Origin of Hydrophilic Surface Functionalization-Induced Thermal Conductance Enhancement across Solid–Water Interfaces

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ABSTRACT: Thermal transport across solid–water interfaces is critical for a wide range of applications such as solar thermal evaporation, nanoparticle-assisted hyperthermia therapeutics, and nanofluids. Surface functionalization using self-assembled hyperthermia monolayers (SAMs) to change the hydrophilicity of the solid surface is a common strategy to improve the thermal conductance of solid–water interfaces. Although it is known that hydrophilic interfaces increase the interfacial bonding, how it impacts the molecular level energy transport across the interface is still not clear. In this paper, we perform molecular dynamics simulations to calculate the thermal conductance of differently functionalized gold (Au)–water interfaces. Combining the heat flux decomposition to different interatomic interactions across interfaces and analyses of water structures close to the functionalized surfaces, we found that there is a collaborative effect from the electrostatic interactions and the Lennard-Jones (L-J) interactions (especially the repulsive part). The electrostatic interactions, which are between the polar functional groups of SAMs and water, will attract water molecules closer to the SAM surface, leading both the electrostatic and L-J interactions to have larger effective forces across the interfaces. This increases the power exchanged between solid and water atoms, enhancing the thermal energy transport. The results from this work will provide new insights to the understanding of thermal transport across solid–water interfaces.

KEYWORDS: hard–soft interface, self-assembled monolayer, interfacial thermal conductance, nonequilibrium molecular dynamics, interfacial heat flux

INTRODUCTION

Thermal transport across solid–water interfaces is critical for a wide range of applications from solar thermal water treatment and nanofluids to nanoparticle-assisted photothermal therapeutics. The interfacial thermal resistance can play important or even dominant roles in heat transfer when the system’s constituent length reaches the nanometer scale. Surface functionalization using self-assembled monolayers (SAMs) to change the interfacial adhesion energy between the hard and soft interface is a common strategy to improve the interfacial thermal conductance (ITC) and it has been studied both computationally and experimentally. Research has found positive relationships between the interfacial adhesion energy and ITC across the solid–water interfaces experimentally. Similar conclusions have also been obtained from a number of molecular dynamics (MD) simulations. Although it is generally known that stronger interfacial adhesion energy can enhance the ITC, how it impacts the molecular level energy transport mechanism across the interface is still not clear. Because interfacial thermal transport is related to the dynamics of atomic motion and interatomic forces between the atoms at each side of the interface, the interfacial adhesion energy, which is a static property, cannot be directly linked to the thermal conductance.

In this study, we use nonequilibrium MDs (NEMD) simulations to study the thermal transport across gold (Au)–water interfaces with the Au surfaces functionalized by different SAMs. Three different SAMs are studied to systematically change the surface hydrophobicity. A positive relation between the interfacial binding energy and the ITC is observed. To understand the underlying mechanism, we characterize the interfacial heat flux via the atomic velocity and interfacial forces. We also decompose the interfacial interaction and thermal transport into contributions from Lennard-Jones (L-J) and electrostatic forces and relate their impacts on the interatomic distances between the SAM head groups and water molecules to ITC. It is found that there is a collaborative effect from the electrostatic interaction and the repulsive part of the L-J interaction. The electrostatic interactions, which are between the highly polarized functional groups of SAMs and water, will attract water molecules closer to the solid surface, leading both electrostatic and repulsive interactions to have larger effective forces across the interface, which in turn result in larger ITC. These results may provide new insights to the understanding of the mechanism of thermal transport across solid–water interfaces.

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In this study, the model consisted of three types of thiol SAM molecules chemically adsorbed on the Au(111) surface with explicitly defined TIP3P model water molecules. Three different terminal groups were studied including 1-hexanethiol (HS–(CH₂)₅–CH₃), 6-mercapto-1-hexanol (HS–(CH₂)₆–OH), and 6-mercaptopentanoic acid (HS–(CH₂)₅–COOH). For brevity, they are denoted as −CH₃ SAM, −OH SAM, and −COOH SAM, respectively (Figure 1). These three types of SAM molecules feature functionalizations with a range of hydrophilicities. The expected hydrophilicity with respect to water increases from the −CH₃ SAM to the −OH SAM and then to the −COOH SAM, as the head groups become more polarized. Specifically, the −COOH SAM has an epoxy and a hydroxyl group, which can both form hydrogen bonds with neighboring water molecules and thus should provide the highest interfacial energy among these three types of SAMs. This trend has been experimentally observed through the hysteresis contact angle experiment from Harikrishna et al.²² We should mention that the interface energy among these three types of SAMs. This trend has been previously used for SAM-on-Au structures. The nonbonded interactions between Au and other atoms are simulated by the L-J potential. The water molecules are modeled using the TIP3P model. The polymer consistent force field, which reproduces water thermodynamic and structural properties well, was used to simulate the gold atoms and the bond interaction between the Au substrate and the sulfur (S) atoms in the thiol SAM molecules. Such a potential has been previously used for SAM-on-Au structures successfully. The water molecules are modeled using the TIP3P model, which reproduces water thermodynamic and structural properties well. The polymer consistent force field is adopted to simulate the SAM molecules, which has been successfully used for thermal transport studies involving SAMs. The nonbonded interactions between Au and other atoms are simulated by the L-J interaction

\[ E(r) = D_0 \left( e^{-2\alpha (r-r_0)} - 2e^{-\alpha (r-r_0)} \right) \]  

where \( D_0 \) is the bond-dissociation energy, \( r_0 \) is the equilibrium bond length, and \( \alpha \) is the parameter characteristic of the atom, which is used to simulate the gold atoms and the bond interaction between the Au substrate and the sulfur (S) atoms in the thiol SAM molecules. Such a potential has been previously used for SAM-on-Au structures successfully. The water molecules are modeled using the TIP3P model, which reproduces water thermodynamic and structural properties well. The polymer consistent force field is adopted to simulate the SAM molecules, which has been successfully used for thermal transport studies involving SAMs. The nonbonded interactions between Au and other atoms are simulated by the L-J interaction

\[ E = 4\epsilon \left( \frac{\sigma}{r_{ij}} \right)^{12} - 2\epsilon \left( \frac{\sigma}{r_{ij}} \right)^6 \]  

where \( \epsilon \) and \( \sigma \) are the representative scales of energy and length and \( r_{ij} \) is the distance between two molecules, \( i \) and \( j \). L-J interaction parameters are chosen from a modified universal force field, except for the S atoms which are bound covalently to Au. All pair coefficients associated with Au atoms are listed in Table 1. A cutoff of 8 Å is used for the Morse potential, and 10 Å is chosen for the L-J interactions. The long-range electrostatic interaction in the entire system is computed by the particle–particle–particle–mesh approach with an accuracy of 1 × 10⁻⁶. Simulations are performed using the large-scale atomic/molecular massively parallel simulator. A time step size of 0.25 fs is chosen because of the presence of lightweight hydrogen atoms. An example simulation setup is shown in Figure 1a. Two solid gold (111) substrates with face-centered cubic lattice were placed at both ends (50 layers on each side) of the simulation box. The SAMs are placed by generating an 8 × 8 array of single thiol chains with the initial tilt angle of 30° and a S–S spacing of 0.497 nm. A total of 2000 water molecules are compacted into the space between the two substrates using Packmol.

NEMD simulations are performed to calculate the ITC. The Langevin thermostats are used to thermostat the heat source at 310 K and the heat sink at 210 K so that one of the interfaces is near room temperature (~300 K). For the simulation procedure, the simulation system is first equilibrated in a canonical ensemble (NVT) at 260 K for 0.5 ns. The system is optimized in an isothermal–isobaric ensemble (NPT) at 1 atm and 260 K for another 1 ns. After the

Table 1. Morse Potential \( D_0, \alpha, \) and \( r_0 \) Parameters for Interactions within Au and between Au and S Atoms

<table>
<thead>
<tr>
<th>pair type</th>
<th>( D_0 ) (kcal/mol)</th>
<th>( \alpha ) (Å⁻¹)</th>
<th>( r_0 ) (Å)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Au–Au</td>
<td>10.954</td>
<td>1.583</td>
<td>3.042</td>
</tr>
<tr>
<td>Au–S</td>
<td>8.763</td>
<td>1.47</td>
<td>2.65</td>
</tr>
<tr>
<td>Au–C₁</td>
<td>0.0634</td>
<td>3.182</td>
<td></td>
</tr>
<tr>
<td>Au–C₂</td>
<td>0.0634</td>
<td>3.182</td>
<td></td>
</tr>
<tr>
<td>Au–H₆</td>
<td>0.0414</td>
<td>2.752</td>
<td></td>
</tr>
<tr>
<td>Au–O₁</td>
<td>0.0414</td>
<td>2.752</td>
<td></td>
</tr>
<tr>
<td>Au–O₂</td>
<td>0.0484</td>
<td>3.026</td>
<td></td>
</tr>
<tr>
<td>Au–O₃</td>
<td>0.0484</td>
<td>3.026</td>
<td></td>
</tr>
<tr>
<td>Au–O₄</td>
<td>0.141</td>
<td>3.6</td>
<td></td>
</tr>
</tbody>
</table>

“L-J potential \( \epsilon \) and \( \sigma \) parameters between Au atoms and the rest atoms. “C₁” represents the carbon atom in the methyl group of the −CH₃ SAM, and “C₂” represents the carbon atom in the backbone of SAMs. “H₆” represents the hydrogen atom connected to carbon atoms. “O₁” represents the oxygen atom in the hydrogen atoms in the hydroxyl and carboxyl groups in the −OH and −COOH SAMs. “O₂” represents the oxygen atoms in the hydroxyl groups in the −OH and −COOH SAMs. “O₃” is the double-bond oxygen atom in the carboxyl group in the −COOH SAM. “O₄” represents the oxygen atom in water molecules.
structures are fully equilibrated, to prevent the heat leakage during the heat-transfer process, we fixed the last three layers of gold atoms at each side and add extra vacuum space. Then the microcanonical ensemble (NVE) is then applied to the simulation, which lasts 3 ns, and the last 1 ns is chosen as the production period to calculate the ITC and the error bars are the standard deviations of the data from the period. Figure 1b shows a typical steady-state temperature profile. When the nonequilibrium state becomes steady, the ITC is calculated using \( G = q/\Delta T \), where \( G \) is the ITC, \( q \) is the heat flux, and \( \Delta T \) is the temperature gap at the higher temperature interface (\( \sim 300 \) K) obtained by extrapolating the linear fits of temperature profiles of Au and water to the interface.

## RESULTS AND DISCUSSION

The calculated ITC values are 51 ± 9, 155 ± 13, and 187 ± 9 MW/m² K for the −CH₃, −OH, and −COOH SAM-functionalized Au−water interfaces, respectively. Our results agree reasonably with time-domain thermoreflectance experiments, which reported ITC values of 65 ± 5 and 190 ± 30 MW/m² K for a −CH₃ SAM- and a −OH SAM-functionalized Au−water interface, respectively. The slight differences between our simulation results and those from experiments may be attributed to the uncertainty in the experimental sample (e.g., defects) and the accuracy of the MD potentials. The interface energies of these three interfaces are, respectively, 68 266 and 340 kcal/mol. As expected, the polarized SAM interfaces (−OH and −COOH SAMs) have much larger interface energies than that of the nonpolar SAM interface (−CH₃ SAM), which agrees with experiments. A couple of experimental measurements show that the −OH SAM and the −COOH SAM have very similar water contact angles within the uncertainty of each other, but the −OH SAM is slightly more hydrophilic than the −COOH SAM. We note that these measurements only measured the advancing contact angle without considering the hysteresis effect. The real contact angle for the SAM surface should range between the advancing and the receding contact angles. Harikrishna et al. indeed find that the receding contact angle of the −COOH SAM is smaller than that of the −OH SAM. In addition, factors such as the pH and surface roughness in Keselowsky’s experiment might also have impacted the measured contact angles. Nevertheless, our simulation results are self-consistent within the scope of this present study, and we believe that the intrinsic interfacial binding energy of the −COOH SAM interface should be larger than that of the −OH SAM interface because −COOH can provide two hydrogen bonding sites.

As expected, we can see that the interface with the most polarized −COOH SAM has the highest value of ITC, whereas that with the nonpolar −CH₃ SAM has the lowest. We plot the ITCs of the three different systems as a function of the corresponding absolute values of the interfacial adhesion energy, calculated as the sum of all interactions between the water and the SAM-functionalized Au substrate, and found a linear relation in the data (Figure 2). Such an observation is consistent with the previous results from MD simulations and experiments. Although this trend is known to be common, the microscopic origin of such observations has never been studied in detail.

At the molecular level, the thermal energy transport across the interface is realized by atoms at either side of the interface doing work to one another across the interface. The rate of this work (i.e., power, \( p \)) exchanged between the SAM-functionalized substrate and water can be calculated as

\[
p = \frac{1}{2} \left( \sum_{i \in \text{sub}/j \in \text{w}} F_{ij}v_{ij} - \sum_{i \in \text{sub}/j \in \text{w}} F_{ji}v_{ji} \right)
\]

where \( F_{ij} \) is the force from atom \( j \) exerted on atom \( i \) and \( v \) is the atom velocity. The indexes \( i \) and \( j \) belong to the SAM-functionalized substrate (sub) and water (w), respectively. We note that this equation for the interface heat flux can be shown equivalent to the equation by Torii et al. who derived a similar equation for bulk heat flux from the Irving and Kirkwood relation. This power normalized by the interface area leads to the heat flux \( q = p/A \), where \( A \) is the cross-sectional area. As can be seen, the heat flux across the interface is related to the dynamics of the atoms (i.e., \( v \)) and interatomic forces. It is not directly related to the interfacial adhesion energy but instead its derivative (i.e., force, \( F \)). Furthermore, the total exchanged power can be separated using the following equation:

\[
p = p^{L-J} + p^{Q}
\]

\[
= \frac{1}{2} \left( \sum_{i \in \text{sub}/j \in \text{w}} F_{ij}^{L-J}v_{ij} - \sum_{i \in \text{sub}/j \in \text{w}} F_{ji}^{L-J}v_{ji} \right) + \frac{1}{2} \left( \sum_{i \in \text{sub}/j \in \text{w}} F_{ij}^{Q}v_{ij} - \sum_{i \in \text{sub}/j \in \text{w}} F_{ji}^{Q}v_{ji} \right)
\]

The ITC can thus also be decomposed into the L-J and the electrostatic portions according to \( G^{L-J} = q^{L-J}/\Delta T \) or \( G^{Q} = q^{Q}/\Delta T \). As shown in Figure 3a, when the surface functionalization changes from the nonpolar −CH₃ SAM to the polar −OH SAM, both the L-J and the electrostatic portions of the ITC increase dramatically. From the −OH SAM to the more polar −COOH SAM, the electrostatic portion of ITC does not show an obvious increase, but the L-J portion is enhanced significantly. Such a finding is slightly surprising, because it would be intuitive to postulate that it is the increased electrostatic interaction due to the more polar SAM groups that contribute mainly to the increase in ITC. However, our data show that as the SAM becomes polarized, the stronger electrostatic interaction will not only enhance the thermal energy transported by the electrostatic forces themselves but also enhance that by the L-J forces. In Figure 3b, the total
interfacial adhesion energy is also decomposed into the L-J and electrostatic portions. Similar to the ITC decomposition, both L-J and electrostatic adhesion energy increase significantly from the $-CH_3$ to the $-OH$ SAM. From the $-OH$ to the $-COOH$ SAM, the increase in L-J energy is obvious, whereas the electrostatic portion only increases slightly. Such effects are consistent with the SAM-induced thermal transport enhancement across interfaces,

Because the atomic velocities are expected to be similar in all three cases because of the same temperature, according to eq 1, the interatomic forces are the only reason for the above observation in ITC (Figure 3a). To analyze the forces across the interfaces, the number distribution of the pairwise forces across the interface as a function of the force magnitude is shown in Figure 4 as a histogram, where electrostatic and L-J forces are plotted separately. The positive force indicates repulsion between two atoms, whereas the negative force means attraction. There is a large population around zero in both Figure 4a,b, which indicates weak forces between atoms that have long distances from each other. Panels c and d in Figure 4 display the same data as those in panels a and b but with different view ranges in the x-axis to enable a clearer comparison. From the nonpolar SAM ($-CH_3$) to the highly polar SAM ($-COOH$), the number of interacting atom pairs increases for both electrostatic and L-J interactions as shown in Figure 4c,d (i.e., higher number distribution of pairwise interactions from the blue line to the red line and then to the orange line). This trend is also confirmed in Table 2, where we calculate the total L-J and electrostatic force and the total number of pair interactions for each interface using $f_{\text{total}} = \sum N f$, where $N$ is the number of interaction pairs and $f$ is the force between the pair.

Table 2 also shows that for the $-CH_3$ SAM interface, the attractive forces (negative) are from the L-J interaction, whereas the electrostatic forces are mainly repulsive. On the contrary, for the $-OH$ and $-COOH$ SAM interfaces, the
interface (SAM changes from the nonpolar density along the in the vicinity of the interfaces. We implication can be proven by characterizing the water structure molecules via both electrostatic and L-J interactions. This thus, there are more water molecules interacting with the SAM should have led to a higher water density at the interface, and groups attracts water molecules closer to the interface, which that the stronger electrostatic interaction due to the polar SAM number of interacting atom pairs. These observations suggest −polar peaks near the interface also increase, which suggests that there −di −e −f −fl −fi −ff −fl −fi - attractive forces are from the electrostatic interaction, whereas the L-J forces are repulsive. Because the repulsive part in the L-J interaction can have a larger gradient than that of the attractive part, the overall repulsive forces can be larger than the attractive forces across the interface. Because the heat flux (eqs 3 and 4) is expressed as the force multiplied by the atomic velocity, the repulsive part in the L-J interaction can contribute more significantly to the ITC. We further calculate the summation of the magnitudes of the repulsive forces and that of the attractive forces across the interfaces and found their ratios to be 1.1, 1.9, and 2.1 for the −CH3, −OH, and −COOH SAM interfaces, respectively. The repulsive forces are indeed larger than the attractive forces in all cases, and this difference increases as the SAM molecules become more polarized.

In addition, Table 2 also shows that the more polar the interface (−COOH > −OH > −CH3) is, the larger the number of interacting atom pairs. These observations suggest that the stronger electrostatic interaction due to the polar SAM groups attracts water molecules closer to the interface, which should have led to a higher water density at the interface, and thus, there are more water molecules interacting with the SAM molecules via both electrostatic and L-J interactions. This implication can be proven by characterizing the water structure in the vicinity of the interfaces. We first plot the water mass density along the z-direction as shown in Figure 5a. When the SAM changes from the nonpolar −CH3 group to the highly polar −COOH group, the water density profile moves closer to the SAM surface. This suggests that the stronger electrostatic interaction can attract water molecules closer to the interface, reducing the interaction distance and thus leading to larger effective forces from both the electrostatic interaction and the L-J interaction. Furthermore, the amplitudes of the water density peaks near the interface also increase, which suggests that there are more atoms participating in the power exchange across the interface. Both factors will lead to more efficient power exchange across the interfaces, thus resulting in higher ITC.

The local water distribution near the interface is also characterized by the radial distribution function (RDF, Figure 5b) calculated as \( g(r) = n(r)/(4\pi\rho \Delta r) \), where \( n(r) \) is the number of atoms in a shell of thickness \( \Delta r \) at a distance \( r \) from the reference atom and \( \rho \) is the average water atom number density. For the −CH3 SAM, we take the C atom in the CH3 end group as the reference atom, whereas for the −OH and −COOH SAMs, the reference atoms are chosen as the oxygen atom in the hydroxyl group. As expected, the RDF shows that from −CH3 to −OH and then to −COOH SAM, the peaks shift from the right to the left, which directly indicate progressively shorter distances between the thiol end groups and water molecules.

**CONCLUSIONS**

In summary, we have calculated the ITC of Au−water interfaces with different SAM functionalizations. We find that when the SAM end group becomes more polar, the ITC becomes larger. Combining the heat flux decomposition to different interatomic interactions across interfaces and analyses of water structures close to the functionalized surfaces, we found that there is a collaborative effect from electrostatic and L-J interactions. The electrostatic interactions, which are between the polar functional groups of SAMs and water, will attract water molecules closer to the solid surface, leading both the electrostatic interaction and the L-J interaction (especially the repulsive part) to have larger effective forces across the interfaces. This increases the power exchanged between solid and water atoms, enhancing the thermal energy transport. The results from this work will provide new insights to the understanding of thermal transport across solid−water interfaces.

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Table 2. Total L-J and Electrostatic Forces and Total Number of Pair Interactions within the Cutoff between the Substrate (Au + SAM) and Water for the Three Interfaces

<table>
<thead>
<tr>
<th>interface</th>
<th>total L-J</th>
<th>total electrostatic</th>
<th>total pair numbers</th>
</tr>
</thead>
<tbody>
<tr>
<td>−CH3</td>
<td>−8.3 × 10^6</td>
<td>2.6 × 10^6</td>
<td>3.7 × 10^4</td>
</tr>
<tr>
<td>−OH</td>
<td>8.9 × 10^5</td>
<td>−9.7 × 10^5</td>
<td>4.3 × 10^4</td>
</tr>
<tr>
<td>−COOH</td>
<td>1.3 × 10^5</td>
<td>−1.5 × 10^6</td>
<td>4.8 × 10^4</td>
</tr>
</tbody>
</table>

“The force unit is kcal mol\(^{-1}\) Å\(^{-1}\).”

**Figure 5.** (a) Water density profiles near the −CH3, −OH SAM, and −COOH SAM surfaces. (b) RDF of the oxygen atoms in water molecules with respect to the carbon atoms in the methyl group of the −CH3 SAM and the oxygen atoms in the hydroxyl groups of the −OH and −COOH SAMs.

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Notes

The authors declare no competing financial interest.

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