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The fluorescent properties of several natural benzofuran derivatives, egonols, isolated from *Styrax officinalis*, were determined. The studied egonol derivatives were found to be effective UV organic luminophores, with quantum yields of near to unity, $\varphi_f = 0.92$. The fluorescence quenching experiments with electron donor molecules of aniline, N,N-dimethylaniline and carbazolocarbazole, presented very high efficiency of quenching at above diffusion quenching rates of 4.5×10^{10} , 2.1×10^8 and 5.6×10^{12} $\text{s}^{-1}\text{mol}^{-1}$, respectively. A strong electron acceptor molecule, tetranitrofluorenone, presented quenching rate of 2.6×10^{11} $\text{s}^{-1}\text{mol}^{-1}$. It is concluded that the first excited singlet state of egonol molecule is a more efficient electron acceptor.

Key Words: Fluorescence emission, *Styrax officinalis*, egonol, benzofuran, fluorescence quantum yields, fluorescence quenching.

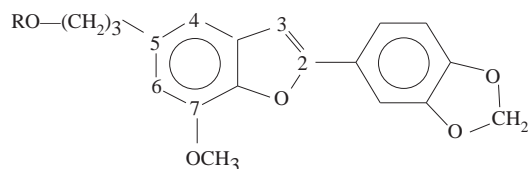
Introduction

Egonol is a natural benzofuran glycoside occurring widely in *Styrax officinalis*¹. First it was isolated by Okada from the seed-oil of *Styrax japonicum*². The structure was suggested as 5-(3-hydroxypropyl)-7-methoxy-2-(3,4-methylenedioxyphenyl)benzofuran, egonol and confirmed in several later publications³. There are also some evidences about the isolation of other derivatives of benzofuran in similar structures⁴⁻⁵. Anil⁶ has reported the isolation and the justification of molecular structure of four benzofuran glycosides from the seeds of *Styrax officinalis*.

Photochemical and photophysical properties of natural products are important as routes to discovery of organic and bio-organic mechanisms under solar conditions. But no investigations are found to be reported on luminescent properties of these natural egonols. In this report we wish to present the results of our studies

on fluorescence parameters of several egonols and fluorescence quenchings with strong π -electron donors and acceptors.

Following three derivatives were investigated for their photochemical properties.



R= **H(I)**, Egonol; R= **Acetyl(II)**, Egonol acetate; R= **Triglycoside(III)**, Egonol triglycoside

Experimental

The separation, identification, and purification of the title compounds were described earlier⁶⁻¹⁶. Aniline and N,N-dimethylaniline were distilled before use at quenching experiments. The electronic absorption spectra were measured on UV-1601 Shimadzu spectrophotometer, fluorescence excitation and emission spectra were determined on PTI QM1 fluorescence spectrometer, at a bandwidth of 1 nm, optical densities on excitation and emission wavelengths during the measurements of quantum yields and quenching experiments were determined on spectrometer Spectronic 601. Quantum yields were measured with respect to quinine sulfate solution in 1N N_2SO_4 ($\varphi_f = 0.546$ ⁷). Quantum-chemical calculations were made by a semiempirical full-valent method AM1⁸. Methanol was supplied from Merck in Uvasol grade.

Correction of fluorescence intensity on inner-filter (the decreasing of fluorescence output with the increasing of absorbance on the excitation wavelength) and reabsorption (the decreasing of fluorescence output with the increasing of absorbance on the emission wavelength) effects during the quenching experiments were made as shown by Miller⁹. The true fluorescence intensity data were calculated from the experimental values according to the following equations:

$$F^0 = F_{exp} \times C^{\text{inner-filter}} \times C^{\text{reabsorption}} \quad (1)$$

$$C^{\text{inner-filter}} = \frac{D_{EX} \cdot \ln(10) \cdot (x_2 - x_1)}{(10^{-D_{EX} \cdot x_2} - 10^{-D_{EX} \cdot x_1})} \quad (2)$$

$$C^{\text{reabsorption}} = 10^{D_{EM} \cdot y_1} \frac{\ln(10) \cdot D_{EM} \cdot y_2}{1 - 10^{-D_{EM} \cdot y_2}} \quad (3)$$

The geometric parameters (x_1 , x_2 , y_1 and y_2), necessary for the calculation of correction factors by eq. (2) and (3), are substantially dependent from the adjustment of optical parts of the fluorescence spectrometer. The evaluations of above parameters for experiments in PTI QM1 device are described below.

Experimental correction factors for inner-filter effects were obtained for quinine sulfate solutions in aqueous 0.1 molar H_2SO_4 at increasing concentration of added acetone which absorbed light in short-wavelength spectral region (Figure 1). Geometric parameters $x_1 = 0.2786$ and $x_2 = 0.6826$ were calculated by the fitting of experimental $C^{\text{inner-filter}}$ data from optical density on the excitation wavelenghtes (DEX). The equation (2) was used with the algorithm of Fletcher-Pauell and by application of a non-linear least squares method (Figure 2).

Experimental correction factors for reabsorption effect were obtained for quinine sulfate solutions in methanol, acidified by a small amount of 0.1 molar aqueous H_2SO_4 , at increasing concentrations of fluorescein, which absorbed and emitted light in long-wavelength spectral region (Figure 3). Absorption and emission data were collected in the spectral region free from fluorescence of added fluorescein. Fluorescence intensities were previously corrected on the inner-filter effect by the equation (2). Geometric parameters $y_1 = 0.2459$ and $y_2 = 0.7462$ were calculated by the fitting of experimental $C^{reabsorption}$ data from optical density on the emission wavelength (D_{EM}). The equation (3) was used with the algorithm of Fletcher-Pauell and by application of a non-linear least squares method (Figure 4).

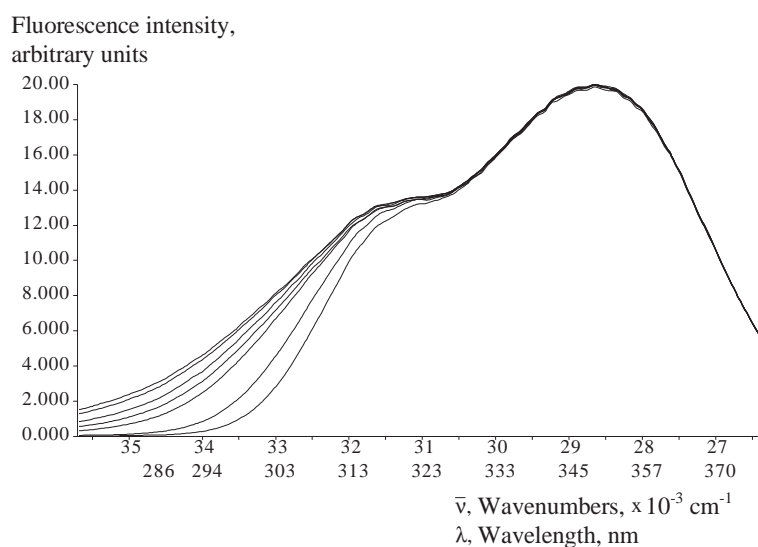


Figure 1. Fluorescence excitation spectra of quinine sulfate with increasing concentrations of added acetone.

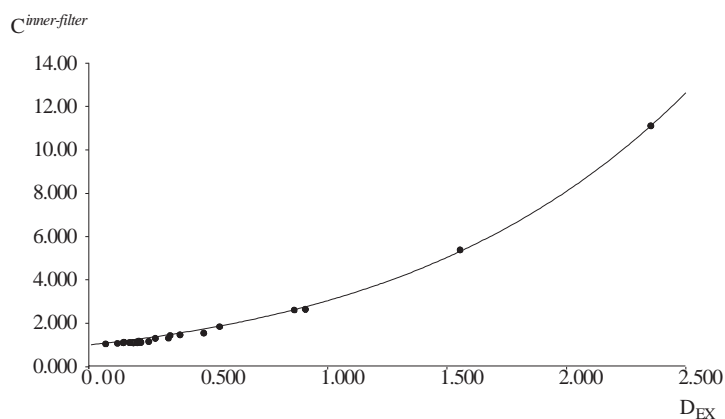


Figure 2. Fitting of the experimental correction factors data for inner-filter effect from the optical density on the excitation wavelength by non-linear least square method.

Results and discussion

In order to be able to interpret the experimental results correctly, we have evaluated the molecular structure of egonol compound at semi-empirical calculations. The structure of the main chromophoric fragment of the

isolated egonol from *Styrax officinalis*, was calculated by the use of AM1 method with a full optimisation of molecular geometry. It was found, that, on the contrary to biphenylic compounds, the egonol molecule is practically planar. The interatomic distance between the hydrogen atom in position 3 of benzofuran cycle and *ortho*-hydrogen of 2-phenyl moiety was calculated to be about 2.3 Å, which is very close to the sum of the corresponding Van-der-Waals radii. The conclusion can be made that no sterical hindrance is present in the natural egonol molecules. Also there are no chromophore groups in egonol molecule, which could cause the appearance of $n\pi^*$ -states. In most cases, effective intersystem crossing with the participation of singlet or triplet states of $n\pi^*$ type results in complete diminish of fluorescence or observation of low fluorescence in conjugated organic molecules^{10–12}. Thus one may predict that natural egonols can be effective organic luminophores.

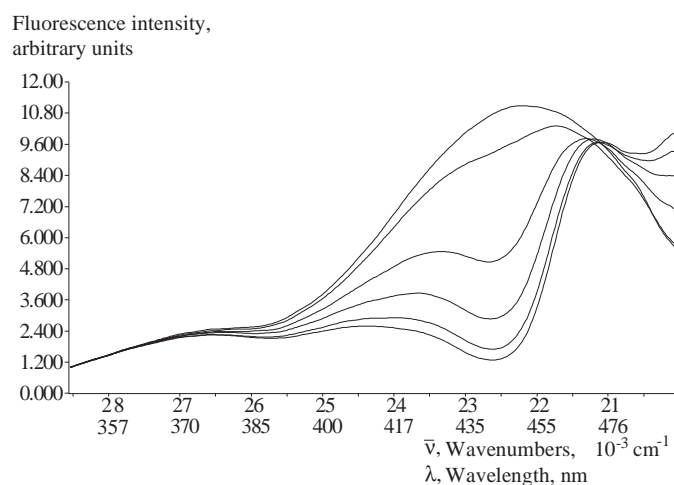


Figure 3. Fluorescence emission spectra of quinine sulfate in acidified methanol with increasing concentrations of added fluorescein.

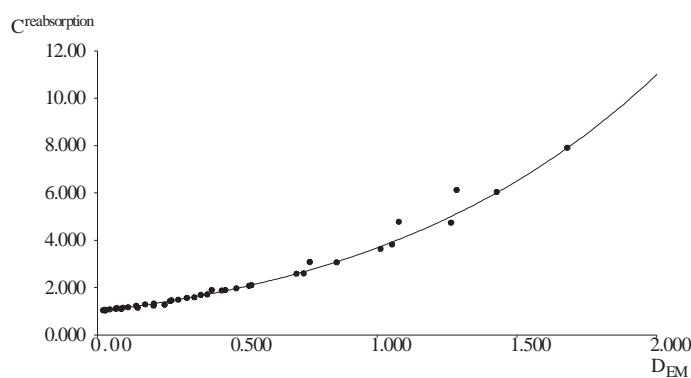


Figure 4. Fitting of the experimental correction factors data for reabsorption effect from the optical density on the emission wavelengths by non-linear least square method.

The spectral characteristics of the studied compounds in methanol solutions are presented in Table 1, the typical fluorescence spectrum is shown in figure 5. In agreement with theoretical approaches, natural egonols are found to fluoresce very effectively, with quantum yield values reaching to unity, $\varphi_f = 0.92$. 5-O-acylation and 5-O-triglycoside substitution on the egonol molecule, do not affect the high fluorescence

emission (Table 1). Egonols are seen to be effective natural organic luminophores for near UV region (Figure 5). Unsubstituted benzofurane is reported to have a fluorescence quantum yield of $\varphi_f = 0.63$ and a phosphorescence quantum yield of $\varphi_{ph} = 0.24$ ¹³. Absorption of egonols, 290-340 nm lies in the short UV end region of solar radiation spectrum. Thus, one can comment that egonols may be one of the organic luminophores in structure of plants, that may transfer the solar light into some photochemical processes.

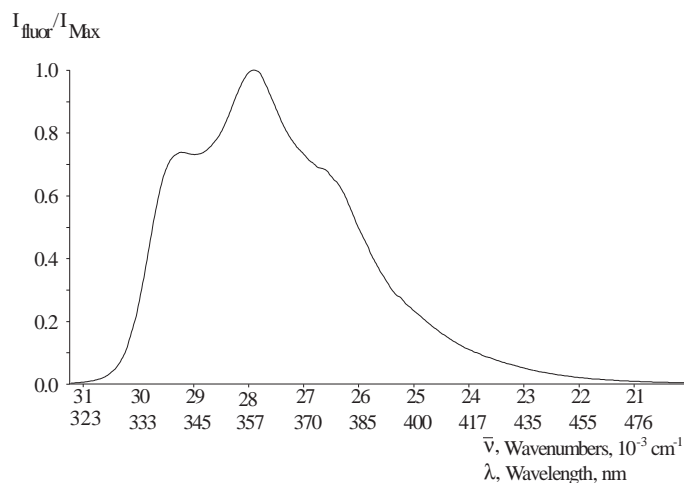
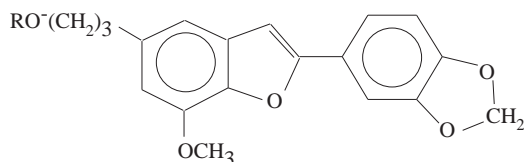


Figure 5. The absorption, **A**, and the fluorescence, **B**, spectra of egonol (I) in methanol, $\lambda_{exc} = 317$ nm, ($[c] = 10^{-5}$ m).

Table 1. The absorption and emission wavelengths (λ), wavenumbers (ν), Stokes shift ($\Delta\nu_{ST}$) and fluorescence quantum yields (φ_{fluor}) of some natural egonol derivatives in methanol.



No	Compound	λ_{abs} (nm)	ϵ_{max} ($\times 10^4$)	ν_{abs} (cm^{-1})	λ_{fluor} (nm)	ν_{fluor} (cm^{-1})	$\Delta\nu_{ST}$ (cm^{-1})	φ_{fluor}
I	Egonol	317	2.75	31550	358	27900	3650	0.923
II	Egonol acetate	317	2.92	31550	358	27900	3650	0.922
III	Egonol triglycoside	317.5	2.95	31500	359	27840	3660	0.918

For a possible application of these compounds in biologically-oriented energy and charge transfer studies, we examined the behaviour of egonol molecule toward the π -electron donors and acceptors. For this aim, theoretical evaluations of the ground and excited states are done in order to find the ionisation potentials and electron affinities. The total energies were calculated in the ground and in the first singlet excited Frank-Condon state for neutral egonol molecule, and for the anion-cation radical forms, by a preliminary optimisation of molecular geometry. The ionisation potentials, calculated as difference of total energy between egonol molecule and its cation-radical, electron affinities, calculated as difference of total energy between egonol molecule and its anion-radical, and presented in Table 2. As seen that ionisation

potential decreases about twofolds from ground state to excited state and electron affinity increases more than threefold from ground state to excited state. Thus one may make the assumption, that the fluorescence of egonol can be quenched more efficiently by electron donors, compared to the electron acceptors.

Fluorescence quenching of egonol emission studies with strong π -electron donor molecule of carbazolocarbazole¹⁴ and also by strong π -electron acceptor molecule of tetranitrofluorenone, prove the above theoretical evaluations¹⁵. Previous quenching experiments by the use of these quenchers, have proven the excessive high quenching abilities¹⁶. The bimolecular rate constants of quenching are reported to be above the limits for diffusion controlled reactions ($> 10^{10} \text{ s}^{-1} \text{ M}^{-1}$), due to formation of charge transfer complexes even in the ground state.

Table 2. Ionisation potentials, $I_p(\text{eV})$ and electron affinities, $E_A(\text{eV})$ of egonol, carbazolocarbazole and tetranitrofluorenone molecules, in the ground and first singlet excited state, S_1^* (only for excited egonol molecule, not for the quenchers) calculated by AM1 method.

	EGONOL	CARBAZOLOCARBAZ.	TETRANITROFLUOR.
I_p at Ground State	7.77	7.54	10.90
I_p at S_1^*	4.40		
E_A at Ground State	1.34	0.92	3.84
E_A at S_1^*	4.72		

The fluorescence lifetime of egonol, which is necessary for calculation of the rate constant of quenching, was estimated on the basis of egonol quantum yield and fluorescence rate constant. The fluorescence rate constant was calculated from the corresponding electronic absorption spectrum¹³. The fluorescence lifetime is calculated to be $\tau_f = \varphi_f / k_f = 2.57 \text{ ns}$, using $\varphi_f = 0.92$, and $k_f = 3.6 \times 10^8 \text{ s}^{-1}$ values. The absorption spectra of both donor carbazolocarbazole and acceptor tetranitrofluorenone quenchers lie at longer wavelengths in comparison with the absorptions of egonols. In addition, the donor quencher, carbazolocarbazole, has a very low fluorescence emission, $\varphi_{\text{fluor}} < 0.1$, that overlaps with the emission band of egonol. The effects described above were taken into account at calculations of Stern-Vollmer plots. The analytical wavelengths for quenchings are selected to be at minimum overlap values for emission band of egonol with the absorption or emission bands of added quenchers (343 nm for quenching by carbazolocarbazole and 400 nm for quenching by tetranitrofluorenone). "Inner filter" and reabsorption effects, aroused with increasing concentrations of quencher, were taking into account by the use described equations of (1)-(3). The Stern-Vollmer plots for quenching of egonol fluorescence by carbazolocarbazole and tetranitrofluorenone are seen in figures 6 and 7. In order to avoid overlapping of emission bands of egonol with the absorption bands of quencher molecule, in addition we have studied the quenching of egonol fluorescence band at $\lambda_{\text{exc}} = 317 \text{ nm}$, with aniline and N,N-dimethylaniline molecules (Figures 8 and 9). The quenching rates with aniline and N,N-dimethylaniline rapidly deviated from linearity above the concentrations of plots in figure 8 and figure 9. Fluorescence quenching rates of egonol, calculated from Stern-Vollmer plots are seen below:

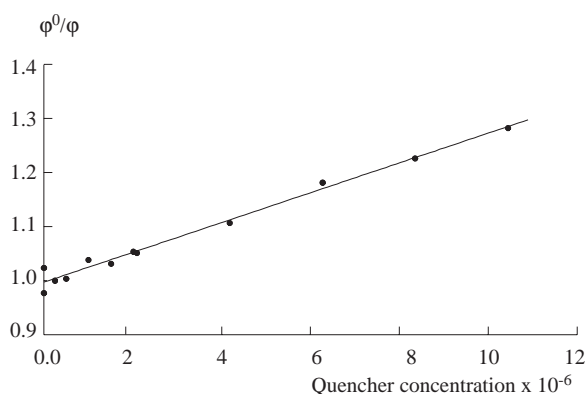


Figure 6. The Stern-Vollmer plot for the quenching of egonol(1×10^{-5} M) fluorescence in the presence of increasing carbazolocarbazole concentrations $(0.1-12.5) \times 10^{-6}$ M in methanol solutions.

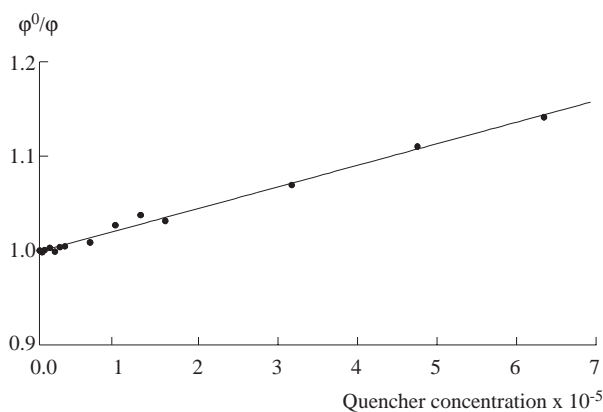


Figure 7. The Stern-Vollmer plot for the quenching of egonol(1×10^{-5} M) fluorescence in the presence of increasing tetranitrofluorenone concentrations $(2-63) \times 10^{-6}$ M in methanol solutions.

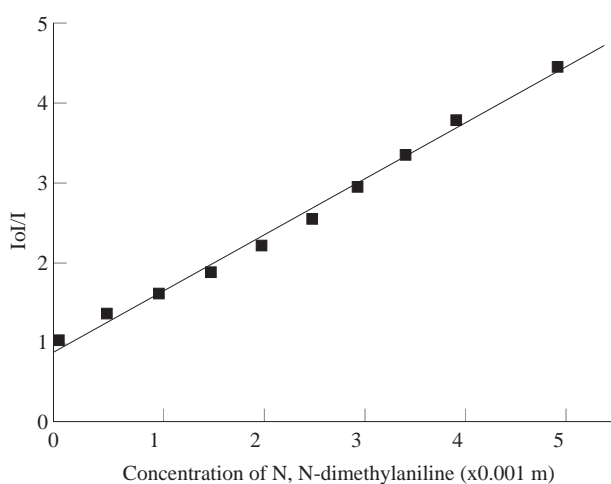


Figure 8. The Stern-Vollmer plot for the quenching of egonol(1×10^{-5} M) fluorescence in the presence of increasing N,N-dimethylaniline concentrations $(0.5-5) \times 10^{-3}$ M in methanol solutions.

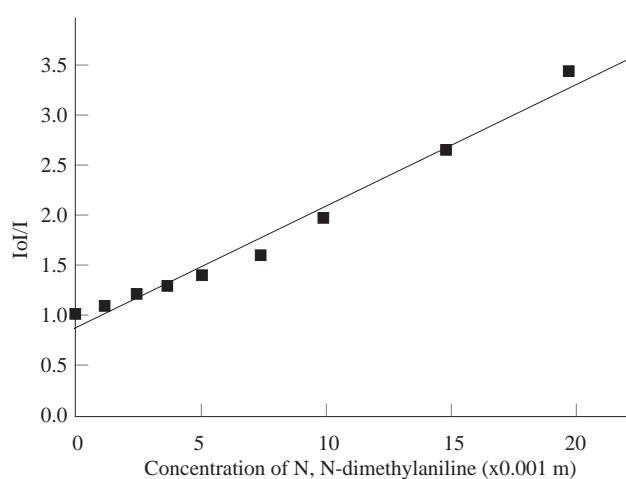


Figure 9. The Stern-Vollmer plot for the quenching of egonol(1×10^{-5} M) fluorescence in the presence of increasing aniline concentrations $(1.3-20) \times 10^{-5}$ M in methanol solutions.

Quencher	Solvent	λ_{exc} (nm)	k_f (s ⁻¹ (mol ⁻¹))
<i>Donors</i>			
Aniline	Methanol	317	4.5×10 ¹⁰
N,N-Dimethylaniline	Methanol	317	2.1×10 ¹¹
Carbazolocarbazole	Methanol	343	5.6×10 ¹²
<i>Acceptor</i>			
Tetranitrofluorenone	Methanol	400	2.6×10 ¹¹

As seen, quenching rates are at or above the diffusion rate limits, for all the quenchers. Increasing π -electron donor capacity in the order of aniline, N,N-dimethylaniline-carbazolocarbazole, accordingly enhances the quenching rates of egonol emission about tenfold. The quenching rate constants are $5.6 \times 10^{12} \text{ s}^{-1} \text{ mol}^{-1}$ for carbazolocarbazole-egonol pair and $2.6 \times 10^{11} \text{ s}^{-1} \text{ mol}^{-1}$ for tetranitrofluorenone-egonol pair. Ionization potentials and electron affinities for egonol, carbazolocarbazole and tetranitrofluorenone is seen in Table 2. The first excited state, $S_1 * -I_p$ and $S_1 * -E_A$ values are calculated for only the egonol, because energy transfer process do not involve the excited states of quenchers. These results are in agreement with semi-empirical calculations, that the excited egonol molecule is a more effective electron-acceptor rather than an electron-donor. But effective fluorescence quenching is seen to be also present toward strong π -electron-accepting compounds, as with tetranitrofluorenone. The evaluated quenching rate constants are found to be above the value diffusion-controlled rate limits, similar observations are reported earlier¹⁴⁻¹⁶. A plausible participation of electron tunnelling between the quencher and the excited fluorescent egonol molecule in liquid state, can be proposed for justification of high efficiency on fluorescence quenching. Such electron tunneling in homogeneous and heterogeneous systems is being well defined by Grätzel¹⁷.

Conclusion

A natural benzofuran type compound, isolated from *Styrax officinalis*, egonol, is found to be effective luminophore, emitting light in the near UV region with a quantum yield values near 0.92. The quenching experiments show very high efficiency of quenching, and that the excited egonol molecules are more efficient electron acceptors, rather than electron donors.

Acknowledgments

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