

Recent advances in understanding carbon nanotube growth through atomic scale simulations

Erik C Neyts

University of Antwerp

Contact e-mail: *erik.neyts@uantwerpen.be*

Atomistic simulations have contributed significantly to our understanding of the metal-catalyzed CNT growth process. Such simulations revealed detailed insights in e.g. carbon dissolution and surface segregation [1, 2], carbon polyynes formation at the surface [3], and cap lift off [4, 5]. First, I shall highlight a selection of simulations which have provided such insights and demonstrate how they have contributed to our understanding of the growth process [6]. Subsequently, a number of recent efforts will be highlighted which address previously uncovered areas, including ion bombardment [7, 8], growth from hydrogen containing growth precursors [9], and the application of accelerated atomic scale techniques for studying the dynamics of the growth process [10].

- [1] H. Amara, PRL 100 (2008) 056105
- [2] F. Ding, CPL 393 (2004) 309
- [3] A. J. Page, Acc. Chem. Res. 43 (2010) 1375
- [4] M. A. Ribas, JCP 131 (2009) 224501
- [5] Y. Shibuta, CPL 382 (2003) 381
- [6] J. A. Elliott, Nanoscale 5 (2013) 6662
- [7] E. C. Neyts, PRL 110 (2013) 065501
- [8] M. Shariat, CPL 590 (2013) 131
- [9] U. Khalilov, Nanoscale 6 (2014) 9206
- [10] K. M. Bal, JCP 141 (2014) 204104