Dicyanobenzene and dicyanopyrazine derived X-shaped charge-transfer chromophores: comparative and structure–property relationship study†

L. Dokládalová,a† F. Bureš,a,b W. Kuznik,b,c I. V. Kityk,c A. Wojciechowski,c T. Mikysek,d N. Almonasy,a Z. Padělková,e J. Kulháneka and M. Ludwig a

A series of novel X-shaped push–pull compounds based on benzene-1,2-dicarbonitrile has been designed, synthesized and further investigated by X-ray analysis, electrochemistry, absorption and emission spectra, SHG experiment and quantum-chemical calculations. The obtained data were compared with those for isolobal 5,6-disubstituted pyrazine-2,3-dicarbonitriles. Structure–property relationships were elucidated. The extension, composition and planarization of the π-linker used as well as the electronic-withdrawing ability of both dicyano-substituted acceptor units affect the linear and nonlinear properties of the target charge-transfer chromophores most significantly.

Introduction

Organic molecules based on the sp2/sp hybridized carbon/hetero atom scaffold end-capped with electron donors and acceptors represent an intensively investigated area of organic chemistry. In these molecules, intramolecular charge-transfer (ICT) from the donor (D) to the acceptor (A) via a π-conjugated system takes place and the D-π-A systems become polarized and gain dipolar character. Due to the ICT, push–pull D-π-A molecules possess distinct linear as well as nonlinear optical (NLO) properties. Charge-transfer chromophores have found widespread applications in modern branches of materials chemistry such as optoelectronics, data processing and storage devices, NLO bioimaging, organic light-emitting diodes (OLEDs), dye-sensitizing solar cells (DSSCs), semiconductors, switches, etc.1–3 Spatial arrangement of a push–pull system involves prevailing linear (D-π-A), quadrupolar (D-π-A-π-D or A-π-D-π-A) and octupolar/tripodal ((D-π)3-A or (A-π)3-D) systems or less common V- (ref. 4), Y- (ref. 5), H- (ref. 6) and X-shaped molecules. X-shaped chromophores based on tetra-substituted ethenes (tetraethynylethenes, TEEs, and cyanooethynylethenes, CEEs) were introduced and extensively studied mainly by Diederich and co-workers.7,8 Besides the ethene central π-linker, the 1,2,4,5-tetrasubstituted benzene pattern represents another class of X-shaped chromophores. Benzene-derived X-shaped chromophores can conveniently be prepared from catechol, o-phenylenediamine or tetrahalogenated benzenes via functionalization with suitable electron donors (e.g. NR2 and OR groups) and acceptors (e.g. NO2 group and π-deficient heterocycles). Such molecules were mainly investigated for their unique absorption/emission properties and chelating abilities towards metal ions.9

Recently, we have designed and synthesized X-shaped chromophores based on the pyrazine skeleton.10,11 Herein we would extend the concept of X-shaped chromophores with two electron withdrawing cyano groups saturated with two electron donors attached opposite to a six-membered (hetero)aromatic. Thus, the previous series of pyrazines P1–P10 was completed with analogous push–pull chromophores B1–B10 featuring the benzene central π-system.

Both chromophore types possess a N,N-dimethylamino donor connected via systematically extended π-linkers that comprise a combination of up to three acetylene and 1,4-phenylene subunits. In contrast to polyenes and polynyes,
(oligo)phenylene ethynylene charge-transfer chromophores possess generally higher thermal and (photo)chemical stability. 4,5-Disubstituted benzene-1,2-dicarbonitrile derivatives are a currently intensively investigated class of compounds due to their easy transformation to (sub)phthalocyanines. Peripherally substituted phthalocyanines found widespread applications in NLO, DSSCs, supramolecular assemblies, discotic liquid crystals, and photodynamic therapy. Recently, Adachi et al. showed highly efficient OLEDs from delayed fluorescence that utilize similar push-pull dicyanobenzene derivatives functionalized with carbazole moieties.}

We report here the design and synthesis of charge-transfer chromophore series B1–B10 with a systematically evaluated π-conjugated path and the elucidation of the structure–property relationships. The extent of the ICT and linear and non-linear optical properties were studied by X-ray analysis, electrochemistry, absorption/emission spectra, semi-empirical computational methods and second harmonic generation (SHG) experiments.

**Results and discussion**

**Synthesis**

The synthesis of target chromophores B1–B10 has been carried out in a modular manner starting from 4,5-diiodobenzene-1,2-dicarbonitrile 11, which can be prepared from commercially available phthalimide. In contrast to Terekhov et al., iodination of phthalimide with I2/oleum (30%) and one recrystallization from acetone afforded directly 4,5-diiodophthalimide in 61% yield without the need for Soxlet extraction and column chromatography. However, it should be noted that regioselective outcome and yield of the iodination step depend strongly on the concentration of the oleum used. Subsequent ammonolysis to 4,5-diiodophthaldiamide and dehydration with (CF3CO)2O/pyridine afforded compound 11 in 42% overall yield (see the ESI† for more details). Similarly to pyrazines, P1–P10, benzene derivatives B1–B10 were obtained by Suzuki–Miyaura and Sonogashira cross-coupling reactions. N,N-Dimethylaminosubstituted π-linkers with boronic acid (ester) function and terminal acetylene moiety were prepared from commercially available or known short π-linkers and their extension/alternation by cross-coupling reactions with [[4-iodophenylethynyl(trimethylsilane). With the extended π-linkers and 4,5-diiodobenzene-1,2-dicarbonitrile 11 in hand, we have carried out twofold Suzuki–Miyaura cross-coupling reactions leading to target molecules B1, B3 and B4 in yields of 83, 87 and 79% (Scheme 1). These chromophores possess the π-linker connected directly to the dicyanobenzene moiety via a 1,4-phenylene unit. This twisted arrangement usually represents a barrier of efficient D–A interaction. Therefore, partial planarization of the π-linker has been achieved via introduction of an additional acetylene unit by Sonogashira reaction. The reaction of 11 with terminal acetylens afforded target chromophores B2 and B5–B10 with the yields ranging from 62 to 92% (Scheme 2). All target compounds B1–B10 were isolated in similar yields as pyrazines P1–P10 and were purified by column chromatography and recrystallization from 1,2-dichloroethane/hexane. In contrast to compounds B7 and B8, pyrazine analogues P7 and P8 could not be prepared. In these cases, the final Sonogashira reactions were dominated by homocoupling of the starting acetylenes and pyrazine derivatives P7 and P8 were only detected in the crude reaction mixture as minor products. Due to the dipolar character and extended π-system of B1–B10, some target compounds are only sparingly soluble in halogenated solvents. However, their...
Scheme 2  Twofold Sonogashira reaction leading to chromophores B2 and B5–B10.
chemical structures have been confirmed by NMR, HR-MALDI-MS and IR spectra as well as elemental analysis and X-ray analysis.

**Crystallography**

Slow diffusion of hexane into solution of chromophores B1 and B4 in CH₂Cl₂/CDCl₃ afforded crystals suitable for X-ray analysis. Fig. 1(a/b) shows the ORTEP plots and side views of both molecules and confirms the proposed molecular structure. The ORTEP plot of previously crystallized pyrazine P1 has also been included as a reference (Fig. 1c). The spatial arrangement of the central dicyano acceptor moieties and the adjacent 1,4-phenylene units in B1, B4 and P1 can be evaluated by the dihedral angles \( \phi \) as shown in Fig. 1 and Table 1.

As can be seen, both donor-substituted \( \pi \)-linkers attached on the benzene-1,2-dicarbonitrile acceptor moiety via a 1,4-phenylene unit are significantly twisted out of the chromophore molecular plane. Pyrazine P1 adopts a similar angular arrangement which is most likely caused by the steric repulsion of the ortho-hydrogens. Whereas in benzene derivatives B1/B4 are both 1,4-phenylene moieties twisted perpendicularly to the adjacent benzene-1,2-dicarbonitrile moiety and adopt mutually almost parallel arrangement with minimal \( \phi_1 \) and high torsion angles \( \phi_3/\phi_6 = 42°-51° \), the derivative P1 possesses these substituents twisted mainly above/below the pyrazine plane with relatively high both torsion angles \( \phi_1 = 25° \) and \( \phi_3/\phi_6 = 30°/29° \). This arrangement results in substantial deformation of the pyrazine ring compared to almost planar benzene rings (see \( \phi_2-4 \) in Table 1). The differences in spatial arrangements of the C4/C5 and C5/C6 substituents in benzene B1/B4 and pyrazine P1 derivatives are visualized in the side views (Fig. 1). The overlap wire model for isolobal compounds B1/P1 is also provided in Fig. 1d.

Bond length alternation (BLA) in the \( N,N \)-dimethylanilino (DMA) and 1,4-phenylene rings can easily be evaluated by the quinoid character \( (\delta r) \). Whereas in benzene, \( \delta r \) is equal to 0, in a fully quinoid ring, \( \delta r \) would be on the order of 0.100–0.120 Å. A comparison with known CT chromophores bearing DMA moieties such as CEEs \( (\delta r \text{ up to 0.070 Å}) \) and TEEs \( (\delta r \text{ up to 0.025 Å}) \) places the nonplanar benzene and pyrazine chromophores B1/B4 and P1 below/at the level of donor-substituted tetraethylenes. Further comparison of the \( \delta r \) values for B1/P1 shown in Table 1 clearly indicates higher BLA in the pyrazine chromophore P1. Estimation of the aromaticity of both central benzene and pyrazine rings by the calculation of the Bird index \( (I_b) \) provides similar insight. Whereas the \( I_b \) values of unsubstituted benzene/pyrazine are 100/88.8, the calculated values for B1, B4 and P1 decrease in the same order of 89.9→88.9→82.2 as a consequence of the \( \pi \)-system extension and replacement of the central benzene by the pyrazine ring.

**1H NMR and IR spectroscopy**

We analysed the \(^{13}C\) NMR spectra of all B and P chromophores measured in CDCl₃ (100 MHz) as well as IR spectra measured neat using an HATR adapter. Electron saturation of the CN group indicated by \(^{13}C\) NMR chemical shift and frequency of vibrational stretching would serve as good indicators of the ICT extent (Table 2).

The chemical shift of C atoms strongly depends on hybridization and the electron density at the nucleus. The higher the electron density, the more shielding occurs and an upfield shift will be observed. The data in Table 2 clearly demonstrate that \(^{13}C\) NMR shifts of the CN group in pyrazine series of compounds (P1–P4) are generally upfielded as a consequence of higher saturation of the CN group compared to compounds B1–B4. Introduction of more electronegative acetylene units as in chromophores B6–B10 reduces the electron saturation of the CN group, which is then engaged similarly in the ICT, and the chemical shifts became steady at around 114 ppm. However, when comparing the particular compounds B1/B2 or P1/P2 that differ in the type of connection of the \( \pi \)-linker to the acceptor moiety (without/with an additional acetylene unit), the chemical shifts of the latter were upfielded.

---

**Table 1** RTG-derived structural parameters of chromophores B1, B4 and P1

<table>
<thead>
<tr>
<th>Compd</th>
<th>( \phi_1 )</th>
<th>( \phi_2 )</th>
<th>( \phi_3 )</th>
<th>( \phi_4 )</th>
<th>( \phi_5/\phi_6 )</th>
<th>( \delta r )</th>
<th>( I_b )</th>
</tr>
</thead>
<tbody>
<tr>
<td>B1</td>
<td>7</td>
<td>2</td>
<td>2</td>
<td>3</td>
<td>51/45</td>
<td>0.011</td>
<td>89.9</td>
</tr>
<tr>
<td>B4</td>
<td>10</td>
<td>5</td>
<td>4</td>
<td>4</td>
<td>48/42</td>
<td>0.010/0.013</td>
<td>88.9</td>
</tr>
<tr>
<td>P1</td>
<td>25</td>
<td>18</td>
<td>13</td>
<td>7</td>
<td>30/29</td>
<td>0.027</td>
<td>82.2</td>
</tr>
</tbody>
</table>

\( ^{a} \) The torsion angles \( \phi \) are in deg. \( ^{b} \) The average quinoid characters \( \delta r \) of the DMA units are given in Å. \( ^{c} \) The Bird index of the central benzene/pyrazine rings.

---

Fig. 1 ORTEP representations and spatial arrangements of chromophores B1 (a), B4 (b) and P1 (c) measured at 150 K and the overlap wire model for isolobal chromophores B1/P1 (d). The thermal ellipsoids are shown at 50% probability with arbitrary spheres for H atoms.
by 3.68 and 2.16 ppm. This observation can be attributed to planarization of the entire π-system and thus resulting better D–A interaction in B2/P2. The chemical shifts δ(C=N) = 112–116 ppm measured for the dicyanobenzene and dicyanopyrazine acceptor moieties are slightly downfielded compared to donor substituted CEEs (δ(C=N) = 108–115 ppm) which further confirms the aforementioned trends seen by the BLA.

The frequency of the stretching vibration of the CN group varied very similar trends, as seen by the NMR. Namely, the C=N stretching appeared generally at lower frequencies for pyrazine series. Extension and (partial) planarization of the π-linker (e.g. 1 vs. 2 or 3 vs. 4) shifted the CN wavenumber to higher energy and became steady at around 2200/2190 cm⁻¹.

### Table 3 ¹³C NMR chemical shifts and frequency of the stretching vibration of the C=N group

<table>
<thead>
<tr>
<th>Compd</th>
<th>δ(C=N)¹</th>
<th>ν(C=N)²</th>
</tr>
</thead>
<tbody>
<tr>
<td>B1/P1</td>
<td>116.27/114.33</td>
<td>2223/2230</td>
</tr>
<tr>
<td>B2/P2</td>
<td>112.59/112.17</td>
<td>2177/2165</td>
</tr>
<tr>
<td>B3/P3</td>
<td>114.19/113.68</td>
<td>2224/2340</td>
</tr>
<tr>
<td>B4/P4</td>
<td>114.62/113.43</td>
<td>2202/2194</td>
</tr>
<tr>
<td>B5/P5</td>
<td>113.83³</td>
<td>2200/2182</td>
</tr>
<tr>
<td>B6/P6</td>
<td>114.29³</td>
<td>2206/2189</td>
</tr>
<tr>
<td>B7/P7</td>
<td>114.29³</td>
<td>2203/—</td>
</tr>
<tr>
<td>B8/P8</td>
<td>114.41/—</td>
<td>2202/—</td>
</tr>
<tr>
<td>B9/P9</td>
<td>114.21³</td>
<td>2202/2188</td>
</tr>
<tr>
<td>B10/P10</td>
<td>114.47³</td>
<td>2202/2190</td>
</tr>
</tbody>
</table>

¹¹³C NMR chemical shift of the CN group given in ppm – measured in CDCl₃ at 100 MHz. ² Frequency of the vibrational stretching of the CN group given in cm⁻¹ – measured by IR (neat, HATR). ³ Not observed due to sparsing solubility in CDCl₃.

### Electrochemistry

Electrochemical measurements of all chromophores B1–B10 and P1–P10 (ref. 10) were carried out under standard conditions (N,N-dimethylformamide containing 0.1 M Bu₄NPF₆) in a three electrode cell by cyclic voltammetry (CV) and rotating disk voltammetry (RDV). The working electrode was a platinum disc (2 mm in diameter) for CV and RDV experiments. As the reference and auxiliary electrodes were used a saturated calomel electrode (SCE) separated by a bridge filled with a supporting electrolyte and a Pt wire, respectively. The acquired data are summarized in Table 3, and representative CV diagrams of chromophores B2 and B10 are shown in the ESI (Fig. S1–S2†).

All chromophores 1–10 in both series B and P showed two close, mostly reversible, one-electron oxidation processes (differences of about 100 mV measured by CV) that were, in most cases, merged into one two-electron wave, as indicated by RDV. This electrochemical behavior reflected the presence of two equivalent N,N-dimethylamino donor groups appended to the dicyanobenzene/pyrazine acceptor. The first oxidation potentials of chromophores in both series B and P range from +0.93 to +0.78 V and +1.02 to 0.81 V, respectively. As can be seen, extension of the π-conjugated path affects the first reduction potentials only negligibly. However, the E°(oxid) values slightly decrease with the extension of the π-system as a result of the gradual N,N-dimethylamino donor isolation. From chromophores B3/P3 onwards, the first oxidation potentials were concentrated in the range of about +0.80 V, and therefore, the N,N-dimethylamino donor in chromophores 4–10 can be considered as similarly involved in the ICT.

In contrast to the first oxidations, the first reductions of chromophores in the series B and P were observed in a wide range from −1.62 to −1.03 and −1.24 to −0.67 V, respectively. The difference between the first reduction potentials of the particular chromophores in both B and P series ranges from 0.36 to 0.44 V. This reflects a higher withdrawing ability of the dicyanopyrazine moiety over the dicyanobenzene caused by the presence of two electronnegative nitrogen atoms. However, when considering the dicyanobenzene/pyrazine acceptor as the main reduction center and the same electron nature and number of appended donors, the observed differences within the particular series B or P must be elucidated as a consequence of the π-linker. The first reduction potentials of the chromophores in the series B or P gradually decrease and become steady at about −1.10 and −0.70 V. When going from the chromophore B1/P1 (one 1,4-phenylene moiety as a π-linker) to B3/P3 (biphenyl π-linker) and B4/P4 (phenylethynylphenyl π-linker), the first reduction potentials decrease from −1.62/−1.24 to −1.48/−1.05 and −1.41/−0.97 V as a result of the π-linker gradual extension. However, chromophores B2/P2 bearing a shorter π-linker showed more positively shifted E°(red) up to −1.28/−0.87 V than extended chromophores B3/P3 and B4/P4. This can be attributed to partial extension of the π-linker by an acetylene unit connected directly to the acceptor and, in particular, to planarization of the entire π-systems. A similar effect can be observed when comparing chromophores B3/P3 and B5/P5 or B4/P4 and B6/P6. Hence, a separation of the π-linker from the dicyanobenzene/pyrazine acceptor by the linear ethynyl spacer plays a very crucial role and affects the electrochemical behavior of X-shaped chromophores considerably. In contrast to this, the first reduction of the chromophore B7 (E°(red) = −1.32 V) is significantly impeded by the propeller-shape of the terphenyl linker. Except B7, chromophores B6/P6–B10/P10 showed almost constant first reduction potentials.
which implies that a combination of two/three 1,4-phenylene moieties in conjunction with acetylene units caused complete isolation of the N,N-dimethylamino donor and the dicyanobenzene/pyrazine acceptor.

The corresponding electrochemical gaps ΔE (Table 3) are mainly affected by the variation in the first reduction potentials and, therefore, mimic the same trends as seen for E_{(red)}^{0}. The lowest electrochemical gaps were recorded for chromophores B10/P10 with the largest and fully planar π-linkers (see also the energy level diagram below).

UV/Vis spectroscopy

Optical properties of all chromophores 1–10 were studied by absorption and emission spectra. The positions of the longest-wavelength absorption and fluorescence maxima λ_{max} and λ_{max}^F as well as the molar absorption coefficients ε and the fluorescence quantum yields q^F are given in Table 4. Representative UV/Vis spectra of selected chromophores are shown in Fig. 2; for complete spectra listing see the ESI (Fig. S3–S4†). All spectra were measured in CH₂Cl₂.

The absorption spectra of chromophores in series B and P showed longest-wavelength absorption maxima (CT-bands) appearing within the range of 335–458 and 354–499 nm, respectively. The absorption spectra of representative B and P chromophores shown in Fig. 2 are very similar. However, the absorption maxima of P chromophores are generally bathochromically shifted, which again confirms the previous conclusion on higher electron withdrawing character of the dicyanopyrazine moiety. Chromophores B1/P1 and B2/P2 showed multiple CT-bands which reflects complex conjugated pathways between the particular N,N-dimethylamino donor and cyano acceptor. The spectra of chromophores B feature more vibronic structure than those of chromophores P. When going from chromophore B1/P1 to B2/P2, thus extending and planarizing the π-linker by one acetylene unit, the absorption maxima were shifted by 48 and 28 nm (Table 4), respectively. However, moving the acetylene unit between two 1,4-phenylene units as in B4/P4 or further π-linker extension as in B6/P6 and B10/P10 led to disappearance of the CT-band and a pronounced hypsochromic shift. This is in accordance with the observation made by Nielsen et al.22 that an extension of the π-system by more nonplanarly arranged 1,4-phenylene units led to a blue-shift of the CT-band measured in a solvent. However, this is also in contrast to gas phase measurement excluding solvation and H-bonding in which a systematic red-shift was observed. Hence, from B7 onwards, the CT-bands were observed as hardly distinguishable shoulders and were merged with the bands appearing at around 350 nm with rising intensity. These bands probably correspond to the transition between the N,N-dimethylamino donor and the extended electronegative π-linker. Thus, insertion of three 1,4-phenylene units in combination with one or more acetylene units can completely quench D–A Interaction and none or very weak CT-band is observed. These observations are in accordance with the aforementioned electrochemical conclusions.

In contrast to all N,N-dimethylamino-substituted π-linkers, which showed emission properties,19 only less extended chromophores B1–B4 showed fluorescent behaviour (Table 4, the ESI Fig. S5†). Moreover, the pyrazine chromophores did not show any fluorescence at all. Thus, extension of the π-system or attachment of a stronger acceptor moiety caused emission disappearance and a gradual reduction of the fluorescence quantum yield.

<table>
<thead>
<tr>
<th>Compd</th>
<th>λ_{max} [nm(eV)]</th>
<th>ε [10³ mol⁻¹ dm³ cm⁻¹]</th>
<th>λ_{max}^F [nm(eV)]</th>
<th>q^F</th>
</tr>
</thead>
<tbody>
<tr>
<td>B1/P1</td>
<td>410(3.02)/471(2.63)</td>
<td>12.82/23.18</td>
<td>548b</td>
<td>0.79b</td>
</tr>
<tr>
<td>B2/P2</td>
<td>458(2.71)/499(2.48)</td>
<td>18.98/24.73</td>
<td>589b</td>
<td>0.76b</td>
</tr>
<tr>
<td>B3/P3</td>
<td>380(3.26)/425(3.92)</td>
<td>19.05/25.83</td>
<td>640b</td>
<td>0.52b</td>
</tr>
<tr>
<td>B4/P4</td>
<td>378(3.28)/433(2.86)</td>
<td>30.29/30.38</td>
<td>658b</td>
<td>0.10b</td>
</tr>
<tr>
<td>B5/P5</td>
<td>422(2.94)/465(2.67)</td>
<td>26.66/35.52</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>B6/P6</td>
<td>424(2.92)/466(2.66)</td>
<td>33.99/36.81</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>B7/—</td>
<td>335(3.70)/—</td>
<td>53.73/—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>B8/—</td>
<td>345(3.59)/—</td>
<td>75.96/—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>B9/P9</td>
<td>342(3.63)/354(3.50)</td>
<td>84.51/87.26</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>B10/P10</td>
<td>362(3.43)/371(3.34)</td>
<td>75.55/93.31</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

a Observed as a shoulder. b Emission maxima/fluorescence quantum yields of the chromophores in series B.

Fig. 2 Representative UV/Vis absorption spectra of chromophores B1/P1, B4/P4 and B2/P2, B6/P6, B10/P10 measured in CH₂Cl₂ (10⁻⁵ M).
B2 molecular orbitals for representative pairs of chromophores measured/calculated for chromophores \( \pi \)-linker, and (vii) the lowest HOMO showed significantly raised LUMO due to its twisted \( \pi \)-system. As expected, the visualizations of frontier \( \pi \)-systems. The calculated data such as the ground stated dipole moments, energies of the HOMO and the LUMO and their differences \( \Delta E \) as well as \( \beta \) coefficients are presented in Table 5.

The first oxidation/reduction potentials measured in DMF were recalculated\(^{23}\) to the corresponding HOMO/LUMO energies and gaps (see also the ESI\(^{\dagger}\)) and were further compared with the AM1 calculated values. Although both HOMO–LUMO gaps differ in their absolute values (differences of 4.6–4.9 eV), they tightly correlate in both series of compounds (see the ESI Fig. S6–S7). Hence, even though they are very time-efficient, the used semi-empirical calculations are obviously capable of properly describing the trends seen by the electrochemical measurements and can be considered as a reasonable tool for the electronic property description of chromophores 1–10.

Fig. 3a shows the energy level diagram derived from the electrochemical measurements. This diagram nicely visualizes the aforementioned trends, namely: (i) the principal changes are observed in the LUMO, (ii) the LUMO is significantly lowered when going from series B (in black) to P (in red), (iii) the HOMO remains practically unchanged throughout the whole series of compounds, (iv) from chromophores B6/P6 onwards, the LUMO as well as the HOMO–LUMO gaps are steady, (v) although it does not contain a large \( \pi \)-linker, the chromophore B2 showed a significantly lowered LUMO level as a result of its planar arrangement, (vi) in contrast, the chromophore B7 showed significantly raised LUMO due to its twisted \( \pi \)-linker, and (vii) the lowest HOMO–LUMO gaps were measured/calculated for chromophores 9 and 10 that feature the longest \( \pi \)-linkers. As expected, the visualizations of frontier molecular orbitals for representative pairs of chromophores B2/B10 and P2/P10 showed acceptor-centered LUMO and donor-centered HOMO (Fig. 3b; for complete listing see the ESI Fig. S20–S29\(^{\dagger}\)). A significant charge-separation is being observed when extending the \( \pi \)-system.

The second-order optical nonlinearities of the newly synthesized chromophores B1–B10 were also determined experimentally by investigation of the Second Harmonic Generation (SHG). The SHG experiment was performed with powdered samples that were oriented by dc-electric field. The studies were performed for the angles that gave the maximal SHG output. A Nd-YAG laser (pulse duration 15 ns) was used as a fundamental laser beam. Powdered BiB\(_2\)O\(_4\) microcrystallites with known parameters of the second order susceptibilities were used as the reference samples. The experimental \( \beta \) coefficients along with the theoretical values are summarized in Table 5. Both hyperpolarizabilities showed sufficiently good correlation (see the ESI Fig. S8–S9). Chromophores bearing short and nonplanar arrangement of the \( \pi \)-linkers as in 1, 3 and 4 showed weak (P series) or no (B series) SHG behavior (see also the SHG readouts in the ESI Fig. S10–S19\(^{\dagger}\)). From the series of compounds 1–5, bearing up to two 1,4-phenylene and

### Table 5 Calculated electronic parameters and experimental hyperpolarizabilities of chromophores 1–10

<table>
<thead>
<tr>
<th>Compd</th>
<th>( \mu ) [D]</th>
<th>( E_{\text{HOMO}} ) [eV]</th>
<th>( E_{\text{LUMO}} ) [eV]</th>
<th>( \Delta E_{\text{HOMO-LUMO}} ) [eV]</th>
<th>( \beta ) (AM1) [esu \times 10^{-30}]</th>
<th>( \beta ) (exp.) [pm V(^{-1})]</th>
</tr>
</thead>
<tbody>
<tr>
<td>B1/P1</td>
<td>8.56/9.91</td>
<td>-8.28/-8.40</td>
<td>-0.92/-1.43</td>
<td>7.36/6.97</td>
<td>245/349</td>
<td>-9/0.53</td>
</tr>
<tr>
<td>B2/P2</td>
<td>9.74/11.89</td>
<td>-8.22/-8.33</td>
<td>-1.07/-1.75</td>
<td>7.15/6.38</td>
<td>649/942</td>
<td>0.9/1.1</td>
</tr>
<tr>
<td>B3/P3</td>
<td>9.67/9.07</td>
<td>-8.15/-8.28</td>
<td>-1.18/-1.78</td>
<td>6.97/6.50</td>
<td>375/408</td>
<td>-9/0.6</td>
</tr>
<tr>
<td>B4/P4</td>
<td>10.43/10.69</td>
<td>-8.20/-8.23</td>
<td>-1.31/-1.69</td>
<td>6.89/6.54</td>
<td>681/787</td>
<td>-9/1.1</td>
</tr>
<tr>
<td>B5/P5</td>
<td>10.77/11.49</td>
<td>-8.21/-8.32</td>
<td>-1.47/-1.92</td>
<td>6.74/6.40</td>
<td>809/956</td>
<td>1.2/1.2</td>
</tr>
<tr>
<td>B6/P6</td>
<td>11.48/11.3</td>
<td>-8.16/-8.38</td>
<td>-1.55/-2.01</td>
<td>6.61/6.37</td>
<td>1263/1444</td>
<td>3.6/1.4</td>
</tr>
<tr>
<td>B7</td>
<td>10.9</td>
<td>-8.18/-</td>
<td>-1.53/-</td>
<td>6.59/-</td>
<td>867/-</td>
<td>2.1/-</td>
</tr>
<tr>
<td>B8</td>
<td>11.3</td>
<td>-8.16/-</td>
<td>-1.61/-</td>
<td>6.55/-</td>
<td>1 268/-</td>
<td>2.8/-</td>
</tr>
<tr>
<td>B9/P9</td>
<td>10.89/11.38</td>
<td>-8.11/-8.15</td>
<td>-1.56/-2.00</td>
<td>6.55/6.15</td>
<td>1218/1484</td>
<td>3.2/3.0</td>
</tr>
<tr>
<td>B10/P10</td>
<td>11.78/11.27</td>
<td>-8.11/-8.29</td>
<td>-1.64/-2.07</td>
<td>6.47/6.22</td>
<td>1701/1912</td>
<td>4.9/3.1</td>
</tr>
</tbody>
</table>

*Below noise level (see the ESI).*

### Calculations and NLO properties

Quantum-chemical calculations and simulation of NLO properties of all target chromophores were performed at the semi-empirical level of theory with the AM1 method and with the time-dependent Hartree–Fock (TDHF) calculations, respectively. All the calculations were performed using the PCGAMESS package. The calculated data such as the ground stated dipole moment \( \mu \), energies of the HOMO and the LUMO and their differences \( \Delta E \) as well as \( \beta \) coefficients are presented in Table 5.
A new series of X-shaped push–pull molecules based on 4,5-disubstituted benzene-1,2-dicarbonitrile has been synthesized. Three compounds B1, B3 and B4 were obtained by twofold Suzuki–Miyaura reactions, and seven target chromophores B2 and B5–B10 with the π-linker separated from the acceptor by an additional triple bond were obtained by Sonogashira cross-coupling. All compounds were compared with isobolog chromophores P1–P10 based on 5,6-disubstituted pyrazine-2,3-dicarbonitrile and subsequent structure–property relationships were further evaluated. X-ray analysis revealed different spatial arrangements of both acceptor moieties and the appended N,N-dimethylamino-substituted π-linkers. The dicyanopyrazine core showed significant out of plane deformation while the dicyanobenzene remained almost fully planar. The extent of the ICT has been studied by BLA (δr) and the Bird index (I_Br). Both quantities indicated higher ICT in the P series of compounds. This has been further confirmed by the 13C NMR and IR spectra that showed higher electron saturation of the CN group in dicyanopyrazine compounds. 13C NMR chemical shift and the frequency of the stretching vibration of the CN group have also been affected by the length, composition and spatial arrangement of the π-linker. Electrochemical measurements carried out by the CV and RDV confirm the observations made above. Principal changes were recorded in the first reduction potential. Significantly positively shifted first reduction potentials have been observed for compounds with fully planar π-linkers connected to the acceptor moiety via triple bonds (e.g. B2/P2, B6/P6 and B10/P10). All chromophores from B6/P6 onwards showed almost constant first reduction potentials which implies that a combination of two/three 1,4-phenylene moieties in conjunction with acetylene units caused complete isolation of the N,N-dimethylamino donor and the dicyanobenzene/pyrazine acceptor. Optical properties of the studied chromophores are dominated by one or more CT-bands, which reflects multiple conjugated pathways between each N,N-dimethylamino and cyano groups. In general, chromophores in the series P showed bathochromically shifted longest-wavelength absorption maxima which reflect higher electron withdrawing character of the dicyanopyrazine moiety. The aforementioned structural changes in particular chromophores affect also their absorption spectra. Extension of the π-system and isolation of the D and A parts of the X-shaped push–pull molecules led to a hypsochromic shift of the CT-bands that are, from B7/P6 onwards, observed as a joint peak at about 350 nm with large molar absorption coefficients. In contrast to P1–P4, chromophores B1–B4 showed also fluorescent properties that gradually decrease with the extension of the π-system.

Quantum-chemical calculations further confirmed the experimental data. In general, from the experiments and calculations carried out for X-shaped molecules 1–10 we can conclude the following:

- replacement of the dicyanobenzene with the dicyanopyrazine acceptor affects the optoelectronic properties most significantly (tuning by the LUMO)
- the HOMO of 1–10 remained almost unchanged independently of the size and structure of the π-linker
- the HOMO is localized on the peripheral donors while the LUMO is placed on the acceptor units
- the LUMO can be further lowered by planarization of the entire π-system (e.g. B2 vs. B7)
- a position of the acetylene unit within the π-linker plays also an important role (e.g. B4/P4 vs. B5/P5)
- π-linker bearing two acetylene units in combination with two 1,4-phenylene moieties completely isolates A and D parts of X-shaped push–pull molecules
- the lowest HOMO-LUMO gap and the highest nonlinearities have been measured/calculated for molecules B9/P9 and B10/P10 with the largest π-system
- planar molecules (e.g. B2, B6 and B10) delivered a strong SHG response
- first-order hyperpolarizability β depends not only on the extent of the ICT but also on the number of polarizable electrons (length of the π-system).

In view of the current interest in push–pull systems and their wide applications mainly in NLO and DSSC (both cyano groups can be hydrolysed to well-anchoring carboxylic acid functions),19 we believe that this structure–property study on X-shaped molecules would serve as a useful guideline.

**Conclusions**

**Experimental section**

For general information and synthesis of 11 see the ESL. Full spectral characterization of pyrazine chromophores P1–P10 is given in our previous publication.10
Organic & Biomolecular Chemistry

General procedure for Suzuki–Miyaura cross-coupling

4,5-Diodobenzene-1,2-dicarbonitrile (152 mg, 0.4 mmol) and an appropriate boronic acid or its ester (0.84 mmol, 2.1 eq.) were dissolved in a mixture of THF–H2O (20 ml, 4:1). Argon was bubbled through the solution for 15 min whereupon [PdCl2(PPh3)2] (28 mg, 0.04 mmol) and Na2CO3 (89 mg, 0.64 mmol) were added and the reaction mixture was stirred at 65 °C for 12 h. The reaction was diluted with water (50 ml) and extracted with CH2Cl2 (2 × 50 ml). The combined organic extracts were dried (Na2SO4), the solvents were evaporated in vacuo and the crude product was purified by column chromatography (SiO2; indicated solvent system).

General procedure for Sonogashira cross-coupling

4,5-Diodobenzene-1,2-dicarbonitrile (152 mg, 0.4 mmol) and an appropriate terminal acetylene (0.84 mmol, 2.1 eq.) were dissolved in THF (20 ml) and Et,N (5 ml). Argon was bubbled through the solution for 15 min whereupon [PdCl2(PPh3)2] (28 mg, 0.04 mmol) and CuI (8 mg, 0.04 mmol) were added and the reaction mixture was stirred at 60 °C for 24 h. The solvents were evaporated in vacuo and the crude product was purified by column chromatography (SiO2; indicated solvent system).

Chromophore B1. General procedure for Suzuki–Miyaura cross-coupling with 4-(N,N-dimethylamino)phenylboronic acid pinacol ester18 (208 mg) gave 121 mg (83%) of chromophore B1 as a bright yellow fluorescent solid. M.p. > 300 °C. Rf = 0.19 (SiO2; CH2Cl2–hexane 6:1). 1H-NMR (400 MHz, 25 °C, CDCl3): δH = 7.88 (s, 2H, 2 × ArH), 7.94 (d, J(H,H) = 8.8 Hz, 4H; 2 × Ph), 6.66 (d, J(H,H) = 8.8 Hz, 4H; 2 × Ph), 5.99 (d, J(H,H) = 8.8 Hz, 4H; 2 × Ph), 5.83 (d, J(H,H) = 8.4 Hz, 4H; 2 × Ph), 6.85 (d, J(H,H) = 8.4 Hz, 4H; 2 × Ph), 6.77 (d, J(H,H) = 8.4 Hz, 4H; 2 × Ph), 3.02 (s, 12H, 2 × N(CH3)2). 13C-NMR (100 MHz, 25 °C, CDCl3): δC = 150.20, 145.60, 135.58, 130.42, 125.83, 112.67, 112.17, 40.42. IR (HATR): νmax/cm⁻¹ = 2920, 2223 (CN), 1647, 1637, 1361, 1356, 1307, 1237, 112.82, 101.43, 86.33, 40.64. IR (HATR): νmax/cm⁻¹ = 2961, 2202 (CN), 1596, 1521, 1358, 1222, 1133, 814. HR-FT-MALDI-MS (DHB) m/z: 566.2466 (M⁺), C40H30N4⁺ requires 566.2465. Anal. Calcd for C40H30N4 (566.71): C 84.51, H 5.51, N 9.89; Found C 84.52, H 5.51, N 9.73.

Chromophore B5. General procedure for Sonogashira cross-coupling with 4′-ethyl-N,N-dimethylbiphenyl-4-amine18 (186 mg) gave 165 mg (73%) of chromophore B5 as a dark red solid. M.p. > 300 °C. Rf = 0.17 (SiO2; CH2Cl2–hexane 6:1). 1H-NMR (400 MHz, 25 °C, CDCl3): δH = 7.91 (s, 2H, 2 × ArH), 7.60 (s, 8H, 4 × Ph), 7.54 (d, J(H,H) = 8.4 Hz, 4H; 2 × Ph), 6.80 (d, J(H,H) = 8.4 Hz, 4H; 2 × Ph), 3.01 (s, 12H, 2 × N(CH3)2). 13C-NMR (100 MHz, 25 °C, CDCl3): δC = 150.64, 143.40, 136.35, 132.7, 131.13, 127.91, 127.53, 126.31, 118.86, 115.10, 113.83, 112.82, 101.43, 86.33, 40.64. IR (HATR): νmax/cm⁻¹ = 2880, 2200 (CN), 1595, 1441, 1360, 1222, 1092, 811. HR-FT-MALDI-MS (DHB) m/z: 566.2450 (M⁺), C40H30N4⁺ requires 566.2465. Anal. Calcd for C40H30N4 (567.1): C 84.78, H 5.34, N 9.89; Found C 84.38, H 5.19, N 9.66.

Chromophore B6. General procedure for Sonogashira cross-coupling with 4′-[4-ethylphenyl]ethyl)-N,N-dimethylaniline18 (206 mg) gave 177 mg (72%) of chromophore B6 as an orange solid. M.p. > 300 °C. Rf = 0.76 (SiO2; CH2Cl2–hexane 6:1). 1H-NMR (400 MHz, 25 °C, CDCl3): δH = 7.91 (s, 2H, 2 × ArH), 7.53–7.48 (m, 8H, 4 × Ph), 7.46–7.40 (m, 4H, 2 × Ph), 6.66 (d, J(H,H) = 8.4 Hz, 4H; 2 × Ph), 3.00 (s, 12H, 2 × N(CH3)2). 13C-NMR (100 MHz, 25 °C, CDCl3): δC = 150.64, 143.40, 136.35, 133.15, 132.11, 131.32, 131.00, 126.46, 120.22, 114.98, 114.29, 112.00, 100.86, 94.56, 87.41, 87.16, 40.39. IR (HATR): νmax/cm⁻¹ = 3004, 2206 (CN), 1605, 1525, 1356, 1222, 1130, 846, 814. HR-FT-MALDI-MS (DHB) m/z: 614.2470 (M⁺), C45H36N4⁺ requires 614.2465. Anal. Calcd for C45H36N4 (614.75): C 85.97, H 4.92, N 9.11; Found C 85.50, H 5.01, N 9.21.
ArH), 7.70–7.66 (m, 16H, 8 × Ph), 7.55 (d, J(H,H) = 9.2 Hz, 4H; 2 × Ph), 6.81 (d, J(H,H) = 8.8 Hz, 4H; 2 × Ph), 3.01 (s, 12H, 2 × N(CH3)2). 13C-NMR (100 MHz, 25 °C, CDCl3): δc = 136.20, 135.73, 135.48, 133.76, 127.62, 127.30, 40.08 (1 signal is missing). IR (HATR): νmax/cm−1 = 3004, 2202 (CN), 1592, 1516, 1357, 1222, 1093, 838, 812. HR-FT-MALDI-MS (DHB) m/z: 766.3062 (M+), C60H38N4+ requires 766.3091. Anal. Calcd for C60H38N4 (814.99): C 88.43, H 4.51, N 7.08. Found C 88.42, H 4.50, N 7.07.

**Computational procedure**

All calculations were performed with a semi-empirical AM1 method implemented in the PCGAMESS package.27 Computational files were prepared and interpreted with the aid of GABEDIT 2.4.6 (ref. 28) including the preparation of graphics.

**Acknowledgements**

This research was supported by the Czech Science Foundation (P106/12/0392). L. and F. B./M. L. are indebted to the Ministry of Education, Youth and Sports of the Czech Republic (CZ.1.07/2.3.00/30.0021 and LG13053). W. K. is grateful to the Balassi Institute for financial support.

**Notes and references**


28 A. R. Allouche, Gabedit is a free Graphical User Interface for computational chemistry packages. It is available from http://gabedit.sourceforge.net/.