

The Role of Neutral and Radical Anionic Organozinc Complexes in the Alkylation Reactions of 1,4-Diaza-1,3-butadienes with Diorganozinc Compounds[†]

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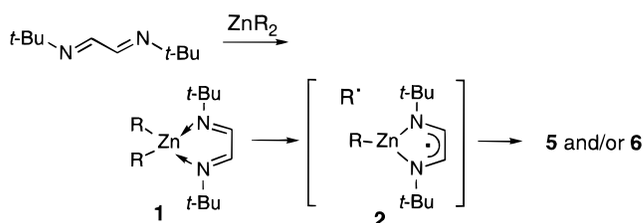
We have earlier postulated the intermediacy of organozinc radical species in the regioselective alkylation reactions of 1,4-di-*tert*-butyl-1,4-diaza-1,3-butadiene (*t*-BuNCHCHN-*t*-Bu) with diorganozinc compounds (ZnR₂). To verify these postulates, we have prepared and studied the neutral organozinc radical complex MeO(CH₂)₃Zn(*t*-BuNCHCHN-*t*-Bu) (**A**) and two diorganozinc radical-anionic complexes, K[R₂Zn(*t*-BuNCHCHN-*t*-Bu)] (R = Me (**8a**), Et (**8b**)). **A** was prepared in situ by the reaction of *t*-BuNCHCHN-*t*-Bu with bis-(methoxypropyl)zinc. Complexes **8** were prepared by reducing the 1:1 coordination complex ZnMe₂(*t*-BuNCHCHN-*t*-Bu) (**1a**) with potassium or by the nucleophilic addition of K(*t*-BuNCHCHN-*t*-Bu) to ZnR₂ (R = Me, Et). The resulting radical-anionic complexes are thermally unstable and readily undergo an intermolecular single-electron transfer, giving a mixture of the metallacyclic heteroleptic zincate complexes K[ZnR(*t*-BuNCHCHN-*t*-Bu)] (**9**) and K[ZnR(*t*-BuNC(R)CHN-*t*-Bu)] (**10**). The ratio in which the latter complexes are formed depends on the R group and on the concentration and the temperature. Alternatively, **9** has been prepared separately from the reduction of [ZnR(*t*-BuNCHCHN-*t*-Bu)]₂ (**4**) with 2 equiv of potassium. The crystal structures of two complexes of **9**, i.e. {**9a**(THF)}_n and {**9c**(Et₂O)}_{1/2}}_n, have been determined. Both complexes form linear coordination polymers containing alternating potassium cations and zinc–diazabutadiene anions.

Introduction

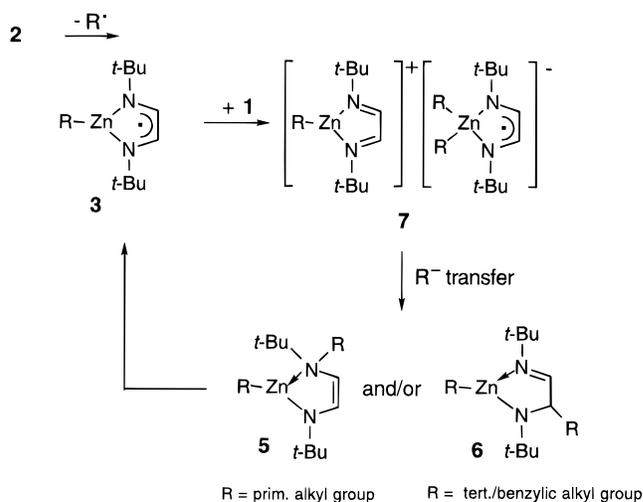
It is well-known that 1,4-di-*tert*-butyl-1,4-diaza-1,3-butadiene (*t*-BuNCHCHN-*t*-Bu) is alkylated by diorganozinc compounds (ZnR₂) with high regioselectivity to either the nitrogen-alkylated product ZnR(*t*-BuN(R)CHCHN-*t*-Bu) (R = primary alkyl group) or the carbon-alkylated product ZnR(*t*-BuNCH(R)CHN-*t*-Bu) (R = tertiary/benzylic alkyl group). Two possible mechanisms for this alkylation reaction have been put forward, i.e. a radical and a polar mechanism (see Scheme 1).² Earlier studies have shown that the initial step common to both mechanisms is the formation of the thermally unstable 1:1 coordination complex ZnR₂(*t*-BuNCHCHN-*t*-Bu) (**1**), which undergoes an intramolecular single-electron-transfer (SET) reaction to give a radical pair, [R[•]/ZnR(*t*-BuNCHCHN-*t*-Bu)[•]] (**2**), in a solvent cage. In the radical mechanism **2** collapses in

Scheme 1

Radical mechanism:



Polar mechanism:



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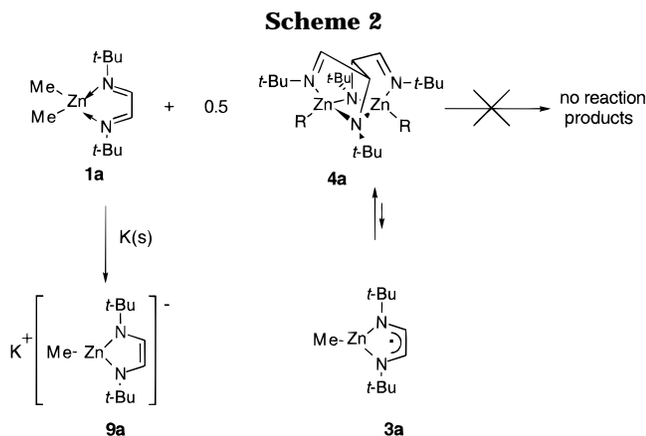
^{||} Address correspondence regarding the crystallography to this author.

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the solvent cage, resulting in the nitrogen- (**5**) and carbon-alkylated (**6**) products, whereas in the polar



mechanism a steady-state concentration of the free organozinc radical is formed. This transfers its unpaired electron to the initial 1:1 coordination complex $\text{ZnR}_2(t\text{-BuNCHCHN-}t\text{-Bu})$ (**1**), to give an organozinc cation/diorganozinc radical-anion pair, $[\text{ZnR}(t\text{-BuNCHCHN-}t\text{-Bu})][\text{ZnR}_2(t\text{-BuNCHCHN-}t\text{-Bu})]^-$ (**7**). Nucleophilic attack of an alkyl group of the diorganozinc radical anion $[\text{ZnR}_2(t\text{-BuNCHCHN-}t\text{-Bu})]^-$ on the organozinc cation $[\text{ZnR}(t\text{-BuNCHCHN-}t\text{-Bu})]^+$ gives the alkylation products with simultaneous regeneration of the free organozinc radical (**3**). In the polar mechanism **5** and **6** can be regarded as 1,2- and 1,4-addition products, respectively.

So far, both mechanisms are supported by the isolation of the 1:1 coordination complex $\text{ZnMe}_2(t\text{-BuNCHCHN-}t\text{-Bu})$ (**1a**)^{1c} and of $[\text{ZnR}(t\text{-BuNCHCHN-}t\text{-Bu})]_2$ (**R** = Me (**4a**), Et (**4b**)). The latter exists in solution in equilibrium with the corresponding neutral organozinc radicals $[\text{ZnR}(t\text{-BuNCHCHN-}t\text{-Bu})]^\bullet$ (**3**) (cf. Scheme 2).^{1,3} Although the radical mechanism has been used as a working hypothesis for the reaction of $t\text{-BuNCHCHN-}t\text{-Bu}$ with ZnR_2 , the proposal of an alternative, polar mechanism is justified by the EPR detection of an unprecedented paramagnetic species during the alkylation reaction in THF. The EPR resonance pattern in THF can only be assigned to a tetrahedrally surrounded organozinc radical complex.^{1c} In this paper we report the preparation of two different types of complexes having this geometry at zinc, i.e. a model for a solvated neutral organozinc radical and representative examples of diorganozinc radical-anionic complexes. To investigate the role of THF in the formation of the former, we prepared a neutral organozinc radical with a potentially C,O-chelating alkyl group, in which the methoxy substituent may be regarded as a suitably-positioned "solvent" donor molecule. For the preparation of the radical-anionic diorganozinc complexes we have made use of the ability of the $t\text{-BuNCHCHN-}t\text{-Bu}$ ligand to stabilize electron-rich zinc α -diimine complexes, as can be concluded from our recent studies on mono- and dianionic homoleptic zinc complexes.⁴

Results

Neutral Organozinc Complexes. The EPR spectrum of the reaction mixture of $t\text{-BuNCHCHN-}t\text{-Bu}$ with

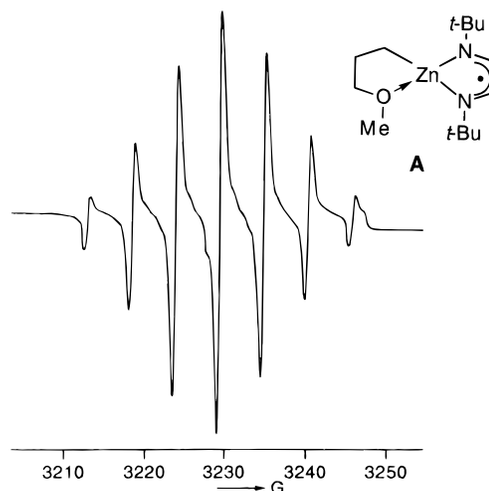


Figure 1. EPR spectrum of the intermediate species (**A**) in the reaction mixture of $t\text{-BuNCHCHN-}t\text{-Bu}$ and $\text{Zn}((\text{CH}_2)_3\text{OME})_2$ in Et_2O (10 G = 1 mT).

$\text{Zn}[(\text{CH}_2)_3\text{OME}]_2$ ⁵ in Et_2O at room temperature consists of a resonance pattern of seven broad lines without a hyperfine structure. This is consistent with the presence of an unpaired electron which couples to two magnetically equivalent nitrogen nuclei ($I = 1$) and two magnetically equivalent protons ($I = 1/2$) with ^1H and ^{14}N coupling constants of 0.49 and 0.59 mT, respectively. We assign this spectrum to the C,O-chelate-bonded neutral organozinc radical complex **A** (see Figure 1).

Radical-Anionic Diorganozinc Complexes. $\text{ZnMe}_2(t\text{-BuNCHCHN-}t\text{-Bu})$ (**1a**) is stable at room temperature, in contrast to the corresponding complexes of other ZnR_2 compounds. In **1a**, homolytic Zn–C bond cleavage can only be induced thermally or photochemically.⁶ The stability of **1a** enables a more detailed study of its reactivity toward various reducing agents that might be used to prepare radical-anionic diorganozinc complexes. First, the reduction of **1a** with the neutral organozinc radical complex $[\text{ZnR}(t\text{-BuNCHCHN-}t\text{-Bu})]^\bullet$ (**3a**) was studied by reacting **1a** with 0.5 equiv of the dimer of **3a**, i.e. $[\text{ZnMe}(t\text{-BuNCHCHN-}t\text{-Bu})]_2$ (**4a**), at room temperature (see Scheme 2). We earlier demonstrated that dimeric complexes **4** in solution are in equilibrium with two coordinatively unsaturated neutral organozinc radical complexes $[\text{ZnR}(t\text{-BuNCHCHN-}t\text{-Bu})]^\bullet$ (**3**).^{1,3} The EPR spectrum of a mixture of **1a** and **4a** in Et_2O did show the presence of the unreacted $[\text{ZnMe}(t\text{-BuNCHCHN-}t\text{-Bu})]^\bullet$ (**3a**).^{1c} Moreover, the ^1H NMR spectrum of the mixture revealed the presence of the starting materials **1a** and **4a** in a 2:1 molar ratio, while no alkylation products were detected. In contrast to this inertness of **1a** to **3a**, the reaction of **1a** with potassium as reducing agent gave, according to the ^1H NMR spectrum, the heteroleptic metallacyclic organo-diamido zincate anion $\text{K}[\text{ZnMe}(t\text{-BuNCHCHN-}t\text{-Bu})]$ (**9a**) (see Scheme 2).

9a has also been obtained by two other procedures. First, the reaction of the potassium salt of $t\text{-BuNCHCHN-}t\text{-Bu}$, $\text{K}(t\text{-BuNCHCHN-}t\text{-Bu})$,⁷ with 1 equiv of ZnMe_2 gave **9a** quantitatively. The application of this proce-

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Table 3. Selected Bond Lengths and Bond Angles for {9c(Et₂O)_{1/2}}_n^a

Bond Lengths (Å)			
K(1)–N(1)	2.990(1)	K(2)–N(1)	3.171(12)
K(1)–N(2)	3.081(13)	K(2)–N(2)	3.248(14)
K(1)–C(5)	2.892(15)	K(2)–C(5)	2.958(15)
K(1)–C(6)	2.917(14)	K(2)–C(6)	3.034(14)
Zn(1)–C(20)	1.974(17)	K(2)–O(30)	2.777(16)
Zn(1)–N(1)	1.943(11)	N(1)–C(5)	1.393(19)
Zn(1)–N(2)	1.944(13)	N(2)–C(6)	1.360(19)
C(5)–C(6)	1.36(2)		
Bond Angles (deg)			
N(1)–Zn(1)–C(20)	135.5(6)	N(1)–C(5)–C(6)	114.4(14)
N(2)–Zn(1)–C(20)	138.6(6)	N(2)–C(6)–C(5)	117.9(13)
N(1)–Zn(1)–N(2)	85.5(5)		

^a The estimated standard deviations of the last significant digits are shown in parentheses.

(360.01 and 357.77°, respectively), close to the expected 360°. Clear differences are observed in the Zn–N, N–C, and C–C bond distances of both units. The Zn–N and C–C bond distances in the Zn(11)NCCN chelate ring are significantly shorter than those found in the Zn(31)NCCN ring. A comparison of the Zn–N, N–C, and C–C bond distances with those found in K(thf)₃[Zn(*t*-BuNCHCHN-*t*-Bu)₂]⁴ suggests a dianionic enamide character for the *t*-BuNCHCHN-*t*-Bu ligand in Zn(11)-NCCN. The distances found in Zn(31)NCCN are comparable to those found in the radical-anion *t*-BuNCHCHN-*t*-Bu ligands in Zn(*t*-BuNCHCHN-*t*-Bu)₂⁹ and K(thf)₃[Zn(*t*-BuNCHCHN-*t*-Bu)₂]⁴.

The crystallographic repeating unit in {9c(Et₂O)_{1/2}}_n consists of two equivalent [Zn(CH₂Ph)(*t*-BuNCHCHN-*t*-Bu)][–] monoanions and two nonequivalent potassium cations, each on a crystallographic 2-fold axis. The nonequivalency of the potassium cations is a result of the nonsymmetrical η³ + η⁴ coordination of the ZnNCCN chelate ring to two potassium cations. K(1) is twice η⁴-coordinated *via* interactions with the N(1), N(2), C(5), and C(6) atoms of two [Zn(CH₂Ph)(*t*-BuNCHCHN-*t*-Bu)][–] units, while K(2) is twice η³-coordinated *via* interactions with the N(1), C(5), and C(6) atoms of these units. K(1) is also coordinated to a molecule of Et₂O to complete its coordination sphere.

The ¹H NMR spectra of **9** show a characteristic singlet resonance in the olefinic region and a singlet for two identical *t*-Bu groups, comparable to the resonances found in, e.g., Li₂(*t*-BuNCHCHN-*t*-Bu),^{10a} M(*t*-BuNCHCHN-*t*-Bu) (M = Si^{II}, Ge^{II}),^{10b,c} and K₂(thf)₂[Zn(*t*-BuNCHCHN-*t*-Bu)₂]⁴. The ¹H NMR spectra of the alkylated zincate complexes **10** show the same characteristic singlet resonance in the olefinic region as was found for **9**, a signal for the alkyl group attached to the olefinic bond, and two singlets for the chemically nonequivalent *t*-Bu groups. The chemical shifts observed for the α-protons of the zinc-bonded alkyl groups in **9** and **10** are at higher field relative to those in related neutral organozinc complexes. In comparison with the starting dimeric organozinc complexes **4** an upfield shift of 0.57–0.81 ppm is observed for the α-protons of the

zinc-bound alkyl groups, probably due to the enhanced negative charges on the alkyl groups in the zincate complexes **9** and **10**.

The EPR spectra of the reaction mixture of K(*t*-BuNCHCHN-*t*-Bu) and ZnR₂ (R = Me, Et) showed the presence of paramagnetic species (a similar reaction with Zn(CH₂Ph)₂ was EPR silent). The relatively broad signals hamper the precise determination of the *g* values. However, after approximately 30 min the EPR signals had disappeared completely. Both species show identical resonance patterns, which is consistent with an unpaired electron being coupled to two magnetically equivalent nitrogen nuclei (*I* = 1) and two magnetically equivalent protons (*I* = 1/2) with ¹H and ¹⁴N coupling constants of 0.45 and 0.64 mT, respectively, without hyperfine splitting. Therefore, we assigned these EPR spectra to the radical-anionic diorganozinc complexes K[ZnR₂(*t*-BuNCHCHN-*t*-Bu)] (R = Me (**8a**), Et (**8b**)). The EPR spectra of **8** are different from that of K(*t*-BuNCHCHN-*t*-Bu); the latter shows ¹H and ¹⁴N coupling constants of 0.43 and 0.56 mT, respectively.⁷

Discussion

Neutral Organozinc Radicals. The EPR spectrum of the reaction mixture of *t*-BuNCHCHN-*t*-Bu with Zn[(CH₂)₃OMe]₂ in Et₂O shows a resonance pattern of seven broad lines without hyperfine structure. This lack of hyperfine coupling in the EPR spectrum indicates the presence of a species with a tetrahedral geometry at the zinc atom. We believe that this is brought about by intramolecular O–Zn coordination to give a C,O-chelated neutral organozinc radical. The intramolecular coordination of the methoxy substituent in the alkyl group illustrates the Lewis-acidic character of the zinc atom in neutral organozinc radicals and leads to a deformation of the planar geometry around zinc. The formation of such a species was investigated to mimic the postulated formation of the solvated tetra-coordinated neutral organozinc radical [ZnR(*t*-BuNCHCHN-*t*-Bu)(THF)][•] in THF solution. The presence of such a solvated neutral organozinc radical species during the alkylation reaction in THF is in agreement with the aforementioned EPR data of the unknown key intermediate² and therefore supports the postulated radical mechanism.

Radical-Anionic Diorganozinc Complexes. The observed stability of ZnMe₂(*t*-BuNCHCHN-*t*-Bu) (**1a**) toward reduction with [ZnMe(*t*-BuNCHCHN-*t*-Bu)][•] (**3a**) demonstrates that the ionization energy of 6 eV for **3a**¹¹ is too high to let it act as a one-electron donor for the reduction of **1a**. However, the reduction of **1a** with potassium *in situ* gives radical-anionic K[ZnMe₂(*t*-BuNCHCHN-*t*-Bu)] (**8a**), which was thermally converted into the diamagnetic zincate species K[ZnMe(*t*-BuNCHCHN-*t*-Bu)] (**9a**). Similar reductions of 1:1 diorganozinc coordination complexes of 2,2'-bipyridine and 3,4,7,8-tetramethylphenanthroline ligands with potassium *in situ* gave radical-anionic diorganozinc complexes, which have been studied with EPR.¹² Counting of π-electrons in **9a** suggested an alternative preparative route involving the reduction of dimeric [ZnR(*t*-

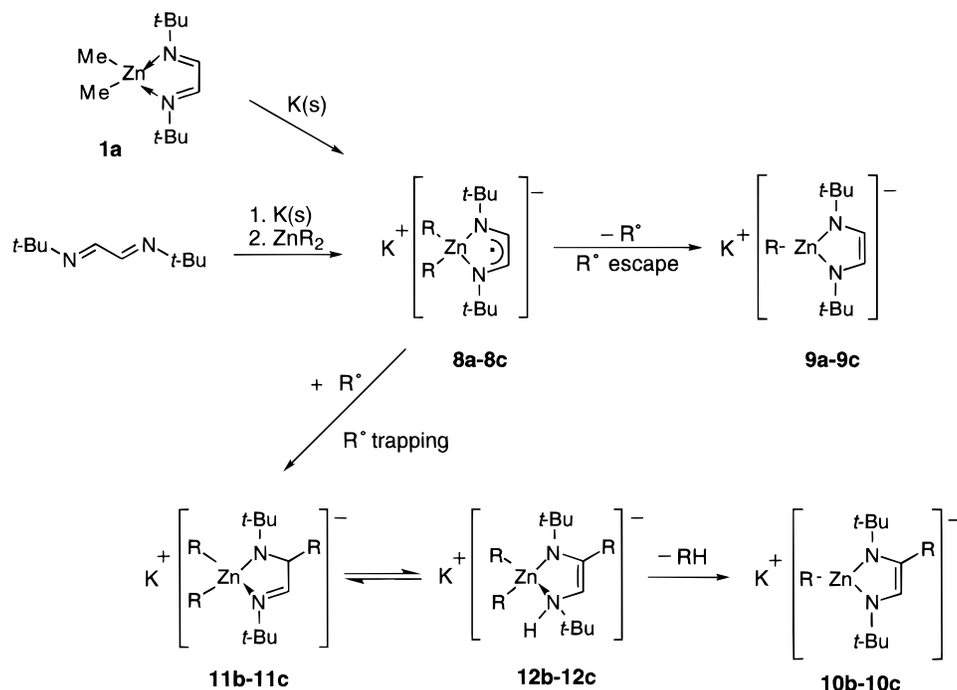
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Scheme 3



$\text{BuNCHCHN-}t\text{-Bu}]_2$ with 2 equiv of potassium. Most likely, the neutral organozinc radical is the active species in this reaction. Compound **9a** was also prepared by the reaction of $\text{K}(t\text{-BuNCHCHN-}t\text{-Bu})$ with ZnMe_2 , while this reaction with ZnR_2 ($\text{R} = \text{Et}, \text{CH}_2\text{Ph}$) gave mixtures of products, from which $\text{K}[\text{ZnR}(t\text{-BuNCHCHN-}t\text{-Bu})]$ (**9**) and $\text{K}[\text{ZnR}(t\text{-BuNC}(\text{R})\text{CHN-}t\text{-Bu})]$ (**10**) have been isolated.

Complexes **9** and **10** are the first examples of heteroleptic organozincate complexes, containing a chelate-bonded dianionic diamide ligand. So far, the only known metallacyclic zincate complexes are $[\text{Li}(\text{L})_2\text{Zn}(-\text{CHR}(\text{CHR})_n\text{CHR}-)]_2$ ($\text{L} = \text{Et}_2\text{O}; 2\text{L} = \text{Me}_2\text{NCH}_2\text{CH}_2\text{NMe}_2, \text{MeOCH}_2\text{CH}_2\text{OMe}; n = 2, 3; \text{R} = \text{H}, \text{Me}$).¹³ Earlier reports on heteroleptic zincates concern the crystal structure of $[\text{KZnEt}_2(\text{O-}t\text{-Bu})]_2$, derived from ZnEt_2 and $\text{KO-}t\text{-Bu}$,¹⁴ and the NMR characterization of $\text{LiZnBr}_2(\text{C}(\text{SiMe}_3)_3)$, prepared from ZnBr_2 and $\text{LiC}(\text{SiMe}_3)_3$.¹⁵

A plausible mechanism for the formation of the zincate complexes **9** and **10** is outlined in Scheme 3. Both the nucleophilic addition reaction of $\text{K}(t\text{-BuNCHCHN-}t\text{-Bu})$ to various ZnR_2 compounds and the reduction reaction of **1a** with potassium result in the *in situ* formation of thermally unstable **8**. When the temperature is raised above its decomposition temperature, **8** undergoes a homolytic cleavage of one of the zinc-carbon bonds to give an alkyl radical (R^\bullet) and a biradical organozinc species. Recombination of these two unpaired electrons gives, after subsequent rearrangement, the diamagnetic zincate complex **9** (the R^\bullet -escape product). For $\text{R} = \text{Et}, \text{CH}_2\text{Ph}$ these alkyl radicals are partly trapped by collapsing with **8** to give $\text{K}[\text{ZnR}_2(t\text{-BuNCH}$

$(\text{R})\text{CHN-}t\text{-Bu}]$ (**11**). Complex **11** is thought to be in equilibrium with its tautomer, $\text{K}[\text{ZnR}_2(t\text{-BuNC}(\text{R})\text{CHN}(\text{H})-t\text{-Bu})]$ (**12**). A subsequent intramolecular hydrolysis of one of the zinc-carbon bonds by the amino hydrogen in **12** finally results in the formation of the zincate **10** (the R^\bullet -trapping product).

Similar imino-enamino tautomerization processes have been reported for the carbon-alkylated organozinc compounds $\text{ZnR}(t\text{-BuNCH}(\text{R})\text{CHN-}t\text{-Bu})$, which are in equilibrium with their enamino tautomers $\text{ZnR}(t\text{-BuNC}(\text{R})\text{CHN}(\text{H})-t\text{-Bu})$.^{2,6} According to this proposed mechanism for the formation of **9** and **10**, the maximum yield of the alkylated zincates **10** is 50% with respect to **8**. Therefore, the formation of **10** in yields lower than 50% must be accompanied by a loss of alkyl radicals. Attempts to trap these alkyl radicals as their secondary products have as yet been unsuccessful. The quantitative formation of **9a** without concurrent formation of **10a**, from both the nucleophilic addition reaction and the reduction reaction, supports the bimolecular reaction pathway for the formation of the latter. The homolytic cleavage of a zinc-carbon bond in thermally unstable **8a** gives **9a** and a methyl radical. The latter are highly reactive at room temperature, which results in radical escape from the solvent cage, rather than a collapse with **8a**. The favored formation of **9b** with respect to **10b** at low temperature and at low concentration also supports the proposed bimolecular process for the formation of **10b**.

An interesting aspect of the solid-state structures of $\{\mathbf{9a}(\text{THF})\}_n$ and $\{\mathbf{9c}(\text{Et}_2\text{O})_{1/2}\}_n$ is that they display three different coordination modes for the monoanionic $[\text{ZnR}(t\text{-BuNCHCHN-}t\text{-Bu})]^-$ units. The significant differences of the Zn-N, N-C, and C-C bond distances of the two nonequivalent $[\text{ZnMe}(t\text{-BuNCHCHN-}t\text{-Bu})]^-$ units in polymeric $\{\mathbf{9a}(\text{THF})\}_n$ are a consequence of the fact that the Zn(11)-containing units act as two-electron donors, while the Zn(31)-containing units act as six-electron donors. The $[\text{Zn}(\text{CH}_2\text{Ph})(t\text{-BuNCHCHN-}t\text{-Bu})]^-$

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units in $\{9c(\text{Et}_2\text{O})_{1/2}\}_n$ act as four-electron donors. The potassium cations in solvent-free **9** are most likely twice η^4 -coordinated to two $[\text{ZnR}(t\text{-BuNCHCHN}-t\text{-Bu})]^-$ units, both acting as six-electron donors. An increase of the electron-donating properties is accompanied by a shortening of the C–N bond distances in the series $[\text{Zn}(11)\text{-Me}(t\text{-BuNCHCHN}-t\text{-Bu})]^- (2 e^-) > [\text{Zn}(\text{CH}_2\text{Ph})(t\text{-BuNCHCHN}-t\text{-Bu})]^- (4 e^-) > [\text{Zn}(31)\text{Me}(t\text{-BuNCHCHN}-t\text{-Bu})]^- (6 e^-)$. The apparent radical-anionic character of the $t\text{-BuNCHCHN}-t\text{-Bu}$ ligand in the $[\text{Zn}(31)\text{Me}(t\text{-BuNCHCHN}-t\text{-Bu})]^-$ unit of $\{9a(\text{THF})\}_n$ is caused by a shift of its π -electron density into the direction of both potassium cations. The ^1H NMR spectra of **9** in THF solution show one singlet for the olefinic protons. This suggests that in THF all $[\text{ZnR}(t\text{-BuNCHCHN}-t\text{-Bu})]^-$ units in the polymer act as symmetrical η^2 -coordinated two-electron donors or, alternatively, that the polymeric structures are completely dissociated into monomeric ion-separated solvated complexes.

Concluding Remarks

The results presented in this paper provide indirect evidence for a radical mechanism in the alkylation reactions of $t\text{-BuNCHCHN}-t\text{-Bu}$ with diorganozinc compounds. The paramagnetic species detected during this alkylation reaction in THF is tentatively assigned to be the solvated neutral organozinc radical complex $[\text{ZnR}(t\text{-BuNCHCHN}-t\text{-Bu})(\text{THF})]^\bullet$. This conclusion is further corroborated by the observed stability of the 1:1 coordination complex **1a** toward reduction by **3a**. Nevertheless, we have shown that radical-anionic diorganozinc complexes of $t\text{-BuNCHCHN}-t\text{-Bu}$ can exist, although they have limited thermal stabilities. Their thermal decomposition gave the corresponding heteroleptic organo(diamido)zincates, alkylated or not alkylated at the carbon atom of the $t\text{-BuNCHCHN}-t\text{-Bu}$ ligand. The nonalkylated derivatives have been prepared independently *via* an alternative route. The synthetic potential of these new complexes in organozinc-mediated reactions is the subject of further study.

Experimental Section

General Data. All experiments were carried out under a dry and oxygen-free nitrogen atmosphere, using standard Schlenk techniques. Et_2O , THF, C_6H_6 , and pentane were dried and distilled from Na/benzophenone prior to use. CH_2Cl_2 was dried and distilled from CaH_2 . All standard chemicals were purchased from Aldrich and Janssen Chimica. $\text{ZnMe}_2(t\text{-BuNCHCHN}-t\text{-Bu})$ (**1a**),^{1c} $[\text{ZnR}(t\text{-BuNCHCHN}-t\text{-Bu})]_2$ (R = Me (**4a**), Et (**4b**)),³ ZnCl_2 ,¹⁶ 1,4-di-*tert*-butyl-1,4-diaza-1,3-butadiene ($t\text{-BuNCHCHN}-t\text{-Bu}$),¹⁷ and $\text{Zn}(\text{CH}_2\text{Ph})_2$ ¹⁸ were prepared according to literature procedures. ^1H and ^{13}C NMR spectra were recorded at 200 and 50 MHz in C_6D_6 , CD_2Cl_2 , or THF- d_6 at room temperature using SiMe_4 as external standard. Coupling (J) constants are in hertz (Hz). EPR spectra were recorded in Et_2O or THF at room temperature. Melting points are uncorrected. Elemental analyses were carried out by Dornis und Kolbe Mikroanalytisches Laboratorium, Mülheim a.d. Ruhr, Germany.

Reaction of $t\text{-BuNCHCHN}-t\text{-Bu}$ with $\text{Zn}(\text{CH}_2\text{CH}_2\text{CH}_2\text{-OMe})_2$. This reaction was carried out according to the

procedure described in ref 5. EPR spectra of this reaction mixture were recorded in Et_2O at room temperature.

Reduction of $\text{ZnMe}_2(t\text{-BuNCHCHN}-t\text{-Bu})$ (1a**).** **Method A:** With **4a**. To a solution of **1a**, prepared *in situ* from $t\text{-BuNCHCHN}-t\text{-Bu}$ (0.28 g; 1.66 mmol) and ZnMe_2 (1.70 mL of a 1.0 M solution in hexane, 1.70 mmol) in Et_2O or THF (30 mL) at room temperature, was added a solution of **4a** (0.42 g; 0.85 mmol) in Et_2O or THF (20 mL). ^1H NMR in C_6D_6 showed this powder to be a 1:1 mixture of **1a** and **4a**.

Method B: With Potassium Metal. To a stirred solution of **1a** in THF (30 mL), prepared *in situ* from $t\text{-BuNCHCHN}-t\text{-Bu}$ (1.25 g; 7.43 mmol) and ZnMe_2 (7.5 mL of a 1.0 M solution in pentane, 7.5 mmol), was added finely divided potassium (0.29 g; 7.42 mmol) in THF (20 mL) at room temperature. After the mixture was stirred for 16 h, the solvent was removed *in vacuo*, leaving an orange-brown residue. The residue was washed with hexanes or Et_2O (50 mL) and dried *in vacuo*, giving **9a** as a yellow-brown powder, yield 2.09 g (7.26 mmol; 98%). ^1H NMR (THF- d_6): δ 5.65 (s, 2, $\text{NCH}=\text{}$); 1.16 (s, 18, $\text{C}(\text{CH}_3)_3$); -0.82 (s, 3, CH_2). ^{13}C NMR (THF- d_6): δ 114.4 ($\text{NCH}=\text{}$); 52.4 ($\text{C}(\text{CH}_3)_3$); 33.9 ($\text{C}(\text{CH}_3)_3$); -11.3 (CH_2). Mp: >180 °C. Anal. Calcd for $\text{C}_{11}\text{H}_{23}\text{KN}_2\text{Zn}$: C, 45.91; H, 8.06; N, 9.73. Found: C, 46.08; H, 8.01; N, 9.64.

General Procedure for the Addition of $\text{K}(t\text{-BuNCHCHN}-t\text{-Bu})$ to ZnR_2 . To a stirred solution of $t\text{-BuNCHCHN}-t\text{-Bu}$ in THF (50 mL) was added 1 equiv of finely divided potassium at room temperature. After the mixture was stirred for 16 h, an equimolar amount of ZnR_2 was added to the brown suspension. The resulting clear solution was stirred for an additional 30 min. The solvent was removed *in vacuo*, leaving an orange-brown sticky residue. The residue was washed with Et_2O (50 mL) and dried *in vacuo*.

(a) To ZnMe_2 . The reaction of $t\text{-BuNCHCHN}-t\text{-Bu}$ (1.12 g; 6.7 mmol), potassium (0.26 g; 6.7 mmol), and ZnMe_2 (6.7 mL of a 1.0 M solution in pentane, 6.7 mmol) gave the product as a yellow-brown powder, isolated yield 1.78 g (6.19 mmol; 93%). ^1H and ^{13}C NMR (*vide supra*) showed this powder to be pure **9a**.

(b) To ZnEt_2 . The reaction of $t\text{-BuNCHCHN}-t\text{-Bu}$ (1.33 g; 7.93 mmol), potassium (0.31 g; 7.90 mmol), and ZnEt_2 (8.0 mL of a 1.0 M solution in hexane, 8.0 mmol) gave the product as a yellow-brown powder, isolated yield 1.84 g. ^1H and ^{13}C NMR showed this powder to be a mixture of **9b** (57%) and **10b** (43%). Identical reaction at 0.15 M $t\text{-BuNCHCHN}-t\text{-Bu}$ concentration and 195 K yielded a mixture of **9b** (73%) and **10b** (27%). Identical reaction at 0.03 M $t\text{-BuNCHCHN}-t\text{-Bu}$ concentration and 298 K gave a mixture of **9b** (77%) and **10b** (23%). We have been unable to separate the two products. **9b**: ^1H NMR (THF- d_6) δ 5.62 (s, 2, $\text{NCH}=\text{}$), 1.21 (t, 3, CH_3), 1.16 (s, 18, $\text{C}(\text{CH}_3)_3$), 0.10 (q br, 2, CH_2); ^{13}C NMR (THF- d_6) δ 114.3 ($\text{NCH}=\text{}$), 52.1 ($\text{C}(\text{CH}_3)_3$), 33.9 ($\text{C}(\text{CH}_3)_3$), 14.7 (CH_3), 3.6 (CH_2); mp >180 °C. **10b**: ^1H NMR (THF- d_6) δ 5.45 (s, 1, $\text{NCH}=\text{}$), 2.45 (q, 2, $^3J = 7.2$ Hz, $\text{NC}(\text{CH}_2)=$), 1.29 (s, 9, $\text{C}(\text{CH}_3)_3$), 1.14 (s, 9, $\text{C}(\text{CH}_3)_3$), 1.21 (t, 3, CH_3), 0.10 (q br, 2, CH_2); ^{13}C NMR (THF- d_6) δ 126.6 ($\text{NC}(\text{CH}_2)=$), 114.9 ($\text{NCH}=\text{}$), 53.0, 52.2 ($\text{C}(\text{CH}_3)_3$), 36.6, 33.9 ($\text{C}(\text{CH}_3)_3$), 26.7 ($\text{NC}(\text{CH}_2)=$), 15.8 (CH_3), 3.6 (CH_2).

(c) To $\text{Zn}(\text{CH}_2\text{Ph})_2$. The reaction from $t\text{-BuNCHCHN}-t\text{-Bu}$ (1.89 g; 11.23 mmol), potassium (0.44 g; 11.25 mmol), and $\text{Zn}(\text{CH}_2\text{Ph})_2$ (2.74 g; 11.06 mmol) gave the product as an orange powder, yield 4.35 g. ^1H and ^{13}C NMR showed this powder to be a mixture of **9c** (70%) and **10c** (30%). We have been unable to separate the two products. **9c**: ^1H NMR (THF- d_6) δ 6.9–6.8 (m, 4H, ArH), 8.52 (dd, 1H, ArH), 5.60 (s, 2, $\text{NCH}=\text{}$), 1.88 (s br, 2, CH_2), 1.15 (s, 18, $\text{C}(\text{CH}_3)_3$); ^{13}C NMR (THF- d_6) δ 153.9 (Ar C_{ipso}), 128.1, 127.0, 119.4 (ArC), 114.1 ($\text{NCH}=\text{}$), 52.2 ($\text{C}(\text{CH}_3)_3$), 33.9 ($\text{C}(\text{CH}_3)_3$), 23.2 (CH_2). **10c**: ^1H NMR (THF- d_6) δ 7.4–6.4 (aryl), 5.61 (s, 1, $\text{NCH}=\text{}$), 3.71 (s, 2, $\text{NC}(\text{CH}_2)=$), 1.89 (s br, 2, CH_2), 1.23 (s, 9, $\text{C}(\text{CH}_3)_3$), 1.09 (s, 9, $\text{C}(\text{CH}_3)_3$).

Preparation of $[\text{Zn}(\text{CH}_2\text{Ph})(t\text{-BuNCHCHN}-t\text{-Bu})]_2$ (4c**).** Complex **4c** was prepared according to a literature procedure,³ with a slightly modified workup, starting from $t\text{-BuNCHCHN}-$

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Table 4. Crystal Data and Details of the Structure Determination for {**9a**(THF)_n} and {**9c**(Et₂O)_{1/2}}_n

	{ 9a (THF) _n }	{ 9c (Et ₂ O) _{1/2} } _n
Crystal Data		
empirical formula	C ₁₅ H ₃₁ KN ₂ OZn	C ₁₉ H ₃₂ KN ₂ O _{1/2} Zn
mol wt	359.91	400.96
cryst syst	orthorhombic	orthorhombic
space group	<i>Pnma</i> (No. 62)	<i>P2₁2₁2</i> (No. 18)
<i>a</i> , Å	15.6658(7)	18.712(2)
<i>b</i> , Å	21.5705(11)	10.634(2)
<i>c</i> , Å	11.3578(9)	10.803(8)
<i>V</i> , Å ³	3838.0(4)	2149.6(17)
<i>Z</i>	8	4
<i>D</i> _{calcd} , g cm ⁻³	1.246	1.239
μ (Mo K α), cm ⁻¹	15.0	13.4
cryst size, mm	0.13 × 0.38 × 0.50	0.08 × 0.38 × 0.38
Data Collection		
temp K	150	150
radiation; λ , Å	Mo K α (graphite); 0.710 73	Mo K α (graphite); 0.710 73
θ _{min} , θ _{max} , deg	1.9, 27.5	1.9, 25
sca type	$\omega/2\theta$	$\omega/2\theta$
data set	-16 to +20, 0 to +27, -14 to 0	-21 to +24, 0 to +3, 0 to +14
total no. of data	6685	3897
total no. of unique data	4526	3421
Refinement		
no. of refined params	280	221
R1, wR2, <i>S</i> ^a	0.0495, 0.1151, 1.02	0.0976, 0.248, 1.14

$$^a R1 = \sum |F_o| - |F_c| / \sum |F_o|; wR2 = [\sum [w(F_o^2 - F_c^2)^2] / \sum [w(F_o^2)^2]]^{1/2}.$$

t-Bu (1.42 g; 8.44 mmol), potassium (0.33 g; 8.44 mmol), and Zn(CH₂Ph)Cl, prepared *in situ* from Zn(CH₂Ph)₂ (1.05 g; 4.24 mmol) and ZnCl₂ (7.0 mL of a 0.62 M solution in Et₂O, 4.34 mmol). Extraction of the product mixture with CH₂Cl₂ (2 × 50 mL) gave **4c** as a yellow powder in 2.15 g (3.31 mmol; 78%) yield. ¹H NMR (C₆D₆): δ 7.45 (d, 3H, N=CH and Ar *H*), 7.39 (dd, 2 H, Ar *H*), 6.96 (dd, 1 H, Ar *H*), 2.58 (d, 1 H, NCH), 2.45 (dd, 2 × 1 H, ²*J* = 10.8, Ar CH₂), 0.94 (s, 9 H, C(CH₃)₃), 0.90 (s, 9 H, C(CH₃)₃). ¹H NMR (CD₂Cl₂): δ 8.15 (d, 1 H, N=CH), 7.2–6.7 (m, 5 H, Ar *H*), 3.14 (d, 1H, NCH), 2.00 (dd, 2 × 1 H, ²*J* = 14.9, Ar CH₂), 1.16 (s, 9 H, C(CH₃)₃), 0.84 (s, 9 H, C(CH₃)₃). ¹³C NMR (CD₂Cl₂): δ 171.3 (N=CH); 152.6 (quaternary Ar); 128.0, 126.8, 119.9 (Ar); 59.7 (NCH); 58.3, 51.9 (C(CH₃)₃); 32.8, 29.6 (C(CH₃)₃); 22.1 (ZnCH₂).

Preparation of K[ZnMe(*t*-BuNCHCHN-*t*-Bu)] (9a**).** To a solution of **4a** (2.38 g; 4.8 mmol) in THF (50 mL) was added finely divided potassium (0.38 g; 9.7 mmol). After the mixture was stirred for 16 h at room temperature, the solvent was removed *in vacuo*, leaving a yellow-brown residue. The residue was washed with Et₂O (2 × 20 mL) and dried *in vacuo*, giving **9a** as a yellow powder, isolated yield 2.73 g (9.5 mmol; 99%). Crystals of {**9a**(THF)_n} were obtained by crystallization from a 1:5 mixture of THF and Et₂O at -30 °C.

Preparation of K[ZnEt(*t*-BuNCHCHN-*t*-Bu)] (9b**).** Complex **9b** was synthesized according to the procedure for **9a**, starting from **4b** (2.72 g; 10.35 mmol) and potassium (0.42 g; 10.74 mmol); isolated yield 2.90 g (9.61 mmol; 93%).

Preparation of K[Zn(CH₂Ph)(*t*-BuNCHCHN-*t*-Bu)] (9c**).** Complex **9c** was synthesized according to the procedure for **9a**, starting from **4c** (2.02 g; 6.22 mmol) and potassium (0.24 g; 6.14 mmol); isolated yield 1.91 g (5.88 mmol; 96%). Crystals of {**9c**(Et₂O)_{1/2}}_n suitable for an X-ray diffraction study, were obtained by crystallization from a 1:5 mixture of THF and Et₂O and were not dried.

X-ray Structure Determination for {9a**(THF)_n} and {**9c**(Et₂O)_{1/2}}_n.** X-ray data were collected on an Enraf-Nonius-CAD4T diffractometer on a rotating anode for transparent, cut to size, inert-oil-covered crystals that were glued on top of a glass fiber and stabilized by the cold dinitrogen stream at 150

K. Pertinent numerical data have been collected in Table 4. The best available crystals of {**9c**(Et₂O)_{1/2}}_n were of poor quality, as indicated by broad and structured reflection profiles and signs of intergrowth. Unit cell parameters were derived from the SET4¹⁹ setting angles. The data were corrected for Lp and absorption (empirical PLATON²⁰) and averaged. The structure was solved with automated Patterson techniques (DIRDIF92²¹) and refined on *F*² by full-matrix least squares (SHELXL93²²). Non-hydrogen atoms were refined with anisotropic displacement parameters. Hydrogen atoms were included at calculated positions, riding on and with isotropic displacement parameters derived from their carrier atoms. One of the [ZnC₁₁H₂₃N₂]⁻ anions of {**9a**(THF)_n} is disordered over an inversion center. The weak data quality of {**9c**(Et₂O)_{1/2}}_n is reflected in the relatively high final *R* value and some unusual displacement ellipsoids. Both structures were checked for higher symmetry and residual voids (PLATON²⁰). The Flack parameter for {**9c**(Et₂O)_{1/2}}_n was refined to -0.04(6).

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Supporting Information Available: Further details on the structure determinations, including tables of atomic coordinates, all bond lengths and angles, and thermal parameters, for {**9a**(THF)_n} and {**9c**(Et₂O)_{1/2}}_n (24 pages). Ordering information is given on any current masthead page.

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