Direct observation and characterisation of 3-azido-2H-azirines: postulated, but highly elusive intermediates†‡

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The azido group, although absent in natural products, is a highly versatile functionality which has found immense applications in various branches of chemistry, as well as biology and materials science. In particular, vinyl azides are useful starting materials for the synthesis of strained compounds, for example azirines, via photolysis or thermolysis and also for ring-enlarged heterocycles of synthetic and biological significance.

Although the parent compound 4 has not yet been reported and only four substituted representatives of 1,1-diazoethenes, 1, 2a, 2b and 3, were mentioned in the literature, a rich chemistry, particularly with reactions of the push–pull olefin 1, was reported by Saalfrank and others (Scheme 1a). The treatment of 1 with different types of amines, for example, led to nitrogen heterocycles, such as 2H-azirines, 1,2,3-triazoles and tetrazoles, as depicted in Scheme 1b.7,9,10 Unimolecular reactions of 1, 2a and 2b induced by thermolysis or photolysis were also investigated intensively.7,8,10,11 In the case of substrate 1, extrusion of two or three molecules of dinitrogen and formation of the products 7 or 8, respectively, were claimed (Scheme 1c).7,10,11 The thermal reaction of 1 was explained with the help of short-lived intermediate 5 and the subsequent cleavage of the C–C bond in the three-membered ring, combined with the loss of a second molecule of dinitrogen. Finally, a two-step substitution of the cyano group by nucleophilic methanol should result in the generation of 7.

Irradiation of a solution of 1 in methanol at −30 °C gave mainly “the thermal product” 7 (70%) and the acrylic ester 8 as a side product (30%).11 The diazo compound 6 and the derived carbene (vinylidene species) were discussed as short-lived intermediates in the surprising formation of 8. It is well...
known in the chemistry of vinyl azides that these substrates generally lead to high yields of the corresponding 2H-azirines under low-temperature irradiation, whereas thermolysis of the same azides can give varying results: in some cases, heating a solution of a vinyl azide also afforded a high yield of the 2H-azirine, but other such substrates avoided the thermal generation of these heterocycles and furnished quite different products. Thus, the reported unimolecular reactions of 1 are in contrast to the general rules of vinyl azide reactivity.4,5

Because 3-azido-2H-azirines are only postulated7,10,11 however, never detected species, and intermediates of types 5 and 6 are obviously not only elusive, but also highly interesting, we judged low-temperature photolyses of diazide 1 and the parent compound 4 to be promising. We became further interested in such experiments, since we previously investigated the low-temperature irradiation of open-chain 1,2-diazidoalkenes,7,20 which led to the first proof of long-sought 2-azido-2H-azirines.12 Such short-lived species turned out to be intermediates in the known transformation of vicinal vinyl diazides into two nitrile fragments, that is accompanied by the cleavage of the C=C bond and loss of two molecules of dinitrogen. In the case of 3-azido-2H-azirines, the electron-donating properties of the azido group should be able to stabilize the azirine ring by reducing the electron deficiency at the sp2 carbon atom. This effect is well documented especially for 3-amino-2H-azirines, which bear, however, a stronger donor.13 On the other hand, it is well known that the direct combination of strain resulting from small rings decreases the stability of azides.

Since light energy circumvents the problem of direct thermal agitation, we planned to irradiate 1 at low temperatures and to detect short-lived intermediate 5 by low-temperature NMR spectroscopy. Surprisingly, substrate 1 was not soluble in methanol at −30 °C, and the same is true even for a 3:1 mixture of chloroform and methanol, although the solubility of 1 in neat chloroform is excellent. Thus, we photolysed a solution of 1 (30 mg) in deuterated chloroform (0.75 mL) with the help of a mercury high-pressure lamp (filter with λ > 320 nm), maintaining the temperature at −60 °C through a cryostat.14 After 1 h, low-temperature NMR analysis showed complete consumption of 1 and the formation of 2H-azirine 5 with 78% yield (Scheme 2 and Fig. 1). Under our conditions, prolonged irradiation (3–4 h) did not lead to a significant decay of azido compound 5, whereas decomposition of this compound occurred when the temperature of its solution was raised above −10 °C. The structural proof of 5 is mainly based on the corresponding 13C NMR data and especially on the characteristic chemical shifts of the 2H-azirine unit with δ (C-2) = 32.8 and δ (C-3) = 150.1. The assignment of these signals was also supported by CIGAR experiments.

To provide further evidence of the structure of 5, we utilised strain-promoted alkyne–azide cycloaddition with the help of cyclooctyne to obtain the 1H-1,2,3-triazole 9.15,16 Although the product 9 showed limited stability at room temperature, most probably because of the electron-withdrawing properties of the triazole unit, which increases the electron deficiency at C-3 of the 2H-azirine, we were able to isolate 9 as a straw-coloured oil in 57% yield, and this allowed complete characterisation of the trapping product.14 We also monitored the reaction of 5 with cyclooctyne in deuterated chloroform by using 1H NMR spectroscopy to get an impression of the rate of this cycloaddition (Fig. 2). It turned out that the consumption of 5 was very slow at −60 °C, whereas a rapid formation of 9 was observed at −20 °C.

Encouraged by successful characterisation of azirine 5, we were highly motivated to generate the parent compound, 3-azido-2H-azirine (16), which requires the synthesis of unknown 1,1-diazidoethene (4). Obviously, the latter compound cannot be prepared from 1,1-dichloroethene by nucleophilic substitution owing to the absence of electron-withdrawing groups that facilitate the access to diazides 1–3. Therefore, it seemed logical to utilise the chloroethane derivative 12 as a precursor for the desired diazide 4 (Scheme 3). However, all our attempts failed when we tried to directly transform chloroacetalddehyde into 12 by using well-established methods17 for the synthesis of geminal diazides from aldehydes. Subsequently, we employed the multifarious chemistry of
α-azido alcohols to prepare 12 in a three-step sequence. Thus, we treated chloroacetaldehyde with hydrazoic acid and reacted the resulting α-azido alcohol 10 with phosphorus tribromide to obtain the intermediate product 11 with 48% yield based on the aldehyde. Selective nucleophilic substitution of 11 with the help of N,N,N′,N′-tetramethylguanidinium azide (TMGA) led to the desired diazide 12, which was transformed into 1,1-diazidoethene (4) in the presence of potassium tert-butoxide in diethyl ether.

The azides 11, 12 and particularly 4 are dangerously explosive and should be handled and characterised in solution only, for example, by IR and NMR spectroscopy. Because of its substitution pattern, 4 is an electron-rich olefin as shown by an upfield shift of the NMR data with $\delta^{(13C)} = 85.3$ and $\delta^{(1H)} = 4.40$ measured for the CH$_2$ group (CDCl$_3$). 1,1-Diazidoethene (4) has a limited stability at room temperature and a half-life period of 220 min at 40 °C in deuterated chloroform. With the help of cyclooctyne, explosive azides 4, 11 and 12 were transformed into the corresponding 1,2,3-triazoles 15, 13 and 14, respectively, and these heterocyclic derivatives were easily isolated and characterised.

A solution of 4 in deuterated chloroform with diethyl ether as an internal standard was irradiated at −55 °C using a mercury high-pressure lamp. After 30 min, NMR low-temperature analysis showed the formation of 3-azido-2H-azirine (16) in 34% yield (Scheme 3). $^1$H NMR spectroscopy at −40 °C indicated a short half-life period of approximately 12 min. Thus, 16 is very short-lived, and attempts to intercept this species in the presence of cyclooctyne did not lead to the desired cycloadduct 17. This unsuccessful experiment exhibits the limit of trapping unstable azides with the help of cyclooctyne.

The photolysis of 4 was also monitored with solutions of the diazide in toluene-d$_8$ at −55 °C by using NMR spectroscopy (Fig. 3). The product 16 was identified by its high-field $^1$H NMR signal at $\delta = 1.49$ (singlet) and the characteristic $^{13}$C NMR signals at $\delta = 27.14$ (C-2) and $\delta = 161.25$ (C-3), and the coupling constant $^1J^{(13C, 1H)} = 182$ Hz is also typical of 2H-azirines.

Finally, we utilized in situ IR spectroscopy to detect the progress of the photochemical step 4 → 16 in deuterated chloroform at −45 °C (Fig. 4). Signals at 2120 cm$^{-1}$ (asym. N$_3$), 1641 cm$^{-1}$ (C=O) and 1311 cm$^{-1}$ (sym. N$_2$) were assigned to the starting compound 4, whereas signals at 2148 cm$^{-1}$ (asym. N$_3$) and 1756 cm$^{-1}$ (C=O) were attributed to the product 16.
Conclusions

In summary, we have demonstrated that the irradiation of 1,1-diazidothene 1 and 4 leads to 3-azido-2H-azirines 5 and 16, respectively. Although these photoproducts are highly unstable, their structures are unequivocally confirmed by low-temperature NMR and IR spectroscopy as well as trapping reactions. This outcome confirms the general rule that photolysis of vinyl azides results in the formation of 2H-azirines as primary products.\(^5\)\(^,\)\(^6\) Thus, we were not able to verify previous reports\(^4\) from 1982 on the irradiation of 1 since we obtained different results.

Compounds 1, 4, 11 and 12 should be considered highly explosive, and effective safety measures should be taken when handling these substances in the pure state. We highly recommend the use of only dilute solutions in appropriate solvents.\(^2\)

Acknowledgements

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Notes and references

2 X. Zhang and Y. Zhang, Molecules, 2013, 18, 7145–7159.


14 See the ESI‡ for further details.


