



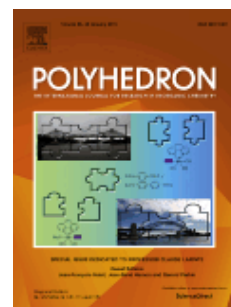
POLYHEDRON

The International Journal for Research in Inorganic Chemistry.

AUTHOR INFORMATION PACK

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DESCRIPTION

Polyhedron publishes original, fundamental, experimental and theoretical work of the highest quality in all the major areas of **inorganic chemistry**. This includes **synthetic** chemistry, **coordination** chemistry, **organometallic** chemistry, **bioinorganic** chemistry, and **solid-state** and **materials** chemistry. Papers should be significant pieces of work, and all new compounds must be appropriately characterized. The inclusion of single-crystal X-ray structural data is strongly encouraged, but papers reporting only the X-ray structure determination of a single compound will usually not be considered. Papers on solid-state or materials chemistry will be expected to have a significant molecular chemistry component, such as the synthesis and characterization of the molecular precursors and/or a systematic study of the use of different precursors or reaction conditions. Polyhedron publishes **full papers**, specially commissioned review articles (**Polyhedron Reports**) and themed issues of the journal (**Polyhedron Symposia-in-Print**). Polyhedron does **not** publish communications or notes.

AUDIENCE

Inorganic chemists, biochemists.

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(iii) Unit cell parameters with esds and X-ray wavelength used.

(iv) Crystal system, space group and number of complete chemical units (see ii) per cell.

(v) Type of diffractometer used and method of data collection. Total number of data collected, excluding any intensity controls, number unique Rint value, number observed (with cut-off parameter), and completeness of data to suitable 2θ or \sin/λ limit, use or otherwise of absorption correction.

(vi) Final results. Give values of R, R_w (and their definitions) and number of parameters. Indicate form of refinement (F or F_2) and treatment of hydrogens.

Note. Easily derived parameters should not be given unless required in discussion of other data (e.g. number of solvent molecules, etc).

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