

Table II. Spectral Studies of the Hydrazidoyl Halides

compd	¹ H NMR		IR (KBr) ^b		UV $\lambda_{\max}$, nm
	δ^a	assignment (no. of protons)	ν , cm ⁻¹	assignment	
6a	2.57, s	methyl (3)	1430, 1130	C-N	364
	7.95, 8.07, s; 8.40, 8.57, 9.07, d; $J = 3.0$ Hz	aromatic (3)	640	C-Cl	
	11.50, s	imino (1)			
6b			1435, 1140	C-N	370
6c			645	C-Cl	
			1420, 1225	C-N	
6d			730	C-Cl	
	1.23, s	methyl (3)	2990, 2940	C-H aliphatic	
	1.34, s	methyl (3)	1620, 1490	C=C aromatic	
6e	2.70-3.15, m	methine (1)	1420, 1140	C-N	280, 390
	7.16, 7.60, 7.77, s; 8.10, 8.29, 8.92, d; $J = 3.0$ Hz	aromatic (3)	645	C-Cl	
	11.15, s	imino (1)			
6f	1.71, d, $J = 6.0$ Hz	methyl (3)			
	4.18-4.50, m	olefinic (1)			
	4.63-4.86, m	olefinic (1)			
6g	7.23-9.06, m	aromatic (3)			
	11.13, s	imino (1)			
			1590, 1510, 1450	C-C aromatic	
6h			1428, 1150	C-N	
			735	C-Cl	
			3110, 830, 860	C-H aromatic	
6i			2960	C-H olefinic	
			1610	C=C olefinic	
			1580, 1510, 1450	C=C aromatic	
6j			1430, 1130	C-N	
			740	C-Cl	
			2985, 2930	C-H aliphatic	
6k			1610, 1485	C=C aromatic	
			1410, 1160	C-N	
			670	C-Cl	
6l	0.65-2.99, m	methyl (9), methylene (4)			260, 850
	4.06-4.40, m	methine (1)			
	5.00-5.21, m	methine (1)			
6m	7.36-8.25, m	aromatic (3)			370
	11.37, s	imino (1)			
	7.41-9.38, m	aromatic (8)			
6n	11.94, s	imino (1)			
			1620, 1560, 1490	C=C aromatic	
			1428, 1160	C-N	
6o			2940	C-H methoxy	
			3430	O-H	
			1580, 1510, 1480	C=C aromatic	
6p			1430, 1130	C-N	
			1670	C=O	
			1610, 1590, 1510	C=C aromatic	
6q			1410, 1315	C-H deformation	
			1450, 1165	C-N	
6r	2.52, s	acetyl (3)			
	7.51-8.40, m	aromatic (4)			
	11.40, s	imino (1)			
6s			2910	C-H aliphatic	
			1700	C=O	
			1600, 1565, 1500	C=C aromatic	
6t	1.42, t, $J = 7.2$ Hz	methyl (3)	1710	C=O	
	4.41, q, $J = 7.2$ Hz	methylene (2)	1610, 1540, 1490	C=C aromatic	
	7.27-8.64, m	aromatic (4)	3000, 2980, 2900	C-H aliphatic	
6u	11.36, s	imino (1)	600	C-Br	

^a s = singlet; d = doublet; t = triplet; q = quartet; m = multiplet. ^b In general, IR spectra show N-H bonds at 3290-3260 cm⁻¹ and C=N bonds at 1610-1590 cm⁻¹, and NO₂ bonds show absorptions at 1530-1520 and 1345-1340 cm⁻¹.

another crop of hydrazidoyl chloride. Products were obtained in ~70-90% yields.

Hydrazidoyl Chlorides 6f,i,j. A mixture of hydrazone (1.0 g) in glacial acetic acid (20.0 mL) was saturated with dry chlorine gas. In the case of hydrazone 5f, the solid went into solution and an orange solution was obtained which upon cooling gradually at room temperature afforded pale yellow crystals of compound 6f, which was then filtered and recrystallized from acetic acid as bright shining crystals.

The solution of hydrazones 5i,j was warmed on a water bath with continuous passage of chlorine gas through the hot solution. After 0.5 h a clear solution was obtained, which upon cooling started to deposit crystals of hydrazidoyl chloride. The products were recrystallized from acetic acid (Table I).

Hydrazidoyl Bromides 6k-p. The dinitrophenylhydrazones (DNPs) 5k-p (0.01 mol) were suspended in glacial acetic acid (10 mL) containing 4% acetic anhydride and vigorously stirred, while bromine (0.03 mol) in 2 mL of the same solvent was

added dropwise with constant stirring. The hydrazones immediately dissolved, and the bromides precipitated out in 80-85% yields after 10-30 min, the time depending mainly on the reactivity of the DNPs. In the case of hydrazidoyl bromides **6n-p**, usually a 4-fold excess of bromine was used, and, after 4 h of stirring, the solution was warmed on a sand bath. The hydrazidoyl bromides thus obtained were separated from the solution, washed with a little acetic acid and then with petroleum ether, and recrystallized from appropriate solvents (Table I).

Hydrazidoyl Bromides 6q-t. Hydrazones **5q-t** (0.04 mol) were added to a mixture of 200 mL of glacial acetic acid and 120 mL of acetic anhydride containing sodium acetate (27 g), and the temperature was lowered to 0-5 °C by means of an ice-salt bath. To this was added over a 1-h period bromine (0.04 mL) dissolved in 5 mL of acetic acid. After additional stirring for 3 h, the solution was poured into 2 L of water. A yellow product so obtained was stirred for some time and then left overnight at room temperature. The product was filtered, washed with water, and dried under vacuum. The products so obtained were generally crystallized from alcohol or aqueous alcohol.

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Synthesis of 2,2-Disubstituted 5-Cyanocyclopentanones and Comparison of Their Enol Contents with the Corresponding Cyclohexanones

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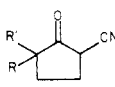
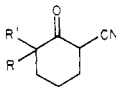
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The enol contents of 14 2,2-disubstituted 5-cyanocyclopentanones and the analogous 2,2-disubstituted 6-cyanocyclohexanones are reported. The Brown-Brewster-Schechter prediction of less enolization for the five-membered allcyclics vs. the six-membered analogues is valid for all but three cases (Ph, *i*-C₃H₇; Ph, CH₂Ph; Ph, *n*-C₄H₉).

We previously reported (2) the development of a procedure for determining the percent enol for 2-cyanocyclohexanones by an infrared technique. Using this method for 14 compounds in the cyclohexanone series and 7 analogous compounds, which were available to us, in the cyclopentanone series (3, 4) gave data which led to a correlation attempt via the Brown-Brewster-Schechter proposal (5) to exo/endo bonds in five- or six-membered rings. The rule predicts less enolization in the five-membered compounds.

We now report the synthesis of the remaining seven 2,2-disubstituted 5-cyanocyclopentanones and their enol contents as well as the necessary 1-cyano-2-methoxy-3,3-disubstituted cyclopentenones (H, H; Me, Me; Ph, *n*-C₅H₁₁; Ph, Ph) employed as conjugated nitrile standards.

Table I. Percent Enol of 2-Cyanocyclohexanones and 2-Cyanocyclopentanones in Dioxane

substituents		mol % enol	
R	R'		
H	H	1	37
Et	Et	2	18
Me	Me	4	20
C ₆ H ₅	<i>i</i> -butyl	6	6
C ₆ H ₅	<i>i</i> -propyl	6	5
C ₆ H ₅	Et	6	10
C ₆ H ₅	cyclohexyl	6	6
C ₆ H ₅	<i>n</i> -amyl	7	10
C ₆ H ₅	CH ₂ C ₆ H ₅	8	4
C ₆ H ₅	<i>n</i> -propyl	8	10
C ₆ H ₅	<i>n</i> -butyl	8	5
C ₆ H ₅	C ₆ H ₅	9	24
C ₆ H ₅	(CH ₂) ₂ C ₆ H ₅	13	13
C ₆ H ₅	Me	13	19

Absorption of the keto nitriles in dioxane at 2210 cm⁻¹ (conjugated CN) gave the concentration of the enol form by reference to a calibration plot for a related 1-cyano-2-meth-