Determination of standard redox rate constants of OLED active compounds by electrochemical impedance spectroscopy

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ABSTRACT

A number of commercial organic compounds (m-MTDATA, PBD, CBP, TAPC, NPB, TPBi, etc) as well as several donor-acceptor-donor (D-A-D) compounds were investigated by cyclic voltammetry and electrochemical impedance spectroscopy. The compounds were chosen as they are promising components of small-molecule-based high efficiency Thermally Activated Delayed Fluorescence (TADF) emitters in Organic Light Emitting Diodes (OLEDs).

Electrochemical impedance spectra of a Pt electrode in CH_2Cl_2 solutions containing the investigated compound and Bu_4NBF_4 as the electrolyte were obtained and analyzed by

electrochemical electrical circuit methods. Charge transfer resistance, double layer capacitance, Warburg constant and other parameters were determined and represented as a function of the potential. Analysis of charge transfer resistance as a function of potential allowed an estimation of standard redox rate constants for the compounds' oxidation and reduction processes.

Two main features concerning the redox reaction rates of OLED-active compounds were revealed: (i) the oxidation and reduction rates of ambipolar compounds, i.e. containing both donor and acceptor parts, were found to be much higher than those of unipolar donor-only and acceptoronly molecules; (ii) the relationship between the oxidation and reduction rate constants was shown to be related to the compounds' conductivity type in the solid state.

KEYWORDS: OLED, standard redox rate constant, electrochemical impedance spectroscopy, donor, acceptor, ambipolar, exciplex, TADF.

1. Introduction

Nowadays one of the most interesting applications of organic compounds is in the rapidly expanding organic electronics area [1]. A common theme in organic electronics is that the organic compounds need to function in an environment where an electric current is passed through them, so whether the compound is a conductor, semi-conductor or insulator, the electrochemical investigation is necessary. Electrochemical methods are usually applied at the first stage of characterisation of the organic compounds. There are several different electrochemical methods which can be used to analyse organic compounds but the mostly widely used one is Cyclic Voltammetry (CV). The main goal of a CV investigation is determination of the compound's stability and an estimation of the Ionization Potential (IP) and Electron Affinity (EA) which correspond to the Highest Occupied and Lowest Unoccupied Molecular Orbital (HOMO and LUMO) energy levels, respectively [2–13]. From an organic electronics point of view, the energy levels are an important criterion for device design, since they determine energy losses during charge

transfer across the layer-layer interfaces. They also determine exciplex formation and its emission energy.

In this study we developed an electrochemical impedance-based technique for estimation not only of thermodynamic but also of kinetic parameters of molecular oxidation and reduction processes. Analysis of electrochemical impedance spectra has enabled the extraction of purely kinetic parameters, charge transfer resistance, which value is not affected by diffusion.

Our new method includes the following stages. Firstly, potential regions corresponding to reversible oxidation and reduction processes are revealed by cyclic voltammetry. Then a number of impedance spectra in increments of 0.1 V are registered in the potential range where a reversible peak is observed in the CV. All the spectra are analysed by an equivalent electrical circuit method, and the parameters (resistance, capacitance, etc) are presented as functions of potentials, and the electrochemical rate constant is calculated.

For our work 10 commercial compounds were chosen: 4,4',4"-Tris[phenyl(*m*-tolyl)amino]triphenylamine (**m**-**MTDATA**); 4,4'-Bis(*N*-carbazolyl)-1,1'-biphenyl (**CBP**); 2-(4-*tert*-butylphenyl)-5-(4-biphenylyl)-1,3,4-oxadiazole (**tBu-PBD**); *N*,*N*'-Di(1-naphthyl)-*N*,*N*'-diphenyl-(1,1'-biphenyl)-4,4'-diamine (**NPB**); 2,2',2"-(1,3,5-Benzinetriyl)-tris(1-phenyl-1-*H*-benzimidazole) (**TPBi**); 4,4'-Cyclohexylidenebis[*N*,*N*-bis(4-methylphenyl)benzenamine] (**TAPC**); 1,3-Bis(*N*-carbazolyl)benzene (**mCP**), Tris(2,4,6-trimethyl-3-(pyridin-3-yl)phenyl)borane (**3TPYMB**); Bis[2-(diphenylphosphino)phenyl] ether oxide (**DPEPO**); 9-(4-*tert*-butylphenyl)-3,6-bis(triphenylsilyl)-9*H*-carbazole (**CzSi**) mostly to show the usefulness of the method. These compounds were chosen because they take an active part in intermolecular exciplex formation (**Table 1**) [10, 12, 14, 15].

	Structure	$E_{\rm EA}$ [eV]	$E_{\rm IP}$ [eV]	k^0_{red} [m/s]	k^0_{ox} [m/s]	Use in OLEDs
m-MTDATA		-2.0	-5.1	_	1.4.10 ⁻³	donor, hole- transport layer [16]
СВР		-2.9	-6.0	_	_	host, hole- transport layer [11]
tBu-PBD		-2.6	-6.2	7.3.10 ⁻⁵	_	acceptor, electron- transport layer [17]
NPB		-2.4	-5.4	_	$8.7 \cdot 10^{-4}$	donor, hole- transport layer [14]
ТРВі		-2.7	-5.9	1.2.10 ⁻⁵	_	acceptor, electron- transport layer [14]
ТАРС	H ₃ C	-2.0	-5.2	_	$2.5 \cdot 10^{-3}$	donor, hole- transport layer [10]
ЗТРУМВ		-3.3	-6.8	_	_	acceptor, electron- transport layer [16]
mCP		-2.4	-6.1	_	_	host, hole- transport layer [18]
DPEPO	Ph_O_O_Ph Ph-P'_P-Ph	-2.0	-6.1	_	_	host [15]
CzSi	Ph, Ph Si Ph' N H	-2.5	-6.0	_	_	host [19]

Table 1. Standard electrochemical rate constants, structures and roles of a number of commercial OLED compounds.

To expand our work, several ambipolar compounds consisting of both donor and acceptor parts (D-A-D) were also studied (**Table 2**): 2,8-bis(10*H*-phenoxazin-10-yl)dibenzo[b,d]thiophene-S,Sinformation for dioxide **(I**) (see supporting details); 3,7-bis(10H-phenothiazin-10yl)dibenzo[b,d]thiophene-S,S-dioxide **(II**) [8]; 2,8-bis(5H-dibenzo[b,f]azepin-5yl)dibenzo[b,d]thiophene-5,5-dioxide (III) see supporting information for details); 2,8-bis(10Hphenothiazin-10-yl)dibenzo[b,d]thiophene-S,S-dioxide (**IV**) [8,9]; 9-[2,8]-9-carbazole-[dibenzothiophene-*S*,*S*-dioxide]-carbazole **(V)** [10]; 2,8-bis(bis(4butylphenyl)amino)dibenzo[*b*,*d*]thiophene-5,5-dioxide (VI) [11].

	Structure	LUMO, eV	HOMO, eV	$k^{0}_{\text{red}}, \text{m/s}$	$k^0_{\text{ox}}, \text{m/s}$
I		-3.15	-5.61	3.7.10 ⁻⁴	$4.6 \cdot 10^{-3}$
п		-3.03	-5.56	1.4.10 ⁻⁴	1.8·10 ⁻³
ш		-2.55	-6.04	2.3.10 ⁻⁴	$1.1 \cdot 10^{-4}$
IV		-3.08	-5.62	1.9.10 ⁻⁴	> 1.10 ⁻³
V	But +Bu +Bu +Bu +Bu	-2.97	-6.07	4.3·10 ⁻³	$9.4 \cdot 10^{-4}$
VI		-2.7	-5.83	7.6·10 ⁻³	$8.4 \cdot 10^{-4}$

Table 2. Standard electrochemical rate constants of a number of D-A-D OLED compounds and estimated values of their HOMO and LUMO levels.

2. Experimental Section

Structural formulas of the compounds are provided in **Tables 1** and **2** along with a summary of the results obtained. Commercial compounds (**Table 1**) were purchased from Sigma-Aldrich, TCI or Molekula and used without further purification. The working CH_2Cl_2 solution in all experiments contained $5 \cdot 10^{-3}$ mol/L of the investigated organic compound and 0.1 mol/L Bu_4NBF_4 as an electrolyte. The solution was initially deaerated by Ar for 10 min, and the atmosphere above the solution was saturated with Ar during whole measurement procedure.

The electrochemical experiments were carried out using a BioLogic SP-150 potentiostat with a built-in frequency response analyser. Pt working electrode ET075 from eDAQ, Pt counter electrode (wire) and Ag/AgCl reference electrode were employed in a three-electrode electrochemical cell. The counter electrode was flame annealed and the working electrode was polished before each experiment. All the potentials in this work are presented versus Fc⁺/Fc (ferrocene) reference potential.

The potential scan rate used in cyclic voltammetry was 100 mV/s. Impedance spectra were obtained in the 10 kHz – 1 Hz frequency range with 20 points per decade in a logarithmic scale (total number of frequencies in one spectrum was 61). The analysis of the electrochemical impedance spectra and the determination of equivalent circuit parameters was performed using «EIS Analyser» software [20].

3. Results and Discussion

The characteristic voltammetric peaks of the compounds in solution are shown in **Figure 1** in comparison with background current of the solvent and electrolyte. The background current significantly complicated recognition of reduction peaks, especially in case of compounds enlisted in **Table 1**, however in order to maintain the same condition throughout the work we did not refuse

of using dichloromethane which appeared to be the best to dissolve all the compounds under investigation up to concentration of 0.05 M required for characterisation of the redox kinetics. The oxidation peaks of the compounds appeared in a range of 0 - 1.3 V while reduction peaks were observed in a range between -1.7 and -2.5 V.



Figure 1. Characteristic cyclic voltammograms showing oxidation and reduction of the compounds. Sample concentration 0.05 M, scan rate 0.05 V·s⁻¹, dichloromethane solution. All potentials vs Fc/Fc^+ redox couple.

As **Figure 1 a, c** shows that most compounds enlisted in **Table 1** underwent either oxidation (m-MTDATA, TAPC, NPB) or reduction (PBD, TPBi) over a broad potential range. Several compounds (DPEPO, CzSi, mCP, 3TPYMB) did not show any noticeable redox behaviour. Since oxidation of all the compounds occurred at rather low potentials (< 1 V) electrolyte oxidation did not mask them and complicate estimation of oxidation potential. CVs of TPBi and PBD did not show any significant difference in comparison with the background curve in broad positive potential range, thus they were considered non-oxidisable.

Two compounds (NPB and TAPC) did not show any noticeable reduction behaviour. Nevertheless a considerable cathodic current within the range of supposed reduction potentials in case of CBP and m-MTDATA was observed. In first experiments involving very high scan rate (up to 0.2 V s^{-1}) and high electrolyte concentration (up to 1.0 M) we observed a small anodic current peak right after the switch of the scan direction which was initially wrongly recognised as a part of quasi-reversible reduction process peak. Supposing that the current responsible for compound reduction may overlap with masking background current, we tried to characterise the reduction process by impedance method anyway within the range of possible reduction potential calculated from reference data about LUMO levels. Finally, no reduction process could be detected and characterised.

The narrow peak potential regions (**Figure 1 b, d**) of the D-A-D compounds enlisted in **Table 2** enabled an estimation of the IP and EA levels, as well as selection of the ranges for registering the impedance spectra. A potential range with a 0.2 V width including a redox peak was chosen and 21 spectra were obtained at 0.01 V increments.

The only reliable information we could certainly extract from CVs were estimated positions of the redox peaks and qualitative description of reversibility of the process. The non-equality of oxidation and reduction peaks observed for **III**, **IV** and **VI** (**Figure 1 b, d**) could not be explained by effect of one single factor. Voltammetric peaks are known to be affected by a set of inseparable factors such as number of electrons transferred, transfer coefficient (α), charge transfer (Faraday process) rate, diffusion of the species and side processes involving reactants, products and other solution components. Registration of characteristic peak was a tricky challenge since immediately after first reversible process all the compounds were found to undergo irreversible redox transformation

accompanied by electrode surface blocking and preventing consequent registration of a reverse scan.

A considerably high solution resistance was a problem usually concomitant with organic solvents. The effect was confirmed by impedance spectroscopy which allowed evaluation of system potential-independent resistance as will be shown below. That fact along with irreversibility disabled direct estimation of number of electrons transferred in one step from the peak potential difference, which equals 0.059/z (V) in case of reversible process and zero ohmic (*I*·*R*) potential drop in solution.

Fortunately impedance spectra analysis, potential dependence charge transfer resistance plots in particular, allowed determination of z value, which was found to be equal to 1 for all the processes considered. Moreover molecular structures of the compounds do not imply possibility of one-step oxidation to bication, that is relatively more stable than preceding radical-cation, as happens in case of symmetrical conjugated compounds (e.g. anthraquinone).

Because of the large number of investigated compounds and the similarity of their impedance spectra we will show and thoroughly discuss the spectra of oxidation of the popular donor **m**-**MTDATA**, reduction of acceptor and electron transfer material **TPBi** and both redox processes that involve emitter **V** (D-A-D compound) that is established as a promising component in the emission layer for OLEDs [10].

The CV of **m-MTDATA** showed a reversible oxidation peak at ca. 0.1 V and no obvious reduction process was observed (**Fig. 1 a**). However we will still consider voltage range -2.7 - -2.9 V as an example of treatment of irreversible process and attempt to describe kinetics of a process masked by high background current. During potential scanning in the positive direction in the 0.03 – 0.23 V range (**Fig. 2a,c**) both real and imaginary parts decrease until 0.08 V is reached and then start increasing until the end of the scan. Since both real and imaginary parts are minimal at the equilibrium potential, so is the impedance module.



Figure 2. Bode (a, b) and complex plane plots (c, d) of **m-MTDATA** impedance spectra in oxidation (a, c) and reduction (b, d) potential ranges. Impedance spectra were registered in increments of 0.01 V in the ranges +0.03 - +0.23 V (oxidation) and -2.7 - 2.9 V (reduction). The thick lines corresponds to oxidation equilibrium potential +0.08 V (a, c).

The Bode plot (**Fig. 2 a**) clearly shows minimization of the impedance at +0.08 V, i.e. at an equilibrium potential, as will be proved below. The phase shift is also minimal at this potential. However, this is not compulsory, as there are no definite kinetic reasons for such a behaviour of the phase shift. A double layer capacitance that affects the imaginary part is controlled by other independent processes. Minimization of the phase shift at the equilibrium potential was a coincidence observed for several compounds including **m-MTDATA**. The evolution of both values

of the real and imaginary impedance during potential scanning in the oxidation peak region is shown in **Figure 3**.



Figure 3. Real and imaginary impedance as functions of electrode potential corresponding to **m**-**MTDATA** reversible oxidation (Fig. 2 a, c) versus the potential at five selected frequencies.

As was proposed by CV analysis, the reduction of **m-MTDATA** is either irreversible or doesn't take place at all. During potential scanning in the negative direction in the -2.7 - 2.9 V range (**Fig. 2 b,d**) a consecutive decrease of impedance is observed without intermediate impedance minima.

The shown example depicts two ways of treatment stipulated by the reversibility of a process. When the process is purely reversible then there is a potential at which the rates of forward and backward redox processes are equal. Then the impedance is minimal and the equilibrium potential can be determined by both CV and EIS methods, the latter being a tool to estimate the rate constant. If the process is irreversible then only the forward electrochemical process is observed and its rate increases monotonically with the potential. In that case there is no distinctive point on the plot and the potential at the start of the current growing is regarded as the redox equilibrium potential. Nevertheless, the rate constant can still be determined from impedance data, but its value would be strongly dependent on the accepted value of the equilibrium potential.

The impedance spectra of **V** shown in **Figure 4** are typical of compounds with both reversible oxidation and reduction profiles. In both cases the impedance passes through a minimum as the potential is scanned in a peak interval. Bode plots (**Fig. 4b,d**) are useful to represent that kind of spectral evolution. The thick lines in both figures (**Fig. 4b,d**) correspond to the equilibrium potential that lies in the middle of the scanning interval. As stated above, the phase shift is not always minimal at the equilibrium potential. That means that minimums on the real and imaginary potential plots do not always coincide, because they may be affected by different processes that occur at the electrode-solution interface.



Figure 4. Bode plots (a, b) and complex plane plots (c, d) of impedance spectra of **V** in reduction (a, c) and oxidation (b, d) potential ranges. Impedance spectra were registered in increments of

0.01 V in the ranges +0.83 - +1.03 V (oxidation) and -2.05 - -2.25 V (reduction). The thick lines correspond to equilibrium potentials: -2.13 V for reduction (a, c) and +0.97 V for oxidation (b, d).

3.1 Impedance spectra analysis

An equivalent electrical circuit shown in **Figure 5** a was found to fit all the experimental spectra obtained in this work. In some cases its simplified modifications (**Fig. 5b,c**) could be used.



Figure 5. Equivalent electric circuits derived to fit the experimental spectra.

The proposed mechanism describing the equivalent circuit (**Fig. 5a**) is illustrated in **Figure 6**. Impedance corresponding to diffusion of a neutral molecule to the electrode surface is described by a specific Warburg element (W). The molecule approaching the electrode surface undergoes fast oxidation (as shown as an example in **Figure 6**) or reduction described by the charge transfer resistance (R_2). Consequently the oxidised molecule may undergo further oxidation or reduction. This stage does not determine the rate of the first oxidation process. Therefore, it is represented by a parallel charge path comprising charge transfer resistance (R_3). The other side-processes, such as solvent oxidation or reduction, contribute to the parallel charge transfer path. Since frequency responses of two parallel resistors are indistinguishable, all relatively slow charge transfer processes are described by one element (R_3). The element of capacitance (C) characterises a double electric layer. Its value is not related directly to the kinetics of the redox process so this element will not be considered in further discussions. The element of resistance (R_1) characterises charge transfer besides the electrode-electrolyte interface, i.e. solution and electrodes.



Figure 6. Correspondence of equivalent electric circuit to the proposed schematic mechanism of the electrochemical processes. Fast and slow stages are assigned by red and green arrows, respectively.

The rates of the processes (**Fig. 6**) determine their contribution to the total system impedance. If the rate of the fast reaction is very high relative to the rate of the parallel slow reaction, i.e. $R_2 \ll R_3$, then the R_3 element would hardly contribute to the system impedance and its value will be non-definable. In that case the equivalent circuit shown in **Figure 5b** would better characterise the system.

If the rate of charge transfer is small in comparison to the rate of diffusion then diffusion impedance is negligible in series connection with R_2 . If there is no Warburg element two resistors are no longer distinguishable and are represented by one element forming an equivalent circuit shown in **Fig. 5c.**

Figure 7 shows the dependencies of the parameters mentioned above on the potential for **m**-**MTDATA** oxidation. R_1 is not changing noticeably, as was expected, because this element is not related to the kinetics of the process. Dependencies of double layer capacitance (*C*), inverse charge

transfer resistance R_2^{-1} (i.e. charge transfer conductivity) and inverse Warburg constant A_W^{-1} have a characteristic peak at potential 0.08 V, indicating that at this potential the system impedance is minimal. We present inverted plots of the charge transfer resistance and the Warburg constant because in that case maximums are more evident than minimums in direct plots. The R_3 value was very high (more than $10^6 \Omega$) and in some points could not be determined with satisfactory precision.



Figure 7. Equivalent circuit parameters: R_1 (a), R_2^{-1} (b), C (c), A_W^{-1} (d), their errors and fitting theoretical curves (b) corresponding to **m-MTDATA** oxidation as functions of electrode potential.

Figure 8 shows the dependencies of the equivalent electrical circuit parameters calculated in a region of possible **m-MTDATA** reduction. This is a case of charge transfer limited reaction and the experimental spectra successfully fitted an equivalent circuit composed of three elements (**Fig. 5c**).

The charge transfer conductance (R_2^{-1}) rises exponentially as the potential is decreased. However it no considerable difference in case of presence and absence of investigated compound was detected. The charge transfer conductivity (**Fig. 5c**) in presence of MTDATA was only three times higher than the conductivity of solvent. This negligible difference is not sufficient for precise estimation of the redox constant. It may be caused by error of charge transfer or zero potential determination. This dependence will be applied to estimate the redox constant.



Figure 8. Equivalent circuit parameters: R_1 (a), C (b), R_2^{-1} (c), determined within the range -2.7 - 2.9 V in presence of **m-MTDATA**. A background curve of charge transfer conductivity obtained for electrolyte solution and theoretical fitting is shown (c).

Nevertheless, for a number of compounds a reduction behavior could be detected. **Figure 9** shows results of impedance analysis if TPBi reduction process.



Figure 9. Equivalent circuit parameters: R_1 (a), C (b), R_2^{-1} (c), A_W^{-1} (d), R_3^{-1} (e), their errors and fitting theoretical curves (c) corresponding to **TPBi** reduction as functions of electrode potential.

In case of **TPBi** the first equivalent circuit (**Fig. 5a**) was applied to describe . An additional parallel resistance R_3 (**Fig. 5a**) was necessary to describe side-processes involving reduction products and the solvent (**Fig. 6**).

Figures 10 and 11 present analogous plots of the equivalent circuit parameters characterising theoxidation and reduction of compound V, respectively.



Figure 10. Equivalent circuit parameters: R_1 (a), R_2^{-1} (b), C (c), A_W^{-1} (d) corresponding to **V** oxidation as functions of potential. Equivalent circuit and the corresponding element are shown in the insets.

In the case of V all the spectra fitted a simplified equivalent circuit shown in **Figure 5b** because of the considerable prevalence of reversible charge transfer rate in respect to the irreversible one.



Figure 11. Equivalent circuit parameters: R_1 (a), R_2^{-1} (b), C (c), A_W^{-1} (d) corresponding to **V** oxidation as functions of potential. Equivalent circuit and the corresponding element are shown in the insets.

3.2 Calculation of the standard rate constant from impedance data

The derivation of the formulas used in this work is presented in fundamental books devoted to electrochemical impedance spectroscopy and electrochemical kinetics [21-24]. The charge transfer resistance R_{ct} depends on electroactive species surface concentrations (c_0^S and c_R^S) and forward and backward rate constants (k_c and k_a) according to (1):

$$R_{\rm ct} = \frac{RT}{z^2 F^2 A} \frac{1}{\alpha k_{\rm c} c_{\rm O}^{\rm S} + (1 - \alpha) k_{\rm a} c_{\rm R}^{\rm S}},\tag{1}$$

where α is charge transfer coefficient (0 < α < 1), *A* – electrode surface area. Rate constants are functions of potential according to the Butler-Volmer equation (2).

$$k_{\rm c} = k^0 \cdot \mathrm{e}^{-\alpha} \frac{zF}{RT} (E - E^0) \qquad (2a) \qquad \qquad k_{\rm a} = k^0 \cdot \mathrm{e}^{\left(1 - \alpha\right)} \frac{zF}{RT} (E - E_0) \qquad (2b)$$

where k^0 is the standard rate constant, E_0 is equilibrium potential, E is electrode potential. Stationary surface concentrations can be found from solution of Fick's partial differential equation [24].

$$c_{\rm O}^{\rm s} = \theta \frac{\xi c_{\rm O}^{\rm b} + c_{\rm R}^{\rm b}}{1 + \xi \theta}$$
 (3a) $c_{\rm O}^{\rm s} = \frac{\xi c_{\rm O}^{\rm b} + c_{\rm R}^{\rm b}}{1 + \xi \theta}$ (3b)

where c_0^{b} , c_R^{b} are bulk concentrations, ξ is a ratio of diffusion coefficients (D_0 and D_R) (4) and θ is potential exponential function (5).

$$\xi = \sqrt{\frac{D_{\rm O}}{D_{\rm R}}} \tag{4}$$

$$\theta = e^{\frac{zF}{RT}(E - E_0)} \tag{5}$$

Substituting (2) - (5) into (1) one gets (6)

$$R_{\rm ct}^{-1} = k^0 \frac{z^2 F^2 c_{\Sigma}}{RT} \frac{\theta^{1-\alpha}}{1+\theta}$$
(6)

A plot of the function (6) is a bell curve having the same form as the experimental points (**Fig. 7b**, **Fig. 9b**, **Fig. 10b**). Theoretical function (6) was fitted to the experimental data by tuning values of E_0 , k^0 and α ; the latter was accepted as being equal 0.5. It allowed adjustment of the equilibrium potential E_0 and an estimation of standard electrochemical rate constant k^0 . Potential E_0 is easily determined as a potential at the maximum on the R_2^{-1} -E plot (**Fig. 7b**, **Fig. 9c**, **Fig. 10b**, **Fig. 11b**).

Considering graph of the theoretical function (6) and its parameters one can easily describe effect of each of them on the form of a graph. Changing E_0 shifts the whole curve along abscises scale. Changing k^0 controls height of the peak. Changing z would cause modification of peak sharpness as demonstrated in **Figure 7b**. Coefficient α regulates symmetry of the peak (at $\alpha = 0.5$ both peak shoulders are equal). An attempt to fit all four parameters with experimental data lead to non-realistic values of α due to non-ideality of real electrochemical system. For that reason α was fixed at 0.5. The value of *z*, when calculated from fitting results, ranged from 0.9 to 1.1. Therefore it was also fixed to be 1.0, two parameters E_0 and k^0 remaining unknown and easy to be calculated.

If the reaction is irreversible then the functions (1) and (6) are modified taking into account the presence of a sole process. It resulted in omitting one of the terms in the denominator in formula (6) (either θ or 1). Thus formula (6) was transformed to (7) and (8) for oxidation and reduction processes, respectively. E.g. formula (8) was used for fitting experimental data in **Fig. 8 c**.

$$R_{\rm ct}^{-1} = k^0 \frac{z^2 F^2 c_{\Sigma}}{RT} \theta^{1-\alpha} \tag{7}$$

$$R_{\rm ct}^{-1} = k^0 \frac{z^2 F^2 c_{\Sigma}}{RT} \theta^{-\alpha}$$
(8)

The calculated values of charge transfer rate constants of all the investigated substances are presented in **Tables 1** and **2**.

Obviously rate constant estimated from formula (8) is strongly dependent on the value of equilibrium potential. In several cases a value calculated from the reference LUMO data can be used. The reference investigation of a solution containing only electrolyte was carried out in order to estimate a minimal limit of measurable kinetic constant. **Figures 8c** and **9e** show charge transfer conductivity plots overlapped with plots obtained in presence of investigated compounds. The apparent redox rate constant referred to the solvent was found to be $4.3 \cdot 10^{-6}$ if the reduction potential assumed to be -2.9 V. We regarded that value as a limit for distinguishing the response of a process involving the investigated molecule from a background signal. Since there was no evidence of occurring redox process one can regard it neither not to take place or to be very slow (rate constant < $1 \cdot 10^{-5}$) to be detected by the impedance method.

On the other hand, rate constants of a very fast charge transfer reaction could not be estimated precisely because in that case the reaction was limited by diffusion. Charge transfer resistance was very small in comparison to diffusion impedance. The higher limit of possible redox constant value was estimated to be $1 \cdot 10^{-2}$. For that reason values which are estimated to be very high are shown in **Table 2** as > $1 \cdot 10^{-2}$. **Figure 12** presents the resulting data from **Tables 1** and **2** in a graphical mode. Negative decimal logarithms (pk = -log(k)) of reduction and oxidation rate constants correspond to abscissa and ordinate of the points, respectively. The approximate regions corresponding to different roles of the compounds are shown on the graph by coloured rectangles.



Figure 12. Comparison of redox standard rate constants of the investigated compounds: hole, electron-transport and host materials (Table 1), intra-molecular exciplex emitters (Table 2). Coloured rectangles designate areas corresponding to roles of the compounds play in diodes.

Kinetic constants that could not be determined were attributed values of $1 \cdot 10^{-5} \text{ m} \cdot \text{s}^{-1}$ to depict on a graph as a minimal value that could be estimated for compound in solution. The real value of a constant must be lower so that the point related to the compound would find it self in the corresponding rectangle region (**Fig. 12**). All the compounds including both donor and acceptor molecule units (**Table 2**) are characterised by much higher both oxidation and reduction rate constants than the other pure donor or acceptor compounds (**Table 1**). Rate constants of the D-A-D compounds exceed the rate constants of single donor or acceptor molecules (**Table 1**) by 2-3 orders. Thus the presence of both donor and acceptor parts together in one molecule enhances the molecule's ability to exchange charge rapidly in both oxidation and reduction directions. Secondly, the ratios of oxidation and reduction rate constants of hole- and electron-transport compounds correlate with their conductivity types although the redox rate constant and the charge carrier mobility refer to different phases (solution and solid). This correlation may be used for prediction of the behaviour of a compound in the solid state based on its behaviour in solution. If a compound is oxidised faster than it is reduced, then charge in the solid state would be predominantly transferred by reversible oxidation, i.e. by M^+/M -states (M = molecule), which means hole-type transport. The opposite case, where charge is predominantly exchanged due to reversible reduction, i.e. between M/M⁻-state, means electron-type transport. Oxidation rate constants of hole-transport materials were found to be higher than oxidation rate constants of electron-transport materials (Fig. 12, Table 1). The same statement can be made in relation to the electron-transport materials and reduction rate constants. The other feature which deserves attention is the very low values of the reduction rate constants. Their values are not only very small, but also they do not differ significantly even between hole- and electron-transport compounds. This means that all the organic compounds under investigation reduce more slowly than they oxidise. This result corresponds to a known fact that hole mobility in organic solid films is in most cases much higher than electron mobility [25, 26]. Finally, the compounds used as host materials were found to be oxidised and reduced very slowly and irreversibly. Therefore, their rate constants could not be estimated.

4. Conclusions

The choice of a compound for a specific role in an OLED is predominantly based on its relative orbital energy levels and its emission spectrum. In this work we have thrown light upon another important aspect – namely, the conductivity properties of an organic material and an easy way to predict these properties by analysis in solution.

The proposed impedance-based technique has proved to be effective for the characterisation of thermodynamic (HOMO and LUMO levels) and kinetic (redox rate constants) parameters of a wide variety of molecules used in OLEDs. Although no direct relationship between structure and numerical data could be revealed, several particular aspects were elucidated:

- (i) the redox rate of ambipolar compounds containing donor and acceptor parts appeared much higher than those of pure donors and acceptors. The rise of rate is very important for organic electronics and electrochromic devices. In the electrochromic devices, it will decrease switching time, current and the possibility of degradation. In the organic electronic devices, it may influence change balance and recombination rate within the active component;
- (ii) the relationship between oxidation and reduction rate constants is responsible for the compound's conductivity type in the solid state and is expected to be an easy tool to predict compound performance when incorporated in an OLED-layer.

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SUPPORTING INFORMATION.

Details of the synthesis and characterisation of compounds **I** and **III**. Electrochemical impedance spectroscopy analysis of ferrocene oxidation kinetics.

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Supporting information for Determination of standard redox rate constants of OLED active compounds by electrochemical impedance spectroscopy

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General experimental details

All reactions were carried out under an argon atmosphere unless otherwise stated. Starting materials were purchased commercially and were used as received. Solvents were dried using an Innovative Technology solvent purification system and were stored in ampoules under argon. Silica gel column chromatography was performed using silica gel 60 purchased from Sigma Aldrich. ¹H and ¹³C NMR spectroscopy was carried out on a Bruker AV400 NMR spectrometer. NMR data was processed in MestReNova V10.Melting points were carried out on a Stuart SMP40 machine with a ramping rate of 4 °C min⁻¹. Videos were replayed manually to accurately determine the melting point. High resolution mass spectrometry was carried out on a Waters LCT Premier XE instrument using ASAP ionisation. Samples were analysed directly as solids using N₂ at 350 °C. Elemental analysis was performed on an Exeter Analytical E-440 machine.

Synthesis of 2,8-bis(phenoxazin-N-yl)dibenzothiophene-S,S-dioxide (I)



The above procedure for III, below, was followed using phenoxazine (516 mg, 2.82 mmol, 2.1 eq.) instead of dibenzazepine. The product (I) was obtained as a yellow solid (669 mg, 86% yield).

¹H NMR (400 MHz, CDCl₃) δ 8.11 (d, J = 8.1 Hz, 2H), 7.80 (d, J = 1.7 Hz, 2H), 7.62 (dd, J = 8.1, 1.7 Hz, 2H), 6.77 – 6.70 (m, 8H), 6.67 – 6.63 (m, 4H), 6.02 – 6.00 (m, 4H); ¹³C NMR (100 MHz, CDCl₃): δ 145.7, 144.2, 137.5, 134.3, 134.0, 133.3, 125.3, 124.9, 123.6, 122.6, 116.2, 113.7; HRMS-ASAP⁺ m/z calculated for C₃₆H₂₂N₂O₄S [M]⁺ 578.1300, found: 578.1301; Anal. Calc. for C₃₆H₂₂N₂O₄S C: 74.73 H: 3.83 N: 4.84, Found: C:74.79 H: 3.82 N: 4.81; m.p. decomp. > 350 °C. An alternative synthesis has been reported [1].

Synthesis of 2,8-Bis(N-dibenzazepin-N-yl)dibenzothiophene-S,S-dioxide (III)



30% Yield

2,8-Dibromo-dibenzothiophene-S,S-dioxide (505 mg, 1.35 mmol, 1 eq.), dibenzazepine (545 mg, 2.82 mmol, 2.1 eq.), $Pd_2(dba)_3$ (80 mg, 87 µmol, 0.065 eq.) and XPhos (83 mg, 174 µmol, 0.13 eq.) were dissolved in dry toluene (20 mL) and were bubbled with argon for 30 minutes, then NaO^tBu

(284 mg, 2.96 mmol, 2.2 eq.) was added and reaction mixture degassed further 15 minutes. The reaction was stirred for 18 h at 110 °C. Upon cooling, reaction was diluted with toluene (50 mL) and washed with water, dried over Na₂SO₄, filtered and solvent removed under reduced pressure. The residue was purified by silica gel column chromatography using CH₂Cl₂:hexane (4:1 v/v) as eluent. Methanol (3 mL) was added to the off-white solid and the liquid was discarded with a pipette to yield product (**III**) (242 mg, 30% yield).

¹H-NMR (400 MHz, DMSO-d₆) δ 7.66 – 7.51 (m, 16H), 7.44 (d, J = 8.7 Hz, 2H), 6.95 (s, 4H), 6.18 (dd, J = 8.7 Hz, 2.3 Hz, 2H), 5.87 (d, J = 2.3 Hz, 2H); ¹³C NMR (100 MHz, DMSO-d₆) δ 152.8, 140.6, 135.1, 131.7, 130.7, 130.3, 130.2, 129.3, 128.2, 127.1, 122.8, 112.3, 101.9; HRMS-ASAP⁺ m/z calculated for C₄₀H₂₆N₂O₂S [M]⁺ 598.1715, found: 598.1728; Anal. Calc. for C₄₀H₂₆N₂O₂S · 1.2 CH₂Cl₂ C: 79.45 H: 4.36 N: 4.62, Found: C:79.29 H: 4.34 N: 4.59.; m.p. decomp. > 350 °C.



Figure S1. ¹H NMR spectrum of **I** in CDCl₃.





Figure S2. ¹³C NMR spectrum of **I** in CDCl₃.



Figure S3. ¹H NMR spectrum of **III** in CDCl₃.



Figure S4. ¹³C NMR spectrum of **III** in CDCl₃.

The method presented in the article can be successfully applied to characterisation of kinetics of reversible and quasi-reversible electrochemical reactions involving organic compounds. Here we demonstrate application of the technique for estimation of redox rate constant of ferrocene, a compound available in most electrochemistry laboratories. The procedure which results are presented here can be used for testing and didactic purposes.

Figure S5 shows sets of impedance spectra of ferrocene within the range of its oxidation potential region.



Figure S5. Bode (a) and complex plane plots (b) of 0.01 M $Fe(C_5H_5)_2$ dichloromethane solution impedance spectra in the range -0.30 - +0.30 V in increments of 0.02 V. The thick lines correspond to starting (solid) and finishing (dashed) scan potentials. Arrows show tendency of spectra change before and after equilibrium potential (0.00 V).

Figure S6 shows results of spectra analysis by impedance spectroscopy along with corresponding cyclic voltammograms. Two concentrations (0.01 M and 0.05 M) were used in order to show independence of the impedance method on compound concentration.



Figure S6. Cyclic voltammograms (a), inverse Warburg constant (b), inverse charge transfer resistance (c, d), capacitance (e) and solution resistance (f) for two different ferrocene concentrations (shown in legend) as functions of electrode potential. Theoretical fitting of charge transfer resistance function is demonstrated in (c) and (d).

CVs (**Fig. S6a**) are very sensitive to compound concentration and selection of the optimal concentration may be required to obtain a characteristic CV curve. However, the numerous CVs experiments are not useful, especially when compounds are expensive and limited in amount. **Figure**

S6b presents inverse Warburg constant potential dependence that characterises diffusion impedance which is minimal at the equilibrium potential and is inversely proportional to the electroactive compound concentration. Charge transfer resistance (**Fig. S6c,d**) is the main parameter used for kinetic characterisation. The redox rate constant value estimated from the potential dependencies was $2.58 \cdot 10^{-5} \text{ m} \cdot \text{s}^{-1}$ (**Fig. S6c**) and $2.24 \cdot 10^{-5} \text{ m} \cdot \text{s}^{-1}$ (**Fig. S6d**). The capacitance potential dependence (**Fig. S6e**) is affected by adsorbtion processes and cannot be used to extract any information about kinetics. The resistance R_1 (**Fig. S6f**) is not dependent on potential. Despite being not reproducible this value still can be used for estimation of solution resistance which appears to be considerably high in case of organic solvent.

Reference for SI.

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