Thiophene/selenophene-based S-shaped double helicenes: regioselective synthesis and structures

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Letter

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Abstract

2,5-Di(trimethylsilanyl)dithieno[2,3-*b*:3′,2′-*d*]thiophene ((TMS)₂-*bb*-**DTT**), 2,5-di(trimethylsilanyl)diseleno[2,3-*b*:3′,2′-*d*]thiophene ((TMS)₂-*bb*-**DST**), and 2,5-di(trimethylsilanyl)diseleno[2,3-*b*:3′,2′-*d*] selenophene ((TMS)₂-*bb*-**DSS**) were used as starting materials to synthesize three S-shaped double helicenes (i.e., **DH-1**, **DH-2**, and **DH-3**) through monobromination, formylation, the Wittig reaction, and double oxidative photocyclization. The photocyclization was a highly regioselective process. The molecular structures of **DH-1** and **DH-2** were confirmed by X-ray single-crystal analysis. Multiple intermolecular interactions, such as C–S, C–Se, S–S, S–Se, and Se–Se, were observed in the crystal packing structures of these compounds. Spectroscopic results and our previous work showed that the combination of molecular structure change and heteroatom replacement from S to Se could precisely modulate molecular energy levels.

Introduction

Given their esthetically pleasing helical structures, inherent helical chirality, and extended π -conjugation, helicenes have attracted extensive research attention. Helicenes are generally divided into carbohelicenes and heterohelicenes. The rapid development of carbohelicenes has led to the synthesis of double, triple, quadruple, quintuple, and sextuple molecules, the chiropotical properties of these molecules, such as their circular dichroism (CD) and circularly polarized luminescence (CPL), have also been widely studied [1-8]. The development of thia-

helicene, a class of typical heterohelicenes, has led to the preparation of symmetric thiophene-based [5]-, [7]-, [9]-, and [11]helicenes [9-16], unsymmetric thiophene-based [7]helicenes [17], and thiophene-based double helicenes with spiro-silicon atoms [18], "saddle" formed 8π annulene [12], and twisted naphthalene as central spacers [19].

As its close analogue, selenophene has properties very similar to those of thiophene. Fused aromatic compounds containing

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selenophene units show favorable optical and electrochemical properties and improved charge transport characteristics in the solid state mainly because such fused aromatic compounds often undergo increased Se–Se interactions, which confer ordering at the molecular scale and, thus, lead to well-aligned solid-state packing and excellent charge-transport properties [20,21].

However, as an important type of heteroacenes, fused selenophenes have rarely been reported in the literature because their synthesis is extremely challenging. The first seven-ringfused heteroacene containing selenophene was synthesized through the intramolecular triple cyclization of bis(ohaloaryl)diacetylene by Yamaguchi in 2005 [22]. Using a similar method, Takimiya reported the synthesis of six-ring-fused and four-ring-fused heteroacenes containing selenopheno[3,2b|selenophene in 2007 and 2009, respectively [23,24]. Five years later, Cheng reported the synthesis of two types of fivering-fused isomers of diselenopheno[2,3-b:7,6-b']fluorene and diselenopheno[3,2-b:6,7-b']fluorene through the cyclization of terminal acetylene as well as six types of biselenophene-based fused tricyclic derivatives [25,26]. In 2017, we reported the first member of diselenoselenophenes (DSSs), 2,5-di(trimethylsilanyl)diseleno[2,3-b:3',2'-d]selenophene ((TMS)₂-bb-**DSS**), from which the TMS group could easily be removed by trifluoroacetic acid and replaced by bromine [27]. Another isomer of **DSS**, diseleno[3,2-b:2',3'-d]selenophene (tt-**DSS**) has been successively synthesized from selenophene [28,29].

Among the limited fused aromatic compounds containing selenophene currently available, helicenes have received relatively little attention [27]. In our previous work, bull hornshaped selenophene-based heteroacenes (1, Figure 1) and selenophene-based [7]helicene (2, Figure 1) were synthesized

from the (TMS)₂-substituted selenophene triacenes, 2,5-di(trimethylsilanyl)diseleno[3,2-*b*:2',3'-*d*]selenophene ((TMS)₂-*tt*-**DSS**) and (TMS)₂-*bb*-**DSS**, respectively. In this work, the novel S-shaped double helicene **DH-3** (Figure 1), which is based on selenophene units, was constructed as a new member of the selenohelicene family (Figure 1).

In this synthetic study, regioselective double oxidative photocyclization was observed during the construction of three S-shaped double helicenes **DH-1–3** based on thiophene/selenophene. From **DH-1** to **DH-3**, sulfur atoms in the molecular framework were gradually replaced by selenium atoms. The crystal structures of **DH-1** and **DH-2** and spectroscopic features of **DH-1–3** were then studied. Finally, the reaction sites of oxidative photocyclization, energy levels, and the electron cloud distribution of the highest occupied molecular orbitals (HOMOs) and the lowest unoccupied molecular orbitals (LUMOs) are predicted.

Results and Discussion Synthesis of **5a–c** and **DH-1–3**

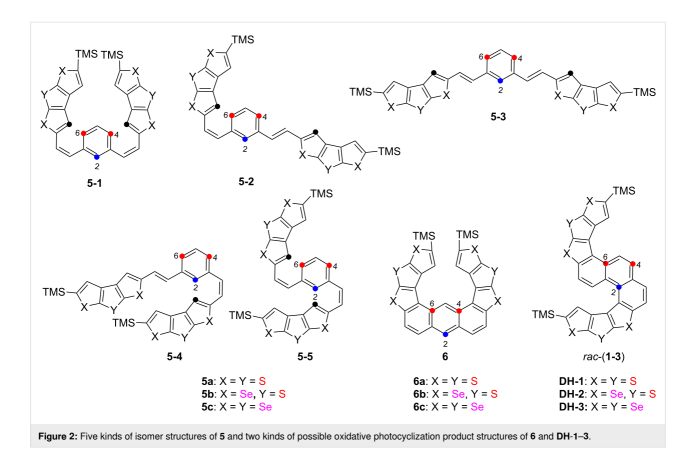
The synthetic route of 1,3-bis(2-(5-(trimethylsilyl)dithieno-[2,3-b:3',2'-d]thiophen-2-yl)vinyl)benzene (**5a**), 1,3-bis(2-(5-(trimethylsilyl)diselenopheno[2,3-b:3',2'-d]thiophen-2-yl)vinyl)benzene (**5b**), 1,3-bis(2-(5-(trimethylsilyl)diselenopheno[2,3-b:3',2'-d]selenophen-2-yl)vinyl)benzene (**5c**), and S-shaped double helicenes **DH-1-3** is shown in Scheme 1. The double oxidative photocyclization of **5a–c** is the key step in the synthesis of **DH-1-3** because oxidative photocyclization induces double radicals on a double bond, which led to the C=C bond rotation along the resulting single C(radical)–C(radical) bond and randomly directed annelated products [30]. Moreover, compounds **5a–c** bear two C=C bonds, which may lead to more complex photocyclization products.

5-(Trimethylsilyl)dithieno[2,3-b:3',2'-d]thiophene-2-carbaldehyde (4a) [19], 5-(trimethylsilyl)diseleno[2,3-b:3',2'-d]thiophene-2-carbaldehyde (4b) [27], and 5-(trimethylsilyl)diseleno[2,3-b:3',2'-d]selenophene-2-carbaldehyde (4c) [27] were prepared via monobromination and formylation reactions with (TMS)₂-bb-DTT, (TMS)₂-bb-DST, and (TMS)₂-bb-DSS as starting materials according to the literature. In (TMS)2-bb-**DTT**, $(TMS)_2$ -bb-**DST**, and $(TMS)_2$ -bb-**DSS**, the notation bb means that the positions of the heteroatoms of the outer two heterocycles are on the same side as those of the middle heterocycle. Taking dithienothiophene as example, according to the positions of sulfur atoms, there are six isomers of dithienothiophene, in which the sulfur atoms on the same side is defined as bb by us. The double Wittig reaction of 4a and 4b with 1,3phenyldimethyltriphenylphosphonium bromide afforded compounds 5a and 5b with yields of 46% and 64%, respectively. After the double Wittig reaction of 4c we obtained a mixture of cis and trans isomers of 5c with the total yield of 50%. According to the results of ¹H NMR the ratio of *cis* and *trans* isomers was approximately 1:0.25 (Scheme 1).

Compounds 5 had five isomers and three reaction sites (2, 4, and 6-positions in the benzene moiety) during oxidative photocyclization. Irradiation of 5a-c resulted in oxidative photocyclization products with two types of configurations wherein two benzene rings were closed in the same and opposite direction, such as DH-1-3 and 6 (Figure 2). However, after

the double oxidative photocyclizations of **5a–c** in the presence of iodine and propylene oxide in dry toluene through irradiation by a 450 W Hg medium-pressure lamp for 1.5 h, only one type of ring-closing product with two benzene rings formed in the same direction, i.e., S-shaped double helicenes **DH-1–3** were obtained in yields of 62%, 30%, and 53%, respectively (Figure 2).

The double oxidative photocyclization reaction sites of 5 were predicted by the orbital-weighted Fukui function in Multiwfn using Gaussian 09 [31,32] at the B3LYP/6-31G** [33] level of theory to verify the reaction-site selectivity of oxidative photocyclization further. Results showed that the conformations of 5a are varied, but the orbital-weighted Fukui function is not connected to conformation. In the orbital-weighted Fukui function, the larger the isosurface distribution, the higher the activity of reactions. Thus the 4 and 6-positions of benzene are the most likely sites for reaction (see Supporting Information File 1, Figure S21). However, after the formation of the naphthalene ring, the α -position is the most likely site for reaction (Supporting Information File 1, Figure S22). Therefore, after the double oxidative photocyclization of compound 5a, product DH-1 is mainly obtained. The predicted result of the double oxidative photocyclizations of 5a is consistent with the experimental result, that is, an S-shaped double helicene can be selectively obtained through the double oxidative photocyclization of compound 5a.



Crystallographic analyses of DH-1 and DH-2

The molecular structures of **DH-1** and **DH-2** were confirmed by single-crystal analysis (Figure 3). Both **DH-1** and **DH-2** belong to the triclinic space group *P*-1. After double oxidative photocyclizations of **5a** and **5b**, **DH-1** and **DH-2** are compressed into S-shaped double helical structures (Figure 3A and B), which consist of one [5]helicene and one [6]helicene. The two helicenes have the same configuration and bend toward the same direction on the same side of the shared naphthalene ring (Figure 3C and D). Both products **DH-1** or **DH-2** feature a pair of enantiomers *MM* and *PP* in their unit cell (see Supporting Information File 2, Figures S2 and S6). The crystal parameters of **DH-1** and **DH-2** are shown in Table 1. The replacement of sulfur with selenium in **DH-1** and **DH-2** leads to turn angles in-plane and helix climbs of [5]helicene and [6]helicene of **DH-1** and **DH-2** significantly change (Table 1, Figure 3).

Each of the two **DH-1** molecules (blue and red molecules in Figure S3 of Supporting Information File 2) in the unit cell interacts with six adjacent molecules. For example, in Figure S3 (Supporting Information File 2), the blue molecule interacts with six adjacent molecules through multiple interactions, such as, S···S (3.44 Å), S···H (2.94 Å, 2.91 Å, 2.97 Å), H···H (2.31 Å), C···C (3.31 Å), S···C (3.45 Å), and C···H (2.90 Å). Each of the two **DH-2** molecules (blue and red molecules, Sup-

porting Information File 2, Figure S7) in the unit cell interact with eight adjacent molecules through multiple interactions, such as, C···C (3.40 Å), Se···H (3.09 Å), H···H (2.26 Å), C···H (2.90 Å), Se···Se (3.62 Å), and Se···S (3.58 Å). However, in contrast to **DH-1**, **DH-2** exhibits multiple S···H (2.88 Å), S···C (3.45 Å), and Se···H (2.94 Å) interactions between two molecules in the unit cell. These multiple interactions confer **DH-1** and **DH-2** with a regular arrangement featuring multi channels (Figures S4 and S8 in Supporting Information File 2), which is a suitable characteristic for helicene compounds used as supramolecular self-assembly units [34-41].

Spectroscopic features of **DH-1–3**

The UV–vis absorption spectra of **DH-1–3** in dichloromethane are shown in Figure 4. In general, the UV–vis absorption spectra of **DH-1–3** are generally similar in shape and exhibit three major absorption bands within 230–280 nm (band-II), 280–330 nm (band-II), and 304–414 nm (band-III) (Figure 4). Progressive red-shifts in the absorption spectra of band-I, band-II, and band-III occur with increasing number of selenium atoms. In band-I, compounds **DH-1–3** show a maximum absorption peak at 232, 240, and 242 nm, respectively. In band-II and band-III, helical distortion and possible conjugation through heteroatoms (e.g., sulfur and selenium atoms) in **DH-1–3** may increase π -electron delocalization, leading to a

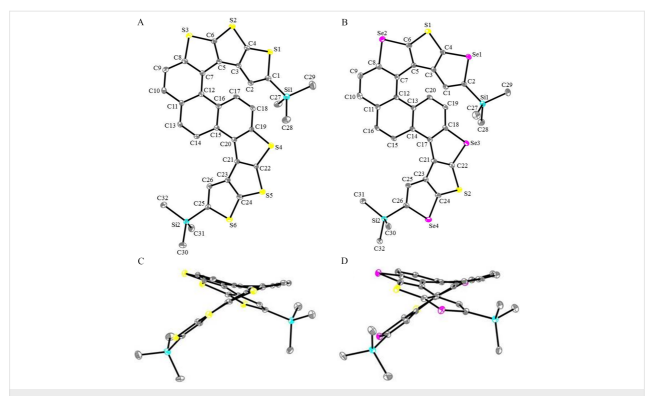


Figure 3: Molecular structures and side view for DH-1 and DH-2. A and B are molecular structures for DH-1 and DH-2. C and D are side views for DH-1 and DH-2. Carbon, sulfur, selenium, and silicon atoms are depicted with thermal ellipsoids set at the 30% probability level and all hydrogen atoms are omitted for clarity.

Compound		Dihedral angle (°) ^a	Torsion angle (°) ^b	Turn angle in-plane (°) ^c	Helix climbs (Å) ^d
DH-1	[5]helicene	25.00	38.64	231.1	0.932
	[6]helicene	50.61	64.38	273.6	2.424
DH-2	[5]helicene	25.01	37.27	230.9	0.879
	[6]helicene	48.16	64.12	279.3	2.462

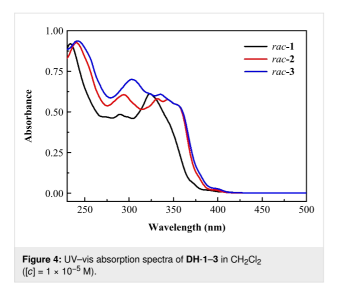
^aDihedral angle between two terminal rings of helicene. ^bThe sum of the three ([5]helicene) or four ([6]helicene) distortion angles. ^cThe angle of turns in-plane. ^dInternal helix climb height.

red-shifted broad absorption. The maximum absorption peaks of **DH-1–3** appear at 268, 275, and 279 nm, respectively, in band-III and at 323, 331, and 336 nm, respectively, in band-III. Thus, the optical band gaps estimated from the absorption edges gradually decrease from **DH-1** to **DH-2** to **DH-3**, and are equal to 3.08, 3.01, and 2.98 eV, respectively. This change trend is consistent with the calculated results, which are 3.97, 3.83, and 3.81 eV for **DH-1**, **DH-2**, and **DH-3**, respectively (see Table S2 in Supporting Information File 1). However, the optical band gaps of **1**, **2**, and **DH-3** obviously differ because of changes in their molecular configuration and equal to 2.86, 3.15, and 3.81 eV, respectively [27,28]. As the number of selenium atoms increases from **DH-1** to **DH-3**, the fluorescence intensity (Figure S19 in Supporting Information File 1) and the fluores-

cence quantum yield (Φ_F , Figure S20, Table S1 in Supporting Information File 1) of the molecules also decrease (Figure 4).

Conclusion

In summary, the key step of regioselective double oxidative photocyclization was successfully employed in the preparation of three S-shaped double helicenes, namely, **DH-1**, **DH-2** and **DH-3** with (TMS)₂-bb-**DTT**, (TMS)₂-bb-**DST**, and (TMS)₂-bb-**DSS** as starting materials. The synthetic method described in this research not only provides a method for the synthesis of S-shaped double helicenes but also enriches the family of selenophene helicenes. Multiple intermolecular interactions and regular arrangement in the crystal packing structures of **DH-1** and **DH-2** indicate that these compounds may be used as supra-



molecular self-assembly units. Changes in molecular structure may substantially modulate the optical band gap of heteroacenes, and the replacement of heteroatoms from S to Se could fine-tune their optical band gap [18,27,28]. Thus, the combination of molecular structure modification and atom replacement could be a viable strategy, for the precise modulation molecular energy levels and yield molecules with strong application potential in organic functional materials, such as OFETs, and CPLs, among others.

Experimental

General procedures and materials

Tetrahydrofuran (THF) for use on vacuum line was freshly distilled from sodium/benzophenone prior to use. *n*-BuLi (hexane) were obtained from Energy Chemical; prior to use, its concentration was determined by titration with *N*-pivaloyl-*o*-toluidine [42]. Column chromatography was carried out on silica gel (300–400 mesh). Analytical thin-layer chromatography was performed on glass plates of silica gel GF-254 with detection by UV. Standard techniques for synthesis under inert atmosphere and Schlenk glassware equipped with an 8 mm PTFE vacuum stopcock, were employed. All starting materials and reagents were commercially available.

¹H NMR and ¹³C{¹H} NMR spectra were recorded on 300 or 400 MHz NMR instruments using CDCl₃ as solvent. The chemical shift references were as follows: (¹H) CDCl₃, 7.26 ppm (CHCl₃); (¹³C{¹H}) CDCl₃, 77.00 ppm (CDCl₃). IR spectra were obtained using an FTIR instrument. MS analysis was carried out on mass spectrometers equipped with EI (70 eV). HRMS analysis was carried out on a mass spectrometer equipped with DART-FT-ICR and MALDI-TOF-CHCA. Melting-point determination was taken on a Melt-Temp apparatus and mp are uncorrected. The X-ray crystallographic

analyses were performed using crystals of compounds **DH-1** and **DH-2** with sizes of $0.14 \times 0.12 \times 0.08$, $0.21 \times 0.17 \times 0.12$ mm³, respectively. The intensity data were collected with the ω scan mode (296 K) on a diffractometer with a CCD detector using Cu K α radiation (λ = 1.54184 Å). The data were corrected for Lorentz and polarization effects, and absorption corrections were performed using the SADABS program [43]. The crystal structures were solved using the SHELXTL program and refined using full-matrix least-squares [44]. Further details are in the deposited CIF files. Slow evaporation of solutions of **DH-1** and **DH-2** in CHCl₃/CH₃OH 5:1 (v/v) were employed for growing single crystals. The fluorescence quantum yields (Φ_F) of **DH-1–3** are characterized in dichloromethane with quinine sulfate in 0.1 N H₂SO₄ as the control.

Synthesis of 5a-c and DH-1-3

Synthesis of 1,3-bis(2-(5-(trimethylsilyl)dithieno[2,3-b:3',2'-d]thiophen-2-yl)vinyl)benzene (**5a**)

n-BuLi (0.06 mL, 2.50 M in hexane, 0.168 mmol, 2.5 equiv) was added dropwise to 1,3-bis[methyl(bromotriphenylphosphonium)] (53.3 mg, 0.067 mmol) in THF (40 mL) at -78 °C. After 2 h at -78 °C, a solution of **4a** (41.0 mg, 0.138 mmol, 2.05 equiv) in THF (5 mL) was added at -78 °C, the mixture kept for 1 h, and then the reaction mixture was warmed up slowly to ambient temperature overnight. The reaction mixture was quenched with CH₃OH and extracted with CH₂Cl₂ (3 × 10 mL). The organic layer was washed with saturated NaCl (20 mL) and water (2 × 20 mL), and then dried over MgSO₄. The residue was purified by column chromatography (eluent: hexane/CH₂Cl₂ 3:1 (v/v) and recrystallized from CHCl₃/ CH₃OH to yield **5a** (20.9 mg, 46%) as a yellow solid; mp > 300 °C; 1 H NMR (400 MHz, CDCl₃) δ (ppm) 7.58 (s, 1H), 7.45 (s, 2H), 7.41–7.36 (m, 3H), 7.34 (s, 2H), 7.31 (d, J =16.0 Hz, 2H), 6.93 (d, J = 16.0 Hz, 2H), 0.39 (s, 18H); ¹³C{¹H} NMR (100 MHz, CDCl₃) δ (ppm) 145.3, 144.7, 143.7, 140.7, 138.4, 137.8, 137.2, 129.1, 127.8, 125.5, 124.9, 124.4, 122.8, 118.0, -0.1; IR (KBr): 3018, 2958, 2849, 1631, 1408, 1360, 945, 837 cm⁻¹; EIMS (70 eV) m/z: [M]⁺ 662.18 (40); HRMS-MALDI (m/z): [M]⁺ calcd for C₃₂H₃₀S₆Si₂, 662.0216; found, 662.0205.

1,3-Bis(2-(5-(trimethylsilyl)diselenopheno[2,3-b:3',2'-d]thiophen-2-yl)vinyl)benzene (**5b**) was synthesized according to the procedure described for compound **5a**.

5b: yellow solid in yield of 64% (46.2 mg); mp > 300 °C; ¹H NMR (300 MHz, CDCl₃) δ (ppm) 7.73 (s, 2H), 7.55 (d, J = 12.9 Hz, 3H), 7.38–7.32 (m, 5 H), 6.79 (d, J = 15.6 Hz, 2H), 0.38 (s, 18H); ¹³C{¹H} NMR (100 MHz, CDCl₃) δ (ppm) 151.6, 149.7, 144.5, 142.6, 141.1, 137.01, 136.98, 129.2, 128.6, 127.5, 125.6, 124.9, 124.4, 121.0, 0.2; IR (KBr): 3011, 2949, 2891, 1627, 1427, 1367, 920, 831 cm⁻¹; DARTMS (m/z): [M + H]⁺ 854.8 (75); HRMS-DART (m/z): [M + H]⁺ calcd for $C_{32}H_{31}S_2Se_4Si_2$, 854.8071; found, 854.8067.

1,3-Bis(2-(5-(trimethylsily1)diselenopheno[2,3-b:3',2'-d]selenophen-2-yl)vinyl)benzene (**5c**) was synthesized according to the procedure described for compound **5a**.

Mixture of cis and trans isomers 5c: yellow solid in yield of 50% (24.4 mg); mp > 300 °C; 1 H NMR (400 MHz, CDCl₃) δ (ppm): 7.76 (s, trans-), 7.73 (s, cis- and trans-), 7.72 (s, cis- and trans-), 7.57 (s, cis- and trans-), 7.56 (s, trans-), 7.49-7.33 (m, trans-, cis- and trans-), 7.30 (d, J = 16.0 Hz, cis- and trans-), 6.90 (d, J = 12.0 Hz, cis- and trans-), 6.79 (d, J = 16.0 Hz, trans-), 6.78 (d, J = 16.0 Hz, trans- and trans-), 6.65 (d, J =12.0 Hz, cis- and trans-), 0.38 (s, trans-), 0.36 (s, cis- and trans-), 0.35 (s, cis- and trans-), the ratio of integral areas of the peaks at 7.76, 7.73, and 7.72 ppm is 1.0:0.25; ¹³C{¹H} NMR (125 MHz, CDCl₃) δ (ppm): 152.78, 152.70, 152.62, 151.28, 151.23, 147.31, 146.49, 146.34, 143.49, 143.47, 141.84, 140.87, 140.81, 140.72, 139.02, 137.66, 137.50, 137.43, 135.39, 135.36, 129.74, 129.30, 129.28, 129.23, 129.05, 128.83, 128.42, 128.35, $127.50,\, 126.97,\, 126.28,\, 125.68,\, 125.20,\, 125.10,\, 125.04,\, 124.85,\,$ 124.63, 122.61, 122.51, 0.43, 0.41; IR (KBr): 3010, 2951, 2889, 1624, 1413, 1369, 908, 833 cm⁻¹; MALDIMS (m/z): [M]⁺ 949.8; HRMS-DART-FT (m/z): [M]⁺ calcd for C₃₂H₃₀Se₆Si₂, 949.6894; found, 949.6887.

Synthesis of **DH-1**

To a solution of 5a (9.6 mg, 0.014 mmol) in dry toluene (6 mL), iodine (7.3 mg, 0.028 mmol, 2.0 equiv) and excess propylene oxide were added. The reaction solution was irradiated with a 450 W unfiltered Hg medium-pressure lamp for 1.5 h. The reaction was quenched with saturated Na₂SO₃ solution (5 mL). The reaction mixture was extracted with CH2Cl2 (3 × 5 mL), washed with H_2O (3 × 10 mL), and then dried over MgSO₄. After removing the solvent in vacuum, the crude product was purified by PTLC with petrol ether (60-90 °C) and hexane/ CH_2Cl_2 5:1 (v/v) as developer to yield **DH-1** (5.9 mg, 62%) as a light-yellow solid; mp > 300 °C; ¹H NMR (400 MHz, CDCl₃) δ (ppm) 9.09 (d, J = 8.8 Hz, 1H), 8.51 (d, J = 8.8 Hz, 1H), 8.22 (s, 1H), 8.11 (d, J = 9.2 Hz, 1H), 8.04 (d, J = 8.4 Hz, 1H), 7.90(d, J = 8.4 Hz, 1H), 7.78 (d, J = 8.8 Hz, 1H), 6.62 (s, 1H), 0.48(s, 9H), 0.14 (s, 9H); ¹³C{¹H} NMR (100 MHz, CDCl₃) δ (ppm) 144.6, 144.1, 143.19, 143.18, 142.1, 141.7, 141.6, 141.4, 141.3, 141.1, 135.0, 133.4, 130.6, 129.8, 129.0, 128.5, 127.53, 127.46, 127.3, 127.0, 126.7, 126.4, 124.3, 123.7, 121.5, 119. 7, 0.0, -0.4; IR (KBr): 3045, 2952, 2860, 1645, 1487, 1410, 972, 837 cm⁻¹; EIMS (70 eV) m/z: [M]⁺ 657.87 (30); HRMS-MALDI (m/z): [M]⁺ calcd for C₃₂H₂₆S₆Si₂ 657.9897; found, 657.9892.

DH-2 was synthesized according to the procedure described for compound **DH-1**.

DH-2: light-yellow solid in yield of 30% (15.8 mg); mp > 300 °C; ¹H NMR (400 MHz, CDCl₃) δ (ppm) 8.92 (d, J = 8.8 Hz, 1H), 8.54 (s, 1H), 8.32 (d, J = 8.4 Hz, 1H), 8.04 (d, J = 8.0 Hz, 1H), 7.95 (d, J = 8.8 Hz, 1H), 7.80 (d, J = 8.0 Hz, 1H), 7.66 (d, J = 8.8 Hz, 1H), 6.66 (s, 1H), 0.47 (s, 9H), -0.01 (s, 9H); ¹³C{¹H} NMR (100 MHz, CDCl₃) δ (ppm) 149.9, 146.9, 145.7, 145.1, 144.8, 144.3, 142.7, 142.6, 139.8, 139.4, 139.2, 137.5, 132.8, 130.8, 130.6, 130.2, 129.59, 129.58, 128.4, 128.1, 126.7, 125.6, 124.5, 124.3, 124.2, 122.2, 0.3, -0.2; IR (KBr): 3060, 2949, 2896, 1632, 1489, 1402, 933, 829 cm⁻¹; DARTMS (m/z): [M + H]⁺ 850.8 (45); HRMS-DART (m/z): [M + H]⁺ calcd for C₃₂H₂₇S₂Si₂Se₄, 850.7753; found, 850.7743.

DH-3 was synthesized according to the procedure described for **DH-1**.

DH-3: light yellow solid in yield of 53% (19.5 mg); mp 265.1–266.9 °C; ¹H NMR (300 MHz, CDCl₃) δ (ppm) 8.85 (d, J = 8.7 Hz, 1H), 8.56 (s, 1H), 8.22 (d, J = 8.7 Hz, 1H), 8.06 (d, J = 8.4 Hz, 1H), 7.92 (d, J = 9.0 Hz, 1H), 7.81 (d, J = 8.7 Hz, 1H), 7.63 (d, J = 8.7 Hz, 1H), 6.68 (s, 1H), 0.45 (s, 9H), -0.11 (s, 9H); ¹³C{¹H} NMR (150 MHz, CDCl₃) δ (ppm) 151.0, 148.0, 147.4, 147.0, 145.9, 143.9, 142.6, 141.1, 140.9, 139.34, 139.26, 138.9, 134.1, 132.3, 131.7, 130.7, 130.2, 129.3, 128.8, 128.0, 126.7, 125.0, 124.9, 124.1, 123.9, 121.9, 0.4, -0.3; IR (KBr): 3047, 2947, 2895, 1633, 1504, 1409, 925, 831 cm⁻¹; DARTMS (m/z): [M]⁺ 945.8; HRMS-DART-FT (m/z): [M]⁺ calcd for C₃₂H₂₆Se₆Si₂, 945.6564; found, 945.6564.

Supporting Information

Supporting Information File 1

Spectral and computational data.

[https://www.beilstein-journals.org/bjoc/content/supplementary/1860-5397-18-81-S1.pdf]

Supporting Information File 2

Supporting crystallographic information for compounds **DH-1** and **DH-2**.

[https://www.beilstein-journals.org/bjoc/content/supplementary/1860-5397-18-81-S2.pdf]

Supporting Information File 3

Title CIF file for compound DH-1.

[https://www.beilstein-journals.org/bjoc/content/supplementary/1860-5397-18-81-S3.cif]

Supporting Information File 4

Title CIF file for compound DH-2.

[https://www.beilstein-journals.org/bjoc/content/supplementary/1860-5397-18-81-S4.cif]

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