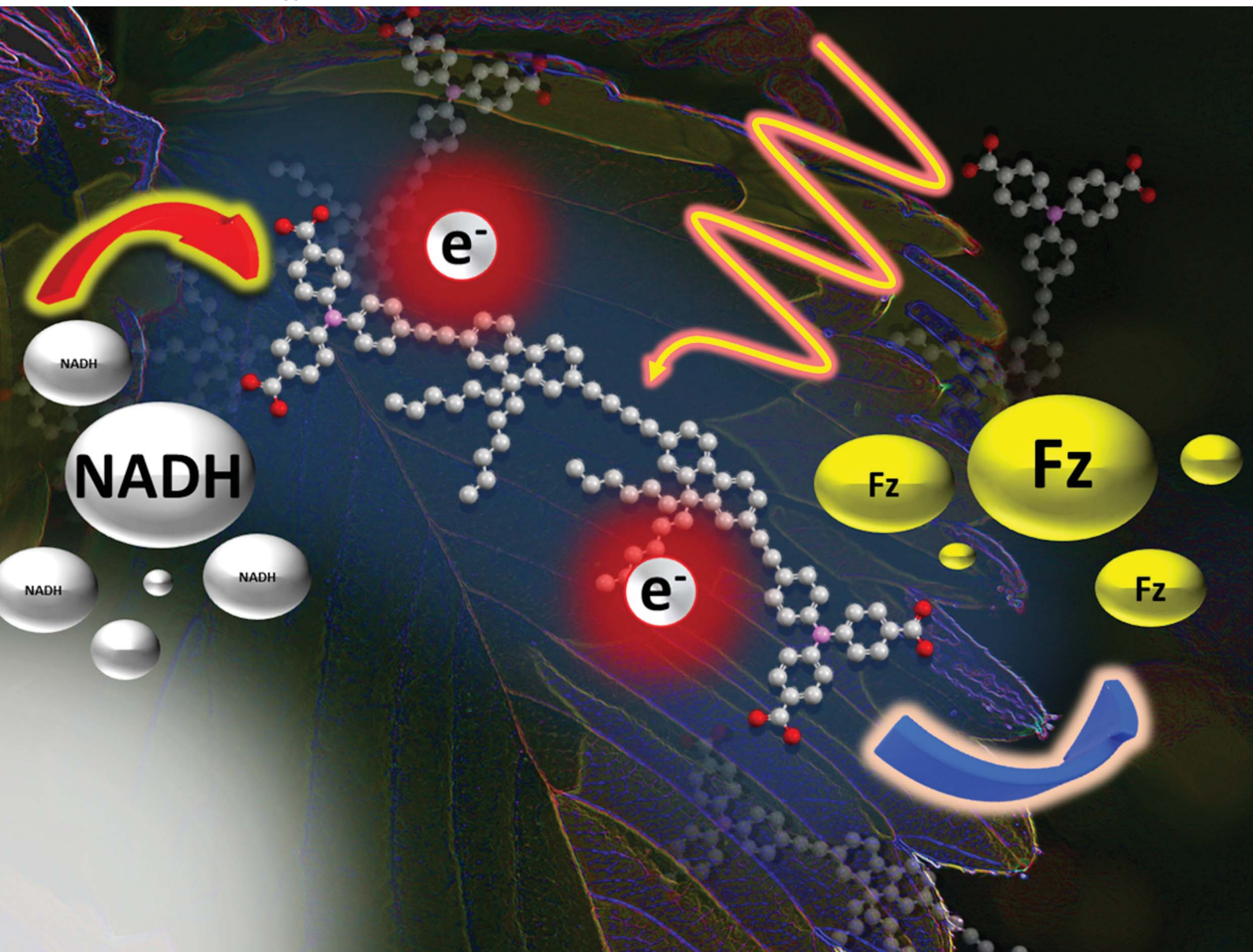


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Light-driven electron transfer in a lipid bilayer with mixed valence molecular wires†

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Triarylamines (TAAs) are one of the most important classes of redox-active organic compounds, which are readily available from modular synthesis, thereby offering the possibility to easily adjust their intrinsic redox potentials. We present herein two bis(triarylamines) (BTAA)s with π -extended 2,7-diethynylfluorene or 2,2'-(1,3-butadiyne-1,4-diyl)-bis(7-ethynylfluorene) bridges and two benzoic acid headgroups per TAA and their (formally) mixed-valent radical cations. Owing to their amphiphilic character and favorable redox properties, these BTAA)s are designed to serve as charge conduits through membranes. The lipid bilayer/BTAA systems are water-soluble, which allowed us to explore their photoactivity in aqueous solution and utilize their mixed valent form for membrane-mediated photoinduced electron transfer. Our findings will be relevant for constructing artificial nanoreactors for solar light energy conversion and light-driven redox chemistry in water.

Introduction

Light-driven long-range electron transfer is relevant for solar energy conversion in the natural photosynthesis of plants and photosynthetic bacteria, where photons drive the movement of electrons across biological membranes. Nature uses sophisticated assemblies of transmembrane protein complexes such as photosystem I and II (PSI and PSII) to convert NADP^+ into NADPH on one side of the membrane and water into O_2 on the other side of the membrane, as shown in Fig. 1. Previous artificial systems attempting to mimic light-driven transmembrane electron transfer across lipid bilayers employed embedded photosensitizers,^{1,2} along with electron carriers, to transfer electrons across membranes *via* diffusion of the photoreduced electron carrier.^{3–5} Additionally, there are some studies of light-driven transmembrane electron transfer using either sophisticated and synthetically challenging supramolecular barrels¹ or rigid oligoaromatic molecular wires with varying levels of

synthetic complexity.^{2,6} In the present study, we explore the concept of mixed valency, *i.e.* electron delocalization over two covalently connected redox centers,⁷ in a π -extended molecular wire for transmembrane electron transfer in lipid bilayers. These are multielectron processes that resemble the complexity of multielectron transfer in the photosynthesis system, which later on can be a future model for performing water-splitting and CO_2 reduction.

Triarylamines (TAAs) are one of the most important classes of redox-active organic compounds. One of their key advantages is their ready accessibility *via* modular synthesis, thereby offering the opportunity to attach one aryl substituent at a time and to generate derivatives with two or three different aryl substituents. This allows fine-tuning of their intrinsic redox potentials^{8–10} and the deliberate introduction of a broad range of chemical functionalities, *e.g.* for the construction of hybrids with two or more chemically different redox-active constituents^{11–21} or for rendering them water soluble.^{22–25} Once protected by substituents to prevent oxidatively induced coupling to benzidines^{26–29} or cyclization to carbazoles,³⁰ one-electron oxidized triarylammonium ions are rather stable, even towards water and air.^{9,22,31} Bridged bis(triarylamines) (BTAA)s, $\text{Ar}^1\text{Ar}^2\text{N-arylene-NAr}^1\text{Ar}^2$, typically undergo two or even more redox processes. Their associated mixed-valent radical cations generated by one-electron oxidation, *i.e.* the removal of an electron from only one of the available redox sites, are outstanding testbeds for probing the effects of the length, the chemical constitution and the conformation of the intervening π -conjugated (poly)arylene bridge and the chemical environment (solvent and counterions)^{11,32–35} on the extent of electronic coupling between the terminally appended redox sites.^{36–42}

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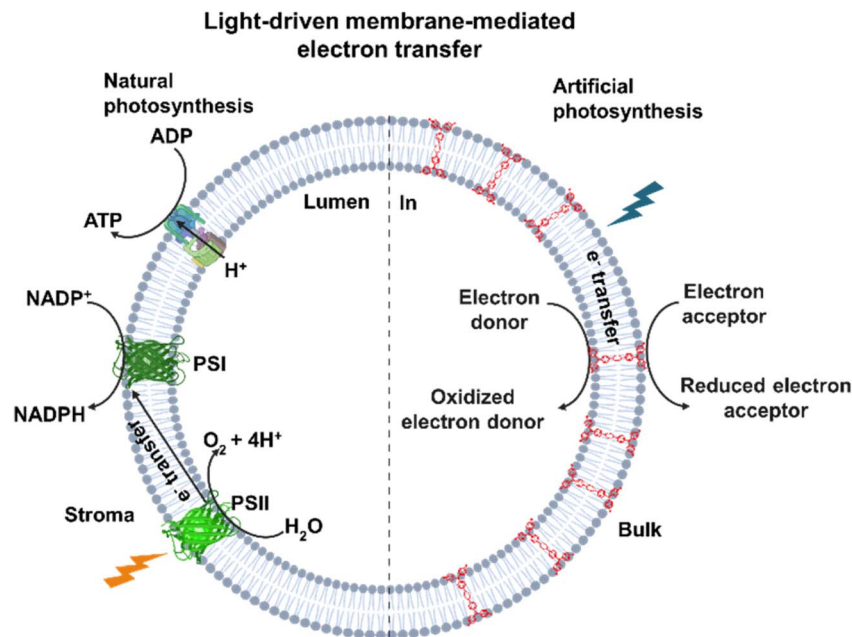


Fig. 1 Comparison of natural and artificial photosynthesis pathways. NADPH = nicotinamide adenine dinucleotide diphosphate, NADP^+ = nicotinamide adenine dinucleotide phosphate, PSII = photosystem II, PSI = photosystem I, ATP = adenosine triphosphate, ADP = adenosine diphosphate, and e^- = electron.

Their proven ability to facilitate efficient through-bond charge delocalization over long distances^{40,43–46} has also made them attractive for applications as single molecule wires in metal–molecule–metal nanojunctions. Recent studies on anchor group-modified bis(diarylamine)-capped oligophenylenes even revealed negative values of the attenuation factor β up to the terphenylene limit, implying that the electron transfer rate increases with increasing molecular length.⁴⁷ The advantageous properties of BTAAAs, *i.e.* their redox activity, modularity, and abundance, thus prompted us to utilize them as charge conduits through membranes. We here convey the results of our work on using BTAAAs with polar head groups for this purpose.

Results and discussion

Design and synthesis

The molecular wires designed here have hydrophobic cores based on fluorene and alkyne units bridging the BTAA redox centers, which are functionalized with carboxylic acids or esters. The carboxylic acids serve as hydrophilic head groups to anchor the molecular wire on both sides of the lipid bilayer so that they span across the lipid bilayer (Fig. 2a).

Fig. 2b summarizes the synthesis and chemical structures of the target BTAAAs. Our synthetic routes afford the methyl esters of the envisioned tetracarboxylic acids and involve cascades of different C–N and C–C cross-coupling reactions, including a Glaser–Hey–Eglinton coupling in the case of 2^{Me} .⁴⁸ Ester cleavage subsequently provided free tetracarboxylic acids. Detailed synthetic protocols and methods are provided in the ESI (Schemes S1 and S2[†]), along with ^1H , $^{13}\text{C}\{^1\text{H}\}$, mass spectra, and the absorption spectra and molar extinction coefficients ϵ

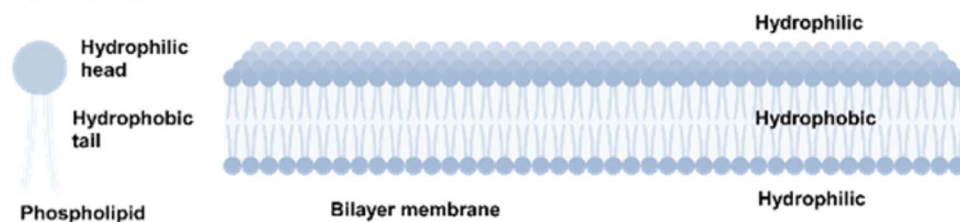
of the target BTAAAs (ESI, Fig. S16 and S17[†]). The molecular design with two polar benzoic acid headgroups at each terminus and nonpolar aliphatic hexyl tails at the core segments was conceived to mimic classical lipid bilayer properties. The carboxylic functionalities render BTAAAs 1^{H} and 2^{H} slightly soluble in aqueous media.

Electrochemistry and spectroscopy

The intended utilization of BTAAAs 1^{H} and 2^{H} as transmembrane redox mediators puts their redox properties into focus. However, voltammetric characterization of free acids 1^{H} and 2^{H} was hindered by the limited solubility of the protonated 1^{H} and 2^{H} in aqueous media, while an increase in pH led to poorly reproducible voltammograms with ill-defined redox waves at Pt, Au and glassy carbon electrodes. We therefore resorted to esters 1^{Me} and 2^{Me} in the $\text{CH}_2\text{Cl}_2/\text{NBu}_4^+ [\text{BAR}^{\text{F}24}]^-$ (0.1 M) electrolyte. Fig. 3 displays cyclic voltammograms of 1^{Me} (panel A) and 2^{Me} (panel B). The very weakly coordinating $[\text{BAR}^{\text{F}24}]^-$ anion ($[\text{BAR}^{\text{F}24}]^- = [\text{B}\{\text{C}_6\text{H}_3(\text{CF}_3)_2-3,5\}_4]^-$) is known for maximizing electrostatic repulsion, thereby maximizing the redox splitting $\Delta E_{1/2}$ between the individual redox couples and hence increasing the thermodynamic stability of the intermediate, mixed-valent radical cation with respect to disproportionation.^{49–51} The expected two oxidations of the TAA constituents nevertheless merge into a single composite wave with (average) half-wave potentials of $E_{1/2}(1^{\text{Me}}) = 675$ mV and $E_{1/2}(2^{\text{Me}}) = 670$ mV on the ferrocene/ferrocenium scale. Only in the case of the shorter 1^{Me} can a small splitting into two separate waves be discerned. Computational simulation of the experimental voltammograms yielded a redox splitting of 80 mV in the case of 1^{Me} and an even smaller value of only 45 mV for



a) Amphiphilic structure of bilayer membrane



b) Design of amphiphilic chromophores

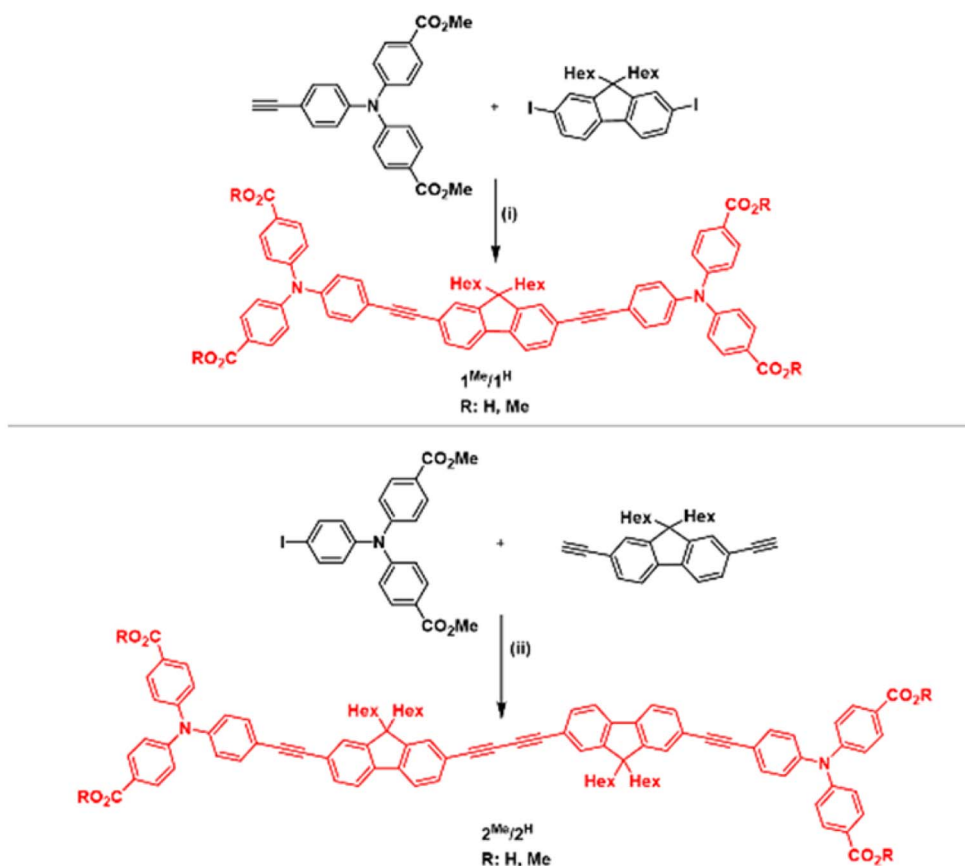


Fig. 2 (a) The structure of a bilayer membrane, built from an amphiphilic phospholipid. (b) Chemical structures and schematic syntheses of target bis(triarylamine) tetracarboxylic acids/methyl esters $1^{\text{H/Me}}$ and $2^{\text{H/Me}}$. (i) $\text{PdCl}_2(\text{PPh}_3)_2$, NEt_3 , CuI , THF , 70°C , and overnight; $1^{\text{Me}} \rightarrow 1^{\text{H}}$: $\text{THF}/\text{H}_2\text{O}$ (2 : 1), LiOH , r.t., and overnight; (ii) $\text{PdCl}_2(\text{PPh}_3)_2$, NEt_3 , CuI , THF , 70°C , and two days; $2^{\text{Me}} \rightarrow 2^{\text{H}}$: $\text{THF}/\text{H}_2\text{O}$ (2 : 1), LiOH , r.t., and overnight.

2^{Me} with roughly twice the spacer length. Simulated cyclic voltammograms are shown as blue circles in panels A (1^{Me}) and B (2^{Me}) of Fig. 3, superimposed with the experimental data. Details of the used software and the fitting procedures are given in the ESI,[†] and the parameters obtained by the simulations are provided in Table S4 of the ESI.[†] The half-wave potential splitting of 2 is barely larger than the statistical contribution of $2 \ln(2RT/F) = 36 \text{ mV}$ to the free energy change for the stepwise oxidations of two non-interacting redox sites.⁵² Similar small half-wave potential separations were also reported for other BTAAAs with large spatial extensions of the connecting linker (Table 1).^{39,44,53–55}

The close proximity of the two redox waves makes it difficult to access and spectroscopically characterize and discriminate

between the one- and two-electron-oxidized radical cations and dications as individual species, which is important in order to elucidate the effective oxidation state when using the present BTAAAs as transmembrane redox mediators.

The half-wave potentials of 1^{Me} and 2^{Me} are appreciably more positive than that of the triarylamine-derived mono- and dicarboxylic acids $(4\text{-MeO-C}_6\text{H}_4)_2\text{N}(\text{C}_6\text{H}_4\text{-COOH-4})$ ($E_{1/2} = 170 \text{ mV}$) and $(4\text{-MeOC}_6\text{H}_4)_2\text{N}(\text{C}_6\text{H}_3(\text{COOH})_2\text{-3,5})$ ($E_{1/2} = 90 \text{ mV}$) reported by Lahti *et al.*,²² which we ascribe to the absence of methoxy donor substituents (literature values were recalculated to the ferrocene/ferrocenium scale using the reported half-wave potential of the $\text{Cp}_2\text{Fe}^{0/+}$ redox couple of 553 mV in CHCl_3 on the Ag/AgCl scale,⁵⁶ *i.e.* the conditions used by these authors). They are nevertheless *ca.* 150 mV lower than that of the 4-



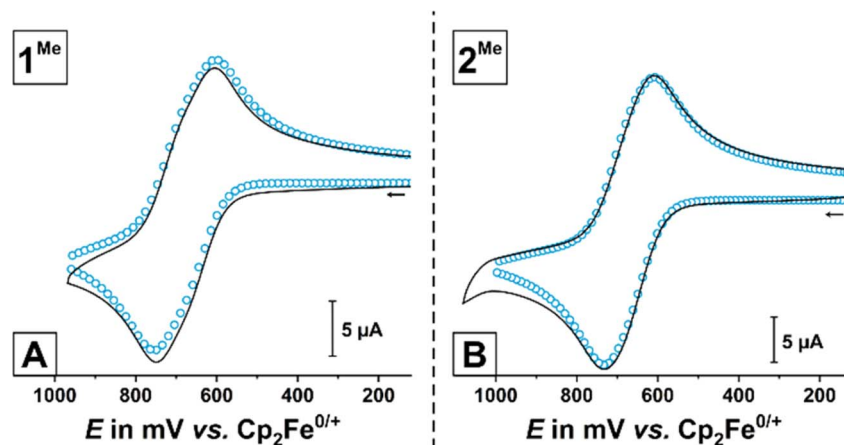


Fig. 3 (A) Overlaid cyclic voltammogram of 1^{Me} in $\text{CH}_2\text{Cl}_2/\text{NBu}_4^+ [\text{BAR}^{\text{F}24}]^-$ (0.1 M, $T = 293 \text{ K} (\pm 3 \text{ K})$, $\nu = 600 \text{ mV s}^{-1}$, black solid line) and corresponding simulation (blue circles). (B) Overlaid cyclic voltammogram of 2^{Me} in $\text{CH}_2\text{Cl}_2/\text{NBu}_4^+ [\text{BAR}^{\text{F}24}]^-$ (0.1 M, $T = 293 \text{ K} (\pm 3 \text{ K})$, $\nu = 600 \text{ mV s}^{-1}$, black solid line) and corresponding simulation (blue circles).

Table 1 Electrochemical potentials of the oxidations of 1^{Me} and 2^{Me} obtained from the cyclic voltammograms in $\text{CH}_2\text{Cl}_2/\text{NBu}_4^+ [\text{BAR}^{\text{F}24}]^-$ (0.1 M, $T = 293 \text{ K} (\pm 3 \text{ K})$) as well as by digital simulation

	$E_{1/2}^{\text{avg } 0/2+}$ (exp.)	$E_{1/2}^{0/+}$ (simulated)	$E_{1/2}^{+/2+}$ (simulated)	$\Delta E_{1/2}$
1^{Me}	675 mV	635 mV	715 mV	80 mV
2^{Me}	670 mV	640 mV	685 mV	45 mV

ethynylphenyl precursor ($E_{1/2} = 820 \text{ mV}$ in CH_2Cl_2 vs. $\text{Cp}_2\text{Fe}^{0/+}$), indicating the extent of electron donation from the dihexyl-substituted fluorenyl core unit.

We next turn to the spectroscopic characterization of the oxidized forms of 1^{Me} and 2^{Me} as models of the oxidized BTAA tetracarboxylic acids that one expects to be the active species in transmembrane hole transfer. To generate these species, we applied UV-Vis/NIR spectroelectrochemistry (SEC), *i.e.* spectroscopic monitoring during *in situ* electrochemical oxidation upon incrementally increasing the applied potential to a value 250 mV positive of the convoluted overall two-electron wave. The latter experiments were conducted inside an optically transparent thin-layer electrolysis cell⁵² with 1,2-dichloroethane (DCE) as the solvent, using again the $\text{NBu}_4^+ [\text{BAR}^{\text{F}24}]^-$ (0.1 M) supporting electrolyte. The higher boiling point of the DCE solvent as compared to CH_2Cl_2 counteracts solvent evaporation in the vicinity of the working electrode during the experiments. Fig. 4 displays the evolution of the UV-Vis/NIR spectra alongside NIR spectra recorded with an IR/NIR spectrometer (for details, see the ESI, Section S3†) as insets; the latter has a superior sensitivity in the low-energy part of the NIR regime. Panels A and B of Fig. 4 show the results on 1^{Me} , while those for 2^{Me} are given in panels C and D.

In both cases, the overall two-electron oxidation proceeds in separate steps *via* an intermediate species (orange lines), which has a characteristic broad NIR band at 1405 nm ($1^{\text{Me}\cdot+}$) or 1400 nm respectively ($2^{\text{Me}\cdot+}$, see panels A and C of Fig. 4 and the corresponding insets). As the oxidation progresses further, the NIR band intensifies and shifts to higher energies (*i.e.* lower

wavelengths) of 1250 nm for both $1^{\text{Me}2+}$ and $2^{\text{Me}2+}$. These observations are indicative of the stepwise formation of first the mixed-valent radical cations and then the dication of BTAAAs with large, π -extended linkers.^{14,39,53,57} Other spectroscopic changes are the gradual and continuous growth of a Vis/NIR band, which is located at 715 nm for 1^{Me} and at 705 nm for 2^{Me} . This band corresponds to a π - π^* transition of a triarylammonium radical cation and is characteristic of the $\text{TAA}^{\cdot+}$ motif. This assignment agrees with the observed rough doubling in intensity upon the second oxidation, which generates an additional triarylammonium chromophore. Also, both oxidized compounds feature one additional band of similar intensity at 550 nm each.

In interpreting the results, one should notice that the close proximity of the individual $E_{1/2}$ values of 1^{Me} and 2^{Me} implies that the intermediate mixed-valent radical cations have only limited stability with respect to disproportionation to the neutral starting compounds and the dioxidized forms, both with isovalent TAA redox sites. This is expressed by the only modest values for the comproportionation equilibrium constant K_c for the comproportionation (*cf.* eqn (1)) of *ca.* 23 (1^{Me}) or only 6 (2^{Me}).

$$K_c = \exp\left(\frac{F}{RT}(\Delta E_{1/2})\right) \quad (1)$$

In eqn (1), R is the universal gas constant, F Faraday's constant, and $\Delta E_{1/2}$ the half-wave potential separation between the two consecutive redox processes. The individual K_c values indicate that a maximum of 70% or 55% of the mixed-valent radical cations are formed as such after the passage of one equivalent of charge, with the remaining 30% or 45% present as equimolar mixtures of the neutral and dioxidized forms. The overlapping NIR bands of the radical cations and the dication in the NIR unfortunately preclude us from deconstructing the spectra and calculating the spectra of the pristine, intermediate radical cations, but it is nevertheless clear that both species



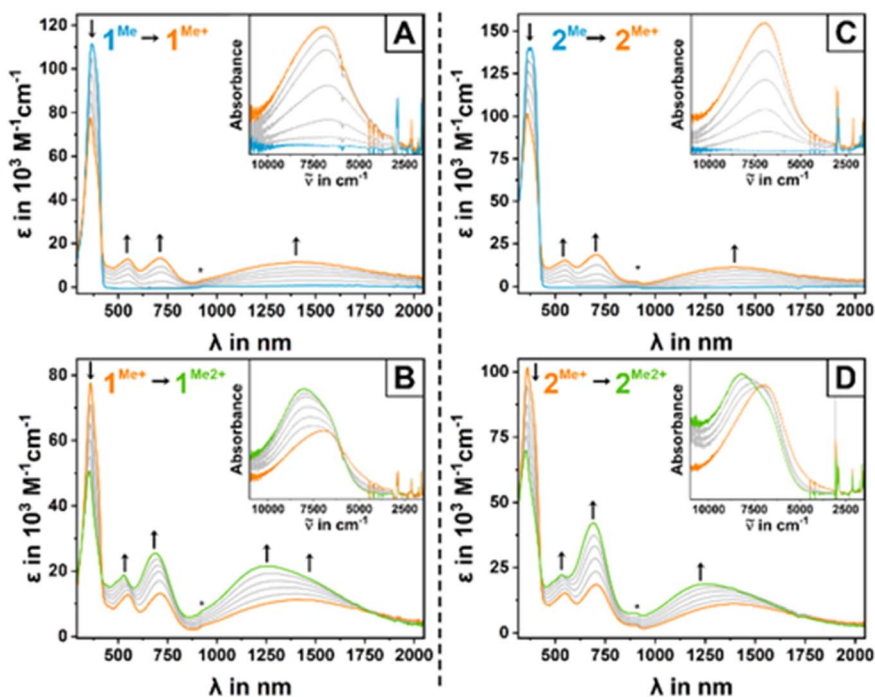


Fig. 4 UV-Vis/NIR spectroelectrochemistry of 1^{Me} (left, panels A and B) and 2^{Me} (right, panels C and D) with the first oxidation shown as blue to orange curves (top, panels A and C) and the second oxidation shown as orange to green curves (bottom, panels B and D), respectively, in CH_2Cl_2 with $0.1 \text{ M NBu}_4^+ [\text{BAR}^{\text{F}24}]^-$ as the supporting electrolyte. IR/NIR spectra are provided in insets in panels A to D. An artifact due to the experimental setup is marked with an asterisk.

have distinct absorptions in the NIR region. The high intensity of the NIR absorption of the radical cations is at first surprising when considering the small half-wave potential splittings and the large $\text{N}\cdots\text{N}$ distances of $23.15 \text{ \AA}$ in $1^{\text{Me}\cdot+}$ or $36.65 \text{ \AA}$ in $2^{\text{Me}\cdot+}$, as obtained from quantum chemical calculations (see below). The intensity gain and the blue-shift upon the second oxidation let us assign the prominent NIR band to charge-transfer (CT) excitation from the respective fluorene linker to the TAA $^{+\cdot}$ acceptor(s) rather than to a pure $\text{TAA} \rightarrow \text{TAA}^{+\cdot}$ intervalence charge-transfer (IVCT) excitation, as the latter would be expected to be specific for the mixed-valent radical cations and to vanish as the oxidation proceeds further to the dications.

To shed more light on the electronic structures of one- and two-electron oxidized $1^{\text{Me}+/2+}$ and $2^{\text{Me}+/2+}$, we resorted to density functional quantum-chemical calculations at the B3LYP+35% exact Hartree–Fock exchange def2SVP level of theory. The latter has proven to provide adequate descriptions of the electronic structures of mixed-valent BTAA's by compensating for the inherent tendency of the DFT method to delocalize charges in extended π -conjugated systems.^{37,58–60} Our calculations employed slightly truncated models with methyl instead of hexyl substituents, which are henceforth denoted as $1^{\text{Me}'}$ and $2^{\text{Me}'}$. Solvent effects were considered within the framework of the polarizable continuum model, applying the standard parameters for CH_2Cl_2 . Further details as well as coordinates of the geometry-optimized structures are provided in Section S11 of the ESI.[†]

Our calculations indeed provided intense transitions in the NIR for both the one- and the two-electron oxidized forms

$1^{\text{Me}'\cdot+/2}$ and $2^{\text{Me}'\cdot+/2+}$, at energies close to the experimental values. Charge density losses on the individual TAA $^+$ sites and the fluorenyl segments of the linkers were evaluated by natural bond orbital (NBO) analysis (see Fig. 5).⁶¹ According to these results, both radical cations possess localized electronic structures where the unipositive charge is confined to one specific TAA $^{+\cdot}$ entity. In contrast to other BTAA's with electron-rich *para*-anisyl (An) substituents NaN_2 , the most electron-rich phenyl group of 1^{Me} and 2^{Me} is the one bonded to the fluorenyl linker. This causes the occupied frontier MOs to spread onto the linker rather than being strictly confined to the periphery.

The calculated NIR transitions in radical cation $2^{\text{Me}'\cdot+}$ are depicted in panel A of Fig. 6, along with graphical representations of the involved molecular orbitals and the corresponding electron density difference maps (EDDMs; blue color = electron density loss and red color = electron density gain). Computational results for its shorter homolog $1^{\text{Me}'\cdot+}$ can be found in Fig. S18 of the ESI.[†] For both compounds, our calculations predict two separate transitions at low energies as highly intense bands ($1^{\text{Me}'\cdot+}$: $\lambda_{\text{calc}} = 1036 \text{ nm}/1675 \text{ nm}$, $f = 0.28/1.07$; $2^{\text{Me}'\cdot+}$: $\lambda_{\text{calc}} = 1212 \text{ nm}/1550 \text{ nm}$, $f = 0.46/0.84$). They both target β -LUSO, which is localized on the oxidized TAA unit, and emanate from the first three occupied orbitals of the β -manifold, *i.e.* β -HOSO to β -HOSO-2. The latter donor MOs are either localized on the other TAA unit or spread over the entire π -conjugated backbone, including the ester-functionalized diarylamine entities. Both NIR excitations therefore assume bridge-to-TAA $^+$ CT, augmented with IVCT contributions. The

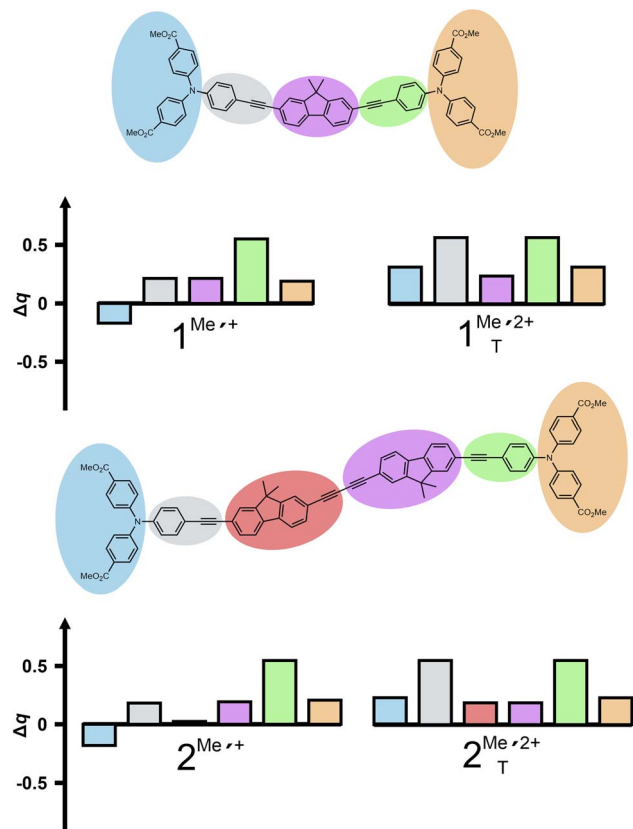


Fig. 5 Calculated electron density losses upon oxidation of model complexes $1^{\text{Me}'}$ and $2^{\text{Me}'}$ to their corresponding one- (left) and two-electron oxidized forms (triplet state), based on natural bond orbital (NBO) analysis. Δq refers to the charge difference of the respective molecular region of the radical monocations (left panels) and the respective dications in their triplet ground state (right panels) relative to the corresponding neutral state. The individual fragments are color-coded according to the graphical representation in the top panels.

absorption bands of one-electron oxidized $1^{\text{Me}'+}$ and $2^{\text{Me}'+}$ generally agree well with the results of investigations on other BTAA's with similarly π -extended linkers.^{39,44,54,55}

In contrast to the plethora of BTAA's with shorter linkers, but in line with important bridge-to-TAA⁺ contributions,^{39,54,55} the NIR transitions of $1^{\text{Me}'+}$ and $2^{\text{Me}'+}$ intensify during further oxidation to the dications while experiencing a shift to higher energies and shorter wavelengths respectively, from 1405 nm to 1250 nm for $1^{\text{Me}'+/2+}$ ($\lambda_{\text{calc}} = 1183 \text{ nm}$, $f = 1.82$), and from 1400 nm to 1220 nm for $2^{\text{Me}'+/2+}$ ($\lambda_{\text{calc}} = 1345 \text{ nm}$, $f = 2.39$); see panels A and B of Fig. 6. The dications were computed to prefer the triplet state by 54.0 kJ mol⁻¹ ($1^{\text{Me}'2+}$) or 42.5 kJ mol⁻¹ ($2^{\text{Me}'2+}$) over the closed and open-shell singlet states, which were calculated to be energetically degenerate. Panels A, B in Fig. 6 and S18 of the ESI† demonstrate the excellent agreement between computed and experimental spectra. A complete collection of all computed electronic absorption bands of $1^{\text{Me}'n+}$ and $2^{\text{Me}'n+}$ ($n = 0, 1, 2$, the latter in their triplet ground and excited singlet states) along with the corresponding TD-DFT calculated transitions on their methyl-substituted models, contour plots of the contributing orbitals, and computed

oscillator strengths are provided in the ESI (see Table S5†). On oxidation of the second TAA entity, the character of the NIR electronic excitation assumes pure bridge-to-TAA⁺ CT character.

In summary, the highly intense NIR absorptions of one- and two-electron oxidized BTAA's $1^{\text{Me}'+/2+}$ and $2^{\text{Me}'+/2+}$ can be traced to bridge-to-TAA⁺ CT augmented by TAA-to-TAA⁺ IVCT excitations in the radical cations. The energetically low-lying bridge-to-TAA⁺ excitations and the bridge participation in the frontier MOs make $1^{\text{Me}'+}$ and $2^{\text{Me}'+}$ (and by inference the one-electron oxidized free carboxylic acids $1^{\text{H}'+}$ and $2^{\text{H}'+}$) interesting candidates for transmembrane electron transfer when incorporated into a lipid bilayer.

Membrane integration

Incorporation of BTAA's in the lipid membrane bilayers was performed with 1^{H} or 2^{H} . At neutral pH, their molecular structures mimic the amphiphilicity of the phospholipid bilayer best by having charged carboxylates at each end of the linear and rigid molecular wire. The charged ends interact with the polar part of the membrane at the water interface, while the entire linker, including the inner phenyl rings, stays inside the hydrophobic part of the membrane. This secures the integration of 1^{H} or 2^{H} into the membrane (Fig. 7) of vesicles. In this study, we used sub-micrometer-sized liposome vesicles for all spectroscopy and electron transfer studies. We also prepared micrometer-sized giant vesicles for microscopic visualization. All vesicles were prepared with lipid bilayers based on 2-dipalmitoyl-*sn*-glycero-3-phosphocholine (DPPC) because DPPC bilayers are known to be impermeable to larger water-soluble molecules at room temperature.^{62,63} Along with the main phospholipid DPPC, we added 1 mol% of 14:0 PEG2000 PE to the lipid bilayer membranes to increase the stability of the liposome, as this lipid creates steric bulk and thereby prevents aggregation of the vesicles (see Fig. 7 for the structures of DPPC and 14:0 PEG2000 PE).^{5,64,65}

Molecular dynamics (MD) simulations of both molecular wires after 100 ns simulation time are shown in Fig. 7a and b. The MD simulations were done using YASARA and the AMBER force field;^{66,67} the details can be found in ESI S4, Fig. S19 and S20.† These MD simulations suggest that both molecular wires 1^{H} and 2^{H} assemble with the lipid bilayer, but 1^{H} assembles at the membrane–water interface, while 2^{H} can align in a trans-membrane fashion. We explain these results as follows: in 2^{H} , the N...N distance is 33.66 Å in the phospholipid bilayer (similar to the N...N distance of 36.65 Å in $2^{\text{Me}'+}$ determined by DFT, see above). The hydrophilic carboxylate groups are even further apart from each other and can therefore easily span a lipid bilayer membrane, being exposed to the membrane's water interfaces, as can be seen in Fig. 7b. Lipid bilayer membranes typically have a thickness of 30 to 50 Å.^{68,69} For the shorter BTAA 1^{H} , the N...N distance is 23.48 Å (similar to the N...N distance of 23.15 Å in $1^{\text{Me}'+}$ determined by DFT, see above). The distance of the hydrophilic carboxylic groups in 1^{H} is 28.51 Å, which is too short to span across the membrane and which can lead to the disruption of the membrane by 1^{H} .

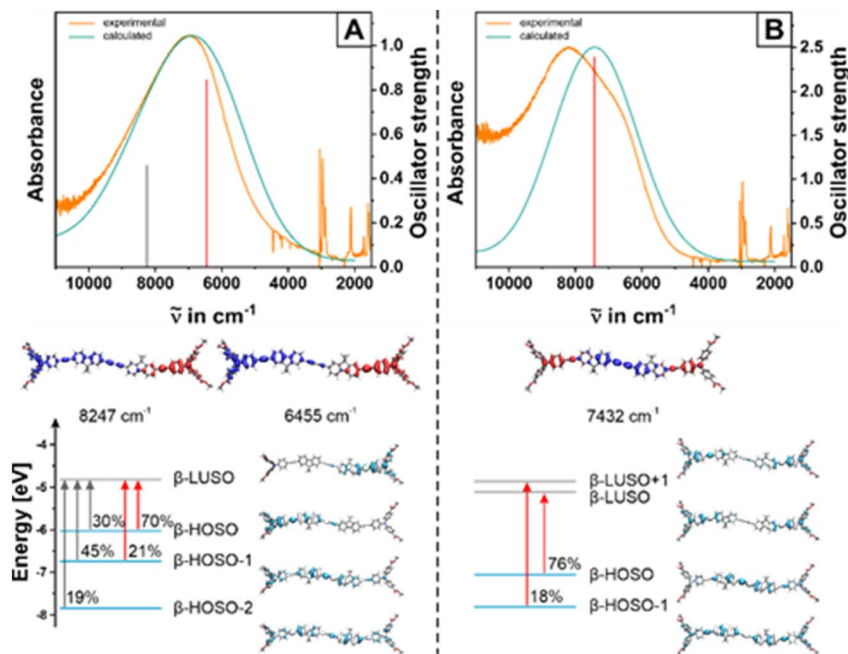


Fig. 6 Top: Comparison between experimental NIR spectra of $2^{\text{Me}+}$ (orange line in panel A) and $2^{\text{Me}2+}$ (green line in panel B) in 1,2- CH_2Cl_2 (0.1 M NBu_4^+ $[\text{BAR}^{\text{F}24}]^-$ as the supporting electrolyte) and TD-DFT calculated electronic spectra of model compound $2^{\text{Me}+}$ (panel A, turquoise line) and $2^{\text{Me}2+}$ (panel B, violet). Computed transitions are provided as red and grey vertical bars. Bottom: Contour diagrams of the acceptor and donor MOs involved in the NIR transitions together with their contributions to the respective transition and corresponding electron density difference maps (EDDMs). Blue color indicates electron density loss and red color an increase in electron density. Isosurface values for the MO plots are set as ± 0.02 .

In order to study how the respective molecular wire affects the membrane, we conducted a calcein leakage test using liposomes. In this test, we quantified the amount of calcein, a fluorescent dye, leaked out of the liposomes into the surrounding solution. For detailed information, please refer to ESI S5, Fig. S21.^{†70,71} We tested both the modified and unmodified DPPC liposomes for three days with and without irradiation using a 470 nm LED. The measurements with irradiation revealed that the modified DPPC liposome with 1^{H} released 93% of calcein, while the modified DPPC liposome with 2^{H} only released 4.5% of calcein. These observations align with our MD simulations, which predicted that the modified liposome with 1^{H} would disrupt the membrane more effectively than 2^{H} ; as 2^{H} is longer in size, it has a higher possibility of aligning in the transmembrane position in the membrane and would not disrupt the membrane.

UV-Vis absorption and emission spectroscopy were used to characterize the photophysical properties of 1^{H} and 2^{H} before and after membrane integration (Fig. 8a and b and Table 2). When measured in a mixture of DCM and MeOH (1 : 1) (Fig. 8a and b, solid black line), 1^{H} and 2^{H} showed absorption bands at 366 nm and 385 nm, respectively. This indicates that the respective LUMO of 2^{H} is lower in energy than that of 1^{H} due to its extended π -system. The absorption spectra shift to the blue for both structures in the membrane environment as can be seen in Fig. 8a and b and in Table 2. While 1^{H} experienced only a 4 nm blue shift, the absorbance of 2^{H} experienced a pronounced blue-shift by ~ 20 nm. The very minor blue-shift

for 1^{H} might be explained by Tyndall scattering of the liposome suspension, similar to other chromophores in liposomes with minor solvent-dependent absorbance.⁷⁵ The very pronounced blue-shift of 2^{H} is likely due to the lower polarity and rigidity around the compound in a lipid bilayer environment.^{72–74} This observation is in line with the MD simulation, which showed that the entire extended π -system of 2^{H} is located in the lipophilic part of the membrane, which leads to destabilizing the LUMO of 2^{H} , resulting in a more pronounced blue shift in the UV-Vis absorption.

The emission spectra of 1^{H} and 2^{H} have their maximum at 422 nm and 430 nm respectively in $\text{DCM}:\text{CH}_2\text{Cl}_2$ (1 : 1), as indicated by the solid green line in Fig. 8a and b. Both compounds have a shoulder at a higher wavelength in their emission spectra, suggesting the existence of unresolved vibrational energy levels in both compounds. Upon incorporation into the membrane, the maximum is slightly red-shifted by 13 nm and 15 nm for 1^{H} and 2^{H} respectively. However, this apparent red shift might also be partly induced by the broadened absorption and respective inner filter effects (Fig. 8a and b, dashed green line).

Furthermore, to show that 2^{H} is successfully integrated into the lipid membrane, we prepared a giant multilamellar vesicle of 2^{H} with the same methodology as that used for preparing the liposomes and performed confocal microscopy. We excited the giant multilamellar vesicle with a laser at $\lambda = 405$ nm and detected the fluorescence of the vesicle in the region of 420–650 nm. A double half-moon-shaped emission profile was



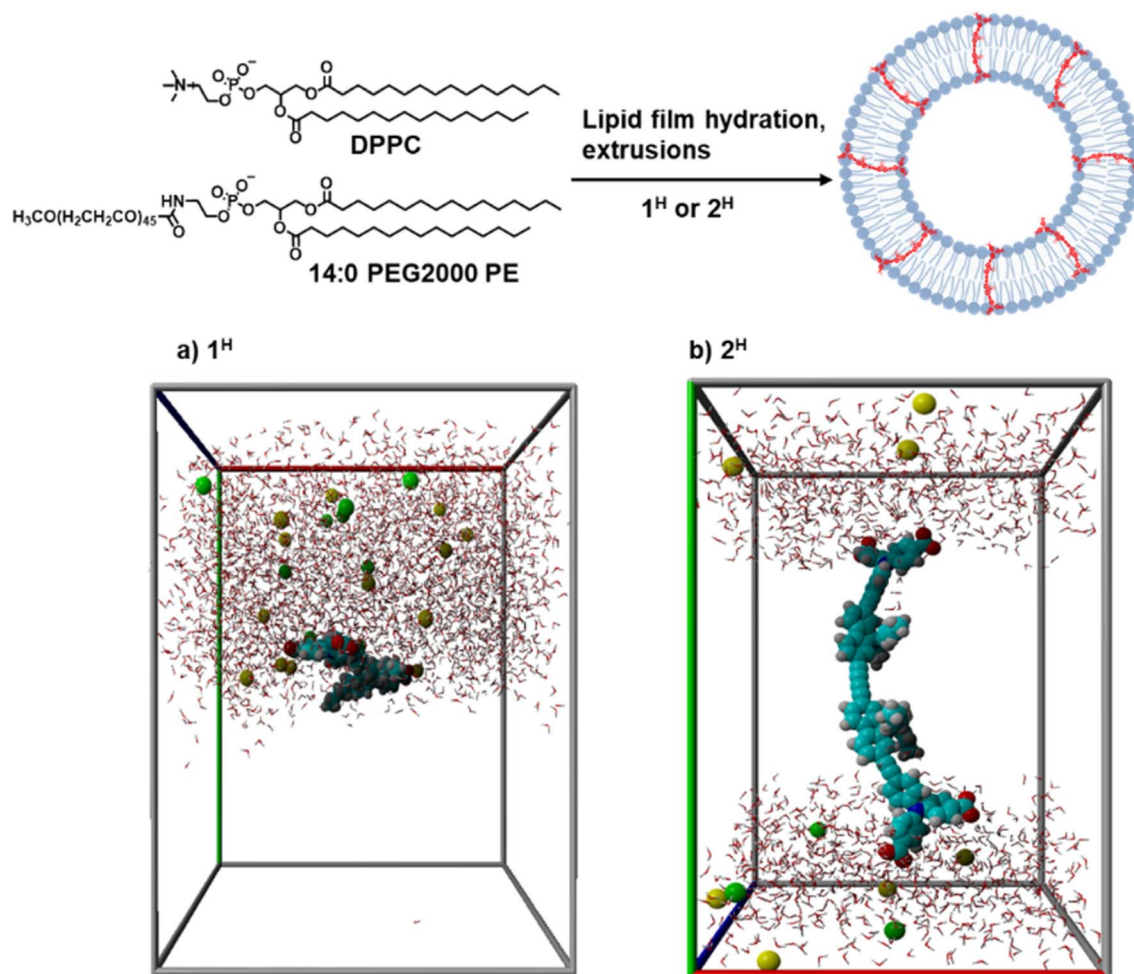


Fig. 7 Integration of 1^H or 2^H into a DPPC (1,2-dipalmitoyl-*sn*-glycero-3-phosphocholine) liposome and a molecular dynamic simulation snapshot of (a) 1^H and (b) 2^H in DPPC after 100 ns simulation time. The lipid bilayer membrane is omitted for clarity.

observed in the microscopic image (Fig. 8c), typical for molecules with a preferred alignment in the lipid membrane.^{65,72,75} It shall also be noted that, apart from the half-moon effect, the molecules are overall evenly distributed within the membrane and no larger macroscopic phase separation can be observed.⁷⁶

Electron transfer at the membrane

To study the photoactive properties of 1^H and 2^H toward electron transfer across the lipid bilayer, we started our studies on photoinduced transmembrane electron transfer by preparing a DPPC liposome solution containing 1 mol% of 1^H or 2^H in the

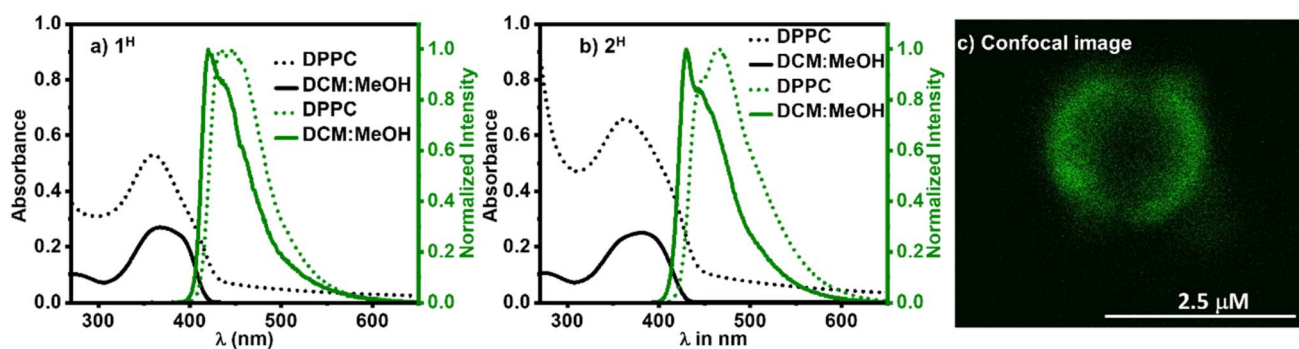


Fig. 8 (a) UV-Vis and emission spectra of 1^H and (b) UV-Vis and emission spectra of 2^H in a mixture of dichloromethane (DCM) : methanol (MeOH) (1 : 1) and in DPPC liposomes with a composition of DPPC : (14:0 PEG2000 PE) : 1^H or 2^H = 100 : 1 : 1, in phosphate buffer at pH = 7.0; (c) confocal microscopy image of 2^H in a giant vesicle with a composition of 100 : 1 : 1 of DPPC : 14:0 PEG2000PE : 2^H in phosphate buffer at pH = 7.0 after size exclusion chromatography, upon fixation in agarose hydrogel.



Table 2 Spectroscopic data of 1^H and 2^H

	Condition	λ_{Abs} (nm)	λ_{Em} (nm)
1^H	DCM:MeOH	366	422
	DPPC ^a	362	435, 446
2^H	DCM:MeOH	385	430
	DPPC ^a	365	445, 467

^a UV-Vis and emission spectra of 1^H and 2^H in DPPC liposomes with a composition of DPPC:(14:0 PEG2000 PE): 1^H or 2^H = 100:1:1, in phosphate buffer at pH = 7.0.

membrane. The water-soluble electron donor NADH was encapsulated, and the water-soluble electron acceptor (2,3-bis-(2-methoxy-4-nitro-5-sulphophenyl)-2H-tetrazolium-5-carboxanilide) (XTT) was added to the bulk solution of the liposome, yielding a composition of NADH/ 1^H /XTT or NADH/ 2^H /XTT (Fig. 9a). A similar system was also prepared with encapsulated XTT and NADH at the bulk solution of liposomes, yielding a composition of XTT/ 2^H /NADH. NADH was chosen due to its water solubility and favorable absorption and emission properties.⁷⁷ XTT was selected because it is a well-known compound to quantify intracellular reducing agents in biological assays.⁷⁸ The details of sample preparation can be found in

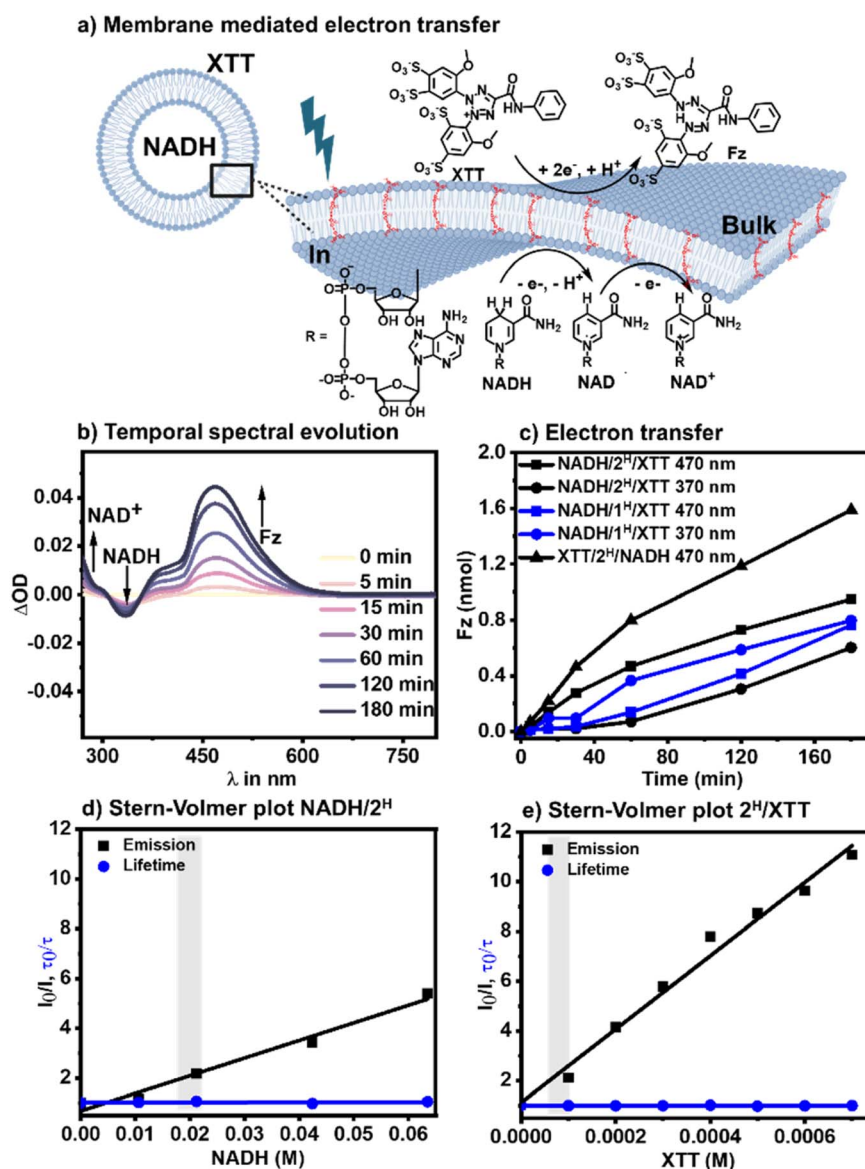


Fig. 9 (a) Schematic illustration of the targeted photoinduced electron transfer at the lipid bilayer with encapsulation of NADH, integration of the chromophore, and presence of XTT in the bulk solution, (b) subtracted temporal evolution of UV-Vis spectra from $t = 0$ min with sample composition NADH/ 2^H /XTT, (c) number of Fz molecules formed under irradiation with 460 nm and 370 nm light sources over time for NADH/ 1^H /XTT, NADH/ 2^H /XTT, and XTT/ 2^H /NADH, (d) Stern–Volmer plots of liposome samples with a composition of 100:1:1 of DPPC:14:0 PEG2000 PE: 2^H with different concentrations of encapsulated NADH, and (e) Stern–Volmer plots of liposome samples with a composition of 100:1:1 of DPPC:14:0 PEG2000 PE: 2^H with different concentrations of XTT in bulk solution. All samples were prepared with 10 mM phosphate buffer at pH 7.0 under ambient conditions. The grey bar corresponds to the used concentration in photoinduced electron transfer experiments.

the ESI Section S6 and Fig. S22.† Based on dynamic light scattering, the liposomes measure 130–140 nm in diameter for all types of samples (the empty liposome with only 1^H or 2^H integrated and with the encapsulated NADH or XTT; see ESI Section S7 and Fig. S23†).

Upon light irradiation, we observed a gradual increase of the absorption band at $\lambda_{\max} = 470$ nm, corresponding to the formation of formazan (Fz), the reduced form of XTT. At the same time, the NAD^+ absorption band at 260 nm increased, the latter corresponding to the oxidized form of NADH and indicating that photoinduced electron transfer had occurred (Fig. 9b). The extent of electron transfer was quantified *via* the intensity of the Fz absorption band (Fig. 9c).

We chose 370 nm and 470 nm LED light sources to irradiate the assembled liposome systems NADH/ 1^H /XTT and NADH/ 2^H /XTT. The applied LED light sources have a full-width at half maximum (FWHM) of ± 24 nm at $\lambda_{\max} = 373$ nm for the 370 nm LED and FWHM ± 22 nm at $\lambda_{\max} = 464$ nm for the 470 nm LED light source. This leads to the excitation of 1^H and 2^H at either the absorption maximum or in the tail of the absorption (see ESI S8, Fig. S25†). Photoexcitation with the 470 nm LED excited the molecular wire only and excluded the simultaneous excitation of NADH, which has an absorption maximum at 340 nm.^{77,79} In the case of the 370 nm LED, both the molecular wire and NADH are simultaneously excited.

Based on the extinction coefficient of Fz, it was calculated in the case of NADH/ 2^H /XTT that 1.00 nmol of Fz formed after three hours with the 470 nm LED irradiation. This corresponds to 4.4 electron transfers per molecule of 2^H . In the case of 2^H with a 370 nm LED, 0.71 nmol of Fz was formed, which is equal to 1.4 nmol of electrons and 3.1 electron transfers per 2^H molecule. This lower performance by the 370 nm light source may be due to the simultaneous absorption of light by both the molecular wire and NADH, as the NADH also absorbs some of the light at 370 nm (see ESI S8, Fig. S25†). In the case of NADH/ 1^H /XTT and after three hours of irradiation with a 470 nm or 370 nm LED, the Fz yield is 0.77 nmol and 0.78 nmol, respectively. The number of Fz is similarly low with both light sources for 1^H , probably due to the lower spectral overlaps of the absorbance of 1^H with the 470 nm LED compared to 2^H (see ESI S8, Fig. S25†). In a previous study, we demonstrated that the light-driven conversion of the water-soluble substrate can be significantly accelerated within the inner aqueous liposome compartment compared to the aqueous bulk.⁷⁷ We therefore tested the reverse location of the XTT and NADH and generated XTT/ 2^H /NADH liposomes. In this system, we observed that by compartmentalization of XTT instead of NADH, we could almost double the production of Fz within three hours, yielding 1.80 nmol, which corresponds to 3.6 nmol electron transfer and 8 electrons transferred per molecule of 2^H . This observation indicates that electron transfer from XTT is a limitation that can be minimized through encapsulation of XTT within the inner aqueous compartment (see ESI S9† for absorption growth of Fz over time, Fig. S26–S28†).

To better understand the photoinduced electron transfer dynamics in NADH/ 2^H /XTT liposomes, we performed a Stern–Volmer luminescence quenching study of 2^H liposomes in the presence of various concentrations of NADH in the inner

Table 3 Stern–Volmer quenching constants of 2^H in DPPC membranes with NADH and XTT as quenchers

Quencher	K_{sv} from I_0/I (M^{-1})	k_q ($M^{-1} s^{-1}$)	τ_0 (ns)	Mechanism
NADH/ 2^H	70.8 ± 6.2	$(8.2 \pm 0.7)10^{10}$	0.86	Static
2^H /XTT	$(14.7 \pm 0.7)10^3$	$(1.7 \pm 0.1)10^{13}$	0.86	Static

compartment and XTT in the bulk, respectively. In the Stern–Volmer experiment, the excited state reaction dynamics of 2^H can be resolved by using steady-state and time-resolved emission spectroscopy, by plotting the intensities and the lifetimes as $\frac{I_0}{I}$ and $\frac{\tau_0}{\tau}$ vs. quencher concentrations $[Q]$,⁸⁰ where I_0 and I are the emission intensities in the absence and presence of the quencher, and τ_0 and τ are the excited state lifetimes in the absence and presence of the quencher, respectively.

$$\frac{I_0}{I} = 1 + K_{sv}[Q] = 1 + k_q\tau_0[Q] \quad (2)$$

The slope from the linear regression of $\frac{I_0}{I}$ vs. $[Q]$ is the Stern–Volmer constant K_{sv} . By applying eqn (2), the quenching constant k_q can be derived. After the linear fitting of both plots, NADH and XTT showed characteristic static quenching behavior with the lifetime remaining independent of the quencher concentration. Quenching by encapsulated NADH takes place with $k_q = (8.2 \pm 0.7)10^{10} M^{-1} s^{-1}$. This value is one order of magnitude faster than that in a related study with encapsulated NADH and a positively charged transmembrane molecular wire⁶ and might indicate that a stronger ground state encounter complex is formed between the here investigated 2^H and NADH. Similarly, the quenching by XTT from the bulk takes place with $k_q = (1.71 \pm 0.1)10^{13} M^{-1} s^{-1}$ (Table 3). This value is three orders of magnitudes higher than typical collision-based quenching constants, which are on the order of 10^{11} and it is two orders of magnitude higher than in a similar case between XTT and a positively charged chromophore within the DPPC lipid bilayer.⁶ Therefore, the k_q values obtained for both quenchers indicate the quenching of 2^H due to a strong association with the quencher already in the ground state.^{76,81} We hypothesize that such strong complex formation is partly due to additional hydrogen bond formation by the carboxylate functionality of 2^H with the quencher. The concentrations of NADH and XTT that were applied during the full electron transfer reaction with NADH/ 2^H /XTT are shown in the highlighted region in the Stern–Volmer quenching plot in Fig. 9d and e. The luminescence and kinetic traces can be found in ESI Section S10 and Fig. S30–S33.†

Conclusion

In summary, we demonstrate that the molecular BTAAAs 1^H and 2^H can form mixed-valence species upon one-electron oxidation and that the amphiphilic BTAAAs 1^H and 2^H can be integrated into the lipid bilayer of liposomes for light-driven electron transfer from NADH to the acceptor XTT or *vice versa*. While the



shorter version **1^H** disrupts the lipid bilayer significantly, the long molecule **2^H** integrates well into the bilayer structure, spanning the membrane. Both compounds are redox-active and generate stable one- and two-electron oxidized cations, thereby enabling transmembrane electron transfer. The mechanism of membrane-mediated electron transfer is likely promoted *via* a mixed valence radical cation state of the one-electron oxidized molecular wire, delocalizing an unpaired electron across the entire molecule. The findings of this study provide a design principle for artificial nanoreactors and are highly relevant for the construction of light-driven redox chemistry systems in water with applications in solar light energy conversion.

Data availability

The data supporting the findings of the article can be found in the ESI.† The raw data of photocatalysis will be made available *via* Chemotion.

Author contributions

N. S. and M. N. contributed equally. N. S. performed the membrane integration, luminescence quenching experiments, transmembrane electron transfer experiments and molecular dynamics simulations. M. N. performed the spectroscopic characterization, cyclic voltammetry (including the simulations), spectroelectrochemistry measurements with data analysis, and all (TD)-DFT calculations on the model compounds, supervised by M. L., R. F. W. and A. P., R. F. W. and A. P. devised the project. P. K. conducted one chemical synthesis. G. H. conducted most chemical syntheses. All authors contributed to scientific discussions and the preparation of the manuscript.

Conflicts of interest

The authors declare no conflict of interest.

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References

- 1 S. Bhosale, A. L. Sisson, P. Talukdar, A. Fürstner, N. Banetji, E. Vauthey, G. Bollot, J. Mareda, C. Röger, F. Würthner, N. Sakai and S. Matile, Photoproduction of Proton Gradients with π -Stacked Fluorophore Scaffolds in Lipid Bilayers, *Science*, 2006, **313**(5783), 84–86, DOI: [10.1126/science.1126524](https://doi.org/10.1126/science.1126524).
- 2 A. Perez-Velasco, V. Gorteau and S. Matile, Rigid Oligoperylenediimide Rods: Anion- π Slides with Photosynthetic Activity, *Angew. Chem., Int. Ed.*, 2008, **47**(5), 921–923, DOI: [10.1002/anie.200703749](https://doi.org/10.1002/anie.200703749).
- 3 T. A. Moore, D. Gust, P. Mathis, J.-C. Mialocq, C. Chachaty, R. V. Bensasson, E. J. Land, D. Doizi, P. A. Liddell, W. R. Lehman, G. A. Nemeth and A. L. Moore, Photodriven Charge Separation in a Carotenoporphyrin–Quinone Triad, *Nature*, 1984, **307**(5952), 630–632, DOI: [10.1038/307630a0](https://doi.org/10.1038/307630a0).
- 4 P. Seta, E. Bienvenue, A. L. Moore, P. Mathis, R. V. Bensasson, P. Liddell, P. J. Pessiki, A. Joy, T. A. Moore and D. Gust, Photodriven Transmembrane Charge Separation and Electron Transfer by a Carotenoporphyrin–Quinone Triad, *Nature*, 1985, **316**(6029), 653–655, DOI: [10.1038/316653a0](https://doi.org/10.1038/316653a0).
- 5 B. Limburg, J. Wermink, S. S. van Nielen, R. Kortlever, M. T. M. Koper, E. Bouwman and S. Bonnet, Kinetics of Photocatalytic Water Oxidation at Liposomes: Membrane Anchoring Stabilizes the Photosensitizer, *ACS Catal.*, 2016, **6**(9), 5968–5977, DOI: [10.1021/acscatal.6b00151](https://doi.org/10.1021/acscatal.6b00151).
- 6 N. Sinambela, R. Jacobi, D. Sorsche, L. Gonzalez and A. Pannwitz, Photoinduced Electron Transfer across Phospholipid Bilayers in Anaerobic and Aerobic Atmospheres, *ChemRxiv*, December 3, 2024, preprint, DOI: [10.26434/chemrxiv-2024-x7wgx](https://doi.org/10.26434/chemrxiv-2024-x7wgx).
- 7 U. Fuerholz, H. B. Buerger, F. E. Wagner, A. Stebler, J. H. Ammeter, E. Krausz, R. J. H. Clark, M. J. Stead and A. Ludi, The Creutz-Taube Complex Revisited, *J. Am. Chem. Soc.*, 1984, **106**(1), 121–123, DOI: [10.1021/ja00313a025](https://doi.org/10.1021/ja00313a025).
- 8 R. I. Walter, Substituent Effects on the Properties of Stable Aromatic Free Radicals. The Criterion for Non-Hammett Behavior, *J. Am. Chem. Soc.*, 1966, **88**(9), 1923–1930, DOI: [10.1021/ja00961a014](https://doi.org/10.1021/ja00961a014).
- 9 S. Dapperheld, E. Steckhan, K.-H. G. Brinkhaus and T. Esch, Organic Electron Transfer Systems, II Substituted Triarylamine Cation-Radical Redox Systems – Synthesis, Electrochemical and Spectroscopic Properties, Hammett Behavior, and Suitability as Redox Catalysts, *Chem. Ber.*, 1991, **124**(11), 2557–2567, DOI: [10.1002/cber.19911241127](https://doi.org/10.1002/cber.19911241127).
- 10 S. Amthor, B. Noller and C. Lambert, UV/Vis/NIR Spectral Properties of Triarylaminines and Their Corresponding Radical Cations, *Chem. Phys.*, 2005, **316**(1), 141–152, DOI: [10.1016/j.chemphys.2005.05.009](https://doi.org/10.1016/j.chemphys.2005.05.009).
- 11 A. Heckmann, C. Lambert, M. Goebel and R. Wortmann, Synthesis and Photophysics of a Neutral Organic Mixed-Valence Compound, *Angew. Chem., Int. Ed.*, 2004, **43**(43), 5851–5856, DOI: [10.1002/anie.200460495](https://doi.org/10.1002/anie.200460495).



- 12 G. Grelaud, M. P. Cifuentes, T. Schwich, G. Argouarch, S. Petrie, R. Stranger, F. Paul and M. G. Humphrey, Multistate Redox-Active Metalated Triarylamines, *Eur. J. Inorg. Chem.*, 2012, **2012**(1), 65–75, DOI: [10.1002/ejic.201100747](https://doi.org/10.1002/ejic.201100747).
- 13 W. Polit, T. Exner, E. Wuttke and R. Winter, Vinylruthenium-Triarylamine Conjugates as Electroswitchable Polyelectrochromic NIR Dyes, *BioInorg. React. Mech.*, 2012, **8**, 85–105, DOI: [10.1515/irm-2012-0005](https://doi.org/10.1515/irm-2012-0005).
- 14 C. Hassenrück and R. F. Winter, Manipulation and Assessment of Charge and Spin Delocalization in Mixed-Valent Triarylamine-Vinylruthenium Conjugates, *Inorg. Chem.*, 2017, **56**(21), 13517–13529, DOI: [10.1021/acs.inorgchem.7b02186](https://doi.org/10.1021/acs.inorgchem.7b02186).
- 15 S. Breimaier and R. F. Winter, Electrochemical and Spectroscopic Studies on Triarylamine-Polychlorotriphenylmethyl Dyads with Particularly Strong Triarylamine Donors, *Eur. J. Org. Chem.*, 2021, **2021**(33), 4690–4700, DOI: [10.1002/ejoc.202100476](https://doi.org/10.1002/ejoc.202100476).
- 16 C.-J. Yao, R.-H. Zheng, Q. Shi, Y.-W. Zhong and J. Yao, 1,4-Benzene-Bridged Covalent Hybrid of Triarylamine and Cyclometalated Ruthenium: A New Type of Organic-Inorganic Mixed-Valent System, *Chem. Commun.*, 2012, **48**(45), 5680–5682, DOI: [10.1039/C2CC32471G](https://doi.org/10.1039/C2CC32471G).
- 17 J.-H. Tang, J.-Y. Shao, Y.-Q. He, S.-H. Wu, J. Yao and Y.-W. Zhong, Transition from a Metal-Localized Mixed-Valence Compound to a Fully Delocalized and Bridge-Biased Electrophore in a Ruthenium-Amine-Ruthenium Tricenter System, *Chem.-Eur. J.*, 2016, **22**(30), 10341–10345, DOI: [10.1002/chem.201601806](https://doi.org/10.1002/chem.201601806).
- 18 J. Zhang, S.-Z. Guo, Y.-B. Dong, L. Rao, J. Yin, G.-A. Yu, F. Hartl and S. H. Liu, Multistep Oxidation of Diethynyl Oligophenylamine-Bridged Diruthenium and Diiron Complexes, *Inorg. Chem.*, 2017, **56**(2), 1001–1015, DOI: [10.1021/acs.inorgchem.6b02809](https://doi.org/10.1021/acs.inorgchem.6b02809).
- 19 K. E. Linton, M. A. Fox, L.-O. Pålsson and M. R. Bryce, Oligo(p-Phenyleneethynylene) (OPE) Molecular Wires: Synthesis and Length Dependence of Photoinduced Charge Transfer in OPEs with Triarylamine and Diaryloxadiazole End Groups, *Chem.-Eur. J.*, 2015, **21**(10), 3997–4007, DOI: [10.1002/chem.201406080](https://doi.org/10.1002/chem.201406080).
- 20 Y.-P. Ou, J. Zhang, A. Wang, A. Yuan, C. Yin and S. H. Liu, Rutheniumethynyl-Triarylamine Organic-Inorganic Mixed-Valence Systems: Regulating Ru-N Electronic Coupling by Different Aryl Bridge Cores, *Chem.-Asian J.*, 2020, **15**(20), 3338–3349, DOI: [10.1002/asia.202000879](https://doi.org/10.1002/asia.202000879).
- 21 Z. Lin, Z. Li, W. Xiao, L. Kong, J. Xu, Y. Xia, X. Zhu, F. Zhang and Y.-P. Ou, Terpyridine Ruthenium-Triarylamine Asymmetrical Mixed-Valence Systems: Syntheses, (Spectro) Electrochemistry and Theoretical Calculations, *J. Organomet. Chem.*, 2023, **993**, 122708, DOI: [10.1016/j.jorganchem.2023.122708](https://doi.org/10.1016/j.jorganchem.2023.122708).
- 22 H. Murata and P. M. Lahti, Synthesis and Oxidation of Triarylamine Derivatives Bearing Hydrogen-Bonding Groups, *J. Org. Chem.*, 2007, **72**(13), 4974–4977, DOI: [10.1021/jo070318a](https://doi.org/10.1021/jo070318a).
- 23 G. Sforazzini, A. Kahnt, M. Wykes, J. K. Sprafke, S. Brovelli, D. Montarnal, F. Meinardi, F. Cacialli, D. Beljonne, B. Albinsson and H. L. Anderson, Synthesis and Photophysics of Coaxial Threaded Molecular Wires: Polyrotaxanes with Triarylamine Jackets, *J. Phys. Chem. C*, 2014, **118**(8), 4553–4566, DOI: [10.1021/jp500624q](https://doi.org/10.1021/jp500624q).
- 24 P. Thamaraiselvi, N. Duraipandy, M. S. Kiran and S. Easwaramoorthi, Triarylamine Rhodanine Derivatives as Red Emissive Sensor for Discriminative Detection of Ag⁺ and Hg²⁺ Ions in Buffer-Free Aqueous Solutions, *ACS Sustain. Chem. Eng.*, 2019, **7**(11), 9865–9874, DOI: [10.1021/acssuschemeng.9b00417](https://doi.org/10.1021/acssuschemeng.9b00417).
- 25 C. A. Swamy P, A. V. Raveendran, N. Sivakrishna and R. P. Nandi, Triarylborane-Triphenylamine Based Luminophore for the Mitochondria Targeted Live Cell Imaging and Colorimetric Detection of Aqueous Fluoride, *Dalton Trans.*, 2022, **51**(40), 15339–15353, DOI: [10.1039/D2DT01887J](https://doi.org/10.1039/D2DT01887J).
- 26 R. R. Nelson and R. N. Adams, Anodic Oxidation Pathways of Substituted Triphenylamines. II. Quantitative Studies of Benzidine Formation, *J. Am. Chem. Soc.*, 1968, **90**(15), 3925–3930, DOI: [10.1021/ja01017a004](https://doi.org/10.1021/ja01017a004).
- 27 R. F. Nelson and R. H. Philp, Electrochemical and Spectroscopic Studies of Cation Radicals. 4. Stopped-Flow Determination of Triarylammonium Radical Coupling Rate Constants, *J. Phys. Chem.*, 1979, **83**(6), 713–716, DOI: [10.1021/j100469a015](https://doi.org/10.1021/j100469a015).
- 28 K. Sreenath, C. V. Suneesh, V. K. Ratheesh Kumar and K. R. Gopidas, Cu(II)-Mediated Generation of Triarylamine Radical Cations and Their Dimerization. An Easy Route to Tetraarylbenzidines, *J. Org. Chem.*, 2008, **73**(8), 3245–3251, DOI: [10.1021/jo800349n](https://doi.org/10.1021/jo800349n).
- 29 M. R. Talipov, M. M. Hossain, A. Boddada, K. Thakur and R. Rathore, A Search for Blues Brothers: X-Ray Crystallographic/Spectroscopic Characterization of the Tetraarylbenzidine Cation Radical as a Product of Aging of Solid Magic Blue, *Org. Biomol. Chem.*, 2016, **14**(10), 2961–2968, DOI: [10.1039/c6ob00140h](https://doi.org/10.1039/c6ob00140h).
- 30 R. Reynolds, L. L. Line and R. F. Nelson, Electrochemical Generation of Carbazoles from Aromatic Amines, *J. Am. Chem. Soc.*, 1974, **96**(4), 1087–1092, DOI: [10.1021/ja00811a024](https://doi.org/10.1021/ja00811a024).
- 31 N. G. Connelly and W. E. Geiger, Chemical Redox Agents for Organometallic Chemistry, *Chem. Rev.*, 1996, **96**(2), 877–910, DOI: [10.1021/cr940053x](https://doi.org/10.1021/cr940053x).
- 32 B. Mladenova, D. R. Kattnig, C. Kaiser, J. Schäfer, C. Lambert and G. Grampp, Investigations of the Degenerate Intramolecular Charge Exchange in Symmetric Organic Mixed Valence Compounds: Solvent Dynamics of Bis(Triarylamine)Paracyclophane Redox Systems, *J. Phys. Chem. C*, 2015, **119**(16), 8547–8553, DOI: [10.1021/acs.jpcc.5b01386](https://doi.org/10.1021/acs.jpcc.5b01386).
- 33 J. Schäfer, M. Holzapfel, B. Mladenova, D. Kattnig, I. Krummenacher, H. Braunschweig, G. Grampp and C. Lambert, Hole Transfer Processes in Meta- and Para-Conjugated Mixed Valence Compounds: Unforeseen Effects of Bridge Substituents and Solvent Dynamics, *J. Am. Chem. Soc.*, 2017, **139**(17), 6200–6209, DOI: [10.1021/jacs.7b01650](https://doi.org/10.1021/jacs.7b01650).



- 34 S. Dümmler, W. Roth, I. Fischer, A. Heckmann and C. Lambert, Excited-State Dynamics in a Neutral Organic Mixed-Valence Compound, *Chem. Phys. Lett.*, 2005, **408**(4), 264–268, DOI: [10.1016/j.cplett.2005.04.045](#).
- 35 K. Tahara, T. Nakakita, A. A. Starikova, T. Ikeda, M. Abe and J. Kikuchi, Small Anion-Assisted Electrochemical Potential Splitting in a New Series of Bistriarylamine Derivatives: Organic Mixed Valency across a Urea Bridge and Zwitterionization, *Beilstein J. Org. Chem.*, 2019, **15**(1), 2277–2286, DOI: [10.3762/bjoc.15.220](#).
- 36 J. Hankache and O. S. Wenger, Organic Mixed Valence, *Chem. Rev.*, 2011, **111**(8), 5138–5178, DOI: [10.1021/cr100441k](#).
- 37 M. Kaupp, M. Renz, M. Parthey, M. Stolte, F. Würthner and C. Lambert, Computational and Spectroscopic Studies of Organic Mixed-Valence Compounds: Where Is the Charge?, *Phys. Chem. Chem. Phys.*, 2011, **13**(38), 16973–16986, DOI: [10.1039/C1CP21772K](#).
- 38 A. Heckmann and C. Lambert, Organic Mixed-Valence Compounds: A Playground for Electrons and Holes, *Angew. Chem., Int. Ed.*, 2012, **51**(2), 326–392, DOI: [10.1002/anie.201100944](#).
- 39 C. Lambert and G. Nöll, The Class II/III Transition in Triarylamine Redox Systems, *J. Am. Chem. Soc.*, 1999, **121**(37), 8434–8442, DOI: [10.1021/ja991264s](#).
- 40 K. Lancaster, S. A. Odom, S. C. Jones, S. Thayumanavan, S. R. Marder, J.-L. Brédas, V. Coropceanu and S. Barlow, Intramolecular Electron-Transfer Rates in Mixed-Valence Triarylaminos: Measurement by Variable-Temperature ESR Spectroscopy and Comparison with Optical Data, *J. Am. Chem. Soc.*, 2009, **131**(5), 1717–1723, DOI: [10.1021/ja808465c](#).
- 41 M. Uebe and A. Ito, Intramolecular Charge Transfer in Kekulé- and Non-Kekulé-Bridged Bis(Triarylamine) Radical Cations: Missing Key Compounds in Organic Mixed-Valence Systems, *Chem.-Asian J.*, 2019, **14**(10), 1692–1696, DOI: [10.1002/asia.201900036](#).
- 42 M. Uebe, K. Kaneda, S. Fukuzaki and A. Ito, Bridge-Length-Dependent Intramolecular Charge Transfer in Bis(Dianisylamino)-Terminated Oligo(p-Phenylene)s, *Chem.-Eur. J.*, 2019, **25**(68), 15455–15462, DOI: [10.1002/chem.201903667](#).
- 43 K. E. Linton, M. A. Fox, L.-O. Pålsson and M. R. Bryce, Oligo(p-phenyleneethynylene) (OPE) Molecular Wires: Synthesis and Length Dependence of Photoinduced Charge Transfer in OPEs with Triarylamine and Diaryloxadiazole End Groups, *Chem.-Eur. J.*, 2015, **21**(10), 3997–4007, DOI: [10.1002/chem.201406080](#).
- 44 A. Heckmann, S. Amthor and C. Lambert, Mulliken–Hush Analysis of a Bis(Triarylamine) Mixed-Valence System with a N···N Distance of 28.7 Å, *Chem. Commun.*, 2006, (28), 2959–2961, DOI: [10.1039/b604603g](#).
- 45 C. Lambert and V. Kriegisch, Heterogeneous Electron Transfer Processes in Self-Assembled Monolayers of Amine Terminated Conjugated Molecular Wires, *Langmuir*, 2006, **22**(21), 8807–8812, DOI: [10.1021/la061404t](#).
- 46 L. G. Heinz, O. Yushchenko, M. Neuburger, E. Vauthey and O. S. Wenger, Tetramethoxybenzene Is a Good Building Block for Molecular Wires: Insights from Photoinduced Electron Transfer, *J. Phys. Chem. A*, 2015, **119**(22), 5676–5684, DOI: [10.1021/acs.jpca.5b03649](#).
- 47 L. Li, J. Z. Low, J. Wilhelm, G. Liao, S. Gunasekaran, C. R. Prindle, R. L. Starr, D. Golze, C. Nuckolls, M. L. Steigerwald, F. Evers, L. M. Campos, X. Yin and L. Venkataraman, Highly Conducting Single-Molecule Topological Insulators Based on Mono- and Di-Radical Cations, *Nat. Chem.*, 2022, **14**(9), 1061–1067, DOI: [10.1038/s41557-022-00978-1](#).
- 48 X. Shi, M. Liu, L. Li, J. Zhang, H. Li, Z. Huang, W. Zhang, Z. Zhang, N. Zhou and X. Zhu, Efficient Synthesis of Discrete Oligo(Fluorenediacetylene)s toward Chain-Length-Dependent Optical and Structural Properties, *Polym. Chem.*, 2021, **12**(17), 2598–2605, DOI: [10.1039/D1PY00165E](#).
- 49 A. Nafady, T. T. Chin and W. E. Geiger, Manipulating the Electrolyte Medium to Favor Either One-Electron or Two-Electron Oxidation Pathways for (Fulvalendiyl)Dirhodium Complexes, *Organometallics*, 2006, **25**(7), 1654–1663, DOI: [10.1021/om051101e](#).
- 50 A. K. Diallo, C. Absalon, J. Ruiz and D. Astruc, Ferrocenyl-Terminated Redox Stars: Synthesis and Electrostatic Effects in Mixed-Valence Stabilization, *J. Am. Chem. Soc.*, 2011, **133**(3), 629–641, DOI: [10.1021/ja109380u](#).
- 51 U. Pfaff, A. Hildebrandt, D. Schaarschmidt, T. Hahn, S. Liebing, J. Kortus and H. Lang, Di- and Triferrocenyl (Hetero)Aromatics: Synthesis, Characterization, (Spectro-) Electrochemistry, and Calculations, *Organometallics*, 2012, **31**(19), 6761–6771, DOI: [10.1021/om300511v](#).
- 52 R. F. Winter, Half-Wave Potential Splittings $\Delta E_{1/2}$ as a Measure of Electronic Coupling in Mixed-Valent Systems: Triumphs and Defeats, *Organometallics*, 2014, **33**(18), 4517–4536, DOI: [10.1021/om500029x](#).
- 53 S. Barlow, C. Risko, S.-J. Chung, N. M. Tucker, V. Coropceanu, S. C. Jones, Z. Levi, J.-L. Brédas and S. R. Marder, Intervalence Transitions in the Mixed-Valence Monocations of Bis(Triarylaminos) Linked with Vinylene and Phenylene–Vinylene Bridges, *J. Am. Chem. Soc.*, 2005, **127**(48), 16900–16911, DOI: [10.1021/ja054136e](#).
- 54 S. Barlow, C. Risko, S. A. Odom, S. Zheng, V. Coropceanu, L. Beverina, J.-L. Brédas and S. R. Marder, Tuning Delocalization in the Radical Cations of 1,4-Bis[4-(Diarylamino)Styryl]Benzenes, 2,5-Bis[4-(Diarylamino)Styryl]Thiophenes, and 2,5-Bis[4-(Diarylamino)Styryl]Pyrroles through Substituent Effects, *J. Am. Chem. Soc.*, 2012, **134**(24), 10146–10155, DOI: [10.1021/ja3023048](#).
- 55 C. Lambert, S. Amthor and J. Schelter, From Valence Trapped to Valence Delocalized by Bridge State Modification in Bis(Triarylamine) Radical Cations: Evaluation of Coupling Matrix Elements in a Three-Level System, *J. Phys. Chem. A*, 2004, **108**(31), 6474–6486, DOI: [10.1021/jp048449s](#).
- 56 I. Noviandri, K. N. Brown, D. S. Fleming, P. T. Gulyas, P. A. Lay, A. F. Masters and L. Phillips, The Decamethylferrocenium/Decamethylferrocene Redox Couple: A Superior Redox Standard to the Ferrocenium/Ferrocene Redox Couple for Studying Solvent Effects on



- the Thermodynamics of Electron Transfer, *J. Phys. Chem. B*, 1999, **103**(32), 6713–6722, DOI: [10.1021/jp991381+](https://doi.org/10.1021/jp991381+).
- 57 K.-T. Wong, T. Hsi Hung, S. C. Kao, C. Hsien Chou and Y. Oliver Su, Synthesis and Properties of Novel Bis(Triarylaminos) Based on a 3,3'-Diphenyl-2,2'-Bithiophene Core, *Chem. Commun.*, 2001, **0**(17), 1628–1629, DOI: [10.1039/B103194P](https://doi.org/10.1039/B103194P).
 - 58 M. Renz, K. Theilacker, C. Lambert and M. Kaupp, A Reliable Quantum-Chemical Protocol for the Characterization of Organic Mixed-Valence Compounds, *J. Am. Chem. Soc.*, 2009, **131**(44), 16292–16302, DOI: [10.1021/ja9070859](https://doi.org/10.1021/ja9070859).
 - 59 M. Renz and M. Kaupp, Predicting the Localized/Delocalized Character of Mixed-Valence Diquinone Radical Anions. Toward the Right Answer for the Right Reason, *J. Phys. Chem. A*, 2012, **116**(43), 10629–10637, DOI: [10.1021/jp308294r](https://doi.org/10.1021/jp308294r).
 - 60 F. S. Gogesch, S. Schwab, A. Rehse, M. Linseis, A. Baksi, G. H. Clever and R. F. Winter, Diruthenium Complexes with π -Extended Bridging Bis(Alkenyl)Arylene Ligands, A Derived Metallamacrocycle, and Their Oxidized Forms, *Organometallics*, 2023, **42**(21), 3085–3098, DOI: [10.1021/acs.organomet.3c00337](https://doi.org/10.1021/acs.organomet.3c00337).
 - 61 E. D. Glendening, C. R. Landis and F. Weinhold, Natural Bond Orbital Methods, *Wiley Interdiscip. Rev.: Comput. Mol. Sci.*, 2012, **2**(1), 1–42, DOI: [10.1002/wcms.51](https://doi.org/10.1002/wcms.51).
 - 62 E. Wu, K. Jacobson and D. Papahadjopoulos, Lateral Diffusion in Phospholipid Multibilayers Measured by Fluorescence Recovery after Photobleaching, *Biochemistry*, 1977, **16**(17), 3936–3941, DOI: [10.1021/bi00636a034](https://doi.org/10.1021/bi00636a034).
 - 63 K. Watanabe, K. Moriya, T. Kouyama, A. Onoda, T. Minatani, S. Y. Takizawa and S. Murata, Photoinduced Transmembrane Electron Transport in DPPC Vesicles: Mechanism and Application to a Hydrogen Generation System, *J. Photochem. Photobiol., A*, 2011, **221**(1), 113–122, DOI: [10.1016/j.jphotochem.2011.04.034](https://doi.org/10.1016/j.jphotochem.2011.04.034).
 - 64 H. Song, A. Amati, A. Pannwitz, S. Bonnet and L. Hammarström, Mechanistic Insights into the Charge Transfer Dynamics of Photocatalytic Water Oxidation at the Lipid Bilayer–Water Interface, *J. Am. Chem. Soc.*, 2022, **144**(42), 19353–19364, DOI: [10.1021/jacs.2c06842](https://doi.org/10.1021/jacs.2c06842).
 - 65 N. Sinambela, R. Jacobi, D. Hernández-Castillo, E. Hofmeister, N. Hagmeyer, B. Dietzek, L. González and A. Pannwitz, Alignment and Photooxidation Dynamics of a Perylene Diimide Chromophore in Lipid Bilayers, *Mol. Syst. Des. Eng.*, 2023, **8**(7), 842–852, DOI: [10.1039/D2ME00243D](https://doi.org/10.1039/D2ME00243D).
 - 66 E. Krieger, T. Darden, S. B. Nabuurs, A. Finkelstein and G. Vriend, Making Optimal Use of Empirical Energy Functions: Force-Field Parameterization in Crystal Space, *Proteins: Struct., Funct., Genet.*, 2004, **57**(4), 678–683, DOI: [10.1002/prot.20251](https://doi.org/10.1002/prot.20251).
 - 67 E. Krieger, J. E. Nielsen, C. A. E. M. Spronk and G. Vriend, Fast Empirical pK_a Prediction by Ewald Summation, *J. Mol. Graphics Modell.*, 2006, **25**(4), 481–486, DOI: [10.1016/j.jmgm.2006.02.009](https://doi.org/10.1016/j.jmgm.2006.02.009).
 - 68 D. Marsh, *Handbook of Lipid Bilayers*, CRC Press, 2013, DOI: [10.1201/b11712](https://doi.org/10.1201/b11712).
 - 69 D. Drabik, G. Chodaczek, S. Kraszewski and M. Langner, Mechanical Properties Determination of DMPC, DPPC, DSPC, and HSPC Solid-Ordered Bilayers, *Langmuir*, 2020, **36**(14), 3826–3835, DOI: [10.1021/acs.langmuir.0c00475](https://doi.org/10.1021/acs.langmuir.0c00475).
 - 70 S. Dutta, B. Watson, S. Mattoo and J.-C. Rochet, Calcein Release Assay to Measure Membrane Permeabilization by Recombinant Alpha-Synuclein, *Bio-Protoc.*, 2020, **10**(14), e3690, DOI: [10.21769/BioProtoc.3690](https://doi.org/10.21769/BioProtoc.3690).
 - 71 B. Maherani, E. Arab-Tehrany, A. Kheirloomoom, D. Geny and M. Linder, Calcein Release Behavior from Liposomal Bilayer; Influence of Physicochemical/Mechanical/Structural Properties of Lipids, *Biochimie*, 2013, **95**(11), 2018–2033, DOI: [10.1016/j.biochi.2013.07.006](https://doi.org/10.1016/j.biochi.2013.07.006).
 - 72 L. E. Garner, J. Park, S. M. Dyar, A. Chworos, J. J. Sumner and G. C. Bazan, Modification of the Optoelectronic Properties of Membranes via Insertion of Amphiphilic Phenylenevinylene Oligoelectrolytes, *J. Am. Chem. Soc.*, 2010, **132**(29), 10042–10052, DOI: [10.1021/ja1016156](https://doi.org/10.1021/ja1016156).
 - 73 B. Wang, M. Wang, A. Mikhailovsky, S. Wang and G. C. Bazan, A Membrane-Intercalating Conjugated Oligoelectrolyte with High-Efficiency Photodynamic Antimicrobial Activity, *Angew. Chem., Int. Ed.*, 2017, **56**(18), 5031–5034, DOI: [10.1002/anie.201701146](https://doi.org/10.1002/anie.201701146).
 - 74 C. M. Legaspi, R. E. Stubbs, M. Wahadoszaman, D. J. Yaron, L. A. Peteanu, A. Kemboi, E. Fossum, Y. Lu, Q. Zheng and L. J. Rothberg, Rigidity and Polarity Effects on the Electronic Properties of Two Deep Blue Delayed Fluorescence Emitters, *J. Phys. Chem. C*, 2018, **122**(22), 11961–11972, DOI: [10.1021/acs.jpcc.7b12025](https://doi.org/10.1021/acs.jpcc.7b12025).
 - 75 A. Pannwitz, H. Saaring, N. Beztsinna, X. Li, M. A. Siegler and S. Bonnet, Mimicking Photosystem I with a Transmembrane Light Harvester and Energy Transfer-Induced Photoreduction in Phospholipid Bilayers, *Chem.–Eur. J.*, 2021, **27**(9), 3013–3018, DOI: [10.1002/chem.202003391](https://doi.org/10.1002/chem.202003391).
 - 76 *Principles of Fluorescence Spectroscopy*, ed. J. R. Lakowicz, Springer US, Boston, MA, 2006, DOI: [10.1007/978-0-387-46312-4](https://doi.org/10.1007/978-0-387-46312-4).
 - 77 R. E. P. Nau, J. Bösking and A. Pannwitz, Compartmentalization Accelerates Photosensitized NADH to NAD⁺ Conversion, *ChemPhotoChem*, 2022, **6**(11), e202200158, DOI: [10.1002/cptc.202200158](https://doi.org/10.1002/cptc.202200158).
 - 78 M. W. Sutherland and B. A. Learmonth, The Tetrazolium Dyes MTS and XTT Provide New Quantitative Assays for Superoxide and Superoxide Dismutase, *Free Radical Res.*, 1997, **27**(3), 283–289, DOI: [10.3109/10715769709065766](https://doi.org/10.3109/10715769709065766).
 - 79 S. Fukuzumi, M. Ishikawa and T. Tanaka, Mechanisms of Photo-Oxidation of NADH Model Compounds by Oxygen, *J. Chem. Soc., Perkin Trans. 2*, 1989, (8), 1037–1045, DOI: [10.1039/P29890001037](https://doi.org/10.1039/P29890001037).
 - 80 V. Balzani, P. Ceroni and A. Juris, *Photochemistry and Photophysics: Concepts, Research, Applications*, John Wiley & Sons, 2014.
 - 81 Y. Rahimi, A. Goulding, S. Shrestha, S. Mirpuri and S. K. Deo, Mechanism of Copper Induced Fluorescence Quenching of Red Fluorescent Protein, DsRed, *Biochem. Biophys. Res. Commun.*, 2008, **370**(1), 57–61, DOI: [10.1016/j.bbrc.2008.03.034](https://doi.org/10.1016/j.bbrc.2008.03.034).

