



Mathematical modelling for the next-generation all-solid-state battery: Nucleation interface

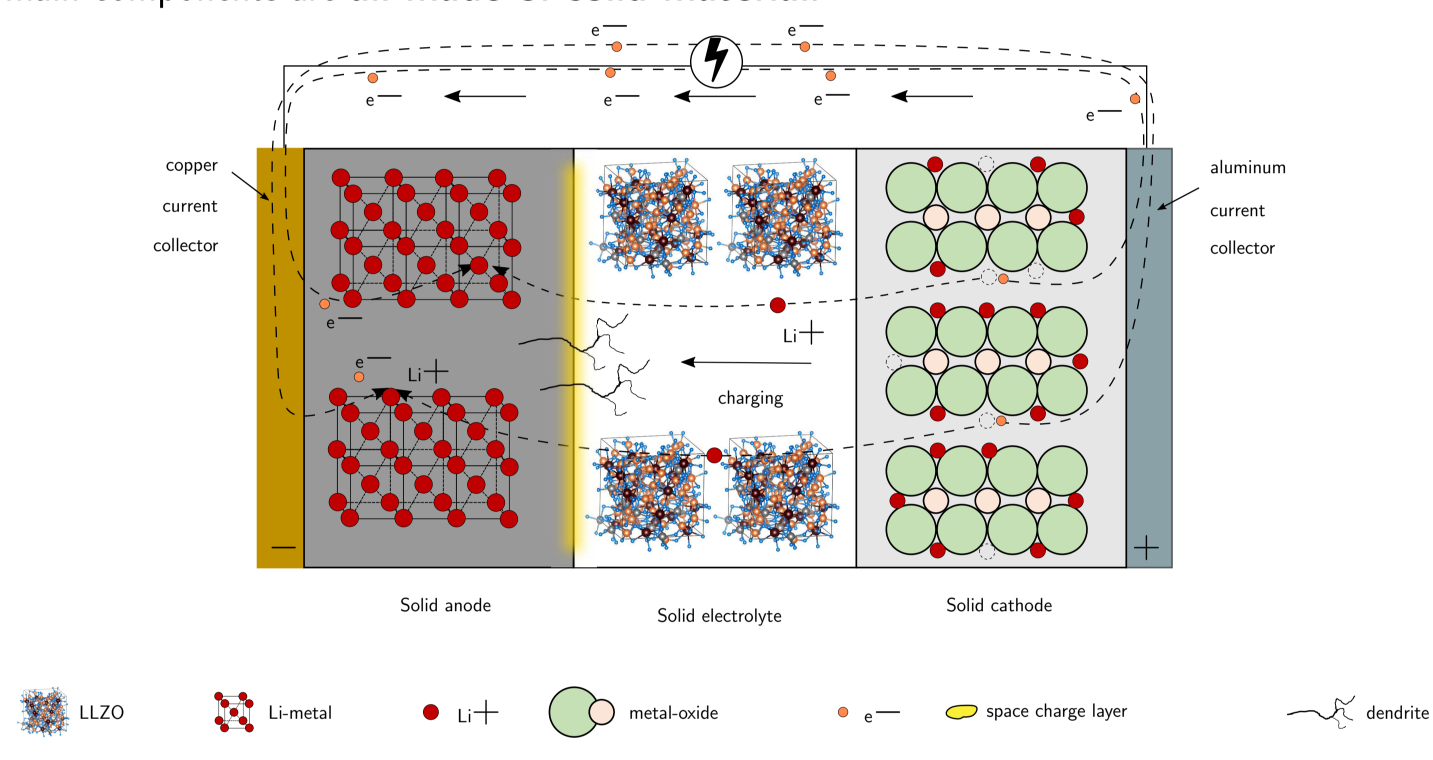
Tuan Vo

ACoM, Applied and Computational Mathematics, RWTH Aachen University

Next-generation All-solid-state battery

Rechargeable Lithium-ion battery (LIB) stays at the heart of every energy storage system and electric vehicle. Undoubtedly, LIB benefits human life efficiently as well as friendlyenvironment. Besides, a more advanced LIB, so-called all-solid-state battery (ASSB), is introduced recently as ASSB is expected with non-inflammation and non-explosion as seen in common LIBs. Yet, defect due to polarization is one natural phenomenon of solid elec**trolyte** (SE) to be tackled.

This poster is aimed to model the polarized SE with the use of **structural tensor**. A typical LIB includes three main components: cathode, anode and electrolyte. Different types of LIB have a variation of constitutive material composed of battery. An ASSB means that the three main components are all made of solid material.



Modelling goal: Interface analysis + Numerical modelling

Two main goals to model the solid electrolyte part of the all-solid-state battery is as follows:

- 1. To capture the **preferred direction** behaviour of the solid electrolyte due to electric potential.
- 2. To satisfy **thermodynamic consistency**:
 - ullet Conservation of mass, linear & angular momentum and energy for the solid electrolyte.
 - Entropy inequality is guaranteed with sharper conditions, which lead to constitutive equation.

$$a_{\mathsf{Griffith}} := a^* = \arg\min_{a \in \mathbb{R}} \left. \iint_{\Omega} f(a, \boldsymbol{u}; \lambda, \mu, \boldsymbol{d} \otimes \boldsymbol{d}) \, d\Omega - \left. \iint_{\Gamma} f(a; \gamma) \, d\Gamma \right|_{\boldsymbol{u}^{(s)}}$$

Contact

Tuan Vo · ACoM · RWTH Aachen University · Email: vo@acom.rwth-aachen.de



Mathematical model

