CONFORMA: A PROGRAM FOR DETERMINING RING CONFORMATIONS AND PUCKERING COORDINATES

INTRODUCTION

Many research areas require precise determination of the conformations assumed by different atom rings, as for example the study of interactions protein-substrate or crystal structure determinations of chemicals. Therefore, a methodology for quantifying the different conformations, allowing an easy and fast comparison between them, is necessary. This kind of problem becomes more complex when one needs to identify or compare hybrid conformations. Several methodologies have been proposed, all aiming at systematizing and quantifying the ring conformations. Among them, Cremer & Pople’s, which depends on the number and type of atoms, is the most widely used, this methodology defines a set of ring puckering coordinates, their number depending on the number of atoms in the ring, which describe the nonplanar character of a ring in a unique way. CONFORMA is a program developed for computers with the MS-DOS operational system, it calculates the puckering coordinates, for five or six member rings as defined by Cremer & Pople, with their standard deviations according to Norrænæ, and the conformation the rings assume. All calculations are performed using atomic coordinates from crystal structure determination or from any other cartesian coordinate system.

PROGRAM DESCRIPTION

For five member rings, CONFORMA calculates the q3 and \( \phi_1 \) coordinates, which are related to the departure from planarity and to the position in the pseudo-rotation pathway, respectively. For six member rings, three coordinates, \( q_2 \), \( q_3 \), \( \phi_2 \) are calculated, and, as described by Cremer \& Pople, also the corresponding spherical polar coordinates \( (Q,\theta,\phi) \), which locate the various possible ring conformations on the surface of a sphere. Thus, a perfect chair has \( \theta \) equal to 0 or 180°; \( \theta \) equal to 90° corresponds to a boat or twisted boat conformation, with \( \phi \) defining the position of the pseudo-rotation pathway. In the case of hybrid conformations, this will be reflected in the values of \( \theta \) and \( \phi \). Q is related to the departure from planarity and is therefore called total puckering amplitude.

Finally, the program performs an analysis and interprets all these puckering coordinates.

EXAMPLES

Two crystal structures will be shown as examples. In figure 17 ring A is in a slightly distorted chair conformation, and in compound of figure 3 the four rings are examined, special attention should be given to ring B which is a highly strained six member ring, whose conformation is not easy to evaluate visually, but puckering coordinates make this evaluation straightforward.

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Figure 1. Atom numbering and ring letter designation scheme of 7-isopropyl-4-methyl-tricyclo[8.4.0.0{2,6}]4,9-tetraacdecadione-3,5.
CONCLUSION

In future versions of the program we intend to implement a graphical facility that will allow the visualization of the departure of hybrid conformed rings from its corresponding pure conformations.

Free copies of the program may be requested from the authors, by letter or e-mail. The program will be sent in a 3.5'' diskette for IBM-PC and compatibles, which contains the following files: readme.1st, with starting information; userregl.doc, the user registration form, which the authors kindly ask users to fill and return; id.coo, which identifies the coordinate files/examples, included in the package; [name].dat, example files with the appropriate format for CONFORMA; CONFORMA.EXE, the executable version of CONFORMA program.

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REFERENCES AND NOTES

5. A program, RING (Cremer, D. QCPE, 1975, 11, 288) written in FORTRAN IV (CDC), only calculates de puckering coordinates.