



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A review of perovskite/Si tandem solar cells: internal and external components toward high efficiency, long-term durability, and commercialization

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The rapid evolution of photovoltaic (PV) technology has made solar modules a key solution to meet growing global energy demands. In this context, achieving higher PV efficiency and reducing energy costs have become paramount objectives. Tandem solar cells, in which perovskite subcells are integrated with silicon (Si) subcells, represent a viable solution to surpass the Shockley–Queisser (S–Q) limit that constrains the efficiency of single-junction solar cells. These tandem configurations have demonstrated remarkable efficiency, reaching up to 34.85%, and are at the forefront of current PV research. This review focuses on recent studies aimed at enhancing the efficiency, stability, and scalability of tandem solar cells, including categorizing key areas of development in tandem solar cells into internal components (e.g., Si and perovskite subcells and interconnecting layers) and external components (e.g., encapsulation and busbars). Additionally, we address the fabrication process and levelized cost of energy (LCOE) of perovskite/Si tandem solar cells for cost-effective mass production. Moreover, we provide an outlook on the technological advancements required for the successful commercialization of tandem solar cells.

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Introduction

Photovoltaic (PV) technology stands at the forefront of addressing the increasing global energy demand. Since the first Si solar cell was reported in 1954,¹ with a power conversion efficiency (PCE) of 6.0%, significant advances in materials development, device architecture, and fabrication processes have pushed Si solar cells to efficiencies exceeding 26.0%.² Meanwhile, metal halide perovskites (MHPs) have emerged as highly promising materials not only for photovoltaics but also for a wide range of optoelectronic applications including light-emitting diodes, photodetectors, and lasers.^{3–7} Their outstanding optoelectronic properties such as high absorption coefficients, long carrier diffusion lengths, tunable bandgaps, together with compatibility with low-temperature solution processing have enabled remarkable efficiency improvements in perovskite solar cells (PSCs). Recent progress in

composition engineering, defect passivation, and interface control has further enhanced device performance and stability, underscoring the potential of MHPs as next-generation semiconductors. Consequently, PSC efficiencies have risen from 3.8% in 2009 to a certified efficiency of 26.95% today.^{2,8}

Despite significant advancements in both Si and perovskite PVs, their PCEs remain inherently constrained by the Shockley–Queisser (S–Q) limit. For Si PVs where the Si has a bandgap (E_g) of 1.12 eV, the maximum achievable PCE is approximately 32%.^{9,10} Similarly, PSCs face a PCE limit of around 30% using perovskite with a E_g of 1.6 eV.¹¹ The efficiency limit of single-junction solar cells arises from transmission of low-energy photons and thermalization of high-energy photons.

Tandem solar cells which consist of multiple cells with light absorbers having different bandgaps can overcome the S–Q limit by ensuring effective coverage across the wide solar spectrum.^{12–14} Perovskite (wide E_g)/Si (narrow E_g) tandem solar cells are a notable example that recently achieved a certified PCE of 34.85% that surpasses the theoretical limit of single junction solar cells.² Fig. 2 shows the certified efficiency evolution of perovskite/Si tandem solar cells, highlighting continuous improvements from 2016 to 2025.² To realize commercialization of the perovskite/Si tandem solar cells, nevertheless, further improvement in both efficiency and stability is required. Thus, we categorize key factors into

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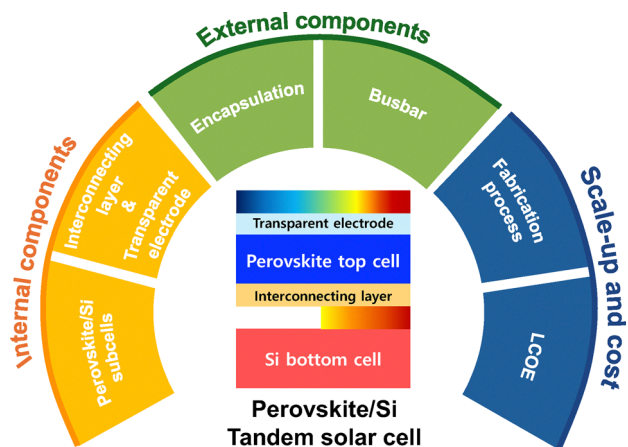


Fig. 1 Schematic overview of the review addressing internal components and external components of perovskite/Si tandem solar cells.

internal and external components, which mutually connect and contribute to the structural and functional characteristics of the cells (Fig. 1), to understand and analyze the current state of perovskite/Si tandem solar cells, gaining insights essential for their further development in the industrial scale.

Internal components of tandem solar cells refer to the elements and materials (e.g., Si and perovskite subcells, and interconnecting layers) that are directly involved in the photo-conversion process within the cell. On the other hand, external components act as the supportive and protective elements around the photovoltaic device (e.g., encapsulation and busbars), crucial for enhancing its overall performance and lifetime.

Among the internal components, the Si bottom cell is typically configured into three types: passivated emitter and rear cell (PERC), tunnel oxide passed contact (TOPCon), and Si heterojunction (SHJ). PERC cells are the most widely adopted due to their compatibility with traditional cell designs and the efficiency boost by the addition of a rear passivation layer. TOPCon cells, regarded as the next generation of PERC technology, also integrate into the conventional cell structure while being further enhanced by a tunnel-oxide layer of SiO_2 , which

improves surface passivation and facilitates selective charge transport. SHJ cells use a hydrogenated amorphous Si (a-Si:H) layer on both sides of the crystalline Si (c-Si) substrate, which provides surface passivation of the c-Si surfaces, thereby improving cell efficiency. Given the unique structures and characteristics of each Si subcell type, further evaluation of their compatibility with perovskite top cells is essential for developing optimal Si/perovskite tandem solar cells.

As a top cell in a tandem configuration, PSCs offer advantages due to their tunable E_g , which allows spectral complementarity with Si bottom cells.^{15,16} However, despite these promising features, PSCs face critical challenges such as open-circuit voltage (V_{OC}) losses and poor stability, which are associated with the presence of defects and their migration.^{17–19}

The interconnecting layer (ICL) is also an essential component for establishing perovskite/Si tandem solar cells as it connects the individual subcells, enabling their operation as a tandem unit. An ideal ICL should possess optimal electrical properties to minimize charge carrier losses between the sub-cells and maintain high optical transparency, particularly in the near-infrared (NIR) region, to reduce light absorption losses in the narrow bandgap subcell.²⁰

For the external components, encapsulation serves as a protective barrier against environmental factors such as moisture, thermal stress and dust, extending durability and operational lifetime of the cells. In addition, busbars play a critical role in ensuring efficient current collection across the large area of the cells.

Lastly, we address the fabrication processes of perovskite/Si tandem solar cells and examine their levelized cost of electricity (LCOE) to assess economic viability—a critical aspect for market adoption.²¹ Here, we compare reported tandem LCOE values of perovskite/Si tandem solar cells with those of conventional c-Si PV technologies. By covering these topics, our review provides a comprehensive analysis of the current state of perovskite/Si tandem solar cells, suggesting pathways to further their enhancement and commercialization.

Part I: internal components of perovskite/Si tandem solar cells

Si bottom cells for perovskite/Si tandem solar cells

Over the past 30 years, Si solar cells have made significant progress, largely driven by advancements in the device structure (Fig. 3(a)).²² A key breakthrough has been the development and optimization of passivating contacts at the metal contact interface, which greatly enhances the PCEs by reducing surface and interfacial recombination losses as well as optical losses (Fig. 3(c)–(e)).²³ In perovskite/Si tandem solar cells where the top perovskite and bottom Si cells are integrated, optimizing the Si bottom cell design is crucial. This includes considerations such as cell structure, surface morphology or texture, and fabrication process.

Below, we introduce the most common types of Si solar cells used in perovskite/Si tandem solar cells —PERC, TOPCon, and

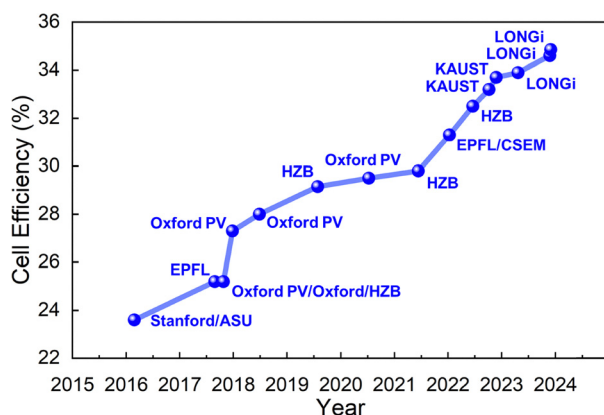


Fig. 2 Certified efficiency evolution of perovskite/Si tandem solar cells (2016–2025).²

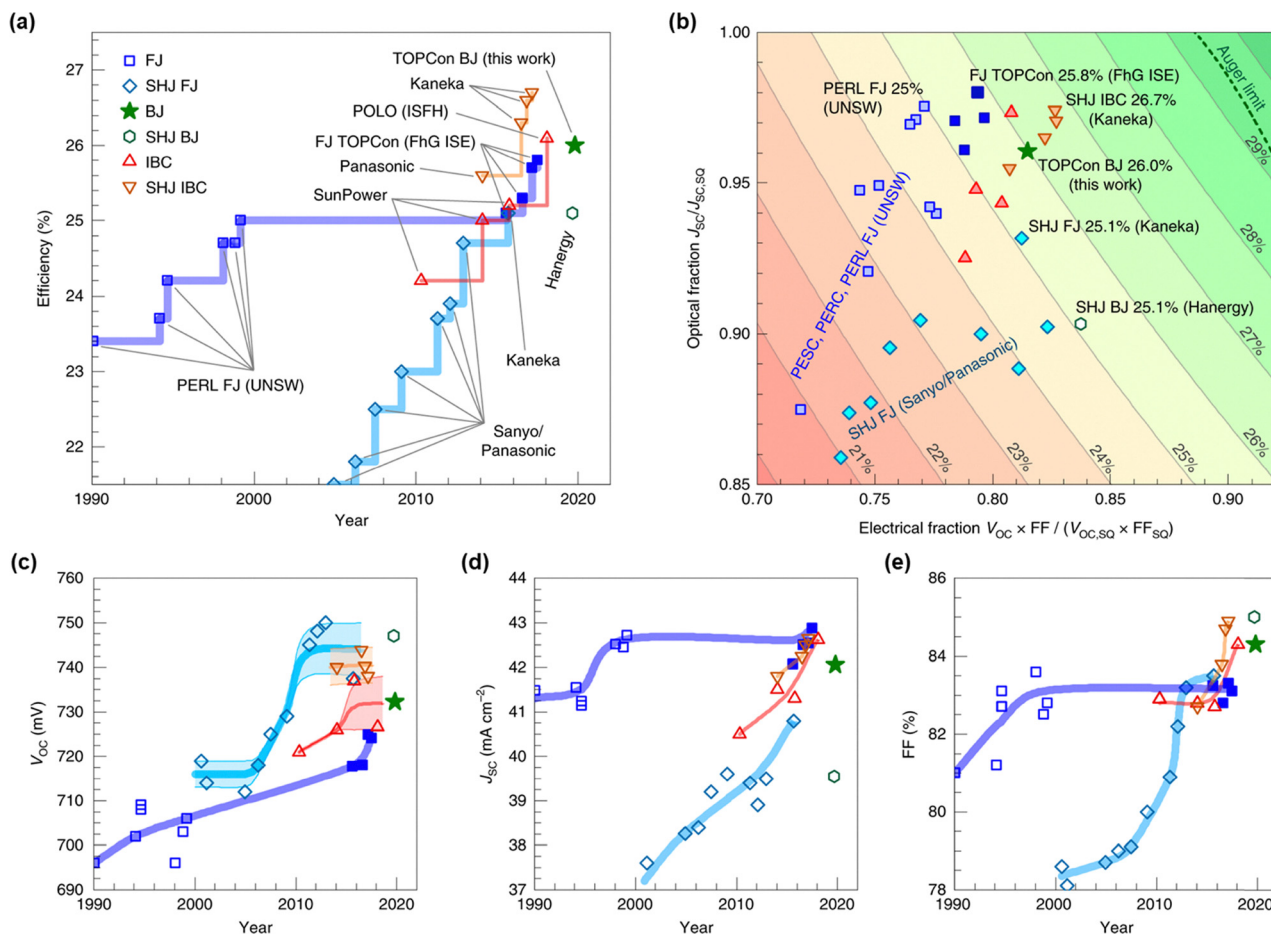


Fig. 3 (a) Efficiency advancement of Si solar cells through cell structure development. (b) Electrical and optical fraction of the various solar cells, parameter change (c) V_{oc} , (d) J_{sc} and (e) FF of various types of Si solar cells for the past 30 years. FJ: front junction, BJ: back junction, IBC: interdigitated back contact, POLO: poly-Si on oxide. Reproduced with permission.²³ Copyright 2021, Springer Nature.

SHJ—and address potential challenges associated with their integration with perovskite top cells.

PERC (passivated emitter rear contact) cells

PERC cells typically employ p-type Si (p-Si) as an absorber layer. The front side consists of an n^+ emitter, a SiN_x :H layer, and a metal grid (Fig. 4(a)). The n^+ emitter is a doped Si layer that

collects charges, while the SiN_x layer reduces light reflection. On the rear side, PERC cells have a local back rear field (BSF), a rear passivation layer, and another metal grid.²⁴ The $\text{AlO}_x/\text{SiN}_x$ stack is a key component of the rear passivation layer to reduce dangling bonds, electrical losses, and carrier recombination, which improves V_{oc} .^{25,26} A high negative fixed charge density of the AlO_x layer ($\sim 10^{13} \text{ cm}^{-2}$) generates an electric field that repels negative charge carriers, reducing recombination at the

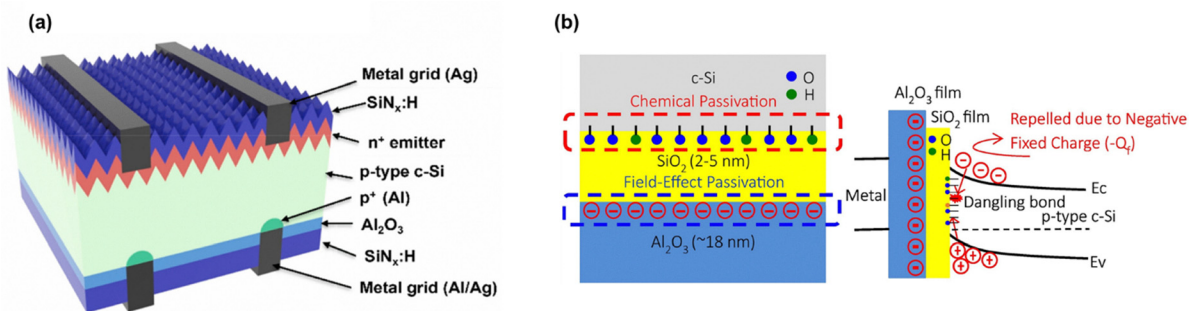


Fig. 4 (a) Structure of PERC cell. Reproduced with permission.²⁵ Copyright 2023, Elsevier Inc., (b) schematic energy band diagram of $\text{AlO}_x/\text{SiN}_x$ interface. Reproduced with permission.³⁰ Copyright 2019, Elsevier B.V.

p-Si surface.^{27,28} On the other hand, SiN_x helps terminate dangling bonds, providing chemical passivation and reducing surface recombination velocity as shown in Fig. 4(b).^{29,30}

TOPCon (tunnel oxide passivated contact) cell

A standard TOPCon cell uses n-type Si (n-Si) as an absorber. The front side consists of a doped p-Si layer, which serves as an emitter, along with a SiN_x layer, and front metal contact (Fig. 5(a) and (b)). On the rear side, an ultrathin tunnel oxide layer, typically made of SiO₂, is paired with a doped n-Si rear contact layer and back metal contact. The SiO₂ layer facilitates the quantum tunneling of electrons while blocking holes, thereby providing surface passivation and enhancing carrier selectivity.^{31–33} The doped p-Si layer exhibits high charge transport properties, facilitating efficient carrier collection.^{34,35} This combination of SiO₂ and p-Si stack mitigates the dangling bonds, suppresses metal contact recombination, promotes charge transport, and increases V_{OC}.³⁵

To achieve a high-performing TOPCon cell, controlling the thickness of the tunnel oxide layer is important.³⁴ For instance, an ultrathin SiO₂ layer allows for high tunneling probability for charge carriers, but pinholes can easily form during fabrication with heat treatment. While the formation of the pinholes may enable efficient charge transport leading to an increase in FF, excessive pinholes can compromise the passivation effect of the SiO₂ layer, creating pathways for leakage currents and increasing recombination at the interface.³⁴ Conversely, a thicker SiO₂ layer limits the transport channels for charge carriers, reducing tunneling efficiency because the probability of quantum tunneling decreases exponentially with increasing thickness.³⁷ Therefore, achieving optimal TOPCon cell performance requires precise control of both thickness and pinhole density of the SiO₂ layer.

In addition to advancements in rear passivation, improvements to the absorber have also contributed to the development of TOPCon solar cells.^{35,38} To date, most research has focused on n-type c-Si wafers, while p-type c-Si wafers have been less explored due to their lower bulk carrier lifetime—primarily caused by boron–oxygen complex formation—and their inferior

interface passivation, which stems from greater sensitivity to surface defect states.³⁹ However, from a mass production perspective, p-type c-Si wafers may be more suitable for TOPCon solar cells, as well-established fabrication processes for p-type wafers align with PERC production lines, potentially reducing production costs.⁴⁰ This compatibility may enable a cost-effective transition from PERC to TOPCon technology with minimal modifications, leveraging the current infrastructure and reducing the need for additional investment.

SHJ (Si heterojunction) cells

The structure of SHJ cells is symmetric with a c-Si layer as an absorber, positioned between intrinsic hydrogenated amorphous Si (a-Si:H(i))/doped a-Si:H(n or p) layers/transparent conducting oxide (TCO) electrodes (Fig. 6(a)).³⁶ The intrinsic a-Si:H layers serve as a passivation layer on c-Si surfaces while the doped a-Si:H layers form p-type or n-type regions, creating a surface potential that promotes charge separation, injection, and collection. The TCO layers, typically deposited *via* sputtering, enhance lateral charge transport and reduce parasitic absorption by a-Si:H layers, especially in the NIR region.⁴¹ Additionally, the TCO layer induces band bending at the a-Si:H/c-Si interface, generating an electric field at the junction. This field enhances carrier selectivity by reducing electron–hole recombination and facilitates efficient extraction of photogenerated carriers.⁴² However, integrating a-Si:H and TCO into the cell reduce light absorption in the range of 300–400 nm and at wavelengths beyond 750 nm. Reducing the thickness of the TCO layer can enhance the passing of more light, minimizing parasitic absorption.

Texturing Si bottom cells for perovskite/Si tandem solar cells

Textured Si bottom cells with micro-sized pyramids improve photon absorption by increasing light trapping and reducing reflection.^{44–46} The surface geometry scatters incident light efficiently, inducing multiple internal reflections that extend the optical path length within the absorber layer. This effect is particularly beneficial in the NIR region where absorption of Si is inherently weaker. The most common texturing method involves anisotropic etching with alkaline solutions such as KOH, NaOH, or TMAH, which selectively etch the Si surface

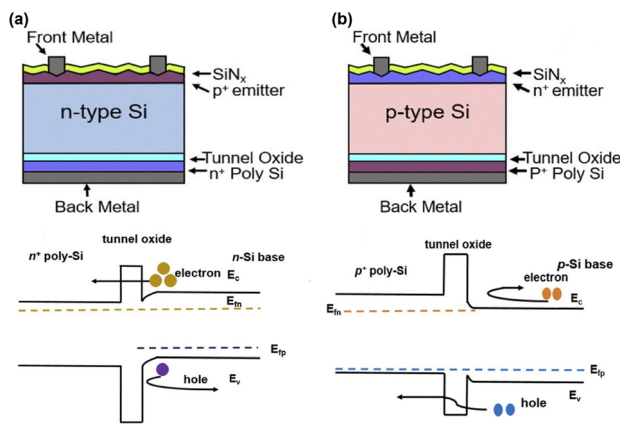


Fig. 5 (a) Structure of n-type TOPCon cell and band diagram, (b) structure of p-type TOPCon cell and band diagram. Reproduced with permission.³⁶ Copyright 2024, The Royal Society of Chemistry.

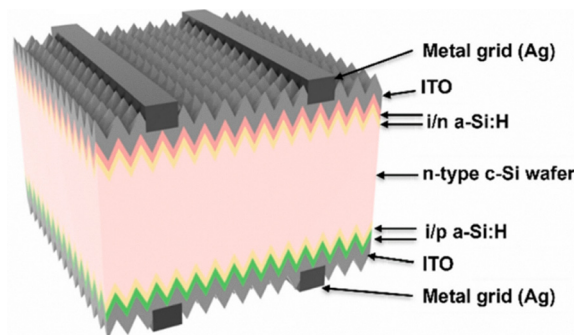


Fig. 6 (a) Structure of SHJ cell. Reproduced with permission⁴³ Copyright 2023, Elsevier Inc.

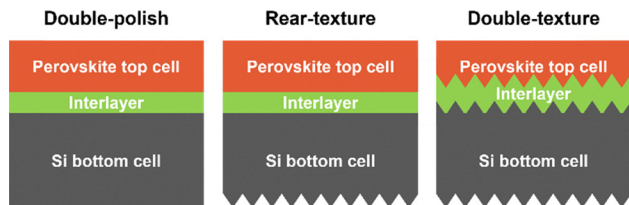


Fig. 7 Types of texturing for Si bottom cells in perovskite/Si tandem solar cells.

along its crystallographic planes to form well-defined pyramid arrays.^{47,48}

In perovskite/Si tandem solar cells, the textured structures are typically classified as double-polish, rear-texture, or double-texture (Fig. 7). The double-polish and rear-texture structures, characterized by a flat top surface of Si were initially developed to simplify thin-film deposition including interlayers and perovskites by providing smoother surfaces for uniform coverage. However, these configurations often increase optical losses due to higher surface reflection and reduced light trapping, resulting in lower J_{SC} values. In contrast, double-textured structures enhance light absorption by light scattering and trapping within the device and have become the standard for high-performance solar cells. Kanda *et al.* investigated the effect of Si surface texturing on the performance of perovskite/Si tandem cells by comparing planar (polished) and textured Si surfaces on the top side (Fig. 8(a)).⁴⁹ Their results showed that texturing the Si top surface significantly reduced reflectance, which increased light absorption and contributed to an improvement in the current density from 13.7 mA cm^{-2} for planar surfaces to 14.8 mA cm^{-2} for textured

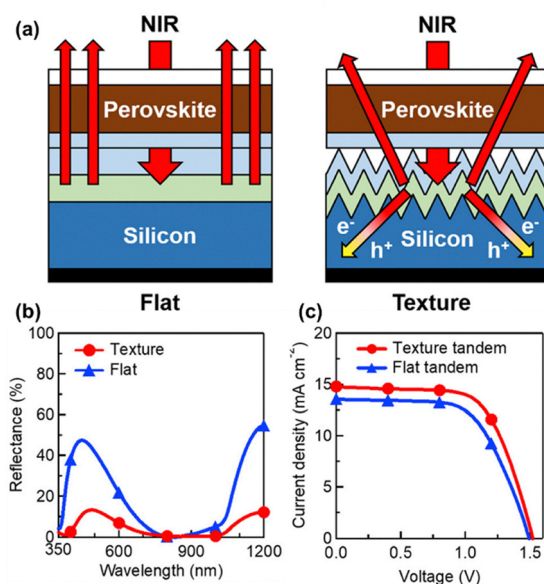


Fig. 8 (a) Structure of perovskite/polished Si (left) and perovskite/textured Si tandem solar cell (right). (b) Reflectance comparison of planar and textured Si solar cells. (c) I - V curves of perovskite/planner and textured Si tandem solar cells. Reproduced with permission.⁴⁵ Copyright 2018, American Chemical Society.

surfaces (Fig. 8(b) and (c)). Achieving a uniform coverage of the perovskite layer on textured Si requires the deposition of a sufficiently thick layer, particularly to cover pyramidal tips. Solution-based methods, such as spin coating and blade coating, often involve challenges in achieving uniformity due to the micro-pyramidal surface features.^{50,51} In contrast, vacuum deposition techniques offer a viable alternative by enabling more uniform film formation on textured surfaces.⁵² However, vapor deposition methods have limitations, including weaker adhesion to textured substrates, which can induce mechanical stress and the formation of defects in the perovskite layer. Therefore, hybrid approaches that combine the scalability of solution-based methods with the precision of vacuum processes are needed to optimize perovskite deposition on textured Si.

PERC cells in perovskite/Si tandem solar cells

PERC cells, widely adopted in commercial single-junction Si solar cells due to their low cost and manufacturing maturity, have also been considered as bottom cells in perovskite/Si tandem configurations. However, their passivation structure, typically composed of $\text{Al}_2\text{O}_3/\text{SiN}_x$ presents challenges for electrical integration. While these layers provide effective surface passivation, their insulating nature prevents the direct formation of a recombination junction between the perovskite and Si subcells. To address this limitation, Wu *et al.* introduced a thin metal stack (Cr/Pd/Ag) as an interfacial contact between the Si emitter and the ITO layer (Fig. 9(a) and (b)).⁵³ By locally penetrating the passivation layers, the metal stack established a low-resistance contact pathway, thereby facilitating efficient carrier transport and resulting in a PCE of 22.5%. Although the metal stack improved carrier transport, PERC cells fundamentally rely on heavy doping at the metal-Si interface to reduce contact resistance.^{54,55} Thus, while effective in single-junction cells, the use of PERC cells in tandem configurations may lead to increased carrier recombination and reduced V_{OC} .

TOPCon cells in perovskite/Si tandem solar cells

With their tunnel oxide and poly-Si passivating contacts, TOPCon cells offer effective surface passivation and compatibility with existing manufacturing processes, making them suitable as bottom cells in perovskite/Si tandem configurations. However, their performance is limited by optical losses arising from the poly-Si layer, which induces parasitic absorption in the near-infrared (NIR) region primarily due to its high free carrier concentration and a refractive index ($n \approx 3.8$) significantly higher than that of c-Si ($n \approx 3.4$). The optical mismatch reduces photon transmission to the Si absorber, disrupting current matching between the subcells.⁵⁷ To mitigate this issue, Ding *et al.* replaced poly-Si with poly-SiO_x ($n \approx 3.2$), which is better matched to that of the Si bottom cell (Fig. 9(c)).⁵⁶ As a result, photon transmission to the Si bottom cell increased, yielding a PCE of 25.12% (Fig. 9(d)). In addition to the optical issue associated with the presence of poly-Si, its interface with the underlying SiO_x layer often suffers from a high defect density, particularly on textured surfaces.⁵⁸ Jiang *et al.* introduced an AlO_x:H capping layer on top of the SiO_x to reduce recombination

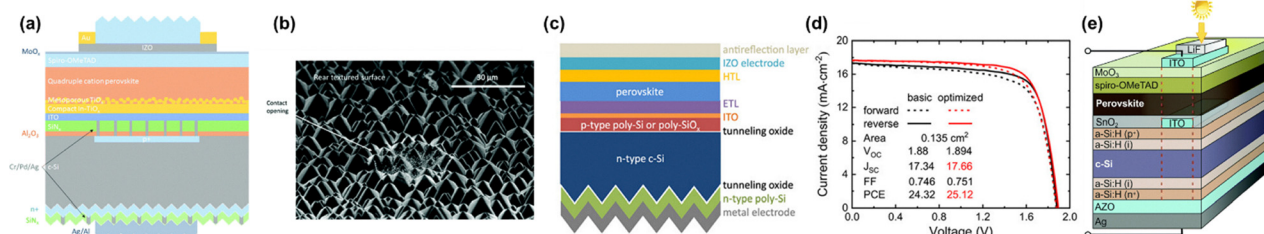


Fig. 9 (a) Structure of perovskite/PERC tandem solar cell with an intermediate contact of a Cr/Pd/Ag metal stack, and (b) SEM image of textured Si surface with local openings using the metal stack. Reproduced with permission.⁵³ Copyright 2017, The Royal Society of Chemistry. (c) Structure of a perovskite/TOPCon tandem solar cell incorporating a poly-SiO_x layer, and (d) *J*-*V* characteristics comparing basic (using poly-Si) and optimized (using poly-SiO_x) configurations of the perovskite/TOPCon tandem solar cell. Reproduced with permission.⁵⁶ Copyright 2024, John Wiley and Sons. (e) Structure of perovskite/SHJ tandem solar cell with SnO₂ layer deposited using ALD. Reproduced with permission.⁴¹ Copyright 2016, The Royal Society of Chemistry.

at the SiO_x/poly-Si interface.⁵⁹ The fixed negative charge in AlO_x:H induces a field-effect that repels minority carriers from the interface, while incorporated hydrogen passivates dangling bonds and reduces the trap density. The combination of field-effect and chemical passivation effectively suppressed recombination at the SiO_x/poly-Si interface, leading to a PCE of 28.67%.

SHJ cells in perovskite/Si tandem solar cells

SHJ cells, consisting of a-Si:H(i) and a-Si:H(n or p) layers, also offer effective surface passivation, having achieved record efficiencies of 27.3% for single junction Si cells and 34.6% for perovskite/Si tandem cells.^{60,61} Nevertheless, their integration into the tandem architecture is limited by the thermal instability of a-Si:H as its exposure to temperatures above 200 °C leads to hydrogen effusion, breaking Si-H bonds and degrading surface passivation.⁶² This issue becomes particularly critical during the fabrication of the perovskite subcell, which involves high-temperature processes such as mesoporous TiO₂ annealing at or above 500 °C.⁶³ Accordingly, low-temperature processing methods have been developed to preserve the integrity of the a-Si:H passivation layers. Albrecht *et al.* employed SnO₂ which can be deposited below 120 °C by atomic layer deposition (ALD) (Fig. 9(e)). Given its high electron mobility and broad optical transparency, SnO₂ served as an efficient electron-selective contact without degrading the a-Si:H interface and thereby enabled integration of the SHJ bottom cell with a thermally compatible perovskite top cell, resulting in a PCE of 19.9%.^{41,64}

Wide-bandgap perovskite top cells for perovskite/Si tandems

Wide-bandgap perovskites have gained considerable attention due to their potential in tandem solar cells, where they can be integrated with lower-bandgap Si cells. These materials typically have a bandgap greater than 1.65 eV and optimization at 1.67–1.75 eV is commonly performed for use with 1.1–1.2 eV Si bottom cells (Fig. 10(a)).⁶⁵ The bandgap can be adjusted through composition engineering of the iodide (I)/bromide (Br) ratio often combined with A-site engineering such as the incorporation of Cs⁺. In this section, key challenges in the development of wide bandgap perovskites are examined, along with strategies to address them.

One of the most critical issues in wide bandgap perovskite is that *V*_{OC} does not increase with an increased bandgap, while current density decreases. This phenomenon can be explained by halide segregation as reported by Hoke *et al.*⁶⁸ The study found that the photoluminescence of I/Br mixed perovskite shows a red-shift with time under constant light soaking, which does not appear in pure halides (*e.g.*, MAFACsPbI₃ and MAPbI₃). Halide segregation is observed only in a mixed halide composition-based wide bandgap perovskite with *V*_{OC} pinning as the Br/I ratio increases (Fig. 10(b)).⁶⁶ Therefore, halide segregation was considered the major cause of *V*_{OC} loss in wide bandgap perovskite solar cells. However, recent studies suggest that non-radiative recombination especially at the interface of the wide bandgap perovskite could be more critical than halide segregations.^{69–71} This was evidenced by the extraordinarily low initial electroluminescence external quantum efficiency (EL-EQE) of mixed halide perovskite devices (10⁻⁶ to 10⁻⁸) prior to the halide segregation as the EL-EQE is directly related to *V*_{OC} as shown in eqn (1).⁷² For example, in a study on a 1.77 eV perovskite, the authors found that a *V*_{OC} loss by trap-assisted non-radiative recombination was 385 mV, which is approximately 5 times larger than a *V*_{OC} loss (75 mV) by halide segregation. Additionally, the top contact of the perovskite layer, such as the electron transporting layer (ETL, *e.g.*, PCBM) is a critical factor for *V*_{OC} loss. The photoluminescence quantum yield (PLQY) of the isolated perovskite film is 10⁻³, but this reduces below the detection limit with an ETL, indicating significant *V*_{OC} loss at the perovskite interface.

$$V_{OC} = V_{OC,rad} + \left(\frac{k_B T}{q}\right) \cdot \ln(EQE_{EL}) \quad (1)$$

The origin of initial traps that cause non-radiative recombination in wide bandgap perovskites was suggested by Huang *et al.*, in the context of the crystallization process (Fig. 10(c)).⁶⁷ According to their findings, the Br-containing precursors have lower solubility than the I-based precursors, resulting in the formation of Br-perovskite nuclei prior to I-perovskite during the spin coating process. Therefore, the I-Br mixed perovskite is not directly formed from a solution state but instead involves solid-state diffusion of I from I-based perovskite into Br-based perovskite species. This halide migration between I and Br during film formation occurs *via* a ‘vacancy-assisted diffusion

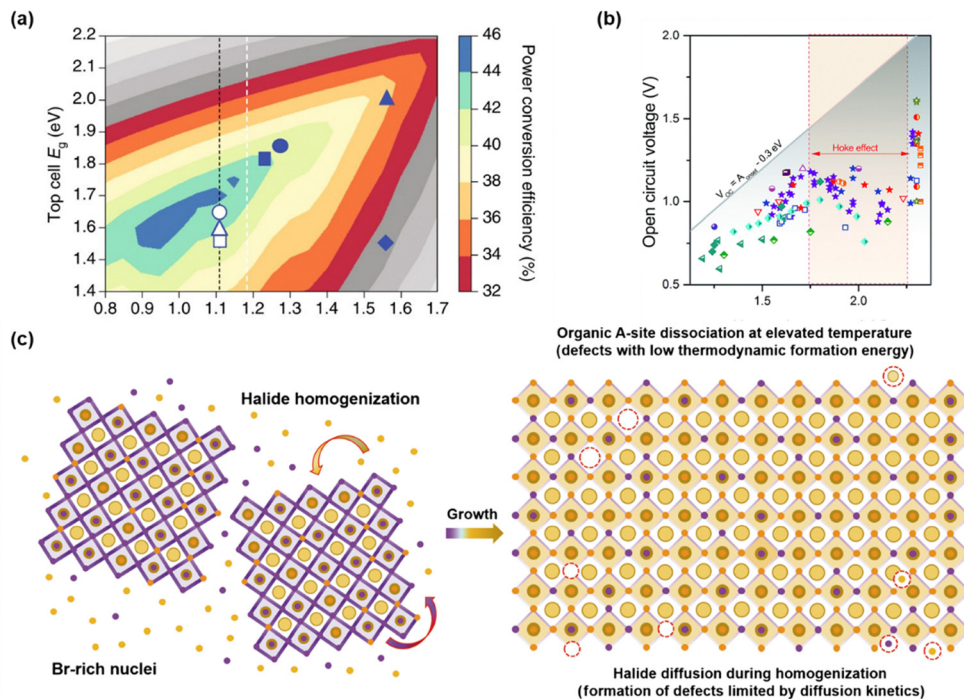


Fig. 10 (a) Theoretical power conversion efficiencies with different bandgaps for 2-terminal tandem solar cells. The black dashed line indicates the 1.12 eV bandgap of Si. Reproduced with permission.⁶⁵ Copyright 2018, Springer Nature, (b) Reported V_{OC} as function of bandgap showing onset of the Hoke effect. Reproduced with permission.⁶⁶ Copyright 2017, The Royal Society of Chemistry, (c) the defect generation model of I–Br mixed perovskite during film growth *via* halide homogenization. Reproduced with permission.⁶⁷ Copyright 2021, American Association for the Advancement of Science.

process”, forming point defects. Recent studies have shown that the positive iodide interstitial defects (I_i^+) generate deep traps in the I–Br mixed wide-bandgap perovskite.⁷³ This defect generation process occurs not only during the growth process, but also during halide segregation, which involves ion migration *via* solid-state diffusion. Hence, the halide segregation may accelerate the defect generation in the I–Br mixed perovskite. Moreover, this segregated film results in the energetic loss due to the different energy levels at the interface between perovskite and charge transport layers (CTLs), since the defects formed during the film growth and segregation could be particularly abundant at the interface.^{73,74} In a recent study by J. Warby *et al.*, it was demonstrated that not only mixed halide, but also cations impact on interface non-radiative recombination at the perovskite interface, such as perovskite/electron transport layer (ETL) interface in a p–i–n structure.⁷⁵ Specifically, the cations were found to affect the broadening of the density of states of C_{60} , resulting in the formation of additional trap states. This finding is consistent with another important aspect of voltage loss, which is the energetic misalignment between the wide bandgap perovskite and CTLs. Typically, CTLs such as SnO_2 , spiro-OMeTAD, nickel oxide (NiO_x), PTAA, and C_{60} are optimized for use with narrow or middle bandgap perovskites ($E_G \leq 1.6$ eV) and have demonstrated record high performances.^{65,76} Therefore, using these same CTLs for wide bandgap perovskites could involve in energetic losses as the energy bands differ from those of lower bandgap perovskites.

In order to minimize V_{OC} loss in wide bandgap perovskite, various solutions have been extensively reported. Compositional

engineering such as replacing A site and X sites with using Cs^+ , K^+ , Rb^+ , and SCN^- ,^{77–80} employing alkylammonium halide additives such as phenformin hydrochloride (PhenHCl), trimethylphenylammonium tribromide (TPABr₃), *n*-butylammonium bromide (BABr), and acetate based molecules lead acetates ($Pb(Ac)_2$), bimolecular (co-incorporation) of PEAI/ $Pb(SCN)_2$.^{73,81–87} To suppress the trap formation during the film growth from rapid crystallization of Br based perovskite, alternative methods for antisolvent dripping such as the gas quenching method is also successfully reported.⁸⁸ In addition, to alleviate halide segregation, which accounts for a partial V_{OC} loss as discussed above, cation engineering using cations such as FA^+ , MA^+ and Cs^+ is widely reported, and the various effects of the ions on the crystal structure and dipole effects are reported.^{89,90} In all of the reported approaches, improvements in V_{OC} were shown, and it is suggested that process changes such as composition engineering, additive engineering, and solvent extraction are needed. On the other hand, this also means that a technique for universally maximizing the V_{OC} of wide bandgap perovskites has not yet been firmly established, as these are necessary to minimize V_{OC} loss.

The stability of perovskite solar cells is a crucial and frequently discussed topic in the perovskite optoelectronics community. This challenge is not limited to wide bandgap perovskites but is relevant to most optoelectronic devices based on perovskite.

One of the main concerns with wide bandgap perovskites is their integration with Si tandem solar cells. To achieve this integration, the top perovskite solar cell’s lifetime must be compatible with that of the Si solar cells, which is around

20 years. If the perovskite top cell's lifetime is shorter, it could compromise the performance of the tandem cell.^{91,92} The halide segregation is commonly indicated as the limiting factor for the stable perovskite.^{68,93} While recent studies have reported stable perovskite solar cells containing MA⁺, this compound is generally considered to be responsible for low thermal and moisture stability. Hence, despite these recent developments, the challenge of achieving stable perovskite remains a critical issue within the perovskite optoelectronics community.^{94,95}

Various approaches have been employed to address the stability issues of perovskite solar cells. These include compositional engineering, which is similar to the methods used to minimize V_{OC} loss, and cation engineering that involves the incorporation of Cs⁺, K⁺, and organic cations.^{77–79} In addition, PEA⁺ and DMA⁺ are examples of large organic cations that have been reported to be effective inhibitors of halide segregation (Fig. 11).^{96,97} Furthermore, the use of triple-halides (I, Br, and Cl) has been suggested to suppress halide segregation. Studies have shown almost no segregation up to 100 sun intensity for 1.67 eV perovskite films when a triple-halide is used.⁹⁸

An alternative approach to obtaining wide bandgap perovskites that are fundamentally free from the halide segregation is CsPbI₃, possessing bandgaps depending on the phases 1.74 eV, 1.73 eV, and 1.64 eV for γ -, α - and β -phases, respectively.^{99,100} Recent studies using CsPbI₃ have shown that it achieves PCEs exceeding 20%, suggesting that this approach holds promise as a solution for wide bandgap perovskite solar

cells free from halide segregation.^{101–103} However, both the thermodynamically unstable phase and the need for further improvements in the material's stability against moisture, light, and heat pose significant challenges that still require attention and ongoing efforts for the use of CsPbI₃ perovskite.

Interconnecting layer (ICL) of perovskite/Si tandem solar cells

The interconnecting layer (ICL) plays a vital role in perovskite/Si tandem solar cells by establishing electrical connection between the two subcells. It should also allow efficient transmission of sub-bandgap photons to the Si bottom cell through high optical transparency to ensure efficient light use. The ICL is expected to have sufficiently high optical transparency while minimizing parasitic absorption and interfacial reflection. Electrically, it should exhibit high vertical conductivity to limit voltage and series resistance losses.

ICLs typically adopt either recombination junctions (RJs) or tunnel junctions (TJs) (Fig. 12 and Table 1). RJs, which are often based on TCO layers, offer high transparency and are relatively easy to fabricate. However, their limited conductivity can cause carrier accumulation, voltage loss, and potential shunting.

TJs, by contrast, consist of heavily doped p⁺⁺/n⁺⁺ layers that form a narrow depletion region, allowing efficient quantum tunneling. While TJs exhibit superior electrical performance, they may also suffer from optical losses caused by free carrier absorption in the doped regions. Mitigating these losses requires precise control over doping concentration and interface passivation. The choice between RJs and TJs involves trade-offs in

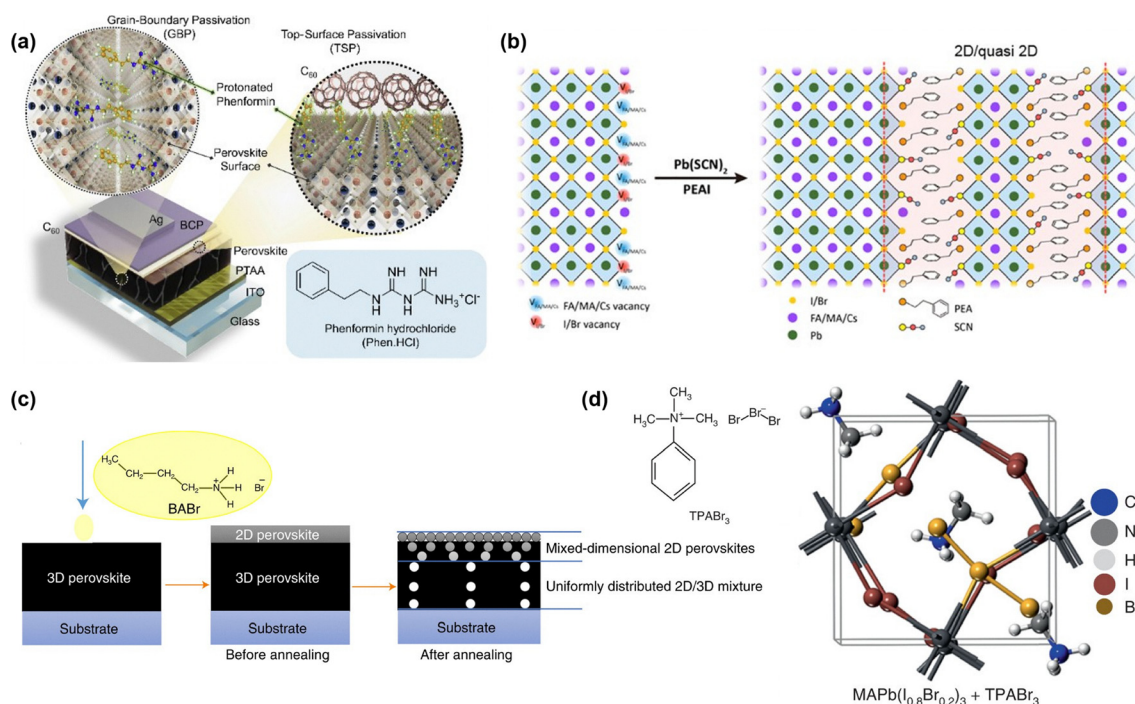


Fig. 11 Various alkylammonium halide additives strategies to reduce V_{OC} loss such as (a) phenformin hydrochloride (Phen.HCl). Reproduced with permission.⁷³ Copyright 2021 Elsevier Inc., (b) $Pb(SCN)_2$ + PEAI. Reproduced with permission.⁸⁷ Copyright 2019, Elsevier Inc., (c) *n*-butylammonium bromide (BABr). Reproduced with permission.⁷⁸ Copyright 2021 Springer Nature, (d) trimethylphenylammonium tribromide (TPABr₃). Reproduced with permission.⁷⁷ Copyright 2022 Springer Nature.

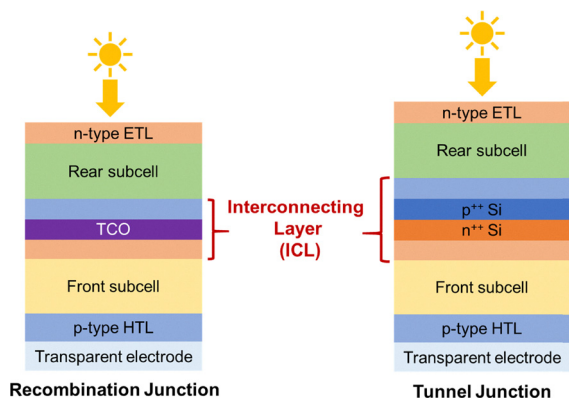


Fig. 12 Structure of perovskite/Si tandem solar cells using recombination junction and tunnel junction.

optical transmission, electrical conductivity, and fabrication complexity, making ICL optimization a key design challenge in perovskite/Si tandem solar cells.

Recombination junctions (RJs)

RJs, most commonly based on TCOs, are widely employed in perovskite/Si tandem solar cells to facilitate carrier recombination between the subcells. In addition to providing electrical connection, they are expected to minimize optical losses through high transmittance in the visible spectrum ($>80\%$) and maintain low resistivity ($\sim 2.0 \times 10^{-4} \Omega \text{ cm}$ at room temperature).^{104,105} Recent efforts have focused on optimizing the junction to reduce parasitic absorption and enhance recombination efficiency, both of which directly impact tandem device performance (Table 2). Among the various TCO-based materials, ITO remains the most widely adopted due to its excellent conductivity and transparency. In a pioneering work, Albrecht *et al.* utilized ITO as the RJ layer, achieving a PCE of 18.1%.⁴¹ However, the large refractive index mismatch between ITO ($n \approx 2.0$) and c-Si ($n \approx 3.4$) led to significant interfacial reflection losses, limiting light transmission to the bottom cell.^{106,107} To address this issue, Mazzarella *et al.* introduced a hydrogenated nanocrystalline SiO_x (nc- $\text{SiO}_x\text{:H}$) interlayer with an intermediate refractive index ($n \approx 2.5$), which improved optical matching, boosting PCE to 25.2%.¹⁰⁸ Another challenge in ITO-based RJs is excessive in-plane conductivity, which can cause lateral leakage and device shunting.^{109,110} In

terms of electrical performance, the sheet resistance of ITO layers used in RJs typically ranges from 10 to 20 $\Omega \text{ sq}^{-1}$, which is necessary to ensure sufficient lateral conductivity for current collection in standard Si-based devices.¹¹¹ However, in tandem structures, this high lateral conductivity ($\sim 10^{-4} \Omega \text{ cm}$) can promote parasitic current pathways, especially in regions of non-uniform perovskite coverage or pinholes, leading to lateral leakage currents and localized shunting.¹¹² Furthermore, the absence of a built-in junction field across the RJ layer necessitates high vertical conductivity to support efficient carrier recombination while minimizing voltage losses. To address these challenges, recent designs increasingly focus on decoupling vertical and lateral conduction paths, for example by incorporating low-conductivity HTLs (*e.g.*, NiO_x), which can suppress lateral leakage while maintaining sufficient vertical recombination efficiency. Xu *et al.* introduced a sputtered NiO_x layer as an HTL, which reduced lateral conductivity while improving interface passivation and energy-level alignment at the NiO_x /perovskite interface (Fig. 13(a)).¹¹³ To further improve interfacial quality and charge extraction, a spiro-TTB buffer layer was introduced at the NiO_x /perovskite interface. The additional buffer layer promoted uniform perovskite nucleation, suppressed pinhole formation, and facilitated efficient charge extraction, collectively contributing to a PCE of 27.43%.

Recent studies have demonstrated the effectiveness of self-assembled monolayers (SAMs) as an interfacial layer in RJs for perovskite/Si tandem solar cells.^{149,152} The molecular-scale thickness of SAMs ensures high optical transparency while allowing dipole-induced energy level alignment to facilitate charge extraction. In addition, SAMs contribute to reducing interfacial recombination by chemically passivating surface charge extraction. In addition, SAMs contribute to reducing interfacial recombination by chemically passivating surface defects. Al-Ashouri *et al.* employed Me-4PACz as a SAM at the RJ, achieving a PCE of 29.05% along with facilitated hole extraction indicated by reduced transient PL lifetime ($\sim 300 \text{ ns}$).¹²² Despite these advantages, conventional SAMs such as Me-4PACz typically exhibit poor wettability and non-uniform coverage, potentially causing device shunting and reduced reproducibility. To address these challenges, Li *et al.* introduced a mixed SAM composed of Me-4PACz and MeO-PhPACz (Fig. 13(b) and (c)).¹⁵¹ The mixed SAM significantly enhanced wettability due to the incorporation of MeO-PhPACz, which contains polar functional

Table 1 Comparison of recombination junction (RJ) and tunnel junction (TJ) interconnecting layers

Feature	Recombination junction (RJ)	Tunnel junction (TJ)
Definition	Interface enabling carrier recombination and electrical connection.	Quantum tunneling through highly doped junction.
Typical materials	ITO, IZO, TCO/ NiO_x , TCO/SAMs.	a-Si:H(p^+/n^+), nc-Si:H, poly-Si, TiO_2 /ITO.
Optical property	High transparency, depending on TCO quality.	High transparency if thin; free-carrier absorption increases with heavy doping.
Electrical property	Moderate resistance, affected by interface defects.	Very low resistance with optimized doping.
Advantages	Simple fabrication, wide material choice.	High conductivity, efficient carrier recombination, low voltage loss.
Disadvantages	Interface traps and long-term stability issues.	Doping control challenges and process complexity.

Table 2 Efficiency of perovskite/Si tandem solar cells with various RJs

Year	Perovskite composition	Perovskite bandgap (eV)	Si type	Recombination junction (RJ)	J_{sc} (mA cm ⁻²)	V_{oc} (V)	FF (%)	PCE (%)	Ref.
2015	FA _{0.83} CS _{0.17} Pb(I _{0.83} Br _{0.17}) ₃	—	SHJ	ITO/SnO ₂	14.00	1.79	79.50	19.90	41
2015	MAPbI ₃	—	SHJ	IZO/PCBM ^a /PEIE ^b	15.80	1.69	79.90	21.40	114
2016	MAPbI ₃	—	PERC	ZTO/c-TiO ₂ /m-TiO ₂	15.30	1.68	68.00	17.40	115
2017	CS _{0.17} FA _{0.83} Pb(Br _{0.17} I _{0.83}) ₃	1.63	SHJ	ITO/NiO	18.10	1.65	79.00	23.60	116
2017	CS _{0.07} Rb _{0.03} FA _{0.765} MA _{0.135} PbI _{2.55} Br _{0.45}	1.62	SHJ	ITO/TiO ₂	17.60	1.75	73.80	22.80	53
2018	CS _{0.17} FA _{0.83} Pb(Br _{0.17} I _{0.83}) ₃	—	PERC	ITO/NiO _x	15.30	1.43	75.00	16.20	117
2018	FA _{0.75} CS _{0.25} Pb(I _{0.8} Br _{0.2}) ₃	1.68	SHJ	ITO/PTAA ^c	18.40	1.77	77.00	25.00	106
2019	CS _{0.15} (FA _{0.83} MA _{0.17}) _{0.85} Pb(I _{0.7} Br _{0.3}) ₃	1.70	SHJ	ITO/PTAA	16.40	1.83	74.50	22.40	118
	CS _{0.15} (FA _{0.83} MA _{0.17}) _{0.85} Pb(I _{0.75} Br _{0.25}) ₃	1.68			17.10	1.81	79.00	24.50	
	CS _{0.15} (FA _{0.83} MA _{0.17}) _{0.85} Pb(I _{0.8} Br _{0.2}) ₃	1.64			17.80	1.80	79.40	25.40	
	CS _{0.15} (FA _{0.83} MA _{0.17}) _{0.85} Pb(I _{0.85} Br _{0.15}) ₃	1.62			17.40	1.77	78.10	24.10	
2019	FA _{0.8} MA _{0.2} PbBr _{0.6} I _{2.4}	1.64	PERC	ITO/PTAA	16.12	1.65	79.92	21.19	119
2019	FA _{0.75} CS _{0.25} Pb(I _{0.8} Br _{0.2}) ₃	1.70	SHJ	ITO/c-TiO ₂ /m-TiO ₂	15.20	1.84	77.30	21.60	120
2019	MAPbI ₃	—	SHJ	ITO/AgNW ^d /TCA ^e /ZnO/PEF ^f /PCBM	14.70	1.78	80.40	21.00	121
2020	(FA _{0.65} MA _{0.2} CS _{0.15})Pb(I _{0.8} Br _{0.2}) ₃	1.70	SHJ	ITO/PTAA	19.20	1.76	79.20	26.70	84
2020	CS _{0.05} FA _{0.77} MA _{0.23} Pb(I _{0.77} Br _{0.23}) ₃	1.67	SHJ	ITO/NiO _x /PolyTPD/PFN ^g	19.12	1.89	75.30	27.13	98
2020	CS _{0.05} (FA _{0.77} MA _{0.23}) _{0.95} Pb(I _{0.77} Br _{0.23}) ₃	1.68	SHJ	ITO/Me-4PACz ^h	19.26	1.90	79.52	29.05	122
2020	FA _{0.75} CS _{0.25} Pb(I _{0.8} Br _{0.2}) ₃	1.68	SHJ	ITO/PTAA/PFN	17.70	1.77	80.30	25.10	123
2022	CS _{0.05} FA _{0.8} MA _{0.15} Pb(I _{0.75} Br _{0.25}) ₃	—	SHJ	IZO/2PACz ⁱ	18.60	1.86	76.00	26.20	124
2022	CS _{0.05} (FA _{0.83} MA _{0.17}) _{0.95} Pb(I _{0.83} Br _{0.17}) ₃	1.63	TOPCon	IZO/MeO-2PACz ^j	19.40	1.80	81.64	28.49	125
2022	CS _{0.05} (FA _{0.74} MA _{0.26}) _{0.95} Pb(I _{0.74} Br _{0.26}) ₃	—	SHJ	ITO/2PACz + MeO-2PACz	19.40	1.86	79.58	28.80	126
2022	FA _{1-x} MA _x PbI _{3-y} Br _y	1.59	PERC	ITO/NiO _x	16.50	1.75	81.80	23.70	127
2022	CS _{0.22} FA _{0.78} Pb(Cl _{0.03} Br _{0.15} I _{0.85}) ₃	1.68	TOPCon	ITO/NiO _x /PolyTPD ^k	19.68	1.79	78.27	27.63	128
2023	CS _{0.05} (FA _{0.83} MA _{0.17}) _{0.95} Pb(I _{0.83} Br _{0.17}) ₃	1.64	SHJ	ITO/PTAA/PFN	19.90	1.73	73.00	25.60	107
2023	MASCN:CS _{0.22} Pb(I _{0.85} Br _{0.15}) ₃	1.68	SHJ	ITO/NiO/2PACz + MeO-2PACz	19.80	1.85	78.90	28.90	129
2023	CS _{0.22} FA _{0.78} Pb(I _{0.85} Br _{0.15}) ₃	1.63	TOPCon	IZO/MeO-2PACz	19.3	1.80	81.9	28.50	130
2023	MASnI _{3-x} Br _x	1.3–2.15	SHJ	ITO/spiro-OMeTAD ^l	16.89	2.14	84.55	30.70	131
2023	CS _{0.05} FA _{0.79} MA _{0.16} Pb(I _{0.75} Br _{0.25}) ₃	—	PERC	ITO/2PACz	18.60	1.69	76.9	24.20	132
2023	CS _{0.22} FA _{0.78} Pb(I _{0.85} Br _{0.15}) ₃ + 5% MAPbCl ₃	1.68	SHJ	ITO/Me-4PACz	20.24	1.98	81.18	32.50	133
2023	CS _{0.15} FA _{0.65} MA _{0.20} Pb(I _{0.80} Br _{0.20}) ₃	1.68	TOPCon	IZO/MoO _x /spiro-OMeTAD	18.80	1.78	81.80	27.40	134
2023	CS _{0.15} FA _{0.65} MA _{0.20} Pb(I _{0.80} Br _{0.20}) ₃	1.68	TOPCon	IZO/Me-4PACz	20.70	1.78	82.00	30.26	135
2023	CS _{0.05} (FA _{0.75} MA _{0.25}) _{0.95} Pb(I _{0.75} Br _{0.25}) ₃	1.55	SHJ	ITO/P3CT-N ^m	20.39	1.83	77.57	28.89	136
2023	CS _x FA _y MA _{1-(x+y)}} Pb(I,Br) ₃ (x + y ≤ 1)	1.65	SHJ	ITO/NiO _x /2PACz	19.56	1.77	79.40	27.51	137
2024	CS _{0.05} FA _{0.80} MA _{0.15} Pb(I _{0.75} Br _{0.25}) ₃	1.67	SHJ	ITO/NiO/2PACz + MeO-2PACz	20.10	1.94	76.70	29.80	138
2024	CS _{0.05} (FA _{0.84} MA _{0.16}) _{0.95} Pb(I _{0.75} Br _{0.25}) ₃	1.67	TOPCon	ITO/NiO	19.58	1.93	81.54	30.09	139
2024	CS _{0.05} FA _{0.80} MA _{0.15} Pb(I _{0.83} Br _{0.17}) ₃	1.55	SHJ	ITO/MeO-2PACz	20.64	1.82	75.92	28.50	140
2024	CS _{0.05} FA _{0.80} MA _{0.15} Pb(I _{0.75} Br _{0.25}) ₃	1.68	SHJ	ITO/Me-4PACz	20.09	1.93	78.43	30.43	141
2024	CS _{0.05} (FA _{0.90} MA _{0.1}) _{0.95} Pb(I _{0.8} Br _{0.2}) ₃	1.65	SHJ	ITO/Me-4PACz/SiO ₂ -NP ⁿ	19.81	1.95	79.90	30.93	142
2024	CS _{0.05} FA _{0.80} MA _{0.15} Pb(I _{0.75} Br _{0.25}) ₃	1.68	SHJ	IZO/Me-4PACz	20.96	1.95	80.50	34.00	143
2024	FA _{0.78} CS _{0.22} Pb(I _{0.82} Br _{0.18}) ₃ and excess 3% MAPbCl ₃	1.65	SHJ	ITO/PolyTPD	20.40	1.96	81.00	32.50	144
2024	FA _{0.8} MA _{0.15} CS _{0.05} Pb(I _{0.76} Br _{0.24}) ₃ and 5% excess PbX ₂ (X = I, Br)	1.69	SHJ	IZO/MeO-4PACz	20.76	1.97	83.00	33.89	145
2025	CS _{0.05} MA _{0.15} FA _{0.8} Pb(I _{0.76} Br _{0.24}) ₃	1.66	SHJ	ITO/NiO _x /Me-4PACz	20.23	1.98	80.76	30.98	146
2025	CS _{0.17} FA _{0.83} Pb(I _{0.80} Br _{0.20}) ₃	1.67	SHJ	ITO/4-PhCz ^o	19.92	1.96	81.00	31.26	147
2025	CS _{0.15} FA _{0.85} Pb(I _{0.77} Br _{0.23}) ₃	1.66	TOPCon	ITO/4PADCB ^p	20.00	1.88	82.60	31.10	148
2025	CS _{0.05} FA _{0.80} MA _{0.15} Pb(I _{0.75} Br _{0.25}) ₃	1.68	SHJ	ITO/2PhPA-CzH ^q	19.92	1.91	84.51	32.19	149
2025	CS _{0.22} FA _{0.63} MA _{0.15} Pb(I _{0.83} Br _{0.14} Cl _{0.03}) ₃	1.66	SHJ	IZO	20.70	1.92	80.90	32.10	150

^a PCBM: [6,6]-phenyl-C₆₁-butyric acid methyl ester. ^b PEIE: polyethylenimine ethoxylated. ^c PTAA: poly[bis(4-phenyl)(2,4,6-trimethylphenyl)amine]. ^d AgNW: silver nanowire. ^e TCA: trichloroacetic acid. ^f PEI: polyethylenimine. ^g PFN: poly[(9,9-bis(3'-(N,N-dimethylamino)propyl)-2,7-fluorene)-alt-2,7-(9,9-dioctylfluorene)]. ^h Me-4PACz: [2-(9H-carbazol-9-yl)ethyl]phosphonic acid methyl ester. ⁱ 2PACz: [2-(9H-carbazol-9-yl)ethyl]phosphonic acid. ^j MeO-2PACz: [2-(3,6-dimethoxy-9H-carbazol-9-yl)ethyl]phosphonic acid. ^k PolyTPD: poly[N,N'-bis(4-butylphenyl)-N,N'-bis(phenyl)benzidine]. ^l Spiro-OMeTAD: 2,2',7,7'-tetrakis(N,N-di-p-methoxyphenylamine)-9,9'-spirobifluorene. ^m P3CT-N: poly[3-(4-carboxylate butyl)thiophene-2,5-diyl] (sodium salt). ⁿ SiO₂-NP: silicon dioxide nanoparticle. ^o 4-PhCz: 4-phenylcarbazole. ^p 4PADCB: N₄,N₄,N₄',N₄'-tetra[[1,1'-biphenyl]-4-yl]-[1,1'-biphenyl]-4,4'-diamine. ^q 2PhPA-CzH: 2-(9H-carbazol-9-yl)ethyl diphenylphosphonic acid.

groups that increase the surface energy, reducing the contact angle from 97° to 76°, and enabling uniform perovskite film formation. It also increased hole conductivity from 5.89 × 10⁻⁴ mS cm⁻¹ to 7.44 × 10⁻⁴ mS cm⁻¹, which can be attributed to the improved molecular packing density and optimized energy-level alignment at the interface while maintaining the high optical transparency (average transmittance >90% in the visible region). Owing to the improved coverage, device reproducibility increased to ~95% yield, significantly higher than the

reproducibility (<50%) obtained with a single SAM. As a result, tandem solar cells using the mixed SAM achieved a PCE of 28.07%. While enhanced coverage improves morphological uniformity, molecular dipole engineering provides an additional pathway for optimizing interfacial energetics in SAM-based RJs. Luo *et al.* developed a series of pyridine-annulated anthracene (PyAA) SAMs with inductive substituents (PyAA, PyAA-Br, PyAA-Me, and PyAA-MeO) on ITO for perovskite/Si tandem solar cells.¹⁵³ By tuning the electron-donating or withdrawing nature

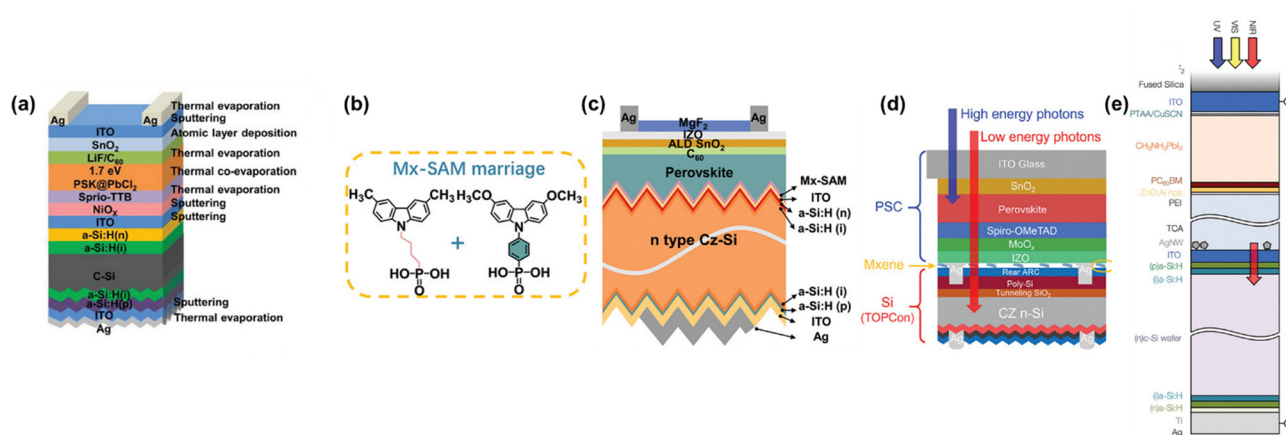


Fig. 13 (a) Structure of a perovskite/Si tandem solar cell using NiO_x interlayers. Reproduced with permission.¹¹³ Copyright 2023, Wiley-VCH GmbH. (b) Mx-SAM configurations, combining Me-4PACz and MeO-PhPACz molecules, and (c) structure of a perovskite/Si tandem solar cell using an Mx-SAM interlayer. Reproduced with permission.¹⁵¹ Copyright 2024, Wiley-VCH GmbH. (d) Structure of an RJ-incorporated MXene. Reproduced with permission.¹³⁵ Copyright 2019, Wiley-VCH GmbH. (e) Structure of perovskite/Si tandem solar cells using a D-PEDOT:PSS RJ. Reproduced with permission.¹²¹ Copyright 2019, Wiley-VCH GmbH.

of the substituents, they modulated the dipole moment of the SAMs, which in turn shifted the vacuum level at the ITO/perovskite interface, improving energy-level alignment and facilitating hole extraction while reducing recombination losses. Among the SAMs, PyAA-MeO featuring a moderately electron-donating methoxy group exhibited an optimal balance between dipole strength and molecular packing, leading to substantial increase in device efficiency with a certified PCE of 30.9%. Despite the remarkable progress achieved with conventional carbazole-based SAMs, their symmetric molecular structure still causes steric hindrance near the anchoring sites, which prevents close molecular packing, thereby limiting both interfacial uniformity and defect passivation. To overcome the structural limitation of SAMs, Jia *et al.* designed an asymmetric carbazole-based SAM in which a spacer and a phosphonic-acid anchoring group flank the phenyl ring of the carbazole core.¹⁵⁴ The asymmetry reduces steric hindrance and enables denser molecular packing and more uniform surface coverage on TCO substrates. Moreover, the modified configuration strengthens coordination with undercoordinated Pb^{2+} ions and induces a larger interfacial dipole, which together enhances charge extraction and suppresses non-radiative recombination. As a result, perovskite/Si tandem solar cells employing the asymmetric SAM achieved a certified PCE of 34.58%, outperforming devices using symmetric Me-4PACz (32.77%) and MeO-4PACz (33.88%).

MXenes are also promising RJ materials due to their high electrical conductivity ($\sim 15\,100\text{ S cm}^{-1}$), broadband optical transmittance ($>90\%$ in the visible spectrum), and tunable work functions, which enable favorable energy-level alignment at the recombination interface.¹⁵⁵ Han *et al.* demonstrated that incorporating MXene interlayer reduced the recombination resistance from $\sim 77.1\ \Omega\text{ cm}^2$ to $\sim 29.5\ \Omega\text{ cm}^2$ and extended the TPV carrier lifetime from $\sim 1.8\ \mu\text{s}$ to $\sim 3.5\ \mu\text{s}$, indicating suppressed recombination and improved interfacial transport (Fig. 13(d)).¹³⁵ As a result, the tandem device achieved a

stabilized PCE of 29.65%. However, MXenes are susceptible to ambient oxidation and hydrolysis, which can compromise long-term operational stability. Another class of candidate RJ materials is that of conductive polymers, which offer solution processability alongside desirable optical and electrical characteristics. PEDOT:PSS, with its high optical transmittance ($\sim 85\text{--}90\%$) and intrinsically high in-plane resistivity ($\sim 10^0\text{--}10^{-1}\ \Omega\text{ cm}$), can suppress lateral leakage currents and has thus been investigated as an RJ material.¹⁵⁵ Ramírez Quiroz *et al.* used a D-sorbitol-doped PEDOT:PSS (d-PEDOT:PSS) that simultaneously served as a transparent conductive adhesive (TCA) (Fig. 13(e)).¹²¹ The layer exhibited high vertical conductivity ($\sim 1000\text{ S cm}^{-1}$), enabling efficient carrier recombination while suppressing lateral leakage. Sorbitol-induced polymer ordering also enhanced interfacial adhesion, improving mechanical stability. The resulting tandem solar cells achieved a PCE of 21.0%. However, the hygroscopic and acidic nature of PEDOT:PSS raises concerns over long-term stability, necessitating additional encapsulation or material modification for practical use.

Tunnel junction (TJ)

Despite substantial advances in RJ design, the use of TCOs often results in optical reflection losses and lateral leakage currents, which limit tandem performance. As an alternative, tunnel junctions (TJs) offer a fundamentally different approach, enabling efficient carrier recombination without relying on TCOs. A typical TJ consists of heavily doped n^{++} and p^{++} layers, with carrier concentrations typically approaching or exceeding 10^{19} to 10^{20} cm^{-3} . The high doping level significantly narrows the depletion region to a few nanometers, enabling quantum-mechanical tunneling of carriers across the junction.¹⁵⁶ Under forward bias, the energy alignment between filled states in the n^{++} layer and empty states in the p^{++} layer enables interband tunneling, either directly or *via* intermediate defect states.

The depletion width W , which critically influences tunneling probability, is given by eqn (2).

$$W = \sqrt{\frac{2\varepsilon(\phi_0 - V)N_A + N_D}{q N_A N_D}} \quad (2)$$

where ε is the dielectric permittivity, q is the elementary charge, N_A and N_D are the acceptor and donor doping concentrations, respectively, ϕ_0 is the built-in potential, and V is the applied voltage. As the depletion width W decreases with increasing doping concentration and forward bias, tunneling efficiency is enhanced by enabling carriers to tunnel across the junction. However, excessive doping may increase free carrier absorption and induce defect states that lead to trap-assisted non-radiative recombination. To mitigate the trade-offs associated with excessive doping, precise doping control and effective interface passivation are required to preserve electrical conductivity without incurring optical or recombination-related losses. Mailoa *et al.* implemented highly doped n^{++} a-Si:H deposited *via* plasma-enhanced chemical vapor deposition (PECVD) to demonstrate TJ integration in perovskite/Si tandem solar cells (Fig. 14(a) and (b)).¹⁵⁷ The highly doped amorphous Si layer resulted in a depletion width narrow enough to allow carrier tunneling, achieving a PCE of 13.7%. The TJ structure exhibited high optical transparency, exceeding 90% in the 700–1200 nm range, thereby facilitating the transmission of sub-bandgap photons to the Si bottom cell.¹⁵⁶ However, the electrical conductivity of a-Si:H is inherently constrained by its disordered atomic structure, and the presence of localized defect states can lead to trap-assisted recombination.^{33,34}

To overcome the limited conductivity and recombination losses induced by structural disorder in amorphous Si tunnel layers, Sahli *et al.* introduced nanocrystalline Si (nc-Si:H) as the TJ material (Fig. 14(c)).⁴⁴ The partial crystallinity of nc-Si:H improved bulk conductivity and carrier mobility by reducing structural disorder. In addition, as nc-Si:H formed conformal contact with textured Si surfaces, it can minimize interfacial voids and defect generation, thereby facilitating efficient vertical charge extraction. The nc-Si:H maintained high near-infrared optical transparency with average transmittance exceeding 85% in the 700–1200 nm range which is comparable to that of a-Si:H. The combined improvements in electrical conductivity and optical transparency enabled a certified PCE

of 25.2%. In addition, nc-Si:H exhibited enhanced thermal and chemical stability. STEM-EDX analysis revealed that conventional ITO/spiro-TTB/perovskite stacks underwent elemental redistribution upon annealing, whereas the nc-Si:H interface retained its compositional integrity, supporting stable long-term operation (Fig. 14(d)).

Shen *et al.* employed an interlayer-free TJ formed directly between ALD-deposited TiO₂ and p⁺-Si to reduce optical losses and simplify processing.¹⁵⁸ At the TJ, carrier recombination occurred *via* defect-assisted tunneling through interfacial trap states, eliminating the need for heavily doped n^{++} layers. The resulting TiO₂/p⁺-Si interface exhibited ohmic behavior after annealing with a low contact resistance of $\sim 30 \text{ m}\Omega \text{ cm}^2$, enabling effective tunneling across the interface. Compared to conventional TJ, due to the absence of interlayer in the TiO₂/p⁺-Si structure reduced parasitic optical absorption, improving photocurrent density by $\sim 1\text{--}1.5 \text{ mA cm}^{-2}$. Also, the process avoided PECVD deposition, improving compatibility with perovskite fabrication steps that are sensitive to plasma or high-temperature exposure. As a result, the tandem device achieved a PCE of 24.1%, validating the effectiveness of defect-mediated tunneling as a simplified and optically transparent TJ strategy. Both conventional heavily doped TJ and alternative designs based on defect-assisted tunneling have demonstrated high vertical conductivity and optical transparency, while preserving interfacial stability and process compatibility. Nonetheless, challenges related to doping-induced absorption, interfacial trap density, and long-term stability remain critical considerations in the design of TJ-based tandem solar cells.

Transparent electrodes

Transparent conductive oxides (TCOs), such as ITO or IZO, are the most used as top electrodes in perovskite/Si tandem cells due to their excellent transparency, conductivity, stability, fast deposition rate, and scalability. In this section, we will provide a brief overview of the challenges associated with using TCOs, with a particular focus on ITO and IZO, as top electrodes in perovskite/Si tandem cells. TCO films can be deposited by various methods, including sol-gel methods, chemical vapor deposition (CVD), e-beam evaporation, and sputtering. Among these, sputtering is the predominant choice for industrial applications due to its high deposition rate and scalability.^{159–163} Pulsed direct current (DC, 40–200 kHz), or radiofrequency

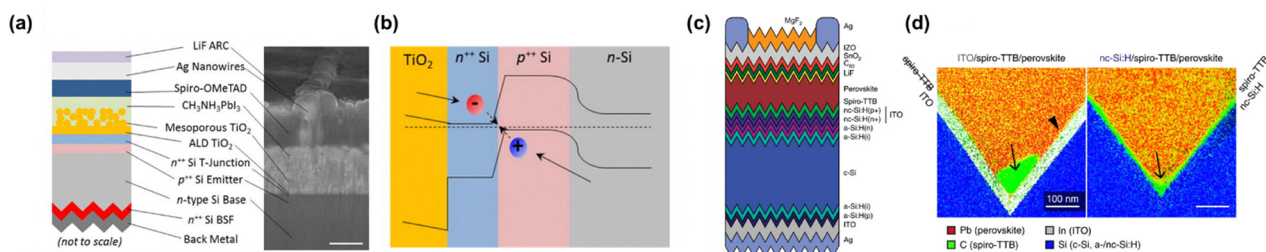


Fig. 14 (a) Structure of the tandem structure with TJ and (b) energy band diagram of tandem solar cells. Reproduced with permission.¹⁵⁷ Copyright 2015, AIP Publishing. (c) Structure of the perovskite/Si tandem solar cells with nc-Si:H TJ and (d) cross-sectional EDX mapping of the different layer stack. Reproduced with permission.⁴⁴ Copyright 2018, Springer Nature.

(RF, ~ 13 MHz) sputtering is commonly adopted for TCO deposition. Pulsed DC sputtering requires high voltage, typically a few hundred volts, to sustain the discharge through secondary electron emission, resulting in fast sputter rates and its widespread adoption in industrial fields. On the other hand, RF sputtering employs oscillating bulk plasma, resulting in a lower discharge voltage. This leads to lower-energy ion bombardment, enabling denser film growth but with reduced sputter rates.¹⁶⁴ Both DC and RF sputtering involve ionized inert gas, usually argon ions (Ar^+), transferring energy to the target material through collisions, which eject target atoms into the vacuum chamber and deposit them onto the substrate (Fig. 15(a)).¹⁶⁵ One of the common challenges associated with the sputter process is the damage inflicted on the underlying layer during the deposition. This is primarily due to the high energy of sputtered atoms, which can reach up to ~ 10 eV depending on the binding energy

of the target material, as well as the positive ions formed in the plasma (~ 15 eV), neutralized ions from the target surface (~ 20 eV), and the luminescence of the plasma.¹⁶⁵ The impact of such damage on optoelectronic devices, including light-emitting diodes and solar cells, has been extensively investigated, including perovskite solar cells. The problems most reported in this regard include the formation of an energetic barrier at the TCO/underlying layer interface due to degradation of the underlying layer during the sputter process, as well as an increase in leakage current, resulting in reduced shut resistance. Kanda *et al.* reported on the effects of sputter damage on the J - V characteristics of perovskite solar cells, whereby the sputtered TCO on the widely used hole-transporting material (spiro-OMeTAD) led to an increase in energetic barrier and the formation of an additional diode path, resulting in an s-shaped J - V curve (Fig. 15(b)).¹⁶⁶ Similar issues were observed in the p-i-n configuration of perovskite solar cells, where the TCO was sputtered on a commonly used ETL buffer layer, bathocuproine (BCP).¹⁶⁷ Hence, to avoid these problem, the most widely adopted method is to introduce a buffer layer on the organic charge transporting layer to prevent the damage from the sputtering process where a few certain characteristics are required such as material durability against the sputter damage and appropriate energy level for charge extraction. The buffer layer for TCO will be discussed in detail in the buffer layer section. To minimize sputter damage, various modifications and improvements have been reported for sputter systems. For example, the face target sputter (FTS) system has been widely employed, where the two targets face each other, confining the plasma to the vicinity of the target and reducing damage. Additionally, a metal and mesh shield can be inserted between the target and substrate to capture high-energy secondary electrons, resulting in improved OLED performance and reduced sputter damage (Fig. 15(c)).¹⁶⁸ The top TCO layer is the first layer that incident light passes through in the device stack, making it crucial for determining the overall amount of light inside the device. To minimize reflection at this layer, an anti-reflection (AR) layer is typically coated onto the TCO, which adjusts the refractive index between air and TCO. Low-refractive index materials such as LiF and MgF_2 are typically used for this purpose. E. Yang *et al.* recently reported that a meso-microporous SiO_2 AR coating prepared *via* sol-gel phase separation can reduce reflectance to below 1% and increase transmittance up to $\sim 99\%$, thereby improving light harvesting in perovskite/Si tandem solar cells.¹⁶⁹ A thinner TCO layer is optically beneficial; however, there is a trade-off between optical transparency and electrical conductivity, as excessively thin TCO layers may not provide sufficient lateral charge transport. To balance these trade-offs, the appropriate thickness of the TCO layer must be carefully determined through a trade-off analysis. Optical calculations, usually using the transfer matrix method, can readily optimize the thickness of both the AR layer and TCO layer (Fig. 16(a)). As previously mentioned, ITO and IZO are commonly used as the top electrode for the perovskite/Si tandem cell. However, Whal *et al.* reported that IZO could have potential benefits for the tandem application.¹⁷⁰ This is due to the fact that IZO does not require high-temperature annealing

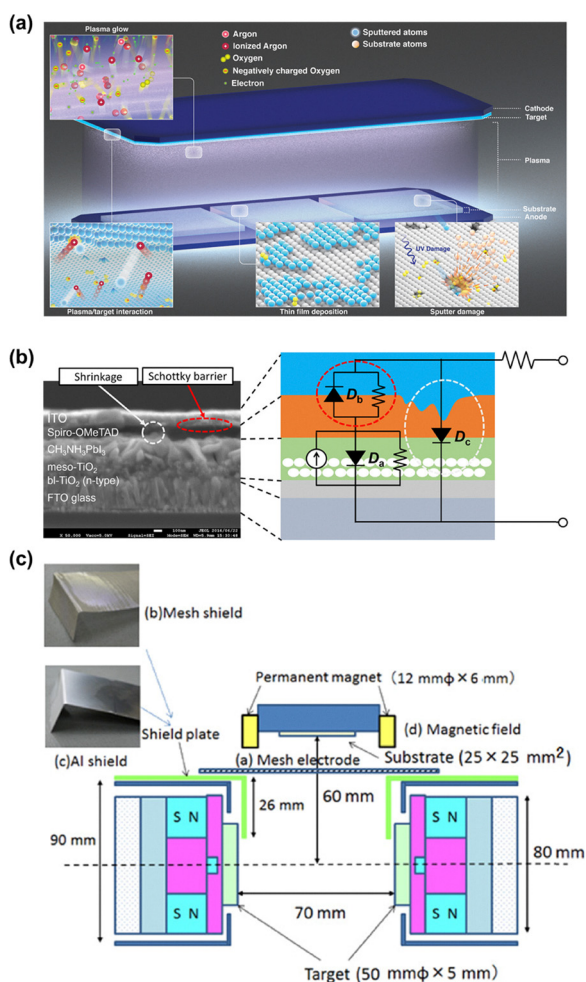


Fig. 15 (a) Schematic illustration describing sputter process. Reproduced with permission.¹⁶⁵ Copyright 2006, Elsevier Inc., (b) cross-sectional SEM image of the perovskite device and its circuit analysis causing additional path due to TCO sputter. Reproduced with permission.¹⁶⁶ Copyright 2016, American Chemical Society shield and shield plates. (c) Low damage sputter system by using face target, mesh shield and shield plates. Reproduced with permission.¹⁶⁸ Copyright 2016 The Japan Society of Applied Physics.

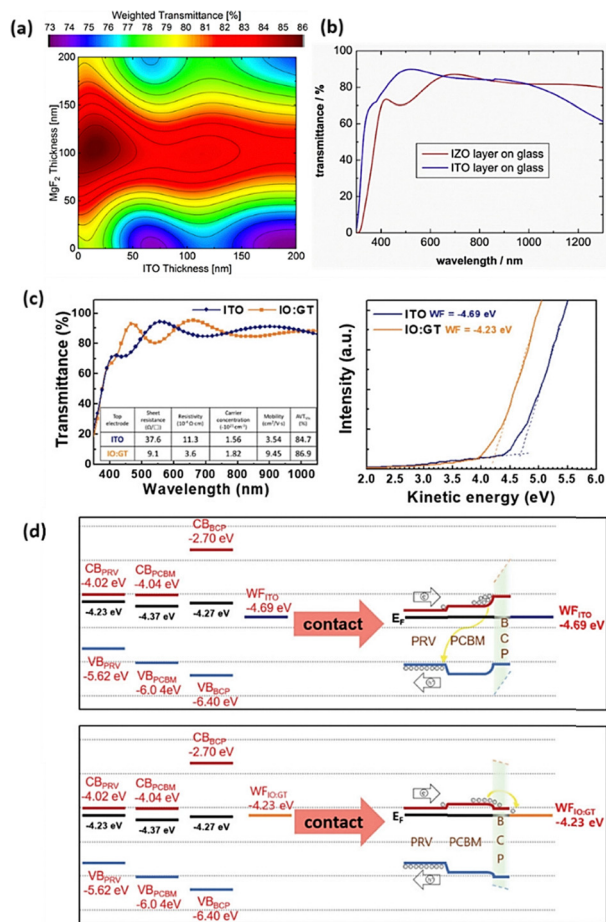


Fig. 16 (a) Simulated transmittance depending on thicknesses of IZO and MgF₂ using the transfer-matrix method. Reproduced with permission.¹⁶⁹ Copyright 2019 American Chemical Society. (b) Transmittance spectra of IZO and ITO. Reproduced with permission.¹⁷⁰ Copyright 2017 Elsevier. (c) Transmittance spectra (inset table: optical and electrical properties) and UPS spectra of ITO and gallium- and titanium-doped indium oxide (IO:GT) thin films. (d) Energy diagram of the perovskite device using ITO and (IO:GT). Reproduced with permission.¹⁷¹ Copyright 2022 Wiley-VCH GmbH.

like ITO does, and it also exhibits higher transmittance at longer wavelengths (above 1000 nm) compared to ITO (Fig. 16(b)). In addition to the use of IZO instead of ITO, a method of applying titanium and tantalum-doped ITO to increase transparency in perovskite solar cells has also been reported (Fig. 16(c) and (d)).¹⁷¹ Therefore, diversifying the materials used can be a good approach to increase the transparency of TCO.

The generated photocarriers are collected *via* lateral current of the TCO, hence the sheet resistance directly affects the performance of the solar cell. This becomes particularly important as the active area increases, as the loss from the TCO is proportional to sheet resistance and active area.¹⁷² To overcome the sheet resistance of the TCO, a variety of alternative materials have been suggested such as carbon-based transparent conducting electrodes (TCE)s, metal nanowires and ultra-thin metal.¹⁷³ However, due to suboptimal optoelectronic characteristics of these alternative materials, TCOs are widely being used as main components in high-efficiency perovskite/Si

tandem.^{122,174} Thus, to achieve high-performing top-perovskite cells in tandem, a practical approach could be to use TCOs in combination with a well-designed metal-mesh or metal busbar to improve the charge collection and alleviate the sheet resistance of the TCOs, particularly for large-area devices. In the following section, a detailed review of the metal-mesh or metal busbar will be provided.

The energy level of TCOs should be carefully considered, for example, a lower work function for the TCO on top of the ETL and a deeper work function for the top of the HTM. One typical method to adjust the energy level of TCOs is by varying the oxygen level, with a higher oxygen flow rate typically resulting in a deeper work function. ITO and IZO have different work functions, with ITO having a slightly shallower work function (4.6–4.7 eV) than IZO (~5.1 eV).^{175,176} Additionally, doping other elements into the indium-based oxide target also affects the energy level. For instance, Yoon *et al.* reported that gallium and titanium-doped indium oxide (IO:GT) showed a work function of –4.23 eV, which is shallower than that of ITO (–4.69 eV).¹⁷¹ Due to the favourable energy level for electron extraction, the modified TCO (IO:GT) allows for sputtering of the TCO on top of the PCBM/BCP layer, where typical ETLs are used in opaque cell structures with much higher PCE (17.9%) than when using ITO (PCE of 8.59%) and comparable PCE when using Ag (19.86%) (Fig. 16(c) and (d)). This study suggests that the additional process, such as ALD for TCO buffer layer, can be replaced by thermal vacuum evaporation *via* TCO engineering, which could be beneficial for mass production in terms of takt time. Recent developments have also explored alternative TCO compositions to enhance transparency and optical management. In particular, indium-gallium-tin oxide (IGTO) has been reported to exhibit improved near-infrared transparency while maintaining sufficient electrical conductivity, offering an effective route to minimize optical losses in perovskite/Si tandem configurations.¹⁷⁷

Part II: external components of perovskite/Si tandem solar cells

Encapsulation

The hygroscopic and volatile nature of organic components in perovskite layers compromises moisture resistance and thermal stability, as exposure to humidity and heat leads to material decomposition. Effective encapsulation is therefore essential for ensuring long-term device performance and requires materials and processes tailored to the sensitivity of perovskite/Si tandem structures. Conventional encapsulation methods, such as heat lamination of polymer sheets, utilize high temperatures to melt and crosslink the encapsulant.¹⁷⁸ Ethylene-vinyl acetate (EVA), widely used in Si photovoltaics for its transparency and mechanical durability (Fig. 17(a)), exhibits a high processing temperature (>140 °C) that can degrade heat-sensitive perovskite layers.^{179,180} Additionally, EVA generates acidic by-products during operation, accelerating perovskite decomposition. Surlyn, another common encapsulant for

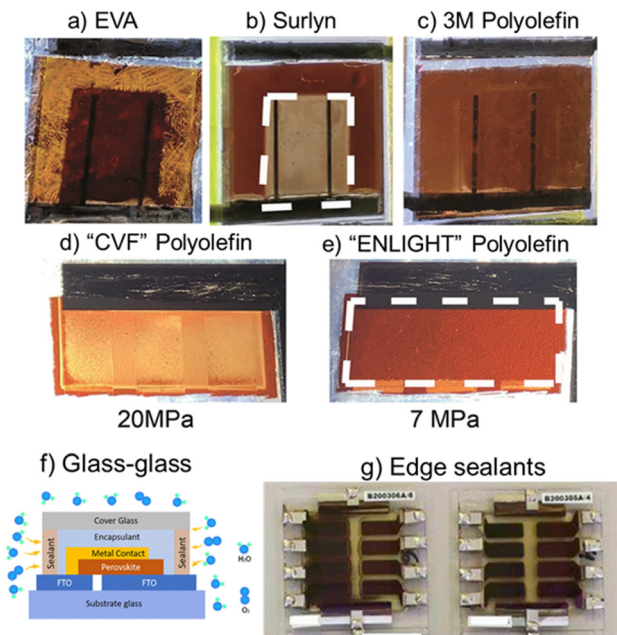


Fig. 17 Example of encapsulation (a)–(e) various polymers. Reproduced with permission.¹⁸¹ Copyright 2018, The Royal Society of Chemistry. (f) Glass–glass,¹⁸⁶ Copyright 2022, American Chemical Society. Licensed under CC BY-NC-ND 4.0. (g) Edge sealants. Reproduced with permission.¹⁸⁴ Copyright 2024 International Solar Energy Society.

Si cells, suffers from poor adhesion to perovskite surfaces and thermal expansion mismatch, which can lead to delamination and moisture ingress (Fig. 17(b)).^{181–183} In contrast, polyolefin has emerged as a promising low-temperature alternative (Fig. 17(c)). Checharoent *et al.* demonstrated that polyolefin, processed at 120 °C, offers improved compatibility with perovskite layers and can effectively replace EVA in perovskite/Si tandem solar cell encapsulation.¹⁶⁷

Mechanical properties are also crucial for maintaining structural integrity during thermal cycling. Encapsulants with higher elastic moduli, such as CVF (20 MPa) and ENLIGHT (7 MPa), are more susceptible to cracking and delamination, leading to discoloration and device degradation after prolonged aging (*e.g.*, 1000 hours at 85 °C) (Fig. 17(d) and (e)). Encapsulants with a lower modulus have shown better mechanical compliance, preserving device integrity under stress. Promising alternatives to EVA include thermoplastic polyurethane (TPU) and polytetrafluoroethylene (PTFE), which offer low-temperature processability and better thermal expansion matching with perovskite layers, reducing the risk of mechanical and thermal failure.¹⁸⁵

Beyond polymer-sheet encapsulants, glass-based packaging has been investigated to further improve durability (Fig. 17(f)).^{186,187} Glass–glass encapsulation provides superior resistance to moisture and oxygen ingress together with high mechanical stiffness, ensuring excellent outdoor stability, although its weight and rigidity limit applications in lightweight or flexible modules.^{188,189} Recent work has suggested low-temperature and low-stress glass–glass lamination schemes tailored for perovskite tandems to reduce thermal stress on the top absorber.¹⁸⁹ In comparison glass–polymer configurations, which replace the

rear glass with a polymeric backsheets, offer reduced weight and greater flexibility.¹⁸⁶ However, their barrier performance strongly depends on edge sealing, as lateral diffusion pathways dominate moisture ingress. To this end, edge sealants play a decisive role. Butyl rubber (PIB) exhibits exceptionally low WVTR and strong adhesion, making it the most effective choice,¹⁹⁰ while epoxy-based sealants offer elasticity and thermal shock resistance at the cost of higher permeability (Fig. 17(g)).¹⁸⁴

Hybrid encapsulation techniques further improve stability by combining the flexibility of organic polymers with the barrier properties of inorganic materials. Multilayer designs incorporating polymers with thin oxide layers (*e.g.*, Al₂O₃ or SiO₂) provide enhanced moisture protection and structural robustness.^{191,192} Ionogel-based encapsulants offer additional tunability in mechanical, optical, and chemical properties through their liquid-phase matrix. Incorporating lead-absorbing monomers into ionogels mitigates lead leakage while enhancing chemical durability and mechanical cohesion.¹⁹³ In summary, these attempts highlight the need for encapsulation strategies that simultaneously account for thermal stress, humidity exposure, and mechanical durability. Polymer-based materials such as TPU, PTFE provide low temperature compatibility, glass–glass structures deliver the most robust environmental protection, and hybrid approaches including inorganic barriers promise to combine durability with additional safety functions. The optimal encapsulant must therefore be selected in close relation to device stack sensitivity and target module applications.

Busbar

Scalability of large-area perovskite/Si tandem modules is constrained by the relatively low current density of perovskite top cells ($\sim 20 \text{ mA cm}^{-2}$), which is approximately half that of conventional Si bottom cells ($\sim 40 \text{ mA cm}^{-2}$). This mismatch complicates current balancing and reduces tandem efficiency, particularly as device area increases. In addition to electrical constraints, the migration of mobile ions and volatile components in perovskite materials can lead to interfacial degradation and layer delamination, further compromising long-term operational stability.^{94,194} To address these issues, optimizing busbar design is essential for efficient current collection, mechanical reliability, and compatibility with thermally sensitive perovskite layers. Metal grids, the primary component of busbar architecture, must provide low-resistance electrical pathways while remaining compatible with the TCO layer and scalable to large-area fabrication.¹⁹⁵ Although thermal evaporation is commonly used to fabricate metal grids in small-area tandem cells, its low throughput limits its suitability for commercial-scale production. In contrast, screen printing and roll-to-roll processing offer higher throughput and scalability but may suffer from increased contact resistance when interfaced with TCO layers. Kamino *et al.* reported that printed grids exhibited elevated series resistance, necessitating further refinement to maintain charge collection efficiency in scaled devices.¹⁹⁶

Additionally, shingled cell interconnection has been adapted to perovskite/Si tandems, reducing optical shading and enabling higher packing density at the module level.¹⁹⁷

Silver paste-based grids are widely adopted due to their high conductivity and proven reliability in current collection. However, their high sintering temperatures ($>800\text{ }^{\circ}\text{C}$) are incompatible with perovskite materials. As a cost-effective alternative, copper offers lower raw material costs and potential for scalable processing. Yet, its susceptibility to oxidation can degrade electrical properties, requiring protective coatings or encapsulation layers. Although such strategies improve stability, they increase process complexity and may limit manufacturability. In addition, copper's higher intrinsic resistivity relative to silver may reduce overall conductivity, though this limitation can be mitigated through grid optimization—such as increasing line thickness or reducing spacing. The fabrication process for Si solar cells employing copper-based busbars typically involves screen-printing silver paste on the front side to form fine fingers and busbars, while a low-temperature Cu paste is applied to the rear side. The Cu rear contact is then co-fired and cured in a nitrogen atmosphere to complete the metallization. A comparison of Ag- and Cu-based tabbing structures reveals that the metal-semiconductor interface presents a critical region for carrier recombination. This area is particularly sensitive to interface passivation quality and doping gradients, which can exacerbate recombination losses. These considerations underscore the importance of careful junction engineering when integrating Cu metallization into perovskite/Si tandem cells (Fig. 18).^{198–200} Recent tandem-oriented metallization studies further emphasize co-optimizing busbar geometry and low-thermal-budget processing to suppress series/contact-resistance growth during scale-up.²⁰¹

Scalability of large-area perovskite/Si tandem modules is constrained by the relatively low current density of perovskite top cells ($\sim 20\text{ mA cm}^{-2}$), which is approximately half that of conventional Si bottom cells ($\sim 40\text{ mA cm}^{-2}$). This mismatch complicates current balancing and reduces tandem efficiency, particularly as device area increases. In addition to electrical constraints, the migration of mobile ions and volatile components in perovskite materials can lead to interfacial

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Beyond printing, scalable low-temperature metallization routes have also been demonstrated, including electroplated Cu busbars on 22.5 cm^2 two-terminal perovskite/Si tandems using an ALD- Al_2O_3 mask with a screen-printed Ag seed.²⁰³

Moreover, the environmental stability of busbars remains a critical issue. Under thermal cycling and humid conditions, metallic contacts are prone to corrosion and the adhesion between electrodes and encapsulant can deteriorate. These effects make pathways that diminish the fill factor and module stability.²⁰⁴ To address this, busbar design should be carefully integrated with encapsulation strategies to guarantee efficient charge collection and improved resistance against moisture and heat induced degradation.^{205,206} Table 3 summarizes representative specifications for screen-printed front-contact designs. Key parameters include a busbar width of $500\text{--}800\text{ }\mu\text{m}$, finger width of $30\text{--}50\text{ }\mu\text{m}$, and aspect ratios between 0.2 and 0.8, with a total metal coverage maintained below 5%. Minimizing finger and contact resistivity while ensuring TCO compatibility is critical to maintaining high fill factors and low series resistance in large-area modules.

Silver paste-based grids are widely adopted due to their high conductivity and proven reliability in current collection. However, their high sintering temperatures ($>800\text{ }^{\circ}\text{C}$) are incompatible with perovskite materials. As a cost-effective alternative, copper offers lower raw material costs and potential for scalable processing. Yet, its susceptibility to oxidation can degrade electrical properties, requiring protective coatings or encapsulation layers. Although such strategies improve stability, they increase process complexity and may limit manufacturability. In addition, copper's higher intrinsic resistivity relative to silver may reduce overall conductivity, though this limitation can be mitigated through grid optimization—such as increasing line thickness or reducing spacing. The fabrication process for Si solar cells employing copper-based busbars typically involves screen-printing silver paste on the front side to form fine

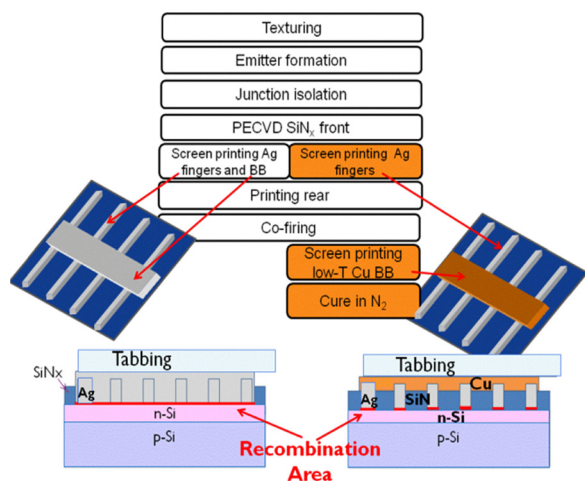


Fig. 18 Example of a Si solar cell using Cu busbar. Reproduced with permission.²⁰² Copyright 2014, Elsevier Inc.

Table 3 Characteristics of a metal grid applied by screen printing

Parameters	Unit	Screen-printed front contacts
Busbar width (w_b)	μm	500–800
Finger width (w_f)	μm	30–50
Average finger height (h_f)	μm	10–15
Aspect ratio (height to width)	—	0.2–0.8
Finger spacing (S)	mm	3–3.5
Metal coverage (finger + busbar)	%	4–5
Finger resistivity (ρ_f)	$\Omega\text{ cm}$	2–3
Contact resistivity (ρ_c)	$\text{m}\Omega\text{ cm}^2$	≤ 1
TCO sheet resistance (R_{sh})	$\Omega\text{ sq}^{-1}$	~ 50
Temperature of contact formation (T)	$^\circ\text{C}$	Low (< 150)

fingers and busbars, while a low-temperature Cu paste is applied to the rear side. The Cu rear contact is then co-fired and cured in a nitrogen atmosphere to complete the metallization. A comparison of Ag- and Cu-based tabbing structures reveals that the metal–semiconductor interface presents a critical region for carrier recombination. This area is particularly sensitive to interface passivation quality and doping gradients, which can exacerbate recombination losses. These considerations underscore the importance of careful junction engineering when integrating Cu metallization into perovskite/Si tandem cells (Fig. 17).^{198–200} Recent tandem-oriented metallization studies further emphasize co-optimizing busbar geometry and low-thermal-budget processing to suppress series/contact-resistance growth during scale-up.²⁰¹

Part III: fabrication, cost and industrial considerations

For the successful commercialization of perovskite/Si tandem solar cells, achieving low production costs is crucial. Production costs are influenced by material expenses and process time, known as takt time.²⁰⁷ Moreover, perovskite/Si tandem modules need to scale up to at least 166 mm by 166 mm, considering the widespread use of industrial Si wafers exceeding M6 size. However, the best power conversion efficiency (PCE) achieved for large area modules is 17.9%, compared to 25.7% for small unit cells, resulting in a PCE gap of approximately 30% in the single-junction perovskite solar cell section.² Furthermore, the scarcity of reports on perovskite/Si modules larger than M6 size suggests that upscaling the top perovskite solar cell is challenging.²⁰⁸ Therefore, overcoming these challenges with a scalable process and appropriate takt time is crucial for the successful commercialization of perovskite/Si tandem solar cells.

Takt-time

The majority of high-efficiency perovskite/Si tandem solar cells reported to date have employed a p–i–n structure for the top perovskite solar cell, with a configuration consisting of ITO/ NiO_x /SAM/Perovskite/Interlayer/ C_{60} / SnO_x /IZO/busbar/AR as shown in Fig. 19(a), albeit with some variations across studies.^{122,174,209,210} To achieve this configuration, a cycle of processes, including sputter-coating, blade coating, annealing, blade coating-heating,

thermal evaporation, ALD-sputter, screen printing, and thermal evaporation, is required as illustrated in Fig. 18(b). However, due to the alternative use of solution-based and vacuum-based approaches, this set of processes could increase the interval time for loading and unloading samples into and out of the vacuum chamber, resulting in significant delays and inefficiency. This issue is especially critical for large-scale manufacturing, where takt time plays a crucial role in the overall efficiency and profitability of the production process. Thus, it is expected that a consistent coating process, such as a fully blade/slot-die/inkjet/or fully vacuum based process would be ultimately required for the successful manufacturing process.

Continuous scalable solution processes

To achieve a consistent coating process suitable for large-scale mass production, the air-processability of the coating method, whether blade, slot-die, spray coating, or ink-jet printing, is crucial. Most studies PSCs have employed spin coating in inert gas-filled glove boxes due to the sensitivity of the perovskite and other layers to ambient environments such as moisture and oxygen. Thus, manufacturing top PSCs for the perovskite/Si tandem applications in ambient or dry-air conditions can cause variations in film characteristics and low reproducibility, posing a great challenge for commercialization.^{211,212} Moreover, the commonly used antisolvent dripping process may not be suitable for industrial mass production due to hazardous vapors and cost. In addition, in order to achieve reproducible and high-throughput processing, precursors in solution stock must have “weeks-long shelf lives” that guarantee identical film properties.²¹³ These issues pose a significant challenge as the direct application of spin coating in glove boxes to upscaled coating processes in ambient air may not be straightforward.

All-vacuum process

As a successful precedent can be observed in the OLED industry, a full-vacuum based process can be a promising solution for the perovskite top cell. The widely used structure for a top PSC in Fig. 18(b), ITO/ NiO_x /SAM/Perovskite/LiF/ C_{60} / SnO_2 /IZO/LiF/Ag, is vacuum processable in principle.^{152,214,215} Since Liu *et al.* reported the vacuum processed perovskite for the solar cell in 2012, numerous groups have reported on vacuum processed PSCs, including all-vacuum processed PSCs.²¹⁶ The advantages of vacuum processing for manufacturing optoelectronic devices include scalability, uniformity of the films, controllability of thickness, and freedom from environmental interference, making it a suitable process for industrial application. An important advantage, particularly for the application in perovskite/Si tandem, is that the top PSC cell can be conformally deposited on the textured Si surface. This characteristic results in optimizing the optical path of the tandem cell without requiring additional polishing for flattening the Si surface.^{46,217}

However, further improvements are required for vacuum-processed perovskite solar cells (PSCs), including those used for the top cell in the perovskite/Si tandem, from various aspects. Firstly, the power conversion efficiencies (PCEs) of vacuum-processed PSCs are generally lower than those of solution-

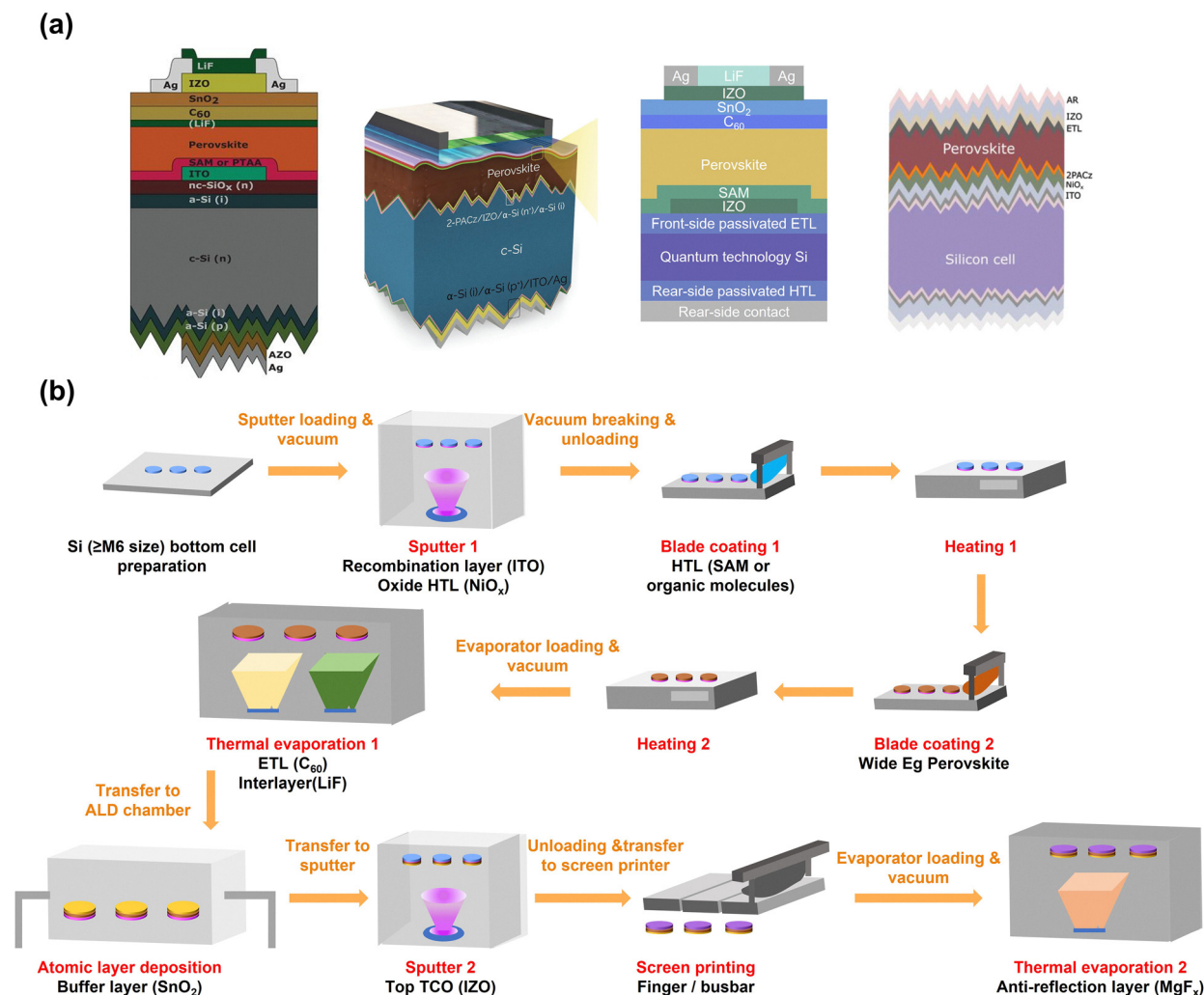


Fig. 19 (a) Schematic illustration of typically reported device structure for perovskite/Si tandem. Reproduced with permission.^{122,174,209,210} Copyright 2020, American Association for the Advancement of Science, Copyright 2022, American Association for the Advancement of Science, Copyright 2022, American Chemical Society, Copyright 2022, Wiley-VCH GmbH respectively. (b) Schematic illustration of an example of a scalable fabrication process for a representative structure of a top perovskite solar cell in Si tandem configuration.

processed PSCs. Considering the breakthrough in PCE achieved through various additive engineering in solution processed PSCs, it could be reasonable to assume that similar efforts will be required to improve the PCE of vacuum processed PSCs. However, due to factors such as the high cost of equipment resulting in a high entry barrier for the vacuum process, there are relatively few papers and researchers in the field, which may have resulted in fundamental technological development may have been insufficient so far. In addition, compared to the solution process, it is challenging to incorporate numerous additives in the vacuum process since the number of precursors is limited by the number of sources (crucibles) in the chamber. Furthermore, it has not been validated whether perovskite precursors, especially organic halides, can be stably evaporated with the given conditions for mass production using linear-type sources that evaporate the sources for a long time at a fast rate. Therefore, numerous challenges and areas for further study exist in the development of vacuum-processed PSCs, particularly for the top cell of perovskite/Si tandems.

Cost

The market viability of PV technologies is heavily influenced by their LCOE, a key metric representing the average cost of electricity generation over a system's lifetime. LCOE is calculated using eqn (3) as the ratio of total lifetime costs to total electricity generated, incorporating the initial investment (I), operating and maintenance costs (OM), discount rate (r), electricity output (E), and annual degradation rate (d).²¹⁸

$$\begin{aligned} \text{LCOE} &= \frac{\text{Total lifetime cost}}{\text{Total lifetime electricity production}} \\ &= \frac{I + \sum_{i=0}^N \frac{OM}{(1+r)^i}}{\sum_{i=0}^N \frac{E(1-d)^i}{(1+r)^i}} \end{aligned} \quad (3)$$

Perovskite/Si tandem solar cells generally exhibit higher PCE than single-junction cells, which enables greater energy yield

per unit area and helps reduce the LCOE despite their relatively more complex and costly manufacturing. While tandem devices currently report an LCOE of approximately \$0.32 and \$0.28 per kWh, respectively (Fig. 20(a)), these estimates assume utility-scale systems operating under standard irradiance conditions over a 25-year lifetime.^{194,195} The superior energy output of tandem configurations offsets their slightly higher production costs, making them more cost-effective over the system's operational period. Importantly, McGovern *et al.* showed through techno-economic modeling that perovskite/Si tandem solar cells can achieve cost competitiveness with c-Si PV when an efficiency of $\geq 30\%$, annual degradation rate below 2% per year, and module manufacturing costs of ~ 62.5 € per m^2 are simultaneously achieved.²¹⁹

Efficiency improvement is one of the most direct strategies to reduce LCOE. For instance, Sofia *et al.* demonstrated that increasing the efficiency of the perovskite top cell from 18% to 24% lowered the LCOE from \$0.32 to \$0.22 per kWh by increasing total electricity output.²²⁰ In addition to efficiency gains, reducing material and processing costs is essential. Expensive charge transport materials such as PCBM, spiro-OMeTAD, and PTAA significantly contribute to overall production costs (Fig. 19(b)).²¹⁸ Among them, PTAA, widely used as a hole transport layer, incurs particularly high material costs. To address this, researchers are developing cost-effective alternatives that retain comparable performance. Furthermore, scalable deposition directly shifts module MSP and thus slot-die coating offers very high material utilization ($\approx 99\%$),²²¹ and

slot-die-processed perovskite/Si tandem solar cells reached 25.9% efficiency over 16 cm^2 area.²²² At the line level, a fully roll-to-roll printed perovskite projects $\approx \$0.7$ per W at 1 000 000 m^2 per year throughput.²²³

Encapsulation technologies also play a role in minimizing LCOE. Advances in hybrid and ionogel-based encapsulants enhance stability by reducing moisture ingress and degradation, thereby extending operational lifetime. For the design targets, robust PSC/PSM encapsulation typically pursues oxygen transmission rate (OTR) $\approx 10^{-4}$ – 10^{-6} $cm^3 m^{-2}$ per day atm and water vapor transmission rate (WVTR) $\approx 10^{-3}$ – 10^{-6} $cm^3 m^{-2}$ per day.²²⁴ Ionogel encapsulation has passed International Electrotechnical Commission (IEC) 61 215 damp-heat/thermal-cycle tests and suppressed Pb leakage to undetectable levels after hail plus 24 h water soak;²²⁵ on large-area devices, 110 cm^2 modules retained $\sim 84\%$ of initial PCE after 1000 h at 85 °C (air), quantitatively lowering Annual Degradation Rate (ADR) inputs in LCOE.²²⁶

Concerns over intellectual property, supply chain, and toxicity regulations

PSCs are advancing rapidly toward industrial deployment, yet their commercialization depends as much on regulatory and economic considerations as on power-conversion efficiency. A central non-technical hurdle is intellectual property. Major research institutes and companies have assembled extensive

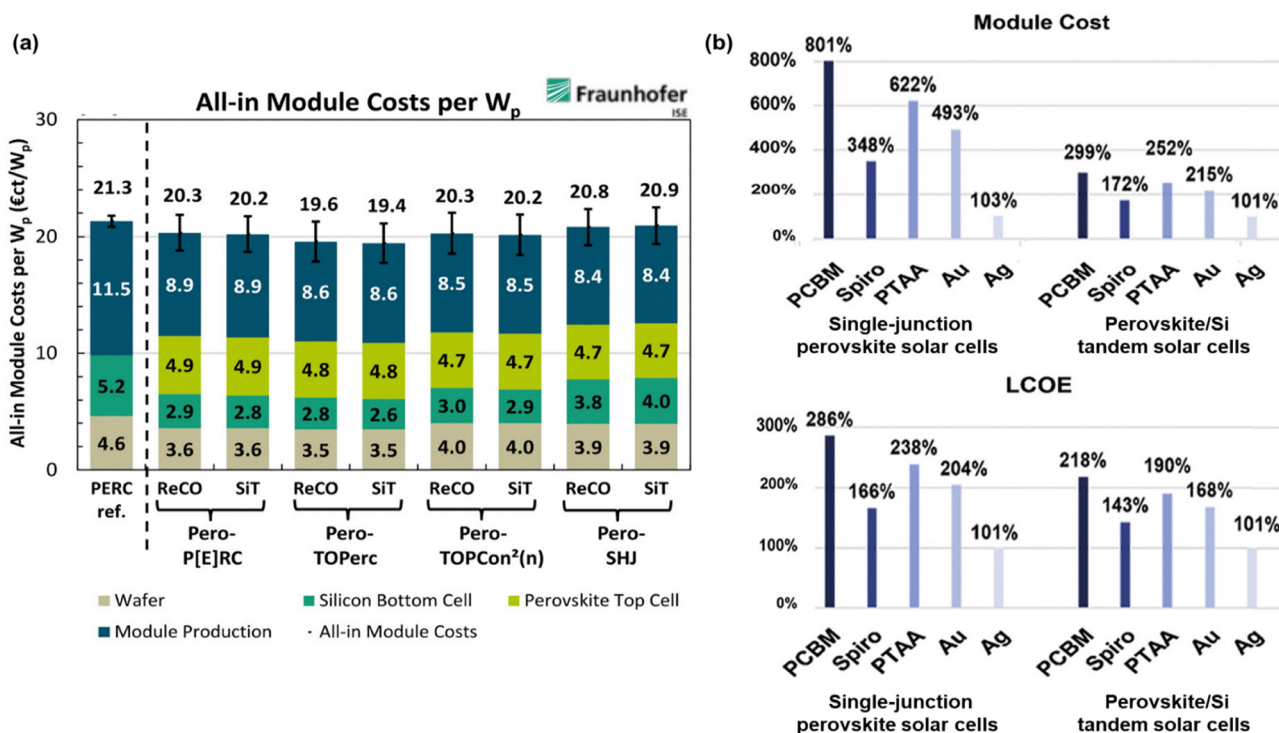


Fig. 20 (a) Module costs for all perovskite/Si tandem solar cells. ReCO: TCO-based RL, SiT: Si-based TJ. Reproduced with permission.¹⁹⁴ Copyright 2020, John Wiley & Sons Ltd, (b) perovskite single-junction and perovskite/Si solar cell module production cost and LCOE ratio by material. Reproduced with permission.²¹⁸ Copyright 2017, The Royal Society of Chemistry. Elsevier Inc.

patent portfolios that cover critical aspects of cell architecture, interfacial engineering, and tandem integration. Oxford PV, for example, has secured broad claims on perovskite–silicon tandem designs and has begun licensing agreements with large manufacturers, illustrating that freedom-to-operate analyses and potential cross-licensing will be essential for any entrant seeking to scale production. Competitive patent filings from firms such as Microquanta, LG, and Panasonic further reinforce the need for early legal strategies to avoid costly infringement disputes.

Commercial viability also hinges on supply chain resilience. Although perovskites can be processed at low temperatures, the materials ecosystem is far from trivial. Cesium, indium, high-purity lead iodide, and sophisticated encapsulation films are required at scale, and several of these precursors face potential scarcity or geographic concentration. A recent U.S. Department of Energy review highlights the vulnerability of global solar manufacturing to regional bottlenecks and price shocks, particularly because glass, substrates, and specialty chemicals remain dominated by a few suppliers. Terawatt-scale modeling likewise identifies cesium and indium as elements requiring careful resource planning for sustainable perovskite expansion.²²⁷

Toxicity associated with lead, the key light-absorbing cation in state-of-the-art perovskites, remains the most visible societal concern. Regulatory assessments show that uncontrolled lead release during module damage or end-of-life disposal can exceed drinking-water action levels, even though well-encapsulated devices can meet U.S. TCLP hazardous-waste limits.²²⁸ Mitigation strategies include robust edge sealing, chemical sequestration of lead within the perovskite lattice, and closed-loop recycling processes capable of recovering both lead and rare transport materials.²²⁹ These intertwined issues are now as decisive as cell efficiency in determining when and where perovskite modules will achieve true commercial impact. In parallel, emerging approaches such as chemisorption with functional groups (thiol, sulfonate, and phosphate) provide promising pathways to immobilize Pb²⁺ at the molecular level, complementing system-level recycling strategies. Together, these intertwined issues highlight that addressing non-technical barriers is as decisive as achieving record efficiencies, and they will strongly influence the timeline and geography of perovskite/Si tandem commercialization.²³⁰

Part IV: outlook and conclusions

Perovskite/Si tandem solar cells stand at the forefront of next-generation photovoltaics as they exceed the efficiency ceiling of single-junction devices. This review has addressed both recent advances and persistent bottlenecks across Si bottom cell configuration, wide-bandgap perovskites, ICLs, TCOs, encapsulation strategies, busbar design, scalable manufacturing processes, and levelized cost. Sustained progress in each of these areas will be crucial for translating lab-scale demonstrations into commercially viable modules.

In tandem integration, the selection of Si bottom cells strongly influences both device performance and

manufacturability. PERC, while dominant in current production, imposes a practical limit on achievable tandem efficiency. TOPCon provides improved surface passivation and maintains compatibility with high-throughput industrial processing while parasitic absorption in poly-Si layers should be minimized. SHJ has achieved record efficiencies through effective interface passivation, but its reliance on a-Si:H-based passivation and doped contact layers leads to thermal instability, which poses a challenge to scale up. Developing passivation schemes with a low thermal budget can provide a viable solution to the scale-up challenge. For the perovskite top cell, efficiency has already reached competitive levels, but stability remains the primary hurdle. Long-term reliability is undermined by halide segregation, defect evolution, and ambient-induced degradation. Defect management in both the bulk and interfaces is therefore critical to addressing the stability challenge: in the bulk, achieving high crystallinity and implementing novel defect passivation strategies are essential to suppress defect formation while at the interfaces introducing passivation layers can enhance material stability and extend device lifetime. In the ICL design, efficient carrier recombination should be achieved while minimizing optical and electrical losses. RJs, including TCOs offer high transmittance, but their limited conductivity can lead to carrier accumulation, voltage loss, and lateral shunting. Introduction of transport layers that can meet optical index matching can limit lateral carrier conduction while preserving transparency. TJs, in contrast, present high vertical conductivity through quantum tunneling, but free-carrier absorption and defect-assisted recombination can cause optical loss. Such losses can be alleviated through precise doping control and effective interface passivation. For the top electrodes of perovskite/Si tandem solar cells, TCOs are the most widely used, but their deposition by sputtering can damage the underlying layers. The sputter-induced damage is usually mitigated by protective buffer layers and optimized sputtering conditions. A further challenge is to balance optical transparency with electrical conductivity, which is often managed by incorporating metal-mesh grids to reduce resistive losses. Moreover, tuning the TCO work function to achieve suitable energy-level alignment with transport layers is essential for effective charge extraction.

For external components, robust encapsulation is vital to ensure long-term durability. In busbar design, adopting Cu-based metallization improves current collection by lowering series resistance and resistive heating, while also supporting scalability for large-area modules through high conductivity and mechanical reliability under thermal cycling, which in turn enables cost reduction. As tandem modules move toward commercial deployment, scalable and low-cost fabrication strategies will be indispensable. Solution-based processes offer high throughput, whereas vacuum-based techniques provide superior uniformity and film integrity. Thus, hybrid manufacturing approaches may integrate these advantages to meet industrial requirements, but continued advances in process engineering will be necessary to achieve cost targets without compromising long-term device performance.

Globally, perovskite solar cell technology has already entered the commercialization stage, led most prominently by

China and followed by the United States, Europe, and other regions in Asia. Microquanta is currently operating a production line with an annual capacity of approximately 100 MW dedicated to building integrated photovoltaic (BIPV) applications. GCL has completed a 1 GW production line and is now expanding toward a 2 GW facility capable of fabricating tandem modules with an area of 2.76 m². Other Chinese companies such as Utmolight and Renshine Solar have each installed production lines of around 150 MW and have begun commercial deployment. In Japan, the government has pledged 1.5 billion USD to support mass production of perovskite solar cells in collaboration with Sekisui Chemical, targeting completion of a 150 MW line by 2027 and a 1 GW line by 2030. In Korea, Hanwha Q Cells has established a 40 MW line and announced plans for full-scale production in the second half of 2026. In the United States, companies including Cubic PV, Swift Solar, and First Solar have launched or announced pilot line projects as they enter the race for commercialization. Taken together, these developments indicate that perovskite solar cells are now firmly on a commercial trajectory, with the decisive phase of widespread market adoption expected to emerge well within the next three years, and potentially sooner. Alongside these advances, rigorous accelerated lifetime testing, including ISOS protocols, damp heat exposure, and thermal cycling, as well as outdoor field validation under fluctuating humidity, temperature, and illumination conditions, remains essential to evaluate long-term operational stability. These studies provide benchmarks for certification and bankability, ensuring reliable projections for industrial deployment.

Conflicts of interest

There are no conflicts to declare.

Data availability

No primary research results, software or code have been included and no new data were generated or analysed as part of this review.

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