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Length-dependent thermopower of selfassembled monolayers of alkanethiolates depends on direction of temperature gradient[†]

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Self-assembled monolayers of alkanethiols on gold are often used as a benchmark for molecular junctions, particularly as a reference for length-dependence. However, their thermopower decreases with molecular length and is bifurcated. While the bifurcation can be explained by the presence of gateway states, the length-dependence directly contradicts theoretical predictions. We reproduced both experimentally using conical tips of eutectic Ga–In (EGaIn) to form junctions. We then show that this unusual length-dependence arrises from the direction of the applied thermal gradient; when the gold substrate is cooled instead of heated, thermopower increases with length monotonically as predicted by theory. This unprecedented relationship between length-dependence and direction of thermal gradient persists over a very large range of absolute temperature. We therefore ascribe it to shifts in level-alignment induced by the electric field in the leads, which is present experimentally, neglected in theoretical modeling and is determined by the direction of the thermal gradient.

Introduction

The thermoelectric effect enables the conversion of waste heat into electrical energy by generating a voltage across a temperature gradient.¹⁻⁹ Thermoelectrics based on organic molecules show promise in expanding this technology by enabling the integration and miniaturization of thermoelectric generators to harness lowgrade waste heat such as that generated by the human body. The Seebeck coefficient *S*, measured in V K⁻¹, provides the most direct quantification of the thermoelectric effect, representing the magnitude of induced thermoelectric voltage V per unit temperature difference ΔT across a material.^{10–15} Self-assembled monolayers (SAMs) of alkanethiolates provide access to highly insulating molecules by operating in the non-resonant tunneling regime when sandwiched between two electrodes.¹⁶⁻¹⁸ The electrode-SAM-electrode structure in these measurements defines molecular ensemble junctions,¹⁹ in which S is heavily influenced by the energy offset between electrodes and molecular orbitals.²⁰ These offsets are nearly invariant for alkanes, meaning S should vary linearly across a series of alkanes due to the length-dependence of conductivity;²¹⁻²⁵ however, Yoon et al. observed two separate

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regimes of length-dependence.²⁶ They ascribed this observation to the presence of gateway states that vanish for longer alkane chains.^{27–29}

Despite serving as a common benchmark for conductance measurements in tunneling junctions due to their straightforward properties, alkanethiols exhibit surprisingly complex thermoelectric properties and a divergence between theory and experiment; theory predicts a uniformly increasing S with increasing molecular length,³⁰ while experimental studies show two different regimes of length-dependence²⁶ and all but $two^{31,32}$ show a monotonic decrease in S with increasing length.³³⁻³⁷ Measurements are typically done in ambient conditions while heating the substrate to which the SAM is bound and holding the top-contact at room temperature because it is straightforward to insert a Peltier device under the substrate and heating avoids issues of condensation. We built an experimental platform, the Rick-9000,³⁸ capable of measuring thermoelectric voltages in SAMs using eutectic Ga-In (EGaIn) top-contacts over a large range of temperatures as illustrated in Fig. 1. Large-area molecular ensemble junctions are formed and measured in a N2 atmosphere at 0% RH to prevent condensation, with 1% O2, necessary for EGaIn to form stable tips.^{39,40} The SAMs are grown on ultra-flat template-stripped⁴¹ Au (Au^{TS}) that also serves as the bottom electrode. We validated the Rick-9000 by measuring S of the two electrodes from -120 to 80 °C and then reproducing the unusual length-dependence reported by Yoon et al. near room temperature. We then measured S down to -80 °C, finding that the length-dependence and magnitude of S are both temperature-

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Fig. 1 Schematic of the experimental setup summarizing its capabilities and the series of even-numbered alkanethiolates measured.

dependent and that *S* decreases with length when the substrate is the hot electrode, but increases when the substrate is the cold electrode.

Results and discussion

Measuring thermopower with the Rick-9000

The Seebeck coefficient can be determined experimentally by measuring the voltage drop that develops along the applied temperature gradient as shown in eqn (1).

$$S = -\frac{\Delta V}{\Delta T} \tag{1}$$

For molecular junctions, this gradient is applied out-of-plane by heating and/or cooling the top and/or bottom electrode(s). The Rick-9000 utilizes conical tips of EGaIn as top-contacts to form and measure junctions⁴² in a controlled atmosphere at 0% RH to prevent condensation; it is capable of operating at absolute temperatures far below the melting point of EGaIn (16 °C). While variable-temperature measurements using EGaIn have been reported, they use cylindrical molds to contain and shape it rather than free-standing tips.^{40,43-47} Thus, to ensure that the determination of *S* over the full temperature range is not impacted by the use of free-standing EGaIn, we contacted Au^{TS} directly with conical tips of EGaIn and measured the thermovoltage (ΔV) at



Fig. 2 Thermal voltage measurement of Au^{TS}//EGaIn contact across the entire range of 80 to -120 °C.

 $T_{\rm Au^{TS}} = -110$ to 80 °C and $T_{\rm EGaIn} = 15$ °C. As shown in Fig. 2, $\frac{\Delta V}{\Delta T}$ is linear across the entire range of temperatures. The resulting value of $S = (1.78 \pm 0.03) \,\mu V \, {\rm K}^{-1}$ is very close to the literature value for Au of 1.94 $\mu V \, {\rm K}^{-1}$.⁴⁸ Plots of the temperature and temperature-gradient stabilities are show in Fig. S1 (ESI†).

A large-area ensemble junction is created when a SAM is introduced between the EGaIn and Au^{TS}. We abbreviate the series of alkanethiolates SCn where n = 2 + 2(m - 1) and $1 \le m \le 9$ for SAMs comprising S(CH₂)_{n-1}CH₃. Assembled junctions are denoted Au^{TS}/SCn//Ga₂O₃/Ga–In where "/" denotes a chemisorbed (covalent) interface and "//" denotes a physisorbed (van der Waals) interface. For simplicity we use EGaIn to represent the entire Ga₂O₃/Ga–In interface because the Ga₂O₃ does not play a significant role in charge-transport.⁴⁹ The Au^{TS} electrode is the bottomcontact (upon which the SAMs are formed) and EGaIn is the topcontact. Using the sign convention of $\Delta T = T_{Au^{TS}} - T_{EGaIn}, \Delta V > 0$ when $\Delta T < 0$ and *S* is determined using eqn (2) where $V_{Contact}$ is the thermovoltage that develops in the leads and electrodes (*e.g.* Fig. 2), which is subtracted from the measured voltage V_{Meas} to isolate the thermovoltage generated by the SAM, ΔV .

$$S = -\frac{\Delta V_{\text{Meas}} - \Delta V_{\text{Contact}}}{\Delta T_{\text{Bottom}} - \Delta T_{\text{Top}}} = -\frac{\Delta V}{\Delta T}$$
(2)

The length dependence of thermopower in Au^{TS}/SCn//EGaIn junctions can be explained using the semiempirical parametric equation eqn (3) proposed by Quek et al.⁵⁰ This equation implies that SSAM linearly depends on the physical width of the energy barrier (number of carbons, n in SCn) and the slope β^{s} , which is the rate of thermopower change with *n* in units of $\mu V K n^{-1}$. It is the thermopower analog to the tunneling decay coefficient β .^{51,52} The term $S_{\rm C}$ is the thermopower of a hypothetical non-shorting junction, where n = 0, which captures the contributions to S that arise from everything other than S_{SAM} and may vary according to junction platform type and anchoring groups. In principle, β^{s} depends only on the shape of the energy barrier of the most insulating component in the junction, but as described below, the energy barrier manifests differently depending on the position of the Fermi level $E_{\rm F}$. In practice, $\beta^{\rm S}$ is an experimentally accessible empirical parameter that captures the relationship between S and the effects *n* has on the energy barrier.

$$S_{\rm SAM} = S_{\rm C} + n\beta^{\rm S} \tag{3}$$

Seebeck coefficients of SAMs of alkanethiolates

The thermopower of a molecular junction *S* can be further separated into the contribution of the anchor group and the molecule theoretically, similarly to the empirical parameters $S_{\rm C}$ and $S_{\rm SAM}$. For example, *ab initio* modeling predicts that S < 0for alkylisocyanates, but S > 0 for alkanethiols.³⁰ And it predicts that *S* hardly varies with length for alkylamines ($\beta^{\rm S} \approx 0$), varies inversely with length for alkylisocyanates ($\beta^{\rm S} < 0$) and varies positively with length for alkanethiols ($\beta^{\rm S} > 0$). The observed trends in $\beta^{\rm S}$ are ultimately determined by the total transmission probability of a junction, which in



Fig. 3 An idealized Lorentzian single-energy level zero-bias transmission probability (T) versus E illustrating how conductance decreases across SCn with a fixed value of E.

turn comprises the contributions from interfaces, anchor groups and molecular backbone. Fig. 3 is a single-level Lorentzian depiction of the zero-bias transmission probability⁵³ $\mathcal{T}(E)$ that illustrates how the level-alignment affects the magnitude and length dependence of S which, for non-resonant tunneling, is proportional to $-\frac{\partial T}{\partial E}$ (e.g. eqn (S2), ESI†).⁵⁴ For π conjugated molecules, the molecular orbital gap varies inversely with (conjugation) length, leading to a modest decrease in conductance with length (small β) and large increase in S (large β^{s}). The molecular orbital gap of alkanes, however, is nearly invariant with length and the transmission probability decreases mainly in the center of the gap with increasing length, resulting in a large drop in conductance (large β) and modest change in S (small β^{s}). Moreover, the trend in S is highly sensitive to the position of $E_{\rm F}$ and that the sign of S reverses at E = 0 in Fig. 3. Experimentally, ΔV also switches sign because the type of charge-carrier switches; holes when transport is facilitated by occupied states and electrons when it is facilitated by unoccupied states. This simple picture explains why the magnitudes of β^{s} and S increase as $\mathcal{T}(E) \to 1$, while the opposite is true of β .⁵⁵ Selzer *et al.* applied this principle experimentally, using Bi instead of Au to eliminate gateway states and optimize the level-alignment to achieve large, positive values of S and β^{S} for alkanethiolates that agreed quantitatively with their model of $\mathcal{T}(E)$.³²

While different trends in *S* have been observed for alkanes with different anchor groups and on different metals and the dependence of *S* on $E_{\rm F}$ is explained well by theory, the disagreement between theory and experiment for SAMs of alkanethiolates on Au persists; for alkanes, *S* ultimately depends on the combination of binding group and metal⁵⁶ and Au–S should produce S > 0 and $\beta^{\rm S} > 0$.³⁰ A common feature of experimental determinations of *S* is that they are performed at $T \approx 300$ K with small values of ΔT achieved by heating the bottom-contact (*e.g.* Au^{TS}) and keeping the top-contact equilibrated with ambient temperature. Theoretical modeling and single-molecule junction measurements are also almost always done using symmetrical chemisorbed contacts in which the direction of the temperature gradient does not matter. Measurements, however, are often performed on SAMs of monothiols with chemisorbed bottom-contacts and physisorbed top-contacts.^{26,34,36,37,57–61} The Rick-9000 is capable of both cooling and heating the bottom-contact while heating the top-contact, enabling not only a large range of absolute temperatures, but control over the direction of the thermal gradient (sign of ΔT) with respect to the disparate electrodes and interfaces. As can be seen in Fig. 2, the sign of the thermo-voltage switches as the bottom electrode temperature crosses 0 °C, satisfying eqn (1); *i.e.* the sign of *S* is determined from the slope, not the sign of the thermovoltage with temperature.

We measured the Seebeck coefficient of Au^{TS}/SCn//EGaIn junction by computing the slope of $-\Delta V$ versus ΔT plots for each SCn SAM. The values of S for each combination of ΔT are summarized in Table S1 (ESI†). The accuracy of these measurements depends on the magnitude of ΔV and its distribution. Similar to previous methods, we obtained S_{SAM} by treating junctions as thermopower circuits.^{20,57,62} In our case, we obtained the S_{SAM} by correcting for systematic bias, which eliminates all systematic voltage contributions V_{Contact} according to eqn (2). To validate S_{SAM} measurements, we measured Au^{TS}/SCn//EGaIn junctions at bottom-electrode temperatures (T_{AuTS}) of 24, 28 and 32 °C while holding the top-contact (T_{EGaIn}) at 20 °C to reproduce the measurement conditions in ref. 26.



Fig. 4 The Seebeck coefficients plotted against number of carbons *n* in the SAMs of SC*n* for Au^{TS}/SC*n*//EGaln junctions. The slopes of these plots are the values of β^{S} that are plotted in Fig. 5. The dashed line between $n \leq 10$ and $n \geq 12$ is to guide the eye. When $T_{Au^{TS}} > T_{EGaln}$, two different values of S < 0 are present for n = 2-10 and n = 12-18, reproducing the findings of ref. 26 (a). When $T_{Au^{TS}} < T_{EGaln}$, there is one value of $S \geq 0$ for n = 12-18 that increases with $|\Delta T|$ (b). The solid lines are linear fits of eqn (3).

The results shown in Fig. 4a perfectly reproduce the two values of β^{s} for $n \leq 10$ and $n \geq 12$ as well as the magnitudes of *S*.

Reversal of the length-dependence

As discussed above, theory predicts $\beta^{S} > 0$ for SC*n*, but $\beta^{S} < 0$ for Au^{TS}/SCn//EGaIn junctions has been reported previously and now reproduced in Fig. 4a.^{26,63} For all alkanethiols and all values of ΔT , S > 1 and the sign of β^{s} is determined by whether S increases or decreases with increasing n in SCn. The explanation for the presence of two separate regimes of lengthdependence is the presence of gap-states created by Au-S bonds that vanish with increasing length of SCn.29,56 The presence and position of these "gateway" states depends on the binding chemistry as well as the electronic structure of the molecule. They create resonances in $\mathcal{T}(E)$ between the resonances associated with the frontier molecular orbitals, reducing the energy gap and affecting the length-dependence of thermopower, β^{s} , but they do not manifest clearly in the length-dependence of conductance, β .^{64,65} The existence of gap-states accounts for the discontinuity in *S* between n = 10 and 12, but it does not explain why the sign of β^{s} differs from theoretical predictions. Importantly, Bi/SCn//EGaIn junctions do not exhibit two different regimes of length-dependence and $\beta^{\rm S} > 0$,³² demonstrating that the presence of Au-S bonds leads to the aberrant lengthdependence rather than something specific to SAMs, asymmetric contacts or EGaIn.

To probe the length-dependence of the thermopower of SC*n* further, we measured it at different values of *T*; specifically, we used the Rick-9000 to cool Au^{TS} instead of heating it, reversing ΔT . Surprisingly, the length-dependence collapsed into a single line that increases with length ($\beta^{\rm S} > 0$) producing the plots shown in Fig. 4b, which are qualitatively indistinguishable from theoretical predictions.³⁰ The numbers in the legend show the temperatures of the EGaIn top-contacts and Au^{TS} bottom-contacts. In all cases, $T_{\rm EGaIn} > T_{\rm Au^{TS}}$ and regardless of which electrode was held constant and which was varied, S > 0 and increases with length. There is a small discontinuity between $n \leq 10$ and $n \geq 12$ (indicated with a dashed line in Fig. 4), but it is within the uncertainty of *S*.

Linear fits to the data in Fig. 4 yielded the values of β^{s} shown in Fig. 5 (and Table S2, ESI⁺) plotted against the average temperature of the electrodes. The values at T = 26 °C show that S crosses zero when ΔT reverses. The magnitude of S then increases very slightly with decreasing temperature, meaning it is nearly constant with the increasing magnitude of ΔT . In general, the magnitude of S should increase with $|\Delta T|$, as is the case in Fig. 4b, but there is no obvious physical reason that $S_{\rm C}$ (the Y-intercept) or β^{s} (the slope) should. From Fig. 5 we cannot conclude if β^{s} is dependent on $|\Delta T|$ or if the variation is due to uncertainty or error in the measurements, e.g. it becomes more difficult to control for fluctuating voltage and temperature drops in the leads as Au^{TS} is cooled far below ambient. As a strictly empirical observation, however, β^{s} passes through zero, which implies the presence of an underlying continuous variable driving the transition from $\beta^{s} < 0$ to $\beta^{s} > 0$. (Note that in all cases S remains positive, meaning transport is always



Fig. 5 The rate of change of the Seebeck coefficients (β^{S}) for the SC*n* series plotted against the average temperature at which the Seebeck coefficients were measured. The dashed lines indicates where the direction of the thermal gradient switched. The red squares (inside dashed lines) show the separate values of β^{S} for the two different regimes of length-dependence (shown in Fig. 4a) when $\Delta T > 0$; the blue squares (outside dashed lines) show the single value of β^{S} obtained for the entire SC*n* series (shown in Fig. 4b) when $\Delta T < 0$.

HOMO-mediated.) Regardless, the restoration of the positive length-dependence at $\Delta T < 0$ is unambiguous and well within the accuracy and precision of the Rick-9000.

The role of electric fields in Seebeck measurements

The room-temperature J/V curves of SCn in Au^{TS}/SCn//EGaIn junctions show a slight asymmetry that is tempting to ascribe to the different work functions of the two electrodes; however, it is an intrinsic property of molecules with asymmetric anchoring groups that is present even in Au/SCn//Au junctions.⁶⁶⁻⁶⁹ The asymmetry arises because the electric field from the applied bias creates a Stark effect that shifts the positions of frontier molecular orbitals. Since transport is dominated by resonances in $\mathcal{T}(E)$ from on frontier orbital, this shift effectively pushes the transport level away from $E_{\rm F}$ at one bias and towards it at the other. This effect is shown schematically in Fig. 6a, where the sign of γ indicates the shift in the position of the frontier orbital that mediates transport E. (The distribution of the density of states should not be confused with the Lorentzian $\mathcal{T}(E)$ shown in Fig. 3.) While both are affected by Γ , $\mathcal{T}(E)$ depends on the difference between $E_{\rm F}$ and the transport level E. For example, if transport is mediated by the HOMO, then it depends on $E_{\text{HOMO}} - E_{\text{F}}$. $\mathcal{T}(E)$ can be described using a variety of different models (e.g. Fig. 3).

As discussed above, the sign of the thermovoltage (which is different from thermopower) depends on the sign of ΔT . Fig. 2 shows the thermovoltage decreasing, crossing zero and then becoming negative as the bottom electrode temperature is raised while keeping the top electrode at a constant temperature, *i.e.* changing the sign of ΔT . Yet the thermopower (Seebeck coefficient) is positive at all values of ΔT . Thus, even though *S* is measured at zero applied bias, the applied temperature gradient creates an applied electric field that includes

 V_{Contact} from the leads that is subtracted when measuring *S*; *i.e.* the total bias can be larger the measured thermovoltage of the junction. We hypothesize that this electric field, which changes sign with ΔT (Fig. 2), leads to a shift in energy that is phenomenologically related to the Stark shift that causes current-rectification under applied bias. As noted above, β^{S} is influenced by the gap-states formed by Au–S bonds, which means they are involved in transport, but their effect on conductance is not resolvable from the tunneling decay coefficient β . We therefore further hypothesize that the Stark effect is more pronounced on the gap-states than the molecular orbitals such that are pushed out of the transport window when $\Delta T < 0$. And because theoretical models do not consider gap-states, β^{S} comports to theoretical predictions only when $\Delta T < 0$.

A single-level Lorentzian to describe $\mathcal{T}(E)$ can be combined with the Mott equation (eqn (S2), ESI^{\dagger}) to compute S using Γ to model the length-dependence of alkanes and π -conjugated molecules at different energy offsets.⁶³ Inserting a γ term into this equation yields the fits shown in Fig. 6b (see Section S2.1 for details, ESI[†]). These fits to experimental data from Fig. 4 account for both directions of ΔT using reasonable values of ε = -0.96 eV and $\gamma = \pm 0.3$ eV. Here, ε is the offset between $E_{\rm F}$ and the transport level (E in Fig. 3) and -0.96 eV is in good agreement with ref. 68. This model also accounts for the two different values of β^{s} at $\Delta T > 0$ by treating the gap-states as smaller effective values of $\varepsilon = -0.40$ eV; *i.e.* the Stark effect acts primarily on the gap-states, pushing them in or out of the transport window. To convert *n* to Γ , we performed a linear fit to experimental values for Au/SCn//Au junctions.68 These values predict larger values of S that our experimental observations. Thus, the computed values of S shown in Fig. 6b are only possible with either very small values of ε or larger values of Γ ; we generated the fits using the latter strategy, adding an offset to the intercept of the linear fit of n to Γ . Furthermore, in order to reverse the length-dependence, the magnitude of the shift has to depend on *n* (see eqn (S3), ESI^{\dagger}), while for the γ parameter (Fig. 6a) for current-rectification is near-constant.⁶⁸ Nonetheless, data for both signs of β^{s} can be fit by reversing the sign of γ and the two length-dependences for $\Delta T > 0$ can be fit by changing $E_{\rm F}$ -E; the dashed lines in Fig. 6b are from $\varepsilon = -0.96$ eV, while the dotted line is $\varepsilon = -0.40$ eV, which is compatible with the hypothesis that gateway states are pushed into the transport window and contribute strongly to the transport level for shorter chains. We also stress that the introduction of a γ parameter is only necessary to fit the data for $\beta^{S} < 0$; as is shown in Fig. 3, S increases with length at a fixed ε because $-\frac{\partial \mathcal{T}}{\partial E}$ increases with length. Therefore, if the gap-states are pushed out of the transport window when $\Delta T < 0$, and transport becomes HOMO-mediated, there does not need to be any Stark effect on the HOMO to explain the behavior of $\beta^{\rm s}$.

Conclusions

We have successfully reproduced the experimental observation that Au^{TS}/SC*n*//EGaIn junctions yield two different thermopower



Fig. 6 A simplified schematic of how the applied electric field induces a Stark shift in the transport level *E* (a). Experimental values of *S* from Fig. 4 plotted with computed values of *n* using the range of *Γ* values from ref. 68 at $\varepsilon = -0.96$ eV (dashed; -0.40 eV dotted) with shifts of ± 0.03 eV (b).

length-dependences, both of which exactly oppose theoretical predictions for alkanethiols. Using the Rick-9000, a custombuilt experimental setup for measuring thermopower with EGaIn top-contacts, we determined that this unusual experimental observation only occurs when the bottom-contact is heated relative to the top-contact. When the bottom-contact is instead cooled, Au^{TS}/SCn//EGaIn junctions yield thermopower length-dependences that agree with theoretical predictions across a large range of temperatures.

The most significant finding in this study is that the thermopower of Au^{TS}/SCn//EGaIn junctions behaves as predicted without any modifications to existing theory. This is an important result because it simultaneously underscores the utility of EGaIn as a platform for studying thermoelectrics in large-area tunneling junctions and for providing fundamental insights into phenomenology that is not captured by theory. We hypothesize that the electric field created by the applied temperature gradient is not innocent and suggest that it should be considered together with the direction of the thermal gradient when modeling junctions with asymmetrical contacts, particularly at technologicallyrelevant magnitudes of thermal gradients, which are considerably larger than those typically used to study molecular junctions in laboratory settings. We were able to fit the experimental data for the aberrant length-dependence by assuming that the influence of this electric field is length-dependent. Future work is required to

	Above 0 °C		Below 0 °C		
	Ambient	Stage 1	Stage 2	Stage 3	Stage 4
Temp range	20-32	12-0	10-(-30)	10-(-60)	10-(-90)
$T_{\rm Abs}$ (K)	299	279	263	248	233
Top T (°C)	20	12	10	10	10
Bottom $T(^{\circ}C)$	24, 28, 32	8, 4, 0	0, -10, -20	-10, -30, -50	-20, -50, -80
ΔT (°C)	4, 8, 12	4, 8, 12	-10, -20, -30	-20, -40, -60	-30, -60, -90

Table 1 Description of temperature conditions

develop quantitative agreements between theory and experiment and a more detailed description of the direction of the thermal gradient in asymmetric junctions, but our results reconcile experiment with the unambiguous theoretical prediction that thermopower is positive and increases with lengths for alkanethiolates on gold.

Experimental

Molecules and materials

All reagents were used as supplied unless otherwise specified. All organic solvents were purchased from Sigma Aldrich and Acros Organics. *n* alkanenethiols (HSC*n*; *n* = 2, 4, 6, 8, 10, 12, 14, 16, 18) purity (95 to 99%) were purchased from Sigma Aldrich and Acros Organics (purity 97%). High – purity eutectic gallium indium (EGaIn; 99.99%) was obtained from Sigma Aldrich and used as supplied. All thiol derivatives were stored under N₂ atmosphere and at less 4 °C. Template stripped gold (200 nm) was thermally deposited onto silicon wafer (100 mm in diameter; 1 10 ohm per cm, 525 ± 50 μ per meter thick) by a custom built metal deposition evaporator. Optical adhesive was purchased from Norland(NOA81) and used as supplied.

Preparation and characterization of SAMs

Working in the glove-box ($O_2 < 1$ ppm, $H_2O < 1$ ppm), we prepared SAMs following the procedure reported previously.⁷⁰ A 3 mM ethanol anhydrous, 99.9% solution of CnSH was added to a vial. The solution was sealed and degassed by bubbling N_2 through the solution for 10 min. A freshly prepared Au^{TS} chip was rinsed with pure ethanol and placed in the solution with the exposed metal face up. The vial was then filled with N_2 from the glove-box environment. After 3 h incubation at room temperature, the SAM-bound Au^{TS} chip was removed from the solution and rinsed by repeatedly dipping the chip into pure ethanol (3×1 mL). The solvent on the SAM was then evaporated in air for a few seconds.

Junction preparation and thermopower measurements

A conical tip of EGaIn for use as a top contact was formed by extruding a small droplet of EGaIn from a 15 μ L syringe onto the substrate and then raising the syringe. The resulting tip was then lowered onto the SAM to form a junction of approximately 20 μ m in diameter. All junction formation and measurements in this work were carried out in an anhydrous 1% O₂ 99% N₂ environment. Thermal gradients were generated by heating the substrate or EGaIn tip and/or cooling the substrate. Voltages for

molecular ensemble junctions were measured after allowing the temperature to equilibrate and stabilize (see Fig. S1, ESI[†]) and reported as $V = V_{\text{meas}} - V_0$ where V_0 is the thermovoltage measured for the EGaIn//Au^{TS} junction (without a SAM present). To analyze thermopower of junctions, we followed the previous method reported by Yoon, Segalman *et al.* Park2019,^{20,57}

Platform temperature conditions

Thermal voltage (ΔV) is created by holding the top electrode temperature and varying the bottom temperature and vise versa. At ambient conditions, the top electrode temperature ($T_{\rm EGaIn}$) was fixed at 20 °C while the bottom electrode temperature ($Au^{\rm TS}$) was increased to 24, 28, 32 °C to create ΔT of 4, 8, 12 (Table 1). The absolute temperature $T_{\rm Abs}$ is defined by taking the average of lowest and highest electrode temperatures *i.e.* at ambient, the lowest $T_{\rm EGaIn} = 20$ + highest $T_{\rm Au^{TS}} = 32$ therefore, ($20 + 32 = 52 \div 2 = 26$ °C ≈ 299 K). At Stage 1, $T_{\rm Abs}$ of 279 K was achieved using liquid nitrogen (N₂) to lower $T_{\rm Au^{TS}}$ to 8, 4, 0 °C while $T_{\rm EGaIn}$ remained fixed at 12 °C creating ΔT of -4, -8, -12. The stability of electrode temperatures is accounted for in Fig. S1 (ESI†).

At Stage 2 ($T_{Abs} = 263$ K), $T_{Au^{TS}}$ was lowered to 0, -10, -20 °C while the T_{EGaIn} remained fixed at 10° creating ΔT of -10, -20, -30. T_{Abs} of 248 K at Stage 3, is a further lowering of $T_{Au^{TS}}$ to -10, -30, -50 while T_{EGaIn} remained fixed at 10 °C which created a ΔT of -20, -40, -60 °C. The lowest T_{Abs} of 233 K at Stage 4, was a further lowering of $T_{Au^{TS}}$ to -20, -50, -80 °C while T_{EGaIn} remained fixed at 10 °C creating a ΔT of -30, -60, -90. The sign + /or – of the gradient is determined from the standard calculation $\Delta T = T_{Au^{TS}} - T_{EGaIn}$. A positive gradient is defined as $+\Delta T = T_{Au^{TS}}^{Hot} - T_{EGaIn}^{Cold}$ and a negative is $-\Delta T = T_{Au^{TS}}^{Cold} - T_{EGaIn}^{Hot}$. The minimum to maximum bottom electrode temperature is defined in Fig. 2 as temperature range.

Data availability

The data supporting this article have been included as part of the ESI.[†] The open-source software used to process raw I/V data can be downloaded *via* this DOI: https://doi.org/10.5281/zenodo.6422417.

Conflicts of interest

There are no conflicts to declare.

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