Review

Peculiarity of methoxy group-substituted phenylhydrazones in Fischer indole synthesis

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Abstract: We found that the Fischer indole synthesis of ethyl pyruvate 2-methoxyphenylhydrazone (5) with HCl/EtOH gave an abnormal product, ethyl 6-chloroindole-2-carboxylate (7), as the main product, with a smaller amount of ethyl 7-methoxyindole-2-carboxylate (6) as the normal product. This abnormal reaction was the result of a cyclization on the side with the substituent (methoxy group) of a benzene ring on phenylhydrazone, which was not previously observed. In this initial investigation, we focused on 1) the application of the above-mentioned abnormal Fischer indole synthesis, 2) the details of this reaction of phenylhydrazone with other kinds of substituents, 3) the mechanism of the first step of the Fischer indole synthesis, 4) the abnormal reaction in methoxydiphenylhydrazones, and 5) a synthetic device to avoid an abnormal reaction. The results of these studies are summarized herein.

Keywords: Fischer indole synthesis, methoxy group, abnormal reaction, phenylhydrazone, reaction mechanism, natural product

1. Introduction

Many natural products having an indole skeleton1) (1, Fig. 1), the compounds derived from them, and numerous purely synthetic indoles have been used as medicines.2) Thus, there are many methods for synthesizing an indole skeleton. Fischer indole synthesis3) (sometimes called Fischer indolization) is one of the oldest and most convenient, and it is currently used for a broad range of applications. Fischer indole synthesis (abbreviated as F.I.) is a method of obtaining indole compounds (4) simply by heating phenylhydrazones (3) in the presence of an acid catalyst (Fig. 2). The detailed reaction mechanism was established only recently, though the basic mechanism was presumed upon the discovery of the reaction. There are some of our contributions in that, too.

We discovered the unique properties of the F.I. of 2-methoxyphenylhydrazones (generally speaking, methoxy-substituted phenylhydrazones) at an initial stage of the synthetic study of mitomycin C4) (2) (Fig. 1) and the relevant compounds. In this report, we provide an overview of our work.

2. Fischer indole synthesis of 2-methoxyphenylhydrazone (5)

Many synthetic studies of mitomycin C and related compounds with an oxygen function at the 7-position of the indole skeleton had been reported before we began the present work. However, these previous investigations did not employ F.I. Therefore, we examined F.I. of ethyl pyruvate 2-methoxyphenylhydrazone (5) to obtain a good starting material for the synthesis of 2. The compound expected from the F.I. of 5 was ethyl 7-methoxyindole-2-carboxylate (6) (Fig. 3). In a previous work5) the compound (6) was prepared from 5 with H2SO4/EtOH, whereas a compound of unknown structure was provided with HCl/EtOH, but a detailed study was not performed.

Therefore, we examined the reaction in detail and obtained the results shown in Fig. 3 and Table 1.6,7) In the reaction with HCl/EtOH (Run 2), a compound (7) in which the chlorine was present instead of the methoxy group was obtained as the main product, but the yield of the expected compound (6) was poor.

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This was a completely unexpected result. We should note that an identical experimental result was reported independently just before our report. However, we will explain the reaction based mainly on our findings, because our study was more detailed and was performed for a longer time.

To clarify the reaction mechanism, we examined the compounds resulting from this reaction using various kinds of acid catalysts. As a result, it was found that various abnormal products were formed along with the normal product (6). The key characteristics of the reaction are the following: 1) as shown in Runs 1 and 2, when the concentration of HCl decreases, the ratio of the 6-Cl product (7) decreases, while formation of the 6-OEt compound (8) increases; 2) an indole having chlorine at a position other than 6 (compounds 10) is formed; 3) 5-substituted indoles (9) are formed in the case of the Lewis acid catalyst (Runs 5, 6) by the substitution of chlorine (by ZnCl₂) or migration of the methoxy group (by BF₃); and 4) the total yield is lower (less than 45%) in some reactions (other products were inseparable tarry ones). We will discuss the matter of the low yield of indoles later in this paper (see chapter 7).

First, we will consider the formation mechanism of the main product 7. Because product (7) has a chlorine atom in place of the methoxy group, it must be concluded that the reaction occurred on the methoxy side rather than the vacant side to give the substitution product (7). Generally, it is apt to be thought that the cyclization and substitution do not advance at the position occupied by a substituent. However, Carlin reported that the F.I. of a 2,6-dimethylphenylhydrazone gave a 4,7-dimethylindole, accompanied with the migration of the substituent (Fig. 4). However, there has been no report on an F.I. in which phenylhydrazone with a substituent side and a vacant one in the two ortho-positions was cyclized on the substituent side.

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| Run | Reagent       | 7-OCH₃(6) | 6-Cl(7) | 6-OEt(8) | 5-X(9) | 3,4,5-Cl(mix.)(10) | H⁺(11) |
|-----|---------------|----------|--------|----------|--------|--------------------|--------|
| 1   | 3M HCl/EtOH   | 15.5     | 9.9    | 4.8      | 0      | 11.6               | 0      |
| 2   | sat.HCl/EtOH  | 4.2      | 34.7   | 0.26     | 0      | 7.7                | 0      |
| 3   | HCl/AcOH      | 17.2     | 20.5   | —        | 0      | 6.1                | 0      |
| 4   | H₂SO₄/EtOH    | 12.8     | —      | 2.0      | 0      | —                  | 0      |
| 5   | ZnCl₂/AcOH    | 17.7     | 0      | —        | 4.3(X=Cl) | 0                    | 0      |
| 6   | BF₃/AcOEt     | 15.0     | —      | —        | 4.7(X=OCH₃) | —                    | 2.7    |

*: no substituent.
mechanism is based on the established mechanism of the F.I. The mechanism of the conversion from 5 to 7 and 8 via intermediates a, b, and c in Fig. 5 explains well the formation of the 6-Cl compound (7) and 6-OEt compound (8). This reaction (5→a→b→c→7) must be an S_N2-type reaction, but in addition, there must be an S_N1-type reaction that proceeds via the cation intermediate which is formed by the initial removal of the methoxy group prior to the introduction of a chloride ion, for a minor root (this latter S_N1 mechanism is not described in our original paper). In the latter S_N1 mechanism, cation rearrangement and the subsequent substitution may allow 10 to form, and an unknown reduction procedure may allow 11 to form.

The above-described mechanism is an example of a reaction using the proton acid catalyst, whereas the Lewis acid catalyst gave a different type of compound. When using the zinc chloride catalyst, on the other hand, substitution by the chlorine atom occurred at the 5-position on the indole nucleus. This fact suggests that chlorine substitution may occur at the 5-position due to the zinc chloride coordinating more strongly with the methoxy group than with the proton acid (Fig. 6). We estimate the Lewis acid-catalyst reaction as follows: In the case of the boron trifluoride catalyst, substitution did not occur but rearrangement of the methoxy group did occur via an intermediate e (Fig. 6), resulting in the weak nucleophilicity of fluoride. In other words, it appeared that the difference in products resulted from the difference between the actions of proton acid and Lewis acid on the oxygen atom of the methoxy group.

Next, we performed the F.I. of 2,6-dimethoxyphenylhydrazone (12) to determine the behavior of
the methoxy group in F.I. more clearly. The result is shown in Fig. 7. The total yield of indole products in this case was lower than that in the case of 2-methoxyphenylhydrazone. The main product in the case of the F.I. by ZnCl₂/AcOH was 5-chloroindole (13), although the yield was as low as in the case of the phenylhydrazone (5). On the other hand, in F.I. by HCl/EtOH, the expected 6-chloroindole (14) was formed in low yield, while 13 was the main product. The reason why it was difficult to substitute by SN2 for the chloride anion at the 6-position in this case is that the 7-methoxy group interfered with the approach of the chlorine. This is why the F.I. of 12 gave 5-chloroindole (13) as the main product by the SN1-type substitution, as seen in the reaction of the phenylhydrazone (5). It was clear that the F.I. of 2,6-dimethoxyphenylhydrazone (12) took place in the root of the substituent, and this fact supported the validity of the reaction mechanism of 2-methoxyphenylhydrazone (5), which is described in Fig. 5.

3. F.I. of naphthylhydrazone

Goldsmith had reported that F.I. of 2-naphthylhydrazone (15) gave benz[e]indole (16) as a result of cyclization at the 1-position, but not at the 3-position (Fig. 8). This result is rational from the viewpoint of the reactivity of the naphthalene. We performed a supplementary examination that provided the same result, and the structure of the product was correct. However, Goldsmith reported:
in the same paper that the F.I. reaction of 17 gave the benz[f]indole (18) as a result of cyclization at the 3-position. Perhaps this was because they thought that the reaction was not caused at the 1-position due to the presence of the 1-methoxy group. In addition, this result is introduced in the review\(^{11}\) as the only example of the synthesis of benz[f]indole by F.I. However, we were skeptical in regard to the structure of 18 based on our results in Fig. 3 and Table 1. We therefore replicated their experiment in our supplementary examination, and obtained a material with the same melting point, but the structure was found\(^{12}\) to be 19, as we had expected. The minor product 20 was also another type of abnormal product. In addition, we found, as in the previous reports\(^{13,14}\) that our reaction advanced equally to the side of the 1-methoxy group with another 1-methoxy-2-naphthylhydrazone. Therefore, our first result that F.I. proceeded to one of the ortho-positions to which a methoxy group was substituted was proved to be universal.

4. Syntheses of 6-alkyl- and 6-arylindole

Among the abnormal indoles described above, we focused on the 6-substituted indoles (7, 8) formed from 2-methoxyphenylhydrazone (5). This is because their relative yields were considerably high, and also because it was hard to synthesize a 6-substituted indole by another method at the time these results were reported.\(^{15}\) Therefore we intended to apply the abnormal F.I. to intentional synthesis of 6-alkyl-substituted indoles.\(^{16}\)

In this experiment we used benzene as a solvent; p-toluenesulfonic acid (TsOH), which has strong acidity but low nucleophilicity, as a catalyst; and acetylacetone, ethyl acetoacetate, and diethyl malonate as nucleophiles. As a result, when acetylacetone and ethyl acetoacetate were used, the expected 6-substituted indoles (21, 22) were obtained. However, when diethyl malonate was used, the expected 6-substituted products were not obtained at all. This was probably because diethyl malonate has low nucleophilicity in acidic media.

Next, when the same F.I. was performed using 7-methoxyindole (6) as a nucleophile, two kinds of the expected indolic dimer (23, 24) were obtained (Fig. 9). The substitution position of the 7-methoxyindole (6) part in 23 is at the 3-position, which is the most highly reactive position in the indole nucleus, and the substitution position at 24 is in the 4-position, which is activated by the methoxy group. At the time of these reports, the synthesis of biaryl compounds such as 23 and 24 was not easy, but these compounds can now be synthesized easily by the cross-coupling reaction\(^{17}\) using Pd metal.

5. Syntheses of naturally occurring 6-alkylindole

As shown in Table 2 and Fig. 9, 6-alkylindoles could be easily synthesized by F.I. Syntheses of 6-substituted indoles, 6-(3-methylbuta-1,3-dienyl)in-
dole (25)\(^{18}\) and 6-(3-methyl-2-butenyl)indole (26)\(^{19}\) have been accomplished using this method.\(^{16},20\) An outline of these reactions is shown in Figs. 10 and 11.

As described in Fig. 10, the compound 22 shown in Table 2 was treated with KOH/EtOH, followed by treatment with pyridine N-oxide/Ac\(_2\)O\(^{21}\) to give the aldehyde 28. Aldol condensation of 28 with acetone gave 29, whose ester group was removed by decarboxylation to give 30. Wittig reaction of 30 finally gave the natural product 25. Another natural product 26 was synthesized by a synthetic route similar to that used for 25 (Fig. 11).

6. Proof for the first cyclization step of F.I.
by a synthetic technique

As for the mechanism (shown in Fig. 5) of the F.I., it was pointed out from the early stages of its discovery that the first cyclization step resembles the Claisen rearrangement reaction, which is known as a typical sigmatropy reaction. In addition, the complete reaction mechanism became clear afterwards, because various intermediates were caught up in various stages of the F.I.\(^3\) However, it was the electron movement in the first cyclization step, which is shown as 35 or 35’ in Fig. 12, that remained a problem until quite recently. The electron movement 35 was assumed on the basis of the nucleophilicity of the benzene nucleus, but 35’ was assumed based on the high nucleophilicity of the enamine part. Electron movement 35 was more reasonable, but the experimental evidence for this movement was lacking.

Therefore we carried out the following experiments to confirm it by the synthetic method. As shown in Fig. 13, F.I. of asymmetric diphenylhydrazone 36 would give indolic products 37 and 38 in a
ratio depending on the electronic effect of substituent X. The results of the experiment are shown in Fig. 14.23)

In the F.I. of diphenylhydrazone (39) having an electron donorative methoxy group at the para position of one of the two benzene nuclei, the indole (40) formed in the cyclization toward the methoxy-substituted benzene was obtained in a larger excess amount than the product (41) formed by the cyclization at the other side. We will discuss the non-indolic product (42) later. On the other hand, the F.I. of diphenylhydrazone (43) having an electron-attractive ethoxycarbonyl at the para-position on one of the two benzene rings mainly occurred in the unsubstituted benzene. In addition, in the case of diphenylhydrazone (46) having two methoxy groups at meta-positions (3,5-dimethoxy), the cyclization to the substituted benzene ring had priority over the cyclization to the unsubstituted benzene. In this case, although there are two methoxy groups at the meta-position which activated the cyclization side, it is mysterious that the ratio of the yield of product (47) to the yield of (48) was small. However, a possible explanation for this result is that the F.I. of m-substituted phenylhydrazone gives23) a larger amount of the p-cyclized indole than the o-cyclized one, because the m-substituent sterically hinders the cyclization. From these results, it was shown that F.I. proceeded readily toward the electron-rich benzene ring (generally an aromatic ring). Therefore, it was proved that the electronic movement of 35 in Fig. 12 had priority over that of 35'.

Next, we carried out the F.I. of other diphenylhydrazone systems, as shown in Fig. 15, to examine the fate of the methoxy group in the ortho-methoxyphenylhydrazones.24)

As in the case of the F.I. of 39 in Fig. 14, the formation rate of indoles was low in both ortho-methoxyphenylhydrazones (49a, b). In the case of dimethoxyphenylhydrazone (49a), the reaction pro-
ceeded in the direction of the electron-rich ring as in the case of NH-hydrazone (12) in Fig. 7. However, the F.I. of monomethoxyphenylhydrazone (49b) gave unexpected products. In other words, the cyclization to an electron-deficient ring took first priority in the case of 49b. This result was not expected based on the results of F.I. of the other ortho-methoxyphenylhydrazones (5, 39, and 49a). We interpreted it as follows: In view of the results, the conformation (49b-2) should have been given priority, although it had been thought that the conformations of phenylhydrazone that took part in the cyclization were 49b-1 and 49b-2. In this case we thought that 49b-3, an enehydrazine tautomer of 49b-2, was stable by hydrogen bonding, and that is the reason why 49b-2 was predominant.

In order to confirm the problem of the conformation fixed with the methoxy group in the enehydrazine structure shown in Fig. 15, the F.I. of 2,2'-dimethoxydiphenylhydrazones (52) (Fig. 16) was conducted.

As a result (Fig. 16), there were only a few indolic products, of which only 53 was obtained in a non-negligible quantity (4%). On the other hand, the non-indolic by-product (described in Table 3) was obtained in 30% yield, and the significance of this result will be explained later. The indole (53) should have been provided by the cyclization at the vacant ortho-position of 52 in the conformation 55a but not 55b. It is thought that this conformation (55a) was more unstable than 55b according to the steric hindrance of the methoxy group. This reaction did not generate the chloroindole (54) at all, although 54 should have been permitted by the cyclization at the methoxy side in the conformation (55b). This result was not expected based on our previous result, and thus this fact shows the difficulty of presuming the conformation at the cyclization. In some cases, the F.I. was found to proceed with difficulty due to the simultaneous presence of a substituent in the N1 (the nitrogen connected directly with the benzene ring) and the ortho-position of the phenylhydrazone as in this experiment, as shown in the following example (Fig. 17) reported in our previous study.

Thus, the F.I. of the 2,6-dichloro-NH-hydrazone (56) reacted smoothly and gave 5,7-dichloroindole (57) at a high yield. On the other hand, the F.I. of the corresponding NCH3 hydrazone (58) did not proceed readily and gave only a curious compound formed by the rearrangement of the methyl group at the 1-position to the 3-position in very low yield with high recovery of the starting material. This result was probably also due to the fact that the phenylhydrazone (58) cannot be converted to the suitable conformation necessary for the F.I. cyclization by the presence of the substituent at N1 (the 1-position).
7. Self-decomposition of methoxy-diphenylhydrazones

As shown in Figs. 14, 15, and 16, F.I. of diphenylhydrazones carrying a methoxy group at the o- or p-position (39, 49a, 49b, 52) of the N atom yielded comparatively poor yields of indole products. In fact, these reactions yielded a large amount of chlorinated diphenylamines other than indolic products, as shown in Table 3.22,24 These diphenylhy-
drazones, which have methoxy group(s) at the α- or p-position from the N atom, should have given the chlorinated diphenylamines by self-degradation before F.I. cyclization. The position of the introduced chlorine atom was the meta-position of the methoxy group. On the other hand, the diphenylhydrazones that had methoxy groups at the m-position of the N atom (46) and had no methoxy group (43) did not yield such chlorinated diphenylamines.

Based on the facts described above, a mechanism for the abnormal formation of chlorinated diphenylamines was proposed as shown in Fig. 18. Because the formation mechanism of every chlorodiphenylamine was fundamentally the same, we can explain the mechanism in the reactions of diphenylhydrazones 39 and 49b as shown in Fig. 18. As can be seen in the figure, the diphenylhydrazones were self-decomposed before proceeding to F.I. due to the resonance effect of the methoxy group.

The results described above led to another consideration. That is, we considered that the decomposition of the diphenylhydrazones would be possible with the general NH-phenylhydrazone having a methoxy group, although such an example has not been reported yet. Sometimes, the F.I. of the methoxyphenylhydrazone may unexpectedly fail to give the corresponding indole in good yield, though the starting materials seem to disappear early in the reaction. Indeed, the total yield of the indoles of the F.I. of phenylhydrazones 5 and 12 did not exceed 45%. This would have been the reason that a decomposition similar to that described in Fig. 18 occurred, even though the corresponding chlorinated aniline did not become trapped.
Recently, a computational calculation was reported\textsuperscript{26} to explain the failure of F.I. to synthesize 3-aminoindoles, during which attempt heterolytic N-N bond cleavage occurred without cyclization due to the substituent effect. This calculation method can be applied to our self-decomposition reaction shown in Table 3 for the mechanistic consideration.

8. The F.I. of 2-substituted phenylhydrazones having substituents other than a methoxy group

As described at the start of this article, in the F.I. of the 2-methoxyphenylhydrazone (5), the cyclization occurred on the methoxy side of the phenylhydrazone to give interesting indole derivatives. Thus, we were also interested in examining the F.I. of phenylhydrazones having other substituents at the 2-position.\textsuperscript{27} The results of these examinations are shown in Table 4.

We obtained several results more unusual than those observed for F.I. of the 2-methoxyphenylhydrazone (5), namely: 1) the F.I. of these phenylhydrazones (70) with HCl/EtOH generally proceeded with difficulty, except in the case of 2-methylthiophenylhydrazone (70b); 2) no substitution in cyclization occurred in any case; 3) the abnormal indolic products were formed only by rearrangement of the 2-substituent of phenylhydrazone; 4) abnormal indolic products were formed only by rearrangement in phenylhydrazones having an ortho-para-orienting substituent, and the position of rearrangement differed according to the substituent; and 5) phenylhydrazones with a stronger electron-attractive substituent (nitro or trifluoromethyl group) gave only a normal indolic product (as a result of the cyclization toward the vacant ortho-position).

9. Synthesis of 7-oxygenated indoles by F.I.

Based on all the above results, F.I. clearly does not realize the efficient synthesis of 7-methoxyindole. However, there is a high demand for the convenient synthesis of 7-oxygenated indole. To resolve this problem, we devised the following synthetic plan based on the results in Table 4. Considering that an abnormal indole product was not formed at all with phenylhydrazones having an NO\textsubscript{2} or a CF\textsubscript{3} group, the F.I. of the phenylhydrazone (73) having an electron-poor oxygen functional group was examined. For electron-poor oxygen functional groups, the tosyl-, mesyl-, and trifluoromesyl groups were chosen in order to convert these into a phenolic or alkoxy group easily later.\textsuperscript{28} The results are shown in Table 5.

PPA and TsOH/benzene were chosen as acid catalysts, since these are strong acids and also poor nucleophiles. The reactions proceeded under the conditions shown in Table 5 and generally gave two kinds of indolic products, namely the normally cyclized product, 7-substituted indole (74), and the rearranged product, 5-substituted indole (75). As expected, however, considerably greater amounts of 7-substituted indoles were formed compared to 5-substituted indoles.
Next, the F.I. of other types of phenylhydrazone was examined as shown in Table 6. Although the overall yield varied, the formation ratio of the rearranged product (78) was low in every case, and thus it is certain that this method will practically give 7-substituted indole (77) in reasonable yield. Heathcock and others\(^{29}\) reported a method for preparing 7-oxygenated indole in which cyclization does not occur physically in the 2-substituent side in F.I. (Fig. 19). Though their method is an excellent one from the viewpoint of yield and positional specificity, the synthesis route is longer. When synthetic convenience is taken into consideration, our method may be better overall.
10. Application of the synthetic method to the syntheses of naturally occurring 7-oxygenated indoles

The above-mentioned synthetic method of 7-oxygenated indole via F.I. was applied to the syntheses of related natural products as follows: The first target was Eudistomidin-A (79), which was isolated from a type of sea sponge in Okinawa, Japan, and the outline is shown in Fig. 20.

The F.I. of 2-tosyloxyphenylhydrazone (80) prepared from 5-bromo-2-aminophenol gave the expected indole (81) as a sole product. After F.I. product (81) had been led to 82 with decarboxylation, the first synthesis of 79 was achieved via 83 based on the method of Hino et al.31) The final detosylation was achieved by the hydrolysis with KOH.

As the next application, Murrayafoline-A (84) and Murrayaquinone-A (85) isolated from a plant were synthesized (Fig. 21). The F.I. of the phenylhydrazone (86) prepared from 5-methyl-2-aminophenol gave the target indole (87) in good yield with a minute amount of the abnormal product (88). The aromatization of 87, followed by hydrolysis, gave the phenolic compound (89). Methylation of 89 gave the natural product (84), and oxidation of 89 with Freny’s salt gave the quinone natural product (85).

11. Role of the enehydrazine part in the F.I. mechanism at the first cyclization step

As described above, we established that the first cyclization step of the F.I. was promoted by substituting the electron-donating substituent on the benzene ring of phenylhydrazone (the opposite is also
true). As a result, the movement of the electron is confirmed as to be 35 but not 35’ in Fig. 12. On the other hand, the electronic effect in the enehydrazine part has been hardly understood until recently.

In the course of these experiments,33) we noticed that the phenylhydrazone prepared from phenylhydrazine (90) and β-diketone (91) was actually in the enehydrazine structure (92) (Fig. 22). The enehydrazine part of 92 seems to have been rendered electron-poor by the carbonyl group. In these compounds, the F.I. of the phenylhydrazone (92, X=OMe) easily gave the corresponding indole product in 43% yield without an acid catalyst. Naito et al.34) reported that the compound (95) whose N atom of the enehydrazine part of the phenylhydrazone (94) was acylated easily was cyclized only by heating to give the corresponding indole (96) (Fig. 22, eq. 2). They explained this result as follows: in this reaction the reactivity of the enehydrazine part was promoted by the enhancement of the LUMO character by the strong electro-attractivity of the trifluoroacetyl group, because in the [3,3]sigmatropic reaction exemplified by Claisen rearrangement, the reaction between two double bonds is well-known to proceed by an appropriate combination of HOMO and LUMO. This reaction showed quite similar reactivity with ours (eq. 1). Naito et al. reported in their series of studies35) that F.I. of α-substituted phenylhydrazone yielded an interesting compound corresponding to the intermediate c in Fig. 5. In these cases, the F.I. of the phenylhydrazone having a more electron-donative substituent at the ortho-position gave the higher yield of the compound which was formed by the cyclization to the substituted ortho-position, as described in Table 4.

On the other hand, we obtained the result that the F.I. proceeded more rapidly if the electronic density of the benzene part of phenylhydrazone increased by the substituent, as described earlier. Therefore, with respect to the first cyclization step of the F.I., it was verified that the electronic movement corresponded to that in 35, but not that in 35’, in Fig. 12.

Moreover, we discovered that the [3,3]sigmatropic reaction might not take place when there is a leaving group in the electron-poor benzene ring of phenylhydrazone (97),36) as shown in eq. 3 in Fig. 22. In this case, the dihydrocinoline derivative (99) was formed in addition to the expected indole (98). The cinoline (99) was surely formed, based on the result that the enehydrazine part attacked the root of the MsO group in a nucleophilic manner, followed by an aromatic addition-elimination reaction but not cyclization, as shown by the intermediate 100. Therefore, this can be considered one of the experiments which prove that the movement of the electron in the first cyclization step in F.I. is shown by 35, but not by 35’.

Fig. 22. The role of the enehydrazine part in F.I.
12. Consideration of F.I. on synthetic chemistry

The experimental methodology of F.I. of the usual phenylhydrazine that has been carried out in acidic media by heating has been proven correct. Because it is a [3,3]sigmatropy reaction, heating is necessary. Although the F.I. can progress even if the acid is not used, the reaction that uses the acid will give the greater result. It is thought that this is because the proton promotes the isomerization from phenylhydrazine to enehydrazine and improves the LUMO character in the part of enehydrazine that promotes the reaction. The electron-donative substituent on the phenylhydrazine makes the F.I. proceed easily, but F.I. of the phenylhydrazine having a methoxy group may cause an abnormal reaction to give the corresponding indole in poor yield.

Recently, a new method for the F.I. was reported\(^{(67)}\) using microwaves instead of a conventional heating procedure. The significant features of this method include its short reaction time and good yield.

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Profile

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