In (1) the rocking angles of the C atoms of the four-membered ring are smaller than unity due to fewer non-bonded repulsions between *trans*-vicinal substituents compared to the repulsion between *cis*-vicinal substituents [$4\cdot3$ (4)° (Allen, 1984)].

As a result of the transannular distance 2.155 (2) Å in the cyclobutane, the seven-membered rings are strained. Both angles at the atoms C(3) and C(4) and the equivalent atoms exceed the angle at sp^2 -C atoms of 120° significantly $[124 \cdot 1 (4) \text{ and } 125 \cdot 7 (2)^{\circ}].$ Therefore the biphenyl groups deviate from a linear arrangement as they do in the tetrabenzocyclododecene derivative (2) and its cis-cis- and cistrans-isomers (Irngartinger, 1972, 1973a,b): the exocyclic bond angles at C(4), C(5), C(10) and C(11) [118.3 (3)°] are smaller than 120°. An angle of 169.9(1) and $169.7(1)^{\circ}$ is given by the axes defined by the atoms C(4), C(27) and C(5), C(22) respectively and the corresponding axes in the second biphenvl group. The 1,4-distances $C(16)\cdots C(17)$ 2.841 (2) and $C(24)\cdots C(25)$ 2.857 (2) Å are much shorter than the corresponding distances $C(3)\cdots C(6)$ 3.167(2) and $C(9)\cdots C(12)$ 3.168 (2) Å [van der Waals distance 3.4 Å (Pauling, 1976)].

In both biphenyl groups there are twists around the central bonds $[C(3)-C(4)-C(5)-C(6) -14.9(2), C(9)-C(10)-C(11)-C(12) 13.3(2)^{\circ}]$ with different sense of rotation, hence (1) is provided with a *meso*-conformation in the crystal and shows only small deviations from S_4 ($\overline{4}$) symmetry. These distortions bring about the different angles 112.0(5) and 116.5(6)^{\circ} between the bonds from the aromatic rings to the cyclobutane system and the two adjacent bonds of the four-membered ring. The bond lengths [1.415(2) Å; Fig. 1] between both substituted C atoms

in the phenyl rings are the longest ones within the six-membered rings.

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