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Room-Temperature Electron Spin Relaxation of Triarylmethyl Radicals at X- and Q-bands

Andrey A. Kuzhelev, ^{†,‡,#} Dmitry V. Trukhin, ^{†,‡,#} Olesya A. Krumkacheva, ^{§,‡} Rodion K. Strizhakov,[†] Olga Yu. Rogozhnikova, ^{†,‡} Tatiana I. Troitskaya, ^{†,‡} Matvey V. Fedin, ^{§,‡,*} Victor M. Tormyshev, ^{†,‡,*} and Elena G. Bagryanskaya^{†,‡,*}

[§] International Tomography Center SB RAS, Novosibirsk 630090, Russia

[†]N.N. Vorozhtsov Novosibirsk Institute of Organic Chemistry SB RAS, Novosibirsk 630090,

Russia

[‡] Novosibirsk State University, Novosibirsk 630090, Russia

these authors contributed equally

ABSTRACT

Triarylmethyl radicals (trityls, TAMs) represent a relatively new class of spin labels. The long relaxation of trityls at room temperature in liquid solutions makes them a promising alternative for traditional nitroxides. In this work we have synthesized a series of TAMs including perdeuterated Finland trityl (D₃₆ form), mono-, di-, and tri-ester derivatives of Finland-D₃₆ trityl, deuterated form of OX63, dodeca-*n*-butyl homologue of Finland trityl, and triamide derivatives of Finland trityl with primary and secondary amines attached. We have studied room-temperature relaxation properties of these TAMs in liquids using pulsed Electron Paramagnetic Resonance (EPR) at two microwave frequency bands. We have found the clear dependence of phase memory time $(T_m \sim T_2)$ on magnetic field: room-temperature T_m values are ~1.5-2.5 times smaller at Q-band (34 GHz, 1.2 T) compared to X-band (9 GHz, 0.3 T). This trend is ascribed to the contribution from g-anisotropy that is negligible at lower magnetic fields but comes into play at Q-band. In agreement with this, the difference between T1 and Tm becomes more pronounced at Q-band compared to X-band due to increased contributions from incomplete motional averaging of g-anisotropy. Linear dependence of $(1/T_m-1/T_1)$ on viscosity implies that g-anisotropy is modulated by rotational motion of trityl radical. Based on the analysis of previous data and results of the present work, we conclude that in general situation where spin label is at least partly mobile, X-band is most suitable for application of trityls for room-temperature pulsed EPR distance measurements.

I. INTRODUCTION

Triarylmethyl (TAM, trityl) radicals appeared recently as a new class of spin labels having long electron spin relaxation on the order of microseconds in liquids at room temperature.¹ This profoundly long relaxation makes trityls a promising alternative for nitroxide spin labels in biological applications of EPR. The development of this research field requires further improvement of trityl-based labels in order to achieve even longer T₁ and T₂ (T_m), as well as identification of the factors affecting relaxation in liquids at 300 K. Apart from low-temperature EPR and general studies of TAMs,²⁻⁶ the nanoscale distances were recently measured at 4 °C in protein⁷ and at 37 °C in DNA duplex⁸ using TAM labels and pulsed dipolar EPR spectroscopy. Physiological temperatures were reached owing to the long phase memory time (T_m) of TAMs.¹ Phase memory time is one of the most critical parameters for dipolar EPR spectroscopy determining the range of available distances.⁹⁻¹⁰ For example, T_m=0.7 µs at 4 °C in TAM-labeled protein allowed measuring distances up to 2.5 nm.⁷ An improvement of T_m to 1.4 µs at 37 °C in TAM-labeled DNA duplex allowed extending the available distances up to 4.6 nm.⁸

Thus, development of strategies for further increase of T_m is a highly topical task. Note, however, that covalent attachment of spin label to biopolymer usually leads to a decrease of roomtemperature T_m value compared to that for free radical in solution. This stems from larger rotational correlation times and the presence of anisotropic interactions. Anisotropies of hyperfine interaction (HFI) and g-factor are the two primary sources causing the decrease of T_m when the mobility of spin label becomes restricted. With respect to free TAM radicals, the multifrequency studies of Finland trityl at X-, S-, L-bands and 250 MHz revealed that HFI anisotropy plays dominant role and the contribution of g-anisotropy is negligible.¹

The available methods for measurements of long interspin distances at physiological temperatures were recently complemented by saturation-recovery EPR technique.¹¹⁻¹² In this approach the spin lattice Relaxation Enhancement ($1/T_1$) of spin labels caused by the paramagnetic metal (Cu²⁺) is studied, and the range of available distances crucially depends on the values of T₁.¹¹⁻¹⁶ Therefore, due to the extraordinary long T₁ values (~10-20 µs) trityl radicals seem to be promising probes for this approach. The frequency dependence of T₁ for Finland trityl and its derivative OX63 was studied in the range of 250 MHz to 9.2 GHz at room temperature.^{1, 17} In particular, it was found that at 250 MHz T₁ is determined by inter- and intra-molecular electron–proton dipolar couplings modulated by molecular tumbling with the rates comparable to the reciprocal of the resonance frequency. The contribution of this mechanism is reduced at higher

 frequencies and becomes negligible at X-band, where spin-lattice relaxation of TAMs is contributed mainly by frequency-independent local mode process with energy of 590 cm⁻¹.¹⁷

As a rule, performing pulsed dipolar EPR experiments at frequencies higher than X-band allows one to increase the S/N ratio, as was demonstrated using nitroxide-based spin labels at Q-band.¹⁸⁻¹⁹ Therefore, one might naturally assume that the same tendency should hold for room-temperature distance measurements on trityl radicals. Up to date, however, there was no clear understanding of the factors affecting electron relaxation times of trityl labels attached to biopolymers. Therefore, the data on electron relaxation times of trityls at X-band and higher frequencies should have a high practical importance for applications of pulsed dipolar EPR.



Scheme 1. Structures of studied trityl radicals.

In this work, we synthesized the broad range of triarylmethyl radicals (TAMs) with varying number and structure of substituents (Scheme 1) and measured their relaxation times at room temperature in liquids. In order to find optimal structures of trityl spin labels and experimental conditions for room-temperature distance measurements in biomolecules we focused on comparison of T_m for different trityls, as well as on field dependence of T_m in the range 9-34 GHz. In addition, the T_1 values for newly synthesized trityls also represent very useful data for applications other than pulsed dipolar EPR, e.g. for distance measurements at physiological temperature using Relaxation Enhancement. Note that we studied different derivatives of Finland trityl including those that were

already used successfully for distance measurements at physiological conditions, namely TAM with piperazine substituent (**FP3**)⁸ and TAM with amide substituent (**FBA3**).⁷

II. EXPERIMENTAL

II.1. Synthesis of Triarylmethyls

The perdeuterated form of Finland trityl (**FD**, see Scheme 2 and Supporting Information) was prepared by the two-step one-pot reaction, i.e. by conversion of the corresponding diamagnenic triarylmethanol to trityl cation with the subsequent reduction of this cation with tin(II) chloride.²⁰

High reactivity of bromomethyl acetate toward anionic nucleophiles was used in synthesis of esters **FDAM3**, **FDAM2** and **FDAM1**. The reaction of triethylammonium salt of **FD** with 1.8 molar equivalents of bromomethyl acetate smoothly afforded a mixture of mono-, di- and triesters. The title products were isolated in acceptable yields (27, 32 and 19 % respectively) using column chromatography technique.

Activation of the triacid **FD** (Scheme 2) with carbonyldiimidazole readily afforded the triester **FDME3** in high yield. It was isolated and characterized in the form of trihydrochloride derivative. Application of alternative methods typical of the acid activation, e.g. with dicyclohexylcarbodiimide or conversion of initial acid to OSU ester or acyl chloride derivatives, were found to be less useful: these reactions resulted in badly contaminated mixtures of products.



Scheme 2. Synthesis of the Finland trityl derivatives FDME3, FDAM3, FDAM2 and FDAM1.

 Contrary to ester **FDME3**, the amides **FP3** and **FBA3** were readily obtained by reaction of piperidine with tris-chloroanhydride form of triacid **IH**, and by coupling of benzyl amine with tris-OSU ester **FOSU3**, respectively (Scheme 3).



Scheme 3. Synthesis of trityls FP3 and FBA3.

The synthesis of deuterated analogue of **OX63** is shown in Scheme 4. The initial triarylmethanol **VID**²¹ was treated with formic acid to remove all the protecting *tert*-butyl groups by substituting them with formate groups. The solid material obtained after removal of volatile components was converted to trityl cation in reaction with trifluoromethanesulfonic acid, the cation was then reduced to intermediate TAM.



Scheme 4. Synthesis of OX63D.

To remove formate groups, the latter was hydrolyzed with excess of aqueous NaOH to give a solution of sodium salt of trityl **OX63D**. The resulting solution was acidified by its slow addition to diluted aqueous hydrochloric acid to afford a fine crystalline precipitate of the title TAM.

The trityl **DBT** (Scheme 5) was obtained by analogy with the protocol focused on Finland trityl.²⁰ A reaction of nonanone-5 with 1,2,4,5-tetra-*tert*-butylthiobenzene in the presence of boron trifluoride readily afforded 2,2,6,6-tetra-*n*-butyl-benzo[1,2-d;4,5-d']bis[1,3]dithiole **VIII.** Triarylmethyl alcohol **IX** was then prepared by treatment of arene **VII** with n-BuLi and the subsequent addition of 0.32 equiv of diethylcarbonate. Purification of the crude product did not require a lengthy and tedious column chromatography, instead it was completed by refluxing the crude product with 1:1 mixture of hexane and carbon tetrachloride.

The arene **IX** was converted to a slurry of tris-lithium derivative by treatment with *n*-BuLi in TMEDA/hexane solution. This organolithium intermediate readily reacted with solid carbon dioxide to afford the triacid **X** in good isolated yield (57 %). The title TAM **DBT** was prepared in quantitative yield by the known literature method.²⁰ The identity of TAMs obtained in this study was confirmed by element analyses and combination of instrumental methods: high-resolution ESI/MS, EPR and FTIR spectroscopies.



Scheme 5. Synthesis of TAM substituted with n-butyl groups at side positions (DBT).

II.2. EPR procedures

All used solvents were preliminary distilled. Concentrations of dissolved trityls were ~100-200 μ M in water, ~50-100 μ M in methanol and chloroform to avoid any effects of the radical-radical collisions on electron spin relaxation. The concentration for each sample / each solvent was adjusted experimentally, starting at 0.2 mM solution and diluting it until the spectrum shape and relaxation times (in evacuated samples) became completely concentration-independent. The

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samples were sealed off in the glass capillary tubes (OD 1.5 mm, ID 0.9 mm, max. volume 50 μ l). In order to prevent the line broadening and relaxation enhancement caused by oxygen, deoxygenation was performed by repeated "freeze-pump-thaw" cycles for at least ten times. For each sample we ensured that the further evacuation does not increase the observed relaxation times T₁ and T₂. The obtained T₁ and T₂ values for **FD** and **FH** in H₂O perfectly agree with those reported previously,^{1, 17} thus confirming the efficiency of deoxygenation.

Pulsed and low temperature CW EPR experiments were carried out at X-band (9 GHz) and Qband (34 GHz) using commercial Bruker ELEXSYS E580 spectrometer equipped with an Oxford flow helium cryostat and temperature control system.

Measurements of electron spin relaxation ($T_m \sim T_2$, T_1) were performed at X- and Q-bands at T=300 K. T_m was measured using two-pulse Electron Spin Echo (ESE) sequence, T_1 was measured using inversion-recovery technique with inversion π -pulse and detecting two-pulse ESE sequence. These experiments were carried out using 2-step phase cycling to eliminate a signal of FID. The pulse lengths were 100/200 ns at X-band and 120/240 ns at Q-band for $\pi/2$ and π pulses, respectively. ESE was measured at field position corresponding to the maximum of the EPR spectrum. Obtained relaxation curves were fitted well using monoexponential function. All measured relaxation times were reproduced for at least 3 times for independently prepared new sample each time. All pulse EPR experiments were performed with repetition time of 300 μ s and 3000 μ s at 300 K and 80 K, respectively. FID-detected EPR spectra were recorded at Q-band at 80 K; the pulse length was 1200 ns.

Experimental CW EPR settings at 140 K were as follows: sweep width 2.5 mT; microwave power $6.142 \cdot 10^{-7}$ W; modulation frequency 100 kHz; modulation amplitude 0.1 mT; time constant 81.92 ms; sweep time 83.89 s; number of points 1024; number of scans 4.

CW EPR experiments at room temperature were performed at X-band using commercial spectrometer Bruker EMX. Experimental CW EPR settings at room temperature were as follows: sweep width 0.1 - 0.5 mT; microwave power 1-6 μ W; modulation frequency 100 kHz; modulation amplitude $2 - 5 \mu$ T; time constant 327.68 – 1310.72 ms; sweep time 167.77-671.09 s; number of points 512; number of scans 1.

In order to determine isotropic g-factor of TAM (g_{iso}) at 300 K, we simultaneously recorded X-band CW EPR spectra of two degassed samples placed in separate sample tubes (**FD** as a reference and the investigated TAM), and thus obtained target g_{iso} value relative to the known g_{iso} of **FD** (2.0026²²).

III. RESULTS

III.1. CW EPR study

Before studying electron spin relaxation of series of TAMs, we investigated all these radicals using continuous wave (CW) EPR in order to identify their purity and HFI couplings. Figure 1 shows X-band CW EPR spectra of studied trityl radicals obtained at 300 K in different solvents. The experimental spectra were simulated using Easyspin software (see SI).²³ The obtained spectroscopic parameters are summarized in Table 1.



Fig. 1. X-band CW EPR spectra of studied trityl radicals at 300 K in different solvents.

CW EPR spectrum of **FD** radical consists of single narrow EPR line of Gaussian shape due to the unresolved hyperfine constants on methyl deuterons of the aryl rings. Similarly, CW EPR spectra of **OX63D** and **DBT** are also simulated by single line, but taking into account both Gaussian and Lorentzian broadenings. The spectra of other radicals **FDME3** and **FDAM1-3** demonstrate additional splitting due to the hyperfine interaction with methylene protons $-O-CH_2$ - of side ester

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moieties. EPR spectra of **FDME3** and **FDAM3** show 7 well or partially resolved lines that were simulated using two groups of three equivalent protons with slightly different hyperfine constants. The observed difference between hyperfine constants should be assigned to the effect of chirality.^{21,}

²⁴⁻²⁵ Due to steric hindrance caused by bulky tetrathioaryl groups attached to central carbon atom, trityl radicals adopt a propeller-like conformation in which all rings have the same direction of twist. This geometrical arrangement allows two conformations to occur, i.e. the right-handed propeller (P-conformation) and left-handed propeller (M-conformation). The propeller-shape conformation of TAM associated with a slow interconversion of conformers is responsible for the diastereotopic character of geminal hydrogen atoms of methylene groups bonded to carboxyl moieties. In addition, the steric hindrance caused by adjacent sulfur atoms creates severe obstacles for free rotation of carboxyl-derived groups.⁶ The combination of hindered rotation of parasubstituents and general chirality of TAM core make the two protons of each methylene group inequivalent. Thus, the total of 6 protons is subdivided into two diastereotopic groups with three protons in each. The A_{iso}(¹H) values of methylene proton of **FDAM3** are two times smaller compared to FDME3. EPR spectra of FDAM2 and FDAM1 consist of 5 and 3 partially resolved lines indicating hyperfine interaction with 4 and 2 methylene protons, respectively, with hyperfine constants close to those in FDAM3. In addition to the main EPR signal of FDAM1, a weak singlet line was observed at g-value corresponding to the signal of FD radical. It was assigned to the product of ester solvolysis, whose amount does not exceed 13% in freshly prepared samples in water (see SI).

The EPR spectrum of triamide derivative of Finland trityl **FBA3** in chloroform consists of 11 partially resolved lines. This spectrum was simulated using 3 equivalent ¹⁴N HFI constants and two groups of three equivalent protons with slightly different hyperfine constants related to protons of methylene groups bonded to amine. The hyperfine splitting on proton of amine group was not observed; presumably, the hyperfine constant for this group is smaller than the linewidth. In methanol only poorly resolved lines of three ¹⁴N HFI splittings were observed, with all other hyperfine constants being hidden within a linewidth.

The EPR spectra of triamide **FP3**, which represents the derivative of secondary amines, do not show any resolved splitting in both methanol and chloroform indicating that nitrogen constant is smaller compared to **FBA3**; according to performed estimation, it does not exceed 0.01 mT.

The radicals **FDME3**, **FDAM2-3**, **FBA3** (that can be dissolved in MeOH and CHCl₃) show the dependence of hyperfine constant $A_{iso}(^{1}H)$ (arising from methylene protons $-O-CH_{2}$ - for **FDME3**, **FDAM2-3** and $-NH-CH_{2}$ - for **FBA3**) on solvent polarity: the $A_{iso}(^{1}H)$ value decreases as solvent polarity grows, indicating a decrease of spin density on methylene protons.

Trityl	Solvent	$\mathbf{A_{iso}}^{a}$		LW ^a	giso ^a
		$^{1}\mathbf{H}\left(n^{b} ight)$	$^{14}N(n^{b})$	(G; L) ^c	
FDAM3	MeOH	5.0 (3), 4.5 (3)	-	5.1; 0	2.00277
	CHCl ₃	6.1 (3), 4.9 (3)	-	5.1; 0	2.00282
FDAM2	MeOH	5.1 (2), 4.5 (2)	-	5.2; 0	2.00271
	CHCl ₃	6.1 (2), 4.8 (2)	-	5.0; 0	2.00281
FDAM1	MeOH	4.7 (1), 4.3 (1)	-	4.4; 1.0	2.00266
	H_2O	4.6 (1), 5.5 (1)	-	5.1; 0	2.00266
FDME3	MeOH	9.4 (3), 9.1 (3)	-	5.2; 0	2.00274
	CHCl ₃	10.6 (3), 10.6 (3)	-	5.6; 0	2.00278
FD	MeOH	-	-	4.5; 0 ^d	2.00260
	H_2O	-	-	4.6; 0 ^d	2.00260
OX63D	MeOH	-	-	6.0; 2.3	2.00267
	H_2O	-	-	7.5; 0	2.00268
DBT	MeOH	-	-	10.0; 3.9	2.00266
FBA3	MeOH	<6.0(6)	18.2 (3)	20.0; 0	2.00255
	CHCl ₃	15.0 (3), 13.3 (3)	15.3 (3)	13.5; 0	2.00261
FP3	MeOH	-	<10.7(3)	39.5; 0	2.00252
	CHCl ₃	-	<10.7(3)	38.3; 0	2.00253

Table 1. Spectroscopic properties of deoxygenated trityl radicals at 300 K. The values of isotropic hyperfine constants (A_{iso}) and peak-to-peak line widths (LW) are given in μT.

^a The simulated spectra were calculated in MATLAB using the EasySpin software package. Isotropic g-factors are close to the literature values of **FD** ($g_{iso}=2.0026$)

^b n - number of equivalent nuclei

^c G and L correspond to the Gaussian and Lorentzian contributions into the linewidth

^d See Supporting Information for comparison with literature data.

III.2. Measurements of electron spin relaxation times T₁ and T_m

III.2.1. Dependence on magnetic field (X/Q-bands)

We have studied relaxation properties of several perspective trityls including **FD** and its derivatives. In all cases we have observed clear dependence of T_m on magnetic field by passing from X- to Q-band (9 to 34 GHz). Figure 2 exemplifies the observed trend using deuterated Finland trityl. The phase memory time T_m remains closely constant below 9 GHz, but it decreases by a factor of ~2.5 at 34 GHz; at the same time, T_1 remains roughly field-independent between 3 and 34 GHz.

In order to understand the underlying mechanisms of T_m and its magnetic field dependence, we have performed a lot of additional measurements of T_1 and T_m relaxations for the series of trityls depending on solvent viscosity, depending on deuteration of radical and/or solvent, depending on polarity of the solvent and, finally, on radical structure (see summary in Table 2). Despite variation

of T_m values on radical structure, the decrease of T_m by a factor of 1.5-2.5 at Q-band compared to X-band was observed for all studied radicals.



Fig. 2. The dependence of relaxation times T_m and T_1 on mw frequency for deuterated Finland trityl (FD) in H_2O at 300 K. The data at 0.25-3 GHz were taken from Ref.1.

Table 2. Electron spin dephasing time, T_m , and electron spin-lattice relaxation time, T_1 for investigated radicals at 300 K at X- and Q-bands. The values of T_m and T_1 are given in μ s. Accuracy of measurements is about 10%.

Trityl	Solvent	X-band		Q-band		
		T _m	T ₁	T _m	T ₁	
FDAM3	MeOH	8.3	11.1	5.6	9.7	
	CHCl ₃	8.4	9.9	6.0	9.6	
FDAM2	MeOH	8.1	11.6	4.9	11.6	
	CH_2Cl_2	9.6	12.3	6.7	11.3	
	CHCl ₃	8.6	11.0	5.6	10.0	
	CH ₂ ClCH ₂ Cl	6.0	10.5	4.3	10.3	
	CH ₃ CH ₂ OH	6.0	10.1	3.1	10.0	
	Tert-butanol	4.0	11.7	1.4	12.2	
FDAM1	MeOH	7.6	14.3	2.8	12.6	
	H_2O	10.1	14.5	4.7		
FDME3	MeOH	8.5	12.2	5.4	11.2	
	CHCl ₃	9.1	11.4	5.4	11.2	
FD	MeOH	9.2	15.6	3.8	15.6	
	H_2O	10.8	17.1	4.5	16.2	
	D_2O	13.6	17.7	4.5	17.0	
FH	MeOH	6.3	16.0	2.0	15.3	
	H_2O	9.2	15.0	4.6	14.4	
	D_2O	10.4	18.0	4.0	16.8	

OX63D	MeOH	5.8	16.5	1.8	15.6
	H_2O	7.3	16.0	2.2	15.3
	D_2O	7.6	16.1	2.0	16.1
DBT	MeOH	5.0	14.9	2.1	14.3
FBA3	MeOH	6.5	19.5	3.6	19.2
	CHCl ₃	7.2	18.0	4.5	17.6
FP3	MeOH	5.0	23.0	2.8	23.0
	CHCl ₃	4.5	26.4	2.3	25.7

III.2.2. Dependence on solvent viscosity

One of the first factors to be established is what motion does induce the observed relaxational (T_m) behavior. T_m values for studied samples are smaller than T_1 at 9 and especially 34 GHz indicating the presence of additional contributions to spin-spin relaxation apart from spin-lattice relaxation. In order to extract these contributions, the values $(1/T_m-1/T_1)\approx(1/T_2-1/T_1)$ were obtained that depend both on magnetic field and on solvent viscosity (Fig.3).



Fig. 3. The dependence of $(1/T_m-1/T_1)$ values on solvent viscosity for FDAM2 at 300 K at X- (black squares) and Q-bands (blue circles). The solid red lines represent the fits obtained using function $(1/T_m-1/T_1)=a\cdot\eta+b$. Viscosity values²⁶ for used solvents are given in SI.

FDAM2 is soluble in broad range of solvents, therefore we obtained the $(1/T_m-1/T_1)$ values as a function of solvent viscosity η and found linear dependence (Fig. 3). Since rotational correlation time of radical τ_c is proportional to η , the data of Fig.3 also implies linear dependence of $(1/T_m-1/T_1)$ on τ_c . This observation agrees well with the Redfield theory's result for relaxation due to anisotropic interaction (*V*) modulated by rotational diffusion: $(1/T_m-1/T_1) \propto \langle V^2 \rangle \tau_c$. The proportionality factor depends on magnetic field and increases ~4 fold at Q-band compared to Xband, i.e. $\langle V^2 \rangle$ turns out to be approximately proportional to magnetic field. Previous studies on **FD** at X-, S-, L-bands and 250 MHz demonstrated that dominant contribution stems from HFI

 anisotropy rather than from g-anisotropy.¹ The results obtained in our work, however, clearly show that in the range 9-34 GHz the field-dependent contributions play a significant role. As will be discussed below, the field-dependent contributions stem from pure g-anisotropy ([g:g] contribution) as well as from inner product of g- and HFI anisotropies [g:A], and these anisotropies are modulated by radical tumbling, as follows from viscosity dependence in Fig. 3.

III.2.3. Role of protons

Protons (of both radical and the solvent) are known to largely contribute to the phase memory time of nitroxides as well as trityls.^{1, 27} We have studied T_1 and T_m at X/Q-bands for Finland trityl depending on (i) deuteration of the radical (compare deuterated **FD** and its protonated analogue **FH**) and (ii) deuteration of the solvent (H₂O vs. D₂O); in addition, we have studied T_1 and T_m of **OX63D** depending on deuteration of the solvent (Table 2 and Fig.4). In all cases T_1 remains nearly the same at X- and Q-bands (variations <10%). Contrary to that, T_m noticeably changes depending on deuteration. For Finland trityl deuteration of radical and solvent increases T_m by 10-25% at X-band, but at the same time the effect at Q-band is negligible.

III.2.4. Influence of solvent polarity and hydrogen bonding

Methanol and chloroform differ in polarity, but they have almost the same viscosity and thus the same rotational correlation time for the same radical. Therefore, comparison of the relaxation times in methanol and chloroform allows one to study the influence of solvent polarity on electron relaxation. For all trityls that are soluble in both methanol and chloroform (FDME3, FDAM3, FDAM2, FBA3, FP3), the obtained $(1/T_m-1/T_1)$ values were found nearly the same within the accuracy (Table 2 and Fig.5), thus excluding the significant dependence of relaxation time on solvent polarity.

The hydrogen bonding is known to be more effective for atoms with large electronegativity. Therefore, one anticipates two trends to be found: (i) hydrogen bonding between methanol and ester derivatives of TAM (**FDAM1-3**, **FDME3**) should be stronger in comparison with triamide derivatives (**FBA3**, **FP3**); (ii) hydrogen bonding between TAM and methanol should be stronger than that between TAM and chloroform. Comparison of the $(1/T_2-1/T_1)$ values in methanol and chloroform obtained for all these radicals (mentioned in (i)) did not reveal any noticeable influence of hydrogen bonding within experimental accuracy of 20%, both at X- and Q-bands (Fig.5). Further detailed studies in series of suitable solvents are required to find out the effects of hydrogen bonding on relaxation rates of TAMs (if any).



Fig. 4. The dependence of relaxation times T_m and T_1 for FD, FH and OX63D at X- and Q-bands in different solvents (indicated on the plot) at 300 K. Accuracy of measurements is about 10%.

III.2.5. Influence of radical structure

 All studied trityls are soluble in methanol that allows obtaining the field dependence of $(1/T_m-1/T_1)$ as a function of trityl structure. **OX63D** and **DBT** (the derivatives of Finland trityl with methyl groups substituted by heavier groups) demonstrate the larger $(1/T_m-1/T_1)$ values compared to **FD**.



Fig. 5. $(1/T_m-1/T_1)$ values (contributions to $1/T_2$ from incomplete motional averaging of anisotropic interactions) for studied trityls in different solvents at 300 K at X- and Q-bands. Accuracy of $(1/T_m-1/T_1)$ values is about 20%.

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FDAM1-3 radicals include ester substituents attached to carboxyl moieties of **FD**; therefore they have additional hydrogen hyperfine constants ~5 μ T resolved in CW EPR (Fig.1). **FDAM1-3** show X-band (1/T_m-1/T₁) values close to those of **FD** in methanol. At Q-band the (1/T_m-1/T₁) value for **FDAM3** (that has 3 ester substituents) is about 2 times lower than that for **FD**. At the same time, trityls **FDAM3** and **FDME3** (differing by HFI constants of CH₂ groups of the ester moiety) have nearly the same (1/T_m-1/T₁) values at X- and Q-bands, respectively.

The amide derivatives of Finland trityl **FBA3** and **FP3** have the additional nitrogen hyperfine constants ~0.01-0.02 mT. They demonstrate 2 and 3 times higher $(1/T_m-1/T_1)$ values compared to **FD**, respectively. At the same time, the difference between $(1/T_m-1/T_1)$ values of amide derivatives (**FBA3**, **FP3**) and **FD** at Q-band is reduced, resulting in practically the same $(1/T_m-1/T_1)$ values for **FBA3** and **FD**.

We have found that T_1 is sensitive to the substituent introduced into carboxyl moiety of trityl **FD**. There are two trends which were found: the decrease of T_1 upon introduction of the "ester groups", and increase of T_1 upon introduction of "amide groups" (resulting ratio of T_1 for **FDAM3** and **FBA3** is about 2). Replacement of the methyl groups of **FD** by more bulky groups (**OX63D** and **DBT**) does not affect the observed T_1 values.

III.3. EPR spectra in frozen solution

The lineshapes of EPR spectra of radicals in frozen solution yield information on anisotropic interactions that are partly (or fully) averaged by motion in liquids. In order to separate field-dependent and field-independent anisotropies (g- and HFI, respectively), we performed low-temperature EPR measurements at X- and Q-bands in glass-forming methanol. We measured CW EPR spectra at X-band at 140 K and FID-detected EPR spectra at Q-band at 80 K. It was technically difficult to employ the same approach at both mw bands due to insufficient sensitivity of FID detection at 80 K at X-band and, at the same time, easy saturation of CW EPR spectra at Q-band for all temperatures below melting point of methanol. For one of the samples (**FD**) we verified that FID-detected and integrated CW EPR spectra coincide at X-band (see SI).



Fig. 6. EPR spectra of studied trityls in frozen methanol: (top) First integral of CW EPR spectra at X-band at 140 K and (bottom) FID-detected EPR spectra at Q-band at 80 K.

The obtained EPR spectra at X-band are similar for all trityls (Fig. 6). The Q-band spectra of trityls exhibit partially resolved g-anisotropy that slightly depends on radical structure. For semiquantitative comparison of spectra at X- and Q-bands we introduced the value of $\gamma_{1/3}$, defined as the width of the spectrum at the 1/3 height of the maximum. The more common value, FWHM, does not account for the shoulders in Q-band spectra. We obtained nearly the same values $\gamma_{1/3}\approx 0.25$ mT at X-band and ~2.5 times higher values at Q-band, indicating field-dependent contribution of ganisotropy to the lineshape of EPR spectra in frozen solution. Note that the obtained increase of $\gamma_{1/3}$ by a factor of 2.5 at Q-band is smaller than the corresponding increase of the microwave frequency (~3.5). This indicates that anisotropy of hyperfine interaction might also contribute to the line width, especially at X-band.

The obtained spectra were simulated using rhombic g-tensor (see SI). Similar values of g-tensor were found for all trityls (except for **OX63D**) that agree well with the previously published data on **FD**.²² The rhombic g-tensor of **FD** was previously explained by distortion from 3-fold symmetry occurring in frozen solution.²²

IV. DISCUSSION

The rotational correlation time of **FD** in water at 300 K is about 0.8 ns,¹ thus the motion regime corresponds to $\omega_0 \tau_c \ge 1$ at both X- and Q-bands, where $T_1 > T_m$ was experimentally found. Therefore, the values of $(1/T_m-1/T_1)$ describe all contributions to spin-spin relaxation apart from spin-lattice relaxation. The obtained linear dependence of $(1/T_m-1/T_1)$ on solvent viscosity at X- and Q-bands indicates incomplete motional averaging of anisotropic interactions, and different slopes in these two mw bands imply significant contribution of field-dependent anisotropies. In agreement with this, for frozen solutions we have found that the EPR linewidth at Q-band is 2-3 times larger compared to that at X-band.

The only field-dependent anisotropic interaction present in studied systems is Zeeman interaction with anisotropic g-tensor. At the same time, field-independent anisotropic HFI interactions are also unavoidable. Not aiming at comprehensiveness, below we would like to demonstrate that our experimentally obtained trends can be rationalized using these two plausible relaxation mechanisms and reasonable sets of parameters. One should consider the following contributions to the phase memory time, taken in a model of radical with one magnetic nucleus I=1/2 of effective anisotropy and neglecting all terms proportional to $1/(1+\omega_0^2 \tau_c^2)$ (because, as was mentioned above, $\omega_0 \tau_c \ge 1$):²⁸

$$\frac{1}{T_m} - \frac{1}{T_1} \approx \frac{2\beta^2 H^2}{15\hbar^2} [g : g] \tau_c \pm \frac{4\pi\beta H}{15\hbar} [g : A] \tau_c + \frac{7\pi^2}{30} [A : A] \tau_c$$
(1)

Here [g:g], [g:A] and [A:A] are the inner products of corresponding g- and A-tensors, generally defined as $[T^1:T^2] = \sum_{ik} T_{ik}^1 T_{ki}^2$. The first term of eq.(1) is proportional to the square of magnetic field (*H*), the second one to the first power of magnetic field, and the third one is field-independent. We do not observe the quadratic dependence of $(1/T_m-1/T_1)$ on magnetic field (Fig.5), thus it is most reasonable to assume that the observed field dependence stems from an interplay of [g:g] term and cross-anisotropy term [g:A]. Given that previous observations indicate negligible contribution of g-anisotropy and dominant role of HFI anisotropy at $v_{mw} < 10$ GHz,¹ one anticipates that going to higher and higher magnetic field will first reveal the contribution of [g:A] term, and only then the term [g:g] might become dominant. Thus, the observed tendency is qualitatively coherent with previous observations.

Based on the available data, we can only estimate the contribution of the first [g:g] term in eq.(1) and its dependence on magnetic field. The values for **FD** g=[$2.0028 \ 2.0020 \ 2.0030$] obtained

from simulation agree well with literature data²² and give $[g:g] \approx 6 \cdot 10^{-7}$ (details in SI). Introducing this [g:g] value into the first term of eq.(1), $\tau_c = 0.8$ ns, and H=0.3 T (X-band) we obtain $(1/T_m-1/T_1) \sim 0.45 \cdot 10^5$ s⁻¹ in water, i.e. very close to the experimental value measured for **FD** (Fig.5). For H=1.2 T (Q-band) the same estimation yields $(1/T_m-1/T_1) \sim 7 \cdot 10^5$ s⁻¹, that is even higher than the experimental value (see Fig.5), despite the second and the third terms of eq.(1) are not taken into account. Since [g:g] term is known to play a minor role at X-band,¹ the obtained value $(1/T_m-1/T_1)\sim 0.45 \cdot 10^5$ s⁻¹ is certainly overestimated. This might be ascribed to the overestimation of [g:g] parameter obtained on the basis of frozen solution data, since structural distortions (and thus g-anisotropy) of trityls in frozen solution were assumed to be larger compared to fluid solutions.²² Summing up, the account of [g:g] term using reasonable parameters is already sufficient (and even excessive) to explain the experimentally observed field dependence of T_m and $(1/T_m-1/T_1)$.

Estimation of [g:A] and [A:A] terms is much more difficult, since there are many small HFI constants whose anisotropies are not known, as well as relative directions of the corresponding tensors. In addition, our experiments show that interactions of electron spin with nuclei of the solvent are not completely negligible. Therefore, we cannot provide meaningful estimation for these two terms, but, as was reasoned above, contribution of [g:A] term is expected to be higher than that of [g:g] term. Overall, the combination of the first and the second terms of eq.(1) can readily provide necessary mechanism to drive noticeable shortening of T_m when passing from X- to Q-band.

The following observations summarize the "Pros" for the proposed above mechanism of T_m in studied trityls:

(i) Clear field dependence of T_m at 0.3-1.2 T (X/Q-band) is found for all studied radicals;

(ii) Viscosity dependence of $(1/T_m-1/T_1)$ shows that anisotropy is modulated by radical tumbling (not by any kind of local modes or internal motions of radical fragments);

(iii) The $1/T_m$ and $(1/T_m-1/T_1)$ values at 9 and 34 GHz are greater for **OX63D** and **DBT** than for **FD**, as one would expect because of the slower tumbling of **OX63D** and **DBT** that have more bulky substituents than the methyl groups;

(iv) Effect of deuteration of radical and/or solvent on T_m is noticeable at X-band, but an increase of the relaxation time T_m is very moderate (<50%), even when all protons are substituted by deuterons. This clearly means that other interactions add to HFI already at X-band. At Q-band the effect of deuteration becomes nearly indistinguishable, meaning that at this frequency contributions from HFIs become negligible, and thus g-anisotropy dominates the relaxation process.

 (v) Introduction of amide substituents (**FBA3** and **FP3**) leads to increase of $(1/T_m-1/T_1)$ values compared to **FD** by a factor of 2-3, most likely due to the two effects: (i) the slower tumbling rate and (ii) additional hyperfine constants, which may well be anisotropic. At Q-band the $(1/T_m-1/T_1)$ for **FBA3** and **FD** become nearly the same, meaning again that at this frequency contributions from HFIs become negligible, and thus g-anisotropy dominates the relaxation process.

Although the central observation of this work, namely the pronounced decrease of T_m at Q-band compared to X-band, is valid for all studied trityls and therefore is general, several smaller details remain unanswered and need to be addressed in the future. The open questions are:

- Introduction of ester into carboxyl moiety of **FD** increased τ_c , yet the decrease of T_m for **FDAM3-1** compared to **FD** was not observed. Furthermore, the $(1/T_m-1/T_1)$ value for **FDAM3** at Q-band is 2 times smaller than that for **FD**. Why does the increase of the number of ester substituents decrease $(1/T_m-1/T_1)$ value and the magnitude of its change with magnetic field is currently unclear.
- The viscosity of water (~1 cP) is two times higher than the viscosity of methanol (~0.5 cP).²⁶ However, the 1/T_m and (1/T_m-1/T₁) values for FD, OX63D, FDAM1 are greater in methanol than in water at both X- and Q-bands.
- **OX63D** has lower g-anisotropy compared to **FD** (as follows from EPR spectra at Q-band in frozen methanol). However, the increase of T_m at Q-band is nearly the same for both radicals.

Note, that the above peculiarities arise, first of all, from the very long relaxation times of trityls, that makes them sensitive to the subtle influences of structure and environment. These specifics of trityls, which at the same time are their most important advantage for EPR, certainly need to be studied in the future; at the same time, the conclusions concerning field dependence of relaxation and optimum conditions for EPR measurements, e.g. distance measurements, are already in hand. We stress that one of the radicals (monoamide analogue of **FP3**) has already been used by us as spin label for room-temperature distance measurements at X-band.⁸ An attractive progress would seem to be using higher magnetic fields (e.g. mw frequencies 17-94 GHz), smaller sample volumes, better sensitivity. However, the present work clearly shows that this will be done at the expense of shortened phase memory time. Possibly, shorter T_m values at 17 GHz ⁷ compared to those at 9 GHz ⁸ found previously have the same origin as the observations of the present work.

V. CONCLUSIONS

In this work, we describe convenient methods for synthesis of broad variety of triarylmethyl radicals (TAMs) with varying polarity, with varying number and nature of substituents attached to carboxyl moieties located in *para*-positions of TAM aryl rings, and finally with varying structure of the TAM core. The primary aim of these developments is the future application of trityls with optimum relaxation properties as spin labels for room-temperature distances measurements in biomolecules. As the first step, in this work we investigated relaxation of this new series of prospective TAMs in solutions at room temperature. Certainly, the next step is the attachment of most promising TAMs to biomolecules, that might shorten their relaxation times. Because of this, an understanding of mechanisms governing T_m is vital. This work shows that T_m shortens at Q-band compared to X-band, due to the contribution of g-anisotropy terms. Although different SDSL strategies, different linkers and immobilization procedures might affect ultimate T_m times of spin-labeled biomolecules, we anticipate that the dependence on magnetic field similar to the present work will be manifested for partly mobile TAM labels. At the current state of research, we believe that, unless spin label is strongly immobilized, X-band is most suitable for room-temperature distance measurements using TAMs.

ASSOCIATED CONTENT

Supporting Information available: Synthesis of TAMs; Simulation of X-band CW EPR spectra at room temperature; Solvolysis of **FDAM1**; Comparison of linewidth of **FD** with literature data; Properties of solvents; Comparison of FID-detected and CW EPR spectra; Simulation of Q-band FID-detected EPR spectra at 80 K; Estimation of contribution from g-anisotropy; Primary relaxation data. This material is available free of charge via the Internet at http://pubs.acs.org.

AUTHOR INFORMATION

Corresponding Authors:

mfedin@tomo.nsc.ru (M. V. F.), torm@nioch.nsc.ru (V.M.T), egbagryanskaya@nioch.nsc.ru (E.G.B)

<u>Notes</u>

The authors declare no competing financial interests.

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