THE REACTION OF ISATIN AZOMETHINE YLIDES WITH (Z)- AND (E)-2-OXOINDOLIN-3-YLIDENE ACETOPHENONES: CONCERTED VS APPARENT NON-CONCERTED 1,3-DIPOLAR CYCLOADDITION#

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Abstract - The reaction of 1-benzyl-5-bromoisatin with proline, sarcosine, or glycine gives azomethine ylides that can be trapped by (Z) - or (E) -2-oxoindolin-3ylidene acetophenones acting as dipolarophiles in a $1,3$ -dipolar cycloaddition. The configuration of the adducts was determined by IH-nmr with nOe experiments. The kinetically controlled adducts always retain the configuration of the reacting dipolarophile but they can rearrange, sometimes under extremely mild conditions. The rearrangements were demonstrated to involve a 1,3-dipolar cycloreversion/ cycloaddition sequence since the intermediates were trapped by more reactive dipolarophiles.

Pursuing a research started by Grigg and coworkers¹ on 1,3-dipolar cycloadditions of azomethine ylides (AY), the reaction of these 1,3-dipoles derived from aminoacids and ninhydrin with (E) - and (Z) -2-oxoindolin-3-yli the reaction of these 1,3-dipoles derived from aminoacids and ninhydrin with (E) - and (Z) -2-oxoindolin-3-ylidene acetophenones was studied.2

*Dedicated to Professor Alan R. Katritzky on the occasion of his 65th birthday.

Since examples were reported3 showing loss of configuration in some reactions with **(E)-2-oxoindolin-3-ylidene** acetates, the cycloaddition with the above stereoisomers was considered a useful tool to test a loss of stereoselectivity. totally unusual in 1,3-dipolar cycloaddition. When ninhydrin-derived AY reacted with **(E)-** or **(a-2-oxoindolin-3-ylidene** dipolarophiles, adducts always retained the configuration of the reagent. Under thermal conditions, nevertheless, some adducts derived from (E) -dipolarophiles rearranged to the thermodynamically stable regioisomer⁴ through a 1,3-cycloreversion-cycloaddition reaction (Scheme 1).

The AYs are known to give adducts loosing the configuration of the dipolarophile³ derived from aminoacids and isatines. This reaction was hence tested with (Z) - and (E) -2-oxoindolin-3-ylidene acetophenones.

RESULTS

Reaction of azomethine ylide from isatin and proline. In order to label oxindole rings deriving from dipole (respectively from dipolarophile), suitable substituents were located on reagents. Thus I-benzyl-5-bromoisatin (1) and proline **(2),** the precursors of AY **(3),** were made to react with **(Z)-1-methyl-2-oxoindolin-3-ylidene** acetophenone (4a). At room temperature the reaction was accomplished within 24 hours and 86% yield of a single adduct was obtained. By nmr it was shown to be 1-benzyl-5-bromo-2-oxoindol-3-spiro-2'-(1'benzoylhexahydropytrolizin)-3'-spiro-3"-(1"-methyl-2"-oxoindole) (5a) and nOe experiments (whose results are reported in the formula in Scheme 2) allowed to determine the relative configuration of the four stereocenters. This resulted to be [3R, 1'S, 2'S, 7'aS],⁵ hence the configuration of 4a was retained in 5a.

When the reaction was performed with **(E)-I-methyl-2-oxoindolin-3-ylidene** acetophenone (4b) again at room temperature (but avoiding the summer season!) the reaction was compmed within *3* days and a single product **(5b:** mp 145.146 "C) was obtained in 73% yield (at 5 "C, same result in 12 days with 82% yield). Its purification must be performed very carefully (see Experimental) otherwise it is converted into a different product (6b: mp 209-21 1 'C) isomer of the former. 6b can be obtained directly from **1, 2,** and **4b** running the reaction at 30 *"C* (78% yield). Nmr experiments allowed to determine their structures and configurations (Scheme 3).

Both 5b and 6b are 1-benzyl-5-bromo-2-oxoindol-3-spiro-2'-(1'-benzoylhexahydropyrrolizin)-3'-spiro-3"-(1"methyl-2"-oxoindole) (isomers of 5a), but the configuration of their chiral centers is [5b: 3R, 1'R, 2'S, 7'aS] and 16b: 3S, l'R, *2'S,* 7'aSl. The crucial nOe effect characterizing 6b is a 5% enhancement of **H-4** obtained when H-**4"** was irradiated. Only a "head to head" facing of the oxindole rings is conceivable to this result, that is further supported by a 2% nOe between N-CH₃ and N-CH₂ oxoindolic groups.

The thermal conversion of 5b into 6b (and the formation of 6b at 30 $^{\circ}$ C) could involve either the cleavage of the C2'-C3 bond with formation of a zwitterion that cyclizes, or a 1.3-dipolar cycloreversion2 to 4b and **3a,** the latter isomerizing into its configurationally distinct isomer (3b), that reacts to give the thermodynamically stable adduct (6b).

The 1.3-dipolar **cycloreversion-cycloaddition** pathway was demonstrated by treating 5b in the presence of (E)-5 **chlom-1-methyl-2-oxoindolin-3-ylidene** acetophenone (4c). This was partly captured by 3b to give the new adduct (6c) in **48%** yield, whereas 4b was released and isolated (Scheme 4).

6c was obtained in high yield at room temperature from 1, 2, and 4c, and nOe experiments, reported in Scheme 4, are in agreement with its [3S, I'R, 2'S, 7'aSl configuration.

Reaction of azomethine ylide from isatin (1) and sarcosine. Sarcosine (7) and isatin (I), through a dipolar intermediate that loses carbon dioxide.2 give AY **(8).** The reactions with either 4a or 4b require 3 days at room temperature and the same adduct (9) is obtained from both isomers. The nmr spectrum reveals that 9 is 1-benzyl-5-bromo-2-oxoindol-3-spiro-2'-(4'-benzoyl-1'-methylpyrrolidin)-3'-spiro-3"-(1"-methyl-2-oxoindole) and ¹H-NOESY experiments do not show the significant relationship between H-4 and H-4" that would suggest a "head to head" facing of the oxindole rings.

If 1,3-dipolar cycloaddition occurs with retention of configuration and 9 is a primary reaction product, either 4a or 4b must isomerize under the experimental conditions and the k_{isom} , must be significantly faster than k_{cyc} of the unstable isomer.

Therefore the reaction was studied kinetically by uv-vis spectroscopic analysis. In the presence of **8,** 4a does not react but isomerizes and the second order rate constant in ethanol at 25 °C is $k_{\text{isom. 4a}} = 8.8 \times 10^{-2}$ (1 mol⁻¹ s⁻¹), while 4b gives 1,3-dipolar cycloaddition with a second order rate constant (ethanol, 25 °C) $k_{\text{cycl. 4b}} = 1.7 \times 10^{-3}$ (I mol⁻¹ s⁻¹). Thus the isomerization of 4a to 4b is 52 times faster than the 1,3-dipolar cycloaddition of the latter. The effect of this significant difference is observed in Figure 1. **A** and B are the times courses of the uv-vis

spectra of two solutions: the former contains 4a and **8** in a ratio of 1:2, the concentrations of 4a and 4b are identical. **A** increases the absorbance (isomerization of 4a to 4b), B decreases the absorbance (cycloaddition of

4b); after 9 h the uv-vis spectra are nearly superimposable and also A begins now to decrease with the same rate of B.

Figure 1. Time course of the reaction of 8 with 4a (A) and 4b (B): the a curve is the uv-vis spectrum at time 0, the b cuve after 560 minutes.

These results are summarized in Scheme 5. The [3R, 3'S, 4'S] configuration of 9, only suggested by the small values of nOe effects observed, was fully confirmed by an x-ray analysis.

Reaction of azomethyne ylide from isatin (1) and glycine. The reaction of 1, glycine (10). and the dipolarophiles may occur with or without loss of carbon dioxide, 6.7 depending on the experimental conditions: higher temperatures favour the formation of the decarboxylated dipole.

The optimum temperature to avoid mixtures again containing undecarboxylated adducts is 60 °C, thus the reactions with 4a and 4b were performed under these experimental conditions that involve the formation of AY (11). Both reactions occur without any isomerization of the dipolarophile: 4a gave a single adduct (12a), 4b gave 1Zb that separated out from the reaction mixture. Both products were found to be I-benzyl-5-bromo-2-oxoindol-3-spiro-2'-(4'-benzoylpyrrolidin)-3'-spiro-3"-(1"-methyl-2-oxoindoles). The nOe experiments, reported in Scheme *6,* assigned the [3R, 3's. 4'Rl configuration to 12a and the [3R, *3's.* **4'Sl** configuration to 12b; thus both adducts retain the configuration of their corresponding dipolarophiles.

If the reactions are run in refluxing ethanol, the result is quite different, and both 4a and 4b gave the same product (12a). Therefore this is the thermodinamically stable adduct of 4b whereas 12b is formed under kinetically controlled conditions. This was easily checked since 12b cleanly converted to 12a when refluxed three days in ethanol.

This rather unusual isomerization may occur either by proton loss and gain of H-4', or by cleavage of the C4'-C5' bond giving rise to a stabilized zwitterion, or by 1,3-dipolar cycloreversion giving 11 and 4b with the latter product equilibrating with its isomer (4a) under the thermal conditions. The first option was excluded because no deuterium incorporation was observed running the isomerization in C_2H_5OD . The last pathway was supported by the following experiment. The unstable isomer (12b) was heated in refluxing ethanol with an equimolecular amount of 4c. After 20 hours the reaction mixture was found to contain (besides a small amount of 12a) 4b, formed by the 1.3-dipolar cycloreversion, and a new adduct (12c), incorporating 4c, that was the result of a new 1.3-dipolar cycloaddition between AY (11) and the new (and more reactive) dipolarophile (4c). The nmr revealed 12c has a [3R, 3's. 4R] configuration, hence a formal loss of the configuration of 4c was accomplished. This is again due to the experimental conditions, since the reaction of 1, 2, and 4c (or its Z isomer 4d) gave the same 12c above described (Scheme 7).

DISCUSSION

Is 9, the product of both 4b and 4a with sarcosine-AY **(8).** an adduct with a stable configuration or is it only the result of a kinetically controlled 1,3-dipolar cycloaddition? The experiment supports its stability (no other product was obtained from 4a or 4b), the above results cast doubt on this.

When 9 was heated into a sealed vial in ethanol (48 h, 120 °C), it cleanly converted into its isomer (13), whose configuration, by nOe experiments reported in Scheme 8, was found to be [3R, 3'R, 4'S], hence with "head to head" facing of the oxindole rings.

The overall balance of l,3-dipolar cycloadditions of AYs with 2-oxoindolin-3-ylidene acetophenones could be rationalized since both E and Z isomeric dipolarophiles were available and their reactivity was hence studied in details.

A loss of configuration of the dipolarophile in an AY-cycloadduct can be the result of an equilibration of the oxoindolidene derivative. This occurs with Z-isomer (4a) when it reacts with sarcosine-AY **(8).** It does not occur either with 3 or with 11. It is nevertheless important to run the reactions when the aminoacid has already given rise to the AY, otherwise isomerizations would be much more diffused: proline causes isomerization of **4a,** its ylide (3) not. Thus l,3-dipolar cycloadditions of AY and oxoindolidene derivatives always occur with retention of configuration of the dipolarophile to give a kinetically controlled product. This can rearrange under more or less severe conditions to a differing isomer through 1,3-dipolar cycloreversion/cycloaddition sequence. The configurations of the stereocenters of 5 and 6 are due to the steric interactions developed by the proline ring in the transition state. Thus, 3a is the kinetically formed $AY¹$ that reacts with 4b through the less congested transition state (14) to give the kinetically conuolled adduct (5b). The reaction of 3b with **4b,** to give the thermodynamically stable isomer (6b), again involves the less hindered transition state that now is 15 (Scheme 9).

The isomerization of sarcosine-AY adducts (9) **vs** (13), though occuring under much more severe conditions, can **be** rationalized with a somewhat similar approach.

Finally, the formation of kinetic **vs** thermodinamic adducts of glycine-AY has a different origin. Again it is the result of a 1.3-dipolar **cycloreversion/cycloaddition** sequence, but involving a thermal equilibration of the dipolarophile also.

In conclusion, these reactions must be carefully run under conditions preventing any isomerization, otherwise adducts can be obtained that, at a first glance, may appear as a proof of a non-concerted character of the cycloaddition.

EXPERIMENTAL

Melting points were determined by the capillary method. Elemental analyses were obtained with a C. Erba mod. 1106 **CHN** analyzer. 'H-Nmr spectra were recorded on a Bruker 3M) spectrometer, **ir** spectra (nujol mulls) on a Perkin Elmer 983 spectrophotometer. The stationary phase for the column chromatography was Merck silica gel (230-400 mesh ASTM).

Reaction of azomethyne ylides and 2-oxoindolin-3-ylideneacetophenones. General method. A mixture of equimolecular amounts (5 mmol) of isatin **(1).** aminoacid (2 or 7 or 10) and 2-oxoindolin-3-ylidene acetophenones (4a-c) in 98% aqueous ethanol (7.5 ml) was stirred under the conditions reported in Table 1. In general the product separated out was filtered, while in the case of 12a the solvent was evaporated and the residue was crystallized. The physical characteristics and the elemental analyses of each adduct are reported in Table 1, their 'H-nmr spectra in Table 2.

Rearrangement reactions of the adducts (5b, 9 and 12b). General method. A solution of the unstable adduct (5b or 9 or 12b) (0.5 mmol) in ethanol (5 ml) was heated under the conditions described in Table 3. Upon cooling or by concentration of the solvent, the isomerized adduct separated, whose analytical and spectral data **are** identical to those of a sample when already described.

Reaction of 5b and 4c. A mixture of equimolecular amounts of 5b (1 mmol) and 4c was stirred in EtOH (25 ml) 20 days at room temperature. The white solid separated (62% yield) was filtered and found (by nmr) to be a mixture of **6c** and 6b in a ratio 3:l. **6c** can be isolated by fractional crystallization with AcOEt and its physical

and spectral data are identical with a sample synthesized as above. The reaction mother liquors were evaporated and the residue was column chromatographed (eluant, CH_2Cl_2) and unreacted 4c (18% yield) and 4b (72% yield) were isolated in the order.

Table 1. Reaction conditions, physical characteristics, and elemental analyses of adducts (Sa,b, 6b,c, and 12a-c)

a) Dissolved at 20 °C, concentrated under vacuum and chilled;^{b)} from 4b; from 4a same conditions, yield about 80%;^{c)} from 4a; from 4b 3 days under reflux (86%);^{d)} dissolved in hot benzene and crystallized upon dilution with cyclohexane;^{e)} analyzed as uncrystallized product.

	5a	5 _b	6b	бc	9b	12a	12 _b	12c	13
$CH2-1$	4.47 d	4.11 d	4.92 d	4.81 d	4.30d	4.58d	4.26d	4.58 d	4.78 d
	5.12d	5.05d	5.00 _d	4.87 d	4.80d	5.01 _d	4.93 d	5.12d	4.90 d
$H-4$	7.84d	7.90d	6.63d	6.64d	7.78d	7.61 d	7.68 d	(c)	6.17d
$Me-1'$	----	----	----	----	2.16s	$---$			2.18s
$H-1$	5.73 d	4.54d	5.43 d	5.45 d	$\overline{1}$	$---$			$---$
$H-4$	----	\overline{a}	----	$\frac{1}{2}$	4.88 dd	5.62 dd	472t	5.64 dd	5.35 dd
$H-5\alpha^d$	(c)	3.90 m	(c)	(c)	4.69t	4.25t	5.01 dd	4.26t	4.26 dd
$H - 5 \betad$	(c)	3.00 _m	(c)	(c)	3.41 dd	3.80 dd	3.75 dd	3.74 dd	3.70 dd
$H-7a$	4.97 m	5.70 m	4.80 m	4.65 _m	\cdots	----		----	$--- -$
$Me-1"$	3.26s	2.83 s	2.40 s	2.81 s	2.80 s	3.26s	3.00 s	3.62 s	2.40 s
$H-4"$	7.27 dd	(c)	7.87 dd	7.80 dd	(c)	7.15 dd	(c)	7.24 d	7.50 dd
$H-7"$	(c)	(c)	$6.38\,dd$						
o -COH e	(c)	(c)	(c)	(c)	(c)	7.90 _m	(c)	7.91 m	7.10 m
$J_{1,7a}$ f	10	9.5	8.0	8.2					
$J_{4,5,\alpha}$ f					9.3	11.0	8.0	11.0	5.5
$J_{4,5,\beta}$ ^f					7.5	9.0	8.0	9.0	10.0
$J_{4\alpha,5\beta}$ f					9.0	11.5	12.0	11.0	9.0

Table 2. 'H-Nmr data of compoundsa

a)CDCl₃ as solvent; b)acetone d-6 as solvent; ^{c)}not determined; ^d) α and β are referred to formulae in their respective schemes; e)*ortho* aromatic protons in the 4'-benzoyl group; D_{Hz} .

Table 3. Experimental conditions and results of the rearrangements.

	Starting product Rearranged product	T/C	Time	Yield $(\%)$
5b	6b	78	$10 \,\mathrm{min}$	70
9	13ª	120	48 h	74
12b	12a	78	3 d	80

mp: 150 °C (from EtOH); ir: $v_{C=0}$; 1724, 1710, 1686 cm⁻¹. Elemental Anal. Calcd for C34H28N303Br: C, 67.33; H, 4.65, N, 6.93. Found: C, 67.02; H, 4.80; N, 6.88.

Reaction of 12b and 4c. A mixture of equimolecular amounts of 12b (0.2 mmol) and 4c was refluxed in ethanol (15 ml) for 20 h. The white solid separated (52% yield) was filtered and found (by nmr) to be a mixture of 1Za and 12c (ratio 5:1); the latter product was separated by fractional crystallization with AcOEt. The mother liquors

were chromatographed as above and, besides a small amount of unreacted 4c, 4b was separated in 57% yield.

Kinetic determinations. The overall reaction rates were measured by following the variation of the uv-vis absorption on a Perkin-Elmer Lambda 16 spectrophotometer provided with a thermostatted cell transport assembly and an automatic multicell programmer. The solutions were measured in 1.00 cm OS Hellma couvettes with 3 ml capacity and the rate constants were determined as follows. Two solutions were prepared by weighing accurately in 10 ml volumetric flasks, as far as possible, identical amounts of 4a and 4b and dissolving them in ethanol. In the specific example the concentrations were 5.06 x 10-3 M. A third solution, containing AY **8** in ethanol (in the same specific example the concentration was 5.44×10^{-3} M) was prepared by weighing accurate equimolecular amounts of isatin (1) and sarcosine (7) in a 10 ml volumetric flask. Four cells were filled with 0.3 solution of 4a or 4b, 0.6 **ml** solution of **8** and 1.5 ml of ethanol, a11 accurately measured with mycrosyinges. A reference cell was filled with 0.6 ml solution of **8** and 1.8 ml of ethanol. At time = 0 both the spectra and the absorbance at 430 nm of the cells, thermostatted at 25 $^{\circ}$ C, were registered. At time intervals of 120 min the variation of the ahsorbances and the change of the spectra (Fig. I) were measured. The second order rate constants were then calculated with the standard procedures.

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