# ChemComm



**View Article Online** 

## COMMUNICATION

Check for updates

Cite this: Chem. Commun., 2022. **58**, 5889

Received 18th March 2022, Accepted 20th April 2022

DOI: 10.1039/d2cc01568d

rsc.li/chemcomm

## Dual-photofunctional organogermanium compound based on donor-acceptor-donor architecture\*

Aleksandra Nyga, 🗓 ‡<sup>a</sup> Takahito Kaihara, ‡<sup>b</sup> Takumi Hosono, <sup>b</sup> Massimiliano Sipala, <sup>a</sup> Patrycja Stachelek,<sup>c</sup> Norimitsu Tohnai,<sup>b</sup> Satoshi Minakata, 回<sup>b</sup> Leonardo Evaristo de Sousa, <sup>[D]</sup> <sup>d</sup> Piotr de Silva, <sup>[D]</sup> \*<sup>d</sup> Przemyslaw Data <sup>[D]</sup> \*<sup>a</sup> and Youhei Takeda 🛈 \*b

A dual-photofunctional organogermanium compound based on a donor-acceptor-donor architecture that exhibits thermally activated delayed fluorescence and mechano-responsive luminochromism has been developed. The developed compound was successfully applied as an emitter for efficient organic light-emitting diodes.

Organogermanes have attracted attention in a wide variety of research fields such as organometallic chemistry,<sup>1</sup> biology,<sup>2</sup> and catalysis.<sup>3</sup> In sharp contrast, the utilization of organogermanium compounds and polymers as optoelectronic materials has been rarely exploited.<sup>4-6</sup> For example, tetraaryl germanes and related polymers have been used as host materials for organic light-emitting diodes (OLEDs), by utilizing the high triplet energy and adequate carrier transport capability.<sup>4</sup> Also, a systematic study of blue-emitters based on phenothiaborinconnected acridan analogues with group 14 elements including germanium for non-doped efficient OLEDs has been reported.5 Given unique characteristics of organogermanes such as a large atomic radius (122 pm), electropositive nature ( $\chi_p$  = 2.01), and capability of  $\sigma$ - $\pi$  conjugation, the development of novel Gecontaining organic functional materials would offer us great opportunities to cultivate design principles for a new class of functional materials. Specifically, the diminished  $\sigma$ - $\pi$  conjugation ability of Ge

element when compared with Si allows for higher-energy triplet excited states than the corresponding silicon compounds.<sup>4</sup> Since the triplet energy is a limiting factor for determining the emission color of thermally activated delayed fluorescent (TADF),<sup>7</sup> donor-acceptor (D-A) type organogermanes are promising for realizing high-energy TADF (blue to green).5

Herein, we disclose the development of a dual-photofunctional Ge-containing donor-acceptor-donor (D-A-D) type compound 1 (Fig. 1a). Notably, the developed compound represents a rare example of luminescent organogermanium compounds,5,8 and it nicely shows dual photofunctionality of TADF and mechanoresponsive luminochromism.9 Furthermore, compound 1 serves as the first example of green-TADF emitter based on organogermanium for an efficient OLED device.

The synthetic route to compound 1 is shown in Fig. 1b (for the details, see the ESI<sup>†</sup>). We initially developed a synthetic method for dihydrophenazagermine 4, starting from N-protected dibromo diarylamine 2. Dilithiation of 2 followed by trapping with Ph<sub>2</sub>GeCl<sub>2</sub> and detaching the N-p-methoxybenzyl (PMB) group with DDQ afforded 4. The X-ray diffraction analysis of the single crystal 4

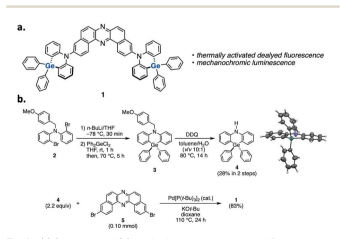


Fig. 1 (a) Structure and (b) synthetic route to compound 1.

<sup>&</sup>lt;sup>a</sup> Faculty of Chemistry, Silesian University of Technology, M. Strzody 9,

<sup>44-100,</sup> Gliwice, Poland. E-mail: przemyslaw.data@polsl.pl

<sup>&</sup>lt;sup>b</sup> Department of Applied Chemistry, Graduate School of Engineering, Osaka University, Yamadaoka 2-1, Suita, Osaka 565-0871, Japan. E-mail: takeda@chem.eng.osaka-u.ac.jp

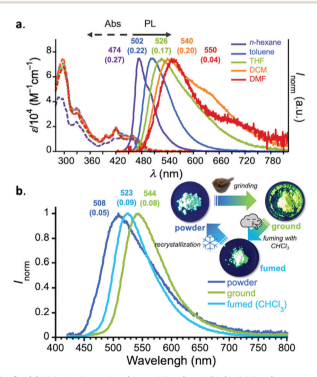
<sup>&</sup>lt;sup>c</sup> Chemistry Department, Durham University, South Road, Durham DH1 3LE, UK

<sup>&</sup>lt;sup>d</sup> Department of Energy Conversion and Storage, Technical University of Denmark, Anker Engelunds Vej 301, 2800 Kongens Lyngby, Denmark. E-mail: pdes@dtu.dk † Electronic supplementary information (ESI) available: Experimental procedures for the syntheses of materials, spectroscopic data of new compounds, single crystal X-ray crystallographic data, cyclic voltammogram, thermogravimetric analysis (TGA) profiles, the copies of NMR spectra of new compounds, and theoretical calculation details. CCDC 2153796. For ESI and crystallographic data in CIF or other electronic format see DOI: https://doi.org/10.1039/d2cc01568d

<sup>‡</sup> These authors contributed equally to this work.

revealed that the angle between two phenylene planes is  $173^{\circ}$  (the inset figure in Scheme S1, for the details, see Table S1, ESI<sup>†</sup>), which is larger than that of its Si analogue (*ca.*  $133^{\circ}$ ) in a D–A system.<sup>5</sup> This planar structure of the donor would be ascribed to the longer Ge–C bond (*ca.* 1.93 Å) when compared to Si–C bond (*ca.* 1.85 Å).<sup>5</sup> D–A–D compound **1** was successfully synthesized through a Pd-catalyzed Buchwald–Hartwig amination of 3,11-dibromo-dibenzophenazine **5**<sup>10</sup> with donor **4** (Fig. 1b).

The steady-state UV-vis absorption spectra of diluted solutions ( $c = 10^{-5}$  M) of 1 displayed a  $\pi - \pi^*$  absorption at  $\lambda_{abs}$  300– 350 nm and weak vibronic absorption ascribed to the acceptor core at  $\lambda_{abs}$  380-420 nm (Fig. 2a). Also, in the lower-energy region, a broad and weak absorption ascribed to chargetransfer (CT) transition was observed. In a non-polar solvent (n-hexane), compound 1 displayed a vibronically-shaped emission spectrum ( $\lambda_{em}$  474 nm) with a moderate photoluminescence quantum yield (PLQY), which is ascribed to the emission from the locally excited state (<sup>1</sup>LE). In contrast to the absorption spectra, the PL spectrum significantly red-shifted as a function of solvent polarity (Fig. 2a), suggesting the CT character of the excited states in those solvents. The Mataga-Lippert plot analysis corroborated the hybridized local and charge-transfer (HLCT) nature of compound 1 (Fig. S3, ESI<sup>+</sup>). An interesting phenomenon other than solvatochromism involves dualemission in a polar solvent such as dichloromethane (DCM) and DMF (Fig. 2a). Comparison with a known D-A system



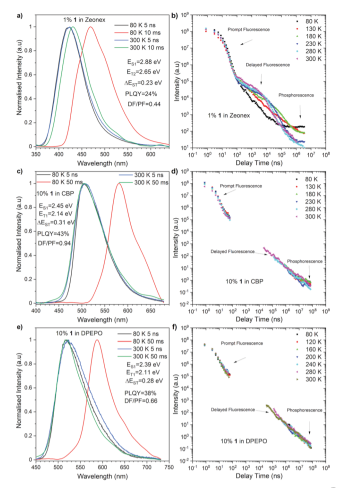
**Fig. 2** (a) UV-vis absorption (dotted lines) and PL (solid lines) spectra of diluted solutions of **1** ( $c = 10^{-5}$  M). Photoluminescence spectra were acquired with the excitation at  $\lambda_{ex}$  400 nm. (b) PL spectra of power (blue), ground (moth green), and fumed sample (light blue) of **1** excited at  $\lambda_{ex}$  400 nm. The photographs represent the image of the emission-color change cycle. All emoji designed by OpenMoji-the open-source emoji and icon project. License: CC BY-SA 4.0.

having a phenazagermin donor allows us to notice that the contribution of the emission ascribed to axial-axial conformer is little in our system.<sup>5</sup> The exclusive population of equatorial-equatorial conformer of compound **1** was supported by the theoretical calculations (*vide infra*).

It is noted that D-A-D compound 1 exhibited a significant change in emission color in the solid states, responding to external stimuli (Fig. 2b). When as-prepared powdery solid ("powder") was ground with a pestle and mortar ("ground"), the emission peak significantly shifted to the lower-energy regime ( $\Delta\lambda$  1303 cm<sup>-1</sup>). In contrast, when the ground was fumed with organic vapor such as CHCl<sub>3</sub> and CH<sub>2</sub>Cl<sub>2</sub>, the emission spectrum slightly reverted back  $(\Delta\lambda 738 \text{ cm}^{-1})$ . The initial powder state was recovered by recrystallization. The powder X-ray diffraction (PXRD) analyses of the solids revealed that only the peak at  $2\theta = 23.4^{\circ}$  (d = 3.8 Å) significantly decreased upon grinding and its intensity was reverted back upon exposing to chloroform vapor (Fig. S4, ESI<sup>+</sup>). This would suggest that the emission color change in response to stimuli would be ascribed to the fluctuation of intermolecular electronic interaction.<sup>11</sup> It is noted that such dual photofunctionality of TADF and mechano-responsive emission color change would offer opportunities for sensing applications.<sup>12</sup>

Time-resolved luminescence spectroscopy of compound **1** was performed in both inert non-polar cyclo olefin polymer host Zeonex<sup>(R)</sup>, 4,4'-bis(*N*-carbazoyl)-1,1'-biphenyl (CBP), and bis[(2-diphenyl-phosphino)phenyl]ether oxide (DPEPO) hosts, the latter two of which were used to mimic the chemical environment within an OLED device. Each material showed emission within two distinct time regions within all the hosts. The first, decaying with a lifetime within the nanosecond time regime in all materials, is attributed to prompt emission from the singlet excited state due to its temperature independence (Fig. 3). In all cases, spectral inspection at time delays (TD) = 5 ns shows a Gaussian charge transfer (<sup>1</sup>CT) singlet peak that decays over longer times.

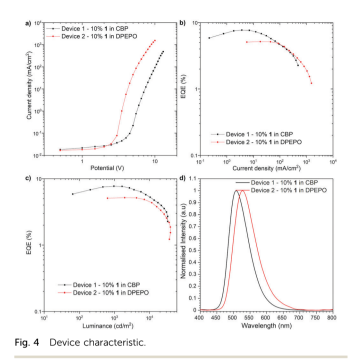
At longer delay times, in the microsecond/millisecond delay time regions, delayed emission was observed (Fig. 3). Depending upon the experimental temperature, both the singlet state delayed emission and triplet state emission was observed on similar millisecond timescales, and therefore, the emission from each state is most easily elucidated upon spectral inspection at different temperatures (Fig. 3). At room temperature (300 K), the delayed emission spectrum had the same shape and onset energy as the prompt emission in both CBP and DPEPO (Fig. 3c and e). Therefore, the delayed emission was identified as TADF. But, the delayed emission in Zeonex® was observed in a slightly lower-energy region (Fig. 3a), probably due to the structural relaxation in the excited state. Intriguingly, the triplet excited state emissions observed at low temperatures showed quite varied energies: 2.65 eV in Zeonex<sup>®</sup>, 2.14 eV in CBP, and 2.11 eV in DPEPO hosts (Fig. 3a, c, and e). Such fluctuation in the triplet energy is unusual for dibenzo[a,j]phenazine-cored D-A-D systems.<sup>13</sup> This observation could suggest a higher triplet state (T<sub>2</sub>) was involved in the emission in Zeonex<sup>®</sup>. Such a scenario was partly supported by the theoretical calculations (vide infra). Such complications cause much weaker TADF emission in the CBP and DPEPO



**Fig. 3** Emission intensity of **1** against delay time measured in (a) Zeonex<sup>(B)</sup>, (c) CBP, and (e) DPEPO at different temperatures. Normalized emission spectra of **1** in (b) Zeonex<sup>(B)</sup>, (d) CBP, and (f) DPEPO at varying delay times at 300 K and 80 K.

matrices and a much lower rISC contribution than it was observed in the Zeonex<sup>®</sup> matrix. The polarity of the host affects the energy of both singlet and triplet excited states. Therefore, as the polarity increases, the  $\Delta E_{\rm ST}$  of the materials increases from 0.23 eV in Zenex<sup>®</sup> to 0.31 eV in CBP, and 0.28 in DPEPO, which weaken the rISC process. When compared with the photophysical properties of a phenothiaborine-connected phenazagermine compound<sup>5</sup> in a phosphineoxide host (diphenyl-phosphoryl)dibenzo[*b,d*]furan: PPF, the  $\lambda_{\rm em}$  for **1** in DPEPO (518 nm) locates at the lower-energy region than that for the D–A dyad (468 nm), while the  $\Delta E_{\rm ST}$  of **1** (0.28 eV) is larger than that for the D–A dyad (0.11 eV).

The OLED devices were fabricated to investigate whether the Gecontaining compound is applicable to optoelectronic applications (Fig. 4). The HOMO and LUMO energy levels of 1 were determined by cyclic voltammetry (CV) to be -5.65 eV and -3.34 eV, respectively (Fig. S1, ESI†). Since the thermogravimetric analysis (TGA) indicated the high thermal stability of compound 1 [ $T_d$  (5 wt% loss under N<sub>2</sub>) = 466 °C] (Fig. S2, ESI†), the devices were fabricated with thermal evaporation technique. As a result of an optimization study, the optimal configuration was obtained as follows: Device 1



[ITO/N,N'-di(1-naphthyl)-N,N'-diphenyl-(1,1'-biphenyl)-4,4'-diamine (NPB) (40 nm)/10% 1 in CBP (25 nm)/2,2',2"-(1,3,5-benzinetriyl)tris(1-phenyl-1H-benzimida-zole) (TPBi) (40 nm)/LiF (1 nm)/Al (100 nm)], Device 2 [ITO/NPB (40 nm)/tris(4-carbazoyl-9ylphenyl)amine (TCTA) (10 nm)/10% 1 in DPEPO (20 nm)/TPBi (60 nm)/LiF (1 nm)/Al (100 nm)] (Fig. 4). The characteristics of the OLED structures revealed a good efficiency of Ge-containing TADF emitter 1 in CBP host OLED device (EQE ca. 7.7%, Fig. 4b), which exceeds the theoretical maximum of the OLED device fabricated with prompt fluorescent emitter (ca. 5%). On the one hand, the DPEPO host-based device showed lower efficiency (EQE ca. 5.1%). The luminance of the device in both hosts is quite high (more than 32 000 cd m<sup>-2</sup> in CBP and 39 000 cd m<sup>-2</sup> in DPEPO), which suggests good charge recombination in the device. The turn-on voltage was around 2.5 V in DPEPO and 4.0 V in CBP. In both structures, the OLED characteristic showed a moderate roll-off dependency (Fig. 4b and c). It is worth noting that the Ge-containing TADF emitter for OLEDs is quite limited so far,<sup>5</sup> and compound 1 represents the first example of green-TADF emitter based on organogermanium compound.

To obtain further insight into the behavior of compound 1, density functional theory (DFT) calculations were performed and the nuclear ensemble method was used to estimate photophysical rates<sup>14</sup> (see the ESI† for details). A conformational analysis of 1 was conducted, revealing that the equatorialequatorial conformation is the most stable one in ground, S<sub>1</sub> and T<sub>1</sub> states. A fluorescence spectrum simulation of compound 1 (Fig. S3, ESI†) predicts an emission peak at 516 nm (2.40 eV) and an emission rate of  $1.2 \times 10^7 \text{ s}^{-1}$  (Table S5, ESI†), which matches well the experimental value (502 nm) in toluene (Fig. 2) but underestimates the S<sub>1</sub> emission energy in Zeonex<sup>®</sup> (Fig. 3a), though toluene and Zeonex<sup>®</sup> share similar dielectric constants (*ca.* 2.3). This suggests that the S<sub>1</sub> emission in this

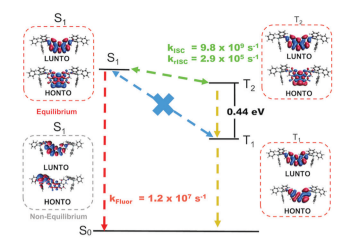


Fig. 5 NTOs for the  $S_1$ ,  $T_1$  and  $T_2$  states and schematics of the TADF mechanism in compound **1** with DFT estimated rates in toluene. A CT  $S_1$  state calculated at a non-equilibrium geometry is also shown.

case could result from another conformer being preferentially locked in place by solid-state effects specific to this host matrix, a hypothesis that is corroborated by emission energies of 2.83 eV and 2.75 eV calculated for the axial-axial and equatorialaxial conformers, respectively.

Estimated rates for ISC from S<sub>1</sub> (Table S4, ESI<sup>+</sup>) indicate that the preferred process takes place between the S<sub>1</sub> and T<sub>2</sub> states, due to a lower average energy gap (0.012 eV between  $S_1$  and  $T_2$ versus 0.421 eV between S1 to T1). The simulated phosphorescence spectra from  $T_1$  and  $T_2$  (Fig. S6 and Tables S4–S6, ESI<sup> $\dagger$ </sup>) predict one order of magnitude higher emission rate from T<sub>2</sub> than from  $T_1$  with a  $T_1$  peak at 579 nm (2.14 eV) and a  $T_2$  peak at 475 nm (2.61 eV). The predicted energy of T<sub>2</sub> matches the phosphorescence energy measured in Zeonex<sup>®</sup> (2.65 eV), providing further evidence for the S<sub>1</sub> to T<sub>2</sub> ISC hypothesis. The T<sub>1</sub> energy, in turn, agrees well with phosphorescence energies in CBP and DPEPO (Fig. 3c and e), suggesting that in these host matrices, transfers to  $T_1$  from  $S_1$  may be more efficient due to higher solvatochromic shift or that internal conversion from T<sub>2</sub> can play an important role. In toluene, calculations indicate an average  $T_1$ - $T_2$  gap of 0.44 eV (Table S7, ESI<sup>+</sup>), enough to prevent rapid depopulation to T1 if the competing processes (rISC, phosphorescence) are efficient enough. Calculated rISC rates from T<sub>1</sub> and T<sub>2</sub> states (Tables S5 and S6, ESI<sup>+</sup>) reveal that rISC to  $S_{1}$  is the preferred transfer for both triplet states. The rISC rate from  $T_2$  to  $S_1$  is orders of magnitude higher than that from  $T_1$ , indicating more efficient TADF when T<sub>2</sub> is the state involved in the triplet harvesting mechanism. This agrees with the observed decrease in TADF performance in CBP and DPEPO, which, as mentioned above, have larger  $T_1$  involvement.

Fig. 5 summarizes the proposed TADF mechanism based on the probabilities of each process (Table S8, ESI<sup>†</sup>). Natural transition orbitals (NTOs) for the  $S_1$ ,  $T_1$  and  $T_2$  states have mostly localized character, explaining the mild red shift observed from CBP to DPEPO. On the other hand, stronger shifts in solution may result from the fact that vibrational effects can alter the electronic character of the excited states,<sup>15,16</sup> (see example of CT S<sub>1</sub> in Fig. 5)

such that the actual picture is not fully captured by NTOs on optimized structures. In fact, this effect should be more prominent in solution than in solid-state, as the vibrational motion may be hindered in the latter case.

In summary, we have developed a dual-photofunctional organogermanium compound that exhibits TADF and mechanochromic behavior. The compound represents the first example of green-TADF emitter using organogermanium scaffold for efficient OLED devices. Theoretical calculations shed light on the importance of the higher triplet excited state to yield TADF *via* rISC process. This work opens up a new avenue for organogermanium-based multiphotofunctional materials in the future.

This work was partly supported by KAKENHI (JP19H05716, JP20H02813, and JP21K18960 to Y. T.), a grant from VILLUM FONDEN (00028053 to P. de Se. and L. E. S.) and the First Team program (POIR.04.04.00-00-4668/17-00 to P. D.). The full list for acknowledgement is included in the ESI.<sup>†</sup>

#### Conflicts of interest

There are no conflicts to declare.

#### Notes and references

- 1 C. S. Weinert, Encycl. Inorg. Bioinorg. Chem., 2015, 1-18.
- 2 (a) J. S. Thayer, Appl. Organomet. Chem., 1987, 1, 227-234;
  (b) L. G. Menchikov and M. A. Ignatenko, Pharm. Chem. J., 2013, 46, 635-638.
- 3 C. Fricke and F. Schoenebeck, Acc. Chem. Res., 2020, 53, 2715-2725.
- 4 (a) C. Yao, Q. Cui, J. Peng, X. Xu, R. Liu, J. Wang, Y. Tian and L. Li, J. Mater. Chem. C, 2015, 3, 5017–5025; (b) C. Yao, Y. Yang, L. Li, M. Bo, C. Peng and J. Wang, J. Mater. Chem. C, 2018, 6, 6146–6152; (c) M.-K. Hung, K.-W. Tsai, S. Sharma, J.-Y. Wu and S.-A. Chen, Angew. Chem., Int. Ed., 2019, 58, 11317–11323; (d) M.-K. Hung, K.-W. Tsai, S. Sharma, J. Lei, J.-Y. Wu and S.-A. Chen, ACS Appl. Mater. Interfaces, 2019, 11, 36895–36904.
- 5 K. Matsuo and T. Yasuda, Chem. Sci., 2019, 10, 10687–10697.
- 6 (a) N. Allard, R. B. Aïch, D. Gendron, P.-L. T. Boudreault, C. Tessier, S. Alem, S.-C. Tse, Y. Tao and M. Leclerc, *Macromolecules*, 2010, 43, 2328–2333; (b) J. Ohshita, Y.-M. Hwang, T. Mizumo, H. Yoshida, Y. Ooyama, Y. Harima and Y. Kunugi, *Organometallics*, 2011, 30, 3233–3236; (c) C. M. Amb, S. Chen, K. R. Graham, J. Subbiah, C. E. Small, F. So and J. R. Reynolds, *J. Am. Chem. Soc.*, 2011, 133, 10062–10065.
- 7 H. Uoyama, K. Goushi, K. Shizu, H. Nomura and C. Adachi, *Nature*, 2012, **492**, 234–238.
- 8 (a) J. Ohshita, K. Murakami, D. Tanaka, Y. Ooyama, T. Mizumo, N. Kobayashi, H. Higashimura, T. Nakanishi and Y. Hasegawa, *Organometallics*, 2014, **33**, 517–521; (b) H. Arii, Y. Iwanami, D. Nakane, H. Masuda, J. Matsumoto, T. Shiragami, K. Mochida and T. Kawashima, *Organometallics*, 2021, **40**, 1363–1370.
- 9 Y. Sagara, S. Yamane, M. Mitani, C. Weder and T. Kato, *Adv. Mater.*, 2016, **28**, 1073–1095.
- 10 Y. Takeda, M. Okazaki and S. Minakata, *Chem. Commun.*, 2014, **50**, 10291–10294.
- 11 S. Goto, Y. Nitta, N. O. Decarli, L. E. de Sousa, P. Stachelek, N. Tohnai, S. Minakata, P. de Silva, P. Data and Y. Takeda, *J. Mater. Chem. C*, 2021, 9, 13942–13953.
- 12 P. Data and Y. Takeda, Chem. Asian J., 2019, 14, 1613-1636.
- 13 Y. Takeda, P. Data and S. Minakata, Chem. Commun., 2020, 56, 8884-8894.
- 14 L. E. de Sousa and P. de Silva, J. Chem. Theory Comput., 2021, 17, 5816–5824.
- 15 L. E. de Sousa and P. de Silva, *ChemRxiv*, 2022, DOI: 10.26434/ chemrxiv-2022-pq978.
- 16 P. de Silva, C. A. Kim, T. Zhu and T. Van Voorhis, *Chem. Mater.*, 2019, **31**, 6995–7006.