Control over emissivity of zero-static-power thermal emitters based on phase-changing material GST

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Controlling the emissivity of a thermal emitter has attracted growing interest, with a view toward a new generation of thermal emission devices. To date, all demonstrations have involved using sustained external electric or thermal consumption to maintain a desired emissivity. In the present study, we demonstrated control over the emissivity of a thermal emitter consisting of a film of phase-changing material Ge₂Sb₂Te₅ (GST) on top of a metal film. This thermal emitter achieves broad wavelength-selective spectral emissivity in the mid-infrared. The peak emissivity approaches the ideal blackbody maximum, and a maximum extinction ratio of >10 dB is attainable by switching the GST between the crystalline and amorphous phases. By controlling the intermediate phases, the emissivity can be continuously tuned. This switchable, tunable, wavelength-selective and thermally stable thermal emitter will pave the way toward the ultimate control of thermal emissivity in the field of fundamental science as well as for energy harvesting and thermal control applications, including thermophotovoltaics, light sources, infrared imaging and radiative coolers.

**INTRODUCTION**

Any object with a temperature above absolute zero gives off light or thermal emission. Black soot generally shows high emissivity, whereas smooth metal exhibits low emissivity. Other natural materials are characterized by their own emissivities. The intrinsic thermal emission of these materials is usually spectrally broad, and their emissivities cannot be easily tuned. However, in certain applications, such as biochemical sensing⁴, light sources⁵,⁶ and thermal emitters with tailored emissivities are highly desirable. For instance, a sharp cut-off in emission wavelength is needed in thermophotovoltaics⁴. Tailored emissivities are highly desirable. For instance, a sharp cut-off in emission wavelength is needed in thermophotovoltaics⁴.

Conventional thermal emission with spectral selectivity is based on micro/nano-structures such as gratings¹⁶, photonic crystals¹⁷, photonic cavities¹⁸–²⁰, nanoantennas²¹ and metamaterials²². By altering the structural parameters of the micro/nano-structures with advanced micro/nano-fabrication, wavelength-selective thermal emitters from THz to visible can be realized¹⁶–³⁰. However, static micro/nano-structures do not offer flexible tunability in terms of thermal emissivity. To date, several strategies have been implemented to engineer control over the thermal emissivity. (i) Electrically tunable materials (such as quantum wells and graphene) have been employed for modulation of thermal emissivity. By electrically tuning intersub-band absorption in n-type quantum wells, the emissivity of a narrow-band mid-infrared thermal emitter based on a photonic crystal slab can be tailored statically¹³,¹⁴ or dynamically¹³–¹⁵. Using an electrostatic gate to control the charge density in graphene, modulation of the emissivity of a narrow-band mid-infrared thermal emitter composed of graphene/SiNx/Au nanoresonators by 3% has been achieved⁵⁶. However, sustained electric consumption is needed to maintain a certain state of the tunable materials. (ii) Materials with high thermal expansion coefficients (such as SiNx) have been introduced in micro-electro-mechanical-system-based emitters²⁷. By manipulating the distance between the top metamaterial pattern and the bottom metallic film by applying heat, control of the temperature-dependent emissivity is achieved. However, the emissivity difference is lower than 35%, and sustained thermal consumption is needed to maintain the mechanical distance so as to control the emissivity. (iii) Phase-changing materials (PCMs) have also been introduced to modulate the thermal emissivity owing to their different optical and infrared properties in their different phases. VOₓ is a typical PCM and exhibits an insulator-to-metal phase transition at a temperature of 67 °C²⁸. By controlling the phases of VOₓ at different temperatures, a tunable thermal emissivity for an emitter composed of a VOₓ layer on a sapphire substrate has been demonstrated²⁸. However, to preserve the metal phase of VOₓ, the temperature of the thermal emitter needs to be maintained above 67 °C. Otherwise, a metal-to-insulator phase recovery is activated. Although significant progress has been made to control the thermal emissivity, there has yet to emerge an energy-efficient solution to maintaining a desired emissivity with zero static power consumption at room temperature.

**Keywords:** absorptivity; emissivity; mid-infrared; switchable; thermal emitters

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Ge$_2$Sb$_2$Te$_5$ (GST) is another PCM material with amorphous and crystalline phases. It has been widely used as a commercialized rewriteable optical storage medium in digital video disks owing to its phase-changing performance. The atom distributions of GST in the amorphous and crystalline phases are shown in Figure 1a. Although the atoms are chaotically arranged in the amorphous phase, they are aligned in an orderly manner in the crystalline phase, leading to distinct infrared properties. The GST show distinct infrared properties, resulting in a sharp contrast between on and off states in the switchable emissivity. When the GST is in the crystalline phase, an emissivity approaching the ideal blackbody maximum (on state) is realized, and the emissivity is significantly suppressed to below 0.2 (off state) in the amorphous phase. By switching between the amorphous and crystalline phases of the GST, a maximum extinction ratio of up to 11 dB can be realized. (ii) The transition between the amorphous and crystalline phases of the GST reveals hybrid phase behaviors, which can be exploited to implement tunable emission by properly controlling the annealing time and the annealing temperature. (iii) Wavelength-selective thermal emission in a broad range (from 3 to 15 μm) can also be enabled by varying the GST thickness. (iv) The fabrication of this thermal emitter involves only simple film deposition; therefore, this thermal emitter benefits from the virtue of large-area and lithography-free fabrication as well as design flexibility.

MATERIALS AND METHODS

Fabrication of the GST–Au samples

The GST–Au samples are fabricated starting with a 120-nm-thick gold film deposition. Because the gold film is sufficiently optically thick, there is low demand for the substrate. The gold film is deposited on the BK7 glass substrate via a thermal evaporation technique. The GST layer is then deposited on the gold film by magnetron sputtering, in which germanium is DC sputtered and the other two components (stibium and tellurium) are radio frequency (RF) sputtered with a 2:2:5 deposition rate ratio. The as-deposited GST alloy is in the amorphous phase, and a 1.5 min annealing process at 180 °C on a hot plate is performed to obtain the crystalline GST in this paper.

Absorptivity measurements

The absorptivities of the GST–Au samples are derived by measuring the reflectivities. Because the gold film is sufficiently optically thick, transmission does not exist in the experiment. Thus, the absorptivity (A) and the reflectivity (R) satisfy $A = 1 - R$. The vertical reflectivities are measured with the Bruker Vertex 70 FTIR and the Hyperion 1000 infrared microscope equipped with a liquid nitrogen refrigerated MCT

![Figure 1](image) (a) Atom distribution diagrams of the two phases (amorphous and crystalline) of the GST. The red and blue dots denote the Ge/Sb atoms and Te atoms, respectively. (b) A 3D schematic of the switchable and tunable thermal emitter composed of a GST film on top of a gold film. (c) An SEM image of a cross-section of the fabricated thermal emitter.
detector, and the oblique reflectivities are measured with a DTGS detector on the Vertex 70 FTIR equipped with the A513 attachment.

**Emittance measurements**

The emitted spectra of the black soot and the GST–Au samples are measured by the Bruker Vertex 70 FTIR equipped with a DTGS detector. The samples are fixed on a temperature controller, and the emitted power is sent into the FTIR and detected by the DTGS detector. The temperatures of both the black soot and the GST–Au samples are controlled over 100 °C to ensure high emitted power and to decrease the noise. Every measurement is repeated 16 times to further reduce the noise. The infrared photographs are taken by the FLIR Tg165 infrared camera.

**RESULTS AND DISCUSSION**

A schematic diagram of the switchable and tunable thermal emitter is shown in Figure 1b. A GST layer is deposited on a gold film (also termed as 'GST–Au sample'). The gold film, which is sufficiently optically thick, can be replaced by other metal (silver, aluminum and so on) films. Figure 1c presents an SEM image of a cross-section of the device with a 450-nm-thick GST on top of a 120-nm-thick gold layer.

According to Kirchhoff's law of thermal radiation, the absorptivity of an object is equal to its thermal emissivity. Before determining the emissivity of the thermal emitter, its absorption properties as an absorber are studied first. The electromagnetic responses of the GST–Au samples are simulated by COMSOL Multiphysics software. In the simulation, the relative permittivity of gold is derived from Rakic's work54. The relative permittivity of the GST (shown in Supplementary Fig. S1) is determined based on the spectrophotometric method from the transmission and reflection measurements55. From the measured permittivity, it can be concluded that the GST in the amorphous phase is transparent in the mid-infrared, whereas the GST in the crystalline phase is highly absorptive.

The absorptivities of the GST–Au samples at normal incidence are investigated via simulations and experiments, which are presented in Figure 2a and 2b, respectively. The absorptivities of the samples at oblique incidence are presented in Supplementary Fig. S2. In the mid-infrared, the metal layer can be approximately regarded as a perfect electric conductor (PEC) owing to the large real and imaginary parts of its refractive index. Thus, nearly complete reflection with a phase shift of $\pi$ at the GST–Au interface is formed. Without the metal layer, fundamental resonance can also occur at a GST thickness of approximately $\lambda/2n$ ($n$ is the real part of the refractive index of GST and $\lambda$ represents the resonant wavelength); however, the maximum absorption is limited to 50%56,57 (Supplementary Fig. S3). When a gold film is placed below the GST film, not only is the GST thickness halved but also a nearly unity absorptivity can be achieved for the cGST–Au sample (see Section 3 of the Supplementary Information for more details). Because the aGST is nearly transparent, a resonant cavity is formed at a GST thickness of approximately $\lambda/4n$ such that a round trip inside the cavity could accumulate a $\pi$ (or $2\pi$) phase shift58.

At resonance, absorption and field penetration can be observed in the metal film in both the aGST–Au and cGST–Au samples because the metal film is not an ideal PEC. For the aGST–Au sample, this small absorption and field penetration in the metal film contribute to the total absorption (below 20%) at resonance; for the cGST–Au sample, nearly unity absorption arises from resonance-induced field enhancement in the highly absorptive cGST layer. Therefore, a sharp contrast between the absorptivities of the aGST–Au and cGST–Au samples is created. By increasing the GST thickness from 360 to 540 nm, the peak absorption wavelength shifts approximately linearly from 9 to 13 µm, demonstrating the wavelength-selectivity of the device. In the simulation, a broad wavelength selectivity from 3 to 15 µm wavelength can be achieved (the permittivity of the GST beyond a wavelength of 15 µm is unknown). Excellent agreement is obtained between the experimental data and the simulations. The slight difference can be attributed to light scattering loss induced by rough film surfaces.

To unveil the physics behind the sharp contrast between absorptivities of the aGST–Au and cGST–Au samples, the specific absorptivity in each layer (GST layer and metal layer) is further investigated (Figure 3a and 3b). Meanwhile, normalized electric field ($|E|$) and resistive loss ($Q$) distributions at resonant wavelengths for the GST–Au samples are presented in Figure 3c and 3d, respectively. It is obvious that the electric field intensity in the device decreases as light propagates into the GST layers and is quite weak at the GST–Au interface (Figure 3c), demonstrating the $\pi$-phase shift due to the reflection in metal and the $\lambda/4n$-thick GST layer. The absorption of light leads to heat generation, which can be characterized by a resistive loss (also termed as 'heat power volume density'). The resistive loss $Q$ is related to the imaginary part of the material relative permittivity ($\varepsilon'$) and the intensity of the electric field ($|E|$) by $Q = \pi c\varepsilon' |E|^2/\lambda$, where $c$ is the speed of light in vacuum and $\varepsilon_0$ is the vacuum permittivity. Because of the transparent property of the aGST, there are no resistive losses in the aGST layer even though its electric field is much stronger than that of the cGST layer. By comparison, large resistive losses occur in the cGST layers for the cGST–Au samples. The highly absorptive GST combined with resonant field enhancement contributes to nearly perfect absorption in the cGST–Au samples. In both cases, there are

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**Figure 2 (a, b)** Simulated and experimental normal-incident absorptivity of the cGST–Au and aGST–Au samples in the mid-infrared with three different GST thicknesses (360, 450 and 540 nm). The dashed and solid lines are for the aGST–Au and cGST–Au samples, respectively. (a) and (c) denote the aGST–Au and cGST–Au samples, respectively.

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resistive losses in the top surfaces of the metal layers due to the non-infinite imaginary part of its relative permittivity and residual electric field penetration into the metal.

The emitted spectra are measured by a Fourier transform infrared spectrometer (FTIR) equipped with a room temperature-doped triglycine sulfate (DTGS) detector. Thermal emission from the central areas (denoted by the red dotted circles in Figure 4a) of the samples is sent into the FTIR for detection. Black soot is conventionally regarded as a perfect reference owing to its high wavelength-independent emissivity. Here the black soot reference is produced by firing a rectangular stainless steel slice with a candle, and its emissivity ($\varepsilon_{\text{ref}}(\lambda, T)$, where $\lambda$ is the wavelength and $T$ is the temperature) is assumed to be 0.97. With the measured emission spectra of the GST–Au samples ($S_{\text{sample}}(\lambda, T)$) and the black-soot reference ($S_{\text{ref}}(\lambda, T)$) (shown in Supplementary Fig. S4a), the normalized emissivities ($\varepsilon_{\text{sample}}(\lambda, T)$) of the GST–Au samples can be derived from
\[
\varepsilon_{\text{sample}}(\lambda, T) = \frac{S_{\text{sample}}(\lambda, T)}{S_{\text{ref}}(\lambda, T)} \varepsilon_{\text{ref}}(\lambda, T) \quad (1)
\]

The derived emissivities of the GST–Au emitters with different GST thicknesses at 100 °C are presented in Figure 4b. The cGST–Au emitter yields a notably high peak emissivity approaching the ideal blackbody maximum. The peak emissivities are 0.92, 0.95 and 0.97 for GST thicknesses of 360, 450 and 540 nm, respectively. The emissivities of the aGST–Au emitters are low, and the emission peaks are below 0.20. Therefore, this thermal emitter shows two distinct states: an on state with a high emissivity when the GST is in the crystalline phase and an off state with a low emissivity when the GST is in the amorphous phase. The extinction ratios and the differential emissivities between the on and off states of the GST–Au emitters are presented in Figure 4c and Supplementary Fig. S4b, respectively. A maximum extinction ratio of 10.15, 11.04 and 10.98 dB can be obtained for GST thicknesses of 360, 450 and 540 nm, respectively. The peak emission wavelength shifts further with increasing cGST film thickness, thereby showing promise as a wavelength-selective thermal emitter. Although the emissivity is measured directly, whereas the absorptivity is characterized through transmission and reflection, excellent agreement between the experimental absorptivity (Figure 2b) and emissivity (Figure 4b) can be observed, verifying Kirchhoff’s law of thermal radiation.

The visible and infrared photographs of the black soot, the aGST–Au and cGST–Au emitters at 100 °C are presented in Figure 5a. The temperature label of the infrared camera is based on the integration of received power from 8 to 14 μm. In the infrared photographs, it is obvious that the black soot shows strong emission, whereas the aGST–Au emitter exhibits weak emission. To reveal angular dependences of emission, the oblique views of the infrared photographs of the samples are presented in Supplementary Fig. S5. The angular-dependent emitted power (from 8 to 14 μm) of the black soot and the cGST–Au emitter can be derived from the displayed temperature on the infrared camera. The normalized angular-dependent emitted power of the cGST–Au emitter with respect to the normally emitted power of the black soot is provided in Figure 5b. As the angle increases, the thermal emission of both the black soot and the cGST–Au emitter decreases slowly, indicating that the thermal emission is robust with respect to the emission angle.

So far, we have mainly focused on the switchable and wavelength-selective properties of the GST–Au emitters. The phase transition of the GST between the amorphous phase and the crystalline phase is

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**Figure 3** (a, b) Simulated absorptivity versus wavelength in specific layers (GST layer and metal layer) for the aGST–Au sample and the cGST–Au sample with different GST thicknesses (360, 450 and 540 nm). m, a and c denote the metal layer, the aGST layer and the cGST layer, respectively. Simulated normalized resonant (c) electric field $|E|$ and (d) resistive loss $Q$ distribution in the GST–Au samples with a 360-nm-thick GST film. The resonant wavelengths are 6.5 and 9.5 μm for the aGST–Au and cGST–Au samples, respectively.
The GST film can be assumed to be at intermediate phases composed of different proportions of amorphous and crystalline molecules. By controlling different intermediate phases of the GST, a tunable thermal emitter can be realized. This tunable property can be investigated by gradually increasing the sample temperature and simultaneously extracting the thermal emission spectrum. When the temperature is raised from 140 to 170 °C in 30 min with a step of 1 °C, the emissivity curves of the GST–Au emitter with a GST thickness of 450 nm are recorded and presented in Figure 6a. The peak emissivity increases with temperature and rises sharply between 150 and 155 °C. The peak emissivity approaches the ideal blackbody maximum at 170 °C. The temperature-dependent emissivities at different wavelengths are explicitly presented in Figure 6b, showing steep upslopes between 150 and 155 °C except for wavelengths of 7 and 8 μm. The temperature-dependent emissivity differences and extinction ratios in dB between the intermediate GST–Au samples and the aGST–Au sample are provided in Supplementary Fig. S6. The relative refractive indices of the GST during the intermediate phases are measured and presented in Figure 6c. Both the real and imaginary parts of the refractive index increase with temperature. The increase in the real part is primarily responsible for the resonance shift to longer wavelength, and the increase in the imaginary part contributes to the increase in the emissivity. The intermediate phases of the GST are also stable at room temperature, and therefore no sustained external thermal consumption is needed to maintain a desired emissivity for the GST–Au emitter. The intermediate phase of the GST is dominated by both annealing time and annealing temperature (Supplementary Fig. S7). By controlling the annealing process, the crystallization level and the intermediate phase of the GST can be correspondingly tailored, thus providing an avenue for tunable thermal emission.

In this paper, the crystallization process of the GST is investigated by a thermal annealing method, while reamorphization (phase transition from the crystalline phase back to the amorphous phase)
is not applied owing to a low-fusion point of the 120-nm-thick gold film. The thermal reamorphization of the GST on a silicon wafer at a temperature above 640 °C is performed, and the infrared properties are presented in Supplementary Fig. S8, demonstrating the reamorphization ability of the cGST. Because the Au film merely plays the role of a PEC, by replacing it with other metal films with higher-fusion points (such as Ti, W and Mo), this emitter can also be reversely switchable from the on state back to the off state. The switching speed depends on the phase transition speed of the GST. The phase transition process takes several seconds with the thermal annealing method in our experiment. This timescale can be decreased by increasing the annealing temperature (Supplementary Fig. S7) or reducing the thermal substrate effect. In addition to thermal annealing, the crystallization and reamorphization of the GST can also be realized using proper laser pulses with high peak power or electrical stimulations. The typical switching time for electrical stimulation is nanoseconds. By stimulating the sample with laser pulses, the switching time can be nanosecond or even improved to femtoseconds. So far, the emission property of the GST–Au emitter has been exclusively demonstrated in the 8–14 μm atmospheric window. The cGST–Au samples have simultaneous switchable, tunable and wavelength-selective emission properties in the 3–5 μm atmospheric window (shown in Supplementary Fig. S9), demonstrating the capability of dual-band emission in both thermal atmospheric windows.

CONCLUSIONS
In summary, control over the emissivity of zero-static-power mid-infrared thermal emitters consisting of a thermally stable GST layer and a metal layer is demonstrated. The GST film does not require sustained thermal consumption to preserve its phase. The metal film plays the role of a PEC, which not only halves the GST thickness but also significantly enhances the resonant peak emissivity of the emitter. Broad wavelength-selectivity is realized simply by controlling the GST thickness. An emissivity approaching the ideal blackbody maximum is achieved when the GST is in the crystalline phase, and a maximum extinction ratio of above 10 dB can be obtained between the two states (on state and off state) of the emitter. Moreover, by controlling the intermediate phases composed of different proportions of the amorphous and crystalline molecules of the GST, the thermal emissivity can be continuously tunable. Moreover, because only layered films are involved in the fabrication, this thermal emitter benefits from the merits of large-area and lithography-free fabrication as well as design flexibility. Combining all these advantages, this switchable, tunable, wavelength-selective zero-static-power thermal emitter will pave the way toward the ultimate control of thermal emissivity in the field of fundamental science as well as for energy-harvesting and thermal control applications, including thermophotovoltaics, light sources, infrared imaging and radiative coolers.

CONFLICT OF INTEREST
The authors declare no conflict of interest.

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Figure 6 (a) Tunable emissivity of the cGST–Au emitter in the mid-infrared with increasing temperature from 100 to 170 °C. (b) Temperature response of the emissivity for the cGST–Au emitter at several typical wavelengths. (c) Measured refractive indices of the GST at different temperatures. n and k denote the real and imaginary parts of the refractive indices, respectively. The GST thickness is 450 nm.

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