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Ring construction via pseudo-intramolecular hydrazonation using bifunctional δ-keto nitrile

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An α-nitro-δ-keto nitrile efficiently reacted with hydrazines at room temperature, even in the absence of a catalyst, to afford the corresponding hydrazones; the reactions proceeded through a pseudo-intramolecular process. The hydrazone derived from hydrazine monohydrate underwent water-assisted cyclization, which yielded the corresponding diazepine. The hydrazones derived from 4-nitrophenylhydrazine and 2,4-dinitrophenylhydrazine were converted to pyridazines upon being heated in DMSO.

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The development of highly efficient synthetic methods for various organic compounds is crucial for the efficient utilization of carbon resources and for low environmental impact. Reactions not requiring protecting groups or special reagents facilitate the simplification of manipulations from the viewpoint of atom economy.1 In order to improve reaction efficiency, the collision frequency of reactants should be increased. Generally, an intramolecular process proceeds faster than an intermolecular process because the spatial proximity increases the collision frequency of reaction sites. With regard to intermolecular processes, the use of reaction fields such as capsules,2 cages,3 bowls,4 and micelles5 has been recognized as being effective because reactive substrates located close to each other make the reaction highly efficient.

In this context, we have demonstrated pseudo-intramolecular processes that proceed under very mild conditions. Indeed, α-arylated and α-nitrated β-keto esters underwent transacylation6,7 with amines very smoothly and rapidly to afford the corresponding amides without any detectable by-products. In these reactions, the highly acidic keto ester reacted to form the corresponding ammonium salt with the amine immediately. Then, under equilibrium, the electrophilic keto ester and the nucleophilic amine were regenerated simultaneously and in close proximity. As a result, the reaction proceeded efficiently like an intramolecular process.

The α-nitro-δ-keto nitrile 18 also satisfies two criteria of a substrate that is to be used in a pseudo-intramolecular reaction: it has an acidic hydrogen and a reactive functional group in the molecule. Indeed, vicinally functionalized 1,4-dihydropyridines were synthesized upon treatment of the keto nitrile 1 with amines.9 Furthermore, the reaction of 1 with a diamine entailed tandem cyclization to afford a 1,7-diazabicyclo[4.3.0]nonane derivative, in which pseudo-intramolecular imination was a key step.10 These results prompted us to study the pseudo-intramolecular reaction of 1 by using a hydrazine instead of a diamine as a dinucleophile (Scheme 1).

Initially, the keto nitrile 1 was allowed to react with unsubstituted hydrazine (2a) at room temperature in CH3CN. After stirring for 48 h, the formation of a spot different from those of 1 and 2a was confirmed on TLC. Structural analysis on the basis of spectral and analytical data clarified the formation of diazepine 7a (Scheme 2).11 The 1H NMR signal of the methyne proton of 1 (Hα) at 6.24 ppm was considerably shifted to 4.88 ppm (Hγ). Moreover, the signals of the carbonyl and cyano carbons shifted from 207 to 138 ppm and from 112 to 172 ppm, respectively, in the 13C NMR. These spectral changes indicate that 2a reacted with both functional groups of 1. The formation of 7a presumably proceeded via pseudo-intramolecular hydrazonation followed by nucleophilic attack on the cyano group. To confirm this hypothesis, the formation process of 7a was monitored by NMR spectroscopy. Just after the addition of 2a to a solution of 1 in CD3CN, small signals of the hydrazinium salt 3a and the diazepine 7a were observed in the 1H NMR, in addition to the signals of 1.12 All the signals of 1 and 3a gradually disappeared, converting to the signals ascribed to 7a.

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On the other hand, methylhydrazine (2b) exhibited different reactivity under the same reaction conditions (Scheme 1). The reaction was considerably more complicated, giving an intractable mixture. Similar complications were also observed in the reactions with bulky t-butylhydrazine (2c) and N,N-dimethylhydrazine (2d). Hence, we monitored the reaction of 1 with 2b by 1H NMR to elucidate the reaction behavior of the substrates. When 2b was added to a solution of 1 in CD$_3$CN, significant changes in the resonance corresponding to both reagents were observed: the singlet signal at 6.24 ppm, attributed to the methyne proton (H$_a$) of 1, completely disappeared, and a couple of singlets with equivalent intensities at around 1.2 ppm and attributed to the two enantiotopic methyl groups (b and c) coalesced into a singlet signal at 1.23 ppm. In addition, the N-methyl group of 2b shifted downfield by about 0.28 ppm. These observations indicate the formation of the hydrazinium salt 3b. However, the reaction mixture became complex within several hours, even at room temperature. When DMSO-d$_6$ was employed as the solvent instead of CD$_3$CN, signals of ring products were not observed, too.

Considering the mechanism previously proposed for the reaction of 1 with 1,2-diamines, the diazepines 7a would be formed via the hydrazinium salt 3a and then the hydrazone 5a. Although the 1H NMR showed no signal for 5a, the observation of the signals of 3a supported the proposed mechanism. For alkylhydrazines 2b-d, the corresponding diazepines 6b,c were possibly formed. However, the isolation of 6b,c failed, probably because of their instability; the N-substituted diazepines 6b,c could not be converted to a stable form such as 7a by tautomerization (Figure 1).

In the present reaction, another ring closure affording a six-membered ring is also possible, as shown in Figure 2. To elucidate a plausible reaction pathway, density functional theory (DFT) calculations for the simplified model compound 8 bearing no methyl group at the β-position were performed. Comparison of the activation energies required to reach transition states showed that the activation energy for the seven-membered ring transition state is lower than that for the six-membered one. In the former case, water plays an important role; a water molecule binds to both the cyano group and the amino group to effect a water-assisted ring construction owing to the close proximity of the groups.

Although it was suggested that the hydrazone 5a was intermediately formed as mentioned above, 5a was too reactive to be isolated. Hence, we attempted the isolation of the hydrazones 5 through the use of aromatic hydrazines 2e-i. In cases of 4-methylphenylhydrazine (2e) and 4-methoxyphenylhydrazine (2f), the reactions were also too complex. In the reaction of 1 with phenylhydrazine (2g), the formation of the hydrazone 5g was confirmed by the 1H NMR of a crude product. However, the isolation of 5g was not achieved because of its rather unstable nature. The use of electron-deficient 4-nitrophenylhydrazine and 2,4-dinitrophenylhydrazine (2h,i) enabled us to isolate the hydrazones 5h,i in quantitative yields: products spontaneously precipitated within 3 h of the addition of 2h,i to a solution of 1 (Scheme 3). The structure of 5h...
was determined on the basis of its spectral and analytical data.\(^{14}\) Finally, the formation of the hydrazones was confirmed by the X-ray single crystal analysis of \(5i\) (see Supporting Information).

Hydrazone formation from the keto nitrile \(1\) and hydrazine \(2h\) (1:1) was monitored by \(^1\)H NMR spectroscopy in methanol-\(d_4\) (used as the solvent), which dissolved the hydrazone homogeneously. As a result, approximately half of \(2h\) was consumed within 1 h to afford \(5h\) in 58% yield. The hydrazonation readily proceeded under mild conditions as a result of the pseudo-intramolecular nature of the reaction; in actuality, a similar reaction of \(2h\) with acetone without involving any functional group to promote a pseudo-intramolecular process resulted in the formation of the corresponding hydrazone only in 28% yield after 1 h (Figure 3).

When the isolated hydrazone \(5h\) was heated in DMSO at room temperature for 1 day, a crystalline product was afforded; its spectral data were quite different from those expected for the corresponding diazepine. The product contained a cyano group which was confirmed by \(^1\)H NMR and IR spectra. Its \(^1\)H NMR spectrum was similar to that of the pyridazine frameworks of \(13\) (Scheme 4). \(^{15}\) The empirical formula \(C_{14}H_{15}N_5O_3\) also indicated that dehydration proceeded affording the nitronic acids \(6a\) and \(6b\) in 58% yield. The hydrazone \(5h\) was heated in a polar solvent, tautomerization of the nitro group occurred, which suggested that the nitrogen atom of the hydrazone attacked the methyne carbon to form a six-membered ring. The empirical formula \(C_{14}H_{15}N_5O_3\) also indicated that dehydration proceeded during the ring construction. On the basis of these spectral and analytical data, the product was determined to be the pyridazine \(13i\) in a similar way (at 70 °C, in 69% yield).

Thus, the reactivity of \(1\) with \(2\) varied with the hydrazine substituent used (Scheme 5). These differences are attributed to the inherent basicities and nucleophilicities of the hydrazines \(2\) and the hydrazones \(5\). The more basic hydrazines \(2a-d\) (unsubstituted and alkyl substituted) easily formed the hydrazinium salts \(3a-d\), which were observed by \(^1\)H NMR. When the hydrazine was liberated under equilibrium, the pseudo-intramolecular hydrazonation proceeded and afforded \(5a-d\). However, the nitrogens of \(5a-c\) were so nucleophilic that ring closure occurred very quickly to afford the diazepines \(6a-c\), among which \(6a\) could be converted to stable \(7a\).

In the case of the aromatic hydrazines \(2e-i\), although the hydrazinium salts \(3e-i\) were the most likely to be formed, they were hardly detected by \(^1\)H NMR because the equilibrium shifted to the intimate-pair \(4\) owing to the low basicity of \(2e-i\). Thus, the hydrazones \(5e-i\) were immediately afforded by a pseudo-intramolecular process just after the salts \(3e-i\) were formed. An aromatic ring with an electron-withdrawing substituent can stabilize the hydrazone and decrease the nucleophilicity of the neighboring nitrogen, which facilitates the isolation of the hydrazones \(5h..\) When the hydrazones \(5h..\) were heated in a polar solvent, tautomerization of the nitro group occurred, affording the nitronic acids \(11h..\) and the \(C_N\)-carbon became sufficiently electrophilic to be attacked by the anilino nitrogen to form the pyridazine frameworks of \(12h..\). The results of DFT calculations (B3LYP/6-31++G*) of model compounds \(14\) and \(15\) showed the considerably higher electrophilicity of the nitronic acid \(C_{15}\) compared to a cyano group, which is in good agreement with the experimental results (Figure 4).

In summary, we successfully synthesized diazepine \(7a\) and pyridazines \(13h..\) by treating keto nitrile \(1\) with hydrazines under mild conditions, which involved pseudo-intramolecular hydrazonation as a key step. In addition, the intermediate hydrazones \(5h..\) were successfully isolated when 4-nitrophenyl- and 2,4-dinitrophenylhydrazines \(2h..\) were used. Moreover, the DFT calculations supported the experimental results reasonably. Consequently, the present pseudo-intramolecular process will be applicable as an efficient protocol in synthetic chemistry, provided suitably designed substrates satisfying the two criteria are employed.
Acknowledgments

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References and notes

7. We consider Ballini’s reactions proceed also in a pseudo-intramolecular process; Ballini, R.; Bosica, G.; Fiorini, D. Tetrahedron, 2003, 59, 1143.
8. The keto nitrile 1 was readily prepared from commercially available ethyl nitroacetate via three steps; Nishiwaki, N.; Nogami, T.; Ariga, M. Heterocycles, 2008, 75, 675.
11. The diazepine 7a. Yellow solid. Mp 113-115 °C. IR (Nujol) 1549, 1549, 1348 cm⁻¹; ¹H NMR (CDCl₃) δ 0.91 (s, 3H), 1.07 (s, 3H), 2.17 (s, 3H), 2.17 (d, J = 19.0 Hz, 1H), 2.52 (d, J = 19.0 Hz, 1H), 4.83 (s, 1H), 6.35 (br s, 2H); ¹³C NMR (CDCl₃) δ 25.8 (CH₃), 26.6 (CH₃), 27.7 (CH₃), 33.1 (C), 41.7 (CH₂), 90.7 (CH), 137.7 (C), 172.2 (C). Analytical data were not given satisfactorily because of instability of 7a.
12. For details about the reaction toward the diazepine 7a monitored by ¹H NMR, see Supporting Information.
13. For details about the formation of the hydrazinium salt 3b monitored by ¹H NMR, see Supporting Information.
14. The pyridazine 13h. Orange plates. Mp 146-148 °C. IR (KBr) 1603, 1560, 1348 cm⁻¹; ¹H NMR (CDCl₃) δ 1.33 (s, 3H), 1.33 (s, 3H), 2.01 (s, 3H), 2.53 (d, J = 16.4 Hz, 1H), 2.66 (d, J = 16.4 Hz, 1H), 6.04 (s, 1H), 7.01 (d, J = 9.2 Hz, 2H), 7.56 (br s, 1H), 8.18 (d, J = 9.2 Hz, 2H); ¹³C NMR (CDCl₃) δ 17.2 (CH₂), 24.3 (CH₃), 24.8 (CH₃), 39.9 (C), 47.0 (CH), 83.7 (CH), 111.5 (C), 111.9 (CH), 126.2 (CH), 140.8 (C), 145.4 (C), 149.6 (C); MS (FAB) 320 (M⁺+1, 100). Anal. Calcd for C₁₄H₁₅N₅O₃: C, 55.81; H, 5.02; N, 23.24. Found: C, 55.74; H, 4.77; N, 23.07.
15. The pyridazine 13h. Orange plates. Mp 146-148 °C. IR (KBr) 1603, 1560, 1348 cm⁻¹; ¹H NMR (CDCl₃) δ 1.29 (s, 3H), 1.46 (s, 3H), 1.81 (s, 3H), 2.34 (d, J = 14.0 Hz, 1H), 2.57 (d, J = 14.0 Hz, 1H), 7.91 (d, J = 9.0 Hz, 2H), 8.37 (d, J = 9.0 Hz, 2H); ¹³C NMR (DMSO-d₆) δ 20.3 (CH₃), 26.5 (CH₃), 27.1 (CH₃), 39.5 (C), 45.0 (CH₂), 103.4 (C), 111.3 (C), 123.5 (CH), 125.0 (CH), 126.3 (C), 140.1 (C), 153.5 (C); MS (FAB) 302 (M⁺+1, 100). Anal. Calcd for C₁₄H₁₅N₅O₃: C, 55.81; H, 5.02; N, 23.24. Found: C, 55.74; H, 4.77; N, 23.07.