Ultrafast Heat Transfer in Nanoscale Materials

David Cahill, Gyung-Min Choi, Jonglo Park, Jingyu Huang, Wei Wang, Rich Wilson

Department of Materials Science and Engineering, Materials Research Laboratory, University of Illinois at Urbana-Champaign

thanks to Cathy Murphy, Byoung-Chul Min, and Kyung-Jin Lee

supported by DOE-BES, ARO and ONR
• Introduction: Heat transfer on length scales of nanometers and time-scales of picoseconds

• How fast can a nanoparticle cool?
  – Transfer of vibrational thermal energy across an interface

• How fast can heat be exchanged between two metals?
  – Transfer of electronic thermal energy across an interface and between electrons and phonons

• What is the largest heat current we can pass through a nanoscale ferromagnetic layer?
  – Thermal generation of spin currents by demagnetization and the spin-dependent Seebeck effect.
Thermal transport coefficients

- Thermal conductivity $\Lambda$ is a property of the continuum

$$\vec{J} = -\Lambda \vec{\nabla}T$$

$$\Lambda = \frac{1}{3V k_B T^2} \int_0^\infty \langle \vec{j}(t) \cdot \vec{j}(0) \rangle \, dt$$

- Thermal conductance (per unit area) $G$ is a property of an interface

$$\vec{J} = G \Delta T$$

$$G = \frac{1}{Ak_B T^2} \int_0^\infty \langle q(t)q(0) \rangle \, dt$$
Thermal transport coefficients

- Thermal conductivity $\Lambda$ appears in the diffusion equation

$$C \frac{dT}{dt} = \Lambda \nabla^2 T$$

$C = \text{heat capacity per unit volume}$

Diffusivity: $D = \frac{\Lambda}{C}$

Effusivity: $\varepsilon = \sqrt{\Lambda C}$

- Interface thermal conductance $G$ is a radiative boundary condition

$$G(T_+ - T_-) = \Lambda \left. \frac{dT}{dz} \right|_{z=0}$$

Kapitza length: $L_K = \frac{\Lambda}{G}$
Thermal conductivities of dense solids span a range of 40,000 at room temperature.

Adapted from Goodson, *Science* (2007)
Interface conductance spans a factor of 60 range at room temperature.

Lyoe and Cahill, PRB (2006)
Heat capacity per unit volume of solids spans only a factor of 4 at room temperature

<table>
<thead>
<tr>
<th>Material</th>
<th>C (MJ m⁻³ K⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>water</td>
<td>4.18</td>
</tr>
<tr>
<td>Ni</td>
<td>3.95</td>
</tr>
<tr>
<td>Al</td>
<td>2.42</td>
</tr>
<tr>
<td>Diamond</td>
<td>1.78</td>
</tr>
<tr>
<td>Polymer (PMMA)</td>
<td>1.8</td>
</tr>
<tr>
<td>PbTe</td>
<td>1.2</td>
</tr>
</tbody>
</table>
Technology drivers for fundamental studies of ultrafast heat transfer at the nanoscale

Proposed next generation phase change memory device

Technology drivers for fundamental studies of ultrafast heat transfer at the nanoscale

Velocity of media $10 \text{ m s}^{-1}$. Size of bit $10 \text{ nm}$. Time-scale $1 \text{ ns}$.

I. How fast can a rapidly-heated nanoparticle cool?

- Limited by interface conductance.
  - Equivalent to discharging of a capacitor through a resistor
    \[
    \tau_G = (VC) \left( \frac{1}{AG} \right) = \frac{1}{3} \frac{rC}{G}
    \]
  - Order of magnitude estimate
    \[
    r = 3 \text{ nm}; \ C = 3 \text{ MJ m}^{-3} \text{ K}^{-1}; \ G = 100 \text{ MW m}^{-2} \text{ K}^{-1}
    \]
    \[
    \tau_G = 30 \text{ ps}
    \]
I. How fast can a rapidly-heated nanoparticle cool?

- Limited by effusivity of the surrounding fluid
  - Solvable but let’s instead approximate by asking “when does the heat capacity of a layer of the thermal diffusion distance in the fluid equal the heat capacity of the particle?”

\[
(4\pi r^2)(\sqrt{D\tau_E}) C_f = \frac{4\pi}{3} r^3 C_p
\]

\[
C_f \approx C_p = C; \quad \tau_E = \frac{r^2}{9D}
\]

\[
r = 3 \text{ nm}; \quad D = 10^{-7} \text{ m}^2 \text{ s}^{-1}
\]

\[
\tau_E = 100 \text{ ps}
\]

Wilson et al., PRB (2002)
Time-domain thermoreflectance

Kang et al., RSI (2008)
Time-domain thermoreflectance

Clone built at Fraunhofer Institute for Physical Measurement, Jan. 7-8 2008
• Optical constants and reflectivity depend on strain and temperature

• Strain echoes give acoustic properties or film thickness

• Thermoreflectance $dR/dT$ gives thermal properties
Thermoreflectance is only one of many methods for ultrafast optical thermometry

- Transient adsorption using plasmon resonances of Au nanostructures is sensitive to both the temperature of the Au and the surrounding dielectric.
  - At a wavelength near the peak absorption, only the Au temperature is important.

Huang et al, ACS Nano (2012)
Ultrafast thermal analysis of surfactant layers surrounding Au nanorods

- Vary surfactant thickness by changing concentration of CTAB in aqueous solution.

- Index of refraction of CTAB surfactant is larger than water and creates a red-shift of the plasmon resonance.

- CTAB forms micelles in water at a concentration of \( \approx 1 \text{ mM} \) (critical concentration)

Huang et al, ACS Nano (2012)
Ultrafast thermal analysis of surfactant layers surrounding Au nanorods

- Measure surfactant thickness by dynamic light scattering to determine the hydrodynamic diameter $d_H$

Huang et al., ACS Nano (2012)

Chen et al., Chem. Soc. Rev. (2013)
Ultrafast thermal analysis of surfactant layers surrounding Au nanorods

- Heat capacity and thermal conductivity of CTAB surfactant layers as a function of CTAB concentration in solution.
- Heat capacity is constant
- Thermal conductivity decreases above critical concentration.

Huang et al, ACS Nano (2012)
II. How fast can a laser pulse heat a metal layer?

- Electrons are heated directly by the laser pulse, i.e., electric fields in the optical pulse create electronic excitations.
  - Heat capacity of electrons is relatively small,
    \[ C_{el} \approx N_{el} k_B \left( \frac{T}{T_F} \right); \ T \ll T_F \]

- Takes time to heat the atomic vibrations. Most of the heat capacity of the solid is in the lattice vibrations, i.e., the phonons. Classical limit: \[ C_{ph} \approx 3Nk_B \]

- Phenomenological two-temperature model treats each system as a thermal reservoir coupled by a thermal conductance (per unit volume) \( g_{ep} \)
II. How fast can a laser pulse heat a metal layer?

Electron thermalization time is on the order of 300 fs but varies over a wide range

\[ \tau \approx \frac{C_{el}}{g_{ep}} \sim \frac{3 \times 10^4 \text{ J m}^{-3} \text{ K}^{-1}}{10^{17} \text{ W m}^{-3} \text{ K}^{-1}} \sim 300 \text{ fs} \]

\[ 0.1 \leq \tau \leq 1 \text{ ps} \]
II. How fast can a laser pulse heat a metal layer?

- Au provides well-defined chemistry for studies of molecular layers, however
  - hot-electron effects are a problem if we want to produce a large change in the Au lattice temperature

II. How fast can a laser pulse heat a metal bilayer?

- Let’s see what happens if we try to avoid the hot electron effects (keep the system closer to equilibrium) by adding a Pt layer with strong electron-phonon coupling and large electronic heat capacity

Wang and Cahill, PRL (2012)
II. How fast can a laser pulse heat a metal bilayer?
Heating of the Au layer is slow because electron-phonon coupling in Au is the smallest conductance in the problem.

- Characteristic time scale for the heating of the Au phonons:
  \[ \tau = \frac{C_{\text{Au,ph}}}{g_{\text{Au,ep}}} \approx 80 \text{ ps} \]

Wang and Cahill, PRL (2012)
Lousy way to heat Au but excellent way to measure $g_{ep}$

- Solid lines are the predictions of the original Kaganov “two-temperature” model of 1957
- Dashed lines are $T^4$ extrapolations of low temperature physics experiments.

Wang and Cahill, PRL (2012)
Extend the story: pump and probe Au/Pt bilayer from different sides

Data and transmission line modeling of Pt (23 nm)/Au(58 nm)

- Attempt to measure the Au/Pt electronic interface conductance
- Increase Au thickness to 60 nm to increase heat flux from Pt to Au
- Can only set a lower limit $G_{ee} > 5$ MW m$^{-2}$ K$^{-1}$

Choi et al., PRB (2014)
• The big picture question: “How can we write magnetic information without resorting to magnetic fields, e.g., with spin currents?”
  – Rapid changes in magnetization and strong temperature gradients in magnetic materials should produce spin currents.
  – Magnitudes of the effects are only beginning to be understood.
  – Create huge heat currents 100 GW m⁻² and detect spin current in real time with 1 ps time resolution.
Subset of an emerging topic of “spin caloritronics”

- Electronic states enumerated by energy, wave-vector, spin

Tri-layer structure to study effects of rapid heating and high heat fluxes on ferromagnets

- Deposit laser pulse energy in Pt film (30 nm) with strong electron phonon coupling
- Heat flows through ferromagnetic (6 nm) layer and into the Cu heat sink (80 nm)

Tri-layer structure to study effects of rapid heating and high heat currents on ferromagnets

- Back-of-the-envelope analysis for size of heat flux
  - Restrict temperature rise to $\Delta T = 100$ K
  - Conductances acting in series
    - Metal-metal interface
    - Thermal conductivity of Co/Pt
    - Electron-phonon coupling
  - Heat current

$$J_Q = G_{\text{series}} \Delta T \sim 200 \text{ GW m}^{-2}$$
Time-resolved magneto-optic Kerr effect (TR-MOKE) to measure spin density or temperature through $M(T)$

Körmann et al., PRB (2011)

http://labfiz.uwb.edu.pl
Schematic of TR-MOKE Setup (Reflectance Configuration)

- Magnet
- Sample
- Objective
- Filter
- Convex lens
- Half wave plate
- Wollaston Prism
- Pump
- PBS
- NBS
- Probe
- CCD
- Balanced detector
Pump Pt-side, probe either Pt-side or Cu side by either TDTR or TR-MOKE

Comparison between experiment and spin diffusion model using spin generation $= dM/dt$

Measured Kerr signal on Cu side

Spin diffusion model
Spin diffusion model

\[ \frac{\partial \mu_s}{\partial t} = D \frac{\partial^2 \mu_s}{\partial^2 z} - \frac{\mu_s}{\tau_s} \]

spin generation rate per unit volume

\[ G_s = -\frac{dM}{dt} \]

\( \mu_s = \mu_\uparrow - \mu_\downarrow \) is the spin chemical potential

\( D \) is the spin diffusion constant

\( \tau_s \) is the spin relaxation time

<table>
<thead>
<tr>
<th></th>
<th>Pt</th>
<th>[Co/Pt]</th>
<th>Cu</th>
</tr>
</thead>
<tbody>
<tr>
<td>( D ) (nm(^2)/ps)</td>
<td>200</td>
<td>100</td>
<td>6500</td>
</tr>
<tr>
<td>( \tau_s ) (ps)</td>
<td>0.5</td>
<td>0.05</td>
<td>25</td>
</tr>
<tr>
<td>( (D\tau_s)^{1/2} ) (nm)</td>
<td>10</td>
<td>2.2</td>
<td>400</td>
</tr>
</tbody>
</table>
Temperature gradient also contributes to spin accumulation

\[ J_S = -\frac{\mu_B}{e} \left( \sigma_{\uparrow \uparrow} S_{\uparrow} - \sigma_{\downarrow \downarrow} S_{\downarrow} \right) \nabla T = -\frac{\mu_B}{e} \frac{\sigma_{\uparrow \uparrow} S_{\uparrow} - \sigma_{\downarrow \downarrow} S_{\downarrow}}{\sigma_{\uparrow} + \sigma_{\downarrow}} \sigma \nabla T \]

Temperature gradient in the Pt/Co layer from thermal modeling

Calculated spin accumulations

\[ J_Q = -\Lambda_{FM} \frac{dT}{dx} \]

\[ \frac{\sigma_{\uparrow \uparrow} S_{\uparrow} - \sigma_{\downarrow \downarrow} S_{\downarrow}}{\sigma_{\uparrow} + \sigma_{\downarrow}} \approx 5 \mu \text{V K}^{-1} \]
Temperature gradient also contributes to spin accumulation

- More refined data with comparison to spin diffusion model including the spin-dependent Seebeck effect

Summary

• How fast does a nanoparticle cool?
  – Scales with $r$ for small $r$, $r^2$ for larger radius.
  – Order of 100 ps for $r=3$ nm.
  – Can independently analyze heat capacity and thermal conductivity of surfactant layers using plasmonic nanoparticles as the temperature sensor.

• How fast can we heat a metal layer?
  – Single layer, 0.1 to 1 ps due to electron-phonon coupling
  – Bilayers are surprisingly complicated and can be surprisingly slow, on the order of 100 ps.

• How large of heat flux can we generate and can you do anything with it?
  – 100 GW m$^{-2}$
  – Exploring thermal generation of spin currents and thermally-enabled manipulation of magnetic information