

Achiral Molecules Move and Order Themselves in a Chiral Form in Host-Guest Inclusion Crystals

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(Received May 22, 1995)

When optically active host compound was mixed with achiral guest compound in the solid state, host-guest inclusion crystals were formed by molecular movement and the achiral molecules became arranged in a chiral form in the inclusion crystals. Mixing of powdered (+)-host crystals with powdered (-)-guest crystals which are formed by a chiral arrangement of achiral guest molecules gave inclusion crystals of the (+)-host and the (+)-guest molecules newly produced through an inversion of the (-)-guest molecules during the complexation in the solid state.

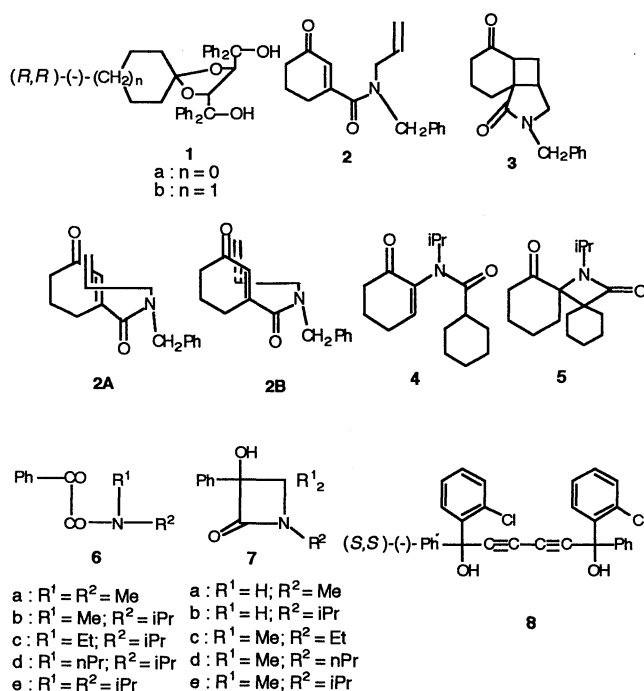
We have reported that irradiation of powdered 2:1 inclusion crystals of (*R,R*)-(-)-trans-2,3-bis(hydroxydiphenylmethyl)-1,4-dioxaspiro[4.4]nonane (**1a**)^{1,2} and *N*-allyl-*N*-benzyl-3-oxo-1-cyclohexenecarboxamide (**2**) which had been prepared by recrystallization of these from ether, in a water suspension which contains alkylsulfate as a surfactant at room temperature for 10 h gave (-)-**3** of 100% ee in 90% yield.³ Similar treatment of 2:1 inclusion crystals of (*R,R*)-(-)-trans-2,3-bis(hydroxydiphenylmethyl)-1,4-dioxaspiro[5.4]decane (**1b**)^{1,2} and **2** prepared by recrystallization from benzene gave (-)-**3** of 100% ee in 87% yield.³ Since the photoreaction of **2** itself in MeCN gives a mixture of *rac*-**3** and an isomeric photocycloaddition product,⁴ it is clear that the steric course of the photoreaction is controlled efficiently by the chiral host compound **1**. It is probable that molecules of **2** are arranged in a chiral form in the inclusion

complex with **1**. The chirality of **2** in the inclusion complex would result, for example, by locating the allyl group above the cyclohexenone ring (**2A**). When the (*S,S*)-(+)-enantiomer of the host **1** is used, the allyl group would be located below (**2B**) and (+)-**3** is produced by the irradiation.

Interestingly, however, we found that the same inclusion complex of **1** and **2** as that prepared by recrystallization can be obtained just by mixing **1** and **2** in the solid state and that photoirradiation of the inclusion crystals obtained by mixing gives optically active photoreaction product. For example, an occasional mixing of powdered **1a** of about 50-100 μm diameter and a half molar amount of oily **2** using an agate mortar and pestle for 1 h in order to avoid solidification gave the IR spectroscopically identical 2:1 inclusion complex crystal of **1a** and **2** with that obtained by the recrystallization method. Photoirradiation of the complex in a water suspension for 10 h gave (-)-**3** of 99% ee⁵ in 48% yield. Similar mixing of **1b** and **2** in the absence of solvent for 1 h followed by irradiation for 10 h gave (-)-**3** of 87% ee in 40% yield. These data clearly show that inclusion crystals of **1** and **2** in which achiral molecules of **2** are arranged in a chiral form produced just by mixing both components. Such molecular movement in the solid state and arrangement in a chiral form are not special for **2** but are rather common. For example, mixing of powdered **1a** and an equimolar amount of powdered *N*-isopropyl-*N*-2-(1-oxocyclohex-2-enyl)cyclohexanecarboxamide (**4**) followed by irradiation gave the photocyclization product (-)-**5** of 81% ee in 37% yield. Photoirradiation of **4** in acetone gives *rac*-**5** together with an isomeric photocycloaddition product.⁶

N,N-Dialkylphenylglyoxylamide **6** also forms an inclusion complex on mixing with **1** and the achiral molecules of **6** are arranged in a chiral form in the inclusion crystal. Occasional mixing of oily **6a** with an equimolar amount of powdered (*R,R*)-(-)-**1a** or the (*S,S*)-(+)-enantiomer of **1a** of about 50-100 μm diameter for 1 h followed by irradiation in a water suspension for 10 h gave (-)-**7a** of 78% ee (54%) or (+)-**7a** of 65% ee (50%), respectively in the yields indicated. Similar treatments of oily **6b** with **1a** or of oily **6d** with **1b** followed by photoirradiation gave (-)-**7b** of 87% ee (45%) or (-)-**7d** of 93% ee (39%), respectively in the yields indicated.

Very interestingly, irradiation of the 1:1 inclusion complex of **1b** and **6a** which had been prepared by mixing both components in the absence of solvent gave (+)-**7a** of 45% ee in 42% yield, although the same irradiation of the complex prepared by recrystallization of **1b** and **6a** from toluene gave the other enantiomer (-)-**7a** of 100% ee in 40% yield. The type of chiral



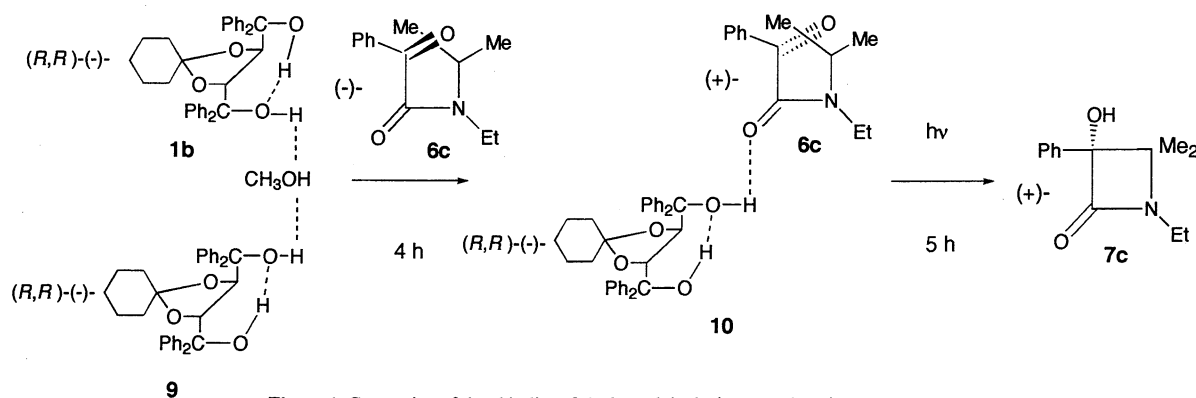


Figure 1. Conversion of the chirality of 6c through inclusion complexation.

arrangement of 6a molecules varies depending on the conditions of the inclusion complexation experiment.

Chiral arrangement of 6a molecules in an inclusion crystal with (S,S)-(-)-1,6-di(o-chlorophenyl)-1,6-diphenylhexa-2,4-diyne-1,6-diol (8)⁷ also depends on the conditions of the inclusion complexation experiment. Recrystallisation of 8 and 6a from ether gives a 1:1 inclusion complex, which upon irradiation affords (-)-7a of 100% ee in quantitative yield.⁸ X-ray structure study of the inclusion complex showed that 6a is arranged in a chiral form.⁸ However, irradiation of the inclusion crystal prepared by mixing 8 and 6a gave *rac*-7a. In the latter crystal, 6a molecules would not be arranged in a chiral form. Interestingly, however, both inclusion crystals are thermally interconvertible. After melting (mp 126-127 °C), the former is solidified by cooling to give the latter, but the latter is converted to the former by heating gradually and melts again at 126-127 °C.

It has been reported that 6e forms chiral crystals in which 6e molecules are arranged in a chiral form produced by a 90° twisting around the single bond between two CO groups,⁹ and that irradiation of the chiral crystals gives optically active 7e of 93% ee in 74% yield.¹⁰ This time, we found that 6c also forms chiral crystals which upon irradiation gives optically active 7c. Although 6e forms inclusion crystals neither with 1 nor 8, 6c forms an inclusion complex with 1 both by recrystallisation and solid state complexation procedures. Especially, 1:1 inclusion crystals of 1b with 6c are easily obtained by mixing a 2:1 MeOH complex (9) of 1b and 6c. The complexation in the solid state can be followed by its IR spectrum. As the complexation proceeds, hydrogen bonded νOH absorption between 1b and MeOH (3550 cm⁻¹) decreases and finally disappears, and a new hydrogen bonded νOH absorption between 1b and 6c (3250 cm⁻¹) appears and increases (Figure 1). The complexation was completed within 4 h. Irradiation of the resulting inclusion complex (10) of 1b with 6c gave (+)-7c of 76% ee.

Very interestingly, by mixing powdered 9 and powdered (-)-crystals of 6c, 1:1 inclusion crystals of 1b and (+)-6c were formed. Firstly, (-)-crystals of 6c were prepared in large quantity by adding one piece of a (-)-crystal of 6c as a seed crystal during recrystallization of 6c from ether. Photoirradiation of powdered

(-)-crystals of 6c in the solid state gave (-)-7c of 80% ee in 54% yield. Secondly, a 1:1 inclusion complex of 9 and (+)-6c (10) was obtained by mixing powdered 9 and (-)-crystals of 6c. The (+)-arrangement of 6c molecules in 10 can be proven by its photoirradiation in a water suspension for 5 h which gives (+)-7c of 76% ee in 72% yield. Interconversion between the molecular arrangement of 6c in the (-)- and (+)-forms easily occurs by the complexation with chiral host 1 in the solid state. In other words, molecules move easily in the solid state and change their chiral arrangement according to the chiral circumstances in the crystal.

This work was supported by the Grant-in-Aid for Scientific Research on Priority Areas No. 06242105 from the Ministry of Education, Science and Culture, Japanese Government.

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