Near-Infrared Fluorescence of Silicon Phthalocyanine Carboxylate Esters

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Seven silicon(IV) phthalocyanine carboxylate esters (SiPcs, 1–7) with non-, partially- and per-fluorinated aliphatic (linear or branched at the alpha-carbon) and aromatic ester groups have been synthesized, their solid-state structures determined and their optoelectronic properties characterized. The SiPcs exhibit quasi-reversible oxidation waves (vs. Fc+/Fc) at 0.58–0.75 V and reduction waves at −0.97 to −1.16 V centered on the phthalocyanine ring with a narrow redox gap range of 1.70–1.75 V. Strong absorbance in the near-infrared (NIR) region is observed for 1–7 with the lowest-energy absorption maximum (Q band) varying little as a function of ester between 682 and 691 nm. SiPcs 1–7 fluorescence in the near-infrared with emission maxima at 691–700 nm. The photoluminescence quantum yields range from 40 to 52%. As a function of esterification, the SiPcs 1–7 exhibit moderate-to-good solubility in chlorinated solvents, such as 1,2-dichlorobenzene and chloroform.

Phthalocyanines (Pcs) are one of the most extensively studied classes of organic functional macrocycles, having four nitrogen-bridged isoindole units. Pcs are known for their robust thermal and chemical properties, and their diverse optoelectronic and magnetic properties, especially their strong absorption in the near Infrared (λabs ~ 700 nm). These attractive optoelectronic properties have catalysed the use of Pcs as dyes and pigments, and as components in electrochromic and optical limiting devices, organic thin-film transistors, organic light emitting diodes (OLEDs), multistage-redox-dependent fluorophores, organic (OSC) and dye-sensitized solar cells (DSSCs). Many of the solid-state properties of Pcs are highly dependent on the extent of intermolecular π-π stacking interactions, especially in aggregates in polar protic solvents such as water or ethanol. Pcs are normally characterised by an intense absorption of the Q-bands in the near infrared (NIR) region of ~700 nm. Dimerization or aggregation between neighbouring Pcs molecules generally leads to fluorescence quenching and a broadened, red-shifted principal Q-band due to electronic/exciton coupling interactions that can normally be observed even in very dilute solutions. Aggregation is an unfavorable property of Pcs since it can not only cause difficulties in purification and characterization but also result in the quenching of luminescence due to enhanced non-radiative excited state decay, and therefore lowers the photoluminescence quantum yields, ΦPL. In the context of achieving non-aggregated Pc derivatives, several approaches have been adopted. These include introduction of bulky peripheral substituents such as tert-butyl groups and rigid spirocyclic fused rings, and the introduction of metal ions such as Sn(IV) and Ln(III) to break the planarity of the Pc-containing compounds such that the metal ion sits outside the plane of the Pc macrocycle. Another viable approach is to introduce elements such as Ru(II), Si(IV) or Ge(IV), that possess more than four valencies. Many Pc derivatives have been explored including main group metals and transition metals such as Al(III), Cu(II), Ni(II), Co(II), Zn(II), Ga (III), In(III), Sn(IV), Ge(IV), Ln(III), Pt(II) exhibiting square-planar, square-pyramidal, or octahedral coordination geometries. Pcs containing Si(IV) are an attractive subclass of Pcs because of the elemental abundance and very low toxicity levels of silicon coupled with their low band gap (~1.7 eV). The SiPcs with axial oxygen-terminated ligands (alkoxide, siloxide, aryloxide, and carboxylate) have been known since the 1960s and they have been applied as photosensitizing agents in photodynamic therapy. Recently, SiPcs with siloxide...
and aryloxide ligands have found use as chromophores and fluorophores in DSSCs and OLEDs\(^{31}\), the latter with external quantum efficiencies of up to 4.5%\(^{13}\). In contrast, to the best of our knowledge, SiPcs with carboxylate ligands have rarely been used in organic electronics. Considering the strong nature of Si–O bond, the wide range of commercially-available carboxylic acids and their capacity to confer increased solubility, silicon phthalocyanine carboxylates are an attractive target for the design of solution-processable electron-accepting materials for organic electronics.

In a recent study, we investigated the optoelectronic properties of two SiPcs bearing axial alkyl/aryl-carboxylate ligands (R1 and R2, Fig. 1) and explored their use in OLED and OSC devices\(^{32}\). Herein, we report a systematic study of the synthesis and optoelectronic characterization of a series of axially substituted SiPc carboxylates, 1–7, bearing perfluorinated (1 and 2) and partially fluorinated (4) carboxylates, alkyl carboxylates (3 and 5) and aryl carboxylates (6 and 7) (Fig. 1). The new SiPcs were comprehensively characterized by \(^1\)H and \(^{19}\)F NMR spectroscopy, NSI\(^{+}\) high-resolution or MALDI mass spectrometry, melting point determination, elemental analysis and single crystal XRD. The effect of different axial substituents on the optoelectronic properties of these complexes is discussed, supported by Density Functional Theory (DFT) calculations, with the results compared to our previously reported reference compounds R1 and R2.

SiPcs 1–7 were synthesized by nucleophilic substitution reactions of commercially available silicon phthalocyanine dichloride, SiPcCl\(_2\), with an excess of the corresponding carboxylic acids, either in DMF (SiPc 3) or in diglyme (SiPcs 1, 2, 4–7) at 140–165 °C (Fig. 1) in poor to good yields (3–64%) based on the carboxylic acids. The SiPcs were purified by column chromatography on silica. The product typically elutes as the first blue band from the column and it is usually well separated from the other fractions. Compounds 1–7 were obtained as air and moisture stable blue solids. It was found that the choice of solvent plays a crucial role in the synthesis of the SiPcs. For instance, 1 and 6 can be prepared from diglyme, but not from DMF. The aliphatic carboxylates were either linear at the alpha-carbon (1–2) or branched (3–5). The aromatic carboxylates possessed solubilizing groups (6 and 7) that were either electron-withdrawing (trifluoromethyl in 6) or electron-donating (octyloxy in 7).

All the compound characterization data are consistent with the proposed SiPc structures and further confirmed by single crystal data. Compounds 2–7 show melting points >300 °C while 1, possessing a long-chain aliphatic carboxylate \(\text{C}_{19}\text{H}_{39}\text{CO}_2\) ligands shows a significantly lower melting point of 168–170 °C. Fluorinated SiPcs 1, 4, and 6 exhibit low solubility of <5 mg mL\(^{-1}\) in chlorinated solvents, such as 1,2-dichlorobenzene and chloroform while the other SiPcs are soluble in those solvents (>10–30 mg mL\(^{-1}\)) but are insoluble in acetonitrile and alkanes. The solubility of SiPcs 2, 3, 5 and 7 in 1,2-dichlorobenzene are comparable to the same reported for reference complexes R1 (>10 mg mL\(^{-1}\)) and R2 (>35 mg mL\(^{-1}\)) in 1,2-dichlorobenzene\(^{35}\).

The axially-substituted SiPcs have characteristic \(^1\)H NMR spectra (Fig. 2). The \(^1\)H resonances of the axial substituents close to the phthalocyanine ring are strongly shifted upfield as they experience a large diamagnetic ring-current shielding effect induced by the macrocyclic ring. This effect is more pronounced in 3–5 where some of the aliphatic protons appear ≤1 ppm. The phthalocyanine ring \(^1\)H resonances appear as two downfield-shifted unresolved multiplets integrating each as 8 H at around 9.8 and 8.5 ppm, for the α and β protons, respectively\(^{36}\). The two axially substituted carboxylate ligands are symmetric on the NMR timescale and exhibit a single set of NMR resonances. This observation in conjunction with the presence of the two sets of phthalocyanine ring \(^1\)H resonances suggest a local \(D_{\infty}\) symmetry in 1–7 in solution state, irrespective of ester ligand. Electrospray ionization mass spectrometry showed the expected isotope distribution pattern for both the [M + H]\(^{+}\) and [M + Na]\(^{+}\) ions for 1–7.

![Figure 1. Synthesis of SiPc complexes 1–7 and benchmark complexes R1 and R2](image-url)
Crystal structures

The crystal structures of 1–7 were determined. Details of structure refinements and some crystallographic parameters are collected in Supplementary Table S1 in the ESI. Selected bond distances of SiPcs 1–7, R1 and R2 are gathered in Supplementary Table S2. The structure of 6 shows two independent molecules in the asymmetric unit, as well as a co-crystallized molecule of acetonitrile. The flexible polyfluorinated chains of the carboxylate ligand in 2 result in a lower quality, but unambiguous, determination of its structure. In all seven structures, the hypervalent silicon(IV) adopts the expected distorted octahedral SiN₄O₂ coordination environment (Fig. 3). Except in the case of 6, the SiPc has a crystallographic inversion center at the silicon atom; therefore, the SiN₄ fragment is planar, irrespective of any distortions within the phthalocyanine ring itself, the SiO₂ fragment is linear, and only two of the Si–N and one of the Si–O bonds are unique. In 6, the same planar SiN₄ fragment (root-mean-squared deviation of the SiN₄ fragment from planarity, 0.0019 and 0.0003 Å, for Si1 and Si51, respectively), and linear SiO₂ fragment [O–Si–O angles, 179.59(9) and 179.40(8)°, for Si1 and Si51, respectively] are seen, despite the lack of a center of symmetry at Si. The Si–N bonds are significantly longer [1.8897(18)–1.9203(11) Å] than the Si–O bonds [1.7358(9)–1.7793(15) Å]. The Si–N bond lengths decrease while the Si–O bond lengths increase when the carboxylate ester becomes more electron-attracting from a non-fluorinated to a (per-)fluorinated one (Supplementary Table S2). All Si–N and Si–O bond distances fell within the range of distances for six-coordinate SiPc compounds with axial O-donor ligands, found in a search of the Cambridge Structural Database (CSD version 5.37). The Si–N distances were seen to lie below the middle of the range of distances (1.89–1.95 Å), whereas the Si–O distances were at the upper end of the range (1.65–1.78 Å).

The complexes show a limited selection of both π···π and C–H···π interactions; however, these do not involve the aryl groups of the carboxylate ligands in 6 and 7, which is likely due to the surrounding sterical bulk. In contrast, the phenyl rings of the phthalocyanines do participate in π–stacking, with centroid–centroid distances ranging from 3.5361(11) to 3.7795(19) Å. The exceptions to this occur in 1 and 5, which show no centroid–centroid distances less than 3.8 Å, although there are phenyl rings positioned appropriately at distances slightly greater than this that show patterns of interactions in line with the other structures. This π···π stacking occurs through either the two trans-benzenes to give an infinite single chain (3 and 4, Fig. 4) or through the two pairs of cis-benzenes to give an infinite double-chain (1, 2, 5, 6, and 7, Fig. 5). As well as these π···π interactions, compounds 5 and 7 also show weak C–H···π interactions between aliphatic C–H groups in the axial ligands and aromatic rings in either the phthalocyanine, or the axial ligand, at distances of 3.00 and 2.71 Å, respectively. The interaction in 5 is at the conventional van der Waals limit, but C–H···π interactions have been suggested to be effective at distances beyond this value.

All the complexes, apart from 5, exhibit weak intermolecular C–H···O hydrogen bonds to the carbonyl oxygen of the carboxylate ligand, which acts as acceptor for two C–H···O interactions simultaneously. These predominantly involve aromatic phthalocyanine C–H groups, but also aliphatic C–H groups in some cases. The H···O separations vary between 2.37 and 2.55 Å, with corresponding C···O separations of 3.265(3) to 3.445(13) Å. For 5, the steric bulk of the adamantyl group may restrict some close-packing interactions with the adjacent carboxylates. The SiPcs with fluorinated carboxylate ligands (1, 2, 4, and 6) additionally exhibit weak intermolecular C–H···F interactions. These show a similar distance spread to the C–H···O hydrogen bonds, with H···F separations varying between 2.36 and 2.62 Å, and C···F separations between 3.169(4) to 3.445(3) Å. Such interactions would generally
be considered too weak to make a significant contribution to the packing, except that they are seen to occur in cooperative pairs; working in conjunction with other C–H···F interactions. The exception to this pattern is in 4, where no other interactions are seen to be operating cooperatively with the C–H···F interactions, suggesting that in this case the close H···F contact may just be an artifact of packing.

The presence of the axial ligands can lead to disruption of intermolecular interaction in the solid state. The linear alkyl chains of the carboxylates in 1 and 2 are arranged parallel to the plane of the phthalocyanine ring (Fig. 6), whereas all the other carboxylate ligands are arranged nearly orthogonal to it, even in the case of 7, containing an aryl octyl ether (Fig. 3). This arrangement of axial ligands is mirrored in other SiPcs, where the presence of aryl carboxylate ligands show positioning orthogonal to the SiPc32,36,37, while alkyl carboxylates show a parallel position26,32,37–40. Potentially arising from these differing chain orientations, and the resulting presence or absence of steric bulk across the face of the SiPc, the phthalocyanine rings pack with the ring-planes arranged in a parallel fashion in 1, 2 and 5, whereas in the rest of the SiPcs they pack in a herringbone pattern, with SiPc planes approximately orthogonal. This pattern of arrangements is not mirrored as closely in other known structures; although those with aryl carboxylates do show packing with orthogonal SiPcs37, those with alkyl carboxylates show a mixture of packing motifs, most showing a parallel arrangement of SiPcs37,39, but some showing herringbone packing37,40, or other motifs with orthogonally arranged SiPcs36,39. The intermolecular separation was estimated using the shortest Si···Si distance in each structure. Surprisingly, the longest separation is found to be in 4 (Table 1), which has the highest solubility in the series, and the shortest in 7.

**Electrochemistry.** The cyclic voltammograms of compounds 1 and 3–7 were obtained in dichloromethane at room temperature and are shown in Fig. 7 and the electrochemical data collated in Table 1. Due to poor yield, the electrochemical measurement of 2 was not possible. At positive potential, the SiPcs exhibit mono-electronic quasi-reversible oxidations in the range 0.58–0.75 V vs. Fc/Fc+. DFT calculations indicate that the highest occupied molecular orbitals (HOMOs) are distributed on the isoindole moieties of the macrocycles (Fig. 8). Based on the DFT calculations, and in line with the literature data, the first oxidation processes

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Figure 3. X-ray structures of 1–7, with ellipsoids drawn at the 50% probability level. Hydrogen atoms, solvent molecules, minor disorder components, and additional independent molecules are omitted for clarity.
of 1 and 3–7 are assigned to the removal of one electron from the indole moiety. The lower energies calculated for the HOMOs of 1 ($E_{\text{HOMO}} = -5.40 \text{ eV}$), 3 ($E_{\text{HOMO}} = -5.27 \text{ eV}$) and 4 ($E_{\text{HOMO}} = -5.30 \text{ eV}$) compared to that of R1 ($E_{\text{HOMO}} = -5.12 \text{ eV}$) are in good agreement with their more positive anodic potentials (Table 1).
The slightly shallower HOMO of 5 (E_{HOMO} = −5.09 eV) bearing the adamantyl moiety compared to that of R1 (E_{HOMO} = −5.12 eV) is in line with the 10 mV cathodic shift of its oxidation potential compared to that of R1. The energy calculated for the HOMO (E_{HOMO} = −5.35 eV) of bis(trifluoromethyl)phenyl carboxylate-substituted compound 6 is significantly lower than that of R2 (E_{HOMO} = −5.07 eV), aligning with the 100 mV anodic shift of the oxidation potential of 6 compared to that of R2. However, the trend observed in the lower HOMO energy
calculated for \( E_{\text{HOMO}} = -5.25 \text{ eV} \) compared to that of \( R_2 \) \( E_{\text{HOMO}} = -5.07 \text{ eV} \) is not in line with the observed 200 mV cathodic shift in its oxidation potential compared to that of \( R_2 \). So, in general, when electron-withdrawing fluorinated groups are added to the carboxylate ligand, the redox potentials are shifted positively, the oxidation becomes harder; e.g., compare (1 and 4) vs. (3 and 5) or 6 vs. 7 (Table 1). A similar anodic shift of 30 mV is also observed on substitution of one methyl group in the \( t\text{Bu} \) group in 3 with a -CF\(_3\) group in 4. Compared to \( R_1 \), all the fluorinated SiPcs, 1, 4, and 6 exhibit anodically shifted oxidation potentials as a function of incorporation of electron-withdrawing -CF\(_3\) groups (Table 1). The similar argument is also valid for the observed 100 mV anodic shift in the oxidation potential of \( \text{SiPC}_6 \) compared to that of \( R_2 \) (Table 1). The most positive oxidation potential in the series is observed for 1, which contains the perfluoroheptanoate axial ligands. Upon scanning to negative potentials, the SiPcs exhibit one or two quasi-reversible monoelectronic reductions, the first at \(-0.97 \) to \(-1.16 \text{ V} \) and the second at \(-1.47 \) to \(-1.63 \text{ V} \), which are typical of Pcs\(^{41}\). For all the SiPcs, DFT calculations point to a lowest unoccupied molecular orbital (LUMO) that has isoindole character coupled with the contribution from the bridging N-atoms. Therefore, and in line with the literature data, the first reduction can be assigned to reduction of the macrocycle itself. Similar to the trend observed in the oxidation of (1 and 4) vs. (3 and 5), the presence of the electron-withdrawing fluorinated groups anodically shifts the first reduction potential of (1 and 4) by 60–190 mV compared to those of (3 and 5) (Table 2). This trend is in good agreement with the more stabilized computed LUMOs of 1 and 4 (\( E_{\text{LUMO}} = -3.33 \text{ eV} \) for 1 and \(-3.21 \text{ eV} \) for 4) compared to those of 3 and 5 (\( E_{\text{LUMO}} = -3.17 \text{ eV} \) for 3 and \(-2.97 \text{ eV} \) for 5). The computed and experimentally determined LUMO energies of the aryl-substituted SiPcs 6 and 7 containing either electron-withdrawing -CF\(_3\) or electron-donating octyloxy groups, respectively, also follow the expected trend. The first reduction potential of 6 is 80 mV anodically shifted compared to that of 7; the LUMO of 6 (\( E_{\text{LUMO}} = -3.27 \text{ eV} \) for 6) is likewise more stabilised compared to the LUMO of 7 (\( E_{\text{LUMO}} = -3.15 \text{ eV} \) for 7) (Table 2). DFT calculations demonstrate that the LUMO + 1 is centred on the isoindole moiety in all the compounds. The observed quasi-reversible reductions

![Figure 8. Calculated frontier MO energies of 1–7, obtained from DFT [(B3LYP/SBKJC-VDZ for Si(IV)) and (6–31 G\(^*\) for C,H,N,O,(F)] with CPCM(MeCN) and 0.5 eV threshold of degeneracy (orbitals are isocontoured at 0.03). Kohn-Sham MOs of 1–7 are also shown.](image_url)

| SiPc | \( \lambda_{\text{abs}}/\text{nm} (\epsilon/10^4 \text{M}^{-1} \text{cm}^{-1}) \) |
|------|--------------------------------------------------|
| 1    | 295 (2.8), 335 (6.9, sh), 356 (7.8), 621 (3.8), 659 (3.3), 691 (29) |
| 2    | 295 (2.8), 335 (6.8, sh), 355 (7.6), 621 (3.7), 659 (3.2), 690 (27) |
| 3    | 290 (2.6), 358 (7.2), 613 (3.9), 651 (3.3), 682 (27) |
| 4    | 293 (2.3), 358 (7.4), 616 (3.7), 654 (3.1), 685 (26) |
| 5    | 290 (2.4), 335 (5.8, sh), 350 (6.2, sh), 360 (6.5), 614 (3.7), 652 (3.1), 683 (26) |
| 6    | 295 (2.4), 358 (7.8), 618 (3.7), 657 (3.2), 688 (27) |
| 7    | 257 (6.1), 291 (2.3), 359 (7.2), 615 (3.8), 653 (3.2), 684 (27) |

Table 2. UV-vis absorption data in dichloromethane at room temperature.
of 3–7 in the range of −1.47 to −1.63 V are assigned to the reduction of the macrocyclic ring system (Fig. 7). Except for 1, the first reduction potentials of 3–7 are cathodically shifted by 40–170 mV compared to axially alkoxy-disubstituted SiPcs ($E_{\text{red}} = −900$ mV) 43.

The redox gap, $\Delta E_{\text{redox}}$, spans only 60 mV across the series of SiPcs and remains nearly constant at 1.69–1.75 V since axial substitution modulates these potentials in the same direction (Table 1). A similar behavior of the redox potentials and the near-constancy of the redox gap have been noted previously for other series of Pc complexes 26.

**Photophysics.** The UV-vis absorption spectra of all the SiPcs were recorded in dichloromethane and values are collated in Table 2. An overlay of the experimental UV-vis absorption spectra of 1 and 7 with their predicted transitions calculated by time-dependent DFT (TD-DFT) at room temperature in dichloromethane is shown in Fig. 9 and similar overlays of the experimental and TD-DFT predicted UV-vis profiles for SiPcs 1–7 are shown in Supplementary Figure S1. The absorption spectra exhibit profiles that are typical of the phthalocyanine chromophore; a B (or Soret) band at 355–360 nm, an intense and sharp near-infrared Q band at 682–691 nm, together with two vibronic bands at 613–621 nm and 651–659 nm. The TD-DFT-assigned characterisations of these bands are tabulated in Table 3. The Q-bands have molar extinction coefficients, $e$, of (26–29) × 10^4 M$^{-1}$ cm$^{-1}$ (Table 3). For all the SiPcs the absorptivity of the Q band strictly follows the Beer-Lambert law as a function of concentration, thereby indicating that the compounds are essentially free from aggregation in dichloromethane 43. Furthermore, the absence of either a red-shifting of the Q band at ca. 750 nm and a blue-shifting Q band at ca. 630 nm, excludes the formation of J and H aggregates, respectively. The retention of monomeric behaviour even at 10$^{-4}$ M demonstrates that the axial ligands are very effective in sterically isolating the chromophoric Pc rings, by preventing intramolecular $\pi$ stacking interactions between the macrocycles in solution 38. The absorption spectra have a window between 420–560 nm, which is typical of phthalocyanines. The absorption cut-off (optical band gap) of the SiPcs is at 720–730 nm (1.72–1.70 eV). The TD-DFT calculations of 1–7 predict that the bands found between 257–350 nm are principally $\pi \rightarrow \pi^*$ locally excited (LE) transitions within the conjugated macrocycle, whereas the $\pi \rightarrow \pi^*$ LE bands of the aromatic carboxylates in 6 and 7 are superimposed with those of the phthalocyanine. The B bands at ca. 355 nm for 1 and 2 are due to the $\pi \rightarrow \pi^*$ based transitions within the isoindole moieties of the macrocycle, whereas, these same bands in 3–7 also contain small contributions from n $\rightarrow \pi^*$ transitions from the bridging N-atoms connecting the four isoindole moieties to the $\pi^*$ orbital of the isoindole (Table 3). For 6, this band contains an additional $\pi \rightarrow \pi^*$ based ligand-to-ligand charge transfer (LLCT) transition emanating from the 3,5-bis(trifluoromethyl)phenylcarboxylate to the macrocycle. The Q-band for all the SiPcs is predicted to originate from $\pi \rightarrow \pi^*$ LE transitions among the isoindole moieties of the macrocycle. Fluorination of the carboxylate ligand from L3 to L4 leads to a modest red-shift of 3 nm in the Q band of 4 compared to that of 3; similar behavior exists between 6 and 7. The position of the Q band, which relates to the HOMO-LUMO gap, remains nearly constant at 682–691 nm across the series. Therefore, the HOMO-LUMO energy gap is invariant to the original of the axial substitution of the carboxylates, an observation corroborated by their electrochemistry. However, the lowest energy absorption maxima of these complexes are red-shifted by ca. 10–20 nm compared to those of alkoxy- and siloxy-substituted SiPcs 43–45.

The SiPcs 1–7 were also characterized by photoluminescence spectroscopy in dilute and degassed dichloromethane solutions at ambient temperature. Relevant photophysical data are shown in Table 4 and the photoluminescence spectra of SiPcs 1 and 7 are shown in Fig. 10, whereas the photoluminescence spectra of all the SiPcs in dichloromethane solutions are shown in Supplementary Figure S2. All the SiPcs emit strongly in the near-infrared (NIR) region with $\lambda_{\text{em}}$ ranging narrowly from 688–700 nm. Two lower intensity bands are observed at between 710–790 nm. The photoluminescence spectrum of each of the SiPcs is a mirror image of the absorption spectrum. The $\lambda_{\text{em}}$ values are typical of carboxy-substituted SiPc compounds and the emission is modestly red-shifted by ca. 8–30 nm compared to alkoxy- and siloxy-disubstituted SiPcs. All the SiPcs exhibit short mono-exponential photoluminescence decay kinetics ($\tau$) in the range 6.7–7.3 ns and high photoluminescence quantum yields ($\Phi_{\text{PL}}$ = 40–52%) in dichloromethane solution. The calculated radiative lifetime, $\tau_{\text{rad}} = \tau \times \Phi_{\text{PL}}^{-1}$, ranges from 13.8–18 ns, the radiative, $k_r$, and non-radiative, $k_{nr}$, decay constants, $k_{nr} = k_r (1 - \Phi_{\text{PL}})$, are in the range of (55.5–72.2) × 10$^4$ and (66.7–83.3) × 10$^4$ s$^{-1}$ for all the compounds. The mirror image behavior between absorption and emission spectra, small Stokes shift ($\lambda_{\text{em}} - \lambda_{\text{abs}}$, of 6–10 nm), short excited-state
The higher fluorescence intensity was observed compared to those with electron-withdrawing groups, Table 4. The higher PLQY of SiPc seems to be associated with a higher rate of radiative decay constant ($k_r$) of 81.8 × 10^6 s⁻¹ compared to the $k_r$ values of the other SiPcs (66.1–80.0 × 10^6 s⁻¹).

### Table 3. Principal transitions by TD-DFT involved in the B and Q bands of SiPcs.

| Cmpd | $\lambda_{ex}$/nm (c/10⁸ M⁻¹ cm⁻¹) | $\lambda_{em}$/nm (f) | Major transition | Character |
|------|----------------------------------|----------------------|------------------|-----------|
| 1    | 358 (7.8)                        | 358 (0.8221)         | H −→ L (88%)     | isoindole(π) to isoindole(π*) |
|      | 691 (29)                         | 679 (0.7111)         | H −→ L (98%)     | isoindole(π) to isoindole(π*) |
| 2    | 355 (7.6)                        | 358 (0.8629)         | H −→ L (80%)     | isoindole(π) to isoindole(π*) |
|      | 690 (27)                         | 680 (0.701)          | H −→ L (98%)     | isoindole(π) to isoindole(π*) |
| 3    | 358 (7.2)                        | 353 (1.1026)         | H −→ L + 1 (90%) | isoindole(π) to isoindole(π*) (major) + bridged-N (l.p.) to isoindole(π*) (minor) |
|      | 682 (27)                         | 672 (0.7208)         | H −→ L (98%)     | isoindole(π) to isoindole(π*) |
| 4    | 358 (7.4)                        | 357 (1.1138)         | H −→ L (83%)     | isoindole(π) to isoindole(π*) (major) + bridged-N (l.p.) to isoindole(π*) (minor) |
|      | 685 (26)                         | 674 (0.7187)         | H −→ L (98%)     | isoindole(π) to isoindole(π*) |
| 5    | 360 (6.5)                        | 354 (1.0757)         | H −→ L + 1 (89%) | isoindole(π) to isoindole(π*) (major) + bridged-N (l.p.) to isoindole(π*) (minor) |
|      | 683 (26)                         | 672 (0.7072)         | H −→ L (98%)     | isoindole(π) to isoindole(π*) |
| 6    | 358 (7.8)                        | 358 (1.1374)         | H −→ L (71%)     | isoindole(π) to isoindole(π*) (major) + bridged-N (l.p.) to isoindole(π*) (minor) |
|      | 688 (27)                         | 677 (0.7206)         | H −→ L (98%)     | isoindole(π) to isoindole(π*) |
| 7    | 359 (7.2)                        | 353 (1.1494)         | H −→ L + 1 (89%) | isoindole(π) to isoindole(π*) (major) + bridged-N (l.p.) to isoindole(π*) (minor) |
|      | 684 (27)                         | 673 (0.7249)         | H −→ L (98%)     | isoindole(π) to isoindole(π*) |
| R1   | 360 (7.1)                        | 356 (0.9739)         | H −→ L (85%)     | isoindole(π) to isoindole(π*) (major) + bridged-N (l.p.) to isoindole(π*) (minor) |
|      | 683 (26)                         | 672 (0.7350)         | H −→ L (98%)     | isoindole(π) to isoindole(π*) |
| R2   | 359 (7.1)                        | 353 (1.1235)         | H −→ L + 1 (89%) | isoindole(π) (major) + isoindole(π) to isoindole(π*) (minor) |
|      | 685 (27)                         | 672 (0.7090)         | H −→ L (98%)     | isoindole(π) to isoindole(π*) |

### Table 4. Photophysical data of complexes 1–7.

| SiPcs | $\lambda_{ex}$/nm | $\Phi_{PL}$/% | $\tau$/ns | $\tau_{rad}$/ns | $k_r / \times 10^6$ s⁻¹ | $k_{nr} / \times 10^6$ s⁻¹ |
|-------|------------------|---------------|-----------|----------------|--------------------------|--------------------------|
| 1     | 700              | 42            | 7.3       | 17.4          | 57.5                     | 79.4                     |
| 2     | 697              | 48            | 6.7       | 13.9          | 71.6                     | 77.6                     |
| 3     | 688              | 52            | 7.2       | 13.8          | 72.2                     | 66.7                     |
| 4     | 692              | 46            | 6.9       | 15.0          | 66.7                     | 78.3                     |
| 5     | 692              | 41            | 7.3       | 17.8          | 56.2                     | 80.8                     |
| 6     | 698              | 40            | 7.2       | 18.0          | 55.5                     | 83.3                     |
| 7     | 693              | 43            | 7.2       | 16.7          | 59.7                     | 79.2                     |

### Table 4. Photophysical data of complexes 1–7.

- aDetermined using relative method using R2 as the reference.
- bCalculated from $\tau_{rad} = \tau \times \Phi_{PL}$. cCalculated from $k_{rad} = (k_{rad}/\Phi_{PL}) - k_{nr}$.
Conclusions

In summary, a series of SiPcs bearing axially substituted carboxylate ligands have been synthesized and structurally and optoelectronically characterized. Their structures were unambiguously characterized by 1H NMR spectroscopy, ESI-MS, elemental analyses and single crystal XRD analyses. The presence of electron-withdrawing or donating groups on the carboxylate ligands does not exert any significant influence on the optoelectronic properties of the SiPcs as the electron-donating/withdrawing groups exert similar effects upon the phthalocyanine ring based HOMO and LUMO. The oxidation and the reduction processes of the complexes were both found to be quasi-reversible. All the SiPcs exhibit NIR fluorescence with high photoluminescence quantum yields and no sign of aggregation in dilute solution. This study demonstrates the wide selection of carboxylate ligands available to render the SiPc compounds solution-processable and thus amenable for use in OLED and OPV devices.

Experimental Section

General Synthetic Procedure. Commercial chemicals were used as supplied. Chromatography was performed on columns with an i.d. of 25–30 mm on silica gel. The reaction was performed under nitrogen using standard Schlenk techniques. Purification and handling of the products were carried out under air. The progress of reactions and the elution of products were followed on TLC plates (silica gel on aluminum sheets, 250 µm with indicator F-254). Compounds were visualized under UV light. 1H and 19F NMR spectra were recorded with Bruker AVANCE 300 (300 MHz for 1H; 282 MHz for 19F) and Bruker AVANCE II 400 spectrometers (400 MHz for 1H; 377 MHz for 19F). The following abbreviations have been used for multiplicity assignments: “s” for singlet, “d” for doublet, “t” for triplet, “m” for multiplet, and “br” for broad. 1H NMR spectra were referenced to the solvent peak. Melting points (m.p.) were recorded in open-end capillaries on an Electrothermal melting point apparatus IA9200 and are uncorrected. The heating rate was 0.5 or 1.0 °C/min. High-resolution nanospray ionization (HR NSI) and MALDI+ mass spectra, were recorded at the EPSRC National Mass Spectrometry Service Centre, Swansea University. Elemental analyses were performed by Mr. Stephen Boyer, London Metropolitan University.

Silicon phthalocyanine dichloride (SiPcCl₂; dye content ~85%) and carboxylic acid (excess) were stirred in 3–4 mL of dry and degassed DMF or diglyme (degassed by bubbling with nitrogen for 5 min) at 140–165 °C overnight to give dark green-blue solution or suspension. The volume of the solvent was chosen to be as small as possible, while still allowing the efficient mixing of the reagents. The temperature of the reaction was as high as possible to help dissolve the starting materials; however, the temperature was lowered for the fluorinated acids to prevent side-reactions involving the electron-deficient C–F bonds. The molar ratio of SiPcCl₂ to carboxylic acid

Figure 10. Typical normalized emission spectra of complexes 1 and 7 in degassed dichloromethane solutions at room temperature (λexc = 360 nm).

Table 5. Photophysical data of complexes 1–7 in PMMA doped film. In PMMA at room temperature (λexc = 612 nm). Determined using an integrating sphere. Calculated from τrad = τ × Φ⁻¹. Calculated from kτ = ΦPL/τ. Calculated from knr = (kτ/ΦPL) − kτ.
ranged from 3.9–7.4. The reaction mixture was cooled to room temperature. Water (50 mL or more) was added to precipitate the product (blue solid). For 2, addition of water precipitated a dark oil; the supernatant solution was decanted and the oil was crystallized on mixing with methanol/water (1/1, v/v). The solid product was filtered. The product was washed on the filter with water for 6; with water and hexane (5–10 mL) for 3; or with water and methanol/water (1/1, v/v) for 1, 2, 4, 5 and 7. The washing was performed to remove the reaction solvents and the excess of the acid. The product was puriﬁed by column chromatography on 10–22 g of silica. The elution was performed with dichloromethane (except for 4) or with 0–0.3% methanol in dichloromethane (for 4) to give green-blue eluate of the product as the ﬁrst band. The combined product fractions were evaporated to dryness. The product was dried under vacuum.

The products are air- and moisture stable blue solids that are soluble in chlorinated solvents (chloroform, dichloromethane, 1,2-dichlorobenzene) and are insoluble in acetonitrile. The exact shade, brightness, and depth of the blue color of the product mainly depend on whether the product is crystalline or not. The crystals of the product may look red in reflection. The products were stored in the dark. Further details are provided below.

Photophysical measurements. The solutions for photophysical investigations were prepared in HPLC-grade dichloromethane. Electronic absorption spectra were recorded with a Shimadzu UV-1800, UV-Vis Spectrophotometer in optical quartz cells of 10 mm path length. The absorption spectra were recorded on a PerkinElmer LAMBDA 950 UV/Vis/NIR spectrophotometer. The photoluminescence spectra were recorded on an Edinburgh Instruments FLS-980 ﬂuorescence spectrometer and were corrected for the spectral sensitivity of the detection system. In dilute solutions used in the photophysical measurements, the photoluminescence spectra were independent of the excitation wavelength. In our earlier work, the absolute photoluminescence quantum yields (PLQY) of SiPc carboxylates R1 and R2 were determined using an integrating sphere on a Hamamatsu C9920–02 absolute PLQY measurement system12. Photoluminescence quantum yields for the SiPcs in this work were determined by a relative method using R2 as the standard. The photoluminescence lifetime was measured using the time-correlated single-photon-counting technique integrated in the FLS-980 ﬂuorescence spectrometer. The samples were excited at 375 nm by pico-second pulsed diode laser head Picoquant LDH-D-C-375. The
luminescence decay was measured at a wavelength corresponding to the emission peak. The emitted light was dispersed in a monochromator with an entrance slit of 2–4 nm and was detected on a microchannel plate detector (<100 ps response). The luminescence decay was fitted to a mono-exponential function with a $\chi^2$ value in the range of 1–1.25.

**Electrochemistry.** Electrochemical experiments were conducted in HPLC grade dichloromethane with 0.1 M tetra-n-butylammonium hexafluorophosphate as the supporting electrolyte, with a PC controlled electrochemical workstation (Electrochemical Analyzer/Workstation CH1600C, CH Instruments, Inc.) at room temperature.

The experiments were carried out under nitrogen in an electrochemical cell through which a stream of nitrogen was passed. The volume of the solvent was 2–3 mL. Glassy-carbon disk (3 mm diameter), platinum spiral, and platinum wire served as working, counter, and quasi-reference electrodes, respectively. Estimated error: ±30 mV. Cyclic voltammetry was performed at scan rates of 0.1 V/s and 1 V/s. The potentials of reversible redox processes were independent of the scan rate.

**DFT calculations.** All calculations were performed with the Gaussian09, revision D.01 suite of programs. All calculations were performed with the Gaussian09, revision D.01 suite of programs, the Becke three-parameter hybrid functional[50], and Lee-Yang-Parr’s gradient-corrected correlation functional (B3LYP)[51]. Singlet ground state geometry optimizations and single point energy calculations for 1–7, R1 and R2 were carried out at the (R)B3LYP levels, using their respective crystallographic structures as starting points. All elements except Silicon were assigned the 6–31G(d,p) basis set[52]. The double-$\zeta$ quality SBKJC VDZ ECP basis set[53] with an effective core potential was employed for the Si(VI)-ion. Vertical electronic excitations based on (R)B3LYP optimized geometries were computed for 1–7, R1 and R2 using the TD-DFT formalism[54,55] in dichloromethane using conductor-like polarizable continuum model (CPCM)[56,57]. Vibrational frequency calculations were performed to ensure that the optimized geometries represent the local minima and there are only positive eigenvalues. The electronic distribution and localization of the singlet excited states were visualized using the electron density difference maps (ED-DMs)[58]. Gaussian 2.2 and Chemissian v3.8[59] were employed to visualize the absorption spectra (simulated with Gaussian distribution with a full-width at half maximum (fwhm) set to 3000 cm$^{-1}$) and to calculate the fractional contributions of various groups to each molecular orbital. All calculated structures and Kohn-Sham orbitals were visualized with ChemCraft[60].

**X-ray Crystallography.** Diffraction data for compounds 1 and 3–7 were collected by using a Rigaku FR-X Ultrahigh brilliance Microfocus RA generator/confocal optics and Rigaku XtaLAB P200 system, using Mo Kα radiation ($\lambda = 0.71073$ Å). Data for compound 2 were collected by using a Rigaku MM-007HF High brilliance RA generator/confocal optics and Rigaku XtaLAB P100 system, using Cu Kα radiation ($\lambda = 1.54187$ Å). Intensity data for all structures were collected at 173 K, using either $\omega$, or both $\omega$ and $\phi$ scans accumulating area detector images spanning at least a hemisphere of reciprocal space. All data were corrected for Lorentz polarization effects. A multiscan absorption correction was applied by using CrystalClear[51]. Structures were solved by direct (SIR2004[62]) or ω, or both ω and ϕ scans. Vizted geometries were computed for 1–7, R1 and R2 using the TD-DFT formalism in dichloromethane using conductor-like polarizable continuum model (CPCM)[56,57]. Vibrational frequency calculations were performed to ensure that the optimized geometries represent the local minima and there are only positive eigenvalues. The electronic distribution and localization of the singlet excited states were visualized using the electron density difference maps (ED-DMs)[58]. Gaussian 2.2 and Chemissian v3.8[59] were employed to visualize the absorption spectra (simulated with Gaussian distribution with a full-width at half maximum (fwhm) set to 3000 cm$^{-1}$) and to calculate the fractional contributions of various groups to each molecular orbital. All calculated structures and Kohn-Sham orbitals were visualized with ChemCraft[60].

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