

Cite this: *Mater. Horiz.*, 2025, 12, 3073Received 7th August 2024,
Accepted 10th January 2025

DOI: 10.1039/d4mh01038h

rsc.li/materials-horizons

Why does silicon have an indirect band gap?[†]

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It is difficult to intuit how electronic structure features—such as band gap magnitude, location of band extrema, effective masses, etc.—arise from the underlying crystal chemistry of a material. Here we present a strategy to distill sparse and chemically-interpretable tight-binding models from density functional theory calculations, enabling us to interpret how multiple orbital interactions in a 3D crystal conspire to shape the overall band structure. Applying this process to silicon, we show that its indirect gap arises from a competition between first and second nearest-neighbor bonds—where second nearest-neighbor interactions pull the conduction band down from Γ to X in a cosine shape, but the first nearest-neighbor bonds push the band up near X, resulting in the characteristic dip of the silicon conduction band. By identifying the essential orbital interactions that shape the conduction band, we can further rationally tune bond strengths to morph the silicon band structure into the germanium band structure. Our computational approach serves as a general framework to extract the crystal chemistry origins of electronic structure features from density functional theory calculations, enabling a new paradigm of bonding-by-design.

Introduction

Silicon has an indirect band gap, with the valence band maximum (VBM) at the Γ point and the conduction band minimum (CBM) at a low-symmetry point $\sim 85\%$ of the way between the Γ and X points. This indirect band gap determines the essential electronic and optical properties of silicon, and thereby its performance in photovoltaic and electronic devices.^{1–3} Although the low-symmetry CBM of silicon is a basic fact of semiconductor

New concepts

Electronic band structure is represented in reciprocal space, but arises from the chemical bonds between atoms in real space. Because bonding environments in crystals are so complex, it has been difficult to isolate which specific atomic orbitals contribute to shaping certain band structure features. For this reason, design and discovery of new semiconductors typically proceeds through a ‘needle-in-a-haystack’ approach, where high-throughput DFT screening approaches are guided by simple and limited chemical heuristics. Here, we present a new conceptual and computable framework to extract chemical bonding insights from DFT-calculated band structures, enabling us to rigorously and intuitively trace the impact of individual crystal bonds on band structure features. As a key example, we explain here the low-symmetry conduction band minimum of silicon, which profoundly impacts its properties for use in photovoltaics and electronics. Even in this basic semiconductor material, our approach leads to new insights to understand and engineer its conduction band minimum position. These calculation techniques can be broadly applied to reveal the crystal chemistry origins of electronic structure features in other optical, electronic and magnetic materials.

physics, it is not so simple to answer why silicon has an indirect band gap. Similar ‘why’ questions can be generally raised about the electronic structures of materials. Why does germanium have a CBM at the L point, despite also being a group IV semiconductor in the diamond structure? Why does zinc blende GaAs have a direct band gap, with such a light electron effective mass? Without a conceptual framework to approach why questions, one must rely on simple heuristics, which may post-rationalize the chemical origins of band structure but fall short in accurately predicting band structure features. Consequently, the search for next-generation thermoelectrics, p-type transparent conducting oxides, topological insulators, and other advanced electronic materials^{4–6} must proceed by brute-force screening *via* a “needle-in-a-haystack” approach,^{7–9} rather than by rational and intuitive design.

Roald Hoffman presented a beautiful theoretical framework to examine how physics and chemistry meet in the solid-state,^{10–12} arguing that chemists approach electronic structure from a bottom-up linear combination of atomic orbitals

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[†] Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d4mh01038h>



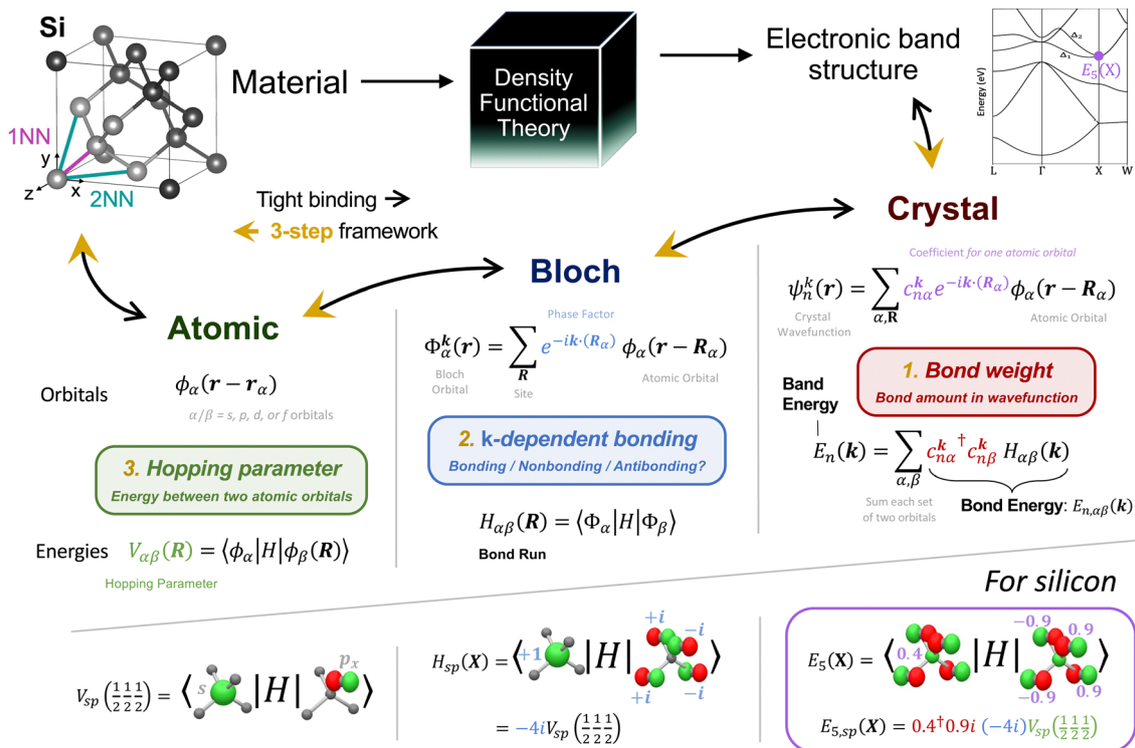


Fig. 1 Tight-binding offers a pathway from crystal chemistry to band structure. Our 3-step framework explains the reverse path—how to interpret a DFT electronic structure with the chemical understanding of tight binding. In the ‘for silicon’ section bright green and red is used for positive and negative isosurfaces for example atomic orbitals, Bloch orbitals, and crystal wavefunctions.

(LCAO). Alternatively, physicists adopt a top-down plane-wave interpretation of electronic structure, often using density functional theory (DFT) for accurate band structure calculations. The tight-binding (TB) model is a periodic version of LCAO that offers a bridge from bonds to bands, and back again.¹³ If a band in the electronic structure is dominated by a single orbital interaction, tight-binding offers a conceptual pathway to interpret how bonding in the wavefunction modulates across reciprocal space, thus explaining the band’s shape. However, when bands in 3D materials are formed by multiple orbital interactions, from multiple neighboring atoms, it becomes difficult to deconvolute how specific orbitals conspire to shape a band structure feature.

Given this complexity, creating tight-binding (TB) models in 3D structures requires a fundamental trade-off between simplicity and interpretability. Simple tight-binding models are usually constrained to first nearest neighbors (1NN), where one asserts *a priori* which bonds are considered. While the resulting models are usually easy to interpret, they may not be physically robust. In the case of silicon, achieving a CBM near the X point using 1NN tight-binding has relied on including an additional s^* and/or d states, which have limited physical relevance as they may not describe the precise physics of the actual excited states.^{14–22} On the other hand, tight-binding models that consider further atomic neighbors, such as by interpolation from DFT band structures or fitting with many-NN, increases the accuracy of a TB model but the model combinatorially

explodes in the number of terms—precluding chemical interpretability.^{23–35}

This paper aims to bridge the chemical intuition of Hoffmann and the practical toolkit of DFT, so that we can better interpret the chemical origins of electronic structure in real materials. To do so, we start from a tight-binding interpolation of the DFT-calculated electronic structure using maximally localized Wannier functions (MLWFs).³⁶ From this tight-binding interpolation, we chemically interpret how band shapes in k -space derive from orbital interactions in real space. We present a three-step process, illustrated in Fig. 1, which proceeds by determining: (1) which orbitals contribute to a band—specifically, what are the orbital characters (coefficients) that contribute to the wavefunction. (2) How orbitals bond across k -space—where the k -dependent phase ($e^{i\mathbf{k} \cdot \mathbf{R}}$) of each orbital changes the bonding/antibonding/non-bonding interactions between the orbitals in real space. (3) How strongly the orbitals bond—determined by the magnitudes of the TB hopping parameters. These three steps systematically sieve through hundreds of TB parameters to build a sparse and physically robust model to interpret chemical bonding contributions to the electronic structure.

Here, we begin with an illustrative 1D model system to emphasize the impact of multi-orbital and $>1\text{NN}$ interactions on band structure. Then we apply our three-step process to build a concise and chemically robust TB interpretation for how multiple orbital interactions combine to form a low-symmetry CBM in silicon. Specifically, we find that this low-symmetry



CBM along the Γ -X line derives primarily from a cosine shape from second nearest neighbor (2NN) p_x - p_x bonds, combined with a linear shape near X from 1NN orbital bonds. Finally, we present a new computational tool to interactively execute our three-step process for any band and k -point in an electronic band structure (<https://viz.whsunresearch.group/tb/>). This tool enables us to precisely identify which bonds affect which segments of the silicon band structure. We then rationally tune these orbital interactions to morph the silicon band structure towards the germanium band structure, in accordance with the actual chemical differences between Si and Ge. Altogether, our work serves as a general blueprint to extract the crystal chemistry origins of electronic band structure, and provides a pathway for rational band-structure engineering by chemical and structural design.

Uncovering the chemistry in band structure

All electronic properties of a material—including band gap, effective mass, band extrema location, *etc.*—are characterized by its band structure. The chemical origin of these electronic properties can be elucidated from accurate TB interpolations of DFT-calculated electronic structure. Tight-binding decomposes a band structure into a summation of terms, similar to a Fourier decomposition, but with basis functions that correspond to physically-relevant orbital interactions.³⁷ This decomposition enables one to trace back which bonds manifest which specific band features of interest. Fig. 1 offers a hierarchical mind-map that captures the tight-binding pathway from crystal chemistry to the band structure.

The central assumption of tight binding is to write the crystal wavefunction, $|\psi_n^k\rangle$, as a linear combination of atomic Bloch orbitals, Φ_α^k , weighted by the coefficient $c_{n\alpha}^k$ as in eqn (1). The indices k , n , and α represent k -point, band, and orbital. An atomic Bloch orbital is the sum over atomic orbitals, ϕ_α , in each cell of the crystal related by a lattice vector translation \mathbf{R} . The complex phase of each atomic orbital is modified by the phase factor $e^{-ik\cdot(\mathbf{R}_\alpha)}$, where $\mathbf{R}_\alpha = \mathbf{r}_\alpha + \mathbf{R}$ and \mathbf{r}_α is the atomic orbital center in the primitive cell.

$$|\psi_n^k\rangle = \sum_\alpha c_{n\alpha}^k |\Phi_\alpha^k\rangle = \sum_{\alpha, \mathbf{R}} c_{n\alpha}^k e^{-ik\cdot(\mathbf{R}_\alpha)} |\phi_\alpha(\mathbf{R}_\alpha)\rangle \quad (1)$$

With an atomic decomposition of the crystal wavefunction, each band dispersion (shape in E - k space) expands as the combination of numerous pairwise bonds between atomic orbitals. In eqn (2), the overall shape of the band, $E_n(\mathbf{k})$ for band n , results from the sum of ‘bond energies’ $E_{n,\alpha\beta}(\mathbf{k})$ —which is the E - k shape of the bond between Bloch orbitals α and β .

$$\begin{aligned} E_n(\mathbf{k}) &= \langle \psi_n^k | \hat{H} | \psi_n^k \rangle = \sum_\alpha \sum_\beta c_{n\alpha}^{k\dagger} c_{n\beta}^k \langle \Phi_\alpha^k | \hat{H} | \Phi_\beta^k \rangle \\ &= \sum_\alpha \sum_\beta E_{n,\alpha\beta}(\mathbf{k}) \end{aligned} \quad (2)$$

This equation substitutes the crystal wavefunction with the middle expression in eqn (1). From this, the bond energy, $E_{n,\alpha\beta}(\mathbf{k})$, is the bond weight, $c_{n\alpha}^{k\dagger} c_{n\beta}^k$, multiplied by the bond run, which we name the Hamiltonian matrix element between

Bloch orbitals α and β , $\langle \Phi_\alpha^k | \hat{H} | \Phi_\beta^k \rangle$. The name ‘bond run’ is inspired by Hoffmann’s discussion¹⁰ that bands made from s orbital bonds ‘run up’ in energy from Γ to the Brillouin zone edge (from bonding to antibonding), while bands from p orbital bonds ‘run down’ from Γ to the Brillouin zone edge (from antibonding to bonding).

To isolate the impact of atomic orbital bonds, the bond run—written as $H_{\alpha\beta}(\mathbf{k})$ —is expanded by substituting the Bloch orbitals with the sum of k -modulated atomic orbitals, as in eqn (1). This results in eqn (3), which is a sum over bonds between an atomic orbital α at \mathbf{r}_α and atomic orbital β at $\mathbf{r}_\beta + \mathbf{R}$. Each interaction is then the TB hopping parameter, $\langle \phi_\alpha | \hat{H} | \phi_\beta(\mathbf{R}) \rangle \equiv V_{\alpha\beta}(\mathbf{R})$, multiplied by a factor which modulates the complex phase based on the phase difference between orbital centers for a given k .

$$\langle \Phi_\alpha^k | \hat{H} | \Phi_\beta^k \rangle \equiv H_{\alpha\beta}(\mathbf{k}) = \sum_{\mathbf{R}} V_{\alpha\beta}(\mathbf{R}) e^{ik\cdot(\mathbf{r}_\alpha - \mathbf{r}_\beta - \mathbf{R})} \quad (3)$$

Here, the k -dependence of a bond run arises from the sum of phase factors $e^{ik\cdot(\mathbf{r}_\alpha - \mathbf{r}_\beta - \mathbf{R})}$. When the bond run is negative for a given k , it indicates the Bloch orbitals α and β are overall bonding, whereas positive indicates net antibonding. While the phase of the coefficients $c_{n\alpha}^{k\dagger} c_{n\beta}^k$ must also be included to determine the precious bond type for each band, this complication is reserved for later when eqn (4) is introduced.

Our three-step process is grounded in these two fundamental equations, offering a way to back solve from a given band structure the chemical bonding contributions, as diagrammed in Fig. 1. Step 1 is bond weight, where $c_{n\alpha}^{k\dagger} c_{n\beta}^k$ quantifies the bonds (between orbitals α and β) that may contribute to the band. While step 2 is the shape of the bond run, which describes how the bond-type (bonding, antibonding, or non-bonding) changes across k -space. Step 3 is the maximum absolute energy of the bond, dictated by the hopping parameter between two atomic orbitals α and β , $V_{\alpha\beta}(\mathbf{R})$.

Eqn (2) is the same theoretical starting point as the crystal orbital Hamiltonian population^{38,39} method, but the implementation of an analytical representation in eqn (3) (instead of a numerical calculation, for example as done through LOBSTER⁴⁰) requires a tight-binding interpolation, which enables us to then break down the contributions from individual bonds. These analytical representations of each orbital wavefunction lets us visualize and further separate into NN or long-range interactions.

To examine how multiple orbitals and further nearest-neighbor interactions manifest in both real space wavefunction and the reciprocal space bands, here we present an illustrative example on a one-dimensional monatomic chain, with two orbitals (s and p) per atom. Traditionally, 2NN and further terms are neglected in simple TB models, but here we show how they could induce major qualitative changes to the band structure. The details of the derivation are in SI.1 (ESI[†]).

Plotted in Fig. 2a, the 1NN bond runs H_{ss} and H_{pp} are cosine curves with extrema at the high symmetry points, but all other bond runs (H_{sp} and 2NN bonds) have extrema at low symmetry points. Using eqn (2), we sum together the bond runs



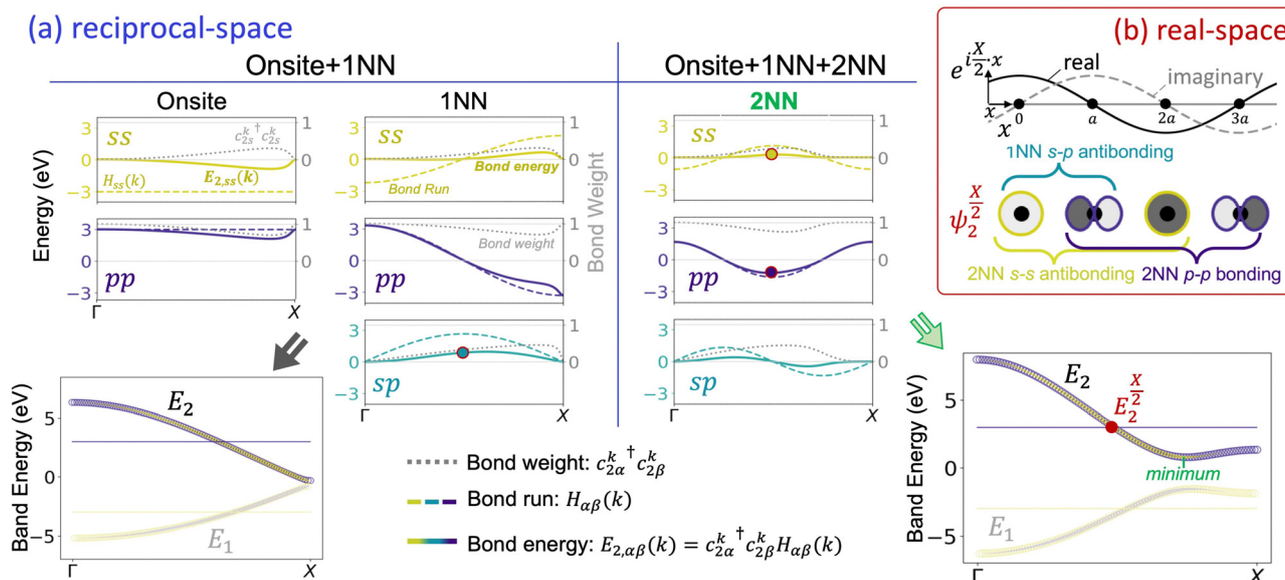


Fig. 2 Multiple bonds combine to form band energy with a monoatomic s + p 1D tight-binding model. When adding 2NNs, band extrema at low-symmetry k -points manifest. In (a), the bond runs, bond weights,⁴¹ and bond energies are plotted for onsite (orbital energy), 1NN, and 2NN bond contributions to the second band, E_2 . The bond energies sum to create the E_2 band energy, where the orange/blue circle size indicates orbital character, $|c_{s/p}^k|$. To make them real, $-iH_{sp}(k)$ and $ic_{sp}^k c_{2p}^k$ are plotted for the sp bonds. On the right, 2NN terms are added, perturbing the bond weights and band energies to create a low symmetry extremum off-X. In (b), the real part of the $\psi_2^{X/2}$ wavefunction is plotted to highlight the real-space bonding implicit in band structure. The orbital at each atom is determined by the phase factor, $e^{i(X/2) \cdot x}$, multiplied by the orbital coefficients, $c_s = 0.49$ and $c_p = -0.87i$. The nonzero bond energies are written and circled in red on the reciprocal-space plots.

multiplied by their bond weight (orbital character) to find the band energy. With only 1NN, $H_2(k)$ is dominated by $H_{pp}(k)$, which has extrema at high symmetry points. We next add 2NN interactions, with an exaggerated bond strength of half the 1NN, which results in a qualitative change in the overall band shape. This additional strong 2NN interaction pulls the bands together near $\frac{X}{2}$ while pushing them apart near Γ and X, creating a low-symmetry band extremum reminiscent of silicon.

When we ground our tight-binding intuition in 1D models, the 2NNs are so far away that their contributions tend to be small. However, in 3D crystals, atoms have much higher coordination numbers. These 2NNs are also much closer in 3D crystals than they are in 1D systems, meaning the 2NN contribution to the tight-binding interactions can be substantial. In the case of silicon, the twelve 2NNs are only $1.6\times$ further than the four 1NNs, with a geometry that allows for strong overlap between p orbitals. By studying silicon, we will show how bonds of similar strength with different frequency of bond runs leads to band extrema away from high symmetry points. Another common cause for low symmetry band extrema is an avoided crossing from s-p orbital mixing. However, avoided crossings will always result in a band inversion (switching of orbital character), which the silicon CBM does not exhibit. Thus, if there is a low symmetry band extremum on a band that does not have a band inversion, long-range interactions beyond 1NN are a likely culprit.

In our ambition to eventually design band structure from the underlying bonds, we need to first elucidate the orbital and

bonding nature at the specific k -point of a band. The key term linking real space chemical bonding with reciprocal space band structure is the phase factor $e^{ik \cdot R}$, manifesting in the crystal wavefunction of eqn (1). For a given k -vector, the combined coefficient $e^{ik \cdot R_\alpha} c_{n\alpha}^k$ for each atomic orbital α dictates the bond type (bonding, antibonding, or non-bonding) between atomic orbitals in real space (step 2). If the complex phases of neighboring orbitals are orthogonal (e.g. real and imaginary), they do not interact and are non-bonding. If they are nonorthogonal (e.g. real and real, or imaginary and imaginary), they are bonding or antibonding, depending on the signs of the wavefunction.

In Fig. 2b, we illustrate the relationship between phase factor and real-space $\psi_2^{X/2}$ wavefunction at $k = X/2$, which requires a 4-unit cell superstructure in real space. Across the 4 atoms in Fig. 2b, the phase factor $e^{i(X/2) \cdot R}$ modulates as $+1$, $+i$, -1 , and $-i$. Because c_s^k coefficients are real and positive for s orbitals, whereas they are imaginary and negative for p orbitals, we see that the real part of $\psi_2^{X/2}$ has s orbitals on atoms 1 and 3; and p orbitals on atoms 2 and 4. Therefore, the only 1NN interaction is s-p antibonding. The phase sign switching between atoms 1 (2) and 3 (4) yields 2NN s-s antibonding (2NN p-p bonding). This example visualizes how chemical bonding in real space implicitly derives from each k -point in reciprocal space.

Detangling the silicon band structure

Although silicon has been studied for decades, the crystal chemistry origins of its low-symmetry conduction band



minimum still lack satisfactory explanation. Tight-binding (TB) models fitted with only 1NN incorrectly produce a conduction band minimum at Γ . While Vogl produced a CBM off-X with an additional excited s^* state, he acknowledges that “the inclusion of some such excited states in any minimal basis set is physically important—although the precise physics of the actual excited states need not be faithfully and quantitatively reproduced.”¹⁶ Indeed, while the sps^* model fits the Γ -X line, it sacrifices the accuracy of the rest of the conduction band structure along nearly every other k -path, as detailed in SI.2 (ESI[†]).

Since band structure arises from the complex interactions between multiple orbitals, it is often possible to have multiple non-unique solutions that fit a singular band feature. Following Vogl, others have included additional orbital states to their TB models—for example Jancu *et al.* and others added d states, producing a reasonable fit of the lowest conduction bands but again with little physical insight regarding the additional parameters.^{17–22} Tight-binding models with >1 NN were also fit (often with s^* states) for silicon and zinc blende semiconductors with varying degrees of 2NN contribution.^{23–32,35}

From a model-building perspective, it is not satisfactory to include terms *ad hoc* just to match a single band structure feature—rather, a term that is physically valid should improve the fit of all band energies throughout the entire Brillouin zone. This is especially important if one aims to later engineer and design the band structure by modifying chemical interactions, which requires one to accurately identify the true chemical origin of band features.

At the other end of the spectrum, one can perform a TB interpolation directly from DFT—which obtains the hopping parameters from a Fourier transformation of the k -dependent orbital Hamiltonian. This TB interpolation circumvents the need to assume which interactions are present, but the resulting many-NN TB models can have hundreds of non-trivial interactions, which is too complex to interpret chemically. Sanchez-Portal, and later Qian *et al.*, applied a TB interpolation which includes many NNs to silicon finding a low symmetry minimum along the Γ -X line with only a sp basis, indicating that s^* and d states are not strictly required to reproduce the minimum off X.^{33,42} Since then, TB interpolations of silicon are frequently achieved using MLWFs and similar methods, but a simple chemical understanding has not yet been detangled from the hundreds of hopping parameters found.

Here, we apply our three-step process to build a chemical interpretation for the conduction band minimum in silicon along the Δ_1 band. Our DFT calculations were done with the Vienna *ab initio* Simulation Package using Perdew–Burke–Ernzerhof pseudopotentials, a plane wave energy cutoff of 520 eV, and k -point density of 0.23 \AA^{-3} .^{43–45} Details of our MLWF parameters and process are discussed in SI.3 (ESI[†]). Our 3-step process is implemented as follows: (step 1) we determine which orbitals contribute to the Δ_1 band—finding that it is $>80\%$ p_x orbital character, with the remaining character being s orbitals. (Step 2) we determine how the orbitals bond across k -space—finding from the bond runs that the second nearest

neighbor p_x - p_x is the only interaction that decreases the band energy at the X point. (Step 3) we determine how strongly the orbitals bond—showing that the 2NN p_x - p_x bond has large hopping parameters and high coordination which makes it a significant influence on the band structure.

Finally, individual chemical bonds are assessed for their contribution to the total shape of the Δ_1 band. From this, we determine that the low-symmetry conduction band minimum of silicon manifests from a linear shape of the 1NNs near X, combined with the cosine shape of the 2NN p_x - p_x bond. Crucially, including the 2NN p_x - p_x bond not only improves the Δ_1 band, but the band structure across all other high-symmetry lines (details in SI.2, ESI[†]), validating its physical significance in creating the low-symmetry Si CBM.

(1) Orbital character of bands. First, we determine which orbitals in the Δ_1 band are present to bond. In a sp model, silicon in the diamond structure has eight orbitals, four for each of the two atoms in the primitive cell. This amounts to 72 Hamiltonian matrix elements—8 onsite, 32 1NN, and 32 2NN interactions. After symmetry and group theory considerations, a wavefunction along the Γ -X line will have either $s + p_x$ orbitals or $p_y + p_z$ orbitals. With only s and p_x orbitals are on the Δ_1 band, the 72 matrix elements can be reduced to 8 unique elements. To separate the character of a general band, it is essential to use atomic orbitals as the momentum-dependent crystal wavefunctions rarely reduce to the hybrid atomic orbitals of simple molecular wavefunctions.

Fig. 3 plots positive (red) and negative (green) isosurfaces for the real part of the complex wavefunctions in one of each of the four doubly-degenerate bands at X. For all the X_1 bands, the first atom in the basis set has only p_x orbitals and the second atom has only (distorted) s orbitals. Away from X, the X_1 degeneracy splits into Δ_1 (CBM band) and Δ_2 bands with s and p_x orbital character, while the X_4 bands remain degenerate. In silicon, tight-binding analysis of the Δ_1 conduction band character shows it is predominantly ($>80\%$) p_x orbital character. The X_1 antibonding wavefunction is mainly 1NN s - p_x antibonding and 2NN p_x - p_x bonding. Importantly, the 2NN p_x - p_x interaction is the only one that is bonding along the x -direction and contributes to lowering the energy at X.

Although these arguments explain the lowering of the Δ_1 band energy at X, the actual CBM is at a low-symmetry point $\sim 85\%$ of the way from Γ to X. In the next two steps, we examine the shape (step 2) and magnitude (step 3) of the participating bonds. Then in the final section, we combine multiple bonds to achieve the Δ_1 band dispersion and observe how the minimum off-X manifests.

(2) Shape of bonds in k -space. Second, we uncover how bonding changes with k -space. The real-space bond type can be visualized in the full wavefunctions by mapping the k -dependent phase modulations onto the atomic orbitals, as in Fig. 2b. For the sake of brevity, we have included this visualization and discussion in SI.4 (ESI[†]). In this section, we focus on how the real-space bonding manifests as higher or lower energies in reciprocal space, forming the shape that each bond has along the Δ_1 band.



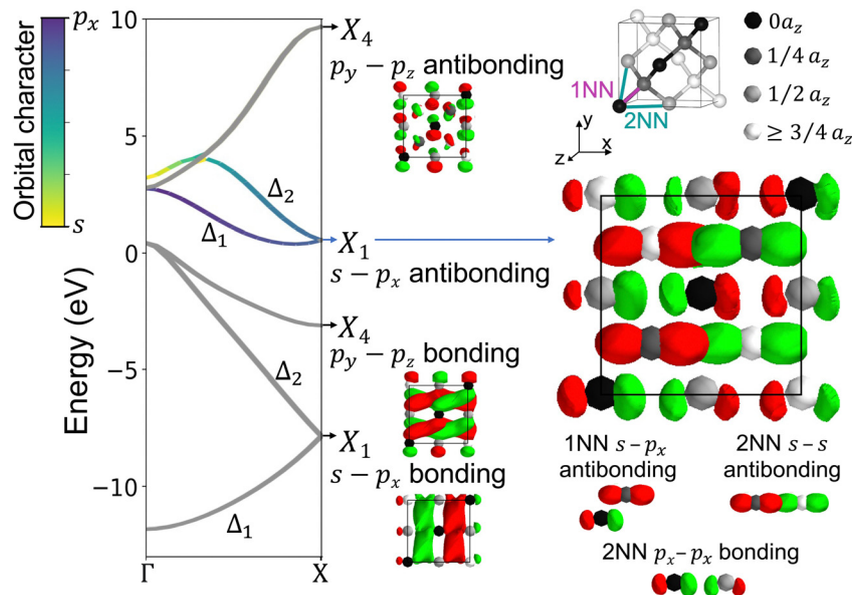


Fig. 3 The crystal wavefunctions at the X point in silicon with the X_1 conduction band highlighted to show orbitals and bond type. Each of the four doubly degenerate bands is accompanied by the present 1NN bond and the plotted real part of a wavefunction. The atom sites are spheres colored to indicate the z-coordinate. The red and green show the positive and negative isosurfaces of the real wavefunction, where neighboring same color lobes are bonding and different color lobes are antibonding. Bonding lobes often mesh together while antibonding lobes are distorted apart. By looking closer at the X_1 band, we determine 1NN and 2NN bond-types, where the 2NN p_x - p_x interaction is the only bonding along the x-direction.

As discussed in the first section, a bond's contributed shape is referred to as the bond energy, which is the bond run multiplied by bond weight. Because the bond run defined in eqn (3) can be any phase and does not vary with band, here we introduce a band-dependent bond run $H_{n,\alpha\beta}^R(k)$, eqn (4). This includes the phase of the orbital coefficients for band n , ensuring that $H_{n,\alpha\beta}^R$ is real and the correct sign for band n . The superscript R is included to indicate that only the primitive cell vectors for onsite, 1NN, or 2NN bonds are included during the sum over R in eqn (3). Finally, using eqn (2) and (4) we write the bond energy, $H_{n,\alpha\beta}^R(k)$, as band-dependent bond run multiplied by the absolute value of the orbital coefficients, eqn (5).

$$H_{n,\alpha\beta}^R(k) \equiv \frac{c_{nz}^k \dagger c_{n\beta}^k}{|c_{nz}^k| |c_{n\beta}^k|} H_{\alpha\beta}^R(k) \quad (4)$$

$$H_{n,\alpha\beta}^R(k) = |c_{nz}^k| |c_{n\beta}^k| H_{n,\alpha\beta}^R(k) \quad (5)$$

In the top of Fig. 3, we plot the band-dependent bond runs $H_{\Delta_1,\alpha\beta}$ and the bond energies $H_{\Delta_1,\alpha\beta}$ for the onsite, 1NN, and 2NN interactions. The bond run shapes are similar to the illustrative 1D example from Fig. 2a, but with half the length in reciprocal space since silicon has a two-atom primitive cell. The bond run magnitudes are dictated by the hopping parameter. The onsite orbital energy terms are most simple, for example the onsite p_x term is $E_{\Delta_1,xx}^0 = \epsilon_p |c_{\Delta_1,x}^k| |c_{\Delta_1,x}^k|$, where ϵ_p is the p_x -like orbital energy, $c_{\Delta_1,x}^k$ is the p_x orbital coefficient. The 1NN and 2NN terms require more derivation, which is left to SI.5 (ESI†). Most importantly, the 2NN p_x - p_x orbital bond energy is

$E_{\Delta_1,xx}^{2NN} = |c_{\Delta_1,x}^k| |c_{\Delta_1,x}^k| \cdot [8V_{xx}(110) \cos(k_x\pi) + 4V_{xx}(011)]$, the right part of which is the bond run $H_{\Delta_1,xx}^{2NN}(k)$, which reveals how the 2NN p_x - p_x interaction changes bond-type throughout the Δ_1 band. Crucially, the dominant 1NN s-s and s-p bonds are higher in energy at X than Γ , encouraging a CBM at Γ . Whereas the positive cosine in the bond run for the 2NN p_x - p_x interaction lowers the energy at X, enabling a minimum near X.

(3) Strength of hopping parameters. Third, we examine the magnitude of atomic orbital interactions using hopping parameters. Based on intuition from 1D models, we would anticipate the hopping parameters for 1NNs to generally be much larger than for 2NNs. However, here we find that the 2NN bonds are very important in silicon, as supported by the hopping parameters from our atomic-like MLWF tight-binding interpolation in Table 1. When including bond multiplicity, the twelve 2NN p_x - p_x parameters sum as $8V_{xx}(110) + 4|V_{xx}(011)| = 2.00$ eV, which is $4\times$ larger than the four 1NN p_x - p_x parameters $V_{xx}\left(\frac{111}{222}\right)$.

Here, $V_{xx}(110)$ indicates the hopping parameter between p_x - p_x orbitals on atoms separated by the vector $1\mathbf{x} + 1\mathbf{y} + 0\mathbf{z}$ (or a symmetrically equivalent vector), where the Cartesian \mathbf{xyz} unit vectors are half the unit cell length. Combined with the dominant orbital character being p_x , the 2NN p_x - p_x contributes significantly to the energy of the Δ_1 wavefunctions. Other tight-binding models which have included 2NN parameters either did not include a $V_{xx}(110)$ term, or they were $\sim 10\times$ smaller than our MLWF-derived result.^{24–32} An exception to this is Grosso & Piermarocchi who fit a $V_{xx}(110)$ about $2\times$ larger than our result in Table 1.³⁵ In all cases, the 2NN contributions were not individually analyzed for their role in shaping the CBM.



Table 1 Silicon hopping parameters (eV) from MLWF used to reconstruct the Δ_1 band. The s and p orbital onsite terms are ϵ_s and ϵ_p . Importantly, the 2NN terms $V_{xx}(110)$ and $V_{xx}(011)$ are similar or larger than the 1NN term $V_{xx}\left(\frac{111}{222}\right)$

ϵ_s	ϵ_p	$V_{ss}\left(\frac{111}{222}\right)$	$V_{sx}\left(\frac{111}{222}\right)$	$V_{xx}\left(\frac{111}{222}\right)$	$V_{xx}(110)$	$V_{xx}(011)$
-5.467	1.650	-1.639	1.075	0.126	0.117	-0.267

The small hopping parameter between 1NN p_x - p_x orbitals can be understood from the geometry as discussed by Slater.⁴⁶ The hopping parameter between two p_x orbitals is given by $l^2 V_{pp\sigma} + (1 - l^2) V_{pp\pi}$, where l is the direction cosine in the x direction. If the p_x - p_x lobes are facing each other (like ∞ - ∞) then there is perfect σ bonding, where $l = 1$, whereas if p_x - p_x lobes are parallel (like 8-8) then $l = 0$ and there is perfect π bonding. Because $V_{pp\sigma}$ and $V_{pp\pi}$ have opposite sign, an intermediate orientation between perfectly aligned (∞ - ∞) and perfectly parallel (8-8), will result in the hopping parameter canceling to zero.

In the tetrahedral coordination environment of the diamond structure, 1NN have $l^2 = 1/3$, such that the $V_{pp\sigma}$ and $V_{pp\pi}$ components nearly cancel. The 2NN have four neighbors with $l^2 = 0$, meaning the $V_{xx}(110)$ parameters are entirely π bonding, while the other eight neighbors have $l^2 = 1/2$, allowing the stronger σ antibonding to dominate the weaker π bonding in the $V_{xx}(110)$ parameters.

Steps 1 + 2 + 3: the low-symmetry Δ_1 minimum in silicon

Finally, we create the total shape of the silicon Δ_1 band dispersion from individual bonds and determine which bonds are necessary to capture the correct qualitative band shape. Each bond contributes a distinct shape over some high-symmetry line of the band structure. To obtain the correct band dispersion that matches a DFT band structure, all significant bonds must be included. As Vogl showed, a 1NN tight-binding model with an sp basis cannot produce a low-symmetry conduction band minimum,¹⁶ whereas a many-NN sp basis can.^{33,42} This indicates that physically significant bonds are missing from the 1NN sp model, which as we have argued, are the 2NN interactions.

The Δ_1 band energy as a function of the bonds can be simplified from the sum over each matrix element, eqn (2), to a sum over unique elements for the s and p_x (written as x) orbitals, eqn (6). As discussed earlier, this reduces the 72 parameters for the Γ -X line to only 8: two onsite (orbital energy) terms, three 1NN terms, and three 2NN terms.

$$E_{\Delta_1}(k) = 2E_{\Delta_1,ss}^0 + 2E_{\Delta_1,xx}^0 + 2E_{\Delta_1,ss}^{1NN} + 4E_{\Delta_1,sx}^{1NN} + 2E_{\Delta_1,xx}^{1NN} + 2E_{\Delta_1,ss}^{2NN} + 4E_{\Delta_1,sx}^{2NN} + 2E_{\Delta_1,xx}^{2NN} \quad (6)$$

Each term has an analytical expression as seen from eqn (5), which is the product of relevant orbital coefficients (a multiplicative factor) with the band-dependent bond run (a cosine or

sine shape). For full derivation and decomposition of eqn (6), see SI.5 (ESI[†]).

To conceptually understand how the CBM arises from multiple orbital interactions, we compare and combine the relevant bond runs and energies in the top panels of Fig. 4 to create the bottom panel. When looking at the bond runs, the onsite and 1NN terms dominate the 2NN, with the 1NN s- p_x and s-s spanning ~ 8 eV each, while the 2NN p_x - p_x only reaches 2 eV. But once the strong p_x orbital character is included with $E_{\Delta_1,\alpha\beta}$, the p_x - p_x terms are nearly unchanged, while the s- p_x decrease significantly, and the s-s drops nearly to zero. This puts the [onsite + 1NN] energy magnitude in the same range as the 2NN p_x - p_x , where each span ~ 1.5 eV (Fig. 4 bottom panel). The shape of onsite + 1NN is a pseudo-linear increase near X, which results primarily from the $E_{\Delta_1,ss}^{1NN}$ shape. The cosine-like shape of $E_{\Delta_1,ss}^{1NN}$ results from the sine curve of $E_{\Delta_1,ss}^{1NN}$ being heavily distorted by the coefficient weight $|c_{\Delta_1,s}|/|c_{\Delta_1,p}|$ increasing from Γ to X.

Finally, the pseudo-linear shape of the onsite + 1NN near X plus the cosine curve of the 2NN p_x - p_x combine to form the conduction band minimum away from the high-symmetry X point in silicon. Importantly, including the 2NN p_x - p_x bond also provides a good band structure fit on all other k -paths (see SI.2 for details, ESI[†]), compared to the s^* state from Vogl *et al.*,¹⁶ which validates the physical importance of 2NN bonding in governing the low-symmetry conduction band minimum of silicon.

Towards bonding-by-design

Band engineering for solar cells, semiconductors, and thermoelectrics frequently requires control over the energy level of bands at specific k -points. Because we now have a theoretical pathway to connect the bonding interactions to the band structure, we can examine the inverse electronic structure design problem—How can I modify chemical interactions to morph an existing band structure to a new band structure with more desirable features? As a representative example, here we will modify the bonding interactions to shift the CBM from the Γ -X line in silicon to the L point as it is in germanium. This illustration paves the way to a vision of bonding-by-design, where instead of searching for pristine materials with a given band structure feature, we can rationally tune the chemistry (by substitutional doping or alloying) to morph a given band structure towards a desired one.

Thoroughly analyzing a band structure feature is arduous, which motivated us to create a computational analysis package (<https://viz.whsunresearch.group/tb/>) which systematically executes our three-step process. Our package features an interactive interface that populates tables with the orbital character and important bonds for any selected point of the $E(k)$ diagram. In addition, the band-dependent bond runs and bond energies for any of the important bonds can be plotted upon selection, allowing a user to rapidly discern how each bond contributes to the band shape. A detailed explanation and tutorial are provided in SI.6 and SI.7 (ESI[†]). Our new utility marks an improvement from tools which plot the atomic orbital character on



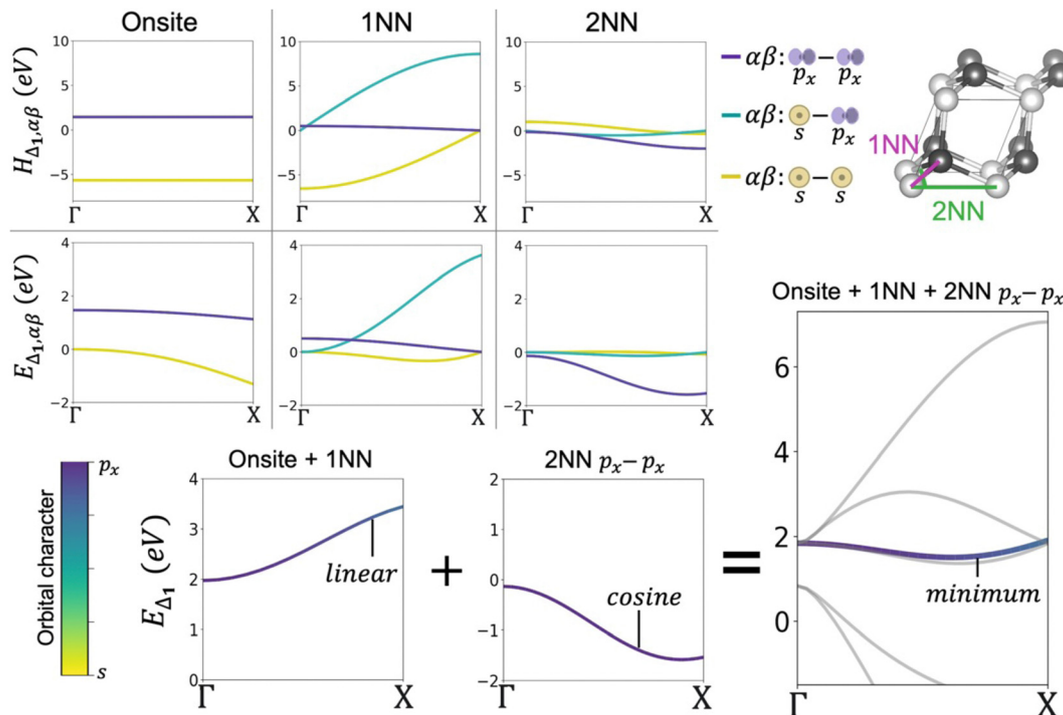


Fig. 4 Deconstructing how each bond contributes to the Δ_1 conduction band in silicon by plotting the band-dependent bond runs $H_{\Delta_1,ij}$ and corresponding energies $E_{\Delta_1,ij}$ for the onsite (atomic orbital energy), 1NN, and 2NN bonds. Interactions between p_x - p_x , s - p_x , and s - s orbitals are colored purple, teal, and yellow, respectively, which corresponds with the orbital character color bar used in the total energy plots H_{Δ_1} . Altogether, the linear behavior near X achieved with onsite + 1NN bonds and the cosine shape of the 2NN p_x - p_x bond combine to form the minimum near X. The gray lines in the bottom right plot show the silicon band structure with all onsite, 1NN, and 2NN bonds, where the small error between the colored and gray Δ_1 band results from including the 2NN s - p_x and s - s bonds.

band structure^{47,48} (only the first step in our 3-step process) by bringing the band energy interpretability of the Extended Hückel method^{49,50} to the higher accuracy of density functional theory. Here we use our package to demonstrate how different segments of a band can be selectively raised or lowered towards a desired shape by modifying a single bond.

Fig. 5a demonstrates a lower off-X minimum is achieved by strengthening the 2NN p_x - p_x hopping parameters which, as described previously, contributes a shifted cosine shape to the

Δ_1 band. As k -point moves along the Γ -L line in the [111] reciprocal-space direction, the orbitals of the nearest-neighbor atom in the (111) real-space direction change phase in accordance with $e^{ik \cdot R}$. The change in phase facilitates bonding at the L point between the p - p orbitals of atoms with a displacement along (111). Thus, in Fig. 5b the L point is lowered by strengthening this 1NN p - p hopping. The lowest conduction band at Γ is an antibonding electron wavefunction of entirely s orbital character. Thus, in Fig. 5c the Γ point is lowered by weakening

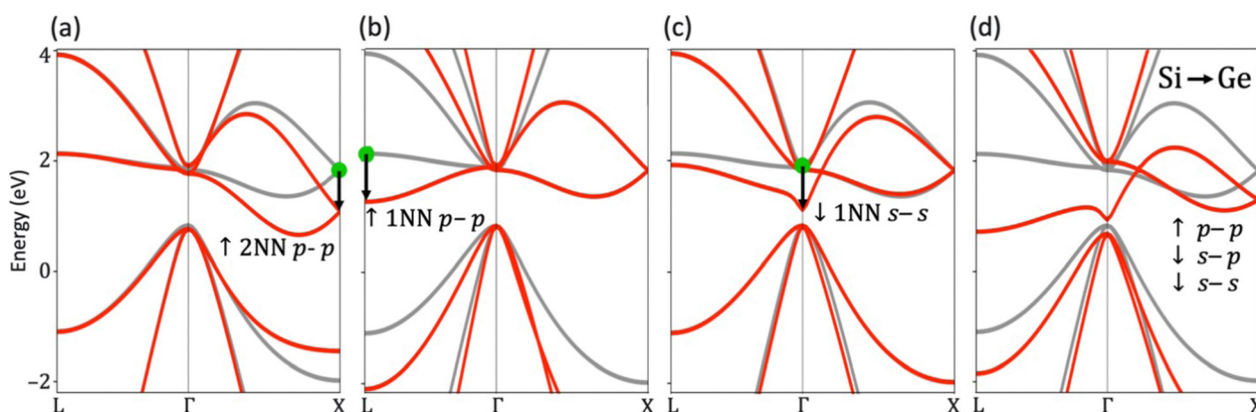


Fig. 5 The role of chemical bonding in band extrema. In (a)–(d) the original Si band structure (gray) is plotted against augmented band structures (red), which changes one bond to selectively lower X, L, or Γ (plots (a), (b), and (c), respectively) or changes bonds based on the chemistry of Ge (plot (d)).



the 1NN *s*-*s* hopping parameters. Notably, the *s* character dramatically decreases from 100% to ~50% at only 0.1 L along the Γ to L line, which creates the sharp curvature near Γ and leads to low effective masses in direct-gap tetrahedral semiconductors like GaAs.

Altogether, it is possible to morph the Si band structure towards the Ge band structure by increasing bonding between *p* orbitals while decreasing bonding of *s* orbitals. This effect is qualitatively consistent with changing chemistry from Si to Ge. In Ge, the occupied *d* shell incompletely screens nuclear charge which attracts the valence shell—an effect called scandide contraction. The additional density near the nucleus in *s* orbitals is disproportionately impacting by the poor screening, thereby reducing the *s* orbital radius of Ge compared to the *p* orbital radii. Yuan *et al.* also found that the *d* orbitals are important in changing CBM location,⁵¹ which we find results from the indirect screening effects of *d* orbitals on *s* and *p* orbitals. Thus, when augmenting Si in Fig. 5d by increasing all 1NN *p*-*p* interactions by 30% and decreasing all 1NN *s*-*p* and *s*-*s* bonds by 15%, we reproduce the characteristic band structure of Ge with a CBM at the L-point. Further discussion of the Si and Ge MLWF band structure can be found in SI.8 (ESI[†]).

The theoretical framework and automated analysis package developed in this manuscript can readily be applied to any tight-binding (TB) model. While our package does not yet directly support spin-orbit coupling, since it is not relevant in silicon, incorporating spin-orbit coupling is theoretically straightforward. One would decompose the bands into bonds that specify both the orbital and spin of each electron, where spin-orbit coupling driven changes would manifest in the interaction between orbitals of different spin.

While our framework is generally applicable, it is limited by whether a reliable TB model can be generated from DFT. For silicon, we have justified the use of maximally localized Wannier functions (MLWFs) to generate a TB model (justification in SI.3, ESI[†]). However, analysis using MLWF may not always yield chemically interpretable results. When the Wannier functions deviate too much from atomic orbitals—becoming combinations of multiple atomic orbitals across several atoms—our three-step process cannot elucidate the atomic origins of electronic structure features. Instead, it reflects the behavior of these hybridized

Wannier functions. Despite these challenges, our method holds promise for application to a broader range of materials. Future work will focus on extending our framework to incorporate spin-orbit effects and developing strategies for constructing suitable TB models in complex systems.

Outlook

Here we presented a computable and chemically motivated framework that considers (1) Which orbitals are in a band, (2) How are they allowed to bond, and (3) How strongly do they bond? This framework produces a sparse and therefore interpretable tight-binding model that can help us intuitively understand the crystal chemistry origins of band structure. When we applied our approach to silicon, we found that the low-symmetry conduction band minimum of silicon originates primarily from 2NN *p_x*-*p_x* bonds, which significantly lowers the energy at X. The significance of the 2NN *p_x*-*p_x* orbital bond compared to the 1NN is explained from the geometry of the bonding angles, in addition to there being 3× as many 2NN atoms than 1NN. This explanation is a revision on Vogl's *sps** model, which captures the CBM position in silicon but at the expense of other conduction bands in the Brillouin zone. Our interpretation provides a clear physical mechanism compared to previous *sp* models with multiple NN.

Broadly speaking, our approach allows us to pinpoint the physical origin of electronic structure features in complex 3D crystals. This framework is general and can be applied to any tight-binding interpolation of a DFT-calculated band structure. By better understanding how crystal chemistry translates to major electronic structure features, we can more intuitively design chemistries and bonding environments to yield a desired band structure feature. A major advantage of this approach is the opportunity to search within the 'perturbation space' of a given material, allowing us to find best-in-class semiconductors which are often minor perturbations (strain, doping, alloying, *etc.*) from their pristine forms. This approach would invert the design paradigm from electronic 'materials-by-design' to the inverse approach of bonding-by-design (Fig. 6)—where instead of searching for materials with specific properties, we can chemically or structurally modify the band

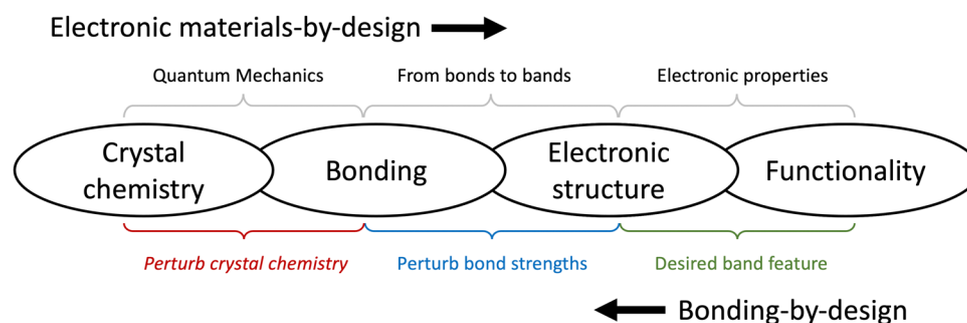


Fig. 6 In materials-by-design, the electronic properties of pristine materials are calculated from DFT. By inverting this paradigm to bonding-by-design, one starts with the desired band feature for a given application, and rationally tunes the crystal chemistry to achieve this band feature.



structure of a given material to tune it towards next-generation electronic, optic, thermoelectric, and correlated quantum materials.

Data availability

All data supporting the conclusions of this paper can be found in the ESI.† The interactive visualization tool used to tune the tight-binding model to morph between the silicon and germanium band structure can be found at <https://viz.whsunre.search.group/tb/>.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

Funding provided by the U.S. Department of Energy, Office of Science, Basic Energy Sciences, Division of Materials Science, through the Office of Science Funding Opportunity Announcement (FOA) Number DE-FOA-0002676: Chemical and Materials Sciences to Advance Clean-Energy Technologies and Transform Manufacturing.

References

- M. A. Green, Silicon photovoltaic modules: a brief history of the first 50 years, *Prog. Photovoltaics Res. Appl.*, 2005, **13**, 447–455.
- A. I. Boukai, *et al.*, Silicon nanowires as efficient thermoelectric materials, *Nature*, 2008, **451**, 168–171.
- Y. Cui, Z. Zhong, D. Wang, W. U. Wang and C. M. Lieber, High Performance Silicon Nanowire Field Effect Transistors, *Nano Lett.*, 2003, **3**, 149–152.
- G. J. Snyder and E. S. Toberer, Complex thermoelectric materials, *Nat. Mater.*, 2008, **7**, 105–114.
- G. Brunin, F. Ricci, V.-A. Ha, G.-M. Rignanese and G. Hautier, Transparent conducting materials discovery using high-throughput computing, *npj Comput. Mater.*, 2019, **5**, 1–13.
- M. Toriyama and G. J. Snyder, *Are Topological Insulators Promising Thermoelectrics?* <https://chemrxiv.org/engage/chemrxiv/article-details/6545ba97c573f893f1c4221d>, 2023, DOI: [10.26434/chemrxiv-2023-3nvl3](https://doi.org/10.26434/chemrxiv-2023-3nvl3).
- G. Hautier, Finding the needle in the haystack: Materials discovery and design through computational ab initio high-throughput screening, *Comput. Mater. Sci.*, 2019, **163**, 108–116.
- S. Curtarolo, *et al.*, The high-throughput highway to computational materials design, *Nat. Mater.*, 2013, **12**, 191–201.
- W. Chen, *et al.*, Understanding thermoelectric properties from high-throughput calculations: trends, insights, and comparisons with experiment, *J. Mater. Chem. C*, 2016, **4**, 4414–4426.
- R. Hoffmann, How Chemistry and Physics Meet in the Solid State, *Angew. Chem., Int. Ed. Engl.*, 1987, **26**, 846–878.
- R. Hoffmann, *Solids and Surfaces: A Chemist's View of Bonding in Extended Structures*, John Wiley & Sons, 2021.
- M. G. Goesten and R. Hoffmann, Mirrors of Bonding in Metal Halide Perovskites, *J. Am. Chem. Soc.*, 2018, **140**, 12996–13010.
- J. K. Burdett, From bonds to bands and molecules to solids, *Prog. Solid State Chem.*, 1984, **15**, 173–255.
- W. A. Harrison, Bond-Orbital Model and the Properties of Tetrahedrally Coordinated Solids, *Phys. Rev. B*, 1973, **8**, 4487–4498.
- D. J. Chadi and M. L. Cohen, Tight-binding calculations of the valence bands of diamond and zinc blende crystals, *Phys. Status Solidi B*, 1975, **68**, 405–419.
- P. Vogl, H. P. Hjalmarson and J. D. Dow, A Semi-empirical tight-binding theory of the electronic structure of semiconductors†, *J. Phys. Chem. Solids*, 1983, **44**, 365–378.
- J.-M. Jancu, R. Scholz, F. Beltram and F. Bassani, Empirical spds * tight-binding calculation for cubic semiconductors: General method and material parameters, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1998, **57**, 6493–6507.
- S. Sapra, N. Shanthi and D. D. Sarma, Realistic tight-binding model for the electronic structure of II-VI semiconductors, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2002, **66**, 205202.
- T. B. Boykin, G. Klimeck and F. Oyafuso, Valence band effective-mass expressions in the sp 3d 5s * empirical tight-binding model applied to a Si and Ge parametrization, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2004, **69**, 115201.
- A. S. Martins, T. B. Boykin, G. Klimeck and B. Koiller, Conduction-band tight-binding description for Si applied to P donors, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2005, **72**, 193204.
- Y. M. Niquet, D. Rideau, C. Tavernier, H. Jaouen and X. Blase, Onsite matrix elements of the tight-binding Hamiltonian of a strained crystal: Application to silicon, germanium, and their alloys, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2009, **79**, 245201.
- D. Soccodato, G. Penazzi, A. Pecchia, A.-L. Phan and M. A. der Maur, Machine learned environment-dependent corrections for a spds * empirical tight-binding basis, *Mach. Learn. Sci. Technol.*, 2024, **5**, 025034.
- Z. Wang, *et al.*, Machine learning method for tight-binding Hamiltonian parameterization from ab-initio band structure, *npj Comput. Mater.*, 2021, **7**, 1–10.
- G. Klimeck, *et al.*, Si tight-binding parameters from genetic algorithm fitting, *Superlattices Microstruct.*, 2000, **27**, 77–88.
- D. N. Talwar and C. S. Ting, Tight-binding calculations for the electronic structure of isolated vacancies and impurities in III-V compound semiconductors, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1982, **25**, 2660–2680.
- T. B. Boykin, Improved fits of the effective masses at Γ in the spin-orbit, second-nearest-neighbor sp 3 s * model: Results from analytic expressions, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1997, **56**, 9613–9618.



- 27 J. P. Loehr and D. N. Talwar, Exact parameter relations and effective masses within sp³s zinc blende tight-binding models, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1997, **55**, 4353–4359.
- 28 K. C. Hass, H. Ehrenreich and B. Velický, Electronic structure of Hg_{1-x}Cd_xTe, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1983, **27**, 1088–1100.
- 29 W. A. Harrison, New tight-binding parameters for covalent solids obtained using Louie peripheral states, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1981, **24**, 5835–5843.
- 30 K. E. Newman and J. D. Dow, Theory of deep impurities in silicon-germanium alloys, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1984, **30**, 1929–1936.
- 31 T. J. Lenosky, *et al.*, Highly optimized tight-binding model of silicon, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1997, **55**, 1528–1544.
- 32 Y. M. Niquet, C. Delerue, G. Allan and M. Lannoo, Method for tight-binding parametrization: Application to silicon nanostructures, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2000, **62**, 5109–5116.
- 33 D. Sanchez-Portal, E. Artacho and J. M. Soler, Projection of plane-wave calculations into atomic orbitals, *Solid State Commun.*, 1995, **95**, 685–690.
- 34 S. Ciraci and I. P. Batra, Electronic-energy-structure calculations of silicon and silicon dioxide using the extended tight-binding method, *Phys. Rev. B*, 1977, **15**, 4923–4934.
- 35 G. Grosso and C. Piermarocchi, Tight-binding model and interactions scaling laws for silicon and germanium, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1995, **51**, 16772–16777.
- 36 N. Marzari, A. A. Mostofi, J. R. Yates, I. Souza and D. Vanderbilt, Maximally localized Wannier functions: Theory and applications, *Rev. Mod. Phys.*, 2012, **84**, 1419–1475.
- 37 G. Dresselhaus and M. S. Dresselhaus, Fourier Expansion for the Electronic Energy Bands in Silicon and Germanium, *Phys. Rev.*, 1967, **160**, 649–679.
- 38 R. Dronskowski and P. E. Bloechl, Crystal orbital Hamilton populations (COHP): energy-resolved visualization of chemical bonding in solids based on density-functional calculations, *J. Phys. Chem.*, 1993, **97**, 8617–8624.
- 39 V. L. Deringer, A. L. Tchougréeff and R. Dronskowski, Crystal Orbital Hamilton Population (COHP) Analysis As Projected from Plane-Wave Basis Sets, *J. Phys. Chem. A*, 2011, **115**, 5461–5466.
- 40 S. Maintz, V. L. Deringer, A. L. Tchougréeff and R. Dronskowski, LOBSTER: A tool to extract chemical bonding from plane-wave based DFT: Tool to Extract Chemical Bonding, *J. Comput. Chem.*, 2016, **37**, 1030–1035.
- 41 In our 1D model, the bond weight ranges between 0–1 and quantifies how much the orbitals in the bond are in the second crystal wavefunction. When the second wavefunction is entirely p orbitals (like at $k = \Gamma$), $c_p = -i$ and the p–p bond weight will be $(-i)^\dagger(-i) = 1$. Because, the bond weight is independent of translation R, the onsite, 1NN, and 2NN weights are the same.
- 42 X. Qian, *et al.*, Quasiatomic orbitals for *ab initio* tight-binding analysis, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2008, **78**, 245112.
- 43 G. Kresse and J. Furthmüller, Efficient iterative schemes for *ab initio* total-energy calculations using a plane-wave basis set, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1996, **54**, 11169–11186.
- 44 G. Kresse and D. Joubert, From ultrasoft pseudopotentials to the projector augmented-wave method, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1999, **59**, 1758–1775.
- 45 G. Kresse and J. Furthmüller, Efficiency of *ab initio* total energy calculations for metals and semiconductors using a plane-wave basis set, *Comput. Mater. Sci.*, 1996, **6**, 15–50.
- 46 J. C. Slater and G. F. Koster, Simplified LCAO Method for the Periodic Potential Problem, *Phys. Rev.*, 1954, **94**, 1498–1524.
- 47 U. Herath, *et al.*, PyProcar: A Python library for electronic structure pre/post-processing, *Comput. Phys. Commun.*, 2020, **251**, 107080.
- 48 L. Lang, *et al.*, Expanding PyProcar for new features, maintainability, and reliability, *Comput. Phys. Commun.*, 2024, **297**, 109063.
- 49 R. Hoffmann, An Extended Hückel Theory. I. Hydrocarbons, *J. Chem. Phys.*, 1963, **39**, 1397–1412.
- 50 Y. Tsuji and K. Yoshizawa, Mixed-Anion Control of C–H Bond Activation of Methane on the IrO₂ Surface, *J. Phys. Chem. C*, 2020, **124**, 17058–17072.
- 51 L.-D. Yuan, H.-X. Deng, S.-S. Li, S.-H. Wei and J.-W. Luo, Unified theory of direct or indirect band-gap nature of conventional semiconductors, *Phys. Rev. B*, 2018, **98**, 245203.

