Theoretical and Experimental Studies of N,N-Dimethyl-N'-Picryl-4,4'-Stilbenediamine

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Abstract N,N-dimethyl-N'-picryl-4,4'-stilbenediamine (DMPSDA) was prepared, purified and crystallised in a form of black lustrous crystals, and its absorption and fluorescence spectra were recorded in cyclohexane, acetonitrile and dimethyl sulfoxide. Non-emissive intramolecular charge transfer state (ICT) was clearly observed in this molecule in all three solvents. Theoretical calculations demonstrating a betaine electronic structure of the trinitrophenyl group in the ground state of the molecule and a charge transfer nature of the long wavelength transition $S_0 \rightarrow S_1$ supported the experimental observations of the ICT formation in the molecule.

Keywords Intramolecular charge transfer · Fluorescence

Introduction

We have previously discussed various time-scale processes occurring with 4,4'-disubstituted stilbenes after their irradiation [1–3]. We found that different 4,4'-disubstituted stilbenes exhibit different sensitivity to intramolecular donor-acceptor effects of substituents and medium polarity. All the investigated 4,4'-disubstituted stilbenes were divided into three groups according to their intramolecular stabilization of the excited $1^t*$ state. Those having strong donor dimethylamino (DMA) group at one phenyl ring and strong acceptor substituent at another phenyl ring constituted the third group called “push-pull stilbenes”. They were characterised by large charge delocalisation between donor and acceptor aromatic moieties, large dipole moment, low polarity $1^p*$ state with preferential stabilisation of $1^t*$ compared to less polar $1^p*$ state, very high activation barrier for the $1^t* \rightarrow 1^p*$ twisted transition, and net increase of the fluorescence quantum yield and fluorescence lifetime with increase of solvent polarity. All these observations indicated completely different non-radiative relaxation pathway than the $1^t* \rightarrow 1^p*$ twisted transition observed in most stilbene compounds, which are not classified as “push-pull stilbenes” [1–3].

Actually, the third group has been formed by three investigated “push-pull stilbenes”, namely 4-dimethylamino-4'-cyanostilbene (DACS), 4-dimethylamino-4'-carbomethoxystilbene (DACMS) and 4-dimethylamino-4'-
nitrostilbene (DANS). These three stilbenes exhibited very large Stock shifts, compared to the first and second groups [1–3]. In that case, the highly polarized excited state, which creates a huge dipole moment, is stabilised extensively by polar interactions (Stock shift value ΔE was found to be approximately 22–23 kcal/mol). Experimental observations of high fluorescence quantum yields and long fluorescence lifetimes in strongly polar solvents indicated a strongly preferred pathway toward the intramolecular charge transfer (ICT) formation in the competing photochemical processes starting from $1^t^*$. Assuming that this point holds in less polar solvents as well, the reduced fluorescence quantum yield and increased non-radiative decay rate is probably due to the insufficient stabilization of the ICT in these solvents. This can easily be understood by assuming that $1^p^*$ is less polar than ICT and $1^t^*$ states, and thus increasing solvent polarity preferentially lowers a highly polar ICT state with respect to $1^p^*$.

Although three aforementioned “push-pull” stilbenes were categorised into one group, they do not necessarily conform to the same relaxation mechanism. In fact, intramolecular charge transfer in these molecules may be observed as an emissive or non-emissive process dependent on the 4,4'-substitution pattern. In case of DANS, we assumed a non-emissive ICT state, which was attributed to a specific interaction of the nitro-group, which quenches the charge-transfer state emission [4, 5]. Actually, non-emissive ICT states were observed in polar solvents when the acceptor group is very powerful and has a low-lying anti-bonding orbital [4]. To generalise this observation, we prepared and studied a new “push-pull” stilbene having a relatively strong acceptor substituent, namely trinitrophenyl (TNP) group. This molecule has been disclosed only once in literature and only its UV absorption spectrum has been previously reported [6].

Herein, we report the synthetic preparation and experimental studies of this molecule, as well as theoretical calculations aimed at elucidating the absolute geometry of the most stable conformer, which might be responsible for the experimentally observed charge-transfer interactions in the ground state of the molecule.

**Experimental**

**Reactants and Solvents**

Reactants were either commercially available or freshly prepared as detailed in the Synthesis section. Commercially available organic solvents, namely cyclohexane (Sigma-Aldrich ACS spectrophotometric grade, ≥99 %), dimethyl sulfoxide (Sigma-Aldrich ACS spectrophotometric grade, ≥99 %) and acetonitrile (Sigma-Aldrich ACS spectrophotometric grade, ≥99 %), used for spectroscopy did not require any additional purification.

**Apparatus and Methods**

Semi-preparative flash chromatography used for purification of the compounds was performed with the Büchi Sepacore® flash chromatography system using 25 g silica gel cartridges (particle size 40–63 μm) and dichloromethane as an eluting solvent. $^1$H NMR spectra were run on 10 % (w/v) sample solutions in CDCl$_3$ with (CH$_3$)$_2$Si as an internal standard at room temperature using a 400 MHz Bruker Fourier transform spectrometer, equipped with a DMX AVANCE I system. UV absorption spectra were measured using an Agilent Cary 300 spectrophotometer, and the steady-state fluorescence spectra were recorded with Horiba Jobin Yvon Fluorolog®-3 modular spectrofluorometer and Agilent Cary Eclipse fluorescence spectrophotometer. The fluorescence excitation and emission spectra were corrected for instrumental sensitivity at different excitation and emission slits using the instrument internal excitation-emission matrix (EEM) correction [7]. A solution of quinine bisulphate in 0.1 N H$_2$SO$_4$ ($\Phi_F=0.52$) was taken as fluorescence standard for the determination of fluorescence quantum yields [8]. Constant-illumination fluorescence intensity decay curves at the photostationary steady-state equilibrium between the trans- and cis-isomers of DMPSDA were recorded with Shimadzu RF-5,301 spectrofluorometer equipped with the 150 W Xenon lamp as a radiation light source. The fluorescence decay at the photostationary steady-state equilibrium was monitored at the emission maximum of DMPSDA after excitation at the excitation maximum using typically 5-nm slit width for excitation and 5-nm slit width for emission. Analysis of the experimental data was performed using Origin® Pro 9.0 for Windows.

Fluorescence intensity decay curves were analysed with a polynomial fit for calculation of the trans-cis isomerization rate constants using a self-written routine within the Origin® Pro 9.0 for analysis of the first-order photochemical reaction rates.

**Synthesis of N,N-Dimethyl-N'-Picryl-4,4'-Stil bendiamine**

The following reactants were prepared by slight modification of our procedure that we published in [9]: trans-4-dimethylamino-4'-nitrostilbene and trans-4-dimethylamino-4'-aminostilbene.

1) **Trans-4-Dimethylamino-4'-nitrostilbene**

An equimolar mixture of 4-nitrophenylacetic acid (Sigma-Aldrich N20204) (9.1 g) and 4-(dimethylamino) benzaldehyde (Sigma-Aldrich 156,477) (7.5 g) in 3 ml of piperidine was heated for 24 h under reflux in an oil bath maintained at
140°. The resulting dark-red crude solid was recrystallized from 100 ml of chlorobenzene. Upon cooling, the product precipitated as lustrous red flakes, then was collected by vacuum filtration on a Buchner funnel, thoroughly washed with petroleum ether (60–80°) to remove the adherent piperidine and dried in a vacuum oven at 60° for about 2 h. Yield of the lustrous red flakes – 7.4 g (53 %). Chemical purity and structural identity of the product was confirmed by ¹H NMR.

(2) Trans-4-Dimethylamino-4′-aminostilbene

Solution of 120 g of potassium hydroxide pellets dissolved in 200 ml of ultrapure water was prepared in 500 ml Erlenmeyer flask and placed in a fridge for storing. This solution is used for decomposition of the prepared stannous complex salt of the product. 15 g of stannous chloride dihydrate was dissolved in 150 ml of concentrated hydrochloric acid at room temperature with stirring. 4 g of fresh trans-4-dimethylamino-4′-nitrostilbene prepared in the previous step was introduced into the stirred solution of SnCl₂. The resulted suspension was refluxed with strong stirring for about 4–5 h until all red flakes and orange viscous particles of the intermediate stannous complex salt formed during the reaction have been completely dissolved, and the yellow reaction mixture has become transparent. The reaction mixture was stirred for additional half an hour and allowed to cool to the point of turbidity. Then it was poured carefully into a cold stirred solution of potassium hydroxide taken from the fridge and placed into an ice bath, in order to decompose the complex stannous salt and to release the product. Caution: the neutralization reaction is highly exothermic and vigour! Well-protective glasses, gloves and clothes should be worn during this step, and all precautions for minimising the reaction mixture splashing should be taken. The crude yellow product was collected by vacuum filtration, thoroughly washed with 10 % sodium bicarbonate solution, ultrapure water and then with petroleum ether (60–80°). The crude yellow product was dissolved in 10 ml of dichloromethane, and subjected to the flash chromatography on the regular silica gel (particle size 40–63 μm). The fractions containing trans-isomer (λ₅₀₀=360 nm) were collected and evaporated to yield the yellow amorphous powder (2.82 g, 77 % reaction yield). Chemical purity and structural identity of the product was confirmed by ¹H NMR.

(3) N,N-dimethyl-N′-picryl-4,4′-stilbenediamine

An equimolar mixture of 0.48 g (2 mmol) of trans-4-dimethylamino-4′-aminostilbene, prepared in a previous step, and 0.50 g (2 mmol) of 2-chloro-1,3,5-trinitrobenzene (Sigma-Aldrich 79,874) in 5 ml absolute ethanol, which yielded initially a clear solution, was left overnight with stirring. The resulted black crystalline precipitate was collected by vacuum filtration on a Buchner funnel, washed with 10 % sodium bicarbonate solution, ultrapure water and then with petroleum ether (60–80°). The crude product was recrystallized from chlorobenzene, and then dried in a vacuum oven at 60° for about 2 h. Yield of the black crystalline product – 0.76 g (85 %). ¹H NMR: δ 3.00 (Me₂N, s, 6H); δ 6.88, δ 6.93 (CH=CH AB, d, vinyl 2H); δ 6.72, δ 7.89 (4-Me₂N-Ar AA′XX′, dd, 4H); δ 7.33, δ 7.83 (4′-picrylamine-Ar AA′XX′, dd, 4H); δ 9.24 (picryl ring protons, s, 2H); δ 3.71 (NH).

Fig. 1 Normalised absorption, excitation and emission spectra of trans-DMPSDA in different solvents: a CH, b ACN and c DMSO, and d the photostationary fluorescence decay curves under constant-illumination conditions in these three solvents.
Results and Discussion

Experimental Studies

We succeeded to synthesize and recrystallize a stable trans-isomer of DMPSDA. Figure 1 shows the absorption, excitation and emission spectra of trans-DMPSDA in different solvents (CH, ACN and DMSO), as well as the photostationary fluorescence decay curves under constant-illumination conditions. Figure 2 demonstrates the fluorescence excitation and emission spectra of trans-DMPSDA in three solvents. Table 1 brings together all the spectroscopic properties of the molecule in these solvents at room temperature including absorption, fluorescence excitation and emission maxima ($\lambda_{\text{abs}}$, $\lambda_{\text{ex}}$, $\lambda_{\text{em}}$), fluorescence quantum yields ($\Phi_f$) and apparent fluorescence decay rate constants ($k_{\text{app}}$) for $N,N$-dimethyl-$N'$-picryl-4,4'-stilbenediamine in acetonitrile (ACN), dimethyl sulfoxide (DMSO) and cyclohexane (CH) at room temperature.

| Solvent | $\lambda_{\text{abs}}$ (nm) | $\lambda_{\text{ex}}$ (nm) | $\lambda_{\text{em}}$ (nm) | $\Phi_f$ | $k_{\text{app}}$ (s$^{-1}$) |
|---------|------------------|----------------|----------------|---------|------------------|
| ACN     | 360              | 360            | 420            | 0.009   | 0.643            |
| DMSO    | 365              | 365            | 425            | 0.013   | 0.614            |
| CH      | 350              | 350            | 390/410        | 0.001   | 0.614            |

The photostationary fluorescence decay of trans-DMPSDA is observed only in relatively polar solvents (ACN and DMSO) and originates from the non-radiative $1^t \rightarrow 1^p$ twisted transition, which is responsible for trans-cis isomerisation of the molecule and acts as a quenching funnel on fluorescence emission. CH is a non-polar solvent, and vibrational relaxation of the Franck-Condon state plays the primary role in its stabilisation of the excited state, thereby considerably slowing the $1^t \rightarrow 1^p$ transition [1, 2].

The measured fluorescence quantum yield values in all three solvents, shown in Table 1, are relatively low compared to the parent trans-4-dimethylamino-4’-aminostilbene [1–3].

Fig. 2 Fluorescence excitation and fluorescence emission spectra of trans-DMPSDA in three solvents at the different excitation and emission wavelengths: a CH, emission range 380–500 nm; excitation 275–350 nm; solid bold line – $\lambda_{\text{em}}=390$ nm, $\lambda_{\text{ex}}=320$ nm; b ACN, emission range 400–520 nm; excitation 300–400 nm; solid bold line – $\lambda_{\text{em}}=420$ nm, $\lambda_{\text{ex}}=360$ nm; and c DMSO, emission range 420–520 nm; excitation 300–400 nm; solid bold line – $\lambda_{\text{em}}=420$ nm, $\lambda_{\text{ex}}=360$ nm

Fig. 3 Stereoisomers 1E, 1E’, 1Z, 1Z’ of DMPSDA studied with the ab initio calculations
We assume that part of the additional non-radiative losses are caused by the electronic energy transfer onto the TNP fragment, which intrinsically shows negligible fluorescence, due to the presence of an efficient conical intersection of the $S_1$ state with the ground state $S_0$ \[10\].

**Theoretical Calculations**

Theoretical calculations were conducted with the following software suites: Gaussian 03 [11], Gaussian 09 [12] and Turbomole 6.1 [13]. The ground state studies were carried out by the density functional theory (DFT) using the Becke’s Three Parameter Hybrid Method and the correlation functional of Lee, Yang, and Parr (LYP) with the correlation potentials VWM (III) (calculated in Gaussian 03 or Gaussian 09) and VWM (V) (calculated in Turbomole 6.1). The optimized structures were controlled by frequency calculations with the vibration analysis using Hessian matrix. The excited states were calculated by the Time Dependent B3LYP hybrid method for the ground state optimized structures.

**Table 2** Absolute energy $E_h$, relative energy $\Delta E$, dipole moment $\mu_g$ and number of negative Hessian of the equilibrium structures for isomers 1E, 1Z and 1E’ calculated by B3LYP/6-31+G (3df,2pd)

| Isomers | $E_h$, Hartree | $\Delta E$, kcal/mol | $\mu_g$, D | $x$ |
|---------|---------------|----------------------|------------|-----|
| 1E      | $-1,574.7860481$ | 0                    | 10.5       | 0   |
| 1E’     | $-1,574.7860106$ | 0.024                | 10.3       | 0   |
| 1Z      | $-1,574.7780105$ | 5.04                 | 7.52       | 0   |
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