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Gertsen, Anders S.; Sørensen, Michael Korning; Fernández Castro, Marcial; Nelson, Jenny; Andreasen, Jens W.

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DTU Energy Department of Energy Conversion and Storage



The "Go Big or Go Home" Club: Molecular Dynamics Modelling of Bulk Heterojunctions

Anders S. Gertsen,^{†,‡} Michael Korning Sørensen,[†] Marcial Fernández Castro,[†]

Jenny Nelson,[‡] Jens W. Andreasen[†]

[†] Technical University of Denmark, Department of Energy Conversion and Storage, 4000 Roskilde, Denmark
 [‡] Imperial College London, Department of Physics, London SW7 2AZ, United Kingdom

Motivation

The "Go Big or Go Home" club is a collaboration within the SEEWHI H2020 project which aims upscale the manufacturing of organic photovoltaics (OPVs) without compromising the device efficiency. Usually, the efficiency is halved when moving from non-scalable spin-coating to roll-to-roll (R2R) deposition techniques, but combining the strengths of molecular dynamics (MD) modelling and *in situ* X-ray scattering enables the analysis of blend microstructure formation during post-deposition drying, in turn allowing us to identify the processing parameters that are key to overcome the scalability lag and move towards cheap, large-scale, and non-toxic solar cells with record efficiencies.

Simulating post-deposition drying

The efficiency of solution processed OPVs is crucially dependent on the 3D mesoscale thinfilm morphology, which in turn is greatly influenced by solvent properties and evaporation rate.

In order to reliably simulate active layer morphologies from MD simulations, we have scripted a range of evaporation schemes incorporating e.g. potential walls mimicking substrate– and air interfaces as well as including a suspended solvent vapour above the film from which solvent molecules are continuously removed.¹

3. R2R coated devices 2. In situ X-ray scattering

Approaching experimental time-scales

Using the MARTINI force field² to coarsegrain our systems, the time- and lengthscales relevant for morphology evolution are within reach of MD simulations:^{3,4}

~1000 times speed-up



I. Molecular dynamics modelling

Figure 2: All-atom and MARTINI coarse-grained models for chloroform (CF), PCBM,⁴ and a dimer of C₁₆-IDTBT.



Simulation time

Figure 1: Solvent evaporation simulation of all-atom O-IDTBR small-molecule acceptors in coarse-grained chloroform.

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Methodology

We have used the GROMACS 2016.3 package⁵ for all MD simulations. The OPLS-AA force field⁶ formed the basis for the all atom simulations of IDTBT and O-IDTBR with our own parameterizations of e.g. most angles and in particular the IDT-BT torsional potential and sidechain couplings – these were based on quantum chemical DFT calculations. The coarse-grained simulations were based on the MARTINI force field² and in part Ref. 4.

Figure 3: Thin-film of C_{16} -IDTBT 12mers (red) and PCBM (semi-transparent grey) annealed at 600 K and cooled down to 300 K. Sidechains are not shown.

References

[1] V. Negi *et al.*, Macromol. Theory Simul. 2016, 25, 550-558.
[2] S. J. Marrink *et al.*, J. Phys. Chem. B 2007, 111, 7812–7824.
[3] C.-K. Lee and C.-W. Pao, J. Phys. Chem. C 2014, 118, 11224–11233.
[4] R. Alessandri *et al.*, J. Am. Chem. Soc. 2017, 139, 3697–3705.
[5] M. J. Abraham *et al.*, SoftwareX 2015, 1–2, 19–25.
[6] W. L. Jorgensen Group, Yale University, Department of Chemistry.

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