Single crystal X-ray data for **A**, **B**, **C** were measured on a Bruker AXS D8 Venture diffractometer (multilayer optics, Mo-Kα and Cu-Kα radiation with λ = 0.71073 Å and 1.54178 Å respectively, Kappa 4-circle goniometer, Photon III C14 CPAD detector).

Single crystal X-ray data for **D**, **E**, **F** were measured on a Bruker AXS Apex II diffractometer (graphite monochromator, Mo-Kα radiation, λ = 0.71073 Å, Kappa 4-circle goniometer, Apex II CCD detector).

All crystals were measured at a temperature of 100 K. Absorption corrections using equivalent reflections were performed with the program SADABS.[1] For **A** a numerical absorption correction was performed using the same program. The crystal of **B** was a non-merohedral twin, and the absorption correction was performed with the program TWINABS.[2]

All structures were solved with the program SHELXS[3] and refined with SHELXL[4] using the OLEX2[5] GUI.

All non-H atoms were refined using anisotropic atomic displacement parameters (ADPs). H atoms bonded to C were located in the difference Fourier maps and placed on idealized geometric positions with idealized ADPs using the riding model.

The crystallographic data can be obtained free of charge from https://www.ccdc.cam.ac.uk/structures/ quoting the CCDC numbers 3000001-3000006.

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