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Germanium- and tin-bridged diazulenylnmethyl cations: effects of group 14 elements on the structure and properties of π -extended cations†

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Diazulenylnmethyl cations bridged with a germanium and tin moiety were synthesized. In these cations, the nature of these elements has impacts on the chemical stability and photophysical properties. Upon aggregation, these cations exhibit absorption bands in the near-infrared region, which are slightly blue-shifted compared to those of silicon-bridged congeners.

Cationic π -conjugated compounds have attracted considerable attention due to their wide range of applications¹ including conducting materials,^{1c,d} dyes for photothermal conversion and bioimaging,^{1g} and sensitizers for photoreactions.^{1e} While various types of cationic π -systems have been developed to date, their properties largely depend on the nature of the π -skeleton employed and fine-tuning of suitable counter anions. A key issue in the design of novel classes of cationic π -skeletons is the need to attain effective charge delocalization that can furnish high chemical stability and unusual electronic structures.² Another important matter for imparting π -skeletons with valuable functionality is controlling the molecular arrangement in the aggregated and solid state.³

In this context, we have recently disclosed silicon-bridged diazulenylnmethyl cations **1** as a new class of stable carbocations (Fig. 1),⁴ where the positive charge is effectively delocalized over the π -skeleton, reflecting the highly polarizable character of azulenyl moieties.^{5,6} Specifically, the connection of two azulenyl groups at the 1-position to the carbocation center, while reinforcing the skeleton in a planar fashion with a Si-bridge, is crucial for the high chemical stability of the system. This stabilization mode is different from that found in conventional cationic π -systems, where nitrogen- and oxygen-based electron-donating groups are usually used.² The Si-

bridge also affects the molecular arrangement in the condensed state.⁷ The observed offset π -stacked arrangement is due to the steric congestion of the substituents on the silicon bridge and the positioning of the counter anions. Consequently, J-type aggregates^{3b} with sharp absorption and emission bands in the near-infrared (NIR) region were formed.⁸ Such NIR chromophores have a promising variety of prospective applications in fields, such as aggregation-induced staining for bioimaging.^{1g,8b,d} Hence, with this prominent π -skeleton in hand, further diversification of its properties became our next subject. A promising approach to achieving this objective would be *via* the introduction of other main group elements at the bridging position.⁹ Herein, we have introduced the heavier group 14 elements, Ge and Sn,^{10,11} into diazulenylnmethyl cations. Relative to silicon,¹⁰ these heavier elements are characterized by larger atomic radii, lower electronegativity, less effective orbital interactions with the π -conjugated skeleton, and heavy-atom effects. We synthesized Ge- and Sn-bridged diazulenylnmethyl cations **2** and **3** (Fig. 1), and investigated the effects of these heavier elements on the stability, structures, and properties in solution as well as in the aggregated and solid states.

Following the synthetic method reported for the Si-bridged diazulenylnmethyl cation **1**,⁴ Ge- or Sn-bridged **2**_{PF₆} and **3**_{PF₆} were synthesized in three steps from 2-iodoazulene (Scheme 1). The key step is the cyclization of diazulenyl-substituted precursors **4** and **5** under Vilsmeier formylation con-

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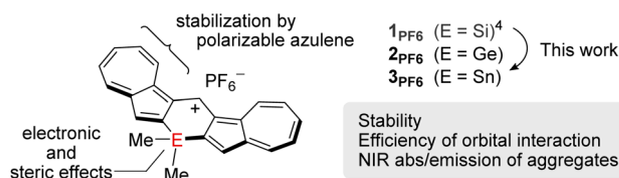
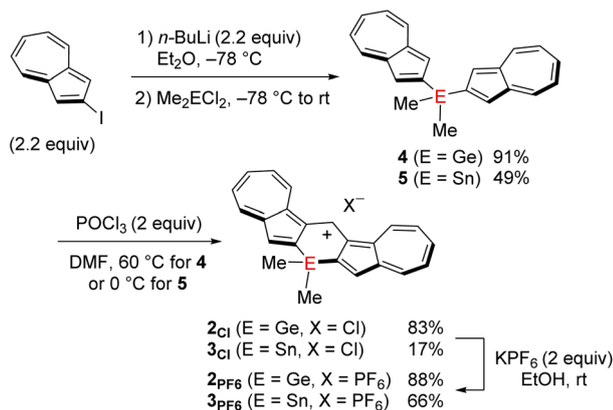


Fig. 1 Design and structures of diazulenylnmethyl cations **1–3** with group 14 elements as bridging moieties.





Scheme 1 Synthesis of 2 and 3.

ditions,⁴ to produce **2_{Cl}** and **3_{Cl}**, respectively. While the reaction of **4** at 60 °C selectively afforded **2_{Cl}**, similar to the synthesis of the silicon congener **1_{Cl}**, a lower reaction temperature was required for the cyclization of **5** in order to prevent the formation of unidentified products that are likely formed *via* cleavage of the C–Sn bond. Subsequently, **2_{Cl}** and **3_{Cl}** were treated with **KPF₆** to afford **2_{PF₆}** and **3_{PF₆}**, which were isolated as deep blue solids.

Whereas **2** and **3** are formally classified as secondary carbocations, they are stable toward air and moisture and they can be stored in air without any precautions irrespective of the counter anions. The chemical stability of **2_{PF₆}** and **3_{PF₆}** was assessed by determining their pK_{R^+} values, which represent their affinity toward a hydroxide ion.¹² The pK_{R^+} values were estimated to be 10.5 and 11.5 for **2_{PF₆}** and **3_{PF₆}**, respectively (Fig. S1†). These values are higher than those of Si-bridged **1_{PF₆}** ($pK_{R^+} = 9.7$)⁴ and trioxatriangulenium hexafluorophosphate ([TOTA][PF₆]; $pK_{R^+} = 9.3$),¹² the latter of which is a representative stable trityl cation (the structure is shown in Fig. S1†). This comparison demonstrates the importance of the nature of heavier elements for chemical stability.

Single crystals of **2_{PF₆}** suitable for X-ray diffraction analysis were obtained by the slow evaporation of a CH₂Cl₂/hexane solution at room temperature (Fig. 2a). In the crystalline state,

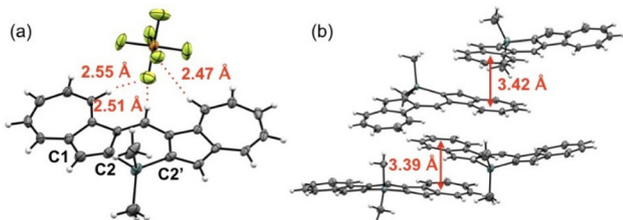


Fig. 2 (a) The molecular structure of **2_{PF₆}** with thermal ellipsoids at 50% probability and (b) molecular arrangement of **2_{PF₆}** in the crystal state. Blue: Ge, orange: P, yellow: F, white: H, gray: C. While the unit cell of **2_{PF₆}** contains two crystallographically independent molecules, only one of them is shown in (a). The relevant distances for the F...H–C hydrogen bonds and the representative C...C distance between the two adjacent seven-membered rings are shown.

2_{PF₆} exhibits a curved π -conjugated skeleton, in which the **PF₆[–]** anion sits in the bay area with hydrogen bonds between the fluorine and the hydrogen atoms of the two azulene and methene moieties. Similar to the structure of **1_{PF₆}**,⁴ the bond length alternation in the seven-membered ring of **2_{PF₆}** is small (1.377(8)–1.403(8) Å), whereby the C1–C2 bond (1.372(9) Å) exhibits a pronounced double bond character (Fig. S2†). This structure stands in contrast to the structure of the parent azulene, which exhibits a more pronounced bond length alternation in its seven-membered ring (1.3679(8)–1.4155(8) Å).¹³ The relatively small bond length alternation in **2_{PF₆}** reflects a large contribution of an aromatic tropylium ion, which is responsible for the effective delocalization of the positive charge and thereby its high chemical stability. The most significant difference is the geometry of the central six-membered ring, wherein the Ge atom sits slightly above/below the plane formed by the remaining five carbon atoms, while **1_{PF₆}** exhibits a more planar geometry. This difference likely results from the longer endocyclic Ge–C bond lengths (Ge–C2, 1.946(6) Å) and the smaller C2–Ge–C2' angle in **2_{PF₆}** (98.5°) compared to those in **1_{PF₆}** (Si–C2, 1.862(3) Å; C2–Si–C2', 99.8°).⁴

In the crystal state, molecules of **2_{PF₆}** adopt an offset π -stacked arrangement, in which the outer seven-membered rings overlap with the closest interfacial distances of 3.39 and 3.42 Å (Fig. 2b). This packing mode is similar to that observed for **1_{PF₆}**,⁴ which implies that this is a common feature of the structurally constrained diazulenomethene skeleton that is probably a result of the location of the **PF₆[–]** anion and the steric repulsion of the vertically oriented substituents on the Ge or Si bridge (Fig. 2 and S6†).⁷

The electronic effects resulting from the Ge- and Sn-bridges were studied by measuring the photophysical and electrochemical properties of **2_{PF₆}** and **3_{PF₆}**. In the UV-visible absorption spectra of **2_{PF₆}** and **3_{PF₆}** in CH₂Cl₂, intense absorption bands were observed with maximum wavelengths (λ_{abs}) at 639 and 637 nm, respectively, which are slightly blue-shifted compared to that of **1_{PF₆}** ($\lambda_{\text{abs}} = 646$ nm) (Fig. 3a). The λ_{abs} of **2_{PF₆}** and **3_{PF₆}** did not significantly change upon changing the solvent from CH₂Cl₂ to THF to DMSO (Table S2†). It should be

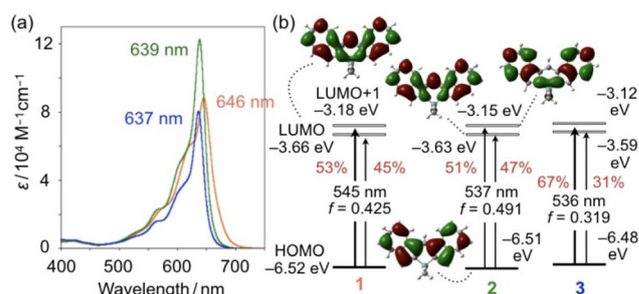


Fig. 3 (a) UV-vis absorption spectra of **1_{PF₆}** (orange), **2_{PF₆}** (green), and **3_{PF₆}** (blue) in CH₂Cl₂. (b) Kohn–Sham molecular orbitals and electronic transitions of **1–3** calculated at the M06/6-311+G(d,p) (for Si, Ge, C, and H) and the LANL2DZ (for Sn) levels of theory with CH₂Cl₂ using the polarizable continuum model. The red values in (b) represent the percentage probability for each S_0 – S_1 electronic transition.



noted that **2**_{PF6} has a larger molar absorption coefficient (ϵ) than **1**_{PF6} and **3**_{PF6}. The cyclic voltammogram of **2**_{PF6} in CH₂Cl₂ showed reversible redox waves for a one-electron reduction, similar to that of **1**_{PF6}, while that of **3**_{PF6} only showed an irreversible reduction wave (Fig. S8†). The cathodic peak potentials (E_{pc}) follow the order **1**_{PF6} (-0.73 V, $E_{1/2} = -0.69$ V) < **2**_{PF6} (-0.79 V, $E_{1/2} = -0.74$ V) < **3**_{PF6} (-0.84 V) relative to the ferrocene/ferrocenium couple (Fc/Fc⁺). These results suggest that the energy levels of the LUMOs of cationic diazulenylmethene skeletons increase upon introducing progressively heavier bridging elements.

To gain insights into the effects of bridging elements on the structures and electronic properties, density-functional theory (DFT) calculations were conducted for **1**, **2**, and **3** without including the counter anions (Fig. S6†) at the M06/6-311+G(d,p) (for Si, Ge, C, and H) and LANL2DZ (for Sn) levels of theory in CH₂Cl₂ using the polarizable continuum model (PCM). The optimized structure for **2** exhibits an endocyclic Ge–C bond length of 1.953 Å and a C–Ge–C angle of 98.3°, both of which are comparable to those obtained from the X-ray diffraction analysis (Fig. S2†). Although a satisfactory level of crystal data could not be obtained for **3**, the structural optimization revealed an endocyclic Sn–C bond length of 2.143 Å and a C–Sn–C angle of 93.5°, which are much longer and smaller than the corresponding values of **1** and **2** (Fig. S3†). This is an important structural effect caused by the bridging elements.

The HOMO and LUMO energy levels increase in the order **1** < **2** < **3**, albeit the differences between **1**–**3** are subtle (Fig. 3b). This tendency itself is consistent with the E_{pc} values obtained from cyclic voltammetry. Notably, while a $\sigma^*-\pi^*$ orbital interaction is observed between the exocyclic SiMe₂ moiety and the diazulenylmethene π -skeleton in the LUMO of **1**,¹⁴ no such orbital interaction is observed in the LUMOs of **2** and **3** when the MOs are represented with a threshold value of 0.017 (Fig. 3b and S4†).

Electrostatic potential (ESP) maps for **2** and **3** obtained at the same level of theory showed that a positive charge is effectively delocalized over the concave area of the molecules (Fig. S5†), which is similar to that for **1**. The natural charge at the methene carbon atom for each compound was obtained by natural population analysis (NPA). The obtained values for **1** (-0.026), **2** (-0.026), and **3** (-0.028) are comparable, which suggests that the impact of the heavier bridging elements on the delocalization of the positive charge is limited. On the other hand, slight differences between the LUMO levels of **1**, **2**, and **3** might be due to the differences in their stabilities as demonstrated by the assessment of their pK_{R_2} values (Fig. S1†), albeit a detailed rationale for these differences remains unclear at this stage (*vide supra*).

Time-dependent (TD) DFT calculations at the same level furnished the energies for the electronic S₀–S₁ transitions for **1**–**3**. The energy values are consistent with the order of the experimentally observed hypsochromic shifts of λ_{max} in the order **1** > **2** > **3** (Fig. 3b). These calculations also show that **2** exhibits the highest oscillator strength (f), which is consistent with the fact that **2**_{PF6} has the largest ϵ value among the three

compounds. It should also be noted here that S₀–S₁ transitions for **1**–**3** consist of HOMO–LUMO and HOMO–LUMO+1 transitions. Given that the contribution of the HOMO–LUMO+1 transition decreases, the oscillator strength value, and thereby the molar absorption coefficient, tend to increase. The HOMO and LUMO+1 of **1**–**3** consist of a combination of the HOMO and LUMO of the two parent azulene units. Thus, the S₀–S₁ transition in **2** has the least azulene character, which is most likely responsible for it having the largest molar absorption coefficient among the three compounds.

Even though the long E–C bonds in **2**_{PF6} and **3**_{PF6} result in distorted structures (Fig. S3†), these compounds retain the J-type aggregation behavior in solution, which is accompanied by absorption bands in the NIR region as observed for **1**_{PF6}.^{3b,4} When the methylcyclohexane (MCH) content of a 1,2-dichloroethane (DCE)/MCH solution of **2**_{PF6} was increased, the intensity of the original absorption band at 646 nm gradually decreased and a new broad band emerged at around 800 nm. The new absorption band became more prominent in a 20 : 80 (v/v) DCE/MCH solution ($\lambda_{max} = 810$ nm) and further blue-shifted to 773 nm in a 10 : 90 DCE/MCH solution (Fig. 4b). The powder X-ray diffraction pattern of the resulting aggregates of **2**_{PF6}, which were collected by filtration from the 10 : 90 DCE/MCH solution, showed good agreement with a pattern simulated from its crystal packing arrangement (Fig. S9†). This result implies that the offset molecular arrangement observed in the crystalline state is also formed, at least partly, in the

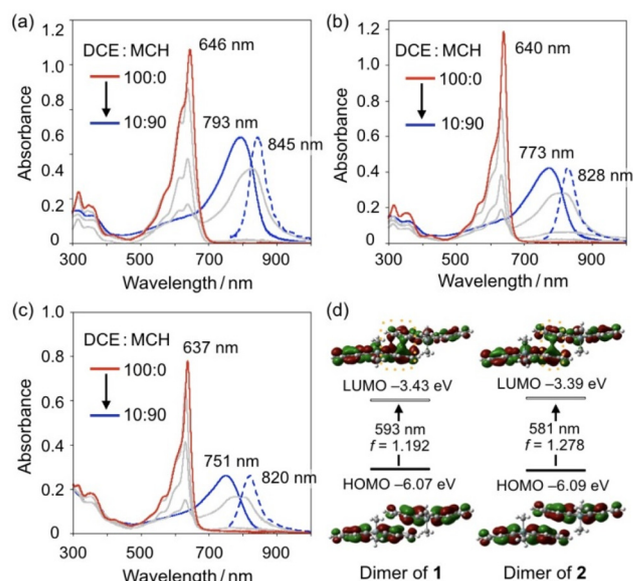


Fig. 4 UV-vis-NIR absorption spectra of (a) **1**_{PF6}, (b) **2**_{PF6}, and (c) **3**_{PF6} in various ratios of 1,2-dichloroethane (DCE)/methylcyclohexane (MCH) (v/v). Fluorescence spectra recorded in a 10 : 90 (v/v) DCE/MCH mixed solvent are shown as blue dashed lines. (d) Comparison of the S₀–S₁ electronic transition for the dimeric motifs of **1** and **2** together with their Kohn–Sham HOMOs and LUMOs calculated at the M06/6-311+G(d,p) level of theory. The geometries for the calculations were taken from their crystal packing structures and the counter ions have not been included.



aggregated state in solution. Compound **3_{PF6}** also exhibited a bathochromically shifted absorption band ($\lambda_{\text{max}} = 751 \text{ nm}$) when the DCE/MCH ratio was increased to 10 : 90 (Fig. 4c). The λ_{max} values of the aggregates exhibited hypsochromic shifts that follow the order **1_{PF6}** > **2_{PF6}** > **3_{PF6}**. Although there is no direct evidence due to the lack of crystal data for **3_{PF6}**, we infer from the crystal structure observed for **2_{PF6}** that the heavier elements distort the central six-membered ring, which reduces the degree of overlap of the adjacent molecules in the aggregated state, thus resulting in a blue-shifted absorption band in the NIR region. Notably, the difference in the λ_{max} in the aggregated state between **3_{PF6}** and **1_{PF6}** is 42 nm, while the difference between their monodispersed states is only 9 nm (Fig. 3a).

To gain insight into the bathochromic shift of the absorption bands upon aggregation, DFT calculations were conducted at the M06/6-311+G(d,p) level for two adjacent molecules of **2**, the coordinates of which were taken from the crystal packing structure of **2_{PF6}** (Fig. 4d). Compared to the monomer (Fig. 3b), the dimeric motif of **2** is characterized by HOMO and LUMO energy levels that are increased by 0.42 eV and 0.24 eV, respectively, and thereby the corresponding transition energy is by 0.198 eV lower, which is in agreement with a bathochromic shift of 44 nm. This calculation also showed a significant increase in the oscillator strength (f) for the dimeric motif. The offset arrangement of the electronic transition dipole of **2**, which is oriented along the long axis of the molecule (Fig. S4[†]), should be relevant to the increased f value, despite the limited overlap of the molecules only in the outer seven-membered ring (Fig. 2b). The difference between the absorption wavelengths of the aggregated states of **1_{PF6}** and **2_{PF6}** can also be reproduced by comparing the calculation results for the dimeric motifs of **1** and **2** (Fig. 4d). In the LUMO of the dimer of **2**, in-phase orbital interactions across the two molecules were observed (see the dotted area in Fig. 4d), the degree of which seems smaller than that of the dimer of **1**.

Although monodispersed **2_{PF6}** and **3_{PF6}** in solution are essentially non-emissive, fluorescence was observed upon the formation of J-type aggregates in their 10 : 90 DCE/MCH solutions (Fig. 4a–c). The emission maximum wavelengths of **2_{PF6}** and **3_{PF6}** were 828 and 820 nm, respectively, which were slightly blue-shifted compared to that of **1_{PF6}** (845 nm). The Stokes shifts for **2_{PF6}** (860 cm^{-1}) and **3_{PF6}** (1121 cm^{-1}) were larger than that for **1_{PF6}** (776 cm^{-1}), which might be related to the loosely packed aggregated structure in **3_{PF6}** that occurs due to the distorted geometry of its central six-membered ring (Fig. S3[†]). The quantum yields of the aggregates of **2_{PF6}** and **3_{PF6}** in 10 : 90 DCE/MCH ($\Phi_{\text{F}} < 0.01$) are lower than the previously reported value of the Si-bridged congeners ($\Phi_{\text{F}} \approx 0.04$).⁴

In conclusion, Ge- and Sn-bridged diazulenylmethyl cations were synthesized and the effects of these main group bridges on the stability and photophysical properties were studied. The incorporation of these heavier elements increases the stability of the resulting cationic π -systems. Among the derivatives, Ge-bridged **2_{PF6}** showed the largest molar absorption coefficient in its UV-visible absorption spectrum. The use of

heavy group 14 elements as bridging moieties also gave rise to a central six-membered ring with a distorted geometry. This geometry, which deviates from planarity, may determine how closely the π -stacked structures align and thus govern the magnitude of the hypsochromic shift of their absorption bands in the aggregated state. Despite the incorporation of heavier elements, aggregates of these cationic π -systems are capable of emitting fluorescence, albeit their emissions are generally faint.

Conflicts of interest

There are no conflicts to declare.

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