

3.7. CRYSTALLOGRAPHIC DATABASES

References

Saulius Gražulis, Justas Butkus and Andrius Merkys. The deposition software performs rigorous checks of syntax and semantics.

The COD website allows searching on COD numerical identifier, unit-cell parameters, chemical composition and bibliographic data. Substructure searches using SMILES and SMARTS strings have been implemented. The free software package *OpenBabel* (O'Boyle *et al.*, 2011; Hutchison, 2007) is used for both the CIF-to-SMILES transformation and the actual search.

The retrieved records can be viewed online or downloaded for further processing. For massive data mining, COD permits downloads and updates of the whole database using Subversion, Rsync or http protocols. The ease of access to the COD data and its open nature has spurred the use of this resource for software testing (Grosse-Kunstleve & Gildea, 2011), teaching (Moeck, 2004) and research (First & Floudas, 2013). Multiple mirrors around the globe (Quirós-Olozabal, 2006; Gražulis, 2007; Moeck, 2007a; Chateigner, 2010) ensure data preservation, provide off-site backups, offer improved search interfaces (Moeck, 2007b) and increase reliability.

For the powder-diffraction community, the COD is interesting not only as an archive of structures solved by powder-diffraction methods, but also as a possibility for use in search/match procedures to identify crystalline compounds. Recently, the development of an open full-pattern search/match internet tool was launched by the COD developers. It allows phase quantifications from X-ray, neutron and electron powder patterns (with high- or medium-resolution instruments) provided that the structures are already in the COD. This tool is particularly suited to nanocrystalline powders, in which severe line broadening appears, precluding phase identification from only peak positions (Lutterotti *et al.*, 2012). COD-derived databases are also offered for software produced by several diffractometer vendors (Rigaku, 2011; PANalytical, 2012a,b; Bruker, 2013). In addition to the COD, searches and matches can be performed against its sister database, the PCOD, which contains structures predicted by the *GRINSP* program (Le Bail, 2005) and hypothetical zeolites (Pophale *et al.*, 2013). The power of such an approach is demonstrated by PCOD entry 3102887 (formulated as SiO₂). It was recently identified as corresponding structurally to a new phosphorus(V) oxonitride polymorph δ -PON (Baumann *et al.*, 2012).

3.7.9. Other internet databases

Other useful databases include the following:

- (i) The American Mineralogist Crystal Structure Database (<http://ruff.geo.arizona.edu/AMScsd.php>).
- (ii) The Mineralogy Database (<http://webmineral.com>).
- (iii) MinCryst (<http://database.iem.ac.ru/mincryst/index.php>).
- (iv) The International Zeolite Association Database of Zeolite Structures (<http://www.iza-structure.org/databases>).
- (v) The Incommensurate Structures Database (<http://webbdcrista1.ehu.es/incstrdb/>).
- (vi) The Nucleic Acid Database (<http://ndbserver.rutgers.edu>).

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