## Photocyclyzation of 2-azidobenzophenone

Andrei Budruev, Darja Sinjagina, Svetlana Scobeleva
Nizhnii Novgorod State University, Gagarin Ave., 23, Nizhnii Novgorod, 603950, Russia.

## E-mail: budruev@gmail.com

### **Abstract:**

Thermolysis of the 2-azidobenzophenone azide group in the course of 1,3-dipolar cyclization reacts with carbonyl group without intermediate formation of nitrene. The reaction gives high yields of 3-phenylanthranil.

Photolysis of 2-azidobenzophenone as well as other aromatic azide gives singlet nitrene as decomposition products. Small singlet-triplet gap wikens triplet forbiddance partly and transition from the singlet into triplet state becames possible by means of intersystem crossing. Thus, two high reactivity intermediates form under photolysis of the azide. It determines the subsequent reactions. The singlet nitrene can insert into carbonyl group giving 3-phenylanthranil, and the triplete nitrene can dimerise to form azocompound or abstract an hydrogen atom from solvent to form amines.

The 2-azidobenzophenone photolysis in acetonitrile we have found high yields of 3-phenylanthranil. The scheme becomes complicated by the 3-phenylanthranil secondary decomposition. No azocompound was found. We have proposed that there has place transition of nitrene into low reactive conjugated byradical having electron density localized on nitrogen atom of nitren and oxygen atom of carbonyl group. The byradical gives 3-phenylanthranil by means of introsystem crossing into the singlet state.

**Keywords:** 2-azidobenzophenone, 3-phenylanthranyl, aromatic azide photochemistry, photolysis, nitrene

**Introduction:** Thermal, catalytic, and photochemical lability of azide group gives us a possibility to use aromatic azides as reagents to form nitrogen – carbon and nitrogen – heteroatom bonds.

Today, catalysis by transitional metal salts is the most actively developed direction in heterocycles synthesis [1]. However, in the reactions with substrates that can easily coordinate such salts (with biopolymers, for instance) their introduction into reaction media turns into irreversible sorption. Biopolymers can loose their biological activity in such circumstances.

The shortcoming is absent in photochemical reactions, when we have an ability to choose proper wavelength of actinic irradiation. However, the formation of radicals and other intermediates

in the excited state complicates the photolysis mechanism and decrease the yield of the desirable product. That is why a search of optimal ways to form the concrete product in the photochemical reactions is very important.

In this paper we described our preliminary results of the 2-asidobenzophenone photolysis in acetonitrile.

## **Experimental Procedure**

## **General information**

# 2-asidobenzophenone synthesis

2.66 g (0.0135 mole) of 2-asidobenzophenone, 15 ml of conc. HCl, and 5 ml of water were place into a glass with volume of 200 ml. The mixture was stirred up to formation of gray homogeneous suspention. After that the reaction mixture was cooled and solution of 1.05 g of NaNO<sub>2</sub> in 10 ml of water was added dropwise into the glass under constant stirring. The precipitate obtained was dissolved to give yellow-brown solution, that was hold in ace in 40 min. That time the solution became brown and precipitate was formed. The diazonium salt was filtered, and 2.69 g of NaN<sub>3</sub> was added into reaction media under constant stirring. After that the reaction media was hold in dark for 12 hours.

The solution decame transparent with dark oil on the bottom. Azide was extracted 3 times using 30 ml and 2 times using 10 ml of  $CH_2Cl_2$ . Extract was dried above calcined  $Na_2SO_4$ , after that the solvent was removed in vacuo. 2-asidobenzophenone was purified with preparative colomn chromatography using silica gel and heptane- $CH_2Cl_2$  (1:1, v/v) as an eluent. After purification the azide became light-yellow. IR spectrum of the azide is shown on Fig. 1.

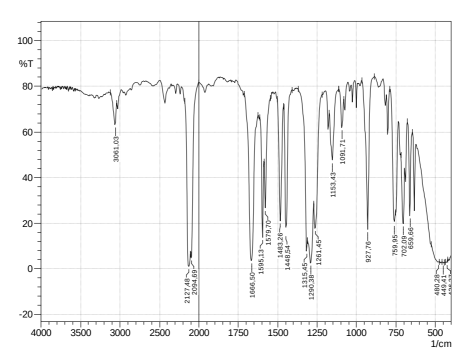


Fig.1. IR spectra of 2-asidobenzophenone in nuiol.

**Synthesis of 3-phenylanthranyl. 0.157 g of 2-azidobenzophenone was boiled in 5 ml of o-cresol** (b.p.=144 °C) in 30 min. This time is sufficient for complete decomposition of azide ( due to decrease of the azidogroup asymmetric stretching vibration). The color of the solution became from yellowish to saturated yellow. The solvent was removed under heating in vacuo. The isolated crystals were thrice recrystalized from ethanol (1-2 ml) with separation of filtered products under the solvent cooling below to 19 °C. Mass-spectrum was measured and it was in closed coincidence with known from literature [2] (Fig.2).

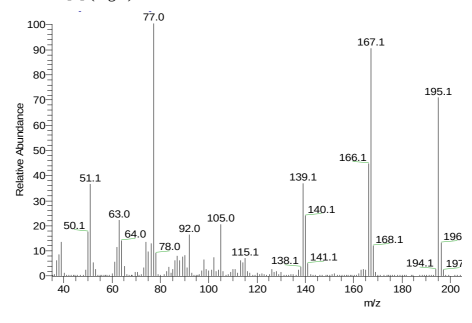


Fig.2. Mass-spectrum of 3-phenilantranil

**Synthesis of acrydone.** 0.157 g of 2-azidobenzophenone was boiled in 10 ml of undecane (b.p. 198 °C) for 10 h. This time is sufficient to decompose the azide completely. The solvent color became from light-yellowish into brown-red. After cooling the dark-brown precipitate was formed. The raw material was recrystalized from boiled undecane. The absorption maxima in IR (Fig. 3) and u.v. spectrum (Fig. 4) are in closed resemblance with the known from literature (SDBS [3] and NIST [4] databases).

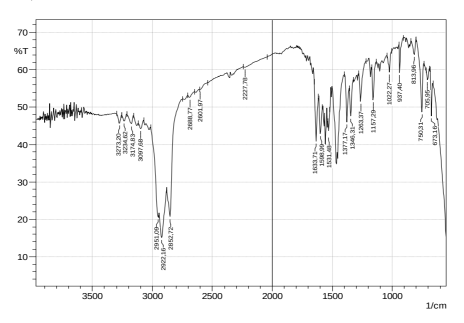


Fig.3 IR spectra of 9(10H)-acridone in nuiol

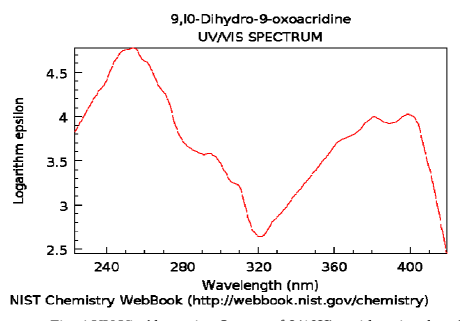


Fig. 4 UV-Vis Absorption Spectra of 9(10H)-acridone in ethanol

**Photolysis of 2-azidobenzophenone solution.** Photolysis of 2-azidobenzophenone dissolved in acetonitrile was performed in 1 cm quartz quevette. The low pressure Hg-lamp (Buf-15) was

used to irradiate the relevant solutions. The concentrations were  $5 \times 10^{-4}$  and  $5 \times 10^{-5}$  M.

**Measurement of u.v. spectra.** The u.v. spectra of the dissolved 2-azidobenzophenone and its photolysis products were measured with Analitik Jena SPECORD-40 in 200-700 nm region.

**Measurement of IR spectra.** IR spectra of the 2-azidobenzophenone and its photolysis products in protic and aprotic solvents were measured with IR-Furrie spectrophotometer Shimadzu IRPrestige-21 in KBr cavities in 4000-400 cm<sup>-1</sup> range.

Chromatographic separation of the reaction products. In order to indentified the 2-azidobenzophenone products we used high efficiency liquid chromatographer Shimadzu LC-20 equipped with C-18 colomn. We used gradient eluting with ethanol:0.01% phosphonic acid with linear declining from 65.5% down to 0% in 20 min and in 25 min in pure ethanol.

### **Discussion**

Thermolysis of 2-substituted phenylazides containing multiple bond in  $\alpha$  and  $\beta$  positions relative to azide group gives rise to formation of the intramolecular cyclization products with quantitative yields. High yields of heterocycles is due to intramolecular 1,3-dipolar cycloaddition when cyclization and nitrogen loss take place simulteniously.

The main 2-azidobenzophenone (1) thermolysis products are 3-phenylanthranyl (2) and thermally stable 9(10H)-acrydone (3). Formation of 3 takes place in more strict conditions (under prolonged warming) (Scheme 1).

$$\begin{array}{c|c}
 & 140 \, {}^{\circ}\text{C} \\
\hline
 & 1 \\
\hline
 & 1 \\
\hline
 & 2 \\
\hline
 & 200 \, {}^{\circ}\text{C} \\
\hline
 & 10 \, h
\end{array}$$

Scheme 1. Thermolysis of 2-azidobenzophenone

Photochemical decomposition of arylazides takes another reaction path. After u.v. irradiation the azide transfers into excited state and then decompose to form singlet nitrene and molecular nitrogen.

Taking into account literature data and using analogy with 2-azidobenzoic acid and its esters

we proposed that nitrene formed under u.v. irradiation of 1 was able to insert into multiple bond of carbonyl group giving 2 (scheme 2).

Scheme 2. Photolysis of 2-azidobenzoic acid and its esters

On the other hand the photochemical formation of **3** is possible in analogy with photochemical cyclization of 2-azidobiphenyl to turn into carbazole that has place with yield close to quantitative one (scheme 3).

Scheme 3. Photochemical formation of acrydone

So, we can concluded that in the course of the photolysis of 1 there is a possibility to form 3 after insertion of nitrene into C=C bond of benzene ring with simultaneous rearrangement.

So, the possible scheme of photolysis of 1 can be presented as (scheme 4):

Scheme 4. Proposed scheme of the 2-azidobenzophenone photolysis

Counter synthesis of the possible photolysis products and their identification by uv-vis spectra measuring gave us a possibility to determine characteristic bands of absorption. In the case of 2 this region is long wavelength maximum (350 nm), and for 3 the bands correspond to absorption with maxima more than 400 nm.

Separation of the photolysis products was performed by means of liquid chromatograph with

diode matrix detector. They were identified with coincidence of their electronic spectra and due to retention times by proposed reaction products in the case of 2 and 3 (scheme 5). Besides, there also were measured UV-VIS spectra of the irradiated solutions of the azides.

Scheme 5. The 2-azidobenzophenone photolysis

In the case of **1**, and of the related ketones there high quantum yield of the intersystem crossing, that is singlet nitrenes obtained in the course of the azide photolysis with large probability transfer into the triplet state. The triplet state of nitrene is greatly different from the singlet state so that their cyclization is possible only as the two-step reaction. At first there has place the triplet biradical intermediate formation; at second there proceeds addition after intersystem crossing in the products.

However, analysis of the UV-VIS spectra of the photolysis products for 1 gives us a possibility to conclude that at small irradiation times of the azide solution in acetonitrile ( $5 \times 10^{-5}$  M) there observes an increase of absorption of 2 with asobestic point preservation that evidence that there has place formation of 2 with yields close to quantitative ones. Prolonged irradiation disturbs the isobestic point preserving, however the increase of absorbance in the region of absorbance of 3 was not observed.

Photolysis of 2 in the experimental conditions used does not form 3.

High yields of 2 from the photolysis (Scheme 6) and the high quantum yield of intersystem crossing of nitrenes forced us to propose the triplet path of their cyclization. It is possible that the triplet nitrene formed isomerizes into the ortho-quinone type biradical product with possible electron density localization on the oxygen atom of carbonyl group and nitreneous nitrogen atom. Due to electron density redistribution this particle is more stable than the triplet nitrene and being not reactive forms 2 after the second change of the spin state.

Scheme 6. The 2-azidobenzophenone photolysis

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