Energy & Environmental Science



CORRECTION

View Article Online



Cite this: Energy Environ. Sci.. 2022, 15, 4000

Correction: Peripheral halogenation engineering controls molecular stacking to enable highly efficient organic solar cells

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DOI: 10.1039/d2ee90045a

rsc.li/ees

Correction for 'Peripheral halogenation engineering controls molecular stacking to enable highly efficient organic solar cells' by Yalu Zou et al., Energy Environ. Sci., 2022, https://doi.org/10.1039/ d2ee01340a

The broader context section for this article was missing. It should have appeared as follows.

Modifying non-fullerene acceptors (NFAs) with peripheral halogenation has been considered as a relatively simple but an effective strategy to boost power conversion efficiencies (PCEs). However, the lack of systematical investigation, especially in the state-of-the-art Y6 series NFAs, makes the bridge between peripheral halogenation in both central units and end groups and molecular stacking, active layer morphology, charge transfer/transport dynamics and device performances of the resulting OSCs a crucial but still unaddressed issue. Herein, based on our previous experience in molecular engineering, a novel series of nonfullerene acceptors (NFAs), CH-6F, CH-4Cl and CH-6Cl, are designed and synthesized, featuring multiple peripheral halogenations in both conjugate extended central units and end groups. With CH-series NFAs, a comprehensive study has been conducted to systematically probe the significant effects of peripheral halogenation on their single crystal packing, disclosing that peripheral halogenation induced completely different crystal systems and quite unique molecular packing modes. Moreover, this superior molecular packing optimizes film morphology, improves photovoltaic performances, and reduces energy losses of the resulting OSCs. Finally, by utilizing CH-series NFAs, a series of highly-efficient OSCs have been afforded with a champion PCE of 18.22% and markedly reduced $\Delta V_{\rm nr}$ of 0.203 V in CH-4Cl-based ternary devices. Our results indicate that controlling molecular stacking modes by peripheral halogenation engineering should be a possible avenue toward OSCs with higher efficiency.

The Royal Society of Chemistry apologises for these errors and any consequent inconvenience to authors and readers.

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