Multi-scale Modelling of Conjugated Polymers to Understand the Role of Side Chain Chemistry in Mixed Ionic-Electronic Conduction

Abstract

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Organic electrochemical transistors (OECTs) have been gathering increasing interest due to their high reported transconductance values,1, 2 high switching speeds3 and low bias operation,4 allowing for operation in aqueous electrolyte. Furthermore they have been shown to operate in a range of biological scenarios5–7 as well as being cheap to

manufacture.8–11 One of the challenges in designing accumulation mode OECTs is understanding the interplay between the ion-penetration into the channel and device performance (mixed ionic-electronic conduction).12 Recently it has been shown that increasing the hydrophilicity of the active material through inclusion of ethelyne-glcyol units

on the side chains can significantly improve device performance due to increased volumetric swelling of the channel.4, 13, 14 Furthermore it has been shown that crystallinity and crystal structure is important for understanding how ion penetration affects hole mobility.15 We model two polymers based on poly-thiophene backbones with either alkoxy- or glycoxy- sidechains (p(aT2) and p(gT2) respectively). We verify molecular dynamics forcefields against monomer crystal structures, allowing us to accurately simulate the respective polymer crystals. Furthermore we extend our simulations to include the aqueous electrolyte, as polymer-ion interactions are critical to understanding the operation of OECTs and how materials could be further tuned through changing the side chain chemistry.

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