

Mizoguchi Research Group

[Understanding the Role of Atom and Electron]

Department of Materials and Environmental Science

Nano-Materials Design Laboratory

Graduate School of Engineering

Department of Materials Engineering

<http://www.edge.iis.u-tokyo.ac.jp>

1

Materials Design

~Paving the Way for Materials Design~

What's kind of Structures?

How to bring about the Properties

Property \longleftrightarrow Structure
Relationship



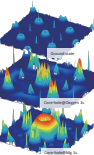
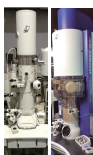
Property-Structure Relationship for Material Design

Much higher performance and higher reliability are now required to the materials to achieve further technology developments. In case of electroceramics, such as multi-layer ceramic capacitor and varistor, the size of grains in devices becomes smaller and smaller, and further property improvements of each grain and grain boundary are desired. To achieve this, clarification of atomic and electronic structures and finding the way to improve their properties are indispensable.

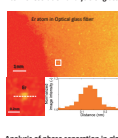
In our group, atomic and electronic structure are investigated by combining electron energy loss spectroscopy (EELS), transmission electron microscopy (TEM), first principles calculation. By combining these methods, atomic and electronic structures and their relationships with materials properties can be unraveled. Particularly, superlattice, ionic liquid, Li-ion battery, photovoltaic cell, electroceramics, and glass are investigated.

2

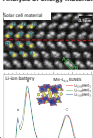
Seeing Atoms & Bonding



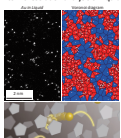
Atomic resolution analysis of glass



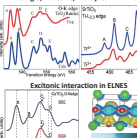
Analysis of energy materials



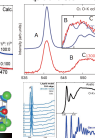
Atomic resolution analysis of liquid



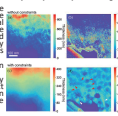
Single, Exotic, Multiplet in EELNES



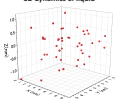
Liquid and Gas EELNES



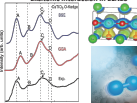
Analysis of phase separation in glass



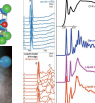
3D dynamics of liquid



Exotic Interaction in EELNES



Liquid and Gas EELNES



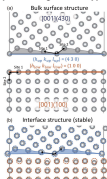
We are investigating electron-energy-loss near-edge structures (EELNES) and X-ray absorption near-edge structures (XANES) from experimental and theoretical viewpoints. EELNES and XANES originate from the electron transition from a core orbital to unoccupied bands. Their spectral features reflect the partial density of states of unoccupied bands, which contains information on the atomic and electronic structures. Experimentally, spatial- and time-resolved EELNES can be obtained at high accuracy by STEM-EELS. On the other hand, theoretical calculation is necessary to interpret the experimental spectrum, including effect of core-hole, which is introduced by electron transition from core-orbital to conduction band, is indispensable to calculate spectrum structures correctly.

K. Nakazawa et al., Ultramicroscopy, 237 (2020) 118077-5; K. Ueno et al., ACS Applied Nano Mater. (2020); Y. Sugimoto et al., ACS Advanc. Sci. (2020); S. Nakazawa et al., Scripta Mater. 184 (2018) 107; T. Miyata et al., Science Adv. 3 (2017) e370154; T. Miyata et al., Ultram. 178 (2017) 81; T. Miyata et al., Ultramicroscopy 171 (2016) 177; K. Nakazawa et al., Sci. Rep. 7 (2017) 14544; S. Nakazawa et al., Ultram. 178 (2017) 101-111; S. Terada et al., J. Phys. Chem. C 120 (2016) 10204-10212; M. Oishi et al., Chem. Phys. Lett. 649 (2016) 102; Y. Yamada, Sci. Rep. 9 (2019) 13003-7; K. Hatakeyama et al., Appl. Phys. Lett. 100 (2012) 023106; T. Mizoguchi et al., ACS Nano 7 (2013) 3008; S. Otsuka et al., Appl. Phys. Lett. 99 (2011) 232102.

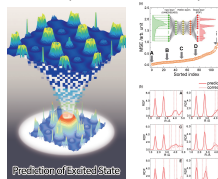
3

Understanding the Role of Atom and Electron in Materials

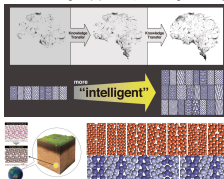
Direct prediction of gain boundary properties



Data-driven prediction of structure from core-loss spectra



Artificial Intelligence (AI) achieve 3,600 times higher efficiency



To design materials properties, we are investigating atomic and electronic structures quantitatively by performing high-precision simulations of atomic structures such as interfaces and lattice defects, which have a great impact on the functions of materials, and core excitation spectra, which show a variety of shapes reflecting electronic states. In addition, from the viewpoint of materials informatics, where information science is applied in materials research, we are applying machine learning methods such as transfer learning, Bayesian optimization, and virtual screening to interfaces, lattice defects, and inner-shell excitation spectra to understand and predict structure-function relationships.

S. Okamoto et al., Materials, 13 (2020) 2020; S. Okamoto et al., High Perform. Mater. 3 (2020) 60; K. Choshi et al., Appl. Phys. Express 13 (2020) 010104; S. Ryohara et al., J. Phys. Mater. 2 (2021) 012001; M. Yoshida et al., J. Phys. Chem. Lett. 9 (2018) 3713; S. Ryohara et al., Sci. Rep. 8 (2018) 12436; S. Ryohara et al., J. Chem. Phys. 148 (2018) 241701; S. Ota et al., J. Phys. Soc. Jpn. 88 (2017) 120401; S. Ryohara et al., Physica B 454 (2016) 11; S. Ryohara et al., Physica B 454 (2016) 10; S. Ryohara et al., Sci. Adv. 2 (2016) e100776; S. Ryohara et al., Jpn. J. Appl. Phys. 55 (2016) 060201; A. S. Kawasumi and T. Mizoguchi, J. Appl. Phys. 119 (2016) 123102; Y. Yamamoto et al., Appl. Phys. Lett. 108 (2016) 231004; Y. Yamamoto et al., J. Ceram. Soc. Jpn. 103 (2016) 460; Y. Yamamoto et al., Appl. Phys. Lett. 108 (2016) 123102; Y. Yamamoto et al., Appl. Phys. Lett. 102 (2013) 023102; Y. Yamamoto et al., Phys. Rev. B 88 (2013) 094117; T. Mizoguchi et al., Adv. Funct. Mater. 23 (2013) 2756.

