HETEROCYCLES VIA ORTHO-LITHIATED BENZAMIDES

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Ortho-lithiated N,N-diethyl benzamides, readily generated by treatment of the corresponding amides with sec-BuLi or t -BuLi in THF or $Et_2O/TMEDA$ at -78° , are important new synthons $(d₁-reagents)$ for the regiospecific construction of highly substituted aromatic derivatives which are difficult to obtain by classical (usually electrophilic substitution) lithiated benzamides chemistry for oniversity of waterloo, waterloo, can
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phthalide and isochroman-1,3-dime heterocycles, phthalideisoquinoline alkaloids, anthraquinone natural products, polycyclic anrhraquinones and corresponding **PAH's,** heterocyclic benzoquinones, and ellipticine alkaloids.

The term directed metalation is defined as the deprotonation of an ${\mathfrak{sp}}^2$ -carbon α to a heteroatom-containing suhstituent on an aromatic or olefinic substrate. In and isochroman-1,3-dione heterocycles, phthalideisoquinoline alkaloids, anthraquinone
natural products, polycyclic anthraquinones and corresponding PAH's, heterocyclic
benzoquinones, and ellipticine alkaloids.
The term di The ortho directing group $2 \quad 1$) anchors the strong base (invariably an alkyllithium reagent) by coordination, 2) provides an inductive electron-withdrawing effect in the deprotonation step, and 3) stabilizes the ortho metalated intermediate by chelation. Systems in which the Z group bears acidic hydrogen require

Directed Metalation

 $Z = NMe_2$, $(CH_2)_n$ ^{NMe}₂, CONHMe, CSNHMe $n=1,2$ OMe, OCH_2 OMe, SO_2 NMe₂, SO_2 NHMe, CF_7 , F, CH=NR, NHCOR, OCH(Me)OEt, CH₂OH

 $Figure 1$ **Figure 1**

two equivalents of base and, by necessity, lead to dimetalated species. Gi_man^1 , and subsequently Hauser², in extensive pioneering efforts, laid the groundwork in this area and defined many of the groups which promote ortho metalation. The industrial development³ of organolithium bases such as $sec-BuLi$ and t-BuLi has provided a strong impetus for the exploitation and advancement of the original discovcries.

In recent years, the list of groups which promote ortha metalation has grown considerably (Fig. 1) and the early findings have been modernized and significantly extended.⁴ For example, Hauser's metalated secondary benzamides have been vigorously applied to the synthesis of heterocycles⁵ and natural products, 6 secondary thioamides have been successfully tested, \overline{z} the methylenemethoxy ether group has received considerable attention. $8,9$ and has served as a model for the development of the analogous OCH(Me)OEt moiety, 9 and even the methyleneoxy anion has been shown to be a moderately efficient ortho metalation director.¹⁰ The criteria for a group Z to be effective in ortho metalation are 1) it must be resistant to nucleophilic attack by the RLi reagent, and 2) it must contain at least one heteroatom which can coordinate with the incipient artho metal atom in a 4-, 5- (most favorable) or 6-memhered intermediate. Nucleophilic attack can be minimized by using sterically demanding (e, g, CONF_2) and charge deactivated (e, g, CONR) Z groups.

In Seebach's terminology, ortho metalated species behave as d_5 reagents.¹¹ In synthesis, this translates into the potential for placing carbon substituents, with and vithout functionality for further modification, into positions normally

Directed Metalation Relative Ortho-Directing Priorities

CONR₂ \geq CONHMe, SO₂NMe₂, SO₂NHMe,

 $CH₂$ NMe₂ OCH, OMe > OMe $CH_2CH_2NMe_2$, NMe_2 , CF_3 , F

- **Figure 2**

not accessible by classical aromatic electrophilic substitution chemistry. The synthesis of contiguously tri- and tetra-substituted aromatics, particularly difficult by classical methodology,¹² becomes a feasible, short range project by directed metalation strategy.

The relative directing ability of various groups will depend upon their inductive, resonance, and coordinating properties.⁴ This question has only recently been addressed (Fig. 2).¹³ The qualitative priorities shown, established by competitive inter- or intra-molecular experiments, 13 are highly dependent on the type of experiment as well as on the solvent and temperature used for the generation of the metalated intermediate. Thus, perhaps not surprisingly, disagreement exists on the order of priorities: the tertiary amide appears to bc better or equal to the 2-oxazolino group, the secondary amide is on par with the tertiary amide, and the secondary and tertiary sulfonamides seem to promote ortho metalation as readily as the tertiary amides. The synergistic effect of two groups to promote metalation in between them has not been systematically studied; 14 numerous recent examples attest to the value of this concept for regioselective aromatic substitution. The determination of the survival rate of groups on an aromatic ring which are not involved in the directed metalation reaction also requires comprehensive study. The problem of kinetic vs thermodynamic ortho metalation in competitive situations has also not been carefully addressed.

Our work in directed metalation chemistry was triggered by the observations of Beak and Brown who showed that N,N-diethylhenzamides (but not the corresponding dimethylamides) are smoothly ortho metalated.¹⁵ We shall show that the resulting ortho metalated benzamides are useful synthons for 1) the construction of highly substituted benzene derivatives and from these, phthalides and isochroman-1,3diones; 2) the total synthesis of **phthalideisoquinoline(henzy1isoquinoline** group) and ellipticine alkaloids; 3) the total synthesis of anthraquinone natural and unnatural products and, from the latter, 4) the construction of polycyclic aromatic hydrocarbons **(PAH's).** These synthetic fragments have the common theme of the diethylcarboxamide group as an excellent promotor of artho metalation. Synthesis of Highly Substituted Aromatics

In connection with a problem in alkaloid synthesis, we first investigated the directed metalation of a series of methoxy- and **dimethoxy-N,N-diethylbenzamides** A which are all easily prepared by standard procedures from commercially available

acids (Scheme 1).¹⁶ We found that a variety of contiguously tri- and tetrasubstituted benzenes $2 \cdot \text{can}$ be prepared (Table 1) most of which are accessible only hy multistep sequences if traditional approaches arc used. The general conditions for this reaction as routinely carried out in our laboratories involve the generation of the ortho-lithiated species using sec-BuLi or, less frequently, t-BuLi in THF solution in the presence of TMEDA at -78° C. In reactions with aromatic aldehydes, $Et₂O$ is used to advantage as a solvent. Following the addition of the electrophile,also at -78' C, the mixture is allowed to warm to room tem perature and processed in a standard manner.

Synthesis of Phthalides

Hauser first showed that dimetalated species of secondary benzamides 3a can be condensed with benzophenone and other ketones to give amide alcohols 4 which upon thermolysis readily undergo cyclization to the phthalides 5^2 (Scheme 2). Additional examples of this type of sequence have been provided by Baldwin¹⁷ and by Raphae1.¹⁸ Recently, Fitt and Gschwend invoked the corresponding thioamide $\mathfrak{Z} \mathfrak{b}$ to achieve the same result⁷ and Meyer and Seebach, as part of an in depth study, showed that the successive metalation and carbonation of benzyl alcohols (3c) also serves as a route to phthalides $\frac{10}{10}$ That N, N-diethylbenzamides $\frac{4d}{10}$ are similarly useful in reactions with carbonyl compounds was first demonstrated by Beak and Brown.¹⁵ We found that reaction of $4d$ with benzaldehydes followed by acidcatalyzed cyclization or with DMF followed by \texttt{NabH}_4 reduction and cyclization leads to phthalides $\frac{5}{5}$ in excellent yield.^{16,19} Using the latter sequence, short syntheses of the isomeric phthalides 7 and 9, of demonstrated utility in anthra-
quinone and anthracyclinone synthesis, and phthalide carboxylic acids 11, useful
in alkeloid cunthosis, howe been offected in our laboratory in alkaloid synthesis, have been effected in our laboratory starting with readily quinone and anthracyclinone synthesis, and phthalide carboxylic acids 11, useful
in alkaloid synthesis, have been effected in our laboratory starting with readil
available benzamides 6, & and 10 (Scheme 3).¹⁶ An alternat

HETEROCYCLES, Vol. 14, No. 10, 1980

l,

TABLE 1

÷.

 $Z = a: \text{CONF}, b: \text{CSNHR}, c: \text{CH}_2\text{OH}, d: \text{CONEt}_2$
Scheme 2

metalated m-methoxybenzyl alcohol, first reported by Uemura,²⁰ has been improved by Trost.²¹ However, this route is inefficient for the synthesis of 3-substituted phthalides.^{10,20}

In search of application for the phthalide synthesis, we noted that there exists a small and not rapidly growing number of natural products which exhibit

Scheme 3

highly substituted phthalide structures $\left(\begin{array}{cc} \text{Fig. 3} \end{array}\right)^2$.
metabolites, and the remaining one, shihunine, is an $\frac{3}{2}$, $\frac{22}{1}$ Three of these are fungal metabolites, and the remaining one, shihunine, is an alkaloid which has recently been synthesized from ortho lithiated benzoate generated by a metal-halogen exchange reaction.²³ We achieved a simple synthesis of iso-ochracinic acid, a product of the parasitic fungus Alternaria kikuchiana which is responsible for black spot disease on Japanese pears (Scheme 4). Formylation of the o-anisamide

12 with DMF proceeds quantitatively to give 13 which upon treatment with acetic acid dianion followed by TsOH affords the phthalide 14. Boron tribromide demethylation proceeds in modest yield to give iso-ochracinic acid (15) (40% overall yield). The previous synthesis of this natural product involved a nonregioselective, low yield Wittig reaction on 3-methoxyphthalic anhydride.²⁴ During the preparation of the present paper, Trost reported the synthesis of 15 (44% overall yield) using carbonation of metalated m -methoxybenzyl alcohol as the key step.²¹

Synthesis of Phthalideisoquinoline Alkaloids

Alkyl substituents ortho to certain directed metalation groups $Z(16)$ are activated by resonance to deprotonation by bases weaker than the alkyllithium reagents (e.g., LDA) (Scheme 5). The resulting chelated benzylic anions 17 may be
attacked by electrophiles to give compounds 18 in an overall synthetically useful process of carbon chain extension. Of particular advantage are those artho alkylated derivatives which arise by the directed metalation regimen themselves, e.g., Z=CONHR, 25 CSNHR, 7 2-oxazolino. 26 The finding by Creger that o-toluic acid undergoes this process²⁷ has been especially exploited by F.M. Hauser in the synthesis of napthalenes and napthalene lactone lignan natural products.²⁸

 $Z = co_2H$, co_2R , CONHR, CONR₂, CSNHR, CH₂NMe₂, 2-Oxazolino

In planning a highly convergent assemblage of the phthalideisoquinoline classical synthesis of the penultimate precursor **22,** proceeding in 9 steps and 28% overall yield was carried out in our laboratory with less than overwhelming enthusiasm.~~ Simplification and abbreviation was achieved by the directed metalaenthusiasm.²⁹ Simplification and abbreviation was achieved by the directed metrion strategy starting with the benzamide derivative 19.29 Metalation of 19. followed by treatment with methyl iodide afforded the toluamide 20 in quantitative

 $\ddot{}$

Scheme 6 \sim

yield. Metalation of 20 resulted in the formation of a deep purple solution of the g-toluoyl anion which upon quenching with dry carbon dioxide gave the homophthalic acid amide 21 (71% yield). The conversion of 19 into 21 may be carried out in a one-pot operation with only minor loss in efficacy. Acidic hydrolysis followed by acetic anhydride cyclization gave 22. This abbreviated and simplified synthesis of 22 proceeds in 46% overall yield. Bromination give the unstable bromaanhydride 23 which upon treatment with the phenethylamine 24 produced the promoanhydride $\frac{23}{20}$ which upon treatment with the phenethylamine $\frac{24}{20}$ produced the
rearranged amide phthalide $\frac{25}{25}$. The utility of such compounds for the synthesis of phthalideisoquinoline alkaloids has been known since the time of Perkin and Robinson. Their availability by this and another directed metalation route⁵⁰ assures synthetic accessibility to most members of the phthalideisoquinoline assures synthetic accessibility to most members of the phthalideisoquinoline
alkaloid group. Bischler-Napieralski cyclization of 25 gave the orange-yellow alkaloid group. Bischler-Napieralski cyclization of 25 gave the orange-yellow
dehydrophthalideisoquinoline 26 which upon catalytic hydrogenation provided the
diasteriarenia alkalaide arrhunting L (27) - 1 - 1 - 1 - 1 - 1 dehydrophthalideisoquinoline 26 which upon catalytic hydrogenation provided the diasteriomeric alkaloids cordrastine I (27) and cordrastine II (28) thereby completing the total synthesis.

An alternate highly convergent approach to the phthalideisoquinoline alkaloids (Scheme 7)³¹ was based on the facile condensation of ortho-lithiated benzamides **With aromatic aldehydes¹⁶ and the large rate enhancement in the anchimerically**assisted hydrolysis of ortho methylenehydroxybenzamides to phthalides.³² Thus,

treatment of the readily but not so efficiently available isoquinoline aldehyde treatment of the readily but not so efficiently available isoquinoline aldehyde
29a with the ortho-lithiated species 30a preferably in ether solution gave com-
nound ^{31a}. This compound was found to be extractable from ac treatment of the readily but not so efficiently available isoquinoline aldehyde
29a with the ortho-lithiated species 30a preferably in ether solution gave com-
pound 31a. This compound was found to be extractable from acid interesting property, presumably the result of strong hydroxyl to isoquinoline nitrogen H-bonding, provided the answer to the initial low yields of the reaction which we observed and reaffirmed a basic credo of the synthetic organic chemist: success depends not on how many reactions you run but on how you work them up. When 31a was subjected to cyclization with TsOH, the phthalide 32a was obtained in high yield. Since 32a has been converted into the cordrastines 27 and $28,$ ³³ this completed an alternate formal synthesis of these natural products. Phthalide has been prepared by the analogous sequence $29k + 30k + 31k + 32k$.³⁴

We have recently applied the directed metalation reaction to the polycyclic aromatic amide 33 (Scheme 8)³⁵ in a projected approach to the antitumor quinolizidine alkaloids.³⁶ Successive metalation and condensation with pyridine 2-aldehyde

 $\frac{34}{2}$

 $Scheme 8$

gave the amide alcohol 34 whose properties were annoyingly similar to those of compound 31. Acid-catalyzed cyclization of 34 gave the phthalide 35. Synthesis of Anthraquinone Natural Products

Anthraquinones have a fascinating history as natural products derived from plants³⁷ and insects³⁸ and as important dye substances.³⁹ The recent renaissance in anthraquinane synthesis is due,in large part, to the discovery of clinically useful antitumor activity in the related anthracyclinone derivatives.⁴⁰ The classical approach to anthraquinones involves Friedel-Crafts coupling of phthalic acids 36 or the corresponding anhydrides $\frac{37}{20}$ with phenols or phenol ethers $\frac{38}{20}$ (Scheme 9). For unsymmetrically oxygenated systems, this approach initially provides at least two regioisomeric o-benzoylbenzoic acids 30. Moreover, it carried the potential of a Hayashi rearrangement, $41+42+43$ via equilibrating o-benzoyl benzoyl cation intermediates (41) in the second Friedel-Crafts step $32*40$. Overall, this route is therefore inefficient and potentially ambiguous.⁴¹ starting materials (Scheme 10). Of these, the disconnections and the potential
the orginism rearrangement, $4l^2+2l^2\lambda\lambda$ via equilibrating q -benzoyl benzoyl cation
intermediates (41) in the second Friedel-Crafts step

Retrosynthetic analysis of the anthraquinone nucleus (44) based on the directed metalation tactic allows four modes of dissection based on initial coupling of two oxygenated, appropriately substituted benzamide 45, 47 and benzaldehyde 46, 48

Directed Metalation Route to Anthraquinones

an the correct positioning of the C-4 methoxy substituent in the target anthraquinone and the regiospecific formation of the new C-C bond. This process, once completed, forces the subsequent Friedel-Crafts central ring closure to proceed regiospecifically. The choice between the &and kmodes is dictated by the relative accessibility of the benzamide and benzaldehyde precursors. natural products (Fig. 4).¹⁹ Our general approach is illustrated by the synthesis of erythroglaucin and catenarin (Scheme 11). Lithiation of 3,5-dimethoxybenzamide of erythroglaucin and catenarin (Scheme 11). Lithiation on the correct positioning of the C-4 methoxy substituent in the target anthra-

unione and the regiospecific formation of the new C-C bond. This process, once

completed, forces the subsequent Friedel-Crafts central ring

Based on these considerations, we have synthesized a number of anthraquinone

Naturally Occurring Anthraquinones Synthesized by Directed Metalation Route

 R^1 = Me, R^2 = R^3 = R^4 = H Islandicin $R = R^3 = R^4 = H$, $R^2 = Me$ Digitopurpone R^1 =Me. R^2 = R^3 =H, R^4 =OMe Erythroglaucin R^1 =Me, R^2 = R³ = H, R⁴ = OH Catenarin R^1 =Me, R^2 = R⁴ = H, R³ = OH Cynodontin

Figure 4

 $-1662-$

Scheme 11 nnnnnn

Scheme 11 (concluded)

(49) followed by quenching with 2,5-dimethoxy-p-tolualdehyde (50) gave the alcohol amide 51 which, without isolation, was treated with TsOH to give the phthalide 52. Hydrogenolysis under standard pressure in acetic acid furnished the benzylbenzoic acid *2.* This compound, unlike the corresponding benzoylbenzoic acid which is deactivated to Friedel-Crafts cyclization and, under vigorous conditions, undergoes the Hayashi rearrangement, was smoothly converted by trifluoroacetic anhydride at room temperature into the anthrone 54. Compound exists in concentration dependent
tautomeric equilibrium with the anthracenol 55 as evidenced by NMR studies.
Tunifying a vell known behavior of anthrones, 54 avecousd equipl tautomeric equilibrium with the anthracenol 55 as evidenced by NMR studies.
Typifying a well-known behavior of anthrones, 54 suffered aerial oxidation upon standing to the anthraquinone 56 . This conversion was more reproducibly carried out by chromium trioxide oxidation. To complete the syntheses, selective demethylation of 56 with BBr₃ yielded erythroglaucin (57) while vigorous treatment with m and the set of the successful syntheses of pyridium hydrochloride delivered catenarin (58). The successful syntheses of naturally occurring anthraquinones (Fig. 4) by short (6 steps) and efficient (20-30% overall yield) routes from readily available amide and benzaldehyde derivativej demonstrates the generality and versatility of the dirccted metalation approach. Worthy of note is the toleration of methyl groups both in the benzamide (e.g. for soranjidiol) and benzaldehyde (50 used in 5 of the 6 syntheses) components in the metalation and condensation reactions respectively. Notice is the generality and versatility of the directed metalation approach.
Northy of note is the toleration of methyl groups both in the benzamide (e.g. for
Soranjidiol) and benzaldehyde (50 used in 5 of the 6 syntheses

Synthesis of Polycyclic Anthraquinones and Polycyclic Aromatic Hydrocarbons (PAH's) While engaged in the synthesis of anthraquinone natural products, we treated

$$
\begin{bmatrix}\n\mathbf{E}_{t,N} \\
\mathbf{I} \\
\mathbf{I} \\
\mathbf{EDA} \\
\mathbf{I} \\
\mathbf{I} \\
\mathbf{I} \\
\mathbf{EDA} \\
\mathbf{I} \\
$$

Scheme 12

of anthraquinone in 44% yield. 42 Pasteur's dictum (chance favors the prepared mind)⁴³ is personally validated in that I had executed, as an undergraduate student, Fieser's preparation of anthraquinone and therefore I was in the position to quickly recognize the insoluble and rock-stable nature of the compound which precipitated from the reaction outlined in Scheme 12. The requirement for benzaldehyde in this reaction could be demonstrated; when it was excluded, only trace amounts of anthraquinone were obtained (most likely by dimerization of ortholithiated benzamide). Furthermore, substituted benzaldehydes were shown to participate in the reaction (Fig. 5). Alkyl and alkoxy benzaldehydes undergo the reaction in poorer yields presumably owing to competitive proton exchange and directed metalation respectively. 1-Naphthaldehyde, 2-napthaldehyde, and 9phenanthraldehyde afforded polycyclic anthraquinones in yields comparable to the parent reaction.

 $(39%)$

 $(41%)$

Fiaure 5

A reasonable mechanism for the reaction outlined in Scheme 12 is depicted in Scheme 13. The expected initial condensation product $\frac{59}{20}$, suffers metalation \sim directed by the alkoxy group to give the dianion 60. Such ortho-metalated species of benzyl alcohol have received credence from the recent work of species of benzyl alcohol have received credence from the recent work of
Seebach.¹⁰ Intermediate 60 in turn undergoes cyclization to the tetrahedral Seebach.¹⁰ Intermediate $\frac{60}{2}$ in turn undergoes cyclization to the tetrahedral
intermediate 61 which, by expulsion of LiEt₂, gives the hydroxyanthrone 62. The
latter undergoes aerial oxidation, also a well preced latter undergoes aerial oxidation, also a well precedented step, to give anthraintermediate 61 which, by expulsion of LiEt₂, gives the hydroxyanthrone 62. The
latter undergoes aerial oxidation, also a well precedented step, to give anthra-
quinone 63. Support for the key step 59+60 of this tandem

 $\frac{59}{100}$

 60

Scheme 13

was derived from the reaction of N,N-diethylbenzamide with acetophenone and HEFROCYCIES, Vol. 14, No. 14
was derived from the reaction of N,N-diethylbenzamide with acetophenone and
benzophenone under identical reaction conditions (Scheme 14).⁴² The formation of
the hydroxyanthrones 64a and 64b i benzophenone under identical reaction conditions (Scheme 14).⁴² The formation of
the hydroxyanthrones $\frac{64a}{44}$ and $\frac{64b}{44}$ is consistent with the involvement of the proposed intermediate $\frac{65}{100}$ in the overall reaction.

Anthraquinones are easily converted into the corresponding aromatic hydrocarbons. For example, the synthesis of benz[a]anthracene (66) and 7,12dimethylbenz[a]anthracene (67) may be achieved in high yield from the anthraquinone 68, (Scheme 15) thus providing a short route to these carcinogenic **PAH'5.** 4 2 rom the reaction of N
under identical react
thrones 64a and 64b is
diate 65 in the overa
ones are easily conve
example, the synthes
alanthracene (67) may
Scheme 15) thus provie
example, the synthes

The one-step synthesis of anthraquinones is less successful for a- and **B**naphthamides and more highly condensed aromatic amides (Fig. $\begin{bmatrix} \text{ess} \text{full} & \text{f} \\ \text{Fig. 6} \\ \text{www.} \end{bmatrix}$ 6). Since the major products are phthalides with only minor amounts of anthraquinones being obtained, it appears that the reaction fails at the second metalation step (59+60, Scheme 13). Nevertheless, the phthalides are also useful precursors to PAH derivatives. Our knowledge concerning the directed metalation capabilities of the CONEt₂ group in more highly condensed aromatic is currently limited. More successful is the combination of certain heterocyclic aldehydes with bencamides and naphthamides, a reaction which allows access in one step to some bizarre major products are phthalides with only minor amounts of anthraquinones being
obtained, it appears that the reaction fails at the second metalation step
(59+60, Scheme 13). Nevertheless, the phthalides are also useful prec tion of heterocyclic aldehydes with heterocyclic amides provides some new and unusual diheterocyclic benzoquinones (Fig. 8). In the formation of thiophens and more integrals and more integrals and more integrals with only minor amounts of anthraquinones being
the formation of the pearst that the reaction fails at the second metalation step
 $\frac{6$

vantage is gained from the inherently high kinetic acidity of the 2-hydrogen in the thiophen and furan rings. On the other hand, the formation of the thiophena-In the formation of thiopheno- and furano-benzoquinones (Figs. 7 and 8), advantage is gained from the inherently high kinetic acidity of the 2-hydrogen in the thiophen and furan rings. On the other hand, the formation of To test the efficacy of this step, we metalated three isonicotinamides under the tion of heterocyclic aldehydes with heterocyclic amides provides some new and
unusual diheterocyclic benzoquinones (Fig. 8).
In the formation of thiopheno- and furano-benzoquinones (Figs. 7 and 8), ad-
vantage is gained f The results show that the N,N-diisopropyl and N-phenyl isonicotinamides incorporate deuterium almost quantitatively whereas the diethyl analogue is less efficient. Unfortunately, tandem metalation reaction with benzaldehyde on the former two compounds did not produce benzoquinonc products. These results, presumably due to hindrance (N,N-diisopropyl) and deactivation (N-phenyl) in the formation of the tetrahedral intermediate $(61, 5$ cheme 13) forced us to use the N.N-diethyl isonicotinamide for these reactions.

 $(81%)$

О

 $(2%)$

HetAr

 $(35%)$

Ar

 Et_2N

 $(37%)$

 $(44%)$

 $\hat{\boldsymbol{\beta}}$

 $\sim 10^{-1}$

 $\sim 10^7$

Synthesis of Ellipticine Alkaloids

The efficient incorporation of an indole 3-aldehyde into the indolo thiopheno **compared (Fig. 3)** and the isonicotinamide deuteration experiments (Synthesis of Ellipticine Alkaloids

The efficient incorporation of an indole 3-aldehyde into the indolo thiopheno

benzoquinone (Fig. 3) and the isonicot suggested that the linear benzoquinones 69 (Scheme 17) representing the ellipticine alkaloid skeleton could be prepared by the tandem directed metalation route. In the event, ⁴² the reaction proceeded to give the desired derivatives $\overset{69}{\sim}$ in yields which are inversely proportional to the utility of the R protective group. yields which are inversely proportional to the utility of the R protective group
The indolopyridobenzoquinones 69 were then transformed in a three-step sequence
without numification of intermaliates into the numidescrhamel The indolopyridobenzoquinones 69 were then transformed in a three-step sequence
without purification of intermediates into the pyridocarbazoles 70 (Scheme 18).
As expected from the potent reductive exidie conditions use As expected from the potent reductive-acidic conditions used, the methylenemethoxy without purification of intermediates into the pyridocarbazoles $\overline{70}$ (Scheme 18).
As expected from the potent reductive-acidic conditions used, the methylenemethoxy
derivative 69, R=CH₂OMe gave ellipticine (70, R=H of the antitumor ellipticine alkaloids furnished the compounds to neutralize the effects of the PAH's obtained earlier (Scheme 15) so that work in our laboratory could continue!

Conclusion

Directed metalation of aromatic substrates is an important strategy for the regiospecific synthesis of highly substituted and condensed aromatics, heterocycles, and several classes of natural products. We have shown that ortho-metalated benz**midesare** useful synthons for the preparation of contiguously tri- and tetrasubstituted benzene derivatives (Table 1), phthalides (Schemes 2 and 4), isochroman-1,3-diones (Scheme 6), phthalideisoquinoline alkaloids (Schemes 6 and 7), anthraquinone natural products (Scheme e preparation
(Table 1), pht)
hthalideisoquin
Scheme 11 and P
mzoquinones (Fi We have shown that ortho-metalated benz

ration of contiguously tri- and tetra-

1), phthalides (Schemes 2 and 4), iso-

eisoquinoline alkaloids (Schemes 6 and 7),

11 and Fig. 4), polycyclic anthraquinones

ones (Figs. 7 (Figs. 5 and 6), heterocyclic benzoquinones (Figs. 7 and 8, Scheme 17), PAH's (Scheme 15), and ellipticine alkaloids (Scheme 18).

The chemical genealogies of the diverse ortho metalated aromatic species (Scheme 15), and ellipticine al

The chemical genealogies of

developed recently (Fig. 1) may

Gilman, Wittig and, of course, 1) may be traced to the original contributions of Hauser, Gilman, Wittig and, of course, Grignard among many others. Wittig discovered phenyllithium and, in recent years, called it his Wünschelrute (divining rod). This quality appears to have been transcribed into many of the ortho metalated aromatics including, on the basis of our experience, the N,N-diethylbenzamides.

Future efforts in the area of directed metalatian chemistry will provide new groups which promote ortho metalation, establish priorities among them, determine kinetic and thermodynamic control parameters in their generation and, undoubtedly, find new applications in organic synthesis.

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1. H. Gilman and J.W. Morton, Jr., <u>Organic Reactions</u>, 8, 258 (1954).
2. W.H. Puterbaugh and C.R. Hauser, J. Org. Chem., 29, 853 (1964).

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- 2. W.H. Gilman and J.W. Morton, Jr., <u>Organic Reactions</u>, 8, 258 (1954).
2. W.H. Puterbaugh and C.R. Hauser, <u>J. Org. Chem., 29, 853 (1964).</u>
- 3. A.W. Langer, ed., Palyamine-Chelated Alkali Metal Compounds, Adv. Chem. Ser., 130, American Chemical Society, Washington, D.C. (1974).
- 4. For a comprehensive review, see H.W. Gschwend and H.R. Rodriguez, Organic Reactions, 26, 1 (1979).
- 5. See, inter alia, A. Marxer, H.R. Rodriguez, J.M. McKenna, and H.M. Tsai, *J.* Org. Chem., 40, 1427 (1975); R.M. Sandifer, C.F. Beam, M. Perkins, and C.R. Hauser, Chem. Ind. (London), 231 (1977); H. Watanabe, C.L. Mao, I.T. Barnish, and C.R. Hauser, $J.$ Org. Chem., 34 , 919 (1969).
- 6. N.S. Narasi and R.S. Mali, Chem. Ind. (London), 519 (1975); B.H. Bhide and $V.P.$ Gupta, Indian J. Chem., Sect. B, 15, 512 (1977).
- 7. J.J. Fitt and H.W. Gschwend, J. Org. Chem., 41, 4029 (1976).
- 8. H. Christensen, Synth. Commun., 5, 65 (1975); R.C. Roland, Tetrahedron Lett., 3973 (1975).
- 9. G.A. Kraus and J.O. Pezzanite, $J.$ Org. Chem., 44, 2480 (1979).
- 10. N. Meyer and D. Seebach, Chem. Ber., 113, 1304 (1980).
- 11. D. Seebach, Angew. Chem. Int. Ed. Engl., 18, 239 (1979).
- 12. For alternate solutions involving aromatic C-H protection, see M. Tashiko, Synthesis, 921 (1979).
- 13. P. Beak and R.A. Brown, J. Org. Chem., 44, 4463 (1979). A.I. Meyers and K. Lutomski, <u>J. Org. Chem</u>., 44, 4464 (1979).
14. D.W. Slocum and C.A. Jennings, <u>J. Org. Chem., 41</u>, 3653 (1976).
- 14. D.W. Slocum and C.A. Jennings, <u>J. Org. Chem</u>., 41, 3653 (1976).
15. P. Beak and R.A. Brown, <u>J. Org. Chem</u>., 42, 1823 (1977).
-
- 16. S.O. de Silva, J.N. Reed,and V. Snieckus, Tetrahedron Lett., 5099 (1978).
- 17. J.E. Baldwin and K.W. Bair, Tetrahedron Lett., 2559 (1978).
- 18. I. Forbes, R.A. Pratt,and R.A. Raphael, Tetrahedron Lett., 3965 (1978).
- 19. S.O. de Silva, M. Watanabe, and V. Snieckus, J. Org. Chem., 44, 4802 (1979).
- 20. M. Uemura, S. Tokuyama, and T. Sakan, Chem. Lett., 1195 (1975).
- 21. B.M. Trost, G.T. Rivers, and J.M. Gold, J. Org. Chem., 45, 1835 (1980).
- 22. T.K. Devon and A.I. Scott, Handbook of Naturally Occurring Compounds. Vol. 1, Acetogenins, Shikimates and Carbohydrates, Academic Press, New York, 1975,p 249 Acetogenins, Shikimates and Carbohydrates, Academic Press
23. G.B. Bodem and E. Leete, <u>J. Org. Chem., 44</u>, 4696 (1979).
24. D.W. Knight and C.D. Portas, Tetrahedron Lett., 4543 (1974).
-
- 24. D.W. Knight and C.D. Portas, Tetrahedron Lett., 4543 (1977).
- 23. G.B. Bodem and E. Leete, <u>J. Org. Chem</u>., 44, 4696 (1979).
24. D.W. Knight and C.D. Portas, <u>Tetrahedron Lett</u>., 4543 (1977).
25. R.E. Ludt, J.S. Griffiths, K.N. McGrath,and C.R. Hauser, <u>J. Org. Chem</u>., 38, 1668 (1973) and references therein.
- 26. H.W. Gschwend and A. Hamdan, J. Org. Chem., 40, 2008 (1975).
- 27. P.L. Creger, J. Am. Chem. Soc., 92, 1396 (1970).
- 28. F.M. Hauser and R.P. Rhee, J. Am. Chem. Soc., 99, 4533 (1977); F.M. Hauser **o,..** and R.P. Rhee, $J.$ Org. Chem., 42, 4155 (1977).
- 29. S.O. de Silva, I. Ahmad, and V. Snieckus, Can. J. Chem., 57, 1598 (1979).
- 30. B.C. Nalliah, D.B. MacLean, R.G.A. Rodrigo, and R.H.F. Manske, Can. J. Chem., $\frac{55}{100}$, 922 (1977).
 $\frac{55}{1000}$, do Silve, I. Abrel, and M. Spieglys, Tatrabadren Lat. 5107 (1978).
- 31. S.O. de Silva, I. Ahmad, and V. Snieckus, Tetrahedron Lett., 5107 (1978).
- 32. K.N.G. Chiong, S.D. Lewis, and J.A. Shafer, J. Am. Chem. Soc., 97, 418 (1975).
- 33. V. Smula, N.E. Cundasawmy, H.L. Holland, and D.B. MacLean, Can. 3. Chem., 51, **A,..** 3293 (1973).
- 34. S.O. de Silva, D.A. Kuntz, and V. Snieckus, unpublished results, 1979.
- 35. S.O. de Silva, M. Watanabe, and V. Snieckus, unpublished results, 1979.
- 36. For recent synthetic work in this area, see G.G. Trigo, E. Galvez, and M.M. Sollhuber, J. Heterocyclic Chem., 17, 69 (1980).
- 37. R.H. Thomson, Naturally Occurring Anthraquinones, 2nd ed., Academic Press, New York, 1971.
- 38. K.S. Brown, Jr., Chem. Soc. Rev., 4, 263 (1975); F.L.C. Baranyovits, $Endeavor, 2, 85 (1978).$
- 39. K.H. Schunderhutte, Chem. Synth. Dyes, 6, 211 (1972).
- 40. Reviews: F. Arcamone, Topics in Antibiotic Chemistry, Vol. 2, P.G. Sammes, ed., Ellis Horwood Ltd., Sussex, England, 1978, p. 89; T.R. Kelly, <u>Annu. Rep</u>.
Med. Chem., 14, 288 (1979).
B.H. Therson, The Chemistry of the Quineneid Corneurds, Bent I. S. Betai. ed.
- 41. R.H. Thomson, The Chemistry of the Quinonoid Compounds, Part I, S. Patai, ed., Wiley, New York, 1974, p. 136.
- 42. M. Watanabe and V. Snieckus, J. Am. Chem. Soc., 102, 1457 (1980).
- 43. Dans le champs de l'observation, I'hasard ne favarise que les esprits pr6 parés. Quoted in A.L. Mackay, The Harvest of a Quiet Eye, The Institute of Physics, Bristol, England, 1977, p. 116.