

A Comparative Study of Thermal- and Radiation-Induced Single Electron Transfer in Reactions of 1,4-Diaza-1,3-butadienes with Dialkylzinc Compounds

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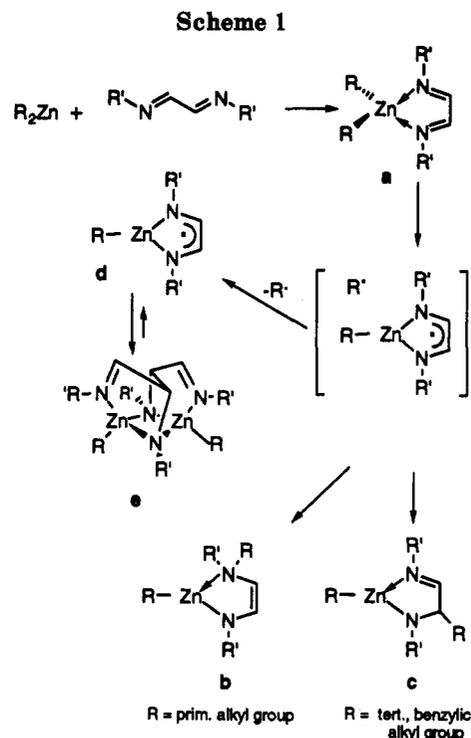
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The reaction of R_2Zn with *t*-BuDAB ($t\text{-BuN}=\text{CH}-\text{CH}=\text{N}-t\text{-Bu}$) results in the quantitative formation of the red 1:1 coordination charge transfer complexes $R_2Zn-t\text{-BuDAB}$ ($R = \text{Me}$ (1a), CH_2SiMe_3 (5a)). Upon heating ($T > 35^\circ\text{C}$), both complexes undergo an inner-sphere single electron transfer (SET) leading to the radical pair $[(RZn-t\text{-BuDAB})^\bullet(R^\bullet)]$, that collapses, depending on R^\bullet , to several alkylation products, *i.e.* the *N*-alkylated product $\text{MeZn}(t\text{-BuN}-\text{CH}=\text{CH}-\text{N}(\text{Me})-t\text{-Bu})$ (1b) and the exocyclic enamines $RZn(t\text{-BuN}=\text{CH}-\text{C}(=\text{CHR}')\text{N}-t\text{-Bu})$ ($R = \text{Me}$ (1f), CH_2SiMe_3 (5f/5g); $R' = \text{H}, \text{SiMe}_3$). Alternatively, the alkyl radical R^\bullet escapes from the organozinc radical $(RZn-t\text{-BuDAB})^\bullet$, that subsequently dimerizes to the dinuclear species $(RZn-t\text{-BuDAB})_2$ ($R = \text{Me}$ (1e), CH_2SiMe_3 (5e)). The charge transfer complex $\text{Me}_2\text{Zn}-t\text{-BuDAB}$ was irradiated in order to induce a SET reaction within the complex. The irradiation reaction resulted in the formation of the symmetric dinuclear species 1e (30%) and the unsymmetric dinuclear species $[\text{MeZn}\{t\text{-Bu}(\text{H})\text{NCH}(\text{Me})\text{CH}(\text{N}-t\text{-Bu})-\}\{-\text{CH}(\text{N}-t\text{-Bu})\text{CMe}=\text{N}-t\text{-Bu}\}\text{ZnMe}]$ (1h) (70%). Product 1h was independently prepared from $\text{MeZn}(t\text{-BuN}-\text{CH}(\text{Me})-\text{CH}=\text{N}-t\text{-Bu})$ (1c). The latter complex is in a tautomeric equilibrium with its enamine $\text{MeZn}(t\text{-BuN}-\text{CH}(\text{Me})=\text{CH}-\text{N}(\text{H})-t\text{-Bu})$, that reacts *via* a condensation reaction with a parent molecule 1c to give 1h.

Introduction

Recently, we reported on the regioselective *N* or *C* alkylation by dialkylzinc compounds (R_2Zn) of 1,4-disubstituted-1,4-diaza-1,3-butadienes ($R'\text{N}=\text{CH}-\text{CH}=\text{NR}'$ (*R'DAB*)).¹ These reactions are the result of an intramolecular electron shift from the R_2Zn moiety to the *R'DAB* ligand in an initially formed 1:1 coordination complex $R_2ZnR'DAB$. The so formed radical pair may collapse within the solvent cage to *N*- (b, cf. Scheme 1) (for primary alkyl groups) or *C*-alkylated (c) (for tertiary alkyl and benzylic groups) organozinc complexes, or the alkyl radical may escape from the organozinc radical. In that case, the remaining organozinc radical (d) is persistent and is in equilibrium with its *C-C* coupled dimer e (see Scheme 1).^{2a,3}

The initially formed coordination complexes in the alkylation reaction are, like $R_2Zn\text{Bpy}$ complexes,^{4,5} strongly colored with widely varying absorption maxima, that



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depend on the nature of R ; see Table 1. It has been suggested that the intense color of the complexes 1a-4a is caused by a $S_0 \rightarrow S_1$ LLCT transition $\sigma(\text{M}-\text{C}) \rightarrow \pi^*\text{DAB}$.²

Theoretical calculations on $R_2Zn-t\text{-BuDAB}$ complexes, $R = t\text{-Bu}, i\text{-Pr}, \text{Et}$, and Me , show that the LUMO's of

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Table 1. Characteristic Features of R_2Zn-t -BuDAB Complexes

R_2Zn-t -BuDAB R group	color ^a	λ_{max} (nm) ^b	conversion temp (°C) ^c
Me (1a)	orange	390	room temp
Et (2a)	red	490	-50
<i>i</i> -Pr (3a)	blue	590	-70
<i>t</i> -Bu (4a)	purple	650	-90

^a In diethyl ether. ^b Extinction coefficient ϵ was determined for Me_2Zn-t -BuDAB and was found to be $1200 M^{-1} cm^{-1}$. ^c Conversion was monitored by ¹H NMR spectroscopy and starts above the temperature given.

these complexes, which are mostly localized within the chelated DAB ligand, have a comparable energy level, whereas the HOMO energies increase in the series $R = Me < Et < i$ -Pr $< t$ -Bu.^{2a,6} This trend is also reflected by the decreasing stability of the coordination complexes, of which t -Bu₂Zn- t -BuDAB (4a) is only stable to -90 °C, whereas the Me_2Zn complex is stable at room temperature; see Table 1. When the temperature is raised above a critical temperature, an intramolecular single electron transfer (SET) from the R_2Zn to the DAB occurs, leading to the radical pair a' , that finally collapses to the alkylated products (see Scheme 1).

ESR experiments have shown that by irradiation in the charge transfer band of the coordination complexes, organozinc-DAB radicals are formed, indicating that the SET within the R_2Zn-t -BuDAB complexes can also be light-induced.^{2a}

Recently, we have found that the bis(trimethylsilyl)methylzinc compound, $(Me_3SiCH_2)_2Zn$, like Me_2Zn forms a stable 1:1 coordination complex at room temperature.⁷ In this paper we describe a study of the product formed by photochemical initiation of Me_2Zn-t -BuDAB and compare the results with those obtained in the thermally induced SET reaction of Me_2Zn-t -BuDAB and $(Me_3SiCH_2)_2Zn-t$ -BuDAB.

Results

Both Me_2Zn and $(Me_3SiCH_2)_2Zn$ react with t -BuDAB in diethyl ether at room temperature to the corresponding 1:1 coordination complexes R_2Zn-t -BuDAB ($R = Me$ (1a), $R = CH_2SiMe_3$ (5a)). Compounds 1a and 5a were crystallized from hexane to afford orange crystals. The UV-vis absorption spectrum of 5a is comparable to that of 1a (see Table 1) with λ_{max} at 390 nm.

Thermal Reaction. Solutions of 1a and 5a in benzene were heated for 6 h at 70 °C. Product analysis revealed that quantitative conversion into several products had occurred. Complex 1a was converted into three products: the C-C coupled dinuclear species $(MeZn-t$ -BuDAB)₂ (1e) (50%), the N-alkylated product $[MeZn(t$ -BuN-CH=CH-N(Me)- t -Bu)] (1b) (25%), and the zinc-enamine $[MeZn(t$ -BuN=CH-C(CH₂)-N- t -Bu)] (1f) (25%) (see Scheme 2, eq 1). Compound 1b is similar to the products which are exclusively formed in the reaction of t -BuDAB with primary dialkylzinc compounds (see compound b in Scheme 1). The ¹H NMR spectrum of 1b shows the same characteristic AB pattern in the olefin region (³ $J_{HH} = 4.4$ Hz) as was observed for comparable N-alkylated prod-

ucts,^{1,8} which is consistent with the presence of *cis* olefin H atoms. The ¹H NMR spectrum of the dimeric compound 1e shows an AX pattern for the N=CH-CH-N moiety with a coupling constant ³ J of 2.7 Hz, characteristic for such dimers.⁷ The ¹H NMR spectrum of the exocyclic zinc-enamine 1f shows three singlet resonances for the N=CH-C=CH₂ moiety at 7.42, 4.59, and 4.18 ppm for the N=CH and the anisotropic C=CH₂ hydrogens, respectively. No geminal coupling was observed for the CH₂ hydrogens.

Thermal conversion of 5a gave the dimeric species $(Me_3SiCH_2Zn-t$ -BuDAB)₂ (5e) (65%) and the exocyclic enamines $Me_3SiCH_2Zn(t$ -BuN=CH-C(CH₂)-N- t -Bu) (5f) (7%) and $Me_3SiCH_2Zn(t$ -BuN=C(CH₂SiMe₃)-N- t -Bu) (5g) (28%) (see Scheme 2, eq 2). The ¹H NMR spectrum of 5e and 5f shows the same characteristic signals as were found in 1e and 1f, respectively, whereas 5g only contains two singlets at 8.11 and 5.04 ppm. The Si-C coupling constant in 5g, ¹ $J_{Si-C} = 24.7$ Hz, is significantly smaller than that of normal Si-C=C bonds in, e.g., trimethylvinylsilane, ¹ $J_{Si-C} = 32.3$ Hz. Also the ¹³C chemical shift of the Si-C=C carbon in 5g, 105.4 ppm, is much lower than that of the corresponding carbon in trimethylvinylsilane, 139.9 ppm, implying that this olefin carbon is strongly nucleophilic.⁹

The thermal decomposition of 5a was followed by ¹H NMR in benzene-*d*₆ which showed the simultaneous formation of the products 5e, 5f, and 5g, as well as a substantial amount of Si(CH₃)₄ and resonances that were ascribed to $Me_3SiCH_2SiMe_3$.

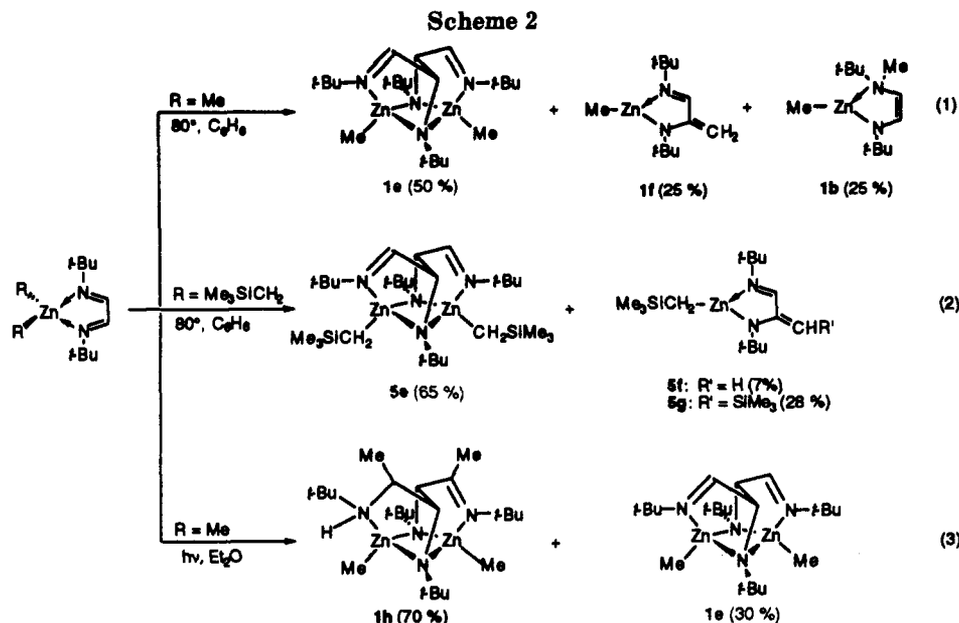
Irradiation Experiment. A solution of 1a in diethyl ether or benzene was irradiated at approximately 20 °C with a high-pressure mercury lamp until no more starting material was detected by ¹H NMR spectroscopy. It appeared that the symmetric dinuclear species $(MeZn-t$ -BuDAB)₂ (1e) (30%) and a new, unsymmetric dinuclear species $[MeZn\{t$ -Bu(H)NCH(Me)CH(N- t -Bu)-\}-CH(N- t -Bu)CMe=N- t -Bu}ZnMe] (1h) (70%) had formed (see eq 3). ¹H NMR measurements during the irradiation experiment showed that the C-alkylated product $[MeZn(t$ -BuN-C(H)Me-CH=N- t -Bu)] (1c) and the dimer 1e are formed first and 1h is formed shortly thereafter. The relative amount of 1h increases during the reaction, whereas the amount of 1e and 1c decreases and the latter eventually disappears.

Characterization and Independent Synthesis of 1h. The dinuclear structure of 1h consists of two differently alkylated DAB skeletons, which are coupled *via* a newly formed C-C bond. The ¹H NMR spectrum shows four t -Bu- and two Me-Zn signals. Furthermore two CH₃ resonances were observed, *viz.* a singlet (1.75 ppm) and a doublet (0.82 ppm), which were assigned to a N=CMe and a N-C(H)Me entity, respectively. A characteristic ABX₃ pattern at 2.87 ppm was observed for the t -Bu(H)-N-C(Me)H-CH-N- t -Bu hydrogen. The N-H resonance of this moiety was not observed in the ¹H NMR spectrum. However, simulation of the ABX₃ pattern (see Figure 1) enabled the determination of ³ J_{NH-CH} , which was found to be 12.0 Hz. Finally, doublet (2.14 ppm) and singlet (3.18 ppm) resonances of the hydrogens bonded to the bridging carbons were found. No coupling between these two anisotropic hydrogens was observed.

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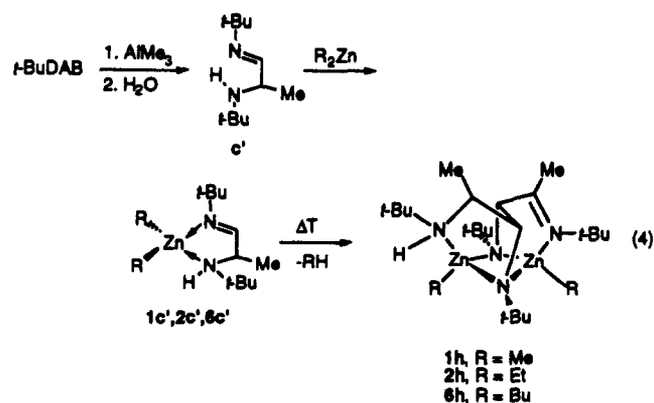
(9) The exocyclic zinc-enamines 1f, 5f, and 5g are reactive toward electrophilic reagents.



Severe disorder made it impossible to refine the X-ray structure of **1h**, and no complete structure can therefore be presented. However a preliminary view of the structure shows that the two alkylated NCCN skeletons are coupled *via* a C-C bond similar to that found in the symmetric dinuclear structure of $(Me_3SiCH_2Zn-t-BuDAB)_2$ (**5e**).⁷ In **5e** the dihedral angle between the hydrogens bonded to the bridging carbons is close to 90° , which would explain the lack of coupling between the two corresponding anisotropic hydrogens bonded to the bridging carbon atoms in **1h**.

We have been unable to separate the two dinuclear products, **1e** and **1h**, but they have each been prepared by independent syntheses. The dinuclear species **1e** was obtained by crystallization from the product mixture of the thermal reaction of $Me_2Zn-t-BuDAB$ (*vide supra*), whereas **1h** was obtained from the reaction of Me_2Zn with $(t-BuN=CH-CH(Me)-N(H)-t-Bu)$ (**c'**).

The independent synthesis of **1h** was based upon the observation that in the irradiation reaction this dinuclear species was formed *via* the C-alkylated product **1c**. The necessary starting material $t-BuN=CH-CH(Me)-N(H)-t-Bu$ (**c'**) was obtained by hydrolysis of $Me_2Al(t-BuN=CH-CH(Me)-N-t-Bu)$, which is formed in the alkylation reaction of Me_3Al with *t*-BuDAB (see eq 4).¹⁰



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The reaction of **c'** with Me_2Zn first gives the 1:1 coordination complex $Me_2Zn(t-BuNH-CH(Me)-CH=N-t-Bu)$ (**1c'**),¹¹ which upon raising the temperature to $50^\circ C$ affords the dinuclear compound **1h** quantitatively. The intermediate $MeZn(t-BuN-CH(Me)-CH=N-t-Bu)$ (**1c**) was never observed. The protonolysis of the R-Zn bond in the initial step of the reaction sequence (see Discussion), producing RH, was established with 1H NMR.

This synthesis of **1h** is also applicable to other dialkylzinc compounds, and the $Et-Zn$ (**2h**) and $Bu-Zn$ (**6h**) analogs of **1h** were obtained from the reaction of **c'** with Et_2Zn and Bu_2Zn , respectively. The $Bu-Zn$ derivative **6h** was prepared to increase the solubility of the product and to enable a molecular weight determination in solution (cryoscopy in benzene) in order to confirm its dinuclear nature (Calcd: 612. Found: 568).

Discussion

When it is accepted that the LUMO energy level of $(Me_3SiCH_2)_2Zn-t-BuDAB$ (**5a**) is at the same level as was calculated for R_2ZnDAB ($R = Me, Et, i-Pr, t-Bu$), then its λ_{max} of 390 nm indicates that its HOMO energy level must be comparable with that of $Me_2Zn-t-BuDAB$ (**1a**). This is in accord with the recent observation that $((Me_3Si)_2CH)_2ZnBpy$ ¹⁴ has about the same λ_{max} as Me_2ZnBpy . Apparently, substitution of one or two of the hydrogens for $SiMe_3$ in Me_2Zn seems to have little or no influence on the LLCT charge transfer band and, accordingly, the stability of **5a** should be similar to that of **1a**. Indeed, upon heating both complexes decompose at the same critical temperature (about $35^\circ C$) to three products, *i.e.*

(11) Dialkylzinc compounds are known to react with secondary amines to form coordination complexes which, however, have only limited stability. Upon heating ($T > 30^\circ C$) these complexes decompose to give zinc amides.¹² The alkyl groups in alkylzinc halides ($RZnX$), however, are much more stable to acidic hydrogens.¹³

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(13) Knoess, H. P.; Furlong, M. T.; Rozema, M. J.; Knochel, P. *J. Org. Chem.* 1991, 56, 5974.

(14) Westerhausen, M.; Rademacher, B.; Schwarz, W. *J. Organomet. Chem.* 1992, 427, 275.

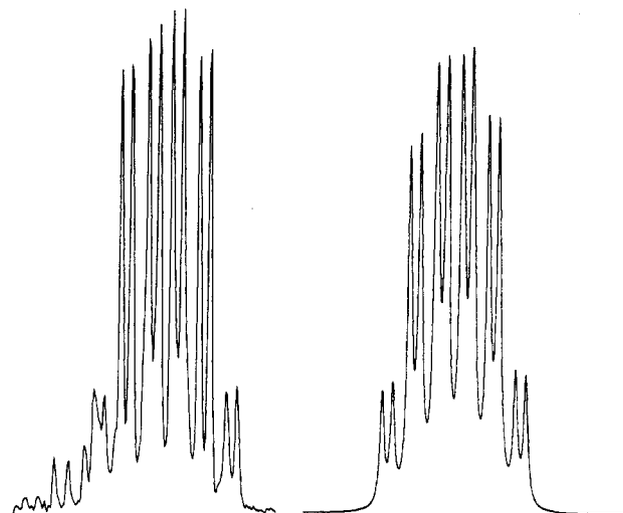
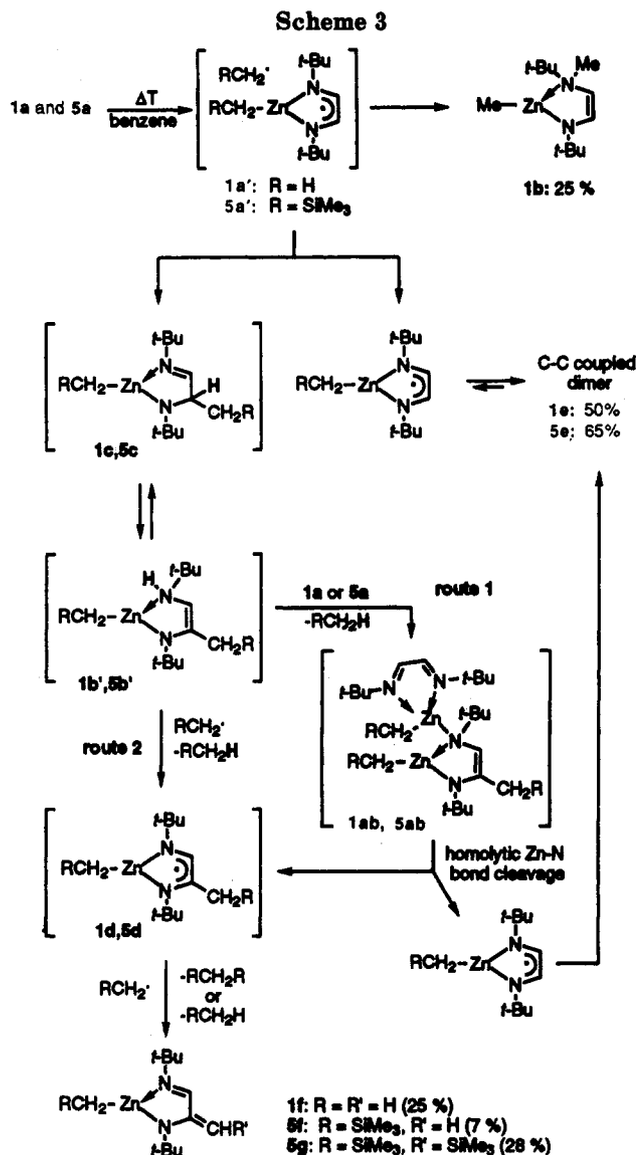


Figure 1. Observed (left) and simulated (right) 300-MHz ^1H NMR resonance for the ABX_3 pattern of the $t\text{-Bu(H)N-C(Me)H-CH-N-t-Bu}$ hydrogen in **1h**.

1b, **1e**, and **1f** for **1a** and **5e**, **5f**, and **5g** for **5a**. Although $(\text{Me}_3\text{SiCH}_2)_2\text{Zn}$ is a primary dialkylzinc compound no N-alkylated product similar to **1b** was found. This was also observed in the reaction of $(\text{Me}_3\text{CCH}_2)_2\text{Zn}$ with $t\text{-BuDAB}$, that resulted in C alkylation only.⁸

The mechanism we propose for the formation of the products **1b**, **1e**, **1f**, **5e**, **5f**, and **5g** is outlined in Scheme 3. Upon heating, the coordination complexes **1a** and **5a** undergo a homolytical cleavage of one of the alkyl-zinc bonds, to give the radical pairs **1a'** and **5a'**, respectively. From this intermediate state the alkyl radicals can add to the NCCN skeleton of the organozinc radical, affording either the N-alkylated compound **1b** or the C-alkylated compounds **1c** and **5c**. Alternatively, the alkyl radicals can escape from the organozinc radical, leaving organozinc radicals that dimerize to **1e** and **5e**. The N-alkylated compound **1b** is the expected product in the reaction of **1a**, whereas the C-alkylated compounds **1c** and **5c** are not observed. However, we believe that **1f**, **5f**, and **5g** are formed from these intermediates. Earlier, we found that C-alkylated compounds like **1c** and **5c** are in equilibrium with their enamine tautomer $\text{RZn}(t\text{-BuN-CR=CH-N(H)-}t\text{-Bu})$ (**1b'** and **5b'**) and that the position of the equilibrium depends on the nature of the group bonded to the C atom of the NCCN skeleton.⁸ When the shifted hydrogen is eliminated from the enamine tautomer, the organozinc radical species **1d** and **5d** are formed. This elimination process may occur in two ways. Firstly, when the tautomerization is faster than the initial C or N alkylation, a carbon-zinc bond in **1a** and **5a** may be hydrolyzed by the amine hydrogen of the enamine to form a N-Zn coupled dinuclear species (**1ab**, **5ab**) (route 1 in Scheme 3). Subsequent homolytic cleavage of the newly formed N-Zn bond leads to the radical species $(\text{RCH}_2\text{-Zn-}t\text{-BuDAB})^\bullet$ and **1d** (**5d**), of which the former dimerizes to **1e** (**5e**). When the rate of tautomerization is slower than the escape of an alkyl radical from the organozinc radical, then the shifted hydrogen in **1b'** (**5b'**) may be eliminated in a free alkyl radical process (route 2). Finally, a hydrogen (or a SiMe_3 group in the case of **5f**) is abstracted from the organozinc radicals **1d** and **5d** by a RCH_2^\bullet radical to give the products **1f**, **5f**, and **5g**. Because each of these RCH_2^\bullet radicals leaves behind a $(\text{RCH}_2\text{Zn-}t\text{-BuDAB})^\bullet$



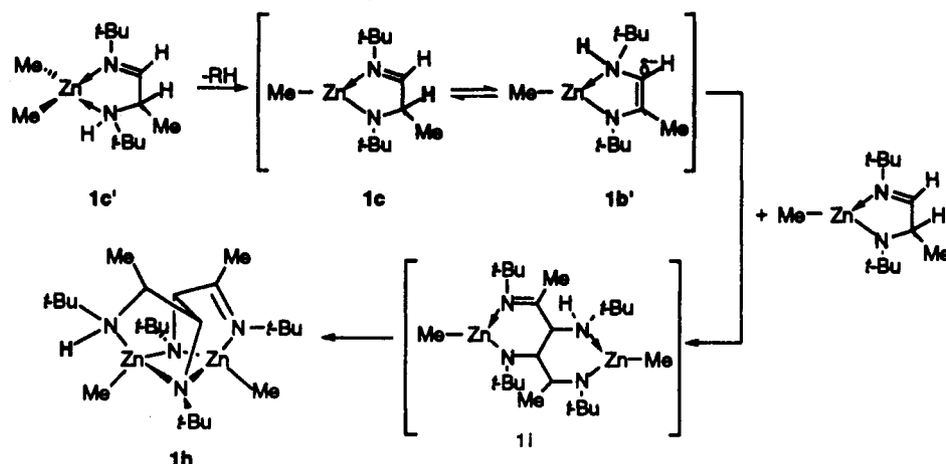
radical that dimerizes to **1e**, these dimers **1e** and **1f** should be formed in a 2:1 ratio, which indeed has been observed (the same ratio was observed in the formation of **5e** and **5f/5g**). A similar reaction has been observed for analogous dimethylaluminum-iminoketone adducts.¹⁵

The formation of SiMe_4 and $\text{Me}_3\text{SiCH}_2\text{SiMe}_3$ in the thermal decomposition of **5a** indicates that the escaped alkyl radicals take up two hydrogens, in the case of **1f** and **5g**, or one hydrogen and a trimethylsilyl group in the case of **5f**, from the initially formed C-alkylated products **1c** and **5c**.¹⁶

Irradiation Reaction. The irradiation of $\text{Me}_2\text{Zn-}t\text{-BuDAB}$ basically gives 30% of the dinuclear species **1e** and 70% of the C-alkylated product **1c**, which was completely converted into the dinuclear species **1h** during the reaction. Since **1c** has been found to be thermally unstable, we believe that this subsequent conversion is caused thermally rather than photochemically. A plausible mechanism for the formation of **1h** would involve a condensation reaction (see Scheme 4). As mentioned above, the C-alkylated compound **1c**, which was independently generated from **1c'**, is in equilibrium with its zinc-enamine tautomer **1b'**; the latter being a nucleophilic

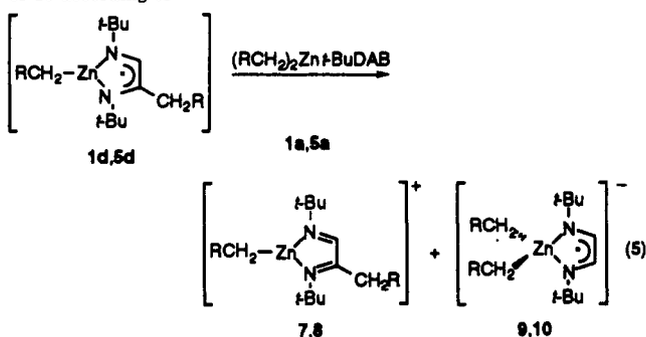
(15) van Vliet, M. R. P.; van Koten, G.; Buysingh, P.; Jastrzebski, J. T. B. H.; Spek, A. L. *Organometallics* 1987, 6, 537.

Scheme 4

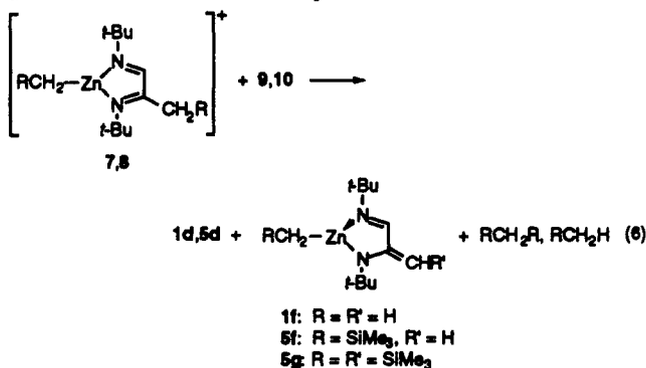


reagent then undergoes a C—C coupling reaction with a parent molecule 1c to give the zinc-aldolate 1i. Finally, a hydrogen shift from one nitrogen to another gives the unsymmetric C—C coupled dinuclear compound 1h. The hydrogen shift could formally be considered as self-rearrangement of the zinc-aldolate in order to reach a conformation wherein four-coordination of the zinc atoms is attained. The formation of 1h is a nice example of double activation of the organic substrates by a zinc reagent. A comparable condensation reaction as mentioned above was found earlier in the regioselective alkylation reaction of

(16) It must be noted that the redox potential of radical 1d or 5d with respect to that of the 1:1 coordination complexes $(RCH_2)_2Zn-t-BuDAB$ 1a or 5a is favorable for one-electron transfer (cf. the low ionization energy of $RZn-t-BuDAB$ radicals ($R = Me, Et$)).¹⁷ This would lead to the formation of cationic 7 or 8 as well as the diorganozinc radical-anion 9 or 10 according to



Reaction of cationic 7 or 8 with 9 or 10 as the nucleophile represents an alternative route to the observed products:



We thank one of the reviewers for this suggestion. However, recently, we found that radical-anion 9 or 10, when generated separately, forms the heterozincate $[RCH_2Zn(t-BuN-CH=CH-N-t-Bu)]^-$ via the escape of a RCH_2^{\cdot} radical.

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Et_2Zn with imino esters ($R'N=CH-COOEt$) resulting in the formation of β -lactams.¹⁸ The latter reaction involves condensation of the formed zinc enolate with the imine functionality of the $Et_2Zn(R'N=CH-COOEt)$ complex.

The photochemically- and or thermally-induced SET reactions 1a differ only in the degree of selectivity of the group transfer: in the thermal reaction a mixture of N- and C-alkylation products is formed, whereas the photochemical reaction resulted in C alkylation only. The highly regioselective C alkylation in the photochemical reaction supports the earlier suggestion that the T_1 state, obtained after a $S_0 \rightarrow S_1$ transition, followed by a fast heavy atom supported ISC process, is the starting point for the photochemistry in 1a.² Calculations have shown an extremely low barrier to Zn—C dissociation in the T_1 state structure. The so formed radical pair, consisting of an organozinc and an alkyl radical, is a triplet and requires a spin flip before alkylation can take place; this increases the lifetime of the alkyl radical and favors alkylation at the least hindered center of the NCCN skeleton of the organozinc radical, e.g. a carbon center. In both reactions (thermal and photochemical) the initially formed C-alkylated product 1c is the starting compound for subsequent reactions. The different amounts of second-order products formed is caused by the absence of large amounts of free alkyl radicals in the photochemical reaction. The higher temperature in the thermal reaction will enhance the formation of free radicals.

We have demonstrated that the β -aminozinc-enamido species 1b are interesting intermediates for further synthesis, i.e. in condensation reactions with other imines and aldehydes, leading to, e.g., indolizines,¹⁹ 2- and 3-pyrrolidinones,⁸ and 2-azetidiones.⁷ Moreover, the dimeric zinc species 1e, 1h, and 5e could be of interest in organometallic chemistry, since the newly formed tetradentate dianionic ligand may have some potential in stabilizing transition metal complexes.

Experimental Section

General Data. All experiments were carried out in a dry and oxygen-free nitrogen atmosphere, using standard

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Schlenk techniques. Solvents were carefully dried and distilled from sodium/benzophenone prior to use. All starting chemicals were purchased from Aldrich Chemical Co. or Janssen Chimica. The starting materials *N,N'*-di-*tert*-butyl-1,4-diaza-1,3-butadiene (*t*-BuDAB),²⁰ Me₂Zn-*t*-BuDAB,² (Me₃SiCH₂)₂Zn,²¹ and Bu₂Zn²² were prepared according to literature procedures. ¹H and ¹³C NMR spectra were recorded on Bruker AC-200 or AC-300 spectrometers in C₆D₆, using SiMe₄ as an external standard (0.0 ppm). Elemental analyses were performed by Dornis und Kolbe, Mikroanalytisches Laboratorium, Mülheim a.d. Ruhr, FRG.

(Me₃SiCH₂)₂Zn-*t*-BuDAB (**5a**). To a stirred solution of *t*-BuDAB (1.68 g, 10 mmol) in diethyl ether (50 mL) was added 1 equiv of (Me₃SiCH₂)₂Zn (2.39 g, 10 mmol). The solvent was removed *in vacuo* at ambient temperature to afford **5a** as an orange solid in quantitative yield; 4.07 g **5a** was crystallized from diethyl ether or hexane. ¹H NMR: δ 7.25 (s, 2H, N=CH), 1.14 (s, 18H, *t*-Bu), 0.27 (s, 18H, SiCH₃), -0.78 (s, 4H, ZnCH₂). ¹³C NMR: δ 152.9 (N=CH), 58.7 (C(CH₃)₃), 30.1 (C(CH₃)₃), 4.0 (Si(CH₃)₃), 0.4 (ZnCH₂). UV-Vis: λ_{max} = 390 nm.

General Procedure for the Thermal Decomposition Reaction of R₂Zn-*t*-BuDAB. The coordination complex R₂Zn-*t*-BuDAB (R = Me, CH₂SiMe₃) (10 mmol) was dissolved in benzene (50 mL) and heated for 6 h at 80 °C. The solvent was removed *in vacuo*, leaving a dark brown sticky residue. The ¹H NMR spectrum of the residue obtained from **1a** showed that the product is a mixture of **1b** (25%), **1e** (50%), and **1f** (25%), whereas the ¹H NMR spectrum of the product obtained from **5a** revealed that the residue was a mixture of **5e** (65%), **5f** (7%), and **5g** (28%). The dimeric species were crystallized from the product mixtures in hexane at -20 °C as white crystals.⁷

¹H NMR of **1b**: δ 6.67 (d, *J* = 4.4 Hz, 1H, CH=CH), 4.29 (d, *J* = 4.4 Hz, 1H, CH=CH), 2.07 (s, 3H, N-CH₃), 1.22, 0.99 (s, 9H, *t*-Bu), -0.22 (s, 3H, ZnCH₃). ¹³C NMR of **1b**: δ 139.7 (CH=CH), 100.3 (CH=CH), 56.2, 50.2 (C(CH₃)₃), 35.7 (N-Me), 31.2, 23.7 (*t*-Bu), -12.5 (Zn-Me). ¹H NMR of **1f**: δ 7.42 (s, 1H, N=CH), 4.59, 4.18 (s, 2 × 1H, C=CHH'), 1.48, 0.88 (s, 9H, *t*-Bu), -0.30 (s, 3H, ZnCH₃). ¹³C NMR of **1f**: characteristic signals δ 168.2 (N=CH), not observed (C=CH₂), 94.7 (C=CH₂), -9.7 (Zn-Me).

¹H NMR of **1e**: δ 7.61 (d, *J* = 2.8 Hz, 2H, N=CH), 2.72 (d, *J* = 2.8 Hz, 2H, N-CH), 1.08, 1.04 (s, 2 × 18H, C(CH₃)₃), -0.01 (s, 6H, ZnCH₃). ¹³C NMR of **1e**: δ 170.8 (N=CH), 60.1 (N-CH), 57.6, 52.2 (C(CH₃)₃), 32.6, 29.5 (C(CH₃)₃), -11.3 (ZnCH₃). Anal. Calcd for C₂₂N₄Zn₂: C, 53.13; H, 9.32; N, 11.26. Found: C, 53.10; H, 9.38; N, 11.18.

¹H NMR of **5e**: δ 7.59 (d, *J* = 2.7 Hz, 2H, N=CH), 2.64 (d, *J* = 2.7 Hz, 2H, N-CH), 1.12, 1.04 (s, 18H, C(CH₃)₃), 0.46 (s, 18H, Si(CH₃)₃), -0.32, -0.44 (dd, *J* = 12.3, 2 × 2H, ZnCHH'Si(CH₃)₃). ¹³C NMR of **5e**: δ 171.42 (N=CH), 59.6 (N-CH), 57.8, 52.4 (C(CH₃)₃), 32.8, 29.9 (C(CH₃)₃), 4.4 (Si(CH₃)₃), -3.4 (ZnCH₂). Anal. Calcd for C₂₈H₆₂N₄Si₂Zn: C, 52.40; H, 9.74; N, 8.73. Found: C, 51.93; H, 9.90; N, 8.85.

¹H NMR of **5f** (only characteristic signals): δ 7.48 (s, 1H, N=CH), 4.64 (s, 1H, C=CHH'), 4.27 (s, 1H, C=CHH'). ¹³C signals not observed.

¹H NMR of **5g**: δ 8.11 (s, 1H, N=CH), 5.04 (s, 1H, C=CHSiMe₃), 1.53, 0.96 (s, 18H, C(CH₃)₃), 0.31 (s, 9H, ZnCH₂Si(CH₃)₃), 0.26 (s, 9H, ZnCH₂Si(CH₃)₃), -0.34 (s, 2H, ZnCH₂Si(CH₃)₃). ¹³C NMR of **5g**: δ 166.4 (N=CH), 155.5 (N-C=CH), 105.0 (C=C(H)SiMe₃), ¹J_{Si-C} = 24.7 Hz) 57.7, 52.6 (C(CH₃)₃), 33.2, 30.6 (C(CH₃)₃), 4.5 (Si(CH₃)₃), -1.8 (ZnCH₂).

Independent Synthesis of the Dinuclear Species 1h (R = Me), 2h (R = Et), and 6h (R = Bu). To a stirred solution of *t*-BuN=CH-CH(CH₃)-NH(*t*-Bu)¹¹ (1.84 g, 10 mmol) in diethyl ether (25 mL) was added 1 equiv of R₂Zn (R = Me, Et, Bu). After stirring for 5 min the solvent was removed *in vacuo* at ambient temperatures, leaving the 1:1 coordination complexes **1c'**, **2c'**, and **6c'** as yellow solids in quantitative yield. (The products were obtained as white solids after crystallization from hexane.) Thermally instable **1c'**, **2c'**, and **6c'** were dissolved in hexane and heated at 60 °C for 30 min. The solvent was removed *in vacuo* to afford white solids. The products were crystallized from hexane to obtain analytically pure compounds.

¹H NMR of **1c'**: δ 6.88 (d, *J* = 2.1 Hz, 1H, N=CH), 2.96 (m, 1H, N-C(CH₃)H), 1.13, 0.93 (s, 2 × 9H, C(CH₃)₃), 0.82 (d, *J* = 7.2 Hz, 3H, N-CH(CH₃)), -0.31 (s, 6H, ZnCH₃). ¹H NMR of **2c'**: δ 6.83 (d, *J* = 2.0 Hz, 1H, N=CH), 2.90 (m, 1H, N-C(CH₃)H), 1.74 (t, 6H, ZnCH₂CH₃), 1.12, 0.93 (s, 2 × 9H, C(CH₃)₃), 0.78 (d, *J* = 7.2 Hz, 3H, N-C(CH₃)₃), 0.39 (q, 4H, ZnCH₂CH₃).

¹H NMR of **1h**: δ 3.18 (s, 1H, N-CH-C(CH₃)-N), 2.88 (ddq, *J* = 2.5 Hz, 6.3 Hz, 12 Hz, 1H, N-CH(CH₃)-CH-N), 2.04 (d, *J* = 2.5 Hz, 1H, N-CH(CH₃)-CH-N), 1.75 (s, 2H, N-CH-C(CH₃)=N), 1.26, 1.20, 1.09, 1.07 (s, 4 × 9H, C(CH₃)₃), 1.01 (d, *J* = 6.3 Hz, 3H, N-CH(CH₃)-CH-N), -0.04, -0.09 (s, 2 × 3H, ZnCH₃). ¹³C NMR of **1h**: δ 183.3 (N=CCH₃), 66.2 (N-CH-CCH₃), 66.0 (N-CH(CH₃)-CH-N), 56.4, 53.4, 54.6, 52.6 (C(CH₃)₃), 53.5 (NCH(CH₃)-CH-N), 33.5, 32.2, 30.9, 30.4 (C(CH₃)₃), 21.7 (N-CH(CH₃)-CH-N), 19.6 (N=CCH₃), -8.4 (ZnCH₃). Anal. Calcd for C₂₄H₅₂N₄Zn₂: C, 54.65; H, 9.94; N, 10.62. Found: C, 54.82; H, 9.79; N, 10.45.

¹H NMR of **2h**: δ 3.11 (s, 21H, N-CH-C(CH₃)=N), 2.88 (ddq, *J* = 2.4 Hz, 6.2 Hz, 12 Hz, 1H, N-CH(CH₃)-CH-N), 2.02 (d, *J* = 2.4 Hz, 1H, N-CH(CH₃)-CH-N), 1.85 (q, 6H, ZnCH₂CH₃), 1.74 (s, 6H, N-CH-C(CH₃)=N), 1.24, 1.20, 1.07, 1.05 (s, 4 × 9H, C(CH₃)₃), 0.98 (d, *J* = 6.2 Hz, 3H, N-CH(CH₃)-CH-N), 0.79 (m, 4H, ZnCH₂CH₃). ¹³C NMR of **2h**: δ 183.2 (N=CCH₃), 66.2 (N-CH-CCH₃), 66.1 (N-CH(CH₃)-CH), 56.1 (C(CH₃)₃), 54.4 (N-CH(CH₃)-CH-N), 53.2, 52.9, 52.0 (C(CH₃)₃), 33.6, 32.3, 30.7, 30.5 (C(CH₃)₃), 19.7 (N=CCH₃), 21.7 (N-CH(CH₃)-CH-N), 14.5, 14.3 (ZnCH₂CH₃), 4.7, 4.3 (ZnCH₂CH₃). Anal. Calcd for C₂₈H₅₆N₄Zn₂: C, 56.63; H, 9.50; N, 10.16. Found: C, 56.53; H, 10.06; N, 9.98.

¹H NMR of **6h**: δ 3.13 (s, 1H, N-CH-C(CH₃)=N), 2.86 (ddq, *J* = 2.3 Hz, 6.3 Hz, 12 Hz, 1H, N-CH(CH₃)-CH-N), 2.07 (m, 4H, butyl), 2.03 (d, *J* = 2.3 Hz, 1H, N-CH(CH₃)-CH-N), 1.79 (m, 4H, butyl), 1.75 (s, 3H, N-CH-C(CH₃)=N), 1.25 (m, 4H, butyl), 1.13 (m, 6H, butyl), 1.24, 1.21, 1.08, 1.07 (s, 4 × 9H, C(CH₃)₃), 0.99 (d, *J* = 6.3 Hz, 3H, N-CH(CH₃)-CH-N), 0.79-0.58 (m, 4H, ZnCH₂). ¹³C NMR of **6h**: δ 183.1 (N=C-CH₃), 66.3 (N-CH-CCH₃), 66.1 (N-CH(CH₃)-CH), 56.1 (C(CH₃)₃), 54.4 (N-CH(CH₃)-CH-N), 53.2, 53.0, 52.1

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(C(CH₃)₃), 33.5 (C(CH₃)₃), 33.2, 33.1 (CH₂), 32.3 (C(CH₃)₃), 31.2, 31.1 (CH₂), 30.7, 30.6 (C(CH₃)₃), 21.7 (N—CH—(CH₃)—CH—N), 19.7 (N=CCH₃), 14.4 (CH₃), 13.9, 13.4 (ZnCH₂). Molecular weight determination in benzene: calcd weight, 610; found, 568.

Irradiation of Me₂Zn-*t*-BuDAB (1a). A stirred solution of 1a (1.32 g, 5 mmol) in diethyl ether (50 mL) in a Pyrex tube at 20 °C was irradiated for 45 min with a 300-W high-pressure Hg lamp. The solvent was removed *in vacuo*, leaving a yellow solid. The ¹H NMR spectrum of the residue revealed that the product was a mixture of the symmetric dinuclear species 1d (30%) and the

unsymmetric dinuclear species 1h (70%). We have been unable to separate the two products (see above for ¹H and ¹³C NMR data for 1d and 1h).

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