NOVEL COMPLEXES OF AMIDES AND OTHER SPECIES WITH A HETEROCYCLIC BORON BETAINE.¹

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Abstract - The heterocyclic boron-containing betaine (J,) **forms** stable 1:l addition complexes with water, amines and, more surprisingly, simple amides. **INTRODUCTION**

We have previously reported that reaction of diphenylbromoborane with di-isopropylcarbamoyl-lithium, followed by treatment with water and chromatography on silica gives the novel heterocyclic betaine (1) in almost quantitative yield.²

Prior to chromatography the product **was** a crude yellow viscous oil vith broad i.r. bands at 3200 cm⁻¹ (weak), 1650 cm⁻¹ (strong) and sharp bands at 1620 cm⁻¹ (s) and 1580 cm⁻¹ (s). This contrasts markedly with (1), for which the characteristic bands are the sharp absorptions at 3550 cm-l (medium) and 1580 cm'l **(5).** By dissolving the crude product in benzene and careful crystallization large prismatic crystals **were** obtained. We **now** report the identification of this product **as** a 1:l complex of **(k)** with di-isapropylfornamide (DIPF). We also wish to report that complexation of (1) is fairly general for other simple amides as well as for amines and water. RESULTS AND DISCUSSION

A crude product **was** obtained by reacting diphenylbromoborane vith diisopropylcarbamoyllithium $(-78^{\circ}C \text{ rising to room temperature})$, treatment with water, extraction into ether and removal of the solvent. The heavy viscous oil **was** dissolved in benzene which **was** concentrated by distillation and then allowed to cool slowly to produce large prismatic crystals.

The i.r. spectrum (KBr disc) of these crystals revealed no sharp band at 3550 cm⁻¹ (characteristic of (l_c)), but showed a very broad, weak band in the region 2800-2300 cm⁻¹. It also showed a sharp, intense absorption at 1620 cm^{-1} and a C=N⁺ absorption similar to that of (1) at 1580 cm^{-1} .

The 1 H n.m.r. spectrum (CDC1₇) contained signals at δ =7.2p.p.m. (ca. 22H, complex multiplet, but showing separation into two groups in the ratio ca. 2:3); 4.00, 3.71, 3.42 and 3.00 (each signal lH, septet, **J** = 7Hz); and 1.52, 0.96, 0.86 and 0.64 p.p.m. (each signal 6H,, doublet, J = 7Hz). In later samples prepared by another method **(see** below) two additional signals separated from the aromatic resonances. One was exchangeable; the other showed variation in chemical shift, sometimes occurring as high as 6.4 p.p.m. The ¹¹B n.m.r. spectrum was virtually identical to that of (l) .

The 13 C n.m.r. spectrum (CDC1₇) showed signals due to four different isopropyl groups $(6 = 19.14, 19.84, 20.89, 22.67 p.p.m., all quarters in the off-resonance decoupled mode; 44.33 ,$ 47.21, 47.75, 52.71 p.p.m., all doublets in the off-resonance mode), two of which correspond closely to those of (l) , and signals due to two different sets of phenyl resonances $(6 = 125.71,$ 126.08 p.p.m., para-carbons; 126.68, 127.08 p.p.m., meta-carbons; 132.40, 134.58 p.p.m., orthocarbons; ca. 148, 150 p.p.m., broad, weak, ipso-carbons). Additionally, the immonium carbon atom which is present in (1) appeared as a weak broad signal at ca. 200 p.p.m. and there was a carbonyl signal at 162.63 p.p.m. Significantly, this last signal was a doublet in the off-resonance decoupled spectrum showing that the fonnyl hydrogen atom of the DIPF unit **was** present. **Re** fomyl proton was not observed as a separate signal in the initial 1 H n.m.r. spectrum, but its location was verified by on-resonance decoupling of the aromatic protons, which caused the ¹³C n.m.r. signal for the carbonyl carbon atom to collapse to a singlet.

These data appeared to indicate that the product was a 1:1 complex between (1) and DIPF, a highly surprising result in view of the structure of (l) . Further support was obtained from the field desorption mass spectrum, $3\text{ which showed substantial peaks at m/e 604 (ion corresponding$ to (1) + Pr₃NCHO), 475 (loss of Pr₃NCHO), and 398 (further loss of Ph). Simple confirmation of this structure was achieved by mixing equimolar amounts of (1) and DIPF in ether, whereupon an identical complex crystallized spontaneously on evaporation.

In view of the surprising ability of (1) to form a stable, crystalline 1:1 complex with DIPF we have investigated its ability to complex other compounds. Compound (1) and the appropriate-substrate were mixed in a 1:1 ratio in ether and the solvent removed at room temperature. Complex formation was indicated by a significant change in the nature of the 0-H stretching frequency in the i.r. spectrum. particularly in the disappearance of the band at 3550 cm-' **(see** above), and the appearance of broad bands at lower frequency. The solid adduct was then dissolved in boiling ether-hexane and allowed to crystallize, if possible.

The compounds fall into three categories: **a)** those which, like DIPF, form stable recrystallizable complexes with Q); b) those which form distinct solid adducts, **as** revealed by change of the 0-H absorption in the i.r. spectrum, but for which the adducts are not sufficiently stable to allow recrystallization from boiling solvents and c) those for which the i.r. spectrum of the 1:1 mixture is merely the sum of the separate spectra of (1) and the compound, indicating little **or** no complex formation. The table summarizes the results. e sum of the
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Table
Pr¹NCH

Nitriles and esters show no evidence for camplexation with **Q)** in contrast to simple amides and amines which form strong complexes. Complexes of amines and mides which are relatively hindered at the site of co-ordination (oxygen for amides - **see** below) are relatively weak and cannot be recrystallized. Indeed, the **even** more hindered amino acid.amide derivatives tested showed **no** evidence for complex formation.

The melting behaviour of all of the crystallizable complexes is similar. They either first melt at ca. 100 $^{\circ}$ C and then rapidly crystallize again or change their crystal shape during heating, finally melting close to the melting point of (1) (179⁰C). This could be consistent with loss of the co-ordinating molecule by evaporation during heating. The transition temperature depends on the rate of heating, and is not easily reproducible.

The boron atoms in Q) in both the solid state **(as** indicated by preliminary X-ray analysis²) and in solution $\binom{11}{B}$ n.m.r.²) are co-ordinatively saturated. Thus, complex formation, especially from relatively poor donors such **as** amides, is rather unusual. However, adduct formation could occur in one of two ways: a) proton transfer from (1) to the donor to give a salt; or b) opening of the heterocyclic ring of Q) to give one boron atom in **^a**3-co-ordinate state capable of camplexation; the other boron atom would remain **as** a hydroxyborate. There is no evidence to suggest that **(JJ** is in any way acidic, and the changes in the spectra of DIPF induced by its complexation with **(k) are** totally different to those induced by its reactions with trifluoraacetic, sulphurie or methanesulphonic acids. We therefore discount mechanism **(a).** The changes induced by complexation of 'DIPF vith boron acceptors such **as** diphenylbromoborane show

some similarity, but even here there **are** substantial differences. For example, the large upfield shift (ca. 1.8 p.p.m.) of the formyl proton on complexation of DIPF with (1) is not matched on camplexation with other acceptors. However, this can be rationalised by assuming complexation via the amide carbonyl' oxygen to one of the boron atoms of (1) (see below).

Complexation of DIPF through oxygen is indicated by the following observations: a)'the carbonyl stretching frequency is lowered on complexation; b) the isopropyl group methyl signals fail to coalesce at temperatures (50°C in CDC1₇, 120°C in chlorobenzene) for which the free amide signals have already coalesced, indicating increased C-N double bond character in the complex.

It is possible that (1) is in equilibrium with open-chain forms (2) and (2) in solution, providing two possible sites for coordination. **However,** models indicate that complexation of DIPF to the three-coordinate baron atom of (2) would be severely hindered, and **we** therefore favour complexation to Q) giving the complex **(9.**

The modeis also reveal that the fornyl proton in complex *(t)* must lie within the shielding zone of one of the four phenyl rings in whatever conformtion it adopts. This could account for the large upfield shift of this proton. By contrast, the formyl carbon atom sits at the edge of, or just outside, this **zone** and is affected much less. % for the large upfield shift of this proton. By contrast, the formyl carbon atom sits a
je of, or just outside, this zone and is affected much less.
It is interesting that a sample of the complex which was freshly prepare

(1) and dry DIPF showed a formyl proton resonance at $\delta = 6.4$ p.p.m.; it is also interesting that addition of either D₂0 or excess DIPF shifted this resonance downfield. Evidently there is a rapid equilibrium between free and complexed aaide. leading to an averaged signal, and water also competes with DIPF for complexation of (1). Indeed, water slowly displaces DIPF or other amides from their complexes with (1) on exposure of the complexes to moist air. Attempts to form an ammonia complex by reaction of (1) with concentrated ammonia solution always resulted in the water complex. Nevertheless, water is easily removed from the complex by passage through a

column of dry silica or by azeotmpic distillation, Passage **over** silica also regenerates **Q)** from other complexes.

In conclusion, the boron-containing betaine (1) forms co-ordination complexes with amides, amines and water which are both unusual and interesting. The upfield shift of the formyl proton of DIPF (or DMF) when complexed to (1) is exceptionally large, but can be explained by its being forced to occupy a position directly **aver** the shielding cone of one of the phenyl rings.

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- 3. We thank Dr. D. Games of University College, Cardiff for running the spectrum for us.

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