

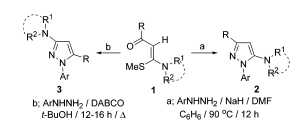
Highly Regioselective Synthesis of 1-Aryl-3 (or 5)-alkyl/aryl-5 (or 3)-(N-cycloamino)pyrazoles

S. Peruncheralathan,[†] A. K. Yadav,[†] H. Ila,^{*,†} and H. Junjappa^{*,‡}

Department of Chemistry, Indian Institute of Technology, Kanpur 208016, India, and BioOrganics and Applied Materials Pvt. Ltd., Peenya, Bangalore 560058, India

hila@iitk.ac.in

Received July 16, 2005

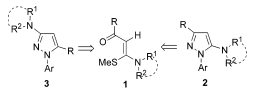


An efficient highly regioselective protocol for the synthesis of isomeric 1,3-diaryl (or 1-aryl-3-alkyl) and 1,5-diaryl (or 1-aryl-5-alkyl)-5 (or 3)-(*N*-cycloamino)pyrazoles has been reported by cyclocondensation of common α -oxoketene *N*,*S*-acetal precursors with arylhydrazines by variation of reaction conditions.

The 1-N-arylpyrazole ring system represents an important heterocyclic template that has attracted considerable interest because of its long history of application in pharmaceutical and agrochemical industry.¹ Numerous compounds containing 1-N-arylpyrazole moiety have been shown to exhibit antihyperglycemic, analgesic, antiinflammatory, sedative, and hypnotic activities.¹⁻³ Some of these compounds have emerged as potent and selective γ-aminobutyric acid (GABA)-gated chloride channel antagonists,² novel ligands for estrogen receptors,³ and agrochemicals of economic importance. One of the most important methods for the synthesis of substituted 1-Narylpyrazoles involves cyclocondensation of 1,3-dicarbonyl compounds and their equivalent 1,3-dienophilic synthons such as propargylic ketone, β -dialkylamino/alkoxy/ chloroketones with arylhydrazines.^{2b,c,3} However the appealing generality of this method is somewhat vitiated as a result of the frequent formation of regioisomeric mix-

tures of unsymmetrical pyrazoles in these reactions.^{3,4}Several elegant methods for the regioselective synthesis of unsymmetrically substituted 1-arylpyrazoles have been developed in recent years;^{1c,d,2} however despite their promising potential, these methods have limited applications in terms of generality and offer only little improvement over classical phenylhydrazine- β -diketone route to this class of compounds. Our own interest in addressing this regiochemistry issue stems from our ongoing research program utilizing α -oxoketene dithioacetals as versatile 1,3-electrophilic building blocks for regiospecific synthesis of substituted and condensed five- and sixmembered heterocycles and aromatic compounds.⁵⁻⁷ During the course of these studies, we became interested in probing the reaction of α -oxoketene *N*,*S*-acetals with an unsymmetrical binucleophile such as phenylhydrazine with a view to achieve synthesis of both 5- and 3-amino-1-arylpyrazoles in highly regiocontrolled fashion by variation of reaction conditions (Chart 1). Although several





5-alkyl/arylamino-1-arylpyrazoles have found applications as pharmaceuticals and agrochemical agents exhibiting a range of biological activities, $^{2,8-10}$ only a few scattered reports are available on the synthesis of 1-aryl-3-(or 5)-*N*,*N*-disubstituted aminopyrazoles, which are not well represented in the literature.¹⁰ The reported methods for the synthesis of 5-alkyl/arylaminoyrazoles require either harsh reaction conditions^{9d,e} or are limited only to a defined set of precursors.^{2,9} In a recent report,¹⁰ a few of the substituted 1-aryl-5-(*N*,*N*-disubstituted)aminopy-

10.1021/jo051478u CCC: \$30.25 © 2005 American Chemical Society Published on Web 10/19/2005

[†] Indian Institute of Technology.

[‡] BioOrganics and Applied Materials Pvt. Ltd.

⁽¹⁾ Reviews: (a) Elguero, J. In Comprehensive Heterocyclic Chemistry; Katritzky, A. R., Rees, C. W., Potts, K. T., Eds.; Pergamon Press: Oxford, 1984; Vol. 5, p 167. (b) Elguero, J. In Comprehensive Heterocyclic Chemistry II; Katritzky, A. R., Rees, C. W., Scriven, E. F. V., Eds.; Pergamon Press: Oxford, 1996; Vol. 3, p 1. (c) Kost, A. N.; Grandberg, I. I. In Advances in Heterocyclic Chemistry; Katritzky, A. R., Boulton, A. J. Eds.; Academic Press: New York, 1966; Vol. 6, p 347. (d) Lee, K. Y.; Kim, J. M.; Kim, J. N. Tetrahedron Lett. **2003**, 44, 6737 and references therein.

⁽²⁾ Sakya, S. M.; Rast, B. Tetrahedron Lett. 2003, 44, 7629 and references therein.

⁽³⁾ Huang, Y. R.; Katzenellenbogen, J. A. Org. Lett. 2000, 2, 2833 and references therein.

^{(4) (}a) Murray, W.; Wachter, M.; Barton, D.; Forero-Kelly, Y. Synthesis **1991**, 18. (b) Kashima, C.; Harada, H.; Kita, I.; Fukuchi, I.; Hosomi, A. Synthesis **1994**, 61. (c) Spivey, A. C.; Diaper, C. M.; Adams, H.; Rudge, A. J. J. Org. Chem. **2000**, 65, 5253.

⁽⁵⁾ Reviews: (a) Junjappa, H.; Ila, H.; Asokan, C. V. Tetrahedron **1990**, 46, 5423. (b) Ila, H.; Junjappa, H.; Mohanta, P. K. Progress in Heterocyclic Chemistry; Gribble, G. W., Gilchrist, T. L., Eds.; Pergamon: New York, 2001; Vol. 13, Chapter 1, p 1. (c) Ila, H.; Junjappa, H.; Barun, O. J. Organomet. Chem. **2001**, 624, 34.

⁽⁶⁾ Purkayastha, M. L.; Ila, H.; Junjappa, H. Synthesis 1989, 21.
(7) (a) Panda, K.; Suresh, J. R.; Ila, H.; Junjappa, H. J. Org. Chem.
2003, 68, 3498. (b) Peruncheralathan, S.; Khan, T. A.; Ila, H.; Junjappa, H. Tetrahedron 2004, 60, 3457. (c) Nandi, S.; Panda, K.; Suresh, J. R.; Ila, H.; Junjappa, H. Tetrahedron 2004, 60, 3663.

^{(8) (}a) Tang, J.; Shewchuk, L. M.; Sato, H.; Hasegawa, M.; Washio, Y.; Nishigaki, N. *Bioorg. Med. Chem. Lett.* **2003**, *13*, 2985. (b) Braibante, M. E. F.; Braibante, H. T. S.; da Roza, J. K.; Henriques, D. M. *Synthesis* **2003**, 1160. (c) Schenone, S.; Bruno, O.; Ranise, A.; Bondavalli, F.; D'Amico, M.; Filippelli, W.; Falcone, G.; De Novellis, V. *Il Farmaco* **1995**, *50*, 179. (d) Cooper, C. B.; Helal, C. J.; Sanner, M. A.; Wagner, T. T. PCT WO 18346 A1, 2002.

^{(9) (}a) Ansel, J. E.; El Kaim, L.; Gadras, A.; Grimaud, L.; Jana, N. K. *Tetrahedron Lett.* **2002**, 43, 8319. (b) Atlan, V.; Buron, C.; El Ka×c0m, L. *Synlett* **2000**, 489. (c) Palacios, F.; Aparicio, D.; de los Santos, J. M. *Tetrahedron* **1996**, 52, 4123. (d) Moreno-Manãs, M.; Sebastián, R. M.; Vallribera, A.; Carini, F. *Synthesis* **1999**, 157. (e) Abdel-Rahman, R. M.; Seada, M.; Fawzy, M.; El-Baz, I. *Pharmazie* **1994**, 49, 729.

⁽¹⁰⁾ Dodd, D. S.; Martinez, R. L. Tetrahedron Lett. 2004, 45, 4265.

SCHEME 1^a

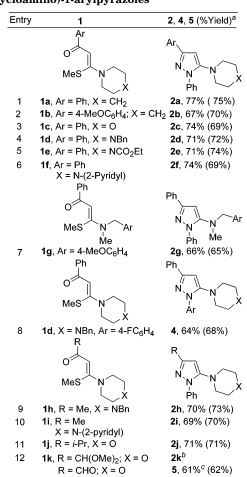


 a Reagents and conditions: (a) ArNHNH2/NaH/DMF/C6H6/90 °C/12 h. (b) ArNHNH2/t-BuOK/t-BuOH/12 h/ Δ .

razoles have been synthesized by treatment of a preformed N-arylhydrazones from appropriate β -ketoamides with Lawesson reagent in a one-pot operation. However this protocol has not been extended for the synthesis of 5-(N-cycloamino)pyrazoles, whereas the corresponding 3-(N-cycloamino)pyrazoles to our knowledge are unknown in the literature. Herein we describe our successful results on regioselective synthesis of substituted 1-aryl-5 (or 3)-N-(cycloamino)pyrazoles from common N,S-acetal precursors.

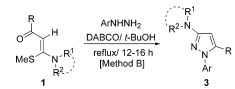
Earlier work from this laboratory has demonstrated synthetic application of α -oxoketene *N*,*S*-acetals as useful three-carbon 1,3-electrophiles in their reactions with symmetrical (hydrazine^{11a} and guanidine^{11c}) and unsymmetrical (hydroxylamine,^{11b} cyanoacetamide,^{11d} and lithioaminocrotononitrile^{11e}) binucleophiles, yielding the corresponding primary or secondary amino-substituted fiveand six-membered heterocycles in highly regioselective fashion.¹² In continuation of these studies, we examined the reaction of ketene N,S-acetals 1a with unsymmetrical binucleophile such as phenylhydrazine under varying conditions (mild basic and neutral). In most of the cases, the corresponding 5-(*N*-piperidino)-1,3-diphenylpyrazole (2a) was isolated in varying yields, whereas under drastic conditions (neat heating at 120 °C), complex mixtures of products were formed. Under optimized conditions, 2a was obtained in 77% yield when 1a was reacted with phenylhydrazine (1.2 equiv) in the presence of sodium hydride in DMF/C₆H₆ at 90 °C (Scheme 1). These reaction conditions were also found to be optimal for the synthesis of 3-(4-methoxyphenyl)-5-(N-piperidino)- and the other 5-(N-cycloamino)- or the acyclic 5-(N-methylbenzylamino)-1,3-diphenylpyrazoles 2b-g in overall high yields (Table 1, entries 2-7). Cycloannulation of the N,S-acetal 1d with 4-fluorophenylhydrazine also yielded the corresponding 1-N-(4-fluorophenyl)-5-N-(piperazino)pyrazole (4) in 64% yield (entry 8). The heterocyclization reaction was found to be equally facile with *N*.S-acetals **1h**-**j** with aliphatic acyl groups yielding the corresponding 3-alkyl-5-(N-cycloamino)pyrazoles 2h-j in good yields (entries 9-11). Similarly the 3-formyl-1-phenyl-5-N-(morphilino)pyrazole 5 was obtained in 61% yield by in situ hydrolysis of the 3-bis(methoxymethyl)pyrazole 2k obtained from 1k under identical conditions (entry 12). During the course of this work, we further observed that use of potassium *tert*-butoxide as base in refluxing *tert*-butyl

TABLE 1.	Synthesis of
5-(N-Cycloamino)-1-arylpyrazoles	



^{*a*} Yields obtained from *t*-BuOK are in parentheses. ^{*b*} Not isolated. ^{*c*} Obtained by in situ acidic hydrolysis of **2k**.

SCHEME 2



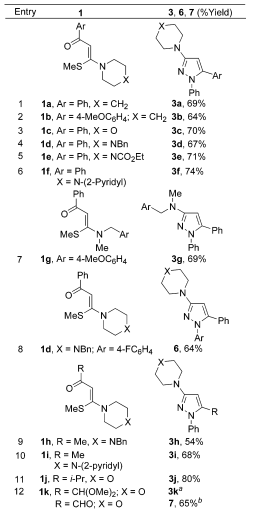
alcohol instead of sodium hydride also gave the 5-aminopyrazoles $2\mathbf{a} - \mathbf{k}$ in comparable yields as shown in the Table 1.

After successfully establishing the reaction conditions for obtention of 1-aryl-5-(*N*-cycloamino)pyrazoles **2**, we were further intrigued to develop a general regiocontrolled route for the 1-aryl-5-aryl/alkyl-3-aminopyrazoles (**3**) from the common α -oxoketene *N*,*S*-acetal precursors **1**. Interestingly, we could achieve this goal after several unsuccessful experiments, when **1a** was reacted with phenylhydrazine in the presence of a weaker base such as DABCO (1.2 equiv) furnishing only 3-(*N*-piperidino)-1,5-diphenylpyrazole (**3a**) exclusively in 69% yield with no trace of regioisomeric pyrazole **2a** (Scheme 2, Table 2, entry 1, Method B). The generality of these reaction conditions is evident from the facile synthesis of other isomeric 3-amino-1-aryl-5-aryl/alkylpyrazoles **3b**-**k** parallel to their 5-amino partners **2b**-**k** in good to excellent

^{(11) (}a) Chauhan, S. M. S.; Junjappa, H. Synthesis **1975**, 798. (b) Rahman, A.; Vishwakarma, J. N.; Yadav, R. D.; Ila, H.; Junjappa, H. Synthesis **1984**, 247. (c) Kumar, A.; Aggarwal, V.; Ila, H.; Junjappa, H. Synthesis **1980**, 748. (d) Aggarwal, V.; Singh, G.; Ila, H.; Junjappa, H. Synthesis **1982**, 214. (e) Satyanarayana, J.; Ila, H.; Junjappa, H. Synthesis **1991**, 889.

⁽¹²⁾ Mahata, P. K.; Venkatsh, C.; Syam Kumar, U. K.; Ila, H.; Junjappa, H. J. Org. Chem. **2003**, 68, 3966.

TABLE 2. Synthesis of 3-(N-Cycloamino)-1-arylpyrazoles



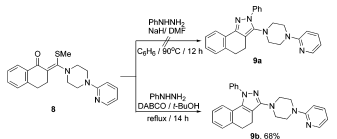
^a Not isolated. ^b Obtained by in situ acidic hydrolysis of **3k**.

yields as shown in Table 2. The regiochemistry of 5- and 3-(N-cycloamino)pyrazoles 2 and 3 was established from the X-ray crystallographic data of one of the regioisomeric pairs 2b and 3b (Figures 1 and 2, Supporting Information).

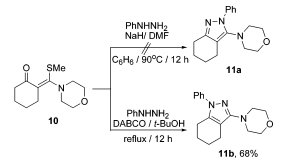
To further extend the scope of this reaction, the N.Sacetals 8 and 10 derived from cyclic ketones such as tetralone and cyclohexanone, respectively, were reacted with phenylhydrazine under earlier described conditions (Methods A and B) with a view to synthesize regioisomeric 3,4- (9a, 11a) and 4,5- (9b, 11b) annulated 3- (or 5-) aminopyrazoles (Schemes 3 and 4). However, these reactions met with only partial success, yielding the corresponding 3-(N-cycloamino)-4,5-annulated pyrazoles 9b and 11b in good yields under DABCO-catalyzed conditions (Method B), whereas the reaction of either 8 or 10 with phenylhydrazine in the presence of sodium hydride/DMF or potassium tert-butoxide/tert-butyl alcohol (Method A) afforded only intractable product mixtures.

The probable mechanism for the formation of the two regioisomeric pyrazoles 2 and 3 from N,S-acetal 1 in the presence of different bases is shown in Scheme 5. Thus in the presence of a stronger base like sodium hydride

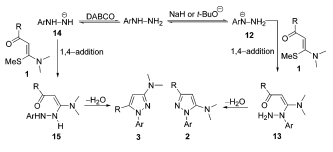
SCHEME 3



SCHEME 4



SCHEME 5



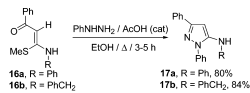
or potassium *tert*-butoxide, the anion **12** is formed by abstraction of more acidic proton of phenylhydrazine.^{1a,13} The anion 12 adds to α -oxoketene dithioacetals 1 in 1,4fashion to give intermediate adduct 13, which on intramolecular cyclization affords 5-(cycloamino)pyrazole 2. In the presence of a weaker and sterically crowded base such as DABCO, phenylhydrazine undergoes abstraction of the less hindered NH₂ proton, resulting in the formation of the anion 14, which undergoes 1,4-additionelimination with N,S-acetal 1 to give the intermediate adduct 15.14 Subsequent intramolecular cyclization of adduct 15 affords the 3-(cycloamino)pyrazole 3. Our attempts to trap either of these adducts 13 or 15 were not successful.

The N,S-acetals **16a,b** carrying a primary amino group were next subjected to cyclization with phenylhydrazine under varying conditions (Scheme 6). Best results were obtained when the reaction of 16a with phenylhydrazine was effected in the presence of a catalytic amount of acetic acid, yielding the corresponding 5-anilino-1,3diphenylpyrazole **17a** exclusively in 80% yield. The N,Sacetal from benzylamine 16b also furnished the 5-aminopyrazole 17b in high yield. Our attempts to get

^{(13) (}a) Coispeau, G.; Elguero, J. Bull. Soc. Chim. Fr. 1970, 2717.

 ⁽b) Elguero, J.; Marzin, C. Bull. Soc. Chim. Fr. 1973, 3401.
 (14) Robinson, M. J. T.; Kenny, P. W. Tetrahedron 1987, 43, 4043.

SCHEME 6



3-anilino-5-phenylpyrazole from **16a** under varying conditions were, however, not successful.

In summary, we have developed a highly efficient protocol for the synthesis of 1,3-diaryl (or 1-aryl-3-alkyl) and 1,5-diaryl (or 1-aryl-5-alkyl)-5 (or 3)-(*N*-cycloamino)pyrazoles from common α -oxoketene-*N*,*S*-acetal precursors in a highly regiocontrolled fashion. Our efforts to extend this regioselective protocol to a wide range of *N*,*S*acetals derived from various cyclic/heterocyclic ketones and primary amines is currently under progress.

Experimental Section

General details are described in Supporting Information. All known N,S-acetals **1a**-**k**, **8**, **10**, and **16a**-**b** were prepared by the earlier reported procedure.^{11b,12}

General Procedure for Preparation of 1-Aryl-3-aryl/ alkyl-5-(N-cycloamino)pyrazoles 2, 4. A solution of the respective N,S-acetal 1 (5 mmol) and arylhydrazine (6 mmol) in benzene (50 mL) was added to a suspension of NaH (0.24 g, 6 mmol) in DMF (10 mL) at room temperature over a period of 0.5 h. The reaction mixture was heated at 90 °C with constant stirring for 12 h (monitored by TLC), poured after cooling into saturated NH₄Cl solution (50 mL), and extracted with benzene (2×25 mL). The combined benzene layer was washed with H₂O (3×50 mL) and brine (1×50 mL), dried over Na₂SO₄, and distilled under reduced pressure to give pyrazoles 2 or 4, which were purified by column chromatography over silica gel using hexane/EtOAc (10:1) as eluent.

1,3-Diphenyl-5-(*N*-**piperidino**)-**1***H*-**pyrazole** (2a). Yield 77% (1.17 g); pale yellow solid; mp 80-81 °C; $R_f 0.80$ (9.5:0.5

hexanes–EtOAc). IR (KBr): 3060, 2944, 1593, 1549, 1498, 1450, 1421, 1305, 1256, 1201 cm⁻¹. ¹H NMR (400 MHz, CDCl₃): δ 7.84 (d, J = 7.6 Hz, 2H), 7.77 (d, J = 7.4 Hz, 2H), 7.35 (t, J = 7.6 Hz, 2H), 7.37 (t, J = 7.6 Hz, 2H), 7.16–7.23 (m, 2H), 6.09 (s, 1H), 2.79 (t, J = 5.4 Hz, 4H), 1.50–1.56 (m, 4H), 1.44–1.47 (m, 2H). ¹³C NMR (100 MHz, CDCl₃): δ 153.8, 150.7, 140.4, 133.5, 128.7, 128.4, 127.6, 126.2, 125.4, 122.5, 91.4, 52.3, 25.4, 23.8. MS (m/z, %): 304 (M + 1, 100); 303 (M⁺, 90). Anal. Calcd for C₂₀H₂₁N₃ (303.40): C, 79.17; H, 6.98; N, 13.85. Found: C, 79.31; H, 6.79; N, 14.01.

General Procedure for Preparation of 1-Aryl-5-aryl/ alkyl-3-(N-cycloamino)pyrazoles 3, 6. A solution of respective N,S-acetal 1 (5 mmol), arylhydrazine (6 mmol), and DABCO (0.67 g, 6 mmol) in 50 mL of t-BuOH was refluxed for 12-16 h with constant stirring, the reaction being monitored by TLC. The reaction mixture was concentrated under reduced pressure and poured into ice-cold water, extracted with DCM (3×50 mL), washed with H₂O (2×50 mL) and brine (1×50 mL), and dried over Na₂SO₄. The solvent was evaporated under vacuum to give pyrazoles 3 or 6, which were purified by column chromatography over silica gel using hexane/EtOAc (10:1) as eluent.

1,5-Diphenyl-3-(*N*-piperidino)-1*H*-pyrazole (3a). Yield 69% (1.05 g); brown solid; mp 105–106 °C; R_f 0.71 (9.5:0.5 hexanes–EtOAc). IR (KBr): 3056, 2930, 1593, 1552, 1512, 1445, 1378, 1348, 1293, 1258, 1236 cm⁻¹. ¹H NMR (400 MHz, CDCl₃): δ 7.04–7.18 (m, 10H), 5.88 (s, 1H), 3.19 (t, J = 5.7 Hz, 4H), 1.58–1.63 (m, 4H), 1.46–1.52 (m, 2H). ¹³C NMR (100 MHz, CDCl₃): δ 159.9, 143.6, 140.3, 131.1, 128.57, 128.54, 128.2, 127.9, 126.1, 124.7, 95.1, 48.8, 25.3, 24.3. MS (*m*/*z*, %): 304 (M + 1, 90); 303 (M⁺, 100). Anal. Calcd for C₂₀H₂₁N₃ (303.40): C, 79.17; H, 6.98; N, 13.85. Found: C, 79.27; H, 7.05; N, 13.63.

Acknowledgment. Financial assistance under CSIR and DST projects is acknowledged. S.P. thanks CSIR for research associateship.

Supporting Information Available: ¹H and ¹³C NMR spectral data for compounds and crystallographic data in CIF format. This material is available free of charge via the Internet at http://pubs.acs.org.

JO051478U