LETTERS TO THE EDITOR

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Non-conventional unit cells

BY RICHARD E. MARSH

Division of Chemistry and Chemical Engineering, California Institute of Technology, Pasadena, California 91125, USA

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As a long-time reader of Acta Crystallographica, I have become used to many of the nonconventional unit cells that some authors foist on me. I have learned to tolerate space groups $P2_1/a$ and A2/a, even though a simple interchange of axes would turn them into the standard $P2_1/c$ and C2/c; I have figured out $I2_1/a$, even though the International Tables do not recognize a bodycentered monoclinic unit cell, and $C\overline{1}$, despite knowing that a triclinic cell must be primitive; I have deduced that *Pnca* is space group No. 60, customarily referred to as Pbcn, and that Pbnm is really *Pnma*. With somewhat less good will I have put up with unit cells which, for no good reason, have angles less than 60° or greater than 120°, resigning myself to carrying out a cell reduction if I wish to compare with another compound or with another investigation of the the same compound. But I feel that I must stand up and complain when I encounter a triclinic structure described in such a way that not one of the three shortest lattice vectors is used as a cell edge.

I refer to an article in the November 1987 issue of Acta Crystallographica Section C - '1,6dioxacyclodeca-3,8-diene', on page 2245. This structure is described as triclinic (space group $P\bar{1}$), with a = 7.263(15), b = 7.683(15), c = 7.225(12) Å, a = 72.62(5), $\beta = 137.37(8)$, $\gamma = 130.36(7)^{\circ}$. (Just look at those last two angles!) It turns out that the three shortest lattice vectors are [110], [101] and [$\overline{1}\overline{1}\overline{1}$]; they define a cell with a' = 6.285, b' = 5.266, c' = 6.903 Å, a' = 103.00, $\beta' = 113.76$, $\gamma' = 99.41^{\circ}$. Couldn't at least one of these axes have been used by the authors?

Is there a purpose in using such an unusual cell? I see none. But neither, in most instances, do I see a purpose in $P2_1/a$, or $C\overline{1}$, or Pbnm. All I see is a *reason*; and the reason I see is laziness. The usual procedure, I believe, is to permit a computer to pick out the unit cell and let it go at that. Rather than going through, once and for all, the very simple (in most software packages that I am aware of) process of interchanging axes so as to produce a convenient unit cell, many investigators leave it to the reader to work things out for himself.

But there may be another reason. Some contributors to Acta Crystallographica may believe that the small amount of time they save by not converting to a convenient unit cell is more valuable than the combined time of all readers who might be enough interested in the results so as to carry out the conversions for themselves.