

Light-induced magnetic waves in materials engineered at the atomic scale

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This image depicts the magnetic ordering, i.e. the antiparallel spin alignment, present in the NdNiO3 thin film that is grown on a LaAlO3 crystal substrate. A short mid-infrared laser pulse triggers vibrations in the substrate, indicated by the smeared oxygen atoms in red. This atomic motion initiates melting of the magnetic order in the functional film, starting locally at the interface between the two materials and progressively moving into the interior of the film. Credit: Max Planck Institute for the Structure and Dynamics of Matter



A new study discovers how the sudden excitation of lattice vibrations in a crystal can trigger a change of the magnetic properties of an atomicallythin layer that lies on its surface.

A research team, led by scientists of the Max Planck Institute for the Structure and Dynamics of Matter at CFEL in Hamburg, the University of Oxford, and the Université de Genève, used extremely short X-ray pulses from a free-electron laser to discover that melting of magnetic order in the thin layer is initiated at its interface with the substrate and progressively moves into the interior of the film on an ultra-short time scale. This novel type of ultrafast light control in materials engineered at the atomic scale may lead to new prospects in magnetic storage technologies. The results are reported online in the journal *Nature Materials*.

Transition metal oxides, such as manganites, cuprates or nickelates, have attracted enormous interest among researchers since their electric and magnetic properties can be altered by even subtle changes of external parameters, such as temperature, and electric or magnetic fields. "But there is also a strong correlation between the atomic arrangement of the crystal lattice and these properties, so that controlled structural modifications allow for tuning the electronic and magnetic state of these materials, too," said Michael Först, scientist at the Max Planck Institute and one of the lead authors of this work.

More recently, researchers started studying heterostructures composed of different oxide materials. An atomically-thin oxide film on a substrate can have properties very different from its bulk form due to a variety of interface effects, including mechanical strain produced at the interface between the substrate and the film. This makes complex oxide heterostructures a versatile tool for engineering materials and devices properties. "In the present work, we studied the possibility to dynamically control the properties of a thin functional film by modifying



the atomic structure of the substrate with light," said Andrea Caviglia, now at the Kavli Institute of Nanoscience at Delft University of Technology.

At cryogenic temperatures, neodymium nickelate (NdNiO₃) is an antiferromagnetic insulator, i.e. the spins of the valence electrons align in an alternating pattern with no resulting net magnetization. Above 200 K, this material becomes a metal and the antiferromagnetic spin ordering disappears concomitantly. When an epitaxial NdNiO₃ thin film is grown on a lanthanum aluminate (LaAlO₃) substrate, the slightly different lattice constants of the two materials induce static strain in the film leading to a reduction of the insulator–metal-transition temperature from 200 K of the bulk material to about 130 K.

Interestingly, the electrical <u>properties</u> of the NdNiO₃ film can be changed on ultrafast time scales by selectively exciting <u>lattice vibrations</u> in the LaAlO₃ substrate. This was shown in an earlier publication of the main authors in *Physical Review Letters*. "In that experiment, a midinfrared laser pulse at 15 μ m wavelength triggered a vibrational mode in the substrate and a drastic change of electrical conductivity in the nickelate film was observed by measuring the change of the sample reflectivity with a terahertz, that is far-infrared, pulse," said Caviglia.

In the current work, the group studied the effect this substrate excitation has on the <u>magnetic properties</u> of the nickelate film. To measure these changes with high spatial and temporal resolution, the team used a technique called time-resolved resonant soft X-ray diffraction at the Linac Coherent Light Source (LCLS) <u>free-electron laser</u> at SLAC, California. The LCLS femtosecond X-ray pulses scatter off the film, carrying time-stamped signatures of the material's spin ordering that the physicists then used to reconstruct the spatiotemporal magnetic dynamics.



First, the researchers found that the magnetic ordering melts on a timescale of few picoseconds, i.e. at similar time scales as the insulator-metal transition observed earlier, thus suggesting that both processes are connected.

"Even more interestingly, the diffraction experiment showed that the magnetic melting in the nickelate starts locally at the interface to the substrate and propagates, comparable to a wave, from there into the NdNiO₃ film," said Först. "The high speed at which this wave front propagates, indicates that these dynamics are driven by local changes of the electronic structure at the interface", he added.

Indeed, a theoretical model, assuming the creation of freely movable charge carriers at the hetero-interface by the <u>substrate</u> lattice vibrations, supports this picture. These charges are likely to scramble the antiferromagnetic order as they propagate into the film.

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