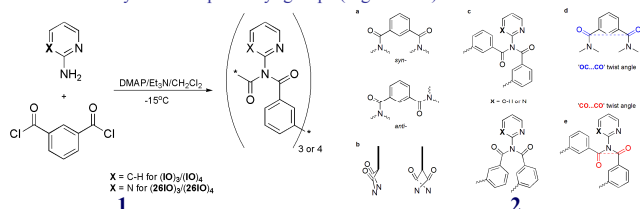


Introduction

Reaction of isophthaloyl dichloride (**1**) with 2-aminopyridine or 2-aminopyrimidine (**2**, **26O**) provides a facile entry into a new class of imide-based '3+3' macrocyclic trimer (**10**)₃, (**26IO**)₃ (as **trezimides**), together with the known tetramer (**10**)₄, (**26IO**)₄ (**tennimide**) scaffold.¹⁻⁵ **Trezimides** can adopt two asymmetric conformations, isolated as (**P**) in (**10**)₃ and (**R**) in (**26IO**)₃.^{2,4} The **tennimide** (**26IO**)₄ structure exhibits three discrete conformations as *cc/cc/oo*, highlighting subtle geometric changes with the **tennimide** channel (pore) open (*o*) and/or closed (*c*). Macrocyclic formation (competing with oligomer/polymer formation) relies on the *ortho*-pyr(im)idine *N* functionality and imide *hinge* ('CO...CO' twist) with the inherent flexibility of the isophthaloyl groups (Figures 1-5).



Scheme 1 Macrocyclic synthesis: the reaction route to (**10**)_{3,4} and (**26IO**)_{3,4}. **Scheme 2** (a) *syn*- and *anti*-conformations of the isophthaloyl groups, (b) side-views of the isophthaloyl conformations, (c) imide conformations. (d) OC...CO isophthaloyl and (e) CO...CO imide hinge torsion angle (°).

Experimental methods

The reaction of isophthaloyl dichloride with 2-aminopyridine (or 2-aminopyrimidine) at -15°C in dry dichloromethane (CH₂Cl₂) with 3 equivalents of triethylamine (Et₃N) and a catalytic quantity of dimethylaminopyridine (DMAP) afforded a mixture of products (and oligomeric/polymeric material) (Scheme 1).²⁻⁴ The single crystal X-ray data (Mo/Cu) were collected on an Oxford Diffraction Gemini S-Ultra diffractometer at 294(1) K: θ range typically from 2-26° (100% data coverage to 25°).

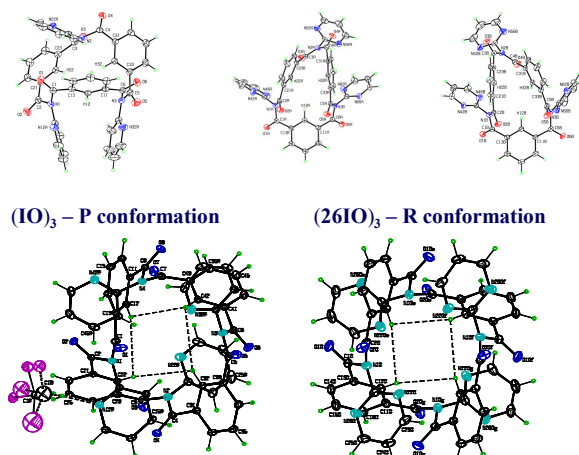


Figure 1: Views of the (**10**)₃ and (**26IO**)₃ **trezimides** (top left/right) and the (**10**)₄ and (**26IO**)₄ **tennimides** (bottom left/right) with displacement ellipsoids drawn at the 20% probability level.

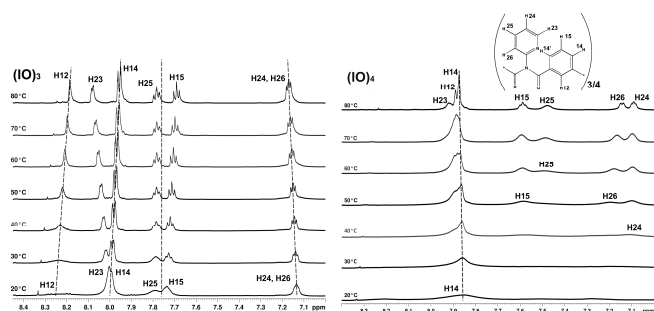


Figure 2: Variable temperature ¹H NMR studies (from 20°C to 80°C, 600 MHz, d₆-DMSO) for (**10**)₃ and (**10**)₄. Dashed lines indicate the displacement of resonances.^{2,4}

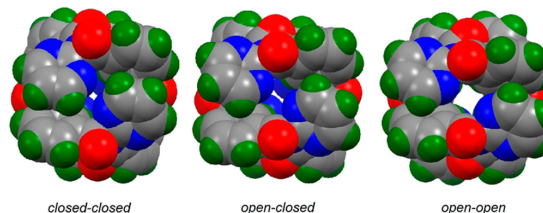
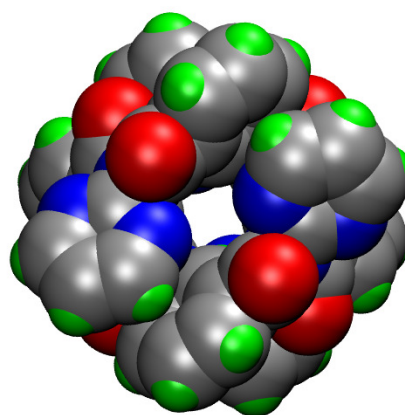


Figure 3. CPK views of the three discrete *cc* ↔ *oc* ↔ *oo* conformational states of (**26IO**)₄.²



Scan to view animations

Figure 4. CPK view of the (**26IO**)₄ molecule (molecule 'O' in the *oo* conformation).²

Halogen bonding directed supramolecular assembly⁵

Brominated macrocycles aggregate via C–Br...O=C_{carbonyl} and/or C–Br...N_{aromatic} halogen bonds (with *N_c* ≤ 0.90) and often augmented by longer C–Br...H/π(arene) contacts. The brominated **trezimides** and **tennimides** (each with 3 or 4 Br atoms) lack classical strong hydrogen bonding donors (*i.e.* N–H, O–H), but contain many acceptors (*i.e.* N_{aromatic}, C=O, arene) participating in a myriad of halogen and weaker hydrogen bonding interactions.⁵ The C–Br groups promote overall macrocyclic aggregation linking macrocycles into 1-D halogen bonded chains. (**BrIO**)₃ exhibits three different types of Br...O=C/π(arene) halogen and Br...H hydrogen bonds, with C–H...O contacts whereas (**26BrIO**)₃ contains Br...O=C halogen bonds (Figure 5).⁵

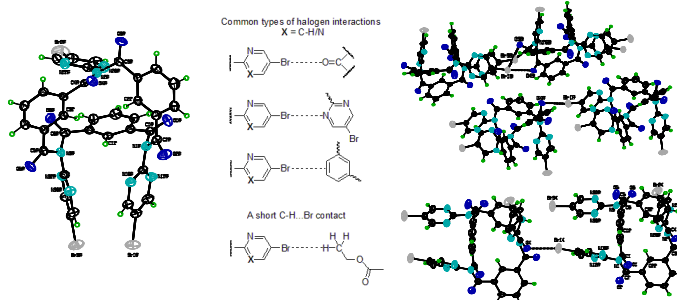


Figure 5: ORTEP diagram of (**26BrIO**)₃ and types of halogen/hydrogen bond interactions in (**26BrIO**)₃.

Results and Conclusions

The one-step formation of **trezimides** and **tennimides** is achieved in modest yields from readily available starting materials [*i.e.* isophthaloyl dichloride and (un)substituted 2-aminopyr(im)idines]. Macrocyclic formation depends on the asymmetric imide *hinge* (*i.e.* the O=C...C=O twist angle – Scheme 2e) and inherent flexibility of the isophthaloyl groups to twist and bend on macrocyclisation. Two distinct **trezimide** conformations (**P**) and (**R**) have been structurally characterised. The macrocyclic scaffold and core has been isolated in three different conformational states in the (**26IO**)₄ **tennimide**. The internal macrocyclic cavity is large to accommodate a small atom/ion. Future work is being directed towards the synthesis of larger macrocycles, coordination chemistry and halogen bonding applications.⁵

References

- Louise S. Evans and Philip Gale, *Chem. Commun.*, 2004, 11, 1286-1287.
- Pavle Mocilac and John F. Gallagher, *J. Org. Chem.* 2013, 78, 2355-2361.
- Pavle Mocilac and John F. Gallagher, *Acta Crystallographica Section B.*, 2013, 69, 62-69.
- Pavle Mocilac, *Structural systematics of drug-like molecules: from benzamides to tennimides*, PhD thesis, Dublin City University, 2012.
- Pavle Mocilac and John F. Gallagher, *CrystEngComm*, 2014, DOI:10.1039/C3CE42168F