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## Recent developments in chemical reactivity of N,Ndimethylenamino ketones as synthons for various heterocycles

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The current review presents recent progress in the utility of N,N-dimethyl enaminones as building blocks for a diverse range of acyclic, carbocyclic, five- and six-membered heterocyclic a broad range of heterocyclic and fused heterocyclic derivatives. Most importantly, these N,N-dimethyl analogues have proven to be of biological interest and provide an access to new class of biologically active heterocyclic compounds for biomedical applications. All of these topics are drawn from the recent literature till 2016.

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#### Introduction

β-Aminovinyl ketones are versatile synthetic intermediates that combine the ambident nucleophilicity of enamines with the ambident electrophilicity of enones. As can be rationalized, these systems have "enamine" character, and can act as building blocks for the synthesis of various heterocycles such as pyridine, pyrimidine, and pyrrole derivatives.1 In addition, they also have

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"enone" character, and may act as acceptors in both 1,2- and 1,4additions. In this way, β-aminovinyl ketones serve as scaffolds for annulation, and can provide access to systems such as pyrroles, indolizidines, quinolizidines, and perhydroindoles, all of which are common motifs in alkaloid structures.1,2

N,N-Dimethyl derivatives of β-aminovinyl ketones are chemical compounds consisting of an amino group linked through a carbon-carbon double bond to a keto group.6 They are typical push-pull ethylenes in which the amine group pushes and the carbonyl pulls electron density. The chemistry of the enamino carbonyl group (1) is potentially an area of considerable scope when one considers that there are present in this moiety two electron-deficient centres (i.e. two electrophilic sites) at C-1 and C-3, while the C-2, carbonyl oxygen and amino functions are electron



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field of nucleotide/oligonucleotide chemistry with support from ICSC-World Laboratory grant in Switzerland; moreover, a Research Fellowship in the UK at Cardiff University in Cardiff for the period of three years (2009–2009), working with Professor Dr M. Bagley on synthetic approaches towards new anti-cancer and antibiotic agents in addition to the synthesis and properties of anticonvulsant enaminones. In 2012, Dr Gaber was appointed Head of the Organic Chemistry Department at NODCAR, then Head of the General Pharmaceutical Chemistry Division (Dean) in 2014. His research interests include heterocyclic and medicinal chemistry with emphasis on the design and synthesis of novel bioactive heterocycles and the study of their biomedical applications with the aim of developing new therapeutic agents.

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rich (*i.e.* three nucleophilic sites) (Fig. 1).<sup>3,4</sup> They can thus function both as nucleophiles and as electrophiles, their versatility in either case being extended by their ability to show ambident reactivity.<sup>2</sup> Accordingly, these  $\beta$ -acyl enamines proved to be suitable for a number of further synthetic transformations, mainly; hydrolysis to diones, reduction and condensation to fused heterocycles.

A few review articles, describing the chemistry of β-aminovinyl ketones, have appeared in the literature, 1,3,5,6 although hitherto with little emphasis on the reactivity of the corresponding N,Ndimethyl derivatives of  $\beta$ -aminovinyl ketones. Since a large number of developments in the use of these N,N-dimethyl derivatives in heterocyclic and medicinal chemistry have been reported recently, and in conjunction with our long-term continuing interest in exploring the synthetic applications of biologically active enamino compounds, 7-19 the current review presents firstly recent progress in the utility of N,N-dimethyl enaminones as building blocks for a diverse range of acyclic, carbocyclic, five- and six-membered heterocyclic as well as condensed heterocyclic compounds, with particular emphasis on the useful chemical transformations of this class of N,N-disubstituted enaminones to the structurally related  $\alpha$ -(arylhydrazono)-β-ketoaldehydes. Structural investigation for these aldehydes and some of other products has also been made. This is finally followed by a consideration of the ever-increasing



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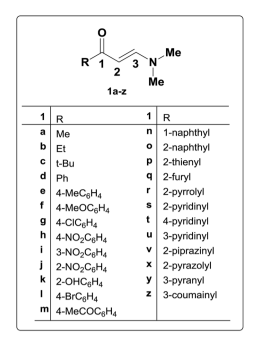


Fig. 1 General formula of N,N-dimethyl enaminone derivatives

chemotherapeutic potentials of these *N*,*N*-dimethyl derivatives and their biomedical applications as valuable synthons on the way to a variety of bioactive heterocyclic compounds. All of these topics are drawn from the recent literature till 2016.

# 2. Synthetic applications of *N*,*N*-dimethylenamino ketones

#### 2.1. Preparation of acyclic compounds

2.1.1. Preparation of  $\alpha$ -(arylhydrazono)- $\beta$ -ketoaldehydes. One of the main route to  $\alpha$ -(arylhydrazono)- $\beta$ -ketoaldehydes is the coupling of arenediazonium salts with aminovinyl ketones. <sup>20–23</sup> However, anomalous behavior has been reported



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Prof. A. S. Shawali in 1996 and he has published 106 scientific papers and reviews all in international journals in the fields of physical organic chemistry, chemistry of hydrazonoyl halides and bioactive heterocyclic chemistry (there are about 755 citations of his work from 2000 until February 2017 (h-index 16).

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for reactions of cyclic aminovinyl ketones and aminovinyl esters with arenediazonium salts.24,25 The structure and chemistry of these \beta-ketoaldehydes has attracted much attention with plenty of uses for these compounds, in heterocyclic synthesis, being reported recently.26-31 It has been noted that in DMSO these molecules exist as a mixture of two conformers E and Z. <sup>22,27,28</sup>

Coupling<sup>21,22,32-34</sup> of diazotized aromatic amines with the aminovinyl ketones 1d,f,g,h and 1q resulted in products of coupling and hydrolysis of the dimethylamine moiety. These products were shown, based on <sup>1</sup>H NMR and <sup>13</sup>C NMR, to exist as a mixture of anti and syn hydrazones with the anti form always prevailing.<sup>22</sup> It is assumed that initially formed arylazo intermediates 2a-k are hydrolyzed, by the action of the aqueous base existing in the medium, into the arylhydrazonoaldehydes 3a-k (Scheme 1).

In addition, 3-(thiophen-2-vl)- or 3-aryl-propional dehydes 4a-k were prepared via coupling of aminovinyl ketones 1p and 1e,g,h with aromatic diazonium salts.28 It is believed that assumed fixation of hydrazones by hydrogen bonding in DMSO solution is least likely although it was observed in almost every case of aroyl derivatives the presence of two NH signals integrating for a total of one proton with varying intensity in each case depending upon the nature of the substituent at the carbonyl moiety (Scheme 2). This may point to the existence of an equilibrium between the two forms 4C and 4D in DMSO solution.28

Furthermore, it was found that diazotized anthranilonitrile (5a) or diazotized methyl anthranilate (5b) couples readily with the aminovinyl ketones 1a, 1d, 1p-s to yield products of coupling and hydrolysis of dimethylamino moiety. The coupling products can thus be formulated as the hydrazone forms 8A,B or potential tautomeric enol azo forms 7,8C or a mixture of one or more of these forms. The hydrazone structure 8A,B was established for all these products and the E-form 8A is believed to be the predominant form except for the product of coupling 1-methyl-3-dimethylaminoprop-2-enone (1a) with diazotized anthranilonitrile (5a) (Scheme 3). <sup>1</sup>H NMR of this product show that it exists in DMSO as an equilibrium mixture of E-form 8A, Z-form 8B and the enol azo form 8C, as <sup>1</sup>H NMR revealed three signals for a total of one proton, and it is

Scheme 1

O ArN=NCI R CHO

N NH

1e, R = 2-Thienyl
1h, R = 
$$C_6H_4Me-p$$
1g, R =  $C_6H_4NO_2-p$ 
1o, R =  $C_6H_4NO_2-p$ 

b, R = 2-Thienyl Ar = Ph
b, R = 2-Thienyl Ar =  $C_6H_4Me-p$ 
c, R = 2-Thienyl Ar =  $C_6H_4Me-p$ 
d, R = 2-Thienyl Ar =  $C_6H_4Me-p$ 
i, R =  $C_6H_4Me-p$  Ar =  $C_6H_4Me-p$ 
d, R = 2-Thienyl Ar =  $C_6H_4Me-p$ 
i, R =  $C_6H_4Me-p$  Ar =  $C_6H_4Me-p$ 
d, R = 2-Thienyl Ar =  $C_6H_4Me-p$ 
i, R =  $C_6H_4Me-p$  Ar =  $C_6H_4Me-p$ 
i, R =  $C_6H_4Me-p$  Ar =  $C_6H_4Me-p$ 
i, R =  $C_6H_4Me-p$  Ar =  $C_6H_4OMe-p$ 
j, R =  $C_6H_4Cl-p$  Ar =  $C_6H_4Cl-p$ 
i, R =  $C_6H_4Cl-p$  Ar =  $C_6H_4Cl-p$ 
ii. R =  $C_6H_4Cl-p$  Ar =  $C_6H_4Cl-p$ 

Scheme 2

interesting to note that the E-form was the major constituent in this equilibrium mixture (70%). All these forms are stabilized by hydrogen bonding.27,35

Similarly, coupling of aminovinyl ketones either 1d,p with benzonitrile diazonium chloride (9), following recently reported procedure,21 produced the corresponding acyclic β-ketoaldehydes **10a,b** (Scheme 4).36

Also, 3-oxo-2-(arylhydrazono)pentanal 11a-h were obtained in 50-80% yields via coupling N,N-dimethyl enaminones either **1b** or **1m** with aromatic diazonium salts. <sup>1</sup>H NMR of the products indicated that they exist at least in DMSO solution as mixtures of the anti-form 11 and syn-form 12 (Scheme 5). The anti-from generally predominated.29

The diazotized 2-aminocyclohexenethiophene (13) coupled readily with N,N-dimethylenamino ketones 1a,d,p-r to yield the corresponding hydrazonopropanals 14a-e, respectively (Scheme 6).27

It has been found that 5-methylisoxazole-3-diazonium chloride (15a) coupled readily with  $\beta$ -acyl enamines 1q,u to produce the expected β-ketoaldehydes 16a,b.33 However, a similar treatment of ketoenamines 1d and 1g resulted in the cyclization into pyrazolo[5,1-c][1,2,4]triazines 18a,b via the assumed intermediacy of acyclic aldehydes 17a,b (Scheme 7).33

Elnagdi et al. 37 reported on the reaction of 1-dimethylamino-5-arylpenta-1,4-diene-3-one 19a-c with diazonium ions in aqueous ethanol, where the dimethylamino group was substituted by the hydroxyl group and thus the corresponding 5aryl-2-arylhydrazono-3-oxopent-4-enals 20a-c were isolated. Compounds 20a-c were presumably formed by hydrolysis of the enamine primary products. The <sup>1</sup>H NMR indicated the presence of a mixture of both E and Z forms of these pentenals in approximately equivalent ratios (Scheme 8).37

Also, enaminophthalimidoacetone derivative 21 readily underwent coupling with aromatic diazonium salts in the presence of sodium hydroxide to yield the analogous aldehydes 22a-d in acceptable yields (Scheme 9).38

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X = CN

X = CN

Scheme 3

f, R = 2-Pyrrolyl

g, R = 2-Pyridyl

Compounds 24a,b were obtained in the same way as aldehydes 22 where the N,N-dimethylenamino ketone 21 coupled with heterocyclic diazonium salts 23a,b to give rise to the corresponding hydrazones 24a,b in good yields (Scheme 10).38

Scheme 5

As recently reported by our research group,<sup>7</sup> pyrazole enaminone derivative 1v also coupled with p-chlorobenzenediazonium chloride to provide the pyrazoloylhydrazone derivative 25, in 82% yield (Scheme 11).

Trials to couple enaminone of the coumarin 1z with benzenediazonium chloride failed. However, p-nitrobenzenediazonium chloride coupled readily with 1z to yield the respective arylhydrazonopropanals that proved to exist as an equilibrium mixture of the anti-form 26A and the syn-form 26B (Scheme 12).25

An interesting synthesis of arylhydrazonals is the formation of 27a-e<sup>39</sup> on coupling of enaminal 27a or enaminoester 27b with various aromatic diazonium salts. Compounds 28a-e were assigned the indicated hydrazone structure in preference to a potentially tautomeric enolazo structure. This assignment is based on <sup>1</sup>H NMR and <sup>13</sup>C NMR spectra which revealed signals for two formyl protons and carbons in 28a-d and one such signal in the spectra of 28e (Scheme 13).40

On the other hand, when dimethylaminomethylene derivative 29 was coupled with benzenediazonium chloride, the phenylhydrazone 32 was formed. It is believed that, as a result of nitrogen lone pair donation to enamine β-carbon, the latter becomes sufficiently nucleophilic and intermediate diazonium salt 30 is initially formed. This then readily hydrolyses into the azo derivative 31 that then undergoes Japp-Klingemann cleavage to yield the final isolated product 32 (Scheme 14).41

2.1.2. Preparation of N-monosubstituted enaminones and related compounds. Dimethylaminomethylidene derivatives 1d and 1q reacted with aniline in ethanol at reflux for three hours to yield products of addition and dimethylamine elimination.

Scheme 6

Scheme 7

NMe<sub>2</sub>

$$\begin{array}{c|c}
\bullet & \bigcirc \\
PhN_2CI & R
\end{array}$$

$$\begin{array}{c|c}
\bullet & \bigcirc \\
N & \\
N & \\
NH \\
Ph
\end{array}$$

$$\begin{array}{c|c}
\bullet & \bigcirc \\
N & \\
NH \\
Ph
\end{array}$$

$$\begin{array}{c|c}
\bullet & \bigcirc \\
N & \\
NH \\
Ph
\end{array}$$

$$\begin{array}{c|c}
\bullet & \bigcirc \\
\bullet & \\$$

a, R = Ph 20 FII  
b, R = 
$$C_6H_4OMe-p$$
  
c, R = 2-Furyl  
Scheme 8

Scheme 9

Scheme 12

a, R = H; X = S; Y = CH; Z = N**b**, R = Me; X = NH; Y = N; Z = CH

Scheme 10

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The formed anilino derivatives proved to be *cis* products **34a,b** and the *trans*-form **33a,b** was completely excluded, since olefinic protons appeared, in each case, as a doublet with J = 7 Hz typical for such *cis* olefinic protons. The predominance of the *cis*-form is attributed to fixation through hydrogen bonding. <sup>42</sup> On the other hand, compound **1d** reacted with *o*-phenylenediamine (**35**) to afford a product that may be

formulated as *cis* enaminone **39a** or stereoisomeric **38** or cyclic products **36** or **37**. Structure **39a** was established through <sup>1</sup>H NMR and <sup>13</sup>C NMR spectra (Scheme 15).<sup>42,43</sup>

In accordance with the previous observation,  $^{42}$   $\beta$ -aminovinyl ketones **1e**,**f** reacted with o-phenylenediamine (**36**) to afford products of condensation via dimethylamine elimination. These were assigned cis structure **41** rather than trans structure **40** based on  $^{1}$ H NMR, which revealed signals for cis olefinic protons in their proper positions with J=9 Hz. The predominance of this form may be due to fixation by hydrogen bonding.  $^{43}$  In contrast to this observation, it has been found that a similar treatment of ketones **1d-f** with benzotriazole (**42**), instead of o-phenylenediamine (**36**), led to the formation of trans enaminones **43**, which were also obtained from the diazotization of cis enaminones **41** with sodium nitrite in acetic acid under stirring at room temperature (Scheme **16**).  $^{43}$ 

On the other hand, tertiary enaminone **1y** reacted with methylamine, ethylamine and aniline in acetic acid under stirring at room temperature to yield the corresponding secondary enaminones, which were found to exist, in each case, as a mixture of the *E*- and *Z*-forms **44a–c** and **45a–c**. The existence of *Z*-form for these compounds is in contrast to the observed predominance of *E*-form for the starting **1y**. This is attributed to the stabilization of *Z*-form for those compounds through hydrogen bonding. Interestingly, when compound **1y** was similarly treated with 2-aminothiazole (**46**) in acetic acid under stirring for three hours at room temperature, only the acyclic *Z*-enaminone **47** was formed in exclusively based on considering *J* value for olefinic protons (J = 9 Hz). While bicyclic pyridine derivative **48** was isolated upon heating of **1y** with **46** under reflux in acetic acid for two hours (Scheme **17**).

Tertiary amines **49a,b** were converted into the corresponding secondary amines **50a,b** when heated at reflux with equivalent amount of aniline in ethanol for two hours in yields of 53% and 37%, respectively.<sup>45</sup> Also, it has been found that a much better

yield (71%) of product **50a** could be obtained on microwave heating of **49** with equimolar amount of aniline in a domestic microwave oven at full power for two minutes.<sup>45</sup> On the other hand, microwave heating of **49** with ammonium acetate for two minutes in a domestic microwave oven at full power produced the primary amine **51** in yield of 91%. The authors<sup>46</sup> failed to

obtain 51 by conventional heating with ammonium acetate in acetic acid (Scheme 18).

Scheme 19

Dimethylaminomethylene thiazolones **52a,b** reacted with equivalent amounts of aromatic amines with reflux in acetic acid for one hour to yield the corresponding secondary amines **53a-e**, respectively.<sup>47</sup> Similar treatment with piperidine in refluxing ethanol for seven hours afforded the piperidino derivatives **54a,b**, which were also obtained on reacting thiazolinones **55a,b** directly with triethyl orthoformate and piperidine in DMF.<sup>47</sup> Utilization of triethyl orthoformate and piperidine in DMF solution was found more economic and safer than using DMFDMA. It is believed that piperidine reacts with acetal forming non-isolable intermediate **56** which then condensed with **52** to give the final isolable products as depicted in Scheme **19**.<sup>47,48</sup>

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60a, X = CN **b**,  $X = CO_2Me$ NMe<sub>2</sub> 1d,g,j,p 1d, R = Ph  $\mathbf{g}$ ,  $R = C_6H_4CI-p$ 62a-f j, R = C<sub>6</sub>H<sub>4</sub>NO<sub>2</sub>-op, R = 2-Thienyl 61,62a, X = CN; R = Ph  $\mathbf{d}$ ,  $X = CO_2Me$ ; R = Ph**b,** X = CN; R =  $C_6H_4CI-p$ **e,**  $X = CO_2Me$ ;  $R = C_6H_4Cl-p$ q, R = 2-Furyl c, X = CN; R = 2-Thienyl  $f_1 X = CO_2Me_1 R = C_6H_4NO_2-0$ 64a, X = CN **b**,  $X = CO_2Et$ 1d,g,j,q 66а-е 65а-е 65,66a, X = CN: R = Ph **b**, X = CN;  $R = C_6H_4CI-p$ MW, 5 min **c,** X = CN;  $R = C_6H_4NO_2-o$ **d,**  $X = CO_2Et$ ;  $R = C_6H_4Cl-p$ N

Scheme 21

Table 1 Relative concentration of cis- and trans-forms of enaminones 61, 65 as indicated from <sup>1</sup>H NMR

e,  $X = CO_2Et$ ; R = 2-Furyl

| Compounds | cis-Form | trans-Form |
|-----------|----------|------------|
| 61b       | 7        | 1          |
| 61e       | 4        | 1          |
| 61f       | 4.2      | 1          |
| 65a       | 6.5      | 1          |
| 65c       | 6        | 1          |
| 65        | 4        | 1          |

Dimethylaminomethylene thiazolone 57 could be converted into the enamine 58 on treatment with p-toluidine in refluxing acetic acid for one hour.47 Nucleophilic displacement of the active dimethylamino group of 57 by cycloaliphatic amines, in refluxing ethanol for seven hours, resulted in the formation of the corresponding derivatives of piperidino and morpholino 59a,b, respectively. Structure 59b was confirmed by X-ray crystal determination (Scheme 20).47

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$$\begin{array}{c} \textbf{1d}, R = Ph \\ \textbf{e}, R = C_6H_4Me-p \\ \textbf{g}, R = C_6H_4Cl-p \\ \textbf{j}, R = R = C_6H_4NO_2-o \\ \textbf{x}, R = 1-(4-\text{chlorophenyl})-3-\text{cyano-}1H-\text{pyrazol-}5-\text{yl} \\ \end{array}$$

$$\begin{array}{c} \textbf{75a}, X = CH_2; R = Ph \\ \textbf{b}, X = CH_2; R = C_6H_4Cl-p \\ \textbf{c}, X = CH_2; R = C_6H_4NO_2-o \\ \textbf{d}, X = CH_2; R = C_6H_4Cl-p \\ \textbf{f}, X = CH_2; R = C_6H_4Cl-p \\ \textbf{f}, X = CH_2; R = C_6H_4NO_2-o \\ \textbf{d}, X = CH_2; R = C_6H_4NO_2-$$

Scheme 23

In addition, interaction of dimethylaminomethylidenes  $\mathbf{1d},\mathbf{g},\mathbf{j}$  and  $\mathbf{1p}$  with anthranilonitrile ( $\mathbf{60a}$ ) or methylanthranilate ( $\mathbf{60b}$ ) yielded the cis enaminones  $\mathbf{61a-f}$ , respectively (Scheme 21). Attempted cyclization of  $\mathbf{61a-c}$  into the corresponding quinolines failed. Successful cyclization of  $\mathbf{61d}$  into the quinolinone  $\mathbf{63}$  could be affected, on heating for five minutes in domestic microwave oven at full power. When compounds  $\mathbf{1d},\mathbf{g},\mathbf{j}$  and  $\mathbf{1q}$  were similarly treated with 2-amino-4,5,6,7-tetrahydrobenzo[b]thiophene-3-carbonitrile ( $\mathbf{64a}$ ) or its 3-carboxylate derivative  $\mathbf{64b}$ , the corresponding cis enaminones  $\mathbf{65a-e}$  were obtained. Again,  $^1$ H NMR indicated the predominance of the cis-form (c.f. Table 1). $^{49}$ 

Reacting enaminonitrile 67 with ethyl glycinate 68 in ethanol/potassium carbonate solution yielded the analogous

ethyl ester **69**, which could not be cyclized into the pyrrole derivative **70** under a variety of conditions. Also, compound **67** reacted with aminothiazole derivative **71** and with aminothiophene derivative **64b** to afford the secondary amines **72B** and **73B**, respectively. Although **72B**, **73B** may also exist as **72A** or **73A**, the hydrogen bonded form **72B** and **73B** seems more stable (Scheme 22). So

On the other hand, when compounds **1d,e,g,j,x** were similarly treated with piperidine (**74a**), only the *E*-enaminones **75a-d**, respectively, were formed in exclusively as indicated from the coupling constant values for olefinic doublets. No trace of *Z*-form was observed in this reaction. Analogously, compounds **75f,g** were formed from the reaction of **1d,x** with morpholine (**74b**) (Scheme 23).<sup>49,51-53</sup>

$$\begin{array}{c} \textbf{NMe_2} \\ \textbf{1e}, \textbf{R} = \textbf{C}_6 \textbf{H}_4 \textbf{Me-p} \\ \textbf{g}, \textbf{R} = \textbf{C}_6 \textbf{H}_4 \textbf{Cl-p} \\ \textbf{p}, \textbf{R} = \textbf{2-Thienyl} \end{array} \\ \textbf{82a}, \textbf{X} = \textbf{C-Ph}, \textbf{Y} = \textbf{C-Me} \\ \textbf{83,84a}, \textbf{X} = \textbf{C-Ph}; \textbf{Y} = \textbf{C-Me}; \textbf{R} = \textbf{C}_6 \textbf{H}_4 \textbf{Me-p} \\ \textbf{b}, \textbf{X} = \textbf{N-Ph}; \textbf{Y} = \textbf{C-Me}; \textbf{R} = \textbf{C}_6 \textbf{H}_4 \textbf{Me-p} \\ \textbf{c}, \textbf{X} = \textbf{C-Ph}; \textbf{Y} = \textbf{C-Me}; \textbf{R} = \textbf{C}_6 \textbf{H}_4 \textbf{Cl-p} \\ \textbf{c}, \textbf{X} = \textbf{C-Ph}; \textbf{Y} = \textbf{C-Me}; \textbf{R} = \textbf{C}_6 \textbf{H}_4 \textbf{Cl-p} \\ \textbf{c}, \textbf{X} = \textbf{C-Ph}; \textbf{Y} = \textbf{C-Me}; \textbf{R} = \textbf{2-Thienyl} \end{array} \\ \textbf{7N} \\ \textbf{N} \\ \textbf{N}$$

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AcOH/r.t.

AcOH/r.t.

AcOH/r.t.

AcOH/
$$\triangle$$

Beac a, R = Ph
b, R = 2-Thienyl
c, R = 2-Furyl

AcOH/ $\triangle$ 

Beac a, R = Ph
b, R = 2-Thienyl
c, R = 2-Furyl

Scheme 26

Interaction of 2-aminobenzothiazole (76) with aminovinyl ketones 1d and 1g provided the heteroaromatic aminoenones 77a,b. Similarly, compounds 1d and 1q,s were transformed to the corresponding acyclic aminoenones 80a-c by treatment with 3-amino-5-methylisoxazole (79) (Scheme 24). These products 77a,b and 80a-c are believed to exist in equilibrium with enols 78a,b and 81a-c, which are stabilized through hydrogen bonding.54

N,N-Dimethylenamino ketones 1e,g and 1p reacted with aminopyrazoles 82a,b to yield the acyclic enaminones 83a-f. The structure of which proved to exist in the cis-form based on

considering J values for olefinic protons (J = 9 Hz). 49 Compound 83a-f readily cyclized into the bicyclic pyrimidines 84a-f on reflux in pyridine solution in the presence of concentrated hydrochloric acid (Scheme 25).49

Treatment of N,N-dimethylamino derivatives 1d-f with thienocoumarin 85 in acetic acid at room temperature furnished products of addition and dimethylamine elimination in 80-85% yields. IR and <sup>1</sup>H MNR spectra of these products indicated involvement of the amino function in this reaction. Therefore, structure 86a-c was suggested for those products. On the other hand, when the reaction of 1d-f with 85 was

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Scheme 28

conducted in refluxing acetic acid, solid products **86a–c** were isolated by filtering the hot solution in 33–43% yields. When the mother liquor was left to stand at room temperature, the C-1 alkylation products **87a–c** were isolated in 32–36% yields. These products were found to be isomeric with **86a–c**. Both IR and  $^1H$  NMR indicated that the amino function was not involved into the reaction. Compounds **86a–c** rearranged into the corresponding  $\alpha,\beta$ -unsaturated ketones **87a–c** on prolonged boiling under reflux in dioxane solution in the presence of diethylamine or in pyridine in 60–70% yields. Better conversion yields could not be achieved under a variety of conditions. It is

thus believed that products **266** are kinetic products, while **71** are the thermodynamic ones. Conversion of **86** into **87** in basic media is believed to proceed *via* base addition across the double bond and elimination of **85**, thus allowing more of the thermodynamic product to be formed (Scheme **26**). <sup>55</sup>

An interesting reaction leading to *cis*-enaminone is the condensation of aminothienopyridazines (88a,b) with aminovinyl ketone 23 to yield products of condensation *via* dimethylamine elimination for which structures 89 or 90 seemed possible. Structure 90 was established for those products based on the <sup>1</sup>H NMR and IR spectral data that revealed involvement of amino function in the reaction. Typical for secondary enaminone, compounds 90a,b existed solely in *Z*-form as this

Scheme 29

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Scheme 31

Scheme 32

O NH<sub>2</sub>

CN

HN S 
$$CO_2Et$$

ab.  $EtOH/K_2CO_3/\Delta$ 

MeO

NMeO

NMeO

NMeO

ACOH/ $\Delta$ 

NMeO

ACOH/ $\Delta$ 

Scheme 34

form is fixed by hydrogen bond and this preferred over sterically and stereoelectronically fixed E-form.54 Further confirmation of this structure assignment was obtained via successful conversion of 90b into the thiophene 91 through hydrolysis of the alkylated amino moiety (Scheme 27).56

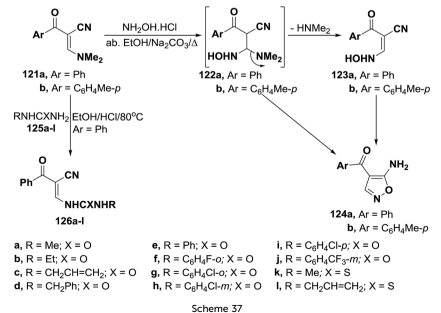
Dienone 19 underwent analogous reactions with aminothienopyridazines 88a,b in a microwave oven at 560 W for 90 seconds to give rise to the expected N-alkylated products 92a,b.56 A similar treatment of 6a with benzothienocoumarin 93 led to the corresponding enaminone 94 (Scheme 28).56

Aminothiophene derivatives 85, 88a,b and 93 reacted with the N,N-dimethylenamino ketones 95a,b in a similar fashion yielding the corresponding secondary enaminones 95a,b, 96ad and 97a,b (Scheme 29).56

In contrast to the behavior of aminothiophene toward N,Ndimethylenamino ketones, interaction<sup>55</sup> of compounds 1d,e with 88a in acetic acid at reflux for three hours gave C-1 alkylation products 99a,b instead of the N-alkylated derivatives 98a,b that would be expected by analogy with the other reports42,56 on the reactivity of nitrogen nucleophiles toward electron poor olefins (Scheme 30).

Similar to the behavior of 73a toward 1d,e enaminoketone 23 reacted with thienocoumarin 85 under microwave heating in the presence of few drops of acetic acid to produce only the C-1 alkylation product 100. This approach was also suitable for the preparation of another C-1 alkylation product 81, where reaction of 23 with benzothienocoumarin 92 under the same reaction conditions led to the corresponding  $\alpha,\beta$ -unsaturated ketone 101 (Scheme 31).<sup>56</sup>

Also, compound **85** reacted with aryl vinyl ketones **103a,b**, generated *in situ* from corresponding dimethylaminopropanone hydrochlorides **102a,b**, in acetic acid/ethanol mixture at reflux for three hours to yield 1:1 adducts. These were stable on reflux in protic solvents and thus possible formation of cycloadducts similar to **104** was ruled out (Scheme 32). Moreover, <sup>13</sup>C NMR of the reaction products indicated presence of only two sp<sup>3</sup> carbons while in cycloadducts **104** four such carbons should have been



Scheme 38

NMe<sub>2</sub> AcOH/NaOAc/ $\triangle$  S CHO
Ph
Ph
N
Me
131
132

observed. Therefore, the products for this reaction could be assigned structure  ${\bf 105.}^{55}$ 

It can thus be concluded that outcome of reactions of amines with electron poor olefins are dependent on nature of reagents used. Amino function, C-1 as well as the diene system are all possible sites of attack.<sup>56</sup>

2.1.3. Miscellaneous. Aminovinyl ketones 1a, 1d, 1p,q reacted with malononitrile or ethylcyanoacetate in ethanol and in the presence of a base, affording 1:1 adducts. These products could be formulated as pyran structure 107 or enamine structure 109. However, structure 109 was assigned on the basis of <sup>1</sup>H NMR spectra, in which two olefinic doublets with a *I* value of 13 Hz were observed indicating that the trans olefinic moiety has not been involved in the reaction and hence this excluded completely the possibility of pyran structure 107 for this reaction products. Consequently, formation of 109 is assumed to occur via the addition of the active methylene reagents to carbonyl groups in 1a, 1d or 1p,q and subsequent water elimination to furnish  $\alpha,\beta$ -unsaturated nitriles 108. Water eliminated in this process then hydrolyses the cyano group in the intermediates 108 affording the final isolated products 109.57,58 This structure was also confirmed by preparing the same reaction products 109c-e via condensation of cyanoacetamide with 1d or 1p,q under the same reaction conditions (Scheme 33).58

Interaction of enaminonitrile **67b** with ethyl thioglycolate in ethanol/potassium carbonate solution yielded the ethyl ester **110**. Trials to effect cyclization of **110** into **111** under a variety of conditions failed. In protic medium **110** decomposed and in aprotic medium **100** was recovered unreacted (Scheme 34).<sup>41</sup> Moreover, compound **109** reacted also with 1-methyl-6,7-dimethoxy-3,4-dihydroisoquinoline (**112**) in refluxing acetic acid to provide the respective dihydroisoquinolinylbutenenitrile derivative **113**. The authors<sup>50</sup> indicated that free donation around the single bond would

allow for a *trans*-form that should in theory experience less steric interaction.

Also, interaction of enaminocarbonyl compound **1p** with pyridazine-3-imine derivative **114** in refluxing ethanol containing a few drops of triethylamine led to the acyclic compound **115** *via* loss of a dimethylamine molecule, while on being heated in a mixture of aqueous acetic acid and hydrochloric acid these afforded the tricyclic product **116** (Scheme 35). The latter could be also obtained *via* cyclization of  $\alpha$ , $\beta$ -unsaturated compound **115** into **116** on boiling in aqueous acetic acid/hydrochloric acid mixture.<sup>59</sup>

Interestingly, reaction of *N*,*N*-dimethylamino derivatives **1d** and **1p** with phenylisothiocyanate in DMF in the presence of potassium hydroxide afforded after acidification with hydrochloric acid two products, the major of which were found to be of molecular formulae corresponding to structure **120** or its tautomeric forms **117–119**. However, thiol structure **120** better agreed with the obtained spectral data for these compounds, thus thione structures **117** and **118** could be ruled out. Structure **119** was also excluded on the basis of <sup>13</sup>C NMR spectra of the isolated products, which revealed the absence of any sp<sup>3</sup>-hybridized carbon atoms. These results strongly support structure **120** for the major products. The minor products for this reaction were assigned structure **34** based on the elemental analyses and spectral data of the isolated reaction products (Scheme **36**).<sup>60</sup>

Unexpectedly, α-(dimethylamino)methylene-β-oxo nitriles **121a,b** reacted with hydroxylamine hydrochloride in absolute ethanol in the presence of anhydrous sodium carbonate under reflux to give the acyclic hydroxylaminopropenonitriles **123a,b**, in excellent yields, based on elemental analyses and spectral data of the isolated reaction products. It is of value to report here that all trials to convert compounds **123a,b** into the corresponding isoxazoles **124** were unsuccessful. This can be attributed to the fact that compounds **123a,b** are mainly existing in the *anti*-form as indicated from their <sup>1</sup>H NMR spectra. <sup>61</sup>

NMe<sub>2</sub> Me<sub>2</sub>N -3 HNMe<sub>2</sub> AcOH/A 20a-c NMe<sub>2</sub> Me<sub>2</sub>N 1d, Ar = Ph e, Ar =  $C_6H_4Me-p$ 133 134 135  $\mathbf{f}$ , Ar = C<sub>6</sub>H<sub>4</sub>OMe-pa. Ar = Ph**b**, Ar =  $C_6H_4Me-p$ c, Ar =  $C_6H_4OMe-p$ 

Scheme 41

NMe<sub>2</sub> AcOH/
$$\Delta$$
NMe<sub>2</sub> AcOH/ $\Delta$ 
NMe<sub>2</sub> NMe<sub>2</sub> NMe<sub>2</sub>
NMe<sub>2</sub> AcOH/ $\Delta$ 
NMe<sub>2</sub> NMe<sub>2</sub> NMe<sub>2</sub> NMe<sub>2</sub> AcOH/ $\Delta$ 
NMe<sub>2</sub> NMe<sub>2</sub> NMe<sub>2</sub> NMe<sub>2</sub> NMe<sub>2</sub> AcOH/ $\Delta$ 
NMe<sub>2</sub> NMe<sub>2</sub>

On the other hand, reaction of 3-dimethylamino-2-benzoylpropenenitrile (**121a**) with *N*-substituted ureas or thioureas **125a–l** in acidic medium yielded ureidopropenenitriles **126a–l**, respectively (Scheme 37).<sup>62</sup>

Reacting enaminoketones 1d,g and 1p,q with hydroxylamine hydrochloride in ethanolic sodium acetate at room temperature resulted in the formation of products of condensation via dimethylamine elimination. These products were found to exist, in each case, as an equilibrium mixture of the hydroxylamino E- and E-forms (127 or 128). Assignment of these forms were based on their E-hydroxylamino hydroxylamino hydroxylamino

When *N*,*N*-dimethyl enaminone **1h** was allowed to react with 2,4-bis(4-methoxyphenyl)-1,3,2,4-dithiadiphosphetane 2,4-disulfide, Lawesson's Reagent (LR), in benzene at room

$$\begin{array}{c} O \\ R \\ NMe_2 \\ 1p, R = 2\text{-Thienyl} \\ q, R = 2\text{-Furyl} \\ s, R = 2\text{-Pyridyl} \\ \end{array}$$

Scheme 43

temperature, the enaminothione **129** was produced in a high yield (Scheme 39).<sup>64</sup>

*N,N*-Dimethyl enamine **131** underwent hydrolysis upon boiling in acetic acid containing sodium acetate to give the bifunctional thiazolidinone- $\alpha$ -carboxaldehyde derivative **132** (Scheme 40).<sup>65</sup>

#### 2.2. Preparation of carbocyclic compounds

As mentioned earlier in this review,  $\beta$ -aminovinyl ketones **1d-f** underwent self-condensation on reflux in acetic acid yielding the **1**,3,5-trisubstituted benzene derivatives **135a-c**. <sup>43,66,67</sup> The electron rich C-2 in one molecule of **1d**, **1e** or **1f** adds to the electron deficient C-3 in another molecule, forming the intermediates **133a-c**. It is most likely that these intermediates **133** reacted, in each case, swiftly with a third molecule of **1**, yielding the intermediates **134** that lose three molecules of dimethylamine, leading eventually to the triaroylbenzene derivatives **135**. It is believed that acidity of the reaction mixture has prompted such trimerization of enaminones into triaroylbenzenes (Scheme 41).

By using the synthetic sequence as was suggested for the synthesis of 135a-c, the trisubstituted benzene derivatives 138a-c were formed from aminovinyl ketones 1m, 1o and 1a, respectively.<sup>66-68</sup> Formation of intermediates 136 and 137 is suggested although direct concerted 2 + 2 + 2 cycloaddition leading directly to 137 can not be overlooked (Scheme 42).<sup>49</sup>

This approach was also suitable for the preparation of the 1,3,5-trisubstituted benzenes **140a-c**, where reaction of the enaminone-derived heterocycles, such as thieno, furo and pyrido enaminones **1p,q,s** in refluxing acetic acid led to the corresponding trisubstituted derivatives **140a-c**. The reaction takes place by condensation of three moles of each enaminone **1p,q** or **1r** to form the corresponding intermediate **139** which loses, in each case, 3 moles of dimethylamine, aromatizes and affords the final isolated product **140a**, **140b** or **140c** (Scheme 43).<sup>49,67</sup>

Furthermore, it has been found that heating a mixture of **141** and an excess of an aryl ethynyl ketone **142** ( $\sim$ 8 equivalents) in toluene resulted in smooth trimerization to afford the linked 1,3,5-triaroylbenzenes **143a–d** in good isolated yields, especially given the fact that six new C–C bonds are formed during the course of the reaction. Both electron rich (X = OMe) and electron deficient (R = NO<sub>2</sub>) aryl ethynyl ketones proved to be suitable reactants, although nitro-substituted linked triaroylbenzenes were isolated in slightly lower yields (Scheme 44).<sup>69</sup>

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Scheme 45

A similar treatment of the bis(enaminone) **144** with aryl ethynyl ketones **142a,b** resulted in the formation of the corresponding 1,3,5-triaroylbenzenes **145a,b** (Scheme 45).<sup>69</sup>

This protocol was also suitable for the preparation of tris(1,3,5-triaroylbenzene) derivative **147** in which the individual cyclotrimers are connected *via* a 1,3,5-trisubstituted phenyl ring (Scheme 46).<sup>69</sup>

Reaction of 3-dimethylaminoacrylaldehyde (28a) with aminothienocoumarin 85 in DMF at reflux for four hours gave a product of cycloaddition and dimethylamine elimination for which structure 148 was established. Similarly, condensed

aminothiophene **91** reacted with 3-dimethylaminoacrylaldehyde (**28a**) under the same applied reaction conditions to afford the analogous cycloaddition product **149**. Structures **138** and **149** were confirmed by preparing the same reaction products *via* alternative synthetic routes involving the reaction of acrylaldehyde (**150**) with condensed aminothiophenes **85** and **92**, respectively, under the same reaction conditions (Scheme **47**).

Lewis acid-catalyzed [3 + 2] cycloaddition of donor–acceptor cyclopropanes **151** and enamines **1d,o** in methylene chloride in presence of MgI<sub>2</sub> yielded nitrogen-functionalized cyclopentane derivatives **152** in good yields (Scheme 48).<sup>70</sup>

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Scheme 46

The use of enaminones as effective synthons for a directed C-H functionalization is reported. Proof-of-concept protocols have been developed for the Rh III-catalyzed synthesis of naphthalenes, based on the coupling of enaminones 1 with either alkynes 153 or  $\alpha$ -diazo- $\beta$ -ketoesters 154. Two inherently reactive functionalities (hydroxy and aldehyde groups) are integrated into the newly formed cyclic framework and a broad

range of substituents are tolerated, rendering target products 155, 156 readily available for further elaboration (Scheme 49).71

#### 2.3. Preparation of heteroaromatic compounds

β-Aminovinyl ketones are important synthetic intermediates, particularly in heterocyclic chemistry. 6,72 Heterocycles prepared from those ketones include carbazolequinone alkaloids,73 tricyclic benzo[a]quinolizines,74 pyrroles,75 benzodizepines,76 pyrimidines, 77 pyridines, 78 isoxazoles, 79 quinolines, 80,81 1,3thiazines, 82 furans, 83 benzothiazoles, 84 pyrazoles, 14,85-87 triazole,88 isochromanes,89 and 1,4-diazepines.90 In this regard, many other recent examples are herewith provided including the preparation of five and six membered heterocycles and some of their condensed derivatives.

#### 2.3.1. Preparation of five membered rings

2.3.1.1. Preparation of furans. Interaction of β-aminovinyl ketones with quinones represents an interesting approach to furan ring system. Many scientific publications in this area of investigation have been found. Thus, e.g., treating vinyl ketone **1b** with *p*-benzoquinone, in acetic acid with stirring at room temperature, yielded a product of addition via dimethylamine and water elimination, this product can thus be formulated as 161 or isomeric 162. However, the benzofuran-3-al structure 162 was assigned for that product on the basis of its elemental analysis and spectra data. Formation of 162, as illustrated in

Scheme 48

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Scheme 111, may be explained to occur *via* initial addition of electron rich C-2 in the enaminone **1b** to the electron-deficient quinone carbon, forming the non-isolable acyclic adduct **157**, followed by subsequent enolization would take place, affording the dihydroxy intermediate **158**. The latter underwent intramolecular ionic heterocyclization at room temperature *via* nucleophilic attack by the OH function on the carbonyl carbon rather than the methylene CH, yielding the *N*,*N*-dimethylaminomethylidene intermediate **160**, accompanied by hydrolysis with the release of dimethylamine and water, leading eventually to the final isolable benzofuranal derivative **162**.<sup>68</sup>

Similar to the behavior of β-aminovinyl ketone **1b** toward *p*-benzoquinone, compounds **1b** and **1p** also reacted with 1,4-naphthoquinone to produce the corresponding furanals **164a,b** through the intermediate Michael adducts **163a,b** (Scheme 50).

In addition, dienones **19a–c** reacted with *p*-benzoquinone, in acetic acid with stirring at room temperature, to yield products of addition *via* dimethylamine and water elimination, for which structure **167a–c** was considered as indicated from <sup>1</sup>H NMR spectra where a formyl-H was observed, in each case, at  $\delta$  9.0 (1H),  $\delta$  9.21 (1H) or  $\delta$  9.17 (1H) ppm. It is thus believed that *p*-benzoquinone initially adds to electron rich C-2, yielding acyclic

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dihydroxy intermediates 165a-c which then cyclize exclusively into benzofuranals 167a-c rather than the isomeric 160a-c. Although cyclization into 160 seems to be kinetically more favored, products 167 are apparently thermodynamically more stable because of their extended conjugated double bond system.37 By using the synthetic sequence as was suggested for the synthesis of 167a-c, the naphthofuran-3-al derivatives 168ad were formed from 1,4-naphthoquinone (Scheme 51).37,60

In contrast to the observed formation of 3-formyl derivatives on reacting β-aminovinyl ketones with quinones, our research group reported8 that pyrazolo enaminone 1x reacted with 1,4naphthoquinone to yield a product of addition and dimethylamine elimination. The naphthofuranoylpyrazole structure 173 was established for that product on the basis of its elemental analysis and spectra data. Formation of 173, as illustrated in Scheme 50, may be rationalized via initial addition of electron rich C-2 in the enaminone 1x to the active double bond of 1,4-

Me<sub>2</sub>N 
$$\stackrel{\bigcirc}{N}$$
  $\stackrel{\bigcirc}{N}$   $\stackrel{}{N}$   $\stackrel{\bigcirc}{N}$   $\stackrel{\bigcirc}{N}$   $\stackrel{\bigcirc}{N}$   $\stackrel{\bigcirc}{N}$   $\stackrel{\bigcirc}{N}$   $\stackrel{\bigcirc}{N}$ 

Scheme 53

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naphthoquinone, followed by subsequent enolization would take place, forming the non-isolable acyclic dihydroxy intermediate **169**. The latter underwent intramolecular ionic heterocyclization at room temperature *via* nucleophilic attack by the OH function on the methylene CH rather than the carbonyl carbon, yielding the dihydronaphthofuran intermediate **171**, accompanied by 1,2-elimination with the release of dimethylamine, leading eventually to the final isolable 2-unsubstituted naphthofuran derivative **173** as described in our earlier report. It seemed that our approach was not suitable for the synthesis of the naphthofuranal derivative **172** that would be expected by analogy with the aforesaid reports <sup>37,60,68</sup> on the reactivity of β-aminovinyl ketones toward quinones (Scheme 52).

In support of this view, pyrano enaminone **1y** also reacted with *p*-benzoquinone to give the corresponding pyranyl benzofuryl ketone **175** and not the pyranylbenzofuran-3-carboxaldehyde derivative **148**. Furthermore, reacting **1y** with **1,4**-naphthoquinone afforded naphthofuryl ketone **177**. The formation of **175** and **177** from **1y**, is assumed to occur *via* initial addition of the aminovinyl ketone, of electron rich C-2, to the active double bond in quinones, forming the intermediate phenolic adducts **175** and **176**, that cyclize *via* loss of dimethylamine to yield the final isolable products (Scheme **53**).<sup>44</sup>

In accordance with the previous observations reported by us<sup>7</sup> and by others,<sup>44</sup>  $\beta$ -ketoenamines **1z**, **23** and **179** reacted with p-benzoquinone to furnish the corresponding benzofuryl ketones **181a–c**, via the intermediacy of dihydroxy compounds **180a–c**.<sup>90,91</sup> The exact structure of products in these reactions has been firmly established on the basis of Heteronuclear Multiple Quantum Coherence spectroscopy (HMQC). Thus, HMQC of compound **181b**, as a representative example, indicated that carbonyl group at  $\delta$  188.9 ppm is not bonded to any hydrogen atoms. This fully supported the proposed structure **181**. A similar treatment of the N,N-dimethylamino derivatives of coumarin **1z** and of benzene **1d** with 1,4-naphthoquinone led to the naphthofuryl ketones **182a**<sup>90</sup> and **182b**,<sup>42</sup> respectively (Scheme 54).

2.3.1.2. Preparation of pyrazoles and isoxazoles. One of the common route to pyrazole and isoxazole ring systems is the interaction of  $\beta$ -aminovinyl ketones with N-nucleophiles. Several contributions have been made to this area of research. Thus, *e.g.*, treatment of vinyl ketone **183b** with hydrazine hydrate and with phenylhydrazine in absolute ethanol at reflux temperature for four hours gave rise to pyrazole products for which two possible structures **188** or **189** can be formulated. However, structure **189a,b** could be established for these products based on the non-identity of reaction product, obtained by reacting **183b** with phenylhyrazine, with a sample of **188b** prepared via initial condensation of acetophenone derivative **182b** with phenylhydrazine in refluxing absolute ethanol

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$$\begin{array}{c} \text{Me} \\ \text{N} \\ \text$$

Scheme 58

in the presence of acetic acid and subsequent treatment of the formed phenylhydrazone **190a** with DMFDMA (Scheme 55).<sup>92</sup> Compound **183b** also reacted with hydroxylamine hydrochloride, in absolute ethanol in the presence of fused sodium acetate under reflux for five hours, to yield a product that may be also formulated as **188c** or **189c**. Attempt to prepare a sample of **188c** from reaction of oxime **190b** with DMFDMA failed. However, structure **189c** could be established for the reaction product based on the fact that this reaction product readily converted, under basic reaction conditions, into the nitrile **191**, which was also prepared *via* direct cyanation of **192b** with *N*-cyanobenzotriazole **193** as an efficient *C*-cyanating reagent.<sup>92,93</sup>

A similar treatment of  $\beta$ -acylated enamine **194** with hydrazines led to 3-unsubstituted derivatives of pyrazole and iso-xazole **196a–c** on treatment with hydrazines and hydroxylamine. The isomeric 5-unsubstituted pyrazole **195b** could be prepared

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by condensing active methylene compound 199 with phenylhydrazine and subsequent condensation of the formed phenylhyrazone 197 with DMFDMA. On the other hand, the 5-unsubstituted isoxazole 196c was converted into the α-cyanoketone 198 on reacting with a little excess of sodium hydride in refluxing dioxane. Alternatively, compound 198 was directly prepared from reaction of 199 with 193 (Scheme 56).92

Scheme 61

O Ar 
$$\frac{NH_2OH.HCl}{ab. EtOH/Na_2CO_3/\Delta}$$
 Ar  $\frac{NH_2}{N}$   $\frac{NH_2}{N}$   $\frac{NH_2}{N}$   $\frac{NH_2}{N}$   $\frac{NH_2}{N}$   $\frac{NH_2}{N}$   $\frac{NH_2}{N}$   $\frac{NC}{N}$   $\frac{NC}$ 

Scheme 62

O CN 
$$NH_2XH$$
  $NH_2NH_2.H_2O$   $NMe_2$   $NH_2NH_2.H_2O$   $NH_2$   $NH$ 

Scheme 63

In accordance with the observed formation of 3-unsubstituted heterocyclic compounds, enaminoketones 1d and 1p,q reacted with hydrazine hydrate in refluxing ethanol to give the corresponding pyrazoles 201a-c, respectively. It is believed that hydrazine adds, in each case, reversibly across the  $\alpha,\beta$ -unsaturated moiety in 1d or 1p,q affording non-isolable intermediates 200a-c. This is followed by deamination and dehydration, leading eventually to the final isolable pyrazoles 201a-c. 55,94 In a similar manner, treatment of N,N-dimethylaminomethylene derivatives 1y and 23 with hydroxylamine hydrochloride led to the isoxazoles 201d,e, respectively (Scheme 57).44,95,96

In addition, heterocyclic enaminone 202 also reacted with each of hyrazine hydrate and methylhydrazine in refluxing ethanol to furnish the anticipated pyrazoles 204a,b, via the intermediacy of 203a,b. On the other hand, when the reaction of compound 202 with acetyl-, phenyl- and benzoyl-derivatives of hydrazine, was conducted in refluxing glacial acetic acid, in all cases, a single tetracyclic product was isolate for which structure 205 was established as indicated from spectral data of that reaction product. Formation of the tetracyclic azepine 205 would involve an initial formation of intermediates 203c-e which underwent nucleophilic cyclization with loss of hydrazine molecules to afford the final single product 205. The same

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compound 205 was also obtained by refluxing enaminone 202 in a mixture of glacial acetic acid and hydrochloric acid (1:1) for four hours (Scheme 58).<sup>97</sup>

In contrast to the observed formation of 3-unsubstituted heterocycles, 5-unsubstituted pyrazoles 207a,b were obtained upon heating of N,N-dimethylamino derivatives of pyridine 1t,u to 60-65 °C under stirring, with hydrazine hydrate through the intermediate formation of acyclic condensation products 206a,b.98 N,N-Dimethylamino derivative of pyrazole 1x underwent analogous reaction in refluxing DMF to yield the respective pyrazolylpyrazole derivative 207c as described by us in a previous communication.7 Compounds 1y also reacted with hydrazine and with phenylhydrazine in acetic acid at room temperature to afford the pyranylpyrazoles 207d,e.44 When compound 1z reacted with hydrazines in refluxing ethanol, the pyrazoles 207f,g were isolated, while the isoxazole 207h was obtained upon heating of 1z with hydroxylamine hydrochloride in ethanol in the presence of anhydrous sodium acetate at reflux for six hours.90 Compound 207f was also obtained in yield of 60%, when the reaction of 1z with hydrazine was carried out under microwave heating for five minutes (Scheme 59).99

An interesting reaction leading to 3-unsubstituted pyrazoles is the interaction of 3-(N,N-dimethylamino)-methylene derivatives **49a** with hydrazines in refluxing ethanol to give the pyrazoles **208a–d.**<sup>100,101</sup> On the contrary, compound **49a** reacted with hydroxylamine hydrochloride in refluxing ethanol to yield

a product that was assigned 5-unsubstituted isoxazole structure 209 based on  $^1H$  and  $^{13}C$  NMR spectra. $^{101}$  Interestingly, it has been found that interaction of 49a with an excess of hydroxylamine hydrochloride, in pyridine at reflux temperature for two hours, led to the 5-aminoisoxazole derivative 212, which is assumed to proceed by reaction of the vinyl ketone 49a with two molecules of hydroxylamine, affording the dioxime 210, that then loses a molecule of water, yielding the  $\alpha$ -cyano oxime 211, followed by spontaneous cyclization to the final isolable aminoisoxazole 212 (Scheme 60). $^{100,102}$ 

Unexpectedly, treatment of aminovinyl ketone 23 with hydrazine hydrate in refluxing absolute ethanol in an attempted to obtain the pyrazole derivative 213 was unsuccessful (Scheme 61). The isolated product was identified as 2,3-dihydrophthalazine-1,4-dione (214) identical to that reported in the literature. 95,103

Interestingly, it has been found that enaminonitriles underwent somewhat different reactions with N-nucleophiles, providing access to cyano or amino heterocyclic compounds, depending on the applied reaction conditions. In the first step of those reactions, the dimethylamino group was substituted by hydrazines or by hydroxylamine. The final products were obtained by ring closure to the carbon atoms either of the carbonyl group, affording cyano compounds, or of the nitrile group, yielding amino analogues. Several data have been reported on this area of study (Scheme 62).

As previously indicated in this review, interaction of enaminonitriles **127a**,**b** with hydroxylamine hydrochloride did not

O CI Bet3N Benzene/r.t.

NHAr

228a, 
$$Ar = Ph$$
b,  $Ar = C_6H_4Me-p$ 

O NMe2

1d,  $R = Ph$ 
p,  $R = 2$ -Thienyl
q,  $R = 2$ -Eyridyl
s,  $R = 2$ -Pyridyl
s,  $R = 2$ -Pyridyl

NAr

NAr

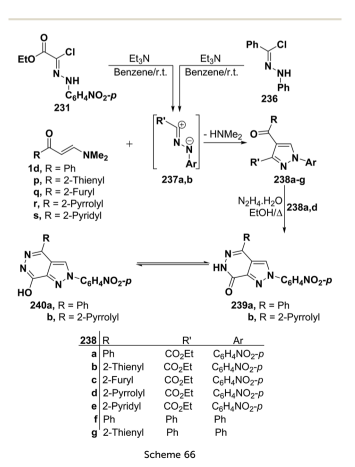
Me

233a-j

N<sub>2</sub>H<sub>4</sub>.H<sub>2</sub>O EtOH/ $\Delta$ 

A R = Ar = Ph
b,  $R = 2$ -Thienyl;  $Ar = Ph$ 
c,  $R = 2$ -Pyridyl;  $Ar = Ph$ 
d,  $R = 2$ -Pyridyl;  $Ar = Ph$ 
c,  $R = 2$ -Pyridyl;  $Ar = Ph$ 
d,  $R = 2$ -Pyridyl;  $Ar = C_6H_4Me-p$ 
d,  $R = 2$ -Pyridyl;  $Ar = C_6H_4Me-p$ 
i,  $R = 2$ -Pyridyl;  $Ar = C_6H_4Me-p$ 
j,  $R = 2$ -Pyridyl;  $Ar = C_6H_4Me-p$ 
j,  $R = 2$ -Pyridyl;  $R = C_6H_4Me-p$ 

Scheme 65



Review

Scheme 67

Scheme 69

give expected isoxazoles 130a,b, but instead afforded the acyclic hydroxylaminomethylenes 215a,b, that could not be evclized into 130a,b under a variety of reaction conditions.61 In contrast, compounds 126a,b reacted, on one hand, with hydrazine hydrate in refluxing absolute ethanol for six hours to yield the cyanopyrazoles 217a,b and, on the other, with phenylhydrazine under the same experimental conditions to produce the aminopyrazoles 219a,b. It is believed that compounds 127a,b react initially with both hydrazines to provide the non-isolable acyclic hydrazino derivatives 216a-d. The intermediates 216a,b cyclize readily via water elimination, leading to pyrazoles 217a,b.61 No traces of the other possible regioisomers 218a,b could be isolated, since the <sup>1</sup>H NMR spectra of the obtained products showed, in each case, the presence of pyrazole H-3 proton as a singlet at  $\delta$  7.9 ppm and this disagree with the tautomeric 218a,b in which the corresponding pyrazole H-5 protons would appear as doublets. 104,105 Similar cyclization of 216c,d into 219a,b is sterically hindered as it could produce pyrazoles with two adjacent bulky substituents. Consequently, intermediates 216c,d cyclize via nucleophilic addition to cyano group, yielding the aminopyrazole derivatives 218a,b as confirmed by elemental analyses and spectral data (Scheme 63).

In contrast to the observed formation of acyclic products **129a,b**, enaminonitrile **220** reacted with hydroxylamine hydrochloride in absolute ethanol in the presence of potassium carbonate anhydrous at reflux temperature to give the aminoisoxazole derivative **223** as confirmed by elemental and spectral data. Compound **223** is assumed to be formed *via* a Michael type addition of the amino group of hydroxylamine to the enamine double bond in **220** with loss of dimethylamine

Scheme 70

molecule, forming acyclic intermediate 221a, which cyclized into the final amino derivative 223 via nucleophilic addition to cyano group.106 On the other hand, it has been found that reacting enaminonitriles 67 and 220 with hyrazines in refluxing ethanol led to the cyanopyrazoles 223a-c, respectively, which is a contradiction to the behavior of compound 220 toward hydroxylamine. The spectral data of the isolated products were

NMe<sub>2</sub> + R<sub>1</sub>-X + NH<sub>2</sub>NH<sub>2</sub> 
$$Cu(OAc)_2 / Cs_2CO_3$$
 N—R<sub>1</sub>

1d-f,o

R<sub>1</sub> = Ph, 4- MeC<sub>6</sub>H<sub>4</sub>, 4-MeOC<sub>6</sub>H<sub>4</sub>, 4-NO<sub>2</sub>C<sub>6</sub>H<sub>4</sub>

X = I, Br

Cu(OAc)<sub>2</sub> / Cs<sub>2</sub>CO<sub>3</sub>

R

272

15 examples up to 87% yield

Scheme 72

Scheme 73

NMMe<sub>2</sub>

1d, R = Ph

g, R = 
$$C_6H_4Cl-p$$
h, R =  $C_6H_4NO_2-p$ 

Nucleophilic

Addition

O NMe<sub>2</sub>
 $C = N$ 
 $C$ 

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in complete agreement with structure 222a-c. Consequently, the acyclic hydrazino intermediates 221b-d, formed in this process, are not subjected to nucleophilic addition, but are stabilized by the evolution of water molecules (Scheme 64). 50,106

An interesting reaction leading to aminopyrazoles is the interaction of enaminonitriles 224a-c with hydrazine hydrate in refluxing dioxane for three hours to furnish the aminopyrazoles 226 and not the cyanopyrazoles 225. Inspection of <sup>1</sup>H NMR spectrum enabled establishing structure 226 for these pyrazole derivatives since the pyrazole H-3 appeared as a singlet at  $\delta$  8.0-8.3 ppm, the authors could not trace in the <sup>1</sup>H NMR any signals for the tautomeric 227 as this would reveal pyrazole H-5 as doublet (Scheme 64).104

It is worthwhile to report here that interaction of β-aminovinyl ketones with hydrazonoyl halides in the presence of bases also provides access to pyrazole ring system. Many examples have been reported on this interesting area of research.

Al-Zavdi and co-workers107 reported that treatment of vinyl ketones 1d and 1p-s with nitrilimines 229a,b (liberated in situ from the corresponding hydrazonovl halides 228a,b by the action of triethylamine in dry benzene solutions at room temperature) led to, in each case, the formation of only one isolable product as tested by TLC analysis. The reaction products were identified as the 5-unsubstituted pyrazole structure 233a-i (Scheme 65) that are assumed to be formed via 1,3dipolar cycloaddition of the nitrilimines 229a,b to the activated double bond in the vinyl ketones 1d and 1p-s to afford the nonisolable dihydropyrazole intermediates 232a-j followed by elimination of dimethylamine yielding the final pyrazole derivatives 233a-j. 94,107-109 The other possible regioisomers 231a-j are not observed through out the reaction course and were excluded on the basis of the spectral data of the isolated products. This conclusion was further confirmed chemically via reacting some examples of the isolated products 233a,b,e,f with

NMe<sub>2</sub> + 
$$R_1$$
 N<sub>3</sub> DES (ChCl : ethylene glycol at 1 : 2)

1d-g,i,o

 $R_1 = Ph,4-MeOC_6H_4, 4-ClC_6H_4, heptyl$ 

283

11 new compounds

Scheme 76

Scheme 77

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$$\begin{array}{c} \text{O} \\ \text{Ar} \\ \text{NMe}_2 \\ \text{1d, Ar = Ph} \\ \text{e, Ar = C}_{6}\text{H}_{4}\text{Me-p} \\ \text{f, Ar = C}_{6}\text{H}_{4}\text{Cl-p} \\ \text{g, Ar = C}_{6}\text{H}_{4}\text{Cl-p} \\ \text{Ar} \\ \text{O} \\ \text{Ar} \\ \text{Ar} \\ \text{Ar} \\ \text{O} \\ \text{Ar} \\$$

hydrazine hydrate in ethanol under reflux to afford the pyrazolo [3,4-d]pyridazine derivatives 234a-d in almost quantitative yields. It is of importance to report here that compounds 234ad can not be prepared by the action of hydrazine on the other possible regioisomers 234a-j under the same experimental conditions as shown in (Scheme 65).

By using the synthetic sequence as was suggested for the synthesis of 233a-j, the 5-unsubstituted pyrazole derivatives 237a,b were formed from nitrilimines 237a,b that generated by the action of triethylamine on hydrazonoyl halides 235 and 236, respectively, under the same experimental conditions.107 Further confirmation of the proposed structure 238 comes from reacting the isolated products 237a,d with hydrazine hydrate in ethanol under reflux to give products that were formulated as the pyrazolo[3,4-d]pyridazinones 239a,b rather than the corresponding hydroxy derivatives 240a,b as confirmed by the spectroscopic data of the isolated products (Scheme 66).107

In accordance with the observed formation of 5-unsubstituted pyrazoles, aminomethylene derivatives 241, 242 and

243 reacted with nitrilimines 245a-h (generated from hydrazonovl halides 244a-k and triethylamine) to furnish the corresponding pyrazole derivatives 246a-k by cycloaddition and dimethylamine elimination. 4,99,110,1111 Structure 246 was further established based on conversion of the isolated products 246a-e with hydrazine hydrate into the corresponding pyrazolo[3,4-d] pyridazines 247 and 249a-e, respectively. Clearly, these products can only be obtained from 5-unsubstituted isomer. It is worth mentioning herein that treatment of pyrazoles 246d-h with hydrazine hydrate under reflux in ethanol led to the expected pyrazolopyridazines 249a-e, respectively. 4,99,110 Similar hyrazinolysis of pyrazoles 246a-c, in refluxing ethanol for two hours, resulted in the formation of only one isolable product. This product was identified as the 7-hydroxypyrazolopyridazine structure 247 that is formed *via* loss of a molecule of methanol. ethanol or aniline (Scheme 67).110

Also, it has been found that reaction of enaminone 23 with nitrilimines 250a-c gave rise to the pyrazoles 251a-c. Refluxing compound 251a,b with hydrazine hydrate for five minutes led to the formation of 4-(aminomethyl)pyrazolopyridazines 253 and 254, respectively, that are assumed to be formed as a result of hydrolysis accompanied by the release of phthalic acid (Scheme 68).112

By analogy with hydrazonoyl chlorides, hydroximoyl chlorides 256 also reacted with vinyl ketones 1d and 1p-s in the

$$\begin{array}{c} \textbf{O} \\ \textbf{NMe}_2 \\ \textbf{1p, 2-Thienyl} \\ \textbf{q, 2-Furyl} \\ \textbf{s, 2-Pyridyl} \\ \textbf{s, 2-Pyridyl} \\ \textbf{a, R} = \textbf{Me} \\ \textbf{b, R} = \textbf{C}_6 \textbf{H}_4 \textbf{COMe}_- p \\ \textbf{c, R} = \textbf{2-Thienyl} \\ \textbf{d, R} = \textbf{2-Pyridyl} \\ \textbf{f, R} = \textbf{2-Naphthyl} \\ \end{array}$$

Scheme 79

Scheme 81

Scheme 82

presence of triethylamine to produce products of condensation via dimethylamine hydrochloride elimination for which three isomeric structures are possible. However, the structure of isolated products was considered to be 5-unsubstituted isoxazole structure 257 rather than other potential isomeric structures 259 and 261 based on spectral data and on the fact that the reaction products 263a,b,d,f were readily converted into the corresponding isoxazolo[3,4-d]pyridazines 264a–d, respectively, upon treating with hyrazine hydrate in refluxing absolute ethanol for 3–4 hours. It is believed that 256a,b initially generate nitrile oxides 257a,b *in situ* and these then undergo 1,3-dipolar cycloaddition to vinyl ketones 1d and 1p–s yielding intermediate cycloadducts 262a–g that aromatize via dimethylamine elimination (Scheme 69). 107,108

In a similar way, aminomethylene derivatives 1p, 239 and 1z underwent analogous reactions with a variety of nitrile oxides

266a–f (generated from hydroximoyl chlorides 265a–f and triethylamine) to give the expected isoxazole derivatives 267a–h by cycloaddition and dimethylamine elimination in a straightforward manner (Scheme 70).<sup>4,99,108</sup> Spectral data as well as chemical behavior of the formed isoxazoles indicated that they are the 5-unsubstituted isomeric structure 267a–h and this agrees with an already established trend in the behavior of  $\beta$ -aminovinyl ketones toward both nitrilimines and nitrile oxides.

An interesting synthesis of 5-unsubstituted pyrazole is the formation of **270** upon fusion of aminomethylene derivative **1d** with phenylhydrazonylpyridinium bromide **269** (Scheme 71).<sup>113</sup>

The synthesis of 1,3-disubstituted pyrazole derivative 272 was initiated with enaminone 1d-f,o, hydrazine hydrate, and aryl halide derivatives 271 in the presence of presence of Cs<sub>2</sub>CO<sub>3</sub> and Cu catalyst in DMF at elevated temperature (Scheme 72).<sup>114</sup>

Scheme 83

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4-Hydrazinothienopyrimidine 273A or 273B and the appropriate (E)-3-(dimethylamino)-1-phenylprop-2-en-1-one (1) were mixed in ethanol containing HCl, and the mixture was heated to 70 °C for 6 h, yielded 4-(5-aryl-1*H*-pyrazol-1-yl)thieno[2,3-*d*] pyrimidine 274A and 4-(5-aryl-1*H*-pyrazol-1-yl)thieno[3,2-*d*] pyrimidine 274B, respectively (Scheme 73).115

2.3.1.3. Miscellaneous. It is also of value to report here that reactions with β-aminovinyl ketones allow an easy access to some other five membered heterocycles such as thiophenes and pyrroles. Some recent reports on this area have been published.

Elnagdi et al.116 reported on the utility of vinyl ketones as aldehyde equivalents in Gewald reactions. It has been found that ketones 1d,g,h reacted smoothly with elemental sulfur and ethyl cyanoacetate in the presence of equivalent amounts of piperidine, under reflux in dry DMF for 6-8 hours, to yield 4-unsubstituted-2aminothiophenes 279a-c in good yields. It is believed that the initial step in the reaction sequence would be the nucleophilic addition of an active methylene nitrile to the α,β-unsaturated moiety in ketones 1d,g,h with subsequent elimination of

dimethylamine to yield 276A or 276B. Alternatively, reaction of adducts 275 with sulfur in the presence of an equivalent amount of piperidine would yield 278 that would cyclize and aromatize to yield 279 via loss of hydrogen sulfide and dimethylamine molecules during the reaction course. This sequence is quite similar to the general mechanism of the Gewald reaction.117 In either case, the reactivity of methylene moiety in either 275 or 276 is essential for the success of reaction (Scheme 74). Consequently, vinyl ketones 1d,g,h proved to fulfill this prerequisite as they contain electron attracting substituents.116

In addition, Al-Zaydi et al.99 reported on the formation of pyrrole derivatives 281 upon treatment with acetic acid and ammonium acetate in a domestic microwave oven at full power. It is assumed that initially formed 2,4-dicoumarinoylpyrroles 280 undergo a Nenitzescu like cyclization118 and decarbonylation; thus yielding the final products 281, respectively (Scheme 75).

Martins and et al. evaluated the effect of type III DES in a reaction between β-enaminones 1d-g,i,o and organic azide 282 for the synthesis of 4-acyl-1-substituted-1,2,3-triazoles 283 via 1,3-dipolar cycloaddition. The products were obtained in high selectivity and good yields (70-84%). The advantages of the method include easy work-up, metal-free conditions, inexpensiveness, and the ability to be used four times without a loss in yield (Scheme 76).119

1,5-Disubstituted 1,2,3-triazole derivatives **286** were prepared via the three-component reactions enaminone 1, amine 284, and tosylhydrazine 285. This metal- and azide-free, regioselective synthetic method proceeds in the presence of only molecular iodine (Scheme 77).120

#### 2.3.2. Preparation of six membered rings

2.3.2.1. Preparation of pyridines and related compounds. As mentioned earlier in this review, refluxing β-aminovinyl ketones 1d-f in acetic acid alone resulted in the formation of 1,3,5-triaroylbenzene derivatives 135a-c by a self condensationelimination route. 43,66,67 In contrast, on being heated in acetic

$$\begin{array}{c} \text{O} \\ \text{Me} \\ \text{NMe}_2 \\ \text{1d, R = Ph} \\ \text{e, R = C}_6\text{H}_4\text{Me}\text{-}p \\ \text{f, R = C}_6\text{H}_4\text{COMe}\text{-}p \\ \text{o, R = 2-Naphthyl} \\ \end{array} \begin{array}{c} \text{Me} \\ \text{O} \\ \text{Me} \\ \text{O} \\ \text{NMe}_2 \\ \end{array} \begin{array}{c} \text{NMe}_2 \\ \text{NMe$$

$$\begin{array}{c} \text{NMe}_2 & \text{Me} & \text{X} & \text{Me} & \text{X} \\ \hline & \text{AcONH}_4/\text{AcOH}/\Delta & \text{AcONH}_4/\text{AcOH}/\Delta & \text{NMe}_2 \\ \hline & \text{NMe}_2 & \text{AcONH}_4/\text{AcOH}/\Delta & \text{NN} \\ \hline & \text{NMe}_2 & \text{AcONH}_4/\text{AcOH}/\Delta & \text{NN} \\ \hline & \text{NMe}_2 & \text{AcONH}_4/\text{AcOH}/\Delta & \text{NN} \\ \hline & \text{NMe}_2 & \text{NC} & \text{NC} & \text{NC} \\ \hline & \text{NMe}_2 & \text{NC} & \text{NC} & \text{NC} \\ \hline & \text{NMe}_2 & \text{NC} & \text{NC} & \text{NC} \\ \hline & \text{NMe}_2 & \text{NC} & \text{NC} & \text{NC} \\ \hline & \text{NMe}_2 & \text{NC} & \text{NC} & \text{NC} \\ \hline & \text{NMe}_2 & \text{NC} & \text{NC} & \text{NC} \\ \hline & \text{NMe}_2 & \text{NC} & \text{NC} & \text{NC} \\ \hline & \text{NMe}_2 & \text{NC} & \text{NC} & \text{NC} \\ \hline & \text{NC} & \text{NC} & \text{NC} & \text{NC} \\ \hline & \text{NC} & \text{NC} & \text{NC} & \text{NC} \\ \hline & \text{NC} & \text{NC} & \text{NC} & \text{NC} \\ \hline & \text{NC} & \text{NC} & \text{NC} \\$$

Scheme 86

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acid in the presence of ammonium acetate, those ketones 1d-g afforded 6-substituted-3-aroylpyridines 289a-d, respectively. 66,67 The electron rich C-2 in one molecule of 1 adds to the electron deficient C-3 of another molecule, followed by dimethylamine elimination, forming the intermediates 21 and 287, respectively. The latter intermediates 287 cyclizes by the action of ammonia into the final isolable products 289, through the intermediacy of 288 (Scheme 78).

By using the synthetic sequence as was suggested for the synthesis of **289**, the 6-substituted-3-aroylpyridines **291a–f** were formed from dimethylaminomethylene compounds **1a**, **1m**, **1o** and **1p**,**q**,**s** respectively, by self cyclocondensation through the intermediate formation of **290a–f** (Scheme 79).<sup>66–68</sup>

Also, treating vinyl ketones 1z in acetic acid/ammonium acetate afforded the corresponding pyridines 277 (ref. 78 and 99) while interaction of 1z with enamino carbonyl compound 1a in a microwave oven in the presence of ammonium acetate and acetic acid for two minutes gave a mixture of 292 and methylpyridine derivative 293 (Scheme 80).<sup>99</sup>

It is of value to report here that reaction of  $\beta$ -aminovinyl ketones with acetyl derivatives also provide access to pyridines. Lots of recent reports on this area have been found. Thus, *e.g.*, reacting aminovinyl ketone **1d** with acetophenone in refluxing acetic acid in the presence of ammonium acetate yielded a product that may be formulated as **294** or isomeric **295**. The cyclization reaction may proceed by two possible mechanisms, which differ in their sequential nucleophilic attack/amine exchange reaction. While initial condensation of the methyl function with the carbonyl group of the vinyl ketone **1d** and subsequent cyclization could lead to structure **294**, initial Michael addition of the methyl ketone across the activated double bond in **1d** and subsequent cyclization might afford compound **295** (Scheme 81).<sup>121</sup>

In a similar way, pyridine enaminones **1t,u** reacted with acetylpyridines in anhydrous tetrahydrofuran in the presence of potassium *t*-butoxide under stirring followed by refluxing in glacial acetic acid in the presence of ammonium acetate to afford the terpyridines **297a–e** in 25.8–29.7% yields through

the intermediate formation of Michael adducts **296a-e** (Scheme 82).<sup>122</sup>

Analogously, aminovinyl ketone 1z reacted with acetyl derivative 298 in refluxing acetic acid in the presence of ammonium acetate to produce the corresponding pyridine derivative 299 (Scheme 83).<sup>78</sup>

An interesting reaction leading to pyridine ring system is the interaction of enaminonitrile **121a** with acetophenone to give a sole isolable reaction product that was assigned structure **301** rather than isomeric **300** based on NOE difference NMR experiments where irradiation of pyridine 3-H at  $\delta$  7.37 ppm enhanced the dimethylamino signal (Scheme 84).<sup>121</sup>

It is also worthwhile to report here that interaction of βaminovinyl ketones with active methylenes allows an interesting access to pyridine ring system. Intensive research work on this area of studies have been described. Elnagdi et al. 42,66,68,121 reported that reaction of aryl enaminones 1d,g, 1m with acetylacetone in refluxing acetic acid in the presence of ammonium acetate afforded pyridine derivatives. Two pathways can be envisioned for formation of these final products (Scheme 85). Thus, initial addition of active methylene moiety to α,β-unsaturated double bond in enaminones would afford Michael adducts 304a,d-f that in the presence of ammonium ion condensed, in each case, with acetylcarbonyl group yielding an enamine that cyclizes into 6-arylpyridines 305a,d-f. In contrast, initial condensation of active methylene moiety with carbonyl function would yield 302a,d-f that in the presence of ammonium acetate cyclizes into the alternative 4-aryl analogues 303. Structure 305 was preferred over possible isomeric structure 303 on the basis of <sup>1</sup>H NMR spectra.<sup>7,42,66</sup> A similar treatment of the aromatic enaminones 1e-g and 10 with ethyl acetoacetate led to the corresponding 4,5-unsubstituted pyridines 306b-d,f as confirmed by the <sup>1</sup>H NMR spectra, which indicated pyridine protons with J = 8.0-8.2 Hz, that is characteristic for pyridine H-5 and H-4.66

In addition, heterocyclic enaminones 1s, 1x, 1z and 23 also underwent analogous reactions with either acetylacetone or ethyl acetoacetate to yield the expected 6-heterocyclylpyridine derivatives 308a-f, respectively, as the sole isolable products (Scheme 86). Formation of these compounds would involve an initial formation of Michael adducts 307a-f, followed by intramolecular cyclization to give the final products 306a-f.<sup>7,25,91,121</sup> 6-

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$$N=N O CO_2Et N=N Me CO_2Et N=N Me CO_2Et N=N Me CO_2Et N=N Me N$$

Substituted pyridine structure 308a-f was assigned for those products based on the presence of pyridyl ring protons, in each case, as doublets with J=8 Hz as reported by us<sup>7</sup> and by others.<sup>25,79,121</sup>

In accordance with the observed formation of 6-heterocyclylpyridine derivatives **308**, enaminonitrile **67** reacted with ethyl acetoacetate in refluxing acetic acid in the presence of ammonium acetate to provide the expected pyridine derivative **310**. In contrast, treating compound **67** with acetylacetone under the same experimental conditions resulted in the formation of an acyclic product **310** that could not be cyclized into the anticipated pyridine derivative **312** (Scheme 87).<sup>50</sup>

Similar to the behavior of β-aminovinyl ketones toward acetylmethylene compounds, those vinyl ketones also reacted with cyanomethylene compounds to furnish pyridines. Thus, reaction of vinyl ketones **95a,b** with malononitrile in refluxing acetic acid and in presence of ammonium acetate yielded products of condensation via dimethylamine elimination. These may thus be formulated as **313–316**. Initial Michael addition to the  $\alpha$ , $\beta$ -unsaturated linkage can afford **314** that would then lose dimethylamine and cyclize into **315** or

isomerize into 317 (Scheme 88). Alternatively, condensation of the carbonyl group in 95a,b with the active methylene moiety would afford 313 that can then cyclize into 316.<sup>25</sup>

An interesting reaction leading also to pyridine ring system is the interaction of phenyl enaminone 1a with benzoylacetonitrile in refluxing acetic acid in the presence of ammonium acetate to furnish a mixture of two products. One of the products was assigned structure 323 and is assumed to be formed via initial addition of the active methylene moiety in benzoylacetonitrile to the enaminone C-3 and subsequent elimination of the dimethylamine yielding the acyclic Michael adduct 322, that would react with ammonia to yield aminopyridine derivative 323 (Scheme 89). Several isomeric structures seemed possible for the other product, depending on the reaction route, namely; the pyridone structure 319 resulting from initial condensation of the active methylene moiety with the enaminone carbonyl function, forming 318, followed by hydrolysis and subsequent cyclization, or an isomeric pyridone structure 321 resulting from initial condensation of the benzoylacetonitrile carbonyl function with the enaminone C-2, affording 320 and subsequent cyclization. Both structures 319 and 321 were excluded

$$\begin{array}{c} \textbf{NMe}_2 \\ \textbf{1d,p} \\ \textbf{NMe}_2 \\ \textbf{a,} \ \textbf{X} = \textbf{S} \\ \textbf{b,} \ \textbf{X} = \textbf{O} \\ \textbf{b,} \ \textbf{X} = \textbf{O} \\ \end{array} \\ \begin{array}{c} \textbf{NMe}_2 \\ \textbf{NMe}_2 \\ \textbf{A}, \ \textbf{X} = \textbf{S} \\ \textbf{b,} \ \textbf{X} = \textbf{O} \\ \end{array} \\ \begin{array}{c} \textbf{NMe}_2 \\ \textbf{NMe}_2 \\ \end{array} \\ \begin{array}{c} \textbf{HN} \\ \textbf{NMe}_2 \\ \textbf{A}, \ \textbf{A} = \textbf{CN} \\ \textbf{NMe}_2 \\ \textbf{A}, \ \textbf{A} = \textbf{Ph}, \ \textbf{X} = \textbf{O}; \ \textbf{b}, \ \textbf{R} = \textbf{Ph}, \ \textbf{X} = \textbf{S}; \\ \textbf{c,} \ \textbf{R} = \textbf{thienyl}, \ \textbf{X} = \textbf{O}; \ \textbf{d}, \ \textbf{R} = \textbf{thienyl}, \ \textbf{X} = \textbf{S} \\ \end{array}$$

Scheme 93

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NMe<sub>2</sub> 
$$\frac{NC}{Pyridine/\Delta, 5h \ or}$$
  $\frac{NH_2}{EtOH/Pip./\Delta, 3h}$   $\frac{NR}{D}$   $\frac$ 

based on the absence of a low field ( $\delta$  > 8.5 ppm) pyridine 6-H signal. Consequently, the pyridone structure **324** was considered for the second reaction product that showed its stability on reflux in acetic acid or mineral acid, confirming structure **324**.<sup>121</sup>

In a similar way,  $\beta$ -aminovinyl ketones  $\mathbf{1d}$  and  $\mathbf{1p}$ ,  $\mathbf{q}$ ,  $\mathbf{s}$  reacted with malononitrile in refluxing ethanolic sodium ethoxide to

yield the ethoxypyridines 327a–d. Initial addition of carbanion of malononitrile across the activated double bond in vinyl ketones would yield the Michael adducts 325a–d. Subsequent addition of ethoxide anion to one of the cyano groups would give the iminoethers 326a–d, which would be cyclized *via* a nucleophilic attack of an NH group on a cyano carbon, affording the intermediates 327a–d. This would be finally followed by deamination and dehydration, leading to the isolated ethoxypyridine products 328a–d (Scheme 90).<sup>94</sup>

In contrast to the behavior of  $\beta$ -aminovinyl ketones toward cyanomethylene reagents, vinyl ketone 329 was treated with malononitrile, in ethanol in the presence of a catalytic amount of piperidine at reflux temperature to give a condensation product for which 6-unsubstituted pyridone structure 333 was considered rather than the alternative 4-unsubstituted structure 330 as indicated from spectral and chemical evidence. <sup>116</sup> In support of the proposed structure, compound 333 was also formed on treating 329 with cyanoacetamide in refluxing pyridine. Based on the above findings, the initial step in the reaction involves condensation of the carbonyl function in 329 with

Scheme 96

malononitrile, yielding the diene **331** which is hydrolysed to an intermediate amide **332** and then cyclizes to **333** with the loss of dimethylamine molecule (Scheme 91).<sup>123</sup>

In a similar manner, vinyl ketone **49a** reacted with ethyl cyanoacetate in refluxing dioxane in the presence of sodium hydride to yield 6-unsubstituted pyridone derivative **335** *via* the intermediacy of the dienes **334a,b** (Scheme 92).<sup>45</sup>

One of the main route to 3-cyanopyridine-2-thiones and -2-ones is the interaction of  $\beta$ -aminovinyl ketones with cyanothioacetamide and with cyanoacetamide. Several data on this research area have been published. For instance, 3-cyanopyridines 337a–d were prepared by treating enaminone 1d,p with cyanothioacetamide in refluxing acetic acid in the presence of ammonium acetate for 1.5 hours, or with cyanoacetamide in sodium ethoxide solution at reflux, yielding the target molecule  $\nu ia$  the intermediacy of Michael adduct 336 (Scheme 93).  $^{42,58,109,124,125}$ 

In accordance with the observed formation of 4,5-unsubstituted pyridines, heterocyclic enaminones **1p**, **242** and **202** reacted with cyanothioacetamide or cyanoacetamide to give the corresponding pyridines **339a–d**, respectively, *via* the intermediacy of Michael adducts **338a–d** (Scheme 94).<sup>67,108,110,126</sup>

Also, aminovinyl ketone **340b** condensed with cyanothioacetamide and with cyanoacetamide via dimethylamine elimination to yield the 3-cyanopyridines that were formulated as **341a,b** rather than isomeric **342a,b** as indicated from the  $^1$ H NMR spectra for the reaction products. In case of X = S,  $^1$ H NMR spectrum showed the pyridine ring CH as singlet at  $\delta$  7.76 ppm. This may suggest a favorable reaction product **342a** formed via initial addition of the active methylene moiety in cyanothioacetamide across the activated double bond in **340b** and subsequent cyclization and aromatization via dimethylamine elimination (Scheme 95). The alternative structure **341a** which could have resulted from initial condensation of the active methylene with the carbonyl function, should display a doublet for the pyridine ring CH.  $^{127}$ 

Treatment of dimethylaminomethylene compound 228a with cyanothioacetamide, cyanoacetamide or anion of malononitrile dimer in dry DMF and sodium hydride led to 5-benzoylpyridines 345a-c rather than isomeric 344a-c based on the mass spectra (MS) of the isolated products. Additionally, the IR spectrum showed the disappearance of the cyano group so that structure 345b became 346 upon hydrolysis (Scheme 96).<sup>128</sup>

Quite surprisingly, interaction of vinyl ketones 329 and 23 with cyanoacetamide or cyanothioacetamide gave, in each case, a sole isolable reaction product that was assigned 4-substituted pyridine structure 333 or 348a,b rather than isomeric 347a-c. Structure 333 or 348, in this case, resulted from initial condensation of the active methylene in cyanoacetamide or cyanothioacetamide with carbonyl function in 329 or 23 and subsequent cyclization and aromatization *via* dimethylamine elimination (Scheme 97). 95,123

On the other hand, it has been found that interaction of vinyl ketones 1d-f with 3-aminocrotononitrile 349 afforded the 5,6-

16 new compounds

Scheme 100

$$\begin{array}{c} O \\ R \\ \hline \\ 1d,f-h \\ \hline \\ 1d,f-h \\ \hline \\ 356 \\ \hline \\ Ar = Ph, \ 4-MeC_6H_4, \ 4-MeC_6H_4, \ 4-NO_2C_6H_4, \\ \hline \\ 3-NO_2C_6H_4, \ 4-ClC_6H_4, \ 4-ReC_6H_4, \ 4-ReC_6H_4,$$

Scheme 101

NMe<sub>2</sub> + 
$$O$$
 1) aq. KOH or NaOH/EtOH, reflux, 12 h  $O$  2) H<sub>2</sub>O  $O$  360a-e

Scheme 102

O NMe<sub>2</sub> 
$$\frac{(NH_2)_2CS}{NaOEt/\Delta}$$
  $\frac{(NH_2)_2CS}{NaOEt/\Delta}$   $\frac{(NH_2)_2CS}{NaOEt/\Delta}$   $\frac{(NH_2)_2CS}{NAOEt/\Delta}$   $\frac{(NH_2)_2CS}{NAOEt/\Delta}$   $\frac{(NH_2)_2CS}{NH}$   $\frac{(NH_2)_2$ 

unsubstituted pyridine derivative 353 *via* the intermediates 351, but not 350 (Scheme 98). Although this reaction can afford 4,5-unsubstituted analogue 352 as well, structure 353 is established based on H-5,6 coupling of pyridine which showed a value of 4 Hz, if the reaction product is 352 those protons would be expected to have a much higher value ( $J \sim 9~{\rm Hz}$ ).  $^{7,42,43,66}$ 

368a-c Scheme 104

 $c, X = NH_2$ 

$$\begin{array}{c} \text{NMe}_2 \\ \text{NMe}_2 \\ \text{1m, R} = C_6 H_4 \text{COMe-} \rho \\ \text{u, R} = 3 \text{-Pyridyl} \\ \text{NH}_2 \\$$

An interesting synthesis of quinolones is the formation of 354 on refluxing aminovinyl ketone 1j in ethanol for 60 minutes under catalytic transfer hydrogenation conditions (CTH) using

O NH2 NH2 .HNO3 369b Ar NH NMe2 121 373

a, Ar = Ph b, Ar = 
$$C_6H_4Me-p$$

NH2 N N NH2 O N

367a-c

Scheme 107

NMe<sub>2</sub> + CI NH<sub>2</sub> (i-pr)<sub>2</sub>NEt<sub>2</sub>/ EtOH R NO<sub>2</sub> i) Oxidative activation ii) displacement with 
$$R_1R_2NH$$
 R NR<sub>1</sub>R<sub>2</sub>  $R_1R_2 = 0$  OMe NH<sub>2</sub> NH

Scheme 108

Scheme 109

cyclohexene as hydrogen source and 10% Pd-C as catalyst (Scheme 99).129

Shankaraiah and et al. was reported a clean, convenient, and highly efficient method for the rapid construction of pyridine 2,6-diarylnicotinaldehydes 355 by an unexpected self condensation of enaminoketones 1 in the presence of NH<sub>4</sub>OAc in acetic acid under conventional conditions and also under microwave irradiation (Scheme 100).130

The three-component sequential reaction of enaminones 1d,f-h, cinnamaldehyde 356 and the appropriate amines 357, proceeded smoothly to give 1,3,4-trisubstituted 1,4-dihydropyridines 358a-t in aqueous DMF (Scheme 101). 131,132

Enaminones 1d-g,i was heated at reflux with isatin 359 in the presence of an aqueous solution of KOH or NaOH, followed by subsequent acidification with dilute hydrochloric acid to give the quinoline-4-carboxylic acids 360a-e in good to excellent yields (75-90%) (Scheme 102).133

2.3.2.2. Preparation of pyrimidines. One of the major routes to pyrimidines is the ring closure of  $\beta$ -aminovinyl ketones by bifunctional reagents. This one-pot cyclization involves both the carbonyl and the dimethylamino groups, leading eventually to the target pyrimidines. Some representative examples have been provided herewith. Thus, treatment of vinyl ketones 1d and 1p,q with thiourea in refluxing sodium ethoxide yielded the pyrimidine-2-thiones 363a-c, respectively. Formation of these products would involve, in each case, an initial Michael addition of the amino group in thiourea to the activated double bond in 1d or 1p,q yielding Michael adduct 361a, 361b or 361c. This is followed by deamination to form the acyclic non-isolable intermediates 362a-c, which then undergo enolization and subsequent nucleophilic cyclization via loss of water, affording the final products 363a-c. A plausible mechanism for the formation of pyrimidines 363a-c is depicted in scheme 163 (Scheme 103).94

In a similar manner, heterocyclic enaminone 202 reacted with thiourea in refluxing DMF/ethanol in the presence of excess anhydrous potassium carbonate to afford the pyrimidine 364, while the S-alkylated product 365 was obtained under similar reaction conditions, except that DMF/ethanol was replaced by methyl glycol.97 On the other hand, treating 202 with guanidine compounds 366a-c in refluxing methyl glycol in

the presence of excess anhydrous potassium carbonate resulted in the formation of the corresponding pyrimidines 368a-c, respectively, via the intermediate formation of acyclic secondary enaminones 367a-c (Scheme 104).97

In addition, reaction of aryl enaminone 1m with guanidine carbonate (369) in refluxing ethanolic sodium ethoxide solution for sixteen hours resulted in the formation of 2-aminopyrimidine derivative 370,68 whereas pyrimidine derivative 372 was isolated on treatment of 3-pyridyl enaminone 1u with guanidine derivative 371 via loss of dimethylamine and water molecules (Scheme 105).134

°CO<sub>2</sub>H 382a.b HNMe<sub>2</sub> 19. R = Ph **b**, R =  $C_6H_4OMe-p$ 383a,b  $c_{1}R = 2$ -Furyl Ŕ' 387a-g 382,383a, R' = Me  $\mathbf{b}$ ,  $\mathbf{R}' = \mathbf{Ph}$ 388a, R = R' = Ph **b**, R =  $C_6H_4OMe-p$ ; R' = Ph c, R = 2-Furyl; R' = Phd, R = Ph; R' = Me $\mathbf{e}$ , R = C<sub>6</sub>H<sub>4</sub>OMe-p; R' = Me f, R = 2-Furyl; R' = Me COR' 388a-g

Scheme 111

Interestingly, reaction of enaminonitriles 121a,b with guanidine nitrate (369) in absolute ethanol in the presence of excess anhydrous potassium carbonate at reflux temperature furnished the pyrimidine-5-carbonitriles 375a,b, via the intermediacy of Michael adducts 373a,b, rather than the 5-aroyl analogues 374a,b as proven from spectral data of the isolated products (Scheme 106).61

Recently, Alinezhad and et al. reported that phenylphosphinic acid or 2-pyrrolidonium bisulphate are found to catalyze the three-component condensation of an enaminone 1d,h, aldehyde 376, and urea or thiourea to afford the corresponding 6-unsubstituted dihydropyrimidinones 377 in high to excellent yields. This methodology is simple and fast synthetic route for the preparation of interesting class of heterocycles (Scheme 107).135,136

Ingham and et al.137 was reported the development of a monolith-supported synthetic procedure of enaminone 1 in the presence of Huunig's base, taking advantage of flow processing and the superior flow characteristics of monolithic reagents over gel-phase beads, to allow facile access to an important family of 2-aminopyrimidine derivatives 379 through formation of thioether derivative 378. The process has been successfully applied to a key precursor on route to Imatinib (Ar = 3-pyridyl,  $R_1$  = 2-methyl-5-nitrobenzyl,  $R_2$  = H) (Scheme 108).

The refluxing reaction of (E)-3-(dimethylamino)-1-(pyridin-3-yl) prop-2-en-1-one (1u) with 1-(2-methyl-5-nitrophenyl)guanidine nitrate 380 was processed in isopropyl alcohol for 12 h, to give N-(2-methyl-5-nitrophenyl)-4-(pyridin-3-yl)pyrimidin-2-amine 381 in 82% (Scheme 109).138

2.3.2.3. Preparation of pyrans. Recently, Kepe et al. 139,140 reported on an one-pot synthesis of different 2H-pyran-2-one derivatives via interaction of either monoactivated methyl

$$\begin{array}{c} \text{OH O} \\ \text{NMe}_2 \\ \end{array} \begin{array}{c} \text{Ph} \\ \text{N} \\ \text{NMe}_2 \\ \end{array} \begin{array}{c} \text{NMe}_3 \\ \end{array} \begin{array}{c} \text{NMe}_4 \\ \end{array} \begin{array}{c$$

Scheme 113

ketones or 1,3-dicarbonyl compounds with DMFDMA and Nacylglycines in acetic anhydride. More recently, the general nature of Kepe's pyranone synthesis was extended to reaction of N-substituted glycines with various β-aminovinyl ketones, with especial emphasis on ketones incorporating heterocyclic moieties. Thus, treating aliphatic enaminones 1a-c with Nacetylglycine (382a), hippuric acid (382b) or N-pyrazinylcarbonylglycine (382c) in acetic anhydride led to the corresponding pyranones 386a-g, respectively. It is believed that Nacylglycines 382a-c are first cyclized to the corresponding oxazolones 383a-g and subsequent reaction of the latter with enaminones 1a-c yielded initially Michael adducts 384a-g. These adducts undergo deamination and enolization to afford the non-isolable enols 385a-g that rearrange into the final isolable products 386a-g, respectively, via an attack of hydroxyl function on the oxazolone ring (Scheme 110).68,141

As an extension to the Kepe acylaminopyranone synthesis,  $^{141,142}$  dienones 19a-c reacted with N-acetyl- and N-benzoyl-glycines 382a,b in acetic anhydride to yield products assumed to be the pyranones 388a-f rather than the isomeric 389a-f. It is thus assumed that *in situ* generated oxazolones

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$$\begin{array}{c} O \\ R \\ \hline \\ OH \\ \hline \\ 1a-j,o \end{array} \qquad \begin{array}{c} AgSCF_3/TCCA, THF, rt \\ \hline \\ Or \quad I_2/toluene, rtt \\ \hline \\ TCCA: trichloroisocyanuric acid \\ \hline \\ X = SCF_3 \text{ or } I \end{array}$$

Scheme 116

Scheme 117

**383a,b** react with **19a-c** *via* initial addition of the active methylene in **383a,b** to the enaminone moiety at C-3, yielding intermediates **387a-g** that further rearrange into the final isolable products **388a-f** (Scheme 111).<sup>37</sup>

Similar reactions of **382a,b** with either aromatic enaminones **1d** and **1o** or with simple heterocyclic enaminones **1p-s** in acetic

anhydride resulted in the formation of the corresponding acylaminopyranones 389a-j, respectively (Scheme 112).<sup>58</sup>

In accordance with the observed formation of pyranones, heterocyclic enaminones **1y**, **1z** and **196** reacted with hippuric acid (**382b**) to give the desired pyranones **390a–d**, respectively (Scheme 113). <sup>44,77,99,126</sup> The formation of compounds **390a–d** starting from the corresponding heterocyclic enaminones and **382b** can thus be considered as an extension to the Kepe pyranone synthesis <sup>141,142</sup> to enable the synthesis of 6-heteroarylpyran-2-ones.

An interesting reaction leading to the formation of a pyranone ring system<sup>78</sup> is the interaction of aminovinyl ketone **1z** with dithiocarboxylic acid **391**, affording the pyranone derivative **394** *via* the intermediate formation of **392** and **393**, respectively, in a manner similar to that suggested to account for formation of pyranone **390b** from reaction of hippuric acid (**382b**) with the same vinyl ketone **1z**.<sup>78</sup> This is also a new extension to the Kepe acylaminopyranone synthesis (Scheme **114**).<sup>141,142</sup>

Unexpectedly, interaction of aminovinyl ketone **1k** with aromatic diazonium salts did not give the anticipated arylhydrazonals **396a–c**, instead cyclic *N*,*O*-acetals **398a–c** were formed in good yields. It is believed that the formed enazo compounds **395a–c** were initially cyclized into **397a–c** that reacted with ethanol to yield the final isolable chromones (4*H*-benzopyran-4-ones, **398a–c**). Both <sup>1</sup>H and <sup>13</sup>C NMR spectra of the reaction products fit completely with the proposed structure. Additionally, the X-ray crystal structure determination of **398a** confirmed the existence of the hydrazono-chromone form **398** rather than the alternative possible enol azo form **399**. <sup>141</sup> Furthermore, the *o*-hydroxy substituted derivative **1k** showed an additional reactivity when treated with acids: it underwent intramolecular ring

O NMe<sub>2</sub> 
$$\frac{1}{412}$$
  $\frac{1}{10}$   $\frac{1}{10}$ 

Scheme 119

closure wherein the aromatic hydroxy group attacked the aminopropenone side chain, leading to chromone **400** in quantitative yield (Scheme 115). 142

A facile and efficient synthetic strategy to 3-((trifluoromethyl) thio)-4*H*-chromen-4-one **401** was developed. AgSCF<sub>3</sub> and trichloroisocyanuric acid (TCCA) were employed here to generate active electrophilic trifluoromethylthio species *in situ*. <sup>143</sup> Or by using iodine in toluene. <sup>144</sup> This reactions could proceed under mild conditions in a short reaction time and be insensitive to air and moisture (Scheme 116).

2.3.3. Preparation of condensed heterocycles. Interaction of  $\beta$ -aminovinyl ketones **1d**,**e** with 4-substituted 5(3)-aminopyrazole derivative **402** in refluxing mixture of equivalent

amounts of ethanol and acetic acid produced the pyrazolo[1,5-*a*] pyrimidine derivatives **404a,b** (Scheme 117). Formation of the latter products, as shown in Scheme 104, may be explained to occur *via* initial nucleophilic attack by the endocyclic nitrogen on the electron deficient C-3 in **1d,e** forming intermediates **403a,b** with loss of dimethylamine, followed by subsequent nucleophilic cyclization with the release of water. <sup>145</sup>

c, R = 2-Thienyl

On the other hand, reaction of vinyl ketones 1d and 121a with 4-unsubstituted aminopyrazole 305 in the presence of zinc chloride and pyridine, respectively, yielded addition products with elimination of dimethylamine and water for which structure 407 was established. Additionally, when compounds 1d and 405 were refluxed in ethanol, compound 406a was isolated in a good yield. It could then be cyclized into 407a on fusion with zinc chloride.146 Likewise, treating vinyl ketones 1d and 121a with aminopyrazolone 408 in refluxing pyridine for three hours led to the corresponding pyrazolo[3,4-b]pyridines 409a,b (Scheme 105).144 In contrast to the observed formation of pyrazolopyridines, 146,147 the same ketones 1d and 121a reacted with 4-unsubstituted aminopyrazole 410 in refluxing pyridine to give the pyrazolo[1,5-a]pyrimidines 412a,b as confirmed by the spectral data of the reaction products. Accordingly, formation of compounds 412a,b, in this case, may be rationalized via initial nucleophilic attack by the exocyclic nitrogen on the electron deficient C-3 of enaminone moiety in 1d and 121a with subsequent elimination of dimethylamine, yielding intermediates **411a,b.** These undergo intramolecular cyclization *via* loss of water, affording the final products 412a,b (Scheme 105).54,148

Similar to the previously described<sup>52</sup> behavior of vinyl ketone **1d** toward aminopyrazole **410**, compound **1d** reacted with 3-aminopyrazolo[3,4-*d*]pyrimidine derivative **413** in refluxing glacial acetic/hydrochloric (1 : 1) mixture for ten hours to give the respective pyrimido[2,3 : 4,3]pyrazolo[1,5-*a*]pyrimidine derivative **414** (Scheme 105). In contrast to the reported behavior of enaminonitrile **121a** toward aminopyrazole **410**, compound **121a** reacted with 3-amino-1*H*-pyrazolo[3,4-*b*]

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Scheme 121

pyridine derivative **415** in glacial acetic acid at reflux temperature for three hours to furnish the tricyclic pyrimidine derivative **416** (Scheme 118).<sup>150</sup>

Refluxing vinyl ketone **1d** or **1d** with 2-aminobenzimidazole (**417**) in pyridine solution afforded the 4-substituted pyrimido[1,2-*a*]benzimidazoles **419a,b**,<sup>54</sup> while 1-substituted 4-cyanobenzimidazo[1,2-*a*]pyridines **422a–c** were obtained on treatment of vinyl ketones **1d,g** or **1p** with 2-cyanomethylbenzimidazole (**420**) in refluxing ethanol containing a catalytic amount of piperidine.<sup>23</sup> Formation of compounds **419a,b** is thus formed *via* addition of the ring nitrogen to the activated double bond in **1d** and **1s** yielding Michael adducts **418a,b** which then cyclize by elimination of water and aromatize *via* loss of dimethylamine, affording the isolable products **419a,b**.<sup>54</sup> On the contrary, compounds **422a–c** are formed *via* addition of active methylene to the activated double bond, leading eventually to the final products through the intermediate formation of **421a–c** (Scheme **119**).<sup>23</sup>

Moreover, heterocyclic enaminones **1z** and **49a** reacted with 3-amino-1,2,4-triazole (**405**) to yield products that may be formulated as **424** or the isomeric **425**. However, structure **425** was established for the reaction products based on <sup>1</sup>H NMR data and NOE experiments (Scheme 120).

Scheme 123

Scheme 124

Scheme 125

It has been reported<sup>68</sup> that vinyl ketones **1b** and **1m** coupled with the diazotized 5(3)-amino-3(5)-phenylpyrazole (**426a**), which exists in equilibrium with an isolable diazobetaine **426b**, to give azolotriazines **429a,b** rather than the expected acyclic  $\alpha$ -hydrazono- $\beta$ -ketoaldehydes **428a,b**. Thus, the inability to isolate the target hydrazones **428a,b** from this reaction may indicate that the reagents react via a direct 4 + 2 cycloaddition mechanism to the activated double bond in **1b** and **1m** leading to the isolated products **429a,b** (Scheme 121). Likewise, coupling compound **1m** with diazotized 2-aminobenzimidazole **430** and with diazotized 3-amino-1H-1,2,4-triazole **431** afforded the analogous [1,2,4]triazino[4,3-a]benzimidazole derivative **432** and triazolo[3,4-c][1,2,4]triazine derivative **433**, respectively, via direct 4 + 2 cycloaddition.<sup>68</sup>

Interestingly, the behavior of cyclic enaminone 95b toward diazotized heterocyclic amine 444 differs also from the well

Scheme 126

Scheme 127

established behavior of acyclic enaminones toward arenediazonium salts, where formation of 2-heteroylhydrazonopropanal 445 has not been observed, but instead a biscoupling product 448 has been obtained. This product is assumed to be formed *via* initial Japp Klingemann type cleavage of the dimethylaminomethylene moiety in 95b, yielding 446 that cyclizes into 447 and then couples further with 444 to yield a hydrazone that cyclizes into the final isolable product 448 (Scheme 122).<sup>25</sup>

478

Multi-component reaction of sulphone **449**, 3-aminotriazole **405** and dimethylformamide–dimethylacetal (DMF–DMA) in DMF under microwave irradiation at 150 °C for 10 min. Afforded compound **450** rather than its isomeric structure **451** (Scheme 110). The conformation of compound **450** was established on the bases of spectral data (MS, IR, <sup>1</sup>H NMR) and elemental analyses.

Furthermore, alternative synthesis of compound 452 was achieved *via* condensation of sulphone 1 with dimethylformamide dimethylacetal (DMF-DMF) to give compound 452, and treatment of the product 5 with 3-aminotriazole 305 under the same reaction condition to yield authentic product 450 (Scheme 123).<sup>151</sup>

With this result in hand, the scope of such multi-component protocol were expanded and derivatives 453–457 were synthesized; sulphone 452 and dimethylformamide–dimethylacetal (DMF–DMA), under the same reaction conditions, afforded, in each case, the corresponding heterocyclic ring systems 458–462, respectively (Scheme 124).<sup>151</sup>

To account for the formation of the products **450** and **458–462**, it was suggested that the studied reactions started with Michael-type addition of the exocyclic amino group of each the amines used to the activated double bond of 5 followed by *in* 

$$\begin{array}{c} O \\ Ph \\ & \\ 1d \\ & \\ R_1 = Me, Ph, 4-MeOC_6H_4 \\ \end{array} \qquad \begin{array}{c} NH_2 \\ Mg-Al \ hydrotalcite \\ R \\ & \\ \end{array} \qquad \begin{array}{c} N \\ N \\ N \\ \end{array}$$

Scheme 128

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16 new compounds up to 80% yield

## Scheme 129

$$\begin{array}{c} O \\ R \\ \hline \\ \mathbf{1d.f, g,p,q} \end{array} \begin{array}{c} H_2N \\ NC \\ \hline \\ \mathbf{486} \\ CN \\ \hline \\ \mathbf{487a-f} \end{array} \begin{array}{c} NC \\ CN \\ \hline \\ \mathbf{487a-f} \\ \end{array}$$

Scheme 130

Scheme 131

situ tandem elimination of dimethylamine and dehydrative cyclization (Scheme 125).151

Also, the reaction of enaminone 465 with active methylene under microwave irradiation. Reaction of enaminone 465 with active methylene derivatives 466-469 gave in glacial acetic acid in the presence of ammonium acetate gave poly-heterocyclic ring systems 470-473, respectively (Scheme 126). The structure of the products was assigned based on the spectral data and elemental analyses.152

The reactivity of the enaminone 474a or the morpholinyl derivative 474b towards some heterocyclic amines were also

examined. Thus, reaction of 474a or 474b with 5-amino-1Hpyrazole-4-carbonitrile 453 in acetic acid under reflux yielded the respective pyrazolo[1,5-a]pyrimidine derivative 475 (Scheme 114). Similar treatment of 474a or 474b with 5-amino-1,2,4triazole (405),5-amino-1*H*-tetrazole (455),2-aminobenzimidazole (451) and 6-amino-2-thioxo-2,3-dihydropyrimidin-4(1H)-one (479) under the same reaction conditions afforded the respective 1,2,4-triazolo[1,5-a]pyrimidine 441, tetrazolo[1,5-a]pyrimidine 477, benzimidazo[1,2-a]pyrimidine 478 and 2,3-dihydropyrido[2,3-d]pyrimidinone 480 derivatives, respectively (Scheme 127).51

The catalytic activity of the as-synthesized and activated hydrotalcites towards the aza-Michael addition reaction was evaluated. Thus, the reaction of an enaminone 1d with the 5amino-1H-pyrazole derivatives 481a-c in the presence hydrotalcite catalysts was carried out without solvent under microwave irradiation, to obtain only one isolable product in each case, (as examined by TLC) which were identified as pyrazolo [1,5-a]pyrimidine derivatives 482a-c (Scheme 128). 153

An environmentally benign multicomponent synthetic method has been realized for the diastereoselective construction of fused tetrahydropyridines 485 from reaction of enaminone 1b,d,f,g, o-aminophenol 483, and cinnamaldehyde 484 in the presence of lactic acid in water-ethanol media (Scheme 129).154

A mixture of enaminone 1d,g,p,q and 3-amino-2-cyanopent-2-enedinitrile (486) in AcOH/NH4OAc was heated under reflux for 2 h to give 7-amino-5-oxo-2-(thienyl)-5,6-dihydro-1,6naphthyridine-8-carbonitriles 487a-d (Scheme 130).155

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The reaction of enaminone 1t with 6-amino-2-thioxo-(1H)pyrimidin-4-one (489) produced 5-(pyridin-4-yl)-2-thioxo-2,3dihydropyrido[2,3-d]pyrimidin-4(1H)-one (490) (Scheme 2). Treatment of thione 490 with hydrazonoyl halides 491a-h in dioxane, in the presence of triethylamine under reflux gave in each case a single product consistent with structure 492 (Scheme 2) based on spectroscopic data (IR, <sup>1</sup>H NMR and MS) and elemental analyses (Scheme 131).16

## 3. Conclusions and future directions

The data considered in this review clearly demonstrate the high synthetic applications of N,N-dimethylenamino ketones in the preparation of acyclic and carbocyclic compounds as well as a broad range of heterocyclic and fused heterocyclic derivatives. Most importantly, these N,N-dimethyl analogues have proven to be of biological interest and provide an access to new class of biologically active heterocyclic compounds for biomedical screening. Presently, the great interest of researchers worldwide in such ketones is confirmed by the fact that most of the articles cited in this review are dated in the last two decades. We are sure that they will generate new and innovative applications with future generations of compounds, and thus providing a valuable adjunct to therapy. Further studies in this direction are currently underway and the results of this research will be reported elsewhere.

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