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Enhanced thermal conductivity and reduced thermal resistance in carbon fiber-based thermal interface materials with vertically aligned structure†

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As electronic devices advance, there's a critical need for thermal interface materials (TIMs) with high thermal conductivity and minimal thermal resistance to address thermal dissipation challenges effectively. Carbon fibers (CFs), known for their axial thermal conductivity, are ideal for creating high-performance TIMs in a vertically aligned structure, aligning with the TIMs' heat transfer direction. Despite advancements in CF alignment for improved thermal conductivity, the high thermal resistance and the need for cost-effective manufacturing remain challenges. We propose a novel sandwich structure integrating vertical heat transfer pathways with surface modification to tackle these issues. This structure features a core of vertically aligned CF composite, achieved through a rolling-press method, flanked by liquid metal-modified layers to reduce contact thermal resistance. Our composites demonstrate an exceptional through-plane thermal conductivity of 51.90 W m⁻¹ K⁻¹ at 73.68 wt% filler content, 323 times higher than that of the PDMS matrix, and a reduced total thermal resistance from 0.55 to 0.32 K cm² W⁻¹ after interface modification. This research offers insights into designing CF-based composites for enhanced thermal management, applicable in cloud computing and autonomous driving.

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Introduction

In recent years, there has been rapid advancement in semiconductor technology, driving silicon-based chips towards more precise manufacturing processes and innovative three-dimensional chip stacking techniques, while introducing new third-generation semiconductor materials like gallium nitride (GaN) and silicon carbide (SiC), applied extensively in 5G base stations, ultra-fast charging stations, and consumer electronics.^{1–3} The advancements in processes and novel materials have led to enhanced chip performance but have also resulted in significant heat accumulation issues. Insufficient dissipation of heat could compromise the reliability and

stability of electronic devices during their operation, potentially threatening their longevity.⁴ To control the operational temperature of chips, ensuring efficient heat transfer from the chips to the heat sink is a crucial step in addressing this challenge. Thermal interface materials (TIMs) have been developed to bridge the mating interface between chips and heat sinks, eliminating air gaps and enhancing effective heat transfer between them.^{5,6} The ideal TIMs not only possess high through-plane thermal conductivity ($\kappa_{\perp}$) but also exhibit low contact thermal resistance (R_{contact}) with the heater and heat sink.⁷ In general, TIMs with low R_{contact} require compressibility and flexibility to ensure a greater effective contact area between the mating surface under normal packaging conditions. Due to their flexibility, unique design freedom, and cost-effectiveness, polymers have become the predominant matrix in the most common TIMs. However, the low thermal conductivity of polymers, due to their amorphous arrangement and molecular chain vibrations, restricts their direct application in thermal management.⁸ The most effective strategy to enhance thermal conductivity is by introducing micro and nanoscale fillers with high intrinsic thermal conductivity into polymer composites.^{9,10} The currently used thermally conductive fillers are generally classified into three categories: ceramics (Al₂O₃,¹¹ SiC,¹² AlN,¹³

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The first step involves preparing a homogeneous mixture solution of PDMS and fillers, where the uniformity and rheological properties of the mixture solution significantly impact the subsequent CF orientation process. Initially, the two components of PDMS, A and B, are mixed in a 1 : 1 ratio using a high-speed mixer (2000 rpm, 2 minutes) to achieve uniformity. Subsequently, a predetermined proportion (55, 60, 65, 70, and 73.68 wt%) of CFs and BN fillers (maintaining a fixed ratio of 8 : 1 for CFs to BN) is added to the prepared PDMS solution. Due to the high filler content, achieving uniform mixing of the solution requires multiple sequential additions of fillers during the mixing process. This step also involves the use of a high-speed mixer operating at 2000 rpm for a duration of 5 minutes to ensure thorough mixing. The second step involves orienting the CFs using a roll press machine. The uncured mixed solution is poured into the hopper of the roll press machine, with a layer of PET film placed above and below (to maintain the shape of the uncured material). The distance between the dual rollers is adjusted to control the degree of fiber orientation, with the rollers operating at a controlled speed of 60 cm min⁻¹.

(UTM, Roell Z030, Zwick, Germany). An infrared camera (Fluke, Ti400, USA) was employed to record temperature evolution.

Preparation and structural characterization of the VCB composites

Preparation of VCB-LM composites

Characterization

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Fig. 1 Preparation and morphology of the VCB TIMs. (a) Schematic diagram of the rolling-press technique for aligning carbon fibers and the production process of VCB composites. (b) The microstructure of continuous single layer composites extruded by the roller. (c and d) Top view and cross-section view of the microstructure of VCB composites.

shows the VCB composite under bending conditions. Even at large bending angles, no delamination or cracking was observed between the layers.

The strategy of this study involves filling polymers with fillers and modifying their arrangement to enhance the thermal conductivity of composites. Consequently, the properties and dimensions of the fillers will directly impact the overall performance of the composites. In our study, high thermal conductivity short chopped pitch-based CFs were chosen as the primary filler to enhance the thermal conductive of the composite. Fig. S7a–c† depict the optical, SEM magnified, and microstructural images of CFs. According to the images, the diameter of the CFs measures 10 μm, with a length of 250 μm (as detailed in the length distribution in Fig. S7d†). Furthermore, the fiber cross-section displays the characteristic radial folded arrangement indicative of mesophase pitch-based CFs. The XRD pattern of the CFs displayed in Fig. S7e† exhibits characteristic peaks associated with graphite, and the peak at 26.4° corresponds to the (002) plane, while the prominent peak at 54.5° is assigned to the (004) plane. These sharp characteristic peaks indicate larger grain sizes, smaller interlayer spacing, and a closely packed arrangement of graphite layers, consistent with the cross-sectional image in Fig. S7c.† The Raman spectrum of the CFs is depicted in Fig. S7f,† wherein distinct peaks at 1353, 1582, and 2704 cm^{−1} labeled as D, G, and 2D, respectively, were observed. These peaks are commonly employed to characterize the structure of carbon materials, and the I_D/I_G ratio of the CFs serves as a pivotal metric for assessing their thermal conductivity. The I_D/I_G value of the CFs utilized in

this study is 0.135, indicating remarkably low defects and superior thermal conductivity. h-BN serves as the second hybrid filler in this study. Its role includes filling the gaps between CFs, bridging these fibers to establish additional thermal pathways. Furthermore, it aids in modulating the rheological properties of the uncured mixed solution. The optical and SEM magnified images of the h-BN are presented in Fig. S8a–c.† The h-BN exhibits a lateral size range of 20–50 μm, as depicted in the lateral size distribution plot in Fig. S8d,† with a thickness of 1–2 μm. Fig. S8e† displays the XRD pattern of BN, showcasing distinct characteristic peaks. Additionally, Fig. S8f† presents the prominent Raman peak at 1365 cm^{−1} for h-BN. These results confirm the excellent crystal quality of the larger-sized BN.

Based on their fabrication method, the produced block-shaped composites exhibit anisotropic characteristics. To explore their internal structure and the influence of fiber orientation on performance, samples were sliced along the direction indicated in the schematic diagram in Fig. 2a. The composites cut along the direction perpendicular to the Cartesian coordinate system's X-axis are denoted as VCB-X, featuring a vertical CF alignment within their interior. Correspondingly, samples cut along the direction perpendicular to the Y and Z axes are denoted as VCB-Y and VCB-Z, respectively. To analyze the internal structure, we used an advanced micro-CT tool to scan the VCB-X, VCB-Y and VCB-Z composites. Within this research, the angle between the fibers and the Z-axis (through-plane direction) was defined as θ , as illustrated in Fig. 2b. Furthermore, the CF skeleton inside the three composites is shown in Fig. 2c (where fibers at different angles are





Fig. 2 Microstructure characterization of VCB composites. (a) Schematic diagram of the anisotropic composite cutting method (VCB-X, VCB-Y and VCB-Z are prepared by cutting vertically along the X, Y and Z axes, respectively). (b) Schematic of the angle (θ) between CFs and the normal vector in the 3D model. (c) Micro-CT model of VCB 73.68 wt%-X, VCB 73.68 wt%-Y and VCB 73.68 wt%-Z composites. (d) Probability statistical plot of CF angle distribution according to the micro-CT model. (e) 3D and 2D WAXD patterns for VCB composites.

represented by distinct colors). ESI Video S1† showcases the angular distribution of CFs in the three specimens. Subsequently, reconstruction of models was conducted using analysis software, generating the fiber angle distribution data within each composite, as depicted in Fig. 2d. By comparing the data from the three specimens, the angular distribution of the VCB-73.68 wt%-X specimen follows a normal distribution centered at 15° . Correspondingly, the model predominantly exhibits blue and green colour, which aligns with our deliberately controlled outcome (smaller angular distributions indicate higher alignment degrees). However, in VCB-73.68 wt%-Y, the distribution appears more random, leading to a relatively chaotic pattern in the model. However, the VCB-73.68 wt%-Z sample exhibits a predominance of larger angular distributions, resulting in a higher uniformity in the model colors. Yet, due to the low radial thermal conductivity of CFs, this orientation in this direction does not favor through-plane heat conduction. Considering the significantly higher axial thermal conductivity of the anisotropic CFs compared to their radial counterpart, we introduced the parameter of “effective fraction”. When θ is less than 45° , it signifies a higher contribution to heat conduction; however, angles greater than 45° contribute less. Through calculations, the VCB-73.68 wt%-X sample exhibited an effective fraction of 59.14%, VCB-73.68 wt%-Y measured 36.87%, while VCB-73.68 wt%-Z was only 1.66%. Wide-angle X-ray diffraction (WAXD) was utilized to assess the degree of orientation in

composites. CFs comprise stacked layers of hexagonal carbon networks preferentially oriented along the fiber axis, with the structural and orientational characteristics revealed by the XRD peak originating from the (002) crystal plane reflection of these carbon network layers.^{38,39} Employing this technique, an analysis was conducted on three variants of VCB composites, and the resultant 3D and 2D patterns are visually presented in Fig. 2e. During the test, X-rays directly pass through the actual heat transfer path of each sample. Due to the vertical distribution of CFs in the VCB-X sample, X-rays passing through the axial direction of the fibers do not exhibit noticeable characteristic peaks. However, in the VCB-Y and VCB-Z samples, X-rays passing through the radial direction of the fibers exhibit distinct peaks that are clustered together. The diffraction peaks of VCB-Z have a relatively narrow half-width, consistent with the micro-CT test data. Fig. S9† also displays the integral intensity of the (002) crystal vs. azimuth angle curves of the three samples. Through the aforementioned testing methodologies, we have successfully fabricated anisotropic composites with highly oriented structures. Among these, VCB-X exhibits the most promising vertical alignment of CFs. This vertical structure establishes the most efficient thermal conduction pathway, greatly enhancing the thermal conductivity efficiency of the thermal interface material.



Thermal conductivity of the VCB composites

To investigate the impact of filler content on the thermal conductivity of anisotropic composites, we prepared samples with varying filler contents (ranging from 55 to 73.68 wt%). Fig. 3a illustrates the influence of filler content on the thermal conductivity in the three directions (X , Y , Z) of the anisotropic VCB composites, and the detailed calculation parameters are available in ESI Table S1.† As the filler content increased, the thermal conductivity of samples in all three directions (X , Y , Z) also increased. However, owing to its structural characteristics,

composites with the same filler content exhibited significant differences in through-plane thermal conductivity across different directions. VCB-73.68 wt%- X attained the highest thermal conductivity, reaching $51.90 \text{ W m}^{-1} \text{ K}^{-1}$. In contrast, at the same filler content, VCB-73.68 wt%- Y registered $17.32 \text{ W m}^{-1} \text{ K}^{-1}$, while VCB-73.68 wt%- Z exhibited only $2.07 \text{ W m}^{-1} \text{ K}^{-1}$. It's worth noting that these test results align with the "effective fraction" depicted in Fig. 2c, confirming that the angular distribution of CFs is the most crucial parameter affecting the thermal conductivity. Next, the anisotropy ratios of the thermal conductivities in different directions, specifically κ_X



Fig. 3 Thermal properties of VCB composites. (a) Effect of filler contents on thermal conductivities of anisotropic VCB composites in the three directions. (b) Effect of filler contents on anisotropic factor ratio (κ_X/κ_Z and κ_Y/κ_Z) of VCB composites. (c) Schematic diagram of the through-plane heat transfer capability test system. (d) Infrared photo and the temperature evolution curves during the heating process for comparing the thermal conductivity of VCB 73.68 wt%- X , VCB 73.68 wt%- Y and VCB 73.68 wt%- Z composites. (e) Effect of filler contents on the thermal conductivity enhancement of anisotropic VCB composites in the three directions. (f) Effect of environmental temperature on the thermal conductivity of the VCB-73.68 wt%- X composite. (g) The thermal conductivity variation of VCB-73.68 wt%- X during heating and cooling at 10 cycles. (h) The comparison of thermal conductivity among various preparation methods for carbon fiber composites reported in the literature. (i) Comparison of thermal conductivity and density between composites with carbon fiber-based, BN-based, and commercial TIMs as reported in the literature.

to κ_Z and κ_Y to κ_Z , were calculated. The results are illustrated in Fig. 3b. A larger numerical value of the anisotropy ratio indicates a greater difference in thermal conductivity. At the highest filler content, κ_X/κ_Z and κ_Y/κ_Z reached 25.07 and 8.37, respectively. To further validate the varying heat transfer capabilities in different directions of the anisotropic composites, we set up a system, as depicted in Fig. 3c, employing an infrared camera to record the temperature rise rates of VCB-73.68 wt%-X, VCB-73.68 wt%-Y, and VCB-73.68 wt%-Z. Fig. 3d displays the infrared images captured at different time intervals and the temperature data curves at the top of the samples under identical test conditions for the three samples. From the IR images, it's evident that VCB-73.68 wt%-X exhibits the fastest heat transfer rate, resulting in a more uniform temperature distribution across the material. Conversely, VCB-73.68 wt%-Z, due to its lower thermal conductivity, impedes heat transfer towards the top, causing most of the heat to concentrate at the interface with the heat source. The disparities in temperature curves further corroborate the superior through-plane thermal transfer performance of VCB-73.68 wt%-X. Moreover, we employed Thermal Conductivity Enhancement (TCE) to express the improvement of the composite's thermal conductivity relative to the pure PDMS matrix. The formula for calculating TCE is as follows:

$$\text{TCE} = \frac{\kappa_c - \kappa_m}{\kappa_m} \times 100\% \quad (1)$$

where κ_c represents the thermal conductivity of the composite, while κ_m stands for the thermal conductivity of the pure matrix. The VCB-73.68 wt%-X exhibits an outstanding TCE of 32 338%, as shown in Fig. 3e, whereas the VCB-73.68 wt%-Z at the same filler content registers only 1194%. This data further validates the crucial significance of controlling the orientation structure of anisotropic fillers.

Beyond thermal conductivity, thermal stability also dictates the applicability of certain materials. Excellent thermal stability allows materials to perform effectively under extreme operating conditions. Fig. 3f illustrates the thermal conductivity change of VCB-73.68 wt%-X across a temperature range of 25 to 115 °C. As expected for many materials, there is a gradual reduction in thermal conductivity with increasing temperature. Fig. 3g records the variations in thermal conductivity during successive heating and cooling cycles, aiming to assess the stability of thermal conductivity under thermal shock conditions. The graph displays ten cycles of heating and cooling. Throughout this phase, VCB-73.68 wt% demonstrates commendable stability, displaying minor fluctuations, both at high and low temperatures. In Fig. 3h, we have compared the thermal conductivity values of previously reported CF-based TIMs with those obtained in this work, considering the preparation techniques employed for oriented CFs. More detailed data can be found in ESI Table S2.† Methods like ice templating^{26,31,40} are suitable for fabricating 3D skeletons, yet they fail to provide high thermal conductivity at low filler concentrations. Techniques such as electrostatic flocking^{27,41,42} and magnetically assisted methods,^{28,34,43} due to their intricate process conditions, have not gained industrial acceptance. Presently,

methods involving 3D printing^{23,30,33} and flow-assisted extrusion,⁴⁴ as well as the pre-arrays^{45–47} of continuous CFs, appear to hold promise as potential industrial-scale manufacturing processes for TIMs. It is worth noting that both 3D printing and flow-assisted extrusion techniques seem to have reached a bottleneck in terms of achieving thermal conductivity improvements, as neither has surpassed 40 W m^{−1} K^{−1}. Moreover, while the approach involving continuous pre-alignment of CFs holds promise for achieving higher values, the cost of continuous CFs remains prohibitively high for manufacturers. Therefore, the rolling-press technique⁴⁸ employed in this study emerges as a potential method for large-scale production of high thermal conductivity CF-TIMs. The TIMs produced through this method achieved thermal conductivity exceeding 50 W m^{−1} K^{−1}, surpassing all previously mentioned fabrication approaches. The density of TIMs is also a crucial parameter under consideration. In Fig. 3i, a comparison of density *vs.* thermal conductivity is drawn among CF-based TIMs,^{29,30,34,45} BN-based TIMs,^{49–53} and commercially available CF-based TIMs from previous literature. The detailed parameters are provided in Table S3.† It's observed that most TIMs exhibit a density below 2 g cm^{−3}, owing to the relatively lower density of CFs and boron nitride fillers. However, the thermal conductivity of BN-based TIMs remains below 15 W m^{−1} K^{−1}, unable to meet the requirements for high thermal conductivity. While commercially available CF-TIMs exhibit commendable thermal conductivity, they tend to have higher densities. Contrastingly, the TIMs developed in this study achieve the highest thermal conductivity alongside relatively lower densities. Therefore, this sample demonstrates significant advantages in applications requiring extreme lightweight characteristics, such as aerospace or portable electronic devices.

Furthermore, the rough surface of CF-based TIMs cannot achieve extensive interfacial contact, resulting in an increase in interfacial thermal resistance. Therefore, a low compression modulus allows the TIMs to deform under pressure, enabling them to conform to rough surfaces. Fig. S10† illustrates the compression stress–strain curve of VCB-73.68 wt%-X. The curve indicates significant deformation occurring at relatively low pressures, implying that this TIM can achieve excellent conformability even at lower packaging pressures. Another point of interest is that the vertically structured TIMs, upon experiencing deformation under pressure, cause a tilt in the angles of the CFs, as depicted in the schematic diagram of Fig. 4a and S11.† While we aim to prevent tilting, the alteration in the anisotropic structure under substantial deformation is unavoidable. Hence, determining the optimal sealing pressure for anisotropic TIMs is an essential consideration. Fig. 4b illustrates the change in thermal conductivity and thickness of VCB-73.68 wt%-X upon deformation under pressures ranging from 10 to 100 psi. It's important to note that the thickness measurements were taken after compression. As depicted in Fig. 4b, with increasing pressure, the thickness of VCB-73.68 wt%-X decreased from its original 1.40 to 0.80 mm after compression at 100 psi, concurrently reducing the thermal conductivity from 50.62 to 19.07 W m^{−1} K^{−1}. As evident in Fig. 4c–e, prior to compression, the CFs in the TIMs were





Fig. 4 The influence of pressure on thermal conductivity and structure of the VCB 73.68 wt%-X composite. (a) Schematic diagram of deformation of the vertical carbon fiber structure under pressure. (b) Effect of compressive stress on the thermal conductivity of the VCB-73.68 wt%-X composite and thickness (samples are measured for thermal conductivity and thickness after pressure release). (c–e) The incline angle of the carbon fiber structure of the sample after different pressure release. (f) Simulation comparison of heat transfer properties of composites with various incline angle CF structures. (g) The temperature gradient distribution curves at the center position calculated by four simulation models.

positioned at a 90° angle to the base. Following compression at 50 psi, the CF angle changed to 51° , further reducing to 38° after compression at 100 psi. Based on the above results, we determined that the optimal packing pressure range for VCB composites is 10–30 psi. Within this range, the inclination angle of the CFs does not change significantly. However, when the pressure exceeds 50 psi, the CFs exhibit excessive inclination, leading to a substantial decrease in thermal conductivity, thus resulting in failure.

To further validate the impact of CF orientation angles on thermal conductivity, finite element analysis was employed for verification. The first step involved establishing four models as depicted in the Fig. S12,[†] positioned at the same heating plate, while the external surroundings were designated as PDMS substrate with a thermal conductivity of $0.16 \text{ W m}^{-1} \text{ K}^{-1}$. Within the substrate, CFs were introduced at angles of 0° , 30° , 60° , and 90° respective to the base. The ambient temperature was set at 22°C , and convective heat transfer conditions were applied to the boundaries of the four models, with a coefficient of $5 \text{ W m}^{-2} \text{ K}^{-1}$. The heating plate was maintained at a constant 80°C , ensuring heat conduction from the bottom of the model to the top. The boundary conditions were set for the models, and subsequently, a mesh generation process was applied to the

configured models, as depicted in Fig. S13.[†] Following this, computations were performed, producing temperature distribution maps for the four models, showcased in detail in Fig. 4f. As indicated by the computational outcomes, the inclined angle of CFs significantly influences the temperature distribution within the models. Fig. 4g illustrates the temperature gradient from the bottom to the top within the models. The model with a 0° orientation of CFs exhibits the most significant temperature differential, reaching only 58.2°C at the top surface. Conversely, the model with a 90° orientation shows the best internal temperature uniformity, reaching 73.9°C at the top surface. Additionally, the model tilted at 60° demonstrates better temperature uniformity compared to the 30° oriented model. This simulation outcome aligns with our experimental results, indicating that an increase in the tilt angle of CFs leads to a decrease in their thermal conductivity.

Thermal resistance of the VCB composites

Generally, the total thermal resistance is a critical parameter affecting the performance of TIMs during practical applications. When TIMs are used to bond the heater and heat sink, the components of the total thermal resistance (R_{total}) are represented as shown in eqn (2).



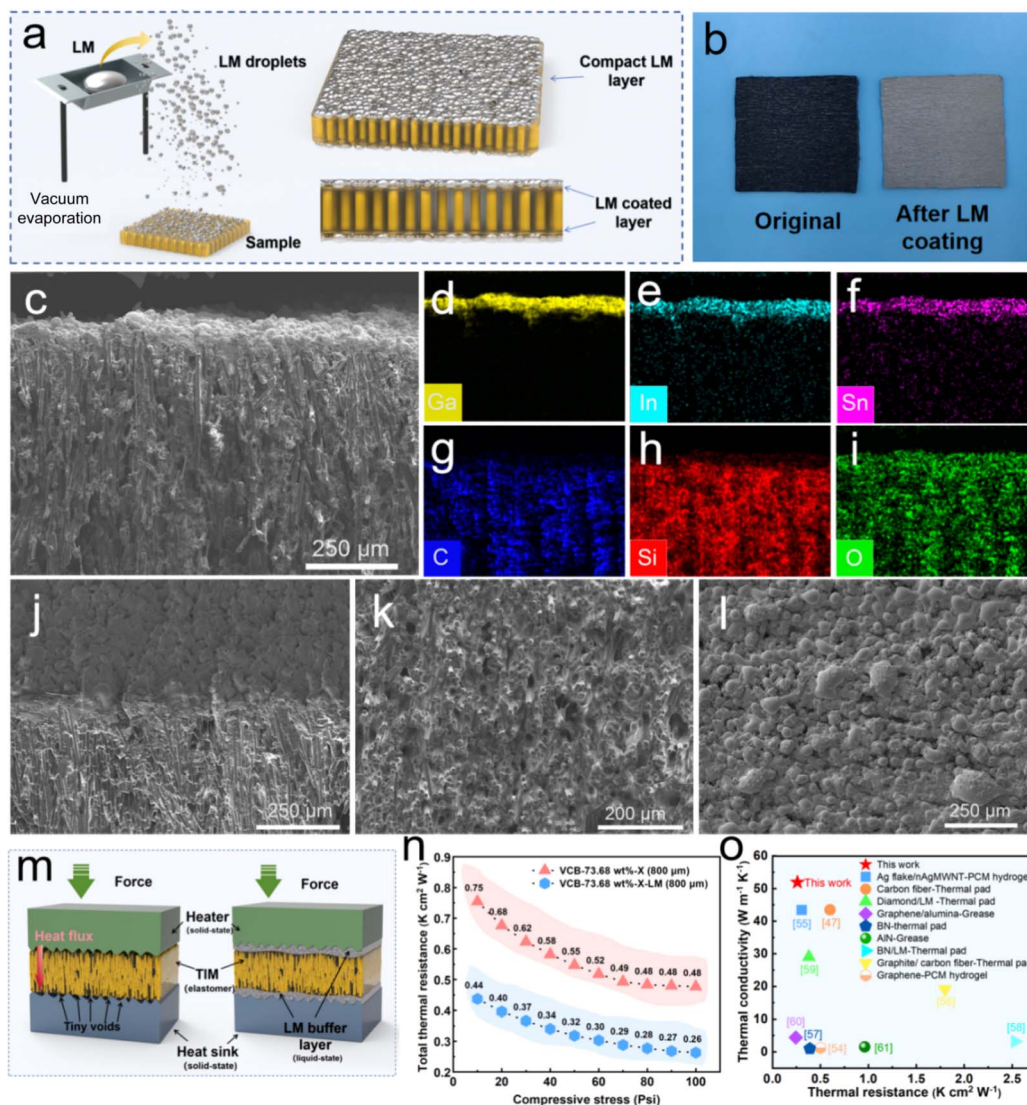


Fig. 5 Preparation and characterization of the functionalized coating. (a) Schematic diagram of the vacuum evaporation technique for coating LM onto composites. (b) Optical photography for the original composite and composite with LM coating. (c) The cross-section view of VCB-73.68 wt%-X-LM with the corresponding element mapping shown in (d)–(i). (j) Cross-section view of the microstructure of VCB-73.68 wt%-X-LM. (k) Top view of the microstructure of the composite without a coating. (l) Top view of the microstructure of the composite with an LM coating. (m) Schematic diagram of the contact state in the original composite and the LM coated composite. (n) Effect of compressive stress on the total thermal resistance of VCB-73.68 wt%-X and VCB-73.68 wt%-X-LM. (o) Comparison of total thermal resistance and thermal conductivity between VCB-73.68 wt%-X-LM and TIMs reported in the literature.

slope in the steady-state temperature *versus* input power plot, indicate values of 0.71 (commercial TIM), 0.81 (VCB-73.68 wt%-X), and $1.88 \text{ W cm}^{-2} \text{ }^{\circ}\text{C}^{-1}$ (VCB-73.68 wt%-X-LM), respectively, showing the highest heat dissipation capability of VCB-73.68 wt%-X-LM. Furthermore, the thermal shock test was conducted on VCB-73.68 wt%-X-LM using the same setup, where the heater's power density was set to 0 and 80 W cm^{-2} . Fig. 6d illustrates the temperature variations of the heating plate over 2000 cycles, demonstrating that VCB-73.68 wt%-X-LM exhibits remarkable thermal stability.

Considering the exceptional heat transfer capabilities of the produced VCB-73.68 wt%-X-LM, we directly integrated it into a commercial desktop computer to validate its thermal behavior

in real dynamic applications. As illustrated in Fig. 6e and depicted in the optical photograph, VCB-73.68 wt%-X-LM was packaged as a TIM between the Central Processing Unit (CPU, Inter core i5) and the air-cooled heat sink. Then, professional system diagnostic software (OCCT) was employed to drive the CPU to run at 100% usage. Meanwhile, the software HWINFO64 was utilized to monitor the temperature evolution of the CPU core and the effective clock frequency. For comparison, we conducted CPU stress tests under the same conditions using VCB-73.68 wt%-X and without using TIMs. As shown in Fig. 6f, under normal state, employing VCB-73.68 wt%-X-LM as the TIM decreased the temperature of the CPU core ($T_{\text{CPU core}}$) by 4°C compared to that without using the TIM, and it decreased by 2°C



Fig. 6 TIM performance and the thermal management demonstration of VCB composites. (a) Schematic diagram of the TIM performance test system. (b) The evolution of heater temperature as a function of different power densities ($10\text{--}80\text{ W cm}^{-2}$) is compared between VCB composites and commercial TIM (Wartime-WT5935C-250-65). (c) The correlation between the steady state temperature and power density of the heater. (d) Thermal shock stability tests in cyclic heating/cooling (2000 cycles) using VCB-73.68 wt%-X-LM as a TIM. (e) Assembly schematic diagram of the computer cooling system (inset photo showcasing the packaged motherboard and VCB-73.68 wt%-X-LM as a TIM applied on the CPU to bond the heat sink). (f) The evolution of CPU temperature during the CPU stress tests (100% CPU usage). (g) CPU effective clock variation during the CPU stress tests.

compared to using VCB-73.68 wt%-X as the TIM. During the stress test stage, the temperature gap increased notably. The $T_{\text{CPU core}}$ with VCB-73.68 wt%-X-LM as the TIM dropped by $8\text{ }^{\circ}\text{C}$ compared to when no TIM was used. It also showed a $2\text{ }^{\circ}\text{C}$ decrease compared to using VCB-73.68 wt%-X as the TIM. Here we noticed a discrepancy between the CPU test results and the TIM performance test results. It's important to note that during the CPU stress test, we utilized the existing commercial heat sink and fan, which have limited convective heat dissipation capabilities. While VCB-73.68 wt%-X-LM efficiently conducts CPU heat to the heat sink, the fan isn't able to promptly remove the heat from the heat sink, hindering the CPU temperature decrease. Despite the impact of the heat sink and fan on the TIM performance, as depicted in Fig. 6g, employing VCB-73.68 wt%-X-

LM as the TIM resulted in an increased effective CPU clock frequency of 52.6 MHz compared to operating without a TIM. The increase in the effective CPU clock frequency directly enhances user experience, validating the potential application of VCB-73.68 wt%-X-LM as the TIM in the realm of consumer electronics.

Conclusions

In summary, we have developed a novel sandwich-structured CF thermal interface material, exhibiting exceptionally high through-plane thermal conductivity and lower thermal resistance. The central layer incorporates vertically aligned CFs to enhance $\kappa_{\perp}$, while the upper and lower layers were modified with liquid metal coatings to minimize the R_{contact} . The aligned



The data underlying this article will be shared on reasonable request to the corresponding author.

Zhenbang Zhang: conceptualization, methodology, writing – original draft; Rongjie Yang: software, data curation, writing – review & editing; Yandong Wang: methodology, formal analysis; Kang Xu: software, investigation, data curation; Wen Dai: formal analysis; Jianxiang Zhang: writing – review & editing; Maohua Li: investigation; Linhong Li: validation; Yingying Guo: validation; Yue Qin: visualization; Boda Zhu: visualization; Yiwei Zhou: writing – review & editing; Xingye Wang: data curation; Tao Cai: project administration; Cheng-Te Lin: project administration; Kazuhito Nishimura: resources; Hao Nan Li: supervision, writing – review & editing; Nan Jiang: supervision, writing – review & editing; Jinhong Yu: supervision, writing – review & editing.

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