## SYNTHESIS OF 7, 12-DIHYDROPYRIDOPYRROLOINDOLES (AZAPYRIDODIINDOLES) via THE FISCHER INDOLE CYCLIZATION. A SEARCH FOR WATER SOLUBLE BENZODIAZEPINE RECEPrOR LIGANDS.

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Abstract- The thermally-induced Fischer indole cyclization of the **3**  acylindole 7 with various substituted hydrazinopyridines provides a simple route to the 7.12-dihydropyridopyrroloindoles **2,** 4 and **5,** respectively. In contrast, the corresponding acid-catalyzed process yielded only 4-amino- $\beta$ carboline 15. Moreover, attempts to prepare the 3-aza-analog 3, from reaction of 7 and 4-hydrazinopyridine **12** under thermal or acidic conditions, provided only 15. The difference between the reactivity of the 4-substituted pyridine **12,** in comparison to **11** or **13,** is discussed in terms of pK, values and intermediates in the Fischer indole cyclization.

During a search for new ligands with which to probe the topography of benzodiazepine receptors **(BzR),** the rigid planar **7.12-dihydropyrido[3,2-b:5,4-b'ldiindolc** 1 was discovered to exhibit high affinity for these receptors.<sup>1</sup> A wide variety of functional groups have been substituted for hydrogen at positions **-1,** -2, -3, -4 and -10 of the pyridodiindole nucleus to determine the effect on binding affinity,  $1.2$  During this work it was proposed to increase the water solubility of these pyridodiindoles and to, simultaneously, alter the electron density distribution of these important rigid pyridodiindole probes without perturbing their topography.

A simple means by which to achieve this goal was to replace C-H at positions -1, **-2,** -3 or -4 of 1 with a nitrogen atom. This provides the new heterocyclic ring system, **7,12-dihydropyridopyrroloindole** also referred to as the **aza-7,l2-dihydropyrido[3,2**  b:5,4-b'ldiindole, in reference to **1.3** The dihydrocbloride salt of this new heterocycle should demonstrate increased water solubility in comparison to **1.** Replacement of



carbon with nitrogen at positions -1 through -4 of 1 would alter the electron density distribution in ring E of the molecule (MNDO calculations).4 More importantly, in accord with a recent model of the pharmacophore<sup>5</sup> for the inverse agonist site at BzR, an electron rich pyridine nitrogen at position -4 (see 4) might well enhance the ability of this azapyridodiindole to interact with a hydrogen bond acceptor atom on the benzodiazepine receptor site, thereby enhancing inverse agonist activity.

The synthesis of azapyridodiindoles 2, 4 and 5 is based on chemistry previously developed in our laboratory<sup>1</sup> in combination with the Fischer indole cyclization, and is outlined in Scheme I for one of the azapyridodiindoles. The oxidation of 2-benzoyl-**1,2,3,4-tetrahydronorharmane** 6 with DDQ, according to published procedures,6,7 provides the 3-acylindole 7 in reasonable yield.<sup>1,7</sup> Treatment of 7 with 2hydrazinopyridine provides the hydrazone 8; however, acid-catalyzed Fischer indole cyclization **(via** 9) was not attempted at this stage. It is known that acid-catalyzed Fischer indolization of pyridylhydrazones either fails or affords low yields of the desired azapyridodiindoles.<sup>8,9</sup> This is due to deactivation of the pyridine ring by the inductive effect of the nitrogen atom and also by protonation of this pyridine nitrogen atom in the acidic media. The Fischer indole cyclization involves a [3,3] sigmatropic rearrangement, therefore the cyclization will take place thermally.<sup>10,11</sup> When the hydrazone 8 was heated. at 160°C for 9 h the Fischer indole cyclization took place, **via** intermediate 9, to provide the azapyridodiindole 10. In keeping with previous work in the diindole area,<sup>1</sup> hydrazine was added to the reaction mixture to cleave the henzamide group, while permitting the disproportionation of the **5,6-dihydropyridodiindole** to take place to provide 2 in 71% overall yield. It is not known whether hydrazine serves as the oxidizing agent during the disproportionation reaction, or whether some other agent (ie.  $O<sub>2</sub>$ ) is responsible for loss of hydrogen across the 5,6-dihydro bond of 2.



The route outlined in Scheme I serves as a general strategy for synthesis of three of the ring-E substituted azapyridodiindoles **(2, 4, 5)** as illustrated in Scheme 11. The 2 hydrazinopyridine **11** is commercially available, while 3-hydrazinopyridine **13** was prepared by the method of Binz and Rath.12 The required 4-hydrazinopyridine **12** was synthesized from 4-chloropyridine by the method of Mann  $et$   $al$ ,<sup>13</sup> The reaction of 7 with **11,** gave the **I-aza-7,12-dihydropyrido[3,2-b:5,4-b']diindole** 23 as illustrated in Scheme I, while heating 7 with 4-hydrazinopyridine **12,** in contrast, provided only **4**  amino- $\beta$ -carboline. On the other hand, treatment of acylindole 7 with the 3hydrazinopyridine **13** resulted in the formation of both 4-aza-7.12-dihydropyrido- [3,2-h:5,4-b'ldiindole 4 and **2-aza-7,12-dihydropyrido[3,2-b:5,4-b']diindole** 5, as expected, in view of the mechanism of the Fischer indole cyclization. $8-11,3$  The latter regioisomer **5** comprised the majority of the material. As anticipated, when **11** was reacted with 7 in refluxing ethanolic hydrogen chloride, the principal product was 4 amino- $\beta$ -carboline 15. This  $\beta$ -carboline derivative was identical to that prepared by an independent route.14 This result serves to confirm further the need to execute the thermally-induced Fischer indole cyclization in the case of hydrazinopyridines<sup>10,11</sup> rather than the use of the acid-catalyzed process.14 The reasons for the failure of **14** to

**Scheme II** 



undergo the Fischer-indole cyclization are several-fold. If the aromatic ring of the phenylhydrazine is substituted with deactivating groups, the reaction will yield **4**  amino- $\beta$ -carboline 15<sup>14</sup> in preference to the product of a Fischer-indole cyclization (Scheme 111). The cleavage of the nitrogen-nitrogen bond in 14 must occur in both



cases to provide either 15 or 3. In the case of 14; however, the nitrogen function of the 4-substituted pyridine (from 12) accepts most of the electron density, rather than permits attack on the  $\beta$ -carboline ring in a Fischer indole cyclization [see 9 (Scheme I)

for an example of a successful cyclization]. In essence, the pyridine nitrogen atom of 14 readily accepts electron density to promote nitrogen-nitrogen bond cleavage of the intermediate 14 to form 15 at the expense of the Fischer indole cyclization. In contrast, both 11 and 13 enter successfully into the Fischer indole cyclization principally because the developing negative charge on the leaving group  $(17c \text{ or } 18c)$  is not delocalized as effectively as it is on 4-aminopyridine 16c. This is supported by the  $pK_{a2}$  values for the three bases 16c (9.17), 17c (5.98) and 18c (6.86).15 Of the three compounds, **4**  aminopyridine 16c is the stronger base (pyridine nitrogen) because the electron density from the amino group is delocalized more effectively onto the pyridine nitrogen atom. This weakens the nitrogen-nitrogen bond of 14 more than would be expected for 17c or 18c (Scheme IV). The situation is more complicated than this, and the  $pK_{a1}$ 's of the substituted aminopyridines can be employed to further illustrate this point. Comparison of the  $pK_{a1}$  values for 16a and 17a clearly indicate that 17a is a stronger base than 16a with regard to proton loss from the aniline "like" nitrogen functions. This illustrates that electron density from the 4-amino function is more readily released and delocalized into the pyridine ring of 16a than into the corresponding ring of 17a. This would facilitate cleavage of the nitrogen-nitrogen bond in 14 (Scheme 111) by stablilization of the developing negative charge in comparison to a hydrazine

Scheme IV



generated from 13. Moreover, this electron release is supported by the  $pK_{a2}$  values for these aminopyridines (Scheme IV). Proton loss from the 4-aminopyridinium species 16b has a  $pK_{a2}$  of 9.17, while the value for 17b (5.98) is indicative of the higher electron density on the pyridine nitrogen atom of 16c relative to 17c.

Comparison of the  $pK_{a1}$  values between 16a and the 2-aminopyridine 18a cannot be employed here because of charge effects between the two closely disposed cationic functions of 18a; however, the ratio of the  $pK_{a2}$  values for 16b and 18b is similar to that for 16b and 17b. It is felt, therefore, that the same effect must be operating in the case of 11 (18c) as in 13c(17c). ţ,

In summary, the pyridine nitrogen atom of 4-aminopyridine 16c accepts charge density more readily than either 17c or 18c, consequently, scission of the nitrogen-nitrogen bond of 14 to expel 16c is more facile than the analogous process (from 11 **or** 13) to furnish 17c or 18c. As a result, the desired Fischer indole cyclization occurred between 7 and hydrazinopyridines 11 and 13, respectively, hut not in the case of 4 hydrazinopyridine 12.

As expected, the three azapyridodiindoles (2, 4 and **5)** demonstrate enhanced water solubility in comparison to 1, moreover, azapyridodiindoles 2 and 4 bind to benzodiazepine receptor with potent affinity (see Table). The biological activity of these compounds will be reported elsewhere.

Table In vitro Binding Affinity For Benzodiazepine Receptors



ACKNOWLEDGEMENT: We wish to thank MMH **(MH** 36644) and NIH (NS 22287) for generous financial support. We also wish to acknowledge Mr. Frank Laib and Mr. Keith Krumnow for technical assistance and Ms. Anjn Gupta for preparation of this manuscript.

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## $3<sub>1</sub>$ Azapyridodiindoles

- 2: **7,12-Dihydropyrid0~3",2":4',5'lpyrrolo[2',3':5,6]pyrido[3,4-b]indole.**  mp >300°C; 'H NMR (DMSO-d6, 250 mHz) 6 7.51 (d, lH, J=5.2 Hz), 7.54 (d, IH, J=5.2 Hz), 7.80 (t, 1H, J=8.1 Hz), 7.87 (t, 1H, J=8.3 Hz), 8.71 (d, 1H, J=1.6 Hz), 8.73 (d, lH, J=1.5 Hz), 9.10 (m, ZH), 9.31 (s, 1H).
- **4: 7,12-Dihydropyrido[2",3":4',5'lpyrrolo[2',3~:5,6lpyrido[3,4-b]indole**  mp >300°C; <sup>1</sup>H NMR (DMSO-d<sub>6</sub>, 250 mHz)  $\delta$  7.52 (t, 1H, J=7.3 Hz), 7.73 (t, 1H, J=6.3 Hz), 7.85 (m, 2H), 8.60 (d, 1H, J=7.95 Hz), 8.76 (d, 1H, J=5.3 Hz), 8.93 (d, lH, J=7.9 Hz), 9.20 (s, lH), 12.80 (s, lH), 13.65 (s, 1H)
- **5: 7,12-Dihydro[4",3":4',5'lpyrrolo[2',3':5,6lpyrido[3.4-b]indole**  mp >300°C; <sup>1</sup>H NMR (DMSO-d<sub>6</sub>, 250 mHz)  $\delta$  7.45 (t, 1H, J=7.5 Hz), 7.65 (t, 1H, J=7.5 Hz), 7.80 (d, 1H, J=8.0 Hz), 8.60 (d, 1H, J=7.3 Hz), 8.65 (d, 1H, J=7.2 Hz), 8.85 (d, lH, J=8.0 Hz), 9.15 (s, lH), 9.35 (s, lH), 12.60 (s, IH), 13.75 (s, 1H).
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**Received, 26th March,** 1988