

29. Radical Anions of Cyclic Azoalkanes: An ESR, ENDOR, and TRIPLE-Resonance Study¹⁾

by Caspar H. Ess²⁾ and Fabian Gerson*

Institut für Physikalische Chemie der Universität Basel, Klingelbergstrasse 80, CH-4056 Basel

and Waldemar Adam

Institut für Organische Chemie der Universität Würzburg, Am Hubland, D-8700 Würzburg

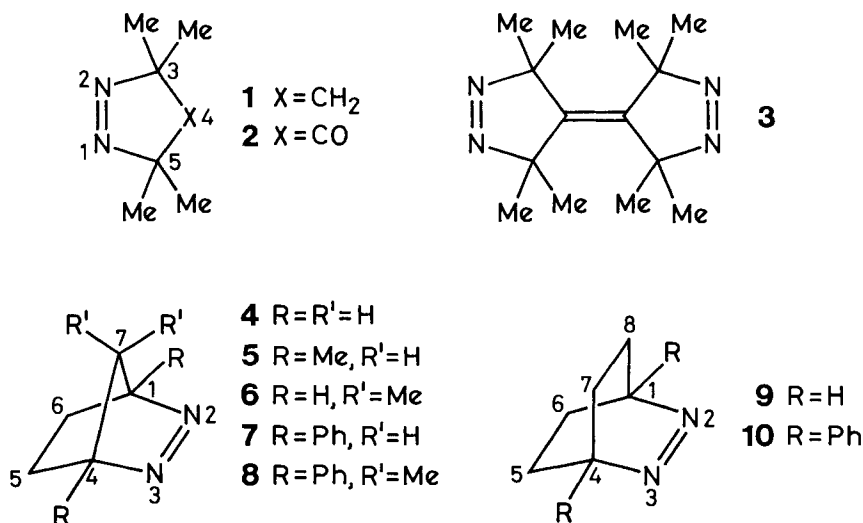
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Radical anions of ten monocyclic and bicyclic azoalkanes containing the azo group in (*Z*)-conformation, have been fully characterized by their hyperfine data with the use of ESR, ENDOR, and general-TRIPLE-resonance spectroscopy. These azoalkanes are represented by 3,3,5,5-tetramethyl-1-pyrazoline (**1**), 2,3-diazabicyclo[2.2.1]hept-2-ene (**4**), and 2,3-diazabicyclo[2.2.2]oct-2-ene (**9**), as well as by their derivatives **2**, **3**, **5–8**, and **10**. For all radical anions **1^{•-}–10^{•-}**, the ¹⁴N-coupling constant, a_N , is in the range of +0.83 to +0.97 mT; this finding indicates that the spin population is essentially restricted to the π system of the azo group. The ¹⁴N-hyperfine anisotropy largely affects the width of ESR lines, particularly at low temperatures. Substantial coupling constants of ⁷Li-, ²³Na-, ³⁹K-, and ¹³³Cs-nuclei point to a close association of the radical anions with their alkali-metal counterions. With the exception of ³⁹K, these nuclei give rise to readily observable ENDOR signals which appear along with those stemming from protons. The prominent hyperfine features of **1^{•-}–10^{•-}** are discussed.

Introduction. – Azoalkanes, in which the azo group is linked to saturated hydrocarbon moieties, have been known since 1909 [1]. Contrary to aromatic azo compounds like azobenzene, they readily lose N₂ upon a thermal or photolytical treatment [2], and this chemical property has been exploited to generate radicals, biradicals [2–4], and novel hydrocarbons [5]. Radical anions of several (*E*)-dialkyldiimines have been studied by ESR spectroscopy [6–8]. Persistent radical anions are, in general, produced from azoalkanes which, like di(*tert*-butyl)diimine, have no H-atoms in the positions neighbouring the azo group, or in which removal of these atoms is impeded by the *Bredt's* rule. Here, we report on such persistent radical anions generated from monocyclic and bicyclic azoalkanes containing the azo group in the (*Z*)-conformation. These compounds are represented by 3,3,5,5-tetramethyl-1-pyrazoline (**1**), 3,3,5,5-tetramethyl-1-pyrazol-4-one (**2**), 3,3,5,5,3',3',5',5'-octamethyl-di(1-pyrazolinyliidene) (**3**), 2,3-diazabicyclo[2.2.1]hept-2-ene (**4**) and its 1,4-dimethyl (**5**), 7,7-dimethyl (**6**), 1,4-diphenyl (**7**), and 1,4-diphenyl-7,7-dimethyl (**8**) derivatives and 2,3-diazabicyclo[2.2.2]oct-2-ene (**9**) and its 1,4-diphenyl (**10**) derivative. The radical anions **1^{•-}–10^{•-}** have been fully characterized by their hyperfine data with the use of ESR, ENDOR, and general-TRIPLE-resonance spectroscopy.

¹⁾ Due to a technical error in the final stage of the printing process, the symbol for radical anion (^{•-}) was replaced by " in the original contribution, *Helv. Chim. Acta* **1991**, *74*, 2078. As this error may cause confusion, the paper is reprinted.

²⁾ Present address: Department of Chemistry, Stanford University, California 94305, U.S.A.



Results and Discussion. – *Generation and Stability of the Radical Anions.* Due to the presence of the electron-attracting azo group, the compounds **1–10** were easily reduced to their radical anions. Cyclic voltammograms of **1**, **4**, and **9** (solvent: CH₃CN; working electrode: Hg; counter-electrode: Ag/AgCl; supporting salt: Bu₄NClO₄; scan: 200mV/s; temperature: 298 K) exhibited reversible waves at half-wave potentials, $E_{1/2}$ (vs. SCE), of -0.82 ± 0.01 , -0.87 ± 0.01 , and -0.82 ± 0.01 V, respectively. For spectroscopic studies, the radical anions **1⁻–10⁻** were prepared from the corresponding neutral compounds by reaction with an alkali-metal mirror in 1,2-dimethoxyethane (DME). Direct use of the metallic K, Na, or Cs led to the radical anions associated with K⁺, Na⁺, or Cs⁺, while adding an excess of LiCl salt to the solution of a radical anion, produced by reduction with K, caused replacement of K⁺ by Li⁺ as the counterion.

As indicated by their moderately negative potentials $E_{1/2}$ and by reversibility of their reduction waves in the cyclic voltammograms, the radical anions **1⁻–10⁻** were thermodynamically and kinetically stable. The kinetic stability (persistence) of **1⁻–10⁻** is not unexpected, considering their molecular structures which meet the requirements stated in the *Introduction*. In the diazabicycloalkene series, the persistence of the radical anions was lowered by dimethyl or diphenyl substitution at the bridgehead atoms C(1) and C(4), presumably because such a substitution promotes exclusion of N₂ by stabilizing thus formed radicals or biradicals [9]. Therefore, **5⁻** and **7⁻** were less persistent than **4⁻**, and the persistence of **10⁻** was lower than that of **9⁻**. By contrast, dimethyl substitution at the methano-bridge atom C(7) of diazabicycloheptene seemed to enhance the persistence of the radical anions, **8⁻** being as stable as **4⁻** despite the presence of Ph groups at C(1) and C(4).

ESR Spectra. The radical anions **1⁻–10⁻** were studied by ESR spectroscopy from 203 K up to the temperature at which their decay set in. The resolution of the spectra was strongly affected by ¹⁴N-hyperfine anisotropy, especially at low temperatures. In

consequence, the line-widths increased on going from the centres of the spectra to the peripheries and, to a lesser extent, on passing from the low- to the high-field halves (see *Appendix*). Temperature-dependent hyperfine splittings from alkali-metal nuclei of the counterions were observed for all radical anions. The pertinent nuclei are ${}^7\text{Li}$, ${}^{23}\text{Na}$, ${}^{39}\text{K}$, or ${}^{133}\text{Cs}$ for 1^- – 3^- associated with Li^+ , Na^+ , K^+ , or Cs^+ , and they are ${}^7\text{Li}$, ${}^{39}\text{K}$, or ${}^{133}\text{Cs}$ for 4^- – 10^- with Li^+ , K^+ , or Cs^+ as the counterions. In the latter case, ${}^{23}\text{Na}$ -hyperfine splittings could not be determined. Brief contact of the solutions of **4**–**10** with Na mirror in DME gave rise to low-intensity spectra identical to those observed with the counterion K^+ (see *Experimental*), while upon prolonged reduction these spectra were superseded by an unresolved S-shaped derivative curve. Interestingly, the ${}^7\text{Li}$ -hyperfine splittings indicated association of 1^- – 10^- with two Li^+ counterions, presumably due to the addition of LiCl salt in a concentration largely exceeding that of the radical anion.

It is noteworthy that the ESR spectra of 3^- were very similar to those of 1^- and 2^- under the applied experimental conditions (solvent: DME; counterion: Li^+ , Na^+ , K^+ , or Cs^+ ; temperature: 203–293 K). Thus, under these conditions, the unpaired electron in 3^- appears to be localized on one 1-pyrazoline moiety, *i.e.*, electron-spin transfer between the two moieties is slow on the hyperfine time-scale ($\sim 10^7 \text{ s}^{-1}$).

The ESR spectra, taken at 203 K are exemplified in *Figs. 1* and *2* by those of 1^- associated with Li^+ , Na^+ , K^+ , or Cs^+ , and of 4^- with Li^+ , K^+ , or Cs^+ as the counterions. Their temperature dependence is illustrated in *Fig. 3* by the spectra of $1^-/\text{K}^+$ at 203 and 273 K. Analyses of all spectra were supported by computer simulations, as demonstrated in *Fig. 4* for $1^-/\text{Na}^+$ at 273 K, and by the use of the ENDOR technique.

ENDOR Spectra. All radical anions 1^- – 10^- gave rise, at 203 K, to ${}^1\text{H}$ -ENDOR spectra, except for 4^- – 10^- associated with Na^+ ; in this case, the unresolved S-shaped ESR absorption did not respond to the ENDOR experiment. The signals appeared pairwise at $\nu_{\text{H}} \pm \frac{1}{2}|a_{\text{H}}|$ where $\nu_{\text{H}} = 14.56 \text{ MHz}$ is the frequency of the free proton, and a_{H} represents the ${}^1\text{H}$ -coupling constant in MHz [10a]. In addition, ENDOR signals from ${}^7\text{Li}$ -, ${}^{23}\text{Na}$ -, or ${}^{133}\text{Cs}$ -nuclei were detected for 1^- – 3^- associated with Li^+ , Na^+ , or Cs^+ , and for 4^- – 10^- with Li^+ or Cs^+ as the counterions. Whereas in the case of ${}^7\text{Li}$ and ${}^{133}\text{Cs}$ high- as well as low-frequency signals could be observed at $\nu_{\text{Li}} \pm \frac{1}{2}|a_{\text{Li}}|$ or $\frac{1}{2}|a_{\text{Cs}}| \pm \nu_{\text{Cs}}$, only the high-frequency signal at $\nu_{\text{Na}} + \frac{1}{2}|a_{\text{Na}}|$ was detected for ${}^{23}\text{Na}$, because of the low sensitivity of our ENDOR system in the range of 0–3 MHz pertinent to $\nu_{\text{Na}} - \frac{1}{2}|a_{\text{Na}}|$. For the same reason, both ${}^{39}\text{K}$ -ENDOR signals escaped detection ($|a_{\text{K}}| \approx 1.4 \text{ MHz}$ or 0.05 mT in the ESR spectra; $\frac{1}{2}|a_{\text{K}}| - \nu_{\text{K}} \approx 0$; $\frac{1}{2}|a_{\text{K}}| + \nu_{\text{K}} \approx 1.4 \text{ MHz}$). Here, $\nu_{\text{Li}} = 5.66$, $\nu_{\text{Na}} = 3.85$, $\nu_{\text{K}} = 0.68$, and $\nu_{\text{Cs}} = 1.91 \text{ MHz}$ are the free frequencies of the ${}^7\text{Li}$ -, ${}^{23}\text{Na}$ -, ${}^{39}\text{K}$ -, and ${}^{133}\text{Cs}$ -nuclei, while a_{Li} , a_{Na} , a_{K} , and a_{Cs} represent the corresponding alkali-metal-coupling constants in MHz [10b]. The ${}^{14}\text{N}$ -ENDOR signals failed to appear in the range of 10.5–15 MHz, expected for $\frac{1}{2}|a_{\text{N}}| \pm \nu_{\text{N}}$, considering the frequency $\nu_{\text{N}} = 1.05 \text{ MHz}$ of the free nucleus and the coupling constant $|a_{\text{N}}| = 23$ – 28 MHz (which corresponds to $|a_{\text{N}}| = 0.83$ to 0.97 mT in the ESR spectra). In this case, the ${}^{14}\text{N}$ -hyperfine enhancement factor [10b] was insufficient to allow observation by the ENDOR technique. *Figs. 5* and *6* reproduce the ${}^1\text{H}$ - and alkali-metal-ENDOR spectra of 1^- associated with Li^+ , Na^+ , or Cs^+ and of 4^- with Li^+ or Cs^+ as the counterions. These spectra were taken under the same conditions as their ESR counterparts in *Figs. 1* and *2*.

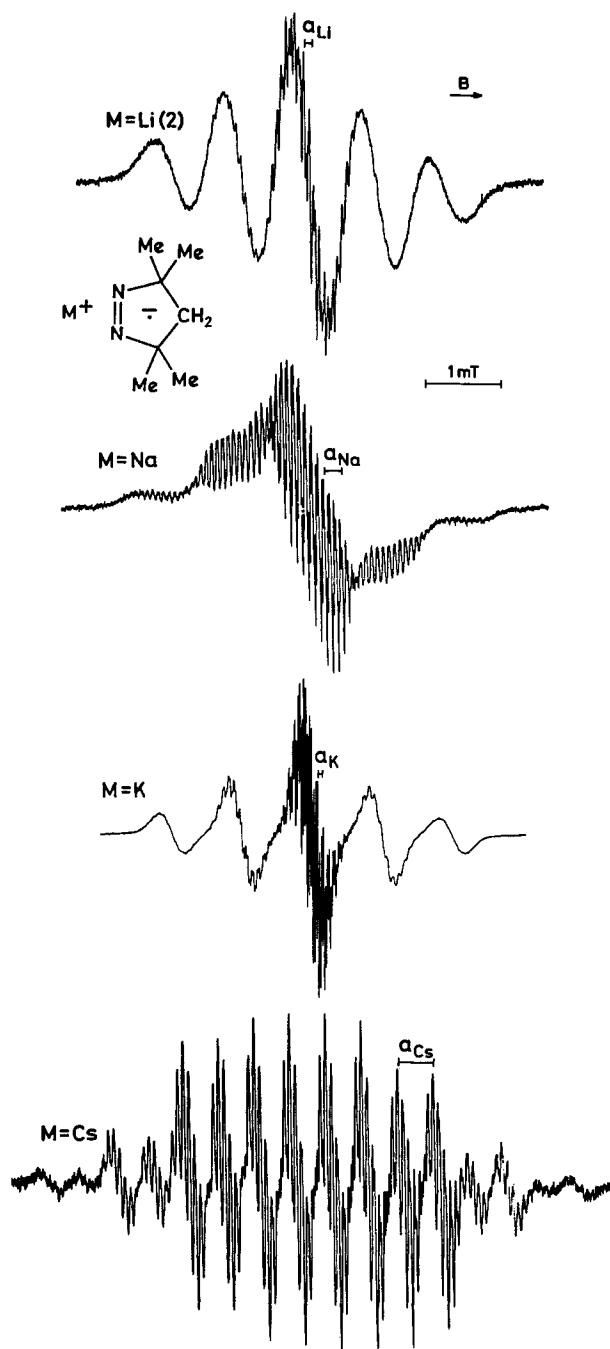


Fig. 1. ESR spectra of $1^{\bullet-}$. Solvent: DME; counterion as indicated; temp.: 203 K. $M = \text{Li}(2)$ signifies that two Li^+ counterions are associated with each radical anion.

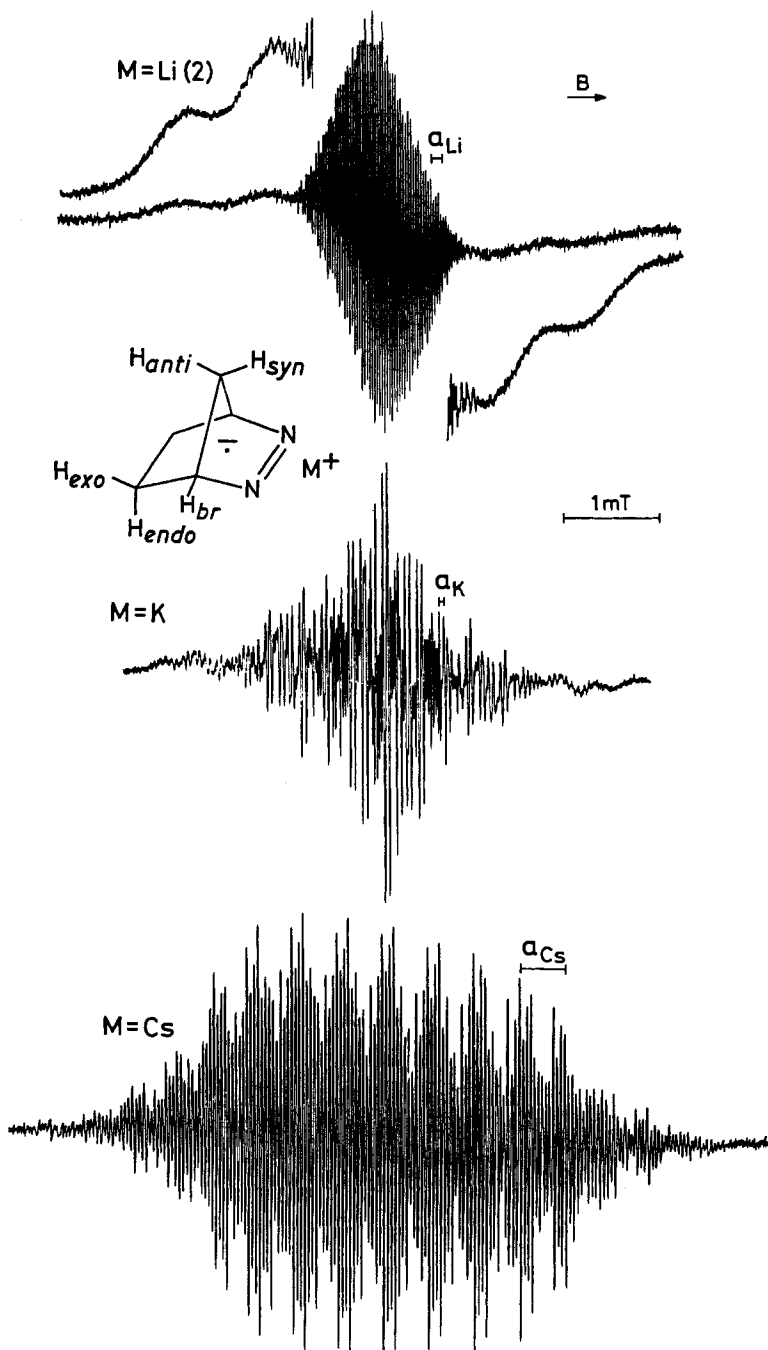


Fig. 2. ESR spectra of $4^{\cdot-}$. Solvent: DME; counterion as indicated; temp.: 203 K. The insets in the uppermost spectrum were recorded using higher modulation. $M = \text{Li}(2)$ signifies that two Li^+ counterions are associated with each radical anion.

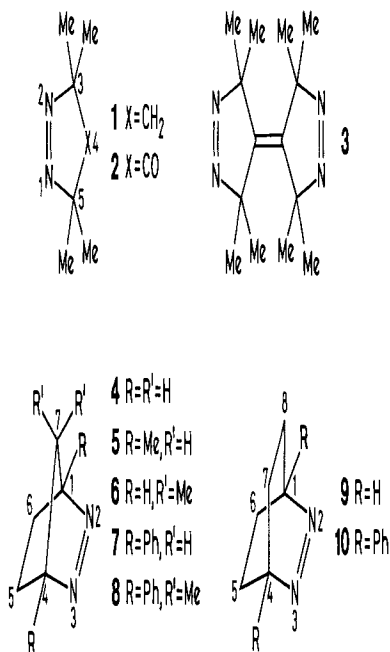


Fig. 3. ESR spectra of $1^{\cdot-}$. Solvent: DME; counterion and temp. as indicated.

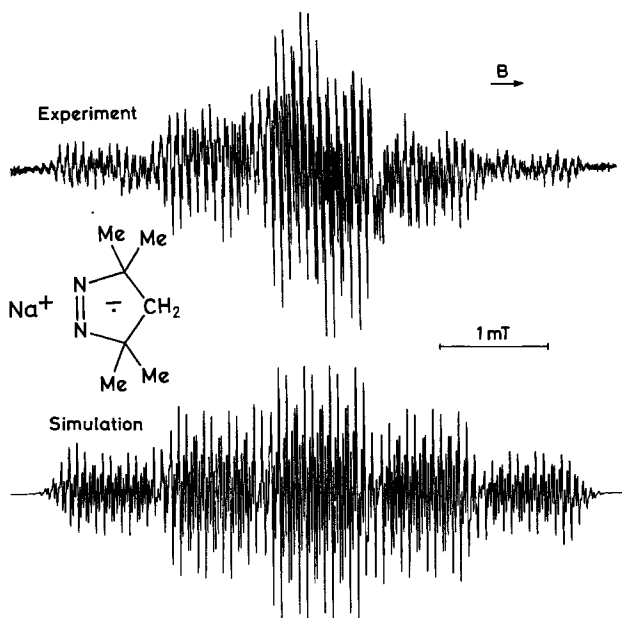


Fig. 4. ESR spectrum of $1^{\cdot-}$ and its simulation. Solvent: DME; counterion as indicated; temp.: 273 K. The simulation makes use of the coupling constants given in Table I; line-shape: Lorentzian; line-width (constant): 0.024 mT, without accounting for the effects of ^{14}N -hyperfine anisotropy (see Appendix).

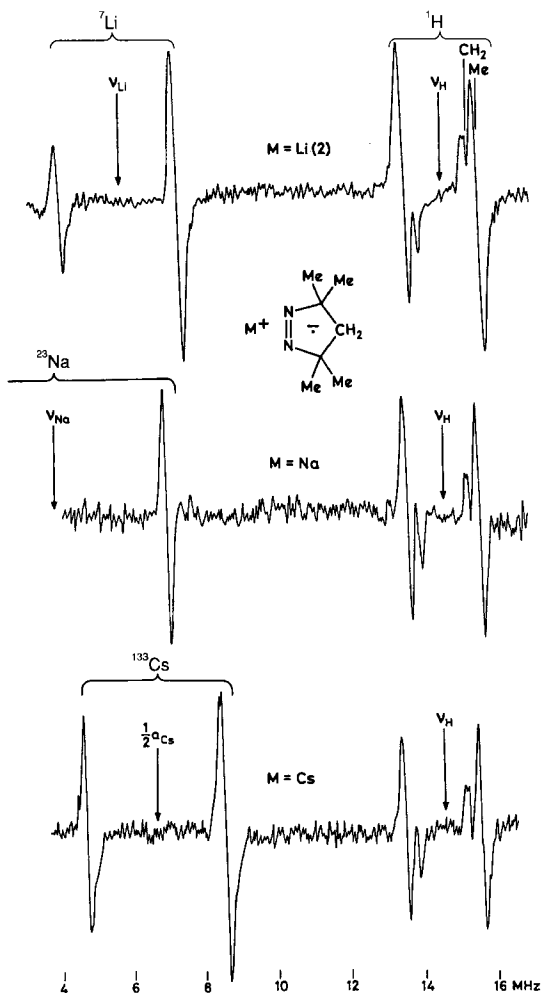


Fig. 5. ^1H -, ^7Li -, ^{23}Na -, and ^{133}Cs -ENDOR spectra of 1^- . Experimental conditions as for Fig. 1.

Signs and Assignments of the Coupling Constants. The hyperfine data for all radical anions 1^- – 10^- and their counterions are listed in Tables 1, 2, and 3 (for 9^- , see also [11]), together with the corresponding g factors. The signs allotted to the coupling constants have been derived by a general-TRIPLE-resonance experiment [10a] combined with theoretical arguments. The TRIPLE-resonance technique yielded relative signs of the coupling constants for all nuclei which gave rise to pairs of ENDOR signals, *i.e.*, for ^1H , ^7Li , and ^{133}Cs . In the case of 4^- – 10^- (Tables 2 and 3), the absolute signs were obtained on the reasonable assumption that the largest values $|a_{\text{H}}|$ due to the protons in the *exo*-positions of the ethano bridges (see below) are positive. As the signs of a_{Li} and a_{Cs} for the counterions Li^+ or Cs^+ associated with 4^- – 10^- are opposite and equal, respectively, to that of $a_{\text{H}_{\text{exo}}}$, a negative sign is indicated for a_{Li} and a positive one for a_{Cs} . These signs must also be shared by ^7Li - and

^{133}Cs -coupling constants for 1^- – 3^- with Li^+ or Cs^+ as the counterions, because *i*) here, too, general-TRIPLE-resonance experiments required opposite signs of a_{Li} and a_{Cs} and *ii*) the values of $|a_{\text{Li}}|$ as well as those of $|a_{\text{Cs}}|$ were very similar in the two series, thus pointing to a closely related mode of association of the radical anions with the same counterions. In turn, using a_{Li} and a_{Cs} as reference, absolute signs could be allotted to the coupling constants $a_{\text{H}}(\text{CH}_2)$ for 1^- and $a_{\text{H}}(\text{Me})$ for 1^- – 3^- with the use of general-TRIPLE-resonance spectroscopy (Table 1).

The signs of a_{Na} (absence of the low-frequency ^{23}Na -signals) and a_{K} (failure to detect both ^{39}K -signals) could not be determined in this way. However, the temperature dependence of a_{Na} and a_{K} indicated that the two coupling constants have the same positive sign as a_{Cs} which is opposite to that of a_{Li} (see below). The sign of the large ^{14}N -coupling constant, a_{N} , is also certainly positive, as it is in other radical ions in which the N-atoms bear a substantial part of the spin population. This sign is consistent with the ESR lines being broader in the high- than in the low-field half of the spectrum [12a][13].

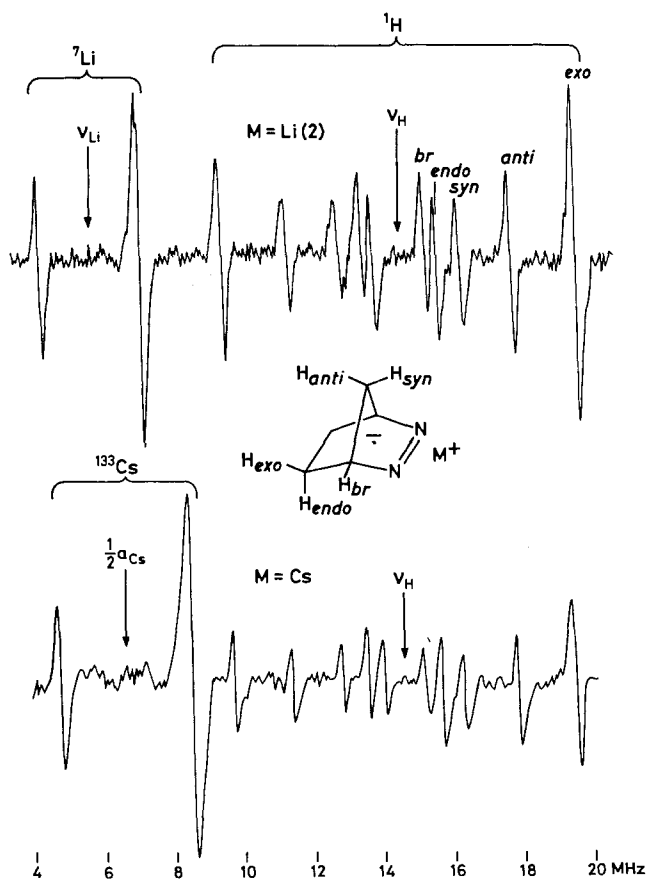


Fig. 6. ^1H -, ^7Li -, and ^{133}Cs -ENDOR spectra of 4^- . Experimental conditions as for Fig. 2.

Assignments of the coupling constants to sets of equivalent nuclei other than protons are unproblematic, due to their specific quantum numbers I , strongly differing values, and characteristic frequencies of the free nuclei in the ENDOR spectra. As for the coupling constants a_{H} , assignments are also in most cases straightforward, owing to different multiplicities of the sets of equivalent protons and to effects of dimethyl or diphenyl substitution. The only protons which cannot be distinguished by experimental evidence are those in the *syn*- and *anti*-positions at C(7) for 4^- , 5^- , and 7^- (or in the corresponding Me substituents for 6^- and 8^-) and in the *exo*- and *endo*-positions at C(5) and C(6) for 4^- – 8^- or at C(5), C(6), C(7), and C(8) for 9^- and 10^- . In these cases, assignments of coupling constants to protons in the individual positions have been guided by analogy with structurally related radical ions (see, e.g., [14][15]) for which always $|a_{\text{Hanti}}| > |a_{\text{Hsyn}}|$ and $|a_{\text{Hexo}}| > |a_{\text{Hendo}}|$. These assignments are supported by theory, as shown in Table 4 by comparison of the ^1H - and ^{14}N -coupling constants, observed for 4^- and 9^- with the corresponding values calculated by the INDO procedure [16] and based on an MNDO-optimized geometry [17]. There is a fair agreement between experiment and theory, except for the sign of a_{Hendo} .

Table 1. Coupling Constants [mT] and g Factors for 1^- – 3^-). Solvent: DME.

1^-									
Counterion, M^+	$\text{Li}^{+\text{b}}$		Na^+		K^+		Cs^+		
	203	293	203	273	203	273	203	293	
Temp.[K]	203	293	203	273	203	273	203	293	
a_{H} (CH_2 , 2 H)	+0.049	^{c)}	+0.051	+0.050	+0.048	+0.048	+0.052	+0.049	
a_{H} (Me, 12 H)	-0.071	^{c)}	-0.072	-0.071	-0.074	-0.072	-0.074	-0.071	
a_{N} (2 N)	+0.888	+0.890	+0.931	+0.933	+0.918	+0.923	+0.930	+0.932	
a_{M}	-0.116	^{c)}	+0.224	+0.236	+0.048	+0.058	+0.472	+0.554	
g factor	2.0037		2.0037		2.0037		2.0041		
2^-									
Counterion, M^+	$\text{Li}^{+\text{b}}$		Na^+		K^+		Cs^+		
	203	273	203	253	203	253	203	293	
Temp.[K]	203	273	203	253	203	253	203	293	
a_{H} (Me, 12 H)	-0.061	-0.060	-0.063	-0.059	-0.062	-0.060	-0.061	-0.061	
a_{N} (2 N)	+0.892	+0.902	+0.938	+0.945	+0.927	+0.938	+0.918	+0.940	
a_{M}	-0.117	-0.097	+0.206	+0.218	+0.047	+0.052	+0.420	+0.470	
g factor	2.0038		2.0039		2.0039		2.0042		
3^-									
Counterion, M^+	$\text{Li}^{+\text{b}}$		Na^+		K^+		Cs^+		
	203	293	203	293	203	273	203	293	
Temp.[K]	203	293	203	293	203	273	203	293	
a_{H} (Me, 12 H)	-0.089	-0.085	-0.094	-0.089	-0.091	-0.088	-0.092	-0.085	
a_{N} (2 N)	+0.872	+0.874	+0.908	+0.910	+0.896	+0.898	+0.888	+0.902	
a_{M}	-0.119	-0.099	+0.208	+0.216	+0.048	+0.053	+0.448	+0.527	
g factor	2.0036		2.0036		2.0037		2.0042		

^{a)} Experimental error: ± 0.001 mT in a_{H} and a_{M} , ± 0.005 mT in a_{N} at 203 K, ± 0.005 mT in a_{H} , a_{M} , and a_{N} at higher temperatures, ± 0.0001 in g .

^{b)} Two Li^+ counterions per radical anion.

^{c)} Not determined.

Table 2. *Coupling Constants [mT] and g Factors for 4⁻–8⁻*. Solvent: DME.

4 ⁻							
Counterion, M ⁺	Li ⁺ b)		K ⁺		Cs ⁺		
Temp. [K]	203	293	203	293	203	293	
a _{Hexo} (2 H)	+0.362	+0.358	+0.355	+0.340	+0.348	+0.337	
a _{Hendo} (2 H)	-0.076	-0.075	-0.076	-0.073	-0.075	-0.074	
a _{Hanti} (1 H)	-0.228	-0.225	-0.232	-0.223	-0.229	-0.221	
a _{Hsyn} (1 H)	-0.122	-0.113	-0.127	-0.122	-0.123	-0.116	
a _{Hbr} (2 H)	+0.054	+0.053	+0.046	+0.044	+0.041	+0.039	
a _N (2 N)	+0.848	+0.846	+0.871	+0.855	+0.856	+0.834	
a _M	-0.104	-0.053	+0.053	+0.055	+0.469	+0.493	
g factor	2.0038		2.0039		2.0042		
5 ⁻							
Counterion, M ⁺	Li ⁺ b)		K ⁺		Cs ⁺		
Temp. [K]	203	c)	203	c)	203	c)	
a _{Hexo} (2 H)	+0.432		+0.422		+0.418		
a _{Hendo} (2 H)	-0.057		-0.057		-0.058		
a _{Hanti} (1 H)	-0.212		-0.215		0.213		
a _{Hsyn} (1 H)	-0.114		-0.115		-0.116		
a _{Hbr} (Me, 6 H)	-0.028		-0.027		-0.026		
a _N (2 N)	+0.843		+0.852		+0.868		
a _M	-0.107		+0.050		+0.450		
g factor	2.0038		2.0039		2.0041		
6 ⁻							
Counterion, M ⁺	Li ⁺ b)		K ⁺		Cs ⁺		
Temp. [K]	203	293	203	293	203	293	
a _{Hexo} (2 H)	+0.421	+0.415	+0.409	+0.406	+0.406	+0.400	
a _{Hendo} (2 H)	-0.067	-0.067	-0.064	-0.065	-0.059	-0.059	
a _{Hanti} (Me, 3 H)	+0.133	+0.136	+0.136	+0.139	+0.139	+0.139	
a _{Hsyn} (Me, 3 H)	+0.013	+0.013	+0.010	+0.012	+0.012	+0.010	
a _{Hbr} (2 H)	<0.010	<0.010	<0.010	<0.010	<0.010	<0.010	
a _N (2 N)	+0.880	+0.872	+0.852	+0.848	+0.864	+0.853	
a _M	-0.102	-0.055	+0.050	+0.058	+0.482	+0.546	
g factor	2.0038		2.0039		2.0042		
7 ⁻							
Counterion, M ⁺	Li ⁺ b)		K ⁺		Cs ⁺		
Temp. [K]	203	c)	203	c)	203	c)	
a _{Hexo} (2 H)	+0.435		+0.434		+0.432		
a _{Hendo} (2 H)	-0.065		-0.062		-0.062		
a _{Hanti} (1 H)	-0.211		-0.216		-0.213		
a _{Hsyn} (1 H)	-0.115		-0.118		-0.117		
a _{Hbr} (Ph, 10 H)	<0.005		<0.005		<0.005		
a _N (2 N)	+0.848		+0.848		+0.867		
a _M	-0.103		+0.046		+0.433		
g factor	2.0038		2.0038		2.0042		

Table 2 (cont.)

8⁻						
Counterion, M ⁺	Li ⁺ b)		K ⁺		Cs ⁺	
Temp. [K]	203	293	203	293	203	293
a _{H_{exo}} (2 H)	+0.514	+0.505	+0.484	+0.482	+0.485	+0.484
a _{H_{endo}} (2 H)	-0.043	-0.043	-0.044	-0.042	-0.044	-0.045
a _{H_{anti}} (Me, 3 H)	+0.111	+0.113	+0.127	+0.121	+0.126	+0.131
a _{H_{syn}} (Me, 3 H)	<0.010	<0.010	<0.010	<0.010	<0.010	<0.010
a _{H_{br}} (Ph, 10 H)	<0.010	<0.010	<0.010	<0.010	<0.010	<0.010
a _N (2 N)	+0.858	+0.840	+0.888	+0.844	+0.972	+0.898
a _M	-0.099	-0.038	+0.049	+0.052	+0.488	+0.529
g factor	2.0038		2.0038		2.0042	

a) Experimental error: ±0.001 mT in a_H and a_M, ±0.005 mT in a_N at 203 K, ±0.005 mT in a_H, a_M, and a_N at higher temperatures, ±0.0001 in g.

b) Two Li⁺ counterions per radical anion.

c) Unstable at higher temperatures.

 Table 3. Coupling Constants [mT] and g Factors for **9⁻** and **10⁻** a). Solvent: DME.

9⁻						
Counterion, M ⁺	Li ⁺ b)		K ⁺		Cs ⁺	
Temp. [K]	203	293	203 ^{c)}	293	203	293
a _{H_{exo}} (4 H)	+0.276	+0.272	+0.274	+0.268	+0.274	+0.272
a _{H_{endo}} (4 H)	-0.073	-0.077	-0.073	-0.071	-0.073	-0.067
a _{H_{br}} (2 H)	-0.019	-0.020	-0.020	-0.020	-0.020	-0.018
a _N (2 N)	+0.858	+0.852	+0.878	+0.876	+0.880	+0.883
a _M	-0.112	<0.015	+0.053	+0.061	+0.481	+0.547
g factor	2.0039		2.0040		2.0042	

10⁻						
Counterion, M ⁺	Li ⁺ b)		K ⁺		Cs ⁺	
Temp. [K]	203	293	203	^{d)}	203	^{d)}
a _{H_{exo}} (4 H)	+0.289	+0.287	+0.301		+0.304	
a _{H_{endo}} (4 H)	-0.066	-0.065	-0.065		-0.062	
a _{H_{br}} (Ph, 10 H)	<0.010	<0.015	<0.010		<0.010	
a _N (2 N)	+0.862	+0.858	+0.867		+0.870	
a _M	-0.066	<0.015	+0.048		+0.435	
g factor	2.0039		2.0039		2.0041	

a) Experimental error: ±0.001 mT in a_H and a_M, ±0.005 mT in a_N at 203 K, ±0.005 mT in a_H, a_M, and a_N at higher temperatures, ±0.0001 in g.

b) Two Li⁺ counterions per radical anion.

c) The values given in this column were quoted in [11].

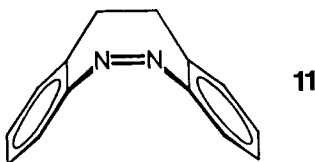
d) Unstable at higher temperatures.

Table 4. Comparison of Observed and Calculated Coupling Constants [mT] for 4^{\ominus} and 9^{\ominus} .

	4^{\ominus}		9^{\ominus}	
	Calc. ^{a)}	Obs. ^{b)}	Calc. ^{a)}	Obs. ^{b)}
$a_{\text{H}_{\text{exo}}}$	+0.278	+0.355	+0.223	+0.274
$a_{\text{H}_{\text{endo}}}$	+0.066	-0.076	+0.037	-0.073
$a_{\text{H}_{\text{anti}}}$	-0.216	-0.232	–	–
$a_{\text{H}_{\text{syn}}}$	-0.082	-0.127	–	–
$a_{\text{H}_{\text{ibr}}}$	+0.019	+0.046	-0.047	-0.020
a_{N}	+0.621	+0.871	+0.671	+0.878

^{a)} See text for the procedure.
^{b)} Solvent: DME; counterion: K⁺; temp.: 203 K.

¹⁴N- and ¹H-Coupling Constants: Spin Distribution. The singly occupied orbital in the radical anions of azo compounds is essentially an antibonding π MO of the –N=N– group. In the azoarenes, there is a substantial conjugation between the π system of this group and those of the aromatic moieties. The ¹⁴N-coupling constant, a_{N} , for the radical anion of (*E*)-azobenzene is +0.45 to 0.50 mT (dependent on the experimental conditions [18]) and it is *ca.* +0.40 mT for the radical anions of (*E*)-azonaphthalenes [19]. In these species, the two N-atoms of the azo group accommodate *ca.* 40–45% of the π -spin population. On passing to the radical anions of azoalkanes, the a_{N} value nearly doubles; irrespective of the conformation of the azo group and the structure of the alkane moieties, it is +0.78 to +0.97 mT, ([6–8], Tables 1–3). The π -spin population in the radical anions of azoalkanes is thus, to the extent of 80–90%, restricted to the azo group. An impressive example is provided by the radical anion of 5,6-dihydrodibenzo[*c,g*][1,2]diazocine (**11**)



which is a bridged derivative of (*Z*)-azobenzene. The a_{N} value for **11**^{•-} is +0.915 mT [19], *i.e.*, it resembles that for the radical anions of azoalkanes, because the non-planar geometry of **11** impairs an effective conjugation between the azo group and the aromatic moieties. Correspondingly, the coupling constants of the protons of these moieties are very strongly reduced relative to corresponding values for the radical anion of (*E*)-azobenzene.

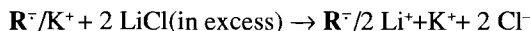
According to ESR nomenclature [12b], bridgehead protons (H_{br}) which are separated from the π system of the azo group in 4^{\ominus} , 6^{\ominus} , and 9^{\ominus} , have to be denoted β , while the methano *syn*- and *anti*-protons (H_{syn} and H_{anti}) in 4^{\ominus} , 5^{\ominus} and 7^{\ominus} , as well as the ethano *endo*-

and *exo*-protons (H_{endo} and H_{exo}) in 4^- – 10^- , should be called γ . The hyperfine interaction of the β -protons with the unpaired electron in a π system mainly implies hyperconjugation and is, thus, proportional to $\cos^2\theta$ where θ represents the dihedral angle between the p_z -axis at the adjacent π centre and the direction of the C– H_β bond [20]. As the sign of the coupling constant $a_{H\beta}$ is the same as that of the π -spin population at this centre, it is usually positive. The small $a_{H\beta}$ value (+0.05 mT) for 4^- complies with the large angle θ (*ca.* 75°) in diazabicycloheptenes. Enlargement of the C-framework to that of diazabicyclooctene increases θ to 90° , *i.e.*, the bridgehead protons come to lie in plane of the π system of the azo group. Hyperconjugation is, thus, ineffective in 9^- , and only spin polarization contributes to $a_{H\beta}$, resulting in a very small and negative value (–0.02 mT). Although coupling to the γ -protons is, in general, considerably weaker than to the β -protons [12b], special arrangements of the C– H_γ bonds relative to the p_z -axis at a π centre can greatly favour the γ -proton-hyperfine interaction with the unpaired electron. This is the case for H_{exo} in 4^- – 10^- and, to a lesser extent, for H_{anti} in 4^- , 5^- , and 7^- , such an arrangement occurring in a W or zig-zag fashion [14]. A long-range electron transfer of this kind is referred to as homohyperconjugation or through-bond interaction [21]. The values of $a_{H_{exo}}$ are relatively large and positive (+0.34 to +0.51 mT for 4^- – 8^- and +0.27 to +0.30 mT for 9^- and 10^-), whereas those of $a_{H_{anti}}$ are smaller in magnitude and negative (–0.21 to –0.23 mT). This sign of $a_{H_{anti}}$ must be due to the location of the CH_2 bridge in the nodal plane of the singly occupied, antibonding MO of the azo group. Substitution of H_{anti} at C(7) by a Me group, on going from 4^- to 6^- and from 7^- to 8^- , replaces the coupling constant of a γ -proton by that of three δ -protons which are separated from the π system by an additional sp^3 -hybridized atom. Interestingly, the coupling constant of these δ -protons still retains half of the $|a_{H_{anti}}|$ value, but its sign changes to positive (+0.11 to +0.13 mT). Molecular models indicate that the C– H_δ bonds in several conformations of the Me group adopt the favourable zig-zag arrangement relative to the p_z -axis at the adjacent π centre. The positive sign of the pertinent coupling constant can be accounted for by the position of the Me δ -protons which are, in general, situated outside the nodal plane of the singly occupied π orbital.

Alkali-Metal Coupling Constants: Association of Radical Anions with their Counterions.

The hyperfine splittings from the magnetic nuclei of alkali-metal cations in the ESR spectra of organic radical anions are the most convincing evidence of the association between these anions and their positively charged counterions Li^+ , Na^+ , K^+ , Rb^+ , and Cs^+ [12c]. The pertinent 7Li -, ^{23}Na -, ^{39}K -, $^{85,87}Rb$ -, and ^{133}Cs -coupling constants have been determined by ESR [12c][22] [23], NMR [24], and (more recently) ENDOR/general-TRIPLE-resonance spectroscopy [10c] [25]. In the few cases studied by the NMR method, absolute signs were allotted to these coupling constants, albeit use of concentrated solutions frequently modified the mode of association relative to that occurring in diluted solutions investigated by ESR/ENDOR technique. Such a reservation is not required for the general-TRIPLE-resonance experiments performed on the ENDOR spectra. Although only relative signs of the coupling constants are obtained by this technique, absolute signs can readily be derived by means of appropriate reference, as has been done in the present paper (see above). The $|a_{Li}|$, $|a_{Na}|$, $|a_K|$, and $|a_{Cs}|$ values observed here are in a ratio of *ca.* 2:4:1:8 which has also been found in several other studies [22]. This ratio roughly follows the order of the hyperfine parameters calculated for a unit *ns*-spin density at the alkali-metal atom [26] ($n = 2, 3, 4$, and

6 for Li, Na, K, and Cs, respectively), thus indicating similar magnitude of the spin populations in the *ns* orbitals of the various counterions. It has been suggested [22] that the sign of the alkali-metal-coupling constants can be deduced from the temperature dependence of its absolute value. The sign is positive (negative), when this value increases (decreases) on warming the solution. According to this criterion, the coupling constants a_{Na} , a_{K} , and a_{Cs} determined in the present work should be positive, whereas a_{Li} is expected to have a negative sign. Such a prediction is in agreement with the results of general-TRIPLE-resonance spectroscopy as far as a_{Li} and a_{Cs} are concerned (see above). Association of the radical anions with Li^+ differs in some aspects from that with Na^+ , K^+ , and Cs^+ . As stated in a preceding section, *two* Li^+ counterions are associated with 1^- – 10^- against a *single one* Na^+ , K^+ , and Cs^+ . The formation of these ion triples $\text{R}^-/2\text{Li}^+$, instead of ion pairs R^-/Na^+ , R^-/K^+ , and R^-/Cs^+ , can be formulated as



where $\text{R}^- = \text{1}^-$ – 10^- . The negative sign of a_{Li} , which contrasts with the positive one of a_{Na} , a_{K} , and a_{Cs} , indicates that, despite similar magnitude of the *ns*-spin populations, the mechanism of electron-spin transfer from the radical anion to the counterion is other for $\text{R}^-/2 \text{Li}^+$ than for R^-/Na^+ , R^-/K^+ , and R^-/Cs^+ . Possibly, two predominantly covalent N–Li bonds are formed in the π plane of the azo group, as the ‘hard acid’ Li^+ is akin to the ‘hard base’ represented by the lone pairs of the N-atoms. The spin transfer through these bonds to the Li-atoms should occur by polarization, like that onto the H-atoms directly linked to the π centres. On the other hand, in R^-/Na^+ , R^-/K^+ , and R^-/Cs^+ , the larger alkali-metal cations being ‘softer acids’ prefer to be attached to the ‘softer base’, which is the π system of azo group; in this case, spin transfer to the counterions should take place by delocalization. A striking example for different modes of association of an organic anion with Li^+ and K^+ has been reported in 1978 [27].

Experimental. – All compounds were synthesized according to procedures described previously: **1** [28], **2** [29], **3** [29][30], **4** [31], **5** [31], **6** [32], **7** [9], **8** [4][33], **9** [34], and **10** [9][35].

The alkali-metal mirrors, used for the reduction of **4**–**10** were generated from K chips and by thermolysis of NaN_3 (*Fluka*) or CsN_3 (*Eastman Kodak*). The appearance of radical anions associated with K^+ upon first contact of **4**–**10** with the Na mirror was due to a KN_3 impurity present in NaN_3 . The K content became larger on thermolysis and sublimation, due to the greater volatility of K as compared to Na, and reaction of **4**–**10** with K was favoured, because of the higher reducing power of K relative to Na.

Cyclic voltammograms were recorded on a *Metrohm Polarecord E506* with a VA Scanner 612/VA Stand 663. ESR spectra were taken on a *Varian-E9* instrument, while a *Bruker-ESP-300* system served for ENDOR and TRIPLE-resonance studies.

Appendix. – The effect of the ^{14}N -hyperfine anisotropy on the line-widths (lw) in the ESR spectra of 1^- – 10^- can be expressed as [12][13]

$$lw = A + BM_1 + CM_1^2$$

where A, B, and C are empirical parameters, and $M_1 = +2, +1, 0, -1, -2$ is the ^{14}N -spin quantum number which decreases with increasing magnetic field. The lw values for the individual M_1 components have to be determined experimentally from the heights of the corresponding lines which are inversely proportional to $(lw)^2$. As an example, the line-widths thus obtained for 2^- associated with 2Li^+ at 203 K amount to 0.214, 0.104, 0.074, 0.124, and 0.254 mT for $M_1 = +2, +1, 0, -1, -2$, respectively, involving the parameters $A = 0.074$, $B = -0.010$, and $C = 0.040$ mT. Fig. 7 shows the pertinent ESR spectrum, along with two computer simulations: one of them (1) makes use of a constant line-width, 0.074 mT, characteristic of the central component with $M_1 = 0$, while the second (2) employs the five different lw values given above. Clearly, the second simulation reproduces the

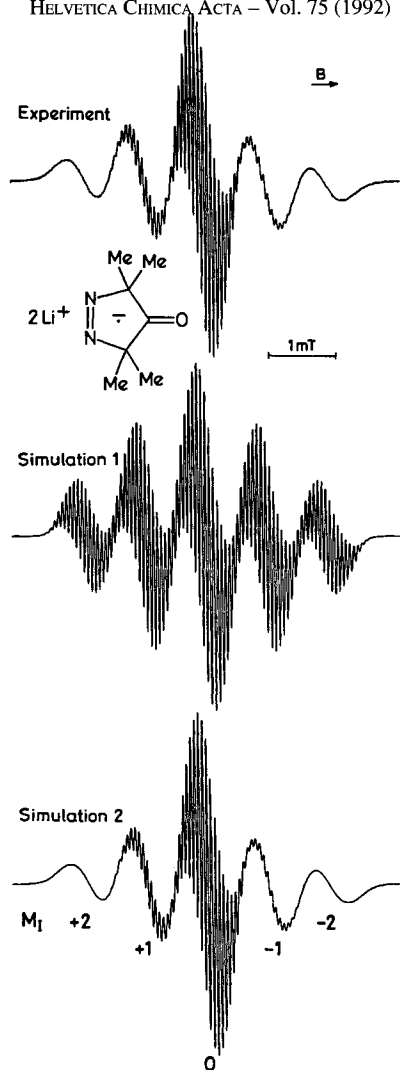


Fig. 7. ESR spectrum of $2^{\bullet-}$ and its simulations. Solvent: DME; counterions as indicated.; temp. 203 K. The simulations made use of the coupling constants given in Table 1. Line-shape: Lorentzian; line-width as indicated in the text: constant for simulation 1 and variable for simulation 2.

experimental spectrum in more details. The computer program for this simulation is presented in [36]. Experimental difficulties in determining the lw values emerge, when the a_n is considerably smaller than the sum of the remaining coupling constants, so that the individual M_I components strongly overlap.

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