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Ligand impacts on band edge energies and excited state splittings of silicane†

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Two-dimensional (2D) silicanes are promising semiconductors for applications in optoelectronics and photochemistry. Covalent functionalization presents a facile strategy to customize silicanes to attain desired properties. A comprehensive understanding of the impacts of ligands on silicanes is thus pivotal. In this study, we perform density functional theory (DFT) and time-dependent DFT (TDDFT) calculations to investigate the effects of three typical classes of ligands: (1) σ -withdrawing and π -donating, (2) σ -withdrawing and π -withdrawing, and (3) σ -donating and π -withdrawing, on the geometric structure, electronic structure, and band edge excited states of silicanes. Covalent functionalization of silicanes enables a wide range of band edge energies. The band gaps can be tuned between indirect and direct, and the underlying mechanism is explained for the first time. Additionally, TDDFT calculations confirm that the band edge optical absorptions can be adjusted by the ligands broadly from the near-infrared to the visible light region. These desirable properties enhance the functionalities of silicanes. Methodologically, we discuss the applicability of using the conventional one-particle orbital model to describe the excited states of silicanes, and nail down the ~ 0.1 eV inaccuracy of the model in describing excited state splittings. We present empirical functions for quick estimations of band edge energies of covalently functionalized silicanes. Through careful comparisons, we justify the use of the carbon-adapted σ/π -donating/accepting indices of typical ligands in silicon chemistry and present a set of silicon-based indices for future prudent applications in inorganic computational studies of silicon-based materials.

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1. Introduction

Over the past two decades, there has been a surge of research in two-dimensional (2D) materials, largely driven by the groundbreaking discovery of graphene in 2004¹ and the subsequent synthesis of silicene.^{2,3} Especially, silicene has found extensive applications in field-effect transistors^{4–6} and spintronics.^{7–9} Silicanes—a group of hexagonal sp^3 -silicon-based networks—can be viewed as functionalized silicenes and have been a focus of research due to their desirable optoelectronic properties, high biocompatibility, and synthetic readiness.^{10–12} These attributes suggest their wide applications in micro- and nano-electronics,¹³ optoelectronics,^{14,15} photocatalysis,^{16,17} and batteries.^{18–20}

Covalent surface functionalization (*i.e.*, replacing surface Si–H by Si–R bonds^{21,22}) is critical in silicon chemistry, particularly for

enhancing solution processability and chemical stability.^{23,24} It is also a handle to modify and control properties such as conductivity, hydrophilicity, lubricity, and biocompatibility.^{25,26} The properties of silicon-based materials change significantly when reduced to the 2D nanoscale.^{27,28} However, compared to other 2D nanostructures such as MXenes and ionic semiconductor nanoplates,^{29,30} ligand functionalization for silicanes is less studied; there have been only a few reports on R being alkenyl,¹⁵ aryl,^{22,31} and amino^{32,33} groups. Consequently, the understanding of the ligand influences on the electronic and optical properties of silicanes is far behind the advance of the functionalization techniques.

Computational chemistry offers atomistic understanding of these influences. Previous computational studies have demonstrated that the band gap is tuned by various ligands from 1.2 to 2.5 eV, from the intrinsic indirect band gap to direct gap,^{21,34,35} and the gap is dependent on ligands coverage,^{35–37} strain,³⁸ and external electric fields.³⁹ The mechanisms underlying these gap modulations, however, remain elusive. Furthermore, the band gap is not the only key property; the band edge energies of silicanes are equally important as they influence the separation, recombination, and migration of photogenerated charge carriers.⁴⁰ These important properties have not yet been studied. In terms of methodology, preceded computational

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studies of the excited states of silicenes have all relied on the conventional one-particle orbital description, *i.e.*, energy gaps between hole and particle orbitals are viewed as excitation energies. While this description is usually applicable for extended systems without local formations, given the spatial confinement of their ~ 4 Å thickness, the excitation energies of silicenes may not be solely determined by the orbital energy gaps. In this respect, silicenes resemble finite-sized molecules, and we may employ similar method and language developed for molecules with explicit electron correlation to investigate and interpret the excited states of silicenes. Surprisingly, no such study has been performed for such an important class of spatially confined materials.

Given these motivations, we performed density functional theory (DFT) and time-dependent DFT (TDDFT) calculations for pristine silicene (SiH) and substituted SiR species with three representative types of ligands: (1) σ -withdrawing and π -donating (sWpD: Cl, SH, OH, OCH₃, and NH₂), (2) σ -withdrawing and π -withdrawing (sWpW: CN), and (3) σ -donating and π -withdrawing (sDpW: BH₂, Li). More sWpD ligands were considered since those are typical oxidative ligands for silicenes. “ σ -withdrawing”, “ π -donating”, *etc.* means that a ligand pulls or pushes electrons from or to its bonding partner through σ - or π -type covalent interaction. As shown below, the results fill the knowledge gaps mentioned above.

II. Methods

All ground state DFT calculations were performed using the Vienna *ab initio* Simulation Package (VASP 5.4.4)⁴¹ and the projector augmented-wave (PAW) pseudopotentials. The Perdew–Burke–Ernzerhof (PBE) functional^{42,43} was used for structural optimizations, while the HSE06 hybrid functional⁴⁴ was used to calculate densities of states (DOS) and band structures. The energy cut-off for the plane wave basis set was set to 500 eV. The monolayer is on the *ab*-plane. A 40 Å high vacuum region was added in the *c* direction to isolate periodic images of the SiR monolayer along the *c*-axis. The first Brillouin zone (FBZ) was sampled using a $9 \times 9 \times 1$ Monkhorst–Pack grid of *k* vectors. The structural optimization convergence criterion was 0.01 eV Å⁻¹. All crystal structures and orbitals were visualized using VESTA.⁴⁵ To compare the calculated orbital energy levels of different SiR species, an inert He atom was placed 20 Å above each SiR layer (Fig. S1, ESI[†]), and the He 1s orbital energies in the different calculations were aligned. This calibration is an alternative to the use of the vacuum energy level, which is also used here to calculate work functions of the investigated species. The effective masses of hole and excited electron ($m_{\text{eff}}^{\text{h}}$ and $m_{\text{eff}}^{\text{e}}$) at band extrema were calculated using VASPKIT.⁴⁶ Throughout this work, symbols such as VBM _{Γ} , VBM_{*M*}, CBM _{Γ} , and CBM_{*M*} are used to denote the valence band maximum (VBM) and the conduction band minimum (CBM) at the subscripted specific *k* points in FBZ. The subscripted VBM and CBM may not be genuine VBM and CBM in the whole FBZ, which are labeled as the unsubscripted “VBM” and “CBM”.

CBM _{Γ} + 1, VBM _{Γ} – 1, *etc.*, are used to label orbital levels that are higher or lower than the explicitly specified ones.

To estimate the Si-adapted σ and π electron donating-accepting (sEDA and pEDA) capabilities of the ligands, natural population analyses were performed at the B3LYP/cc-pVTZ level.^{47,48} The sEDA and pEDA values of the ligands studied are listed in Table S2 (ESI[†]), defined as per the pioneering work of Oziminski and Dobrowolski (OD) for carbon chemistry, in which the same functional was employed.⁴⁹ Additionally, Si₁₃R₁₃H₉ models (Fig. S2, ESI[†]) were used to evaluate the natural atomic charge of Si atoms in SiR at the ω B97X-D/cc-pVDZ level.^{48,50} The central Si–R unit in the molecular model closely mimics each Si–R unit in the SiR monolayer. All these molecular calculations were performed using the GAMESS-US program.^{51,52}

TDDFT calculations for the SiR species were carried out using the CP2K-8.2 package⁵³ at the B3LYP⁴⁷/GPW⁵⁴ level with the Goedecker–Teter–Hutter (GTH) pseudopotential.^{55,56} The GPW hybrid basis set contains the TZVP-MOLOPT atomic basis⁵⁷ and the plane waves up to 350 Ry. The use of the B3LYP functional is motivated by its success in TDDFT calculations of semiconductors and confined system such as Si nanoparticle.^{58,59} Our confidence on this functional also comes from the good agreement between the TDDFT-calculated lowest absorption energy of SiH and the experimental value (*vide infra*). The SCF convergence criterion was 1×10^{-6} atomic unit (a.u.). $4 \times 4 \times 1$ supercells were employed to partially counteract the shrinkage of the FBZ to the Γ point in our TDDFT calculations. Two Fortran codes were developed to process the cube files of the orbitals from the supercell calculations: one returns the orbitals' *k* vectors in the FBZ of the genuine SiR unit cell, and the other returns two-electron integrals of four read-in orbitals: $\int dr_1 \int dr_2 \frac{1}{r_{12}} \phi_1(r_1) \phi_2(r_1) \phi_3(r_2) \phi_4(r_2)$, through numerical integration. Setting $\phi_1 = \phi_2$ and $\phi_3 = \phi_4$ ($\phi_1 = \phi_3$ and $\phi_2 = \phi_4$), we can calculate Coulomb (exchange) integral between two orbitals. The two-electron integrals mentioned below were obtained using the code. The two codes are available upon request.

III. Results and discussion

1. Geometrical structures

Fig. 1(a) shows a generic SiR structure with a hexagonal unit cell defined by the lattice vectors *a* and *b*. The SiR plane is buckled to the height *h* defined there. The first Brillouin zone (FBZ) is shown in Fig. 1(b), with the reciprocal lattice vectors \tilde{a} and \tilde{b} , high-symmetry points and their connecting paths. The ligands are categorized in Fig. 1(c) according to their σ - and π -donating and withdrawing capabilities, quantified by the Oziminski–Dobrowolski (OD)⁴⁹ sEDA and pEDA (*s*: σ , *p*: π , EDA: electron donator–acceptor descriptor) values. The sDpD quadrant in Fig. 1(c) is empty. Actually, we considered Ag atom as the sDpD ligand: SiAg was found to share similar structure with regular silicenes but exhibit no band gap. This hypothetical non-silicene metallic species is thus not discussed. SiLi, a salt with the Li⁺ cations located under (above) the buckled up (down) Si[−] anions (Fig. S1, ESI[†]), is also not of silicene type.



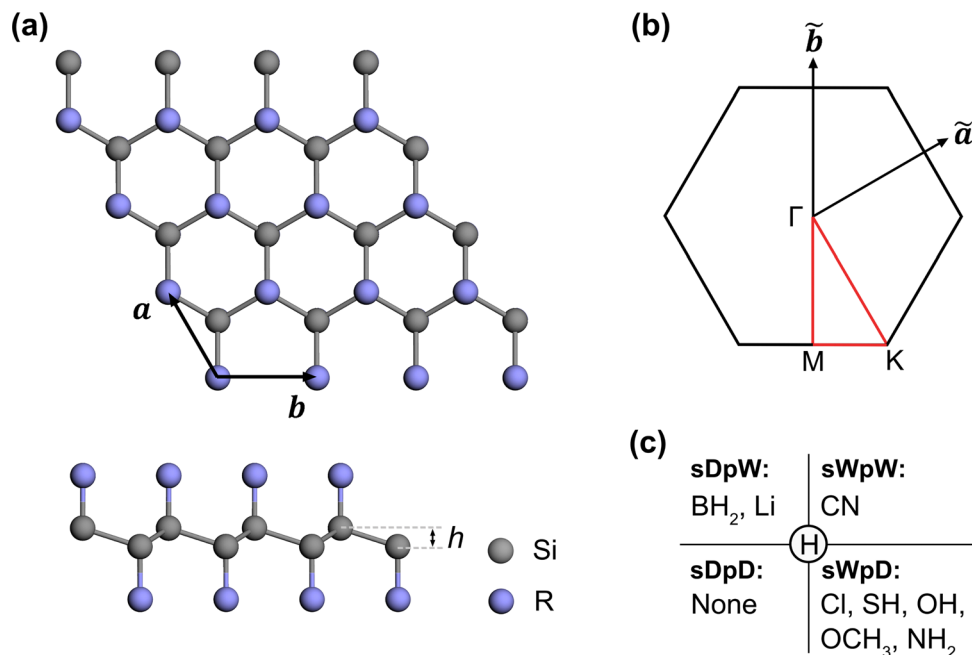


Fig. 1 (a) Top and side views of the honeycomb structure of SiR, and (b) its first Brillouin zone with the special k points being labeled. (c) Classification of the investigated ligands in the four quadrants defined by σ/π -withdrawing/donating capabilities.

However, this well-studied species is included in this work to highlight the differences between its anionic silicide framework and the regular silicane frameworks.

The OD sEDA and pEDA were obtained for the C–R bonds, *i.e.*, they are carbon-adapted. The use of the carbon-adapted OD EDA values for SiR may sound questionable in the beginning, but is fully justified in Section S2 (ESI[†]), where we calculated the Si-adapted sEDA and pEDA indices and compared them with the C-adapted ones proposed by OD. A good agreement between the two sets is seen. For comprehensiveness, the full list of ligands in the OD paper were considered,⁴⁹ much more than the a few ligands studied in this work. Since the σ/π -withdrawing/donating are local covalent effects, in both finite-sized and extended systems, the EDA values obtained using molecules can be used

to explain the ligands' impacts to silicanes, in which they are covalently bonded to the Si framework.

The optimized structural parameters are summarized in Table 1. The parameters of SiH agree well with previous theoretical and experimental results.^{39,60–62} The hexagonal Si framework is maintained for SiCl, SiCN, and SiOCH₃, with essentially $a = b$, and is slightly broken for SiSH ($a = 3.909$ Å, $b = 3.979$ Å), SiOH (3.882, 3.917 Å), SiNH₂ (3.832, 3.903 Å), and SiBH₂ (3.923, 3.884 Å). The Si–Si bond lengths in all SiR species are marginally elongated (<2.5%) from the SiH analogue, within the tolerable range of 17% elongation under equibiaxial strain.⁶³ The bulky OCH₃ leads to additional transversal stress and the largest a and b . The sWpD ligands slightly increase h from the SiH analogue, due to their π^* antibonding (π -donating) interactions with the Si–Si bonds; their

Table 1 Summary of DFT calculated properties of the investigated systems

	SiH	SiCl	SiSH	SiOH	SiOCH ₃	SiNH ₂	SiCN	SiBH ₂	SiLi
$a/\text{Å}$	3.888	3.918	3.909	3.882	3.977	3.832	3.935	3.923	3.730
$b/\text{Å}$	3.888	3.921	3.979	3.917	3.975	3.903	3.933	3.884	3.730
Space group	$P\bar{3}m1$	$P\bar{3}m1$	$P1$	$P1$	$P1$	$P1$	$P1$	$P1$	$P\bar{3}m1$
Average Si–Si bond length/Å	2.357	2.378	2.391	2.374	2.411	2.374	2.381	2.393	2.393
Average Si–R bond length/Å	1.501	2.075	2.169	1.694	1.683	1.780	1.831	1.992	2.628
$h/\text{Å}$	0.719	0.732	0.749	0.763	0.723	0.787	0.713	0.791	1.043
Si charge/atomic unit	+0.071	+0.305	+0.195	+0.520	+0.623	+0.430	+0.319	+0.003	−0.769
Work function Φ/eV	4.98	5.57	4.58	3.93	3.47	3.42	9.08	5.06	2.36
m_{eff}^c/m_0^a	0.170	0.247	0.194	0.274	0.273	0.296	0.175	0.190	0.862
	0.169	0.247	0.199	0.289	0.271	0.284	0.175	0.205	0.861
	2.996							1.155	
	0.122							0.659	
m_{eff}^c/m_0^a	0.570	1.789	1.196	0.690	0.487	0.314	0.772	0.153	0.465
	0.540	1.435	0.553	0.474	0.660	0.583	0.856	0.235	0.451

^a The first (second) entry was for the effective mass at CBM_Γ or VBM_Γ along the Γ -to- M (Γ -to- K) direction. For SiH (SiBH₂), the third and fourth entries was obtained at CBM_M (CBM_K) for the M -to- Γ (K -to- Γ) and M -to- K (K -to- M) directions. All the effective masses were calculated using HSE06 functional. m_0 is the atomic unit of mass, *i.e.*, the electronic mass.



h values correlate well with their pEDA values, except for the bulky OCH_3 , whose extra transversal stress suppresses the buckling. The sWpW CN ligand slightly reduces h due to the lack of π -donating capability. The sDpW BH_2 ligand, through its dominating σ -donating capability, makes the Si atoms less siliconium-like (see the +0.003 Si charge in Table 1) and hence significantly increases h . Naturally, the h value is even much larger in the purely silicide framework in SiLi.

The atomic charge certainly reflects the electronegativity difference between Si and R. Shown in Fig. S3 (ESI[†]) is the anticorrelation between the quantities, where the electronegativity of the atoms directly connected to Si is chosen to represent the electronegativity of R. The -0.769 atomic unit (a.u., *i.e.*, the amount of electronic charge) Si charge in SiLi is close to the -1 formal oxidation state of Si in the LiSi salt and reflects the ionicity of the Si–Li interaction. This ionicity is consistent with the negligible pDOS of Li in the valence band (*vide infra*, Fig. 2(i)). On the other hand, the $+0.305$ a.u. Si charge in SiCl is far from the $+1$ formal oxidation state of Si in this compound, reflecting the polar yet covalent nature of the Si–Cl bond. Despite the overall anticorrelation, the Si charge depends on the detailed covalent interaction, and the electronegativity of the atom in contact with Si may not fully represent the electronegativity of R. For instance, given the same O atom connected to Si, the Si charge is higher in SiOCH_3 than in SiOH. Overall, atomic charge is only a qualitative indicator.

2. Electronic structures

The ligand-induced changes in band structure and density of state (DOS) are far more significant (Fig. 2) than in geometric structure. The valence band maximum (VBM) and conduction band minimum (CBM) of SiH are located at Γ and M in FBZ (Fig. 2(a)), *i.e.*, VBM_Γ and CBM_M . VBM_Γ is doubly degenerate. The degeneracy arises from the e-type $\sigma_{\text{Si-Si}}$ bonding orbitals (BOs), a result of the C_{3v} site symmetry of each Si–H moiety (Fig. 3(a) and 5(a), (b)). CBM_Γ is slightly higher than CBM_M by 0.21 eV, which is the true CBM. Consequently, SiH features a 2.93 eV Γ -to-M indirect band gap and a 3.14 eV Γ -to- Γ direct gap, consistent with previous experimental and computational results.^{31,63–65} The atom-projected density of states (pDOS) of SiH in Fig. 2(a) shows almost no H contribution at the band edges. The CBM_Γ orbital exhibits antibonding (AB) character between adjacent SiH units (see the red lines in Fig. 3(a)), while primarily nonbonding (NB) character for each Si–H bond. Our numerical orbital analysis indicates that the periodic modulation function of the CBM_M orbital is a 45:55 mixture of the a_1 -type CBM_Γ and the e-type degenerate $\text{CBM}_\Gamma + 1,2$ orbitals (the second and third lowest CB orbitals at Γ , plotted in Fig. S4, ESI[†]). Since the $\text{CBM}_\Gamma + 1,2$ orbitals are delocalized on the hollow centres of the 6-membered rings, CBM_M is transversally polarized towards the ring centers. The effective mass of excited electron and hole for Γ -to-M (M-to- Γ) were calculated to be 0.170 (2.996) and 0.570 m_0 (Table 1), respectively. These values

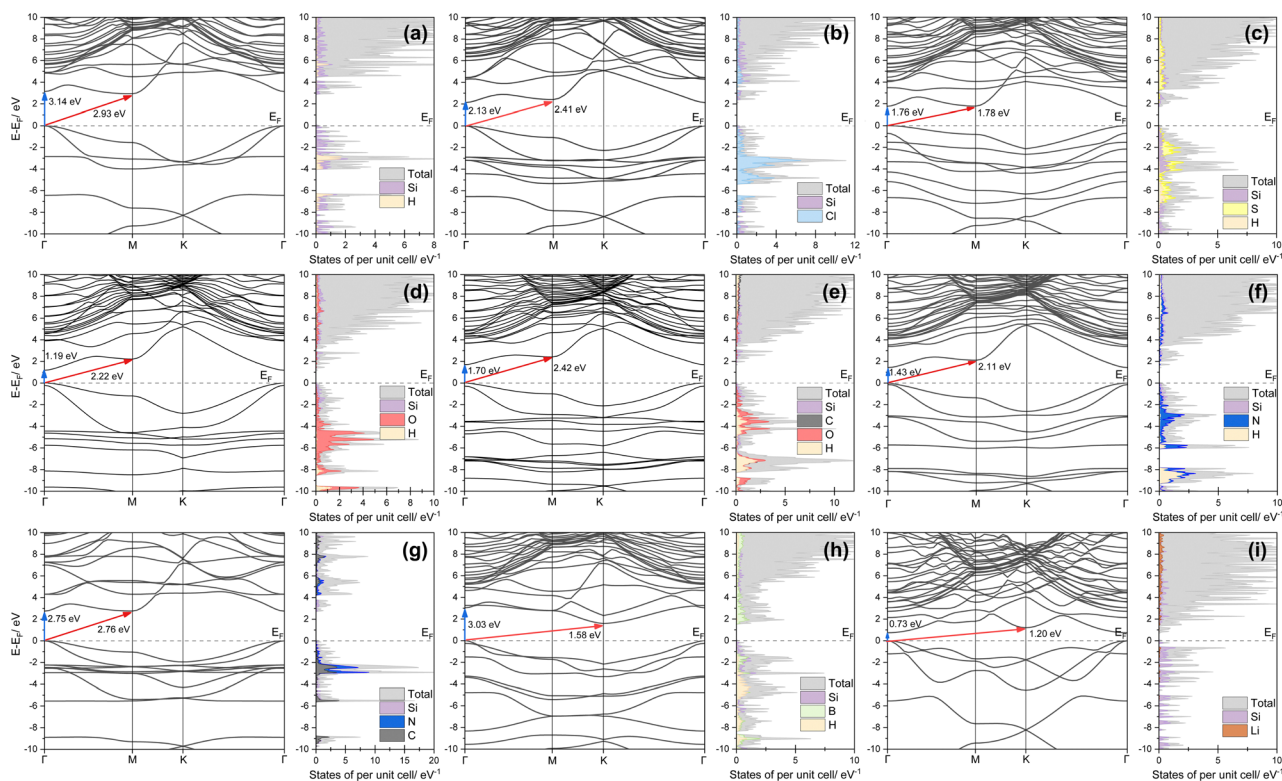


Fig. 2 The electronic band structures, densities of states, and atom-projected densities of states of (a) SiH, (b) SiCl, (c) SiSH, (d) SiOH, (e) SiOCH_3 , (f) SiNH_2 , (g) SiCN, (h) SiBH_2 , and (i) SiLi. The red and blue arrows indicate the indirect and direct band gaps, respectively. All band structures and densities of states have been shifted so that the Fermi energy is at 0 eV.



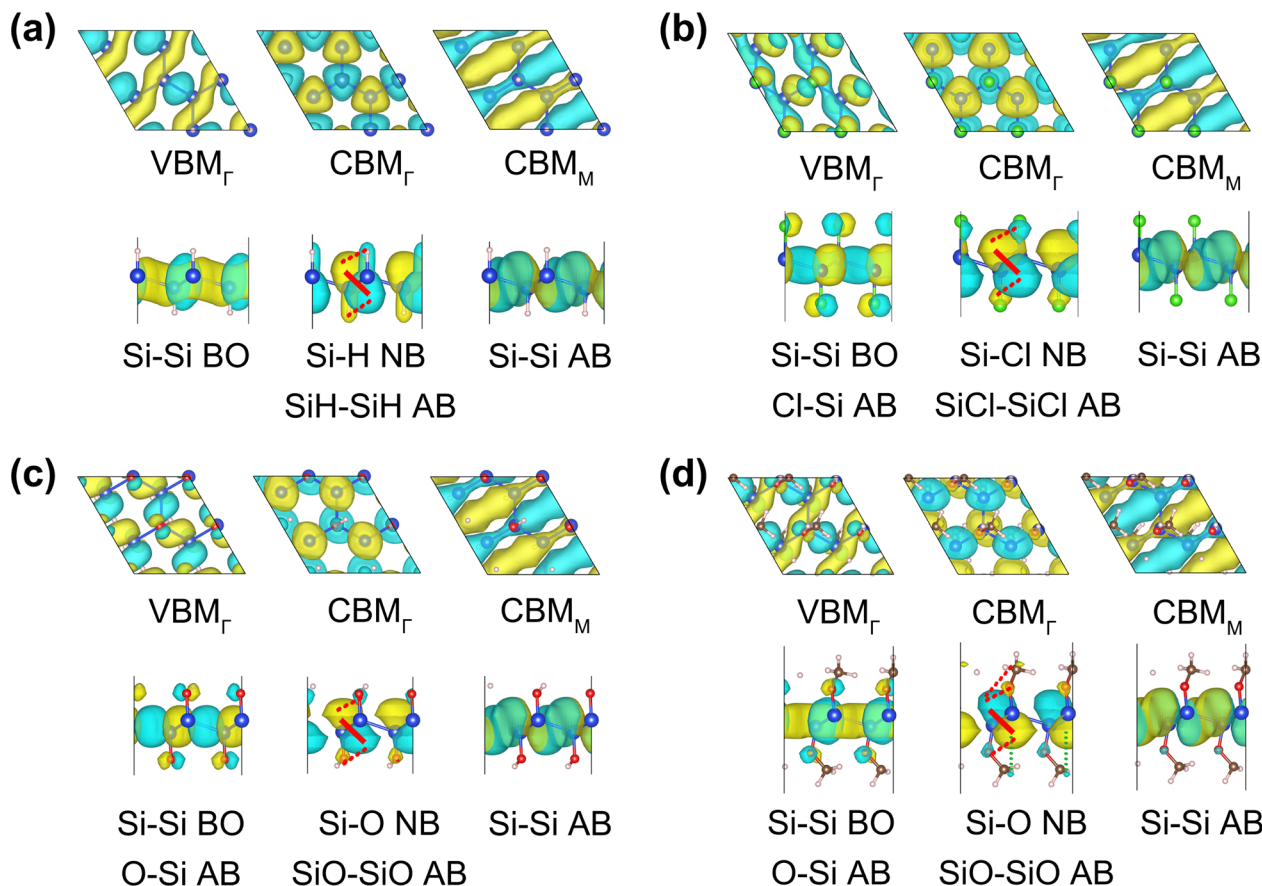


Fig. 3 The top and side views of the VBM_Γ, CBM_Γ, and CBM_M orbitals of (a) SiH, (b) SiCl, (c) SiOH, and (d) SiOCH₃. The characters of the orbitals with respect to certain pairs of atoms are specified: BO = bonding orbital, NB = nonbonding, and AB = antibonding.

imply higher mobility of the excited electron at Γ than at M , and higher electron mobility than hole mobility.

The five sWpD ligands reduce the Γ -to- Γ gaps more than the Γ -to- M gaps (the blue vs. red arrows in Fig. 2(a)–(f)), leading to direct band gaps for these functionalized SiR species. Their band edge orbitals resemble the SiH counterparts but exhibit π lone pair lobes on the π -donating ligands in the VBM_Γ orbitals (Fig. 3(b)–(d)). The doubly degenerate VBM_Γ level in SiH and SiCl is split to the VBM_Γ and VBM_Γ – 1 levels in all SiR species with non-atomic or non-linear ligands in the lack of the 3-fold site symmetry. The energy inversion between CBM_M and CBM_Γ results from their different spatial distributions. The CBM_M orbital, distributed in the hollow space of the Si 6-membered rings, is less stabilized by the more positively charged Si framework induced by the oxidation (see Table 1 for the Si charges). On the contrary, the CBM_Γ localized on the Si atoms is more stabilized. The pDOS of the five substituted SiR species in Fig. 2(a)–(f) show greater contributions of the ligands to the band edges than in SiH, due to the aforementioned π^* interaction around VBM (Fig. 3 and Fig. S5, ESI[†]), which raises or prevents further decrease of VBM and contributes to the band gap reductions in these SiR species. The effective masses of excited electron at CBM_Γ of the five SiR species only slightly change from the SiH analogues. This holds for all SiR species

under consideration, except SiLi. The effective hole masses of the five are all larger than the corresponding effective excited electron masses. This holds for all species except SiBH₂ and SiLi. The effective hole mass at VBM_Γ becomes the most anisotropic for SiSH, with twice difference in the Γ -to- M direction vs. the Γ -to- K direction. This is consistent with the significant anisotropy of the π^* interaction at VBM_Γ (*vide infra*, Fig. 6(g) and (h)). The most significant anisotropy in effective mass of excited electron is seen at CBM_M of SiH, with ~ 30 times difference in the M -to- Γ vs. M -to- K directions. This is attributed to the especially flat M -to- Γ CBM section in Fig. 2(a), which is responsible for the 2.996 m_0 large effective mass of excited electron. This flatness reflects the substantial interaction between the periodic modulation functions of CBM_Γ and CBM_Γ + 1,2 (*vide supra*) along the path between Γ and M in FBZ.

With the sWpW ligand, the VBM and CBM of SiCN are both at Γ , giving a 2.75 eV direct band gap (Fig. 2(g)). However, CBM_M is only 0.01 eV higher, suggesting a competition between direct and indirect band gaps. As mentioned above, the periodic modulation functions of CBM_M feature a mixture of CBM_Γ and CBM_Γ + 1,2. The CBM_Γ + 1,2 component brings Si-C π character into CBM_M, which is stabilized by the π -withdrawing CN (Fig. S6 and S7, ESI[†]) to be pseudo-degenerate



with CBM_Γ . The effective masses of excited electron in the Γ -to- M and Γ -to- K directions are similar to the SiH analogues, while the effective masses of hole are noticeably higher than the SiH analogues. SiCN can serve as an electron transport layer like SiH.

BH_2 and Li were chosen as representative sDpW ligands. The SiLi VBM and CBM are located at Γ and are localized separately on Si^- and Li^+ (Fig. S6(c), ESI[†]), a feature of salts, with a direct gap of 0.73 eV. For SiBH_2 , the CBM is shifted to K, resulting in an indirect band gap of 1.58 eV (Fig. 2(h)). The VBM_Γ and CBM_Γ orbitals of SiBH_2 resemble those of the other SiR species (Fig. S6(b), ESI[†]). CBM_K is the genuine CBM of SiBH_2 , as the bonding interaction between the 2p vacant orbitals of adjacent BH_2 ligands is oriented along the b direction and is maximized by this specific k vector. The effective masses of excited electron at CBM_K in the K -to- Γ and K -to- M directions differ by about twice, which is consistent with the anisotropy of the bonding interaction mentioned above.

3. Excited states, the applicability of the one-particle description

Because of the Born-von Karman boundary condition imposed by running the TDDFT calculations only at the Γ point of the $4 \times 4 \times 1$ supercell (*vide supra*, see also Section S3, ESI[†]), only the k vectors of $\frac{m}{4}\tilde{a} + \frac{n}{4}\tilde{b}$ in FBZ of the true SiR unit cell were sampled. The TDDFT-DOS of the SiR species are shown in Fig. 4. For SiH, the lowest excited state (labeled ① in Fig. 4(a)) is a doubly degenerate optically bright state, with excitation from the doubly degenerate VBM_Γ (Fig. 5(a) and (b)) to CBM_Γ (Fig. 3(a)). The calculated absorption energy of 3.00 eV is in agreement with the experimental absorption onset at 3.0 eV,³¹ which reflects the accuracy of using the B3LYP functional in describing this class of materials. In Fig. 4(a), the doubly degenerate Γ state is immediately followed by a triply degenerate M state (the lower ② state in the panel) with excitations from the doubly degenerate VBM_Γ to the triply degenerate CBM_M . Another triply degenerate state of the same excitation scheme (the higher ② state) lies 0.10 eV higher. According to the one-particle orbital description, a 6-fold degeneracy should have arisen from the VBM_Γ -to- CBM_M ($2 \times 3 = 6$) transitions, which is however split to the two 3-fold degenerate ② levels. The degenerate VBM_Γ component orbitals can be linearly combined into one symmetric and one antisymmetric orbital with respect to the symmetry plane containing the long diagonal axis of the rhombic unit cell, which we call $\text{VBM}_{\Gamma'}$ and $\text{VBM}_{\Gamma''}$ orbitals (Fig. 5(a) and (b)). The prime (double-prime) denotes evenness (oddness) with respect to the symmetry plane. The $\text{CBM}_M(1/2, 1/2)$ orbital shown in Fig. 5(c) has $k = \frac{1}{2}\tilde{a} + \frac{1}{2}\tilde{b}$ and is also a prime orbital. One state in the lower (higher) ② branch purely consists of the $\text{VBM}_{\Gamma''}$ -to- $\text{CBM}_M(1/2, 1/2)$ ($\text{VBM}_{\Gamma'}$ -to- $\text{CBM}_M(1/2, 1/2)$) excitation; it is an M'' (M') state. The other two states in the respective ② branch are related to the M'' or M' state by C_6 and C_3 rotations, and the relevant $\text{CBM}_M(1/2, 0)$ and $\text{CBM}_M(0, 1/2)$ orbitals are shown in (Fig. S9(e) and (f), ESI[†]). The splitting of the two ② branches is actually the M'' - M' splitting.

According to the theory of configuration interaction with single excitations (CIS) for molecular systems, the energy ($E(\text{H} \rightarrow \text{P})$) of one singlet excitation scheme equals the energy gap between the hole (H) and particle (P) orbitals ($\varepsilon_P - \varepsilon_H$), subtract by the Coulomb integral between the two orbitals (J_{HP} , which is the electrostatic attraction between the positively charged hole and the negatively charged electron), and plus twice their exchange integral ($2K_{\text{HP}}$, which is the explicit correlation energy between the electron left in the hole orbital and the electron excited to the particle orbital),⁶⁶ *i.e.*, $E(\text{H} \rightarrow \text{P}) = \varepsilon_P - \varepsilon_H - J_{\text{HP}} + 2K_{\text{HP}}$. The M' and M'' states involve the same orbital energy gap. Numerical integration yields the same value of the $\text{VBM}_{\Gamma'}\text{-CBM}_M(1/2, 1/2)$ and $\text{VBM}_{\Gamma''}\text{-CBM}_M(1/2, 1/2)$ Coulomb integrals. However, the exchange integral is 0.053 eV for the $\text{VBM}_{\Gamma'}\text{-CBM}_M(1/2, 1/2)$ hole-particle pair, and 0.013 eV for the $\text{VBM}_{\Gamma''}\text{-CBM}_M(1/2, 1/2)$ pair. Twice the 0.04 eV difference qualitatively explains the 0.10 eV M'' - M' splitting. Certainly, the TDDFT formalism is similar but not identical to the CIS formalism. The use of the CIS language to explain the TDDFT result is of qualitative nature, but insightful. This splitting of the extended SiH's excited state can be interpreted using the language of molecular systems. This is because the splitting here and the splittings of molecular electron configurations to term symbols are both driven by spatial confinement of orbitals.

The ③ states at the midpoint of the Γ -to- M path feature a similar magnitude of prime-double-prime splitting. Each ③ branch contains 6 degenerate states due to the 6-fold degeneracy of $\text{CBM}_{1/2\Gamma_M}$. The states grouped under ④, ⑤, and ⑥ in Fig. 4(a) can be viewed as the upshifted analogues of those under ①, ②, and ③, respectively, now with the hole on the lower-lying 6-fold degenerate $\text{VBM}_{1/2\Gamma_M}$. The second and third lowest state under ⑥ are two closely-lying optically bright Γ states. It is noteworthy that if we fully sample the FBZ by using even larger supercell, *e.g.*, $16 \times 16 \times 1$ and further broaden the already denser TDDFT-DOS peaks, the resultant continuous DOS will blur the splittings induced by the explicit electron. This justifies the use of the relatively small supercell and the discrete TDDFT-DOS as qualitative tools to deepen our understanding of the excited states. Now we have nailed down the ~ 0.1 eV splitting of transitions involving degenerate hole and particle orbitals as the magnitude of explicit electron correlation effect. If the ~ 0.1 eV energy difference does not matter for a specific study of SiR species, then the one-particle description is still applicable.

The TDDFT-DOS of SiCl in Fig. 4(b) shares features in Fig. 4(a): (1) a low-energy optically bright VBM_Γ -to- CBM_Γ transition; (2) the prime-double-prime splittings; (3) a high-energy optically bright $\text{VBM}_{1/2\Gamma_M}$ -to- $\text{CBM}_{1/2\Gamma_M}$ transition; (4) all low-energy excited states having hole and particle orbitals on the Γ -to- M path. Point (4) is shared by all SiR species considered in this work except SiBH_2 and SiLi, since the Γ -to- M path contains all band edge orbitals in all species except the two. In Fig. 4(b), the VBM_Γ -to- CBM_M excitation lies substantially higher than the VBM_Γ -to- CBM_Γ excitation, as the CBM_M does not compete for the true CBM. This is also seen in Fig. 4(c)–(g) for the same reason.

The rhombic primitive unit cell is slightly distorted to a parallelogram in SiOH, SiOCH_3 , SiSH, and SiNH_2 . Still, we use



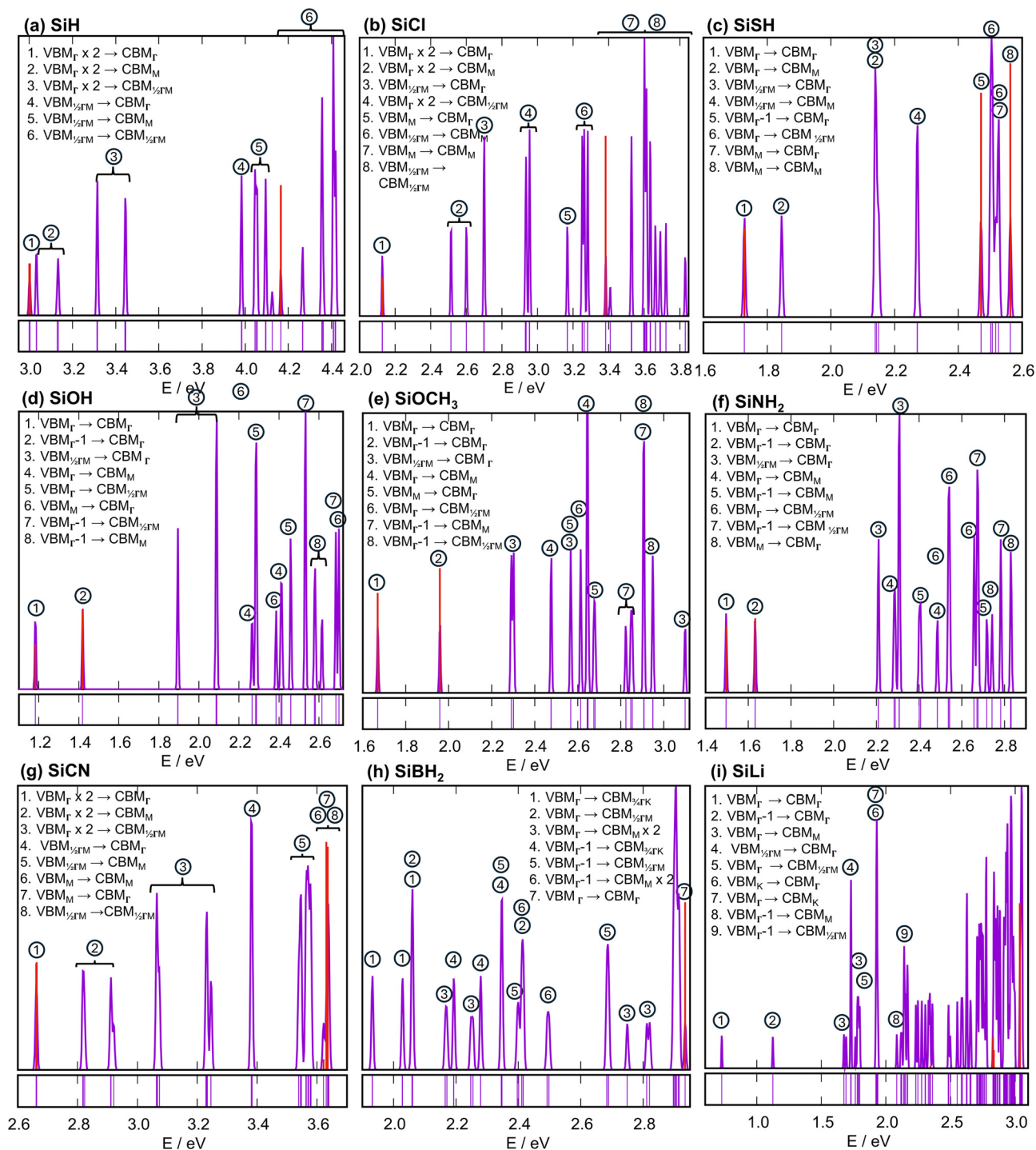


Fig. 4 TDDFT-DOS of the six labeled silicane species obtained using their $4 \times 4 \times 1$ supercells. They were obtained by Gaussian broadening with $\sigma = 0.007$ eV to each state's spike in the lower minor panels and summing over the Gaussians, and oscillator strengths are plotted as red spikes. States involving the same set of hole and particle orbitals are grouped and labeled by circled numbers. " $VBM_{\Gamma} \times 2$ " in (a) and (b) indicates the doubly degenerate VBM_{Γ} . Equal-height spikes in the lower minor panels indicate the existence of states at specific energies.

M (which is strictly applicable only to rhombic) to label the states at $\frac{1}{2}\tilde{a}$, $\frac{1}{2}\tilde{b}$, and $\frac{1}{2}\tilde{a} + \frac{1}{2}\tilde{b}$ in their FBZs. Certainly, the 3-fold degeneracy of the M states is lifted, and the six Γ -to-M paths are no longer symmetrically related. Correspondingly, the TDDFT-DOS peaks grouped under the same numeric label are more

broadly distributed in Fig. 4(c)–(g) than in Fig. 4(a), (b). Another feature in Fig. 4(d)–(f) is the splitting of the band edge optically bright state of 0.24 eV in SiOH, 0.29 eV in SiOCH₃, and 0.14 eV in SiNH₂ (Table S1, ESI[†]). They arise from the splitting of the originally doubly degenerate VBM_{Γ} level due to the loss of the 3-fold site symmetry, which leads to the 0.25, 0.30, and 0.14 eV





Fig. 5 Frontier orbitals of the $4 \times 4 \times 1$ SiH supercell. The orbitals are plotted with the same isosurface value of ± 0.006 a.u. and are normalized in the supercell. The red dashed lines and the green dash line in (a) highlight a unit cell and the symmetry plane that cuts through the long diagonal axis of the super cell.

respective $\text{VBM}_{\Gamma} - 1 - \text{VBM}_{\Gamma}$ gaps (Fig. 2). The splittings in orbital energy levels match the splittings in excited states, *i.e.*, the one-particle orbital description explains the splittings in those Γ excited states.

The comparison of the VBM_{Γ} and $\text{VBM}_{\Gamma} - 1$ orbitals of SiOH in Fig. 6(a) and (b) attributes their splitting to the different O lone pairs involved in their π^* interactions. In the $\text{VBM}_{\Gamma} - 1$ (VBM_{Γ}) orbital, the O $\sigma(\pi)$ lone pairs that are in plane with (perpendicular to) the OH bonds provide the π^* interactions on both sides of the Si framework. The σ lone pair is well known to be more stable and thus exerts weaker π^* interaction to $\text{VBM}_{\Gamma} - 1$, resulting in the $\text{VBM}_{\Gamma} - 1 - \text{VBM}_{\Gamma}$ splitting and also the splitting of the band edge bright state. This explanation also applies to the similar splitting in SiOCH_3 .

The NH_2 and SH ligands differ from OH and OCH_3 by having only one active lone pair in exerting the π^* interactions. The two themselves differ: the NH_2 ligands above and under the Si framework are neither *trans* nor *cis* to each other, while the SH ligands on two sides are *trans* to each other (see the red lines in Fig. 6(f) and (i)). Thus, the antibonding interaction in SiNH_2 cannot be concentrated in one VBM_{Γ} orbital as in SiSH. Also, unlike in a free NH_2 radical, the lone pair lobe of the NH_2 ligand is not symmetric with respect to the plane that bisects the H–N–H angle. It can be polarized through mixing with one of the N–H bonding lobes to adjust its π interaction with the Si framework. This flexibility, absent in OH, OCH_3 , and SH ligands, allows the NH_2 ligand to exert similar π^* interactions in $\text{VBM}_{\Gamma} - 1$ and VBM_{Γ} , resulting in their smaller splitting, and

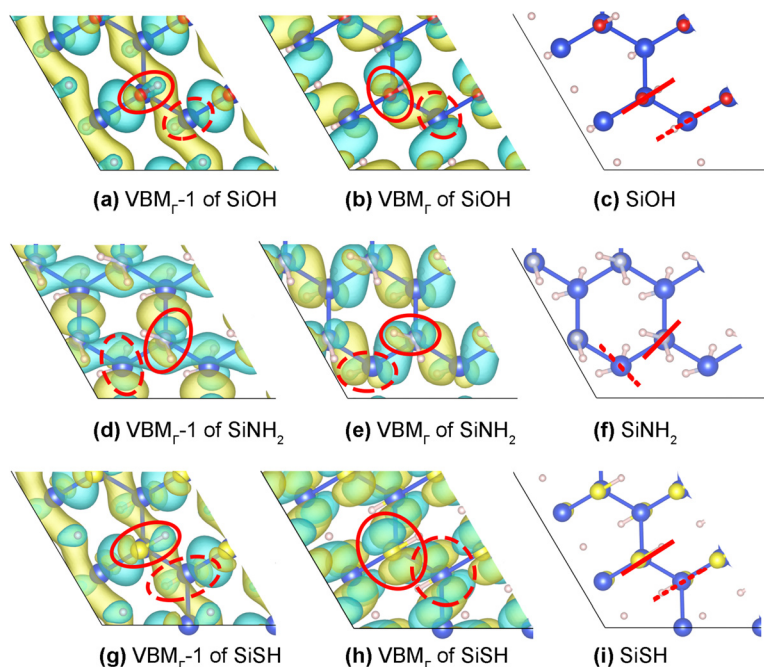


Fig. 6 The $\text{VBM}_{\Gamma} - 1$ and VBM_{Γ} orbitals of SiOH ((a) and (b)), SiNH_2 ((d) and (e)), and SiSH ((g) and (h)). The solid (dashed) ovals highlight the orientations of lone pair lobes of the ligands above (under) the Si framework. Red solid (dashed) lines in (c), (f), and (i) represent the planes that bisect the H–N–H angles or contain the O–H or S–H bonds above (below) the Si framework.



the corresponding smaller splitting of band edge bright excitations.

The SiSH TDDFT-DOS in Fig. 4(c) shows a distinct difference from the other panels: the $\text{VBM}_{\Gamma} - 1$ -to- CBM_{Γ} excitation (labeled ⑤ in Fig. 4(c)) is not a band edge excitation, as it lies 0.75 eV higher. This is due to the 0.70 eV substantial $\text{VBM}_{\Gamma} - 1$ - VBM_{Γ} splitting in SiSH, which pushes the $\text{VBM}_{\Gamma} - 1$ orbital deep into the VB (the band structures in Fig. 2(c) vs. Fig. 2(d), (f)). The large splitting is ascribed to: (1) each SH ligand mainly exerts π^* interaction on VBM_{Γ} through its π lone pair. The σ lone pair with S 3s and SH bonding character is inactive in pushing up $\text{VBM}_{\Gamma} - 1$ (Fig. 6(g) vs. Fig. 6(h)); (2) the SH ligands above and under the Si framework are *trans* to each other, like the OH ligands in SiOH (Fig. 6(c) and (i)). The SH π lone pair lobes above and below are hence parallel and concertedly push up the VBM_{Γ} energy.

Except for the red-shifts due to its smaller band gaps, the TDDFT-DOS of SiCN in Fig. 4(g) shares similar features as the SiH analogue in Fig. 4(a). The splittings of the ② states and ③ states in SiCN are for the same reason as the analogous splittings in SiH. In SiBH_2 , the BH_2 $2p_{\pi}$ vacant ($\sigma_{\text{B-H}}$) orbital lobes form bonding (antibonding) interaction with the $\sigma_{\text{Si-Si}}$ bonding orbital lobes from both above and under the Si framework (see Fig. S10(a) and (b), ESI[†]) and stabilize (destabilize) the $\text{VBM}_{\Gamma} - 1$ (VBM_{Γ}) orbital, leading to their 0.25 eV orbital energy gap. Correspondingly, there is a 0.26 eV gap between the lowest excited states with the hole on the VBM_{Γ} orbital and the $\text{VBM}_{\Gamma} - 1$ orbital, *i.e.*, the lowest ① and ④ states in Fig. 4(h). All these band edge states are optically dark states since the electron is excited to the $\text{CBM}_{3/4\Gamma\text{K}}$ orbital without conserving the crystal momentum. The true lowest unoccupied orbital is the CBM_{K} orbital, which is not captured by our $4 \times 4 \times 1$ supercell model. The $\text{CBM}_{3/4\Gamma\text{K}}$ (1/4, 1/2) orbital is plotted in Fig. S10(c) (ESI[†]) as an approximate of the CBM_{K} orbital. The 1/4 index determines the largely nonbonding interaction along the *a*-axis, while the 1/2 index determines the maximum bonding interaction between the BH_2 $2p_{\pi}$ vacant lobes along the *b*-axis, as all the vacant lobes, above or under the Si framework, are pointing in the *b*-direction. The maximum bonding character along *b* and largely nonbonding character along *a* explain the shift of the true CBM from Γ or M in the other SiR species to K in SiBH_2 .

The lowest optically bright state (⑦ in Fig. 4(h)) lies 1 eV above the lowest band edge state and has the CBM_{Γ} orbital as the particle orbital, which is similar to the analogue of SiH. Since both the CBM_{Γ} and CBM_{K} orbitals are mostly not within the Si framework, they have substantial spatial overlap, as illustrated by their product in Fig. S8 (ESI[†]). This spatial overlap should exist between CBM_{Γ} and the other CBM orbitals on the Γ -to-K path. The substantial spatial overlaps suggest that after the VBM_{Γ} -to- CBM_{Γ} optical excitation, the electron can non-adiabatically relax to the CBM_{K} orbital, resulting in a long-living electron on the BH_2 group and a long-living hole in the Si framework. The long-living spatially separated holes and electrons will find extensive use in photovoltaics and photochemistry.

SiLi is a well-studied species for its extensive use in Li-ion battery (LIB).^{18,67} It is a purely ionic system and adopts a

completely different structure compared to the other aforementioned species. In a range of 0.69 eV, the bottom of CB is all dominated by Li atomic orbitals (Fig. 2(i) and Fig. S6, ESI[†]), while the VBM is Si-dominated. Consequently, all low-lying excited states labeled in Fig. 4(i) are optically dark states with an electron being transferred from the Si^- framework to the Li^+ cationic layer, which will be easily returned to the Si framework due to the electropositivity of Li. Therefore, SiLi is immune to photodegradation and is a promising LIB material, as already well known.⁶⁷⁻⁶⁹

The DFT and TDDFT band gaps are compared in Fig. S11 (ESI[†]). The excellent agreement between the two sets of independently obtained gaps corroborates their respective accuracies.

4. Band edge variation

We used the 1s energy of an extra added He atom to calibrate orbital energy levels of the nine SiR species (Fig. S12, ESI[†]). The so calibrated band edge energies are compared in Fig. 7. Replacing H with Cl downshifts both VBM and CBM due to the ~ 4 -fold more positively charged Si framework (compare the Si atomic charges of SiH and SiCl in Table 1). The VBM does not downshift as much due to the Cl π^* interaction (Fig. 3(b)). The σ -withdrawing capability of SH is almost half that of Cl (Table S2, ESI[†]), downshifting VBM and CBM less in SiSH. This less VBM downshift is even counteracted by the stronger π -donating capability of SH, which raises the VBM higher than in SiH. The π -donating capabilities of OH, OCH_3 , and NH_2 are nearly twice as strong as that of Cl (Table S2, ESI[†]), and the π -donating capabilities dominate over the σ -withdrawing capabilities of the three ligands. Consequently, the VBMs of SiOH, SiOCH_3 , and SiNH_2 are pushed up by 0.93, 1.63 and 1.53 eV higher than in SiH, despite their more positively charged Si atoms (Table 1). The higher VBM of SiOCH_3 vs. SiOH can be attributed to: (1) the higher pDEA of OCH_3 ; (2) the elongation of the Si-Si bond induced by the transversal stress of the bulky OCH_3 groups, whose $\sigma_{\text{Si-Si}}$ bonding character is hence weakened.

The antibonding character in the CBM_{Γ} orbitals in Fig. 3 and Fig. S5, S6 (ESI[†]) are represented by the three red lines (1 solid + 2 dashed). This π^* interaction is stronger in SiOH, SiOCH_3 , and SiNH_2 than in SiCl due to the shorter Si-O and Si-N bonds (Table 1), leading to the higher CBMs of the former three. Additionally, the CBM of SiOCH_3 is further raised by the extra antibonding interaction between the CH_3 moieties and the Si atoms (the green dotted lines in Fig. 3(d)), which originates from steric hindrance of the CH_3 moieties.

In summary, the sWpD ligands pull down (push up) band edges through their σ -withdrawing (π -donating) capabilities, and the balance between the opposite effects determines the final VBM and CBM energies. On the contrary, the σ - and π -withdrawing capabilities of CN synergistically downshift the VBM and CBM of SiCN to exceptionally low. The BH_2 ligand raises (lowers) the SiBH_2 VBM through its σ -donating (π -withdrawing) capability. The two effects largely cancel and result in the VBM marginally changed from the SiH analogue.





Fig. 7 Comparison of band edge energies of the nine investigated species.

The CBM of SiBH₂ is significantly lower than SiH due to the bonding interaction between the BH₂ vacant lobes (Fig. S6(b) and S10(c), ESI[†]). The VBM of SiLi is much higher than the other SiR species because of its purely anionic Si framework.

We also calculated the work functions (Φ) for the SiR species as the difference between the vacuum and Fermi levels. Their data in Table 1 are in excellent agreement with the He-calibrated VBM energies shown in Fig. 7. The He calibration provides an alternative method for comparing energy levels of 2D materials obtained in individual calculations.

The qualitative understanding the influence of the σ/π -withdrawing/donating capabilities of ligands on the band edge energies motivated us to quantify this impact. We fitted the calibrated VBM and CBM energies of the silicane species (excluding SiLi) using 2D polynomial functions of the sEDA (x) and pEDA (y) values of the R ligands. The resultant functions (in eV) for silicanes with sWpD R reads:

$$f(x,y) = \text{VBM/CBM}(\text{SiH}) + a(\cos\theta x + \sin\theta y) + b(-\sin\theta x + \cos\theta y) + c(\cos\theta x + \sin\theta y)^2 + d(-\sin\theta x + \cos\theta y)^2,$$

with parameters $\theta = 1.3706$, $a = 194.095$, $b = -14.877$, $c = -2613.28$, $d = 24.996$ for VBM and $\theta = 1.3685$, $a = 242.098$, $b = -19.264$, $c = -3544.81$, $d = 27.185$ for CBM. The VBM/CBM(SiH) term means taking the VBM or CBM of SiH as the reference. The details of the fittings are given in Section S4 (ESI[†]). We chose this fitting function since it is the lowest order polynomial expansion that can be fitted against the limited data set. The assumption under this choice is the continuity of the ligand's impact on the band edge energies. This is a reasonable assumption since the fundamental factors that determine the covalent impact: (1) overlap between orbital lobes on the ligands and the Si framework; (2) energies of the frontier orbitals of the ligands, are continuous functions. These empirical functions allow quick prediction of band edge energies of a

silicane species, given the readily obtainable sEDA and pEDA values of ligands. The functions are plotted in Fig. S15 (ESI[†]). For possible future applications of the functions, or more accurate functions to be developed based on a similar idea, we present the Si-adapted sEDA and pEDA values for a much larger set of ligands than the currently investigated nine in Table S2 (ESI[†]).

IV. Conclusions

We systematically investigated the geometrical structures, electronic structures, and band edge excited states of silicanes with ligands that fall into three classes based on their electron-withdrawing/donating capabilities. The sWpD ligands reduce the band gap of the substituted silicanes and also tend to change the mixture of direct and indirect gap in SiH to purely direct gaps. This is attributed to the different spatial distributions of the CBM_F and CBM_M orbitals and consequently their different responses to the more positively charged Si framework induced by the oxidative ligands. The sWpW ligand reduces and maintains the mixture of direct and indirect gaps. The sDpW ligand leads to an indirect band gap. These ligands, through their rich combinations of sW/D and pW/D capabilities, modulate the band edge energies in an enormous range of 6 eV. Empirical functions are developed for the quick prediction of band edge energies of silicanes using the readily obtainable sW/D and pW/D indices of ligands. The explanation of the indirect-to-direct gap conversion, the demonstration of the large range of band edge energies, and their predictive functions are unprecedented contributions of this work. The wide range of band gaps of the substituted silicanes can regulate the band edge absorption from the near-infrared to the visible spectrum. The wide range of band edge energies allow for all three types of hetero-stackings between silicane species: straddling, staggered, and



broken. Silicenes with sDpW ligands also feature far separated conducting holes and electrons in both real and reciprocal spaces, which are likely to have longer lifetime decays of the excited charge carriers. These findings will inspire extensive applications of silicene-based materials in optoelectronics and photocatalytic applications.

We performed TDDFT calculations to assess the applicability of the one-particle orbital description for band edge excited states in silicenes. Excited states with degenerate hole and degenerate particle orbitals feature ~ 0.1 eV splitting, which quantifies the inaccuracy of the one-particle description. This inaccuracy is understandable since the splitting is induced by explicit electron correlation, which is pronounced when hole and particle orbitals are spatially confined. On the other hand, the one-particle description is satisfactory in explaining the band edge bright excited state splittings induced by orbital level splittings. The different ligands lead to different splittings of the frontier orbital levels and the associated different splittings of the band edge bright states, and this is attributed to the different π interactions between the ligands and the Si framework. Detailed analyses of frontier orbitals in the extended systems are indispensable to unravel the mystery in those splittings that can or cannot be described by the one-particle picture. Another methodological contribution is the justification of the use of the C-adapted σ/π -donating/accepting indices proposed by Ozimiński and Dobrowolski in silicon chemistry. We calculated the Si-adapted indices, and they are in close agreement with the C-adapted indices. Despite the close agreement, the Si-adapted indices are presented for future prudent examinations of ligand effects in silicon chemistry.

Author contributions

Conceptualization, G. Y., T. Z., Z. Y.; model construction and calculation, G. Y.; results analysis G. Y., T. Z., Z. Y., and E. P.; writing: G. Y., T. Z., and Z. Y.; funding acquisition, Z. Y., T. Z., and G. Y.; all authors read and agreed to the published version of the manuscript.

Data availability

The data supporting this article have been included as part of the ESI.† The codes for generate unit cell's k vectors for orbitals obtained from supercell calculation performed on Γ point only, and for numerical calculation of 2-electron integral for orbitals in cube format are available on request.

Conflicts of interest

There are no conflicts to declare.

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