
M.Ce. identified the research needs, reviewed literature, planned and troubleshooted the experimental methods, prepared samples, collected data, analyzed results, prepared figures, and wrote the manuscript. N.P. planned and performed the emissivity measurements. M.B. initiated and supervised the project and secured funding. All authors have reviewed the manuscript.

Competing interests
The authors declare no competing interests.

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