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Broader context

The realization of a sustainable hydrogen economy hinges on the ability to produce green hydrogen efficiently and affordably. Solar-driven water splitting represents an ideal solution, yet its widespread adoption is currently hindered by the high fabrication costs and mechanical rigidity of conventional inorganic photovoltaics. This review underscores the transformative potential of solution-processable tandem solar cells — comprising organic and perovskite light absorbers — as a next-generation power source for water electrolysis. By offering a synergistic combination of tunable optoelectronic properties, light weight, and compatibility with high-throughput manufacturing, these materials address the dual challenges of performance and cost. We provide a comprehensive roadmap linking fundamental material innovations with practical system engineering, such as catalyst integration and stability management. This work identifies the critical steps required to transition from laboratory-scale efficiency records to scalable, durable, and cost-competitive solar-to-hydrogen platforms; thereby accelerating the deployment of renewable fuel technologies in the global energy landscape.



1 **Solution processed tandem solar cells for water electrolysis**

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7



1 Abstract

2 Green hydrogen production *via* solar-driven water splitting stands as a pivotal technology for
3 transitioning to a carbon-neutral economy. While photovoltaic-electrochemical (PV-EC) systems offer
4 a promising pathway by driving water electrolysis as a practical route to solar hydrogen production,
5 single-junction solar cells typically fail to provide the sufficient photovoltage required to drive water
6 electrolysis without external bias. Tandem solar cells, particularly utilizing solution-processable
7 materials, have emerged as a viable solution to overcome this thermodynamic barrier while offering the
8 benefits of low-cost fabrication and mechanical flexibility. This review critically examines recent
9 advancements in solution-processed tandem architectures for unassisted solar water splitting,
10 classifying them into three primary categories: all-organic, all-halide perovskite, and perovskite-organic
11 hybrid tandem systems. We elucidate how bandgap engineering and spectral splitting strategies in these
12 architectures enable high open-circuit voltages (>1.6 V) and improved solar-to-hydrogen efficiencies,
13 reaching up to 17.8%. Beyond material optimization, this article highlights the importance of system-
14 level integration, including the engineering of interconnecting layers, geometric area matching between
15 PV and catalysts, and the development of earth-abundant electrocatalysts for minimizing kinetic
16 overpotentials. Finally, we provide a forward-looking perspective on overcoming critical bottlenecks
17 such as long-term stability in aqueous environments and scalability, aiming to guide future research
18 toward practical and economically viable solar fuel production.

19



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3 efficiently and affordably. Solar-driven water splitting represents an ideal solution, yet its widespread
4 adoption is currently hindered by the high fabrication costs and mechanical rigidity of conventional
5 inorganic photovoltaics. This review underscores the transformative potential of solution-processable
6 tandem solar cells — comprising organic and perovskite light absorbers — as a next-generation power
7 source for water electrolysis. By offering a synergistic combination of tunable optoelectronic properties,
8 light weight, and compatibility with high-throughput manufacturing, these materials address the dual
9 challenges of performance and cost. We provide a comprehensive roadmap linking fundamental
10 material innovations with practical system engineering, such as catalyst integration and stability
11 management. This work identifies the critical steps required to transition from laboratory-scale
12 efficiency records to scalable, durable, and cost-competitive solar-to-hydrogen platforms; thereby
13 accelerating the deployment of renewable fuel technologies in the global energy landscape.

14



1 Introduction

2 The escalating global climate crisis and the urgent need for energy decarbonization necessitate
3 a profound shift toward sustainable energy carriers. Among various options, green hydrogen, produced
4 *via* renewable energy, has emerged as a cornerstone for achieving a sustainable hydrogen economy.^{1,2}
5 Green hydrogen is essential because it offers a clean, high energy density medium for long-term energy
6 storage and a direct replacement for fossil fuels across hard-to-abate sectors like industry and heavy
7 transport.^{1, 3-6} Among the methods for producing green hydrogen, a representative approach involves
8 water splitting utilizing solar energy. This method directly converts the most abundant renewable
9 energy source, solar energy, into a chemical fuel. The conversion of solar energy into hydrogen is
10 primarily realized through the integration of a photovoltaic (PV) device and an electrochemical (EC)
11 cell, known as the PV-EC system. For water electrolysis, single-junction solar cells are fundamentally
12 limited by the Shockley–Queisser efficiency threshold,^{7, 8} and consequently suffer from insufficient
13 photovoltage output to drive the reaction, motivating increasing interest in tandem architectures,^{7, 9}
14 where subcells with complementary bandgaps are connected in series to deliver enhanced photovoltage
15 and overall conversion efficiency. In 2016, a notable milestone was achieved with the report of over
16 30% solar-to-hydrogen (STH) conversion efficiency using a triple-junction InGaP/GaAs/GaInNAsSb
17 solar cell, underscoring the potential of tandem devices for solar fuel production.¹⁰

18 However, despite their high efficiencies, incumbent tandem technologies face major
19 challenges, including high fabrication costs, material scarcity, and mechanical rigidity, which limit their
20 suitability for large-scale deployment.¹¹⁻¹⁴ These limitations have spurred increasing interest in solution-
21 processable semiconductors, particularly organic and metal halide perovskite light absorbers. Both
22 classes of materials offer important advantages — such as low-cost fabrication, facile bandgap
23 tunability, and compatibility with flexible substrates — making them attractive for the design of tandem
24 solar cells with distinctive manufacturing and mechanical benefits.¹⁵⁻²⁰ Tandem solar cells composed of
25 organic and/or perovskite subcells have therefore emerged as promising candidates for next-generation
26 PV technologies, particularly in applications requiring high photovoltage, lightweight structures, and

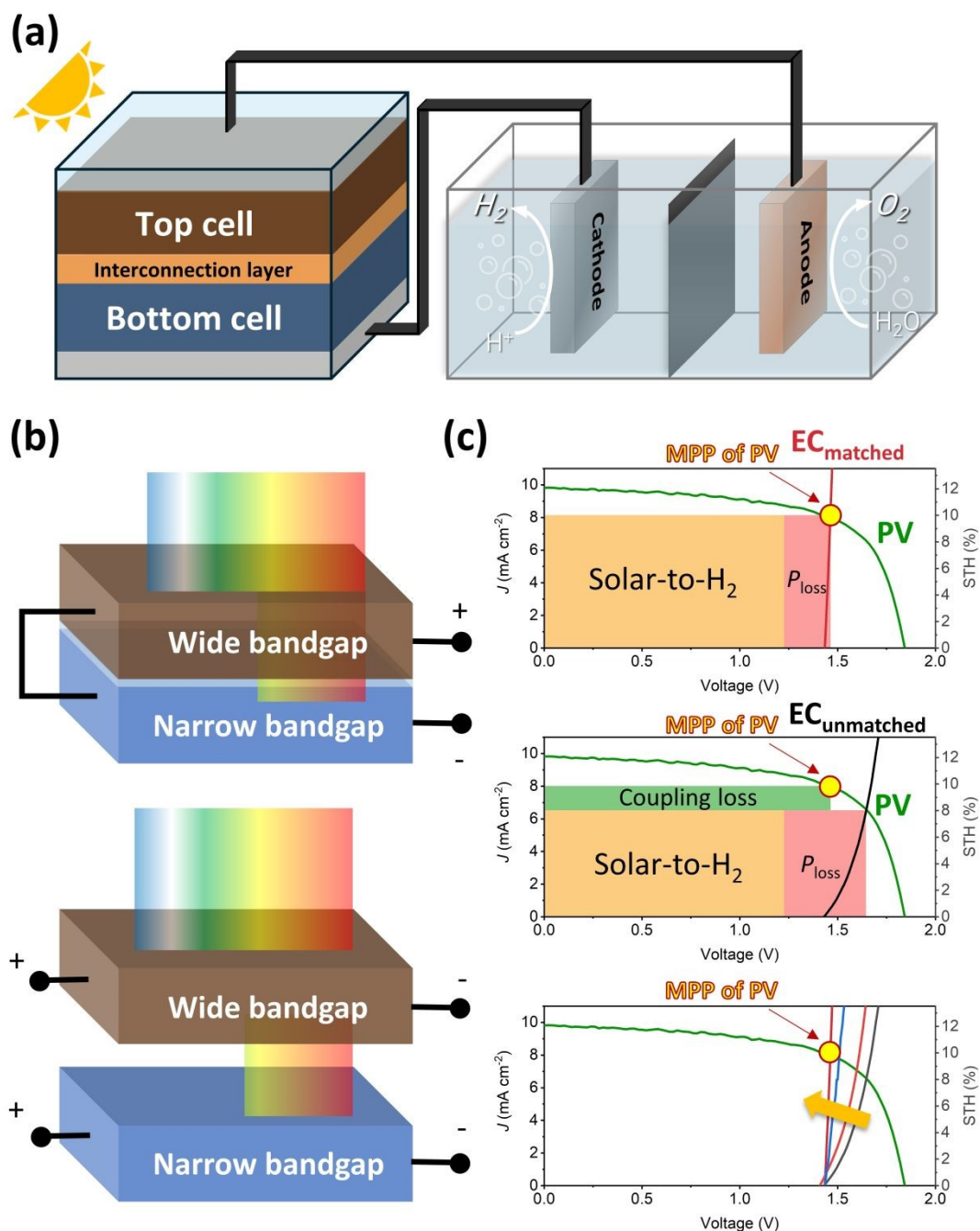


1 mechanical flexibility.^{21, 22} Nonetheless, these emerging solution-processable tandems face their own
2 significant hurdles that must be overcome for practical deployment. Key challenges include intrinsic
3 material instability — particularly the notorious sensitivity of perovskites to moisture and oxygen, and
4 the photodegradation of organic absorbers.²³⁻²⁷ Furthermore, achieving efficient charge recombination
5 at the interconnecting layer, minimizing interfacial voltage losses, and ensuring robust encapsulation
6 for long-term operation in aqueous environments remain critical bottlenecks.

7 This minireview summarizes recent progress in solution-processed tandem architectures
8 specifically aimed at solar-driven water electrolysis. We categorize the relevant systems into three
9 primary categories: organic, halide perovskite, and perovskite-organic hybrid tandems. By focusing on
10 advances in materials design, interfacial engineering, and system integration strategies for PV-EC
11 configurations, we elucidate the prevailing challenges for each technology. Finally, we provide a
12 forward-looking perspective on the research directions required to enable efficient, durable, and
13 scalable solar-to-hydrogen conversion.

14



1 **2 Water electrolysis and tandem solar cells: solution processable PV materials**

2

3 **Fig. 1** (a) Schematic illustration of a PV-EC water electrolysis system. (b) Tandem solar cell
 4 configurations: two-terminal (top) and four-terminal (bottom). (c) J - V characteristics of tandem PV
 5 (green) and EC in a two-electrode configuration. The MPP of PV is marked with a yellow circle and
 6 the operating point of EC is defined as the point where the J - V curves of OPV and EC intersect. The
 7 amount of power stored as hydrogen (Solar-to- H_2) and that wasted as kinetic loss (P_{loss}) are indicated.
 8 The coupling loss is generated (green box), when the MPP of PV and the operating point of EC is not
 9 matched. Adjusting this operating point to coincide with the MPP of PV (yellow arrow at the bottom)
 10 allows the generated power to be fully transferred to EC with a minimal loss.

11

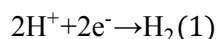


1 2.1 Water electrolysis systems for hydrogen production

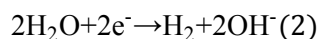
2 The PV-EC system is a modular approach for solar-to-hydrogen conversion, where a PV cell
3 is electrically connected to an EC cell (electrolyser) to drive water electrolysis using sunlight (**Fig. 1a**).
4 The PV cell in the PV-EC system only needs to provide a sufficient voltage difference across its external
5 terminals, regardless of the absolute potential alignment of its internal bands with the electrolyte. As
6 illustrated in **Fig. 1a**, the hydrogen evolution reaction (HER) takes place at the hydrogen evolution
7 catalyst (HEC)-loaded cathode, while the oxygen evolution reaction (OER) occurs at the oxygen
8 evolution catalyst (OEC)-loaded anode. In acidic and alkaline media, these half-cell reactions are
9 represented as follows:²⁸⁻³⁰

10 Cathodic reaction (HER)

11 In acidic electrolyte,

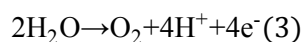


13 In alkaline electrolyte,

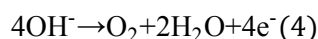


15 Anodic reaction (OER)

16 In acidic electrolyte,



18 In alkaline electrolyte,



20 The overall water electrolysis reaction can be expressed as:



22 The Gibbs free energy required to split water under standard conditions (25°C, 1 atm) is 237.2 kJ mol⁻¹,
23 corresponding to a thermodynamic potential of 1.23 V. However, in practical systems, the applied



1 voltage must exceed the thermodynamic limit of 1.23 V owing to kinetic constraints, often requiring
2 1.6–2.0 V for sustained water electrolysis.³¹ This increase in required voltage under non-ideal
3 conditions can be explained using the Nernst equation.

5 **2.2 Voltage requirements and detailed-balance rationale for tandem operation**

6 Unassisted water electrolysis requires not only the thermodynamic minimum voltage of 1.23
7 V but also an additional potential to overcome catalytic overpotentials and resistive losses. As a result,
8 a high photovoltage of approximately 1.8 V or higher is required for practical operation.^{32, 33} Under
9 these conditions, a single-junction photoabsorber inevitably faces a fundamental trade-off between
10 photocurrent generation and photovoltage output. Specifically, a narrow bandgap is advantageous for
11 achieving high photocurrent, whereas a wide bandgap is necessary to provide sufficient voltage for
12 unassisted water splitting.^{34, 35} From the perspective of the detailed-balance or Shockley-Queisser limit,
13 this trade-off imposes a strict upper bound on the efficiency of single-absorber systems and provides a
14 fundamental rationale for tandem architectures. By enabling broader utilization of the solar spectrum,
15 tandem configurations serve as an essential design strategy to simultaneously achieve high photocurrent
16 and high photovoltage.^{36, 37}

17 When intrinsic optical limitations such as thermalization and transmission losses are
18 considered together with practical operational losses including overpotential and power mismatch, the
19 STH conversion efficiency of single-junction PV-EC systems is generally limited to about 20%. The
20 tandem structure, proposed as a means to overcome this limitation, can improve the practical efficiency
21 to around 30%, and theoretical studies predict efficiencies exceeding 40%.^{9, 10} In particular, solution-
22 processed perovskite and organic absorbers have recently attracted considerable attention, although
23 their intrinsically low photovoltage restricts STH efficiency in single-junction systems to below 10-
24 15%. However, the incorporation of perovskite-based tandem configurations can increase the practical



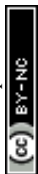
1 STH efficiency to approximately 18-20%, and further bandgap optimization is expected to yield
2 theoretical efficiencies approaching 30%.^{38, 39}

3 From this perspective, the practical advantages of tandem devices, including monolithic two-
4 terminal (2T) configurations, lie not merely in their structural simplicity. The essential significance of
5 tandem architectures is that it enables the voltage required for unassisted water splitting to be achieved
6 without excessive photocurrent loss. Therefore, the distinction between two-terminal and four-terminal
7 (4T) tandem architectures should be viewed not as a question of whether tandem architecture is
8 necessary, but rather as a matter of how to implement tandem configurations most effectively at both
9 the device and system levels.⁴⁰

11 2.3 Tandem structures for unassisted water electrolysis

12 In light of the foregoing discussion, the central challenge in unassisted PV-EC water
13 electrolysis lies not in whether tandem architectures are necessary, but rather in how tandem
14 functionality can be most effectively realized at both the device and system levels. By incorporating
15 two photoabsorbers with complementary bandgaps, tandem photovoltaic configurations provide a
16 viable pathway toward satisfying the photovoltage requirement for bias-free water splitting while
17 maintaining efficient utilization of the solar spectrum.⁴¹ Depending on the mode of electrical
18 interconnection between the top and bottom subcells, tandem devices are broadly categorized into 2T
19 monolithic and 4T architectures (**Fig. 1b**). These two configurations differ fundamentally in their
20 approaches to voltage generation, current matching, optical management, fabrication complexity, and
21 system integration, and thereby present distinct opportunities and trade-offs for PV-EC water
22 electrolysis systems.

23 In the 2T configuration, the two subcells are monolithically integrated and electrically
24 connected in series via an intermediate recombination layer, such as a tunnel junction or a transparent
25 recombination junction. The overall open-circuit voltage (V_{OC}) approximates the sum of the V_{OC} values



1 generated by each subcell, while the operating current is limited by the subcell producing the lowest
2 photocurrent as a consequence of the series connection. This architecture offers practical advantages in
3 terms of wiring simplicity, ease of current matching, module integration, and direct coupling with
4 electrolyzers. However, 2T tandems are inherently sensitive to current mismatch between subcells; any
5 perturbation of the current balance established under standard test conditions, whether caused by
6 spectral or thermal variations during operation, can lead to significant performance degradation.
7 Moreover, the monolithic stacked structure demands high-quality intermediate recombination layers
8 and strict process compatibility between subcells, while optical absorption and resistive losses at the
9 interconnecting layers can further compromise device performance.⁴²

10 In contrast, in the 4T configuration, the top and bottom subcells operate electrically
11 independent of one another, allowing each subcell to operate closer to its maximum power point (MPP)
12 without the current-matching constraints imposed by a series-connected tandem. Consequently, 4T
13 tandems generally exhibit greater tolerance to spectral variations and offer higher design flexibility in
14 terms of bandgap selection, material combinations, and subcell optimization.⁴³ However, realizing these
15 advantages requires additional design elements, including transparent electrodes, individual electrical
16 contacts, and, in many cases, external power management circuitry to combine the outputs of the two
17 subcells.⁴⁴ These requirements introduce additional optical interfaces, parasitic absorption, resistive
18 losses, and increased system complexity. Thus, the 2T configuration is advantageous when compact
19 integration and simple direct coupling are prioritized, whereas the 4T configuration is more suitable
20 when high design flexibility and stable operation under variable illumination conditions are required.⁴⁰

21 Despite this, from the perspective of practical unassisted solar hydrogen production systems,
22 the 2T configuration is often regarded as more significant than the 4T architecture. This is because a 2T
23 tandem integrates two subcells into a single high-voltage photovoltaic unit, thereby enabling direct
24 coupling with an electrolyzer without the need for complex power management circuitry, while also
25 reducing wiring connections, contact interfaces, packaging complexity, and system-level losses.³⁸ In
26 this regard, although the 4T architecture provides a versatile platform for material exploration and



1 performance optimization, the 2T configuration is more closely aligned with practical device
2 implementation, large-area fabrication, and stack-matched series scaling. In such a configuration,
3 scalability is achieved not by increasing the PV voltage alone, but by proportionally extending both the
4 PV module voltage and the electrolyzer stack voltage through series interconnection, thereby preserving
5 direct voltage-matched coupling across scales without additional power conversion electronics.⁴⁵
6 Furthermore, the 2T architecture is highly compatible with the monolithic integration and series-
7 connected module fabrication technologies established in the conventional photovoltaic industry, which
8 offer important advantages for the scalable deployment of low-cost, high-reliability hydrogen
9 production systems. For these reasons, a growing body of research has focused on 2T tandems, driven
10 not only by the pursuit of high photovoltage, but also by the simplicity, integrability, module
11 compatibility, and potential for balance-of-system cost reduction required for practical applications.⁴⁶

12 The key challenge for future research therefore lies not simply in achieving a high V_{OC} , but in
13 realizing a 2T tandem architecture capable of simultaneously delivering sufficient current density and
14 long-term stability at the actual water electrolysis operating point. To this end, several critical
15 requirements must be addressed. First, the bandgaps and thicknesses of the top and bottom subcells
16 must be optimized with due consideration of the actual operating spectrum, so as to minimize current
17 mismatch losses under diverse outdoor conditions. Second, optical absorption and resistive losses in the
18 tunnel junction or recombination layer must be mitigated to minimize voltage and fill factor losses
19 arising from series integration. Third, device design should be guided by the actual water-splitting
20 operating point from a holistic PV-EC coupling perspective that encompasses not only the photovoltaic
21 efficiency but also the performance of the catalyst and electrolyzer. Fourth, particularly in the case of
22 perovskite-based tandems, long-term durability and encapsulation stability under photo-, thermal-,
23 moisture-, and electrochemical stress must be ensured, as these represent essential prerequisites for
24 practical deployment. Finally, rigorous engineering validation is required that extends beyond the
25 pursuit of record device efficiencies at the laboratory scale to encompass large-area module fabrication,



1 manufacturing yield, process reproducibility, system cost, and the ultimate levelized cost of hydrogen
2 production.⁴⁷

3

4 **2.4 PV-EC integration from the standalone systems**

5 The precise synchronization of PV and EC should be considered for efficient solar fuel
6 production from their integration. The operating point of PV-EC is determined by the graphical
7 intersection of the photovoltaic and the electrochemical J - V curve, representing the specific voltage and
8 current at which the PV supply matches the demand of the electrolyzer (**Fig. 1c**).⁴⁸ Single-junction
9 organic and perovskite PVs are now recorded an efficiency of 20%,⁴⁹ and 27%,⁵⁰ respectively. In the
10 case of tandem configurations (to generate photovoltage 1.7-2.1 V for practical water electrolysis), the
11 state-of-art record efficiency is 21.5%, 29.1%, and 26.4% for organic-organic,⁵¹ perovskite-
12 perovskite,⁵² and perovskite-organic tandem PVs.⁵³ However, the PV efficiency is not directly
13 translated into the EC efficiency in the PV-EC configuration, it is because: electrochemical
14 overpotential losses, voltage/operating-point mismatch, and fill factor losses. To minimize energy
15 waste, impedance matching (or coupling efficiency) must be optimized so that this operating point
16 aligns as closely as possible with the maximum power point (MPP) of PV. Thus, unlike grid-connected
17 electrolyzer, minimizing the loss can be achieved through the rational selection of the sub-cells of
18 tandems to provide the necessary potential/and highly active catalysts to induce a steeper J - V or by the
19 geometric matching between the PV active area and the catalyst electrode area. It indicates that coupling
20 PV and EC directly creates a new operating regime, and design rules of optimal PV-EC should be
21 approached in different ways with standalone PV or EC systems.

22 In grid-connected PV, a MPP tracker (MPPT) continuously forces the cell to operate at the
23 voltage and the current at the MPP (V_{mp} and I_{mp}) regardless of load. The design target of PV therefore
24 shifts from maximising the output power density (P_{max} at the MPP) to matching the operating voltage
25 window of electrolyzers across all insolation conditions. In particular, the voltage match is not just at



1 standard test condition (1 sun, AM 1.5G). The operating voltage under 0.1 sun illumination can drop
2 100-150 mV from its 1 sun value, because the V_{OC} of PVs shifts logarithmically with the light
3 intensity.⁵⁴ If the electrolysis onset voltage is 1.8 V and the tandem V_{OC} at 0.1 sun falls to 1.75 V, the
4 system simply stops operating: solar-to-hydrogen efficiency collapses to zero, not merely decreases.
5 The voltage margin at low insolation, not peak efficiency at/over 1 sun, is often the binding design
6 constraint. The $J-V$ intersection of the high-FF PV with the electrolyser load line sits close to MPP; but
7 the intersection is pulled toward lower current and lower voltage than MPP with the low-FF PV,
8 indicating that the operating point is doubly penalised. In particular to the series connected tandems,
9 the lowest-FF subcell limits the whole device. In addition, high series resistance (R_S) of PVs causes a
10 faster, non-linear drop in current and voltage, pushing the intersection point far below the optimal
11 operating point, rapidly degrading system efficiency. In PV-EC, the cell spends substantial time near
12 V_{OC} by the lower insolation than 1 sun. For perovskite-based tandems specifically, ion migration is
13 field-dependent and accelerated at high forward bias (near V_{OC}).⁵⁵ A cell that appears stable under
14 continuous 1 sun MPP illumination (as is standardly measured) can degrade faster in PV-EC cycling.
15 Therefore, stability tests for the PV cell will be more appropriate at the high forward bias for PV-EC
16 qualification, not at the fixed voltage or MPP for grid-tied systems.

17 Grid electrolyzers (such as proton exchange membrane (PEM) electrolyzers) are designed for
18 steady DC input (1.8-2.1 V) and high current density (1-3 A cm⁻²),¹⁰ while the PV-EC electrolyzers
19 operate at low current density (~ 10 mA cm⁻² or below, set by PV current) and variable power by
20 insolation. If the PV delivers 15 mA cm⁻² at 1 sun, the electrolyser must be sized to absorb that current
21 (for a 500 mA cm⁻² PEM design, you need ~ 33 times more PV area than electrolyser area); thus, PV-
22 EC electrolyzers should be designed with a lower nominal current density than grid PEM. While, thanks
23 to the low current density, membrane resistance (ohmic losses) is essentially negligible in PV-EC.
24 However, start-stop cycling (by the insolation) can cause mechanical stress at the catalyst
25 layer/membrane interface in PV-EC electrolyzers, leading to delamination.⁴⁸

26



1 2.5 Advantages of solution processable PV materials

2 The primary motivation for shifting to solution-processable semiconductors is to overcome the
3 significant drawbacks of incumbent tandem technologies. While highly efficient, conventional III-V or
4 silicon-based tandems rely on high-cost, energy-intensive fabrication methods such as vacuum
5 deposition and epitaxial growth.¹¹⁻¹⁴ This leads to high manufacturing costs and creates mechanically
6 rigid devices, limiting their suitability for large-scale, cost-sensitive PV-EC deployment.

7 Solution-processable PV materials offer a paradigm shift by enabling low-cost fabrication.⁵⁶
8 This advantage stems from their compatibility with low-temperature processing, which reduces the
9 overall energy and capital expenditure of manufacturing. In particular, solution-processable PV
10 materials present a distinct set of practical advantages for integrated PV-EC systems, particularly in
11 terms of device form factor, fabrication accessibility, and deployment versatility. Andrei et al.
12 demonstrated that replacing conventional glass substrates and bulk encapsulants with flexible, thin-film
13 alternatives, enabled directly by the low-temperature solution-based deposition of perovskite absorbers
14 onto plastic substrates, reduces device mass to just 30-100 mg cm⁻², a 15-fold reduction relative to
15 traditional photoelectrochemical architectures.⁵⁷ This dramatic reduction in gravimetric footprint is not
16 merely an aesthetic advantage: it enables emergent device functionalities that rigid systems cannot
17 access, most strikingly the spontaneous flotation of the assembled artificial leaf driven by gas bubbles
18 evolving during operation, which was validated during outdoor testing on open water. The compatibility
19 of solution-processed perovskites with scalable deposition routes, including the roll-to-roll and doctor-
20 blading techniques already established in the organic PV industry, further suggests that lightweight PV-
21 EC devices could be manufactured and deployed at scales and in environments, such as open water
22 surfaces, that avoid direct competition with land use. In this sense, the processability of the light
23 absorber is not a secondary manufacturing consideration but a primary enabler of the deployment
24 paradigm itself.

25



1 3 Water electrolysis driven by solution processable tandem PVs

2 3.1 Organic tandem PV driven water electrolysis systems

3 **Table 1** Organic tandem PV driven water electrolysis systems

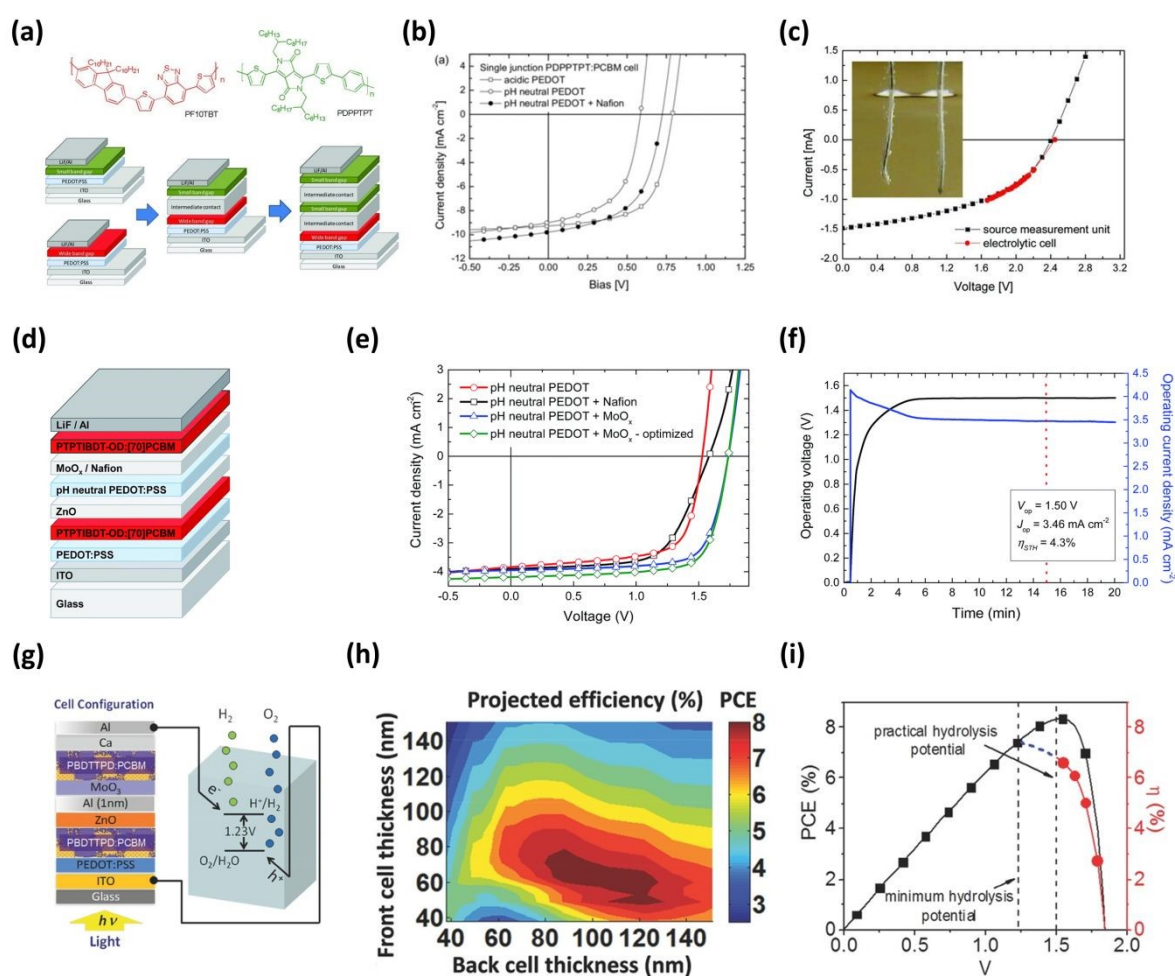
Type	Tandem PV subcells	PV area (cm ²)	Catalyst (HEC OEC)	Electrolyte	Stability	Photocurrent (mA cm ⁻²)	V _{OP} (V)	STH (%)	Year	Ref.
2T	PF10TBT:PCBM/ PDPPPT:PCBM/ PDPPPT:PCBM	0.09 ~ 0.16	Pt Pt	1 M KOH	-	-	1.70	3.1	2013	[58]
2T	PCDTBT:PC ₇₀ BM/ PMDPP3T:PC ₆₀ BM/ PMDPP3T:PC ₆₀ BM	0.0676	RuO ₂ RuO ₂	1 M KOH	-	4.4	1.49	5.41	2015	[59]
2T	PCDTBT:PC ₇₀ BM/ PMDPP3T:PC ₆₀ BM/ PMDPP3T:PC ₆₀ BM	0.0676	Co ₃ O ₄ /NiMoZn	0.1 M KBi	-	3.98	1.56	4.89	2015	[59]
2T	PCDTBT:PC ₇₀ BM/ PMDPP3T:PC ₆₀ BM/ PMDPP3T:PC ₆₀ BM	1.7	RuO ₂ RuO ₂	1 M KOH	-	2.94	1.67	3.61	2015	[59]
2T	PTPTIDBT- OD:PC ₇₁ BM/ PTPTIDBT- OD:PC ₇₁ BM	0.0676	RuO ₂ RuO ₂	1 M KOH	-	3.46	1.50	4.3	2016	[60]
2T	PTB7:PC ₇₁ BM/ PTB7:PC ₇₁ BM/ PTB7:PC ₇₁ BM	0.09	SST-NiMoZn GC-RuO ₂	0.1 M KP _i	50 h (79%) ^a	4.53	1.75	6.0	2016	[61]
2T	PBDTPD:PC ₇₁ BM/ PBDTPD:PC ₇₁ BM/ PBDTPD:PC ₇₁ BM	0.1	Pt Ni foam	1 M NaOH	-	5.4	1.5	6.1	2016	[62]
2T	PM6:IT-M/ PM6:Y6	0.56	Pt NiFeO _x (OH) _y	1 M KOH	2 h (80.8%)	8.2	1.46	10.0	2022	[63]
4T	PBQx-TF:FPCC-Br/ PBQx-TF:FPCC-Br	1 and 1 (parallel illumination)	Pt Pt	1M KOH	600 s (72%)	5.62	1.93	6.91	2025	[64]

^a Tested under day/night cycles (16 h light/8 h dark) with UV filter.

6 The state-of-art organic photovoltaic (OPV) cells have the bulk-heterojunction structure
7 (mixed polymer donor and small molecular acceptor materials) as a photoactive layer. Typically, the
8 magnitude of the V_{OC} loss is determined by the energy offset between donor and acceptor materials for
9 free charge generation in such BHJ devices.⁶⁵ Their single junctions therefore are difficult to produce

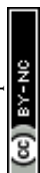


1 sufficient photovoltage to drive unassisted water electrolysis that thermodynamically requires 1.23 V
 2 and practically demands 1.6–2.0 V due to kinetic overpotentials. To overcome this thermodynamic
 3 barrier without external bias, stacking multiple photoactive layers in a tandem configuration has been
 4 the primary strategy. Organic tandem solar cells offer a unique advantage in this regard: the bandgaps
 5 of organic semiconductors can be easily tuned through molecular design, allowing for precise spectral
 6 complementarity between subcells. Furthermore, their compatibility with solution processing enables
 7 the fabrication of lightweight and flexible hydrogen production devices, distinguishing them from rigid
 8 inorganic counterparts.



10

11 **Fig. 2** (a) Molecular structures of the wide bandgap (PF10TBT, $E_g = 1.95$ eV) and narrow bandgap
 12 (PDPPTPT, $E_g = 1.53$ eV) polymers (top), and layouts of the single, tandem, and 1 + 2 type triple
 13 junction solar cells (bottom). (b) Effect of pH-neutral PEDOT and Nafion on the J - V curves of single
 14 junction PDPPTPT:PCBM solar cells. (c) Comparison of the I - V curves of the triple junction cell
 15 measured using a water electrolysis cell with different sized contacts and using a source-measurement



1 unit. Adapted with permission.⁵⁸ Copyright 2013, Wiley-VCH GmbH. (d) Layout of the PTPTIBDT-
2 OD:PCBM homo-tandem solar cell with Nafion or MoO₃ layers in the intermediate contact. (e)
3 Comparison of the $J-V$ characteristics of PTPTIBDT-OD:PCBM homo-tandem solar cells with and
4 without Nafion or MoO₃ layers in the intermediate contact. (f) Simultaneous measurement of the
5 operating voltage and current density of the device during light-driven electrochemical water splitting
6 using RuO₂ catalysts in 1.0 M KOH. Adapted with permission.⁶⁰ Copyright 2016, The Royal Society
7 of Chemistry. (g) Schematic PV-driven electrochemical water splitting study. (h) PCE of homo-tandem
8 solar cells predicted from optical and electrical modelling. (i) PCE of the homo-tandem solar cells
9 (measured *via* source-measurement unit) and estimated STH conversion efficiency (denoted as η) at
10 various operating voltages. Adapted with permission.⁶² Copyright 2016, WILEY-VCH Verlag GmbH.

11

12 3.1.1 Multi-junctions to drive water electrolysis

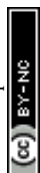
13 The development of organic tandem architectures for water splitting has evolved from
14 increasing the number of junctions to optimizing the quality of each junction. In a pioneering 2013
15 study, Esiner et al. employed a triple-junction organic solar cell consisting of PF10TBT:PC₆₁BM wide
16 bandgap (1.95 eV) and two PDPPTPT:PC₆₁BM narrow bandgap (1.53 eV) subcells to achieve a V_{OC} of
17 2.33 V and STH efficiency of 3.1%.⁵⁸ This device architecture (**Fig. 2a**) introduced the concept of
18 spectral splitting through bandgap engineering within an all-organic system and demonstrated the
19 feasibility of achieving high voltage output using solution-processable organic materials. A critical
20 enabler for this fully solution-processed stack was the development of a robust intermediate connecting
21 layer comprising ZnO, pH-neutral PEDOT, and Nafion. While the use of conventional acidic
22 PEDOT:PSS typically damages the underlying ZnO layer, substituting it with pH-neutral PEDOT:PSS
23 reduces the work function from 5.05 eV to 4.65 eV, resulting in voltage losses. To address this limitation,
24 the authors adopted a strategy of spin-coating a Nafion solution onto the PEDOT:PSS layer. This
25 specific architecture prevented the acidic corrosion of the underlying ZnO layer during processing and
26 minimized voltage losses, ensuring efficient charge recombination between subcells (**Fig. 2b**). When
27 the triple-junction solar cell was coupled with Pt electrodes for water electrolysis, an STH efficiency of
28 3.1% was achieved (**Fig. 2c**).

29 To overcome the voltage limitations of single-junction devices, Esiner et al. developed a
30 homo-tandem solar cell utilizing a newly synthesized wide bandgap polymer, PTPTIBDT-OD ($E_g =$
31 2.04 eV), which is intrinsically capable of delivering high V_{OC} .⁶⁰ However, realizing the full potential



1 of such high-voltage tandem devices requires precise engineering of the interconnection layer. **Fig. 2d**
2 illustrates the specific device architecture employed to address this challenge. A conventional
3 interconnection layer based on ZnO and pH-neutral PEDOT:PSS often suffers from a work function
4 mismatch; the low work function of pH-neutral PEDOT:PSS (4.65 eV) creates a non-ohmic contact
5 with deep-HOMO polymers, leading to significant voltage losses. To mitigate this, the authors
6 introduced a thin layer of molybdenum oxide (MoO_x) on top of the pH-neutral PEDOT:PSS layer. This
7 MoO_x layer, with its high work function (5.40 eV), serves as an efficient hole collector, ensuring ohmic
8 contact and preventing potential losses at the recombination interface. The impact of this interfacial
9 modification on photovoltaic performance is evident in the J - V characteristics shown in **Fig. 2e**. The
10 control tandem device without any modification exhibited a limited V_{OC} of 1.53 V, confirming the
11 severity of the interfacial barrier. While the incorporation of a Nafion layer resulted in a marginal
12 improvement to 1.58 V, the introduction of the thermally evaporated MoO_x layer dramatically boosted
13 the V_{OC} to 1.74 V. This optimized interconnection layer configuration not only maximized the voltage
14 output but also maintained a high FF of 0.73, which is critical for minimizing power losses during
15 operation. The robust performance of this high-voltage tandem cell enabled efficient bias-free water
16 splitting when coupled with RuO_2 electrocatalysts. **Fig. 2f** displays the simultaneous measurement of
17 the operating voltage and current density of the integrated system over time. The device exhibited stable
18 operation with a stabilized operating voltage (V_{OP}) of 1.50 V, which remarkably aligns with the voltage
19 at the maximum power point ($V_{\text{MPP}} \approx 1.48$ V) of the tandem PV. This precise matching between the
20 photovoltaic power output and the electrochemical load resulted in a STH conversion efficiency of
21 4.3%, demonstrating that interface engineering is a decisive factor in constructing high-efficiency,
22 unassisted solar water splitting systems.

23 To address the limited photovoltage of single-junction organic solar cells and achieve efficient
24 unassisted water splitting, Gao et al. proposed a homo-tandem organic solar cell architecture utilizing a
25 wide-bandgap polymer. **Fig. 2g** illustrates the schematic of this integrated PV-driven electrochemical
26 water splitting system. The device connects two identical PBDTTPD:PC₇₁BM photoactive layers in



1 series to generate sufficient voltage to overcome the thermodynamic potential of water splitting and
2 kinetic overpotentials. A key enabling feature of this architecture is the robust interconnection layer
3 composed of solution-processed ZnO, ultrathin Al, and MoO₃, which facilitates efficient charge
4 recombination between the subcells while minimizing voltage losses. In a homo-tandem configuration
5 where subcells share identical absorption profiles, balancing the photocurrents is critical for maximizing
6 device performance. To optimize this current matching, the authors employed combined optical and
7 electrical modelling to predict the power conversion efficiency (PCE) as a function of subcell
8 thicknesses. **Fig. 2h** presents the simulation results, identifying that a specific asymmetric thickness
9 combination — a thinner front cell (60–80 nm) and a thicker back cell (90–130 nm) — is required to
10 achieve PCEs exceeding 8.0%. This configuration ensures that sufficient light is transmitted through
11 the front cell to be absorbed by the back cell, effectively balancing the photocurrents despite the
12 identical bandgaps. Guided by this modelling, an optimized homo-tandem device (70 nm front / 120
13 nm back) was fabricated, yielding a high V_{OC} of 1.84 V. **Fig. 2i** displays the PCE of the tandem device
14 and the estimated STH conversion efficiency relative to the operating voltage. Notably, the V_{MPP} of the
15 solar cell (≈ 1.54 V) aligns well with the practical voltage required for water electrolysis in 1 M NaOH
16 electrolyte (~ 1.5 V). Consequently, this optimized homo-tandem system achieved an estimated STH
17 efficiency of 6.1% without any external bias, demonstrating that precise optical and structural
18 engineering can enable efficient solar fuel production even when using homo-tandem architectures.



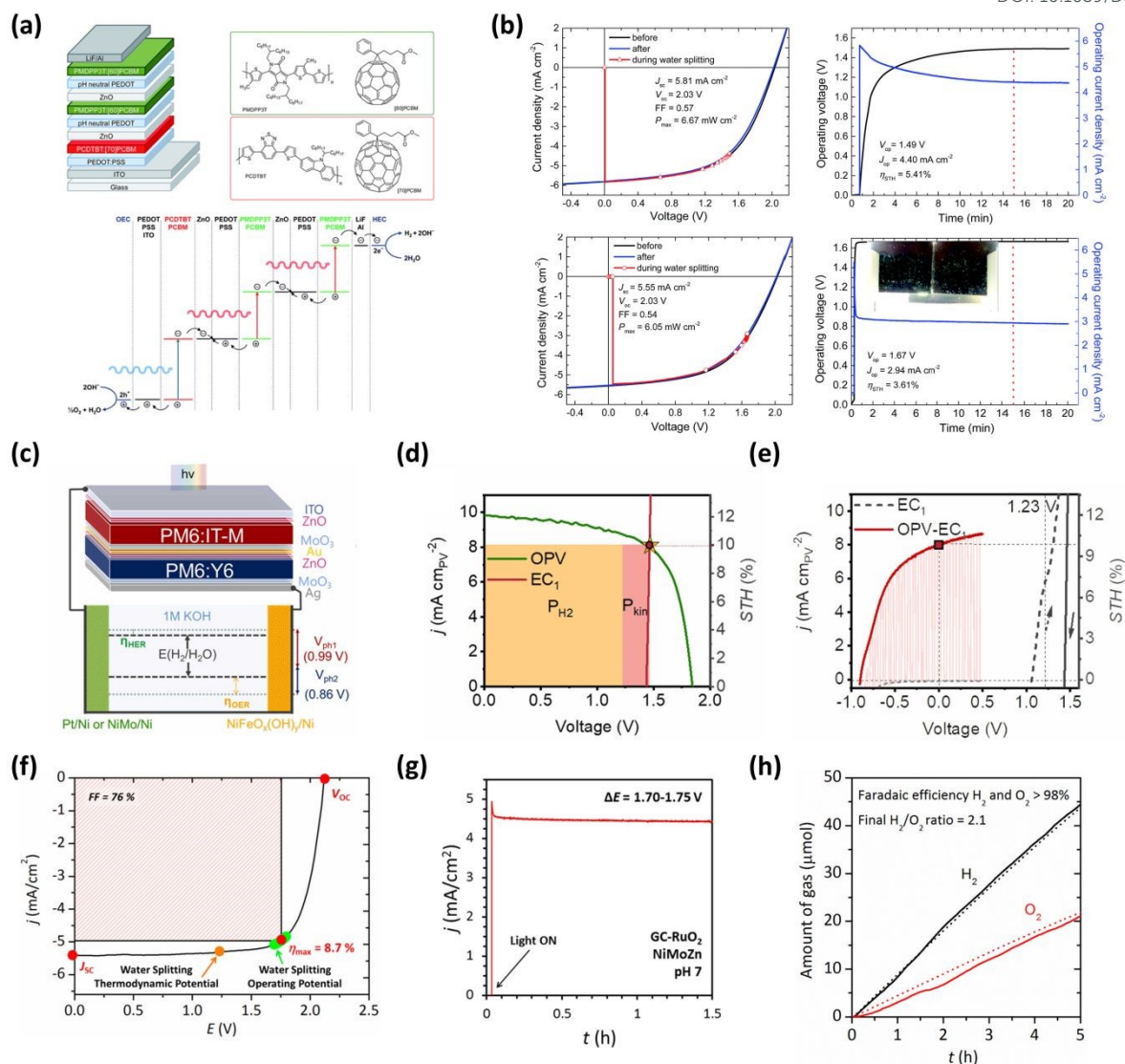


Fig. 3 (a) Device layout of triple junction organic solar cell. (b) J - V curves of triple junction solar cells before, during, and after water splitting measurement of 20 min. Adapted with permission.⁵⁹ Copyright 2015, The Royal Society of Chemistry (c) Schematics of the PM6:IT-M/PM6:Y6 tandem PV-EC system. (d) J - V curves of tandem OPV (0.56 cm²) and EC₁ in a two-electrode configuration (2.0 cm²). The MPP of PV and the operating point of EC₁ are marked with a star and a dot, respectively. The amount of power stored as hydrogen (P_{H_2}) and that wasted as kinetic loss (P_{kin}) are indicated. (e) LSV of EC₁ in a two-electrode configuration and OPV-EC system with its STH efficiency. Adapted with permission.⁶³ Copyright 2022, Elsevier B.V (f) J - V curve of the triple junction solar cell. (g) Current density vs time profile of a water splitting experiment using a triple junction solar cell (a PCE of the PV was 8.5%), GC-RuO₂ anode, and NiMoZn cathode in a two-electrode configuration and a two-compartment cell containing 0.1 M phosphate buffer, pH = 7.0, under AM 1.5 G illumination with a GG400 filter. (h) Gas evolution of the water splitting experiment using the triple junction solar cell under the same condition. Adapted with permission.⁶¹ Copyright 2016, American Chemical Society.

3.1.2 System integration and catalytic optimization



1 To address the high voltage requirements of water electrolysis, Esiner et al. developed an
2 "organic artificial leaf" based on a solution-processed triple-junction organic solar cell.⁵⁹ **Fig. 3a**
3 illustrates the device architecture and energy level diagram, comprising a wide bandgap
4 PCDTBT:PC₇₁BM top cell and two identical narrow bandgap PMDPP3T:PC₆₁BM bottom cells
5 connected *via* ZnO/pH-neutral PEDOT interconnecting layers. This series-connected configuration was
6 designed to generate an open-circuit voltage exceeding 2.0 V, providing sufficient potential to drive the
7 water splitting reaction without external bias. The performance of this triple-junction cell when coupled
8 with RuO₂ electrocatalysts was critically dependent on the geometric surface area ratio between the
9 catalyst and the solar cell. Above **Fig. 3b** present the *J-V* characteristics and transient operation of a
10 small-area device (0.0676 cm²) connected to large-area catalysts (~1.2 cm²), resulting in a high catalyst-
11 to-PV area ratio of approximately 15. Under these conditions, the low local current density on the
12 catalyst surface minimized the kinetic overpotential (0.26 V total), allowing the system to stabilize at
13 an operating voltage (V_{OP}) of 1.49 V. Notably, this operating point closely matched the voltage at the
14 maximum power point ($V_{MPP} = 1.44$ V) of the solar cell, enabling the device to deliver a high
15 photocurrent density of 4.40 mA cm⁻² and achieve a STH conversion efficiency of 5.4%. However,
16 scaling up the device revealed the critical impact of kinetic constraints. Below **Fig. 3b** show the
17 performance of a larger-area solar cell (1.7 cm²) coupled with catalysts of similar size (~1.2 cm²),
18 effectively reducing the area ratio to ~0.7. Consequently, the current density on the catalyst surface
19 increased significantly, raising the required overpotential by approximately 0.18 V. This shift forced
20 the system to operate at a higher voltage of 1.67 V, which is far beyond the V_{MPP} (1.40 V) of the solar
21 cell, pushing the device into a region of lower photocurrent extraction. As a result, the operating current
22 density dropped to 2.94 mA cm⁻², and the STH efficiency decreased to 3.6%, demonstrating that
23 matching the geometric areas of the PV and catalyst components is as crucial as material selection for
24 efficient solar-to-fuel conversion.

25 To push the boundaries of OPV-driven water splitting, Kim et al. developed a high-efficiency
26 hetero-tandem organic PV-EC system by combining a wide bandgap PM6:IT-M front cell and a narrow



1 bandgap PM6:Y6 back cell.⁶³ **Fig. 3c** illustrates the integrated device architecture, where this hetero-
2 tandem OPV, capable of delivering a high V_{OC} of 1.84 V, is coupled with an alkaline electrolyzer
3 consisting of earth-abundant catalysts (Pt or NiMo for HER and NiFeO_x(OH)_y for OER). This
4 configuration was strategically designed to overcome the voltage deficit of single-junction cells while
5 maximizing the spectral utilization of solar energy through complementary absorption. A critical
6 innovation in this work lies in the optimization of the operating point through geometric area
7 engineering. **Fig. 3d** displays the linear sweep voltammetry (LSV) curves of the OPV and the
8 electrolyzer (EC₁). By adjusting the operating area ratio of the electrolyzer to the PV to approximately
9 3.6, the authors successfully aligned the operating voltage of the coupled system ($V_{OP} \approx 1.46$ V) with
10 the maximum power point voltage (V_{MPP}) of the tandem solar cell. This precise impedance matching
11 minimized power transfer losses, allowing the system to operate at a high current density of 8.2 mA
12 cm⁻², which directly translates to a benchmark STH conversion efficiency of 10%. The robust
13 performance of this optimized system is further highlighted in **Fig. 3e**, which presents the LSV scan of
14 the integrated OPV-EC device. Even without any external bias (at 0 V), the system maintained a high
15 operating current density of 8.0 mA cm⁻², corresponding to an STH efficiency of 9.8%. This result
16 underscores that synergistically combining a high-voltage hetero-tandem absorber with rational system
17 engineering — specifically, matching the PV and catalyst areas — is a viable pathway to realize highly
18 efficient, unassisted solar hydrogen production using organic photovoltaics.

19 To address the kinetic challenges of water electrolysis at neutral pH, Elias et al. focused on
20 optimizing the FF of the photovoltaic component to maximize power transfer.⁶¹ As aforementioned in
21 Section 2.4, the FF losses in PV are critical to determine the STH efficiency in PV-EC, but the organic
22 PVs and their tandems in particular have low FF, because of bimolecular recombination as a result of
23 low carrier mobility in organic semiconductors.⁵⁴ They developed a triple-junction organic solar cell
24 using identical subcells based on the PTB7:PC₇₁BM blend. **Fig. 3f** presents the J - V characteristics of
25 this optimized device. The solar cell achieved a high V_{OC} of 2.13 V and a remarkable FF of 76%. They
26 argued that the simplified fabrication and the homogeneous electrical performance throughout the



1 device can result in high FF by using the homo-tandem of the same three sub-cells. As indicated in the
2 figure, the operating potential required by the catalysts (≈ 1.7 V) intersects the J - V curve very close to
3 the voltage at the maximum power point of the PV, thereby minimizing electrical losses that typically
4 occur when the operating point deviates from the optimal power output. This high-performance
5 photovoltaic device was integrated with a catalyst system designed for neutral pH operation, comprising
6 a glassy carbon-supported RuO_2 anode (GC- RuO_2) and a stainless steel-supported NiMoZn cathode.
7 **Fig. 3g** illustrates the operational stability of this integrated PV-EC system. The device maintained a
8 stable current density of approximately 4.53 mA cm^{-2} over 1.5 hours, with a V_{OP} stabilizing between
9 1.70 and 1.75 V. This stable performance confirms that the high voltage and FF of the triple-junction
10 PV cell are sufficient to overcome the thermodynamic and kinetic barriers of water splitting in a
11 phosphate buffer solution at pH 7. The efficacy of the system was further validated by quantitative gas
12 analysis. **Fig. 3h** shows the time-dependent evolution of hydrogen and oxygen gases over a continuous
13 5-hour operation. The system demonstrated near-unity Faradaic efficiency (greater than 98%) for both
14 gases, with a production ratio of H_2 to O_2 close to the theoretical value of 2:1. Consequently, this
15 configuration achieved a STH conversion efficiency of approximately 6%, demonstrating that
16 optimizing the fill factor of the light absorber is a decisive strategy for enabling efficient, bias-free water
17 splitting under neutral conditions.

18 While early-stage efforts established fundamental design strategies such as bandgap
19 engineering and interfacial tuning, recent work has emphasized system-level optimization, including
20 electrode area ratio control and catalyst selection. However, progress remains limited by the inherent
21 performance constraints of organic PV materials. The highest STH efficiency reported for an organic
22 tandem is 10%, achieved with the PM6:IT-M/PM6:Y6 hetero-tandem.⁶³ This figure remains well below
23 the 18% or higher STH values already attained with perovskite-based or III-V-based PV-EC
24 platforms.⁶⁶ Consequently, organic tandem PV cells are still regarded as proof-of-concept systems
25 whose STH metrics lag behind those of more mature inorganic counterparts.

26



1 3.1.3 Cost-effective wide-bandgap acceptors for solar cells enabling bias-free underwater

2 hydrogen production

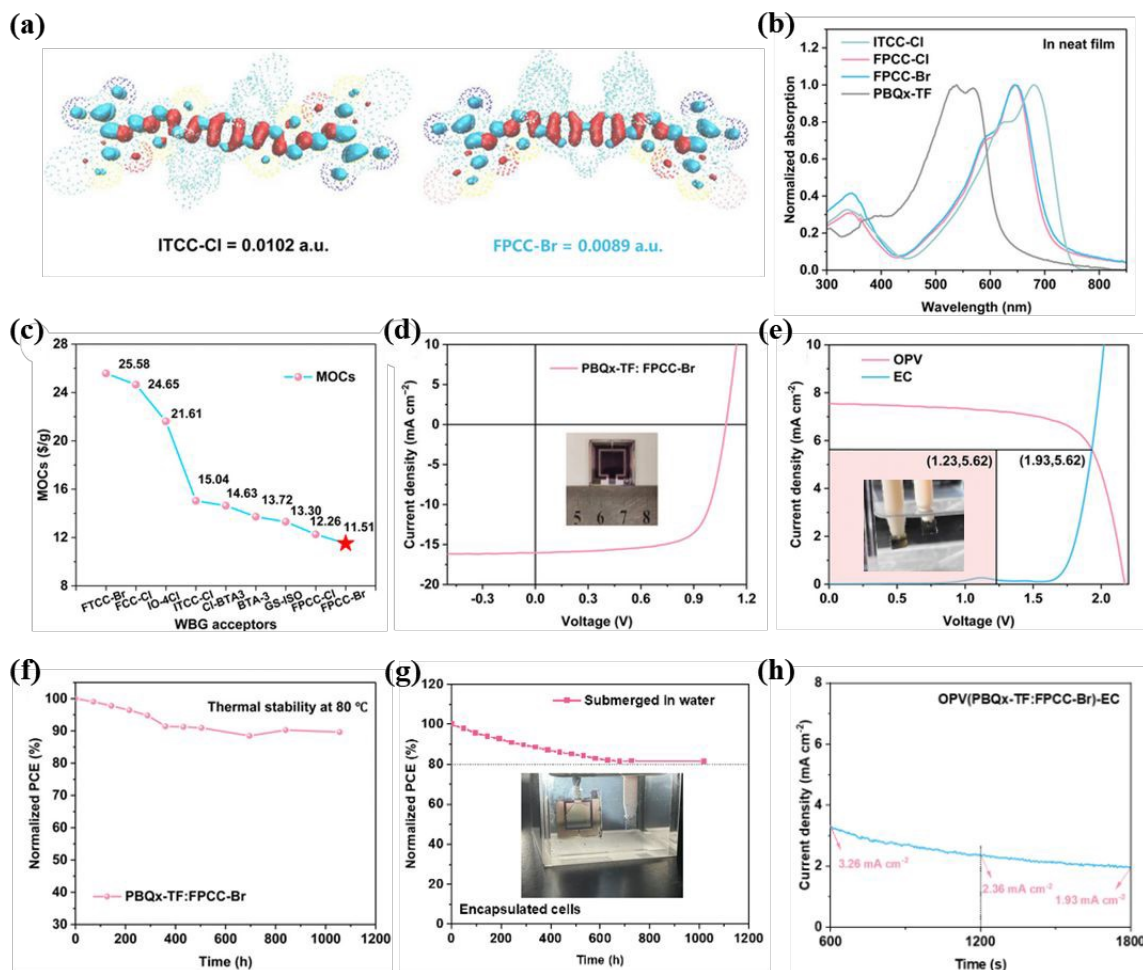
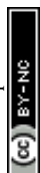


Fig. 4 (a) Overlap and the exchange integrals of HOMO (red) and LUMO (sky blue) distributions. (b) UV-vis absorption spectra of neat films. (c) Material-only cost (MOC) values of typical wide band gap acceptors. (d) J - V curves of a 1 cm² device using blade-coating. (e) J - V curves of two OPV cells connected in series and the physical diagram of two series connected cells directly used for underwater photovoltaic electrolysis. (f) Thermal stability of FPCC-Br-based devices at 80 °C. (g) The underwater photovoltaic stability with an FPCC-Br-based cells. (h) The chronoamperometric measurement of two FPCC-Br-based cells connected in series. Adopted with permission.⁶⁴ Copyright 2025, The Royal Society of Chemistry.

The development of high-performance wide-bandgap acceptors plays a pivotal role in optimizing the front cell of tandem architectures and extending the application scope of PV-EC systems to non-conventional environments. Xiao et al. designed and synthesized a novel wide-bandgap acceptor,



1 FPCC-Br, based on a molecular design strategy that minimizes the spatial overlap between the HOMO
2 and LUMO (exchange integral) (**Fig. 4a**). This approach induced a blue-shift in the absorption spectrum
3 and enhanced V_{OC} (**Fig. 4b**), yielding a PCE of 13.6% in PBQx-TF:FPCC-Br-based single-junction
4 cells (a record high for bandgaps below 720 nm), 19.3% in ternary cells, and 20.1% in tandem front
5 cells, while effectively mitigating the long-standing trade-off between synthetic cost and device
6 performance.⁶⁴

7 Owing to its streamlined synthetic route and high yield, FPCC-Br possesses the lowest raw
8 material cost among wide-bandgap acceptors reported to date, and maintains a PCE of 12.2% in large-
9 area (1 cm²) blade-coating fabrication, underscoring its strong commercialization potential (**Fig. 4c-d**).
10 Furthermore, this wide-bandgap material is particularly well-suited for applications beyond terrestrial
11 environments, including underwater photovoltaic electrolysis. Xiao et al. experimentally demonstrated
12 this by connecting PBQx-TF:FPCC-Br-based organic solar cells in series with Pt electrodes to achieve
13 bias-free underwater hydrogen production. At a water depth of 5 cm, the system delivered a current
14 density of 5.62 mA cm⁻² at an operating voltage of 1.93 V, achieving a STH efficiency of 6.91% (**Fig.**
15 **4e**).

16 Despite being fabricated via a solution-processed approach, the FPCC-Br-based system
17 exhibited excellent operational and environmental stability. Under thermal stress at 80 °C, it retained
18 approximately 90% of its initial PCE after 1000 hours (**Fig. 4f**). Even under continuous operation in
19 water, the device maintained 81.5% of its initial efficiency after 1000 hours, demonstrating the
20 remarkable water resistance and long-term stability of the organic semiconductor (**Fig. 4g**).
21 Furthermore, during electrolysis at a depth of 5 cm, the system sustained 72% of its initial current
22 density for 600 s, while the decay rate of the current density gradually stabilized over time (**Fig. 4h**).
23 These results highlight that the FPCC-Br-based wide-bandgap acceptor represents a promising
24 candidate for realizing highly reliable PV-EC systems.

25



1 3.2 Perovskite tandem PV driven water electrolysis systems

2 **Table 2** Perovskite tandem PV driven water electrolysis systems

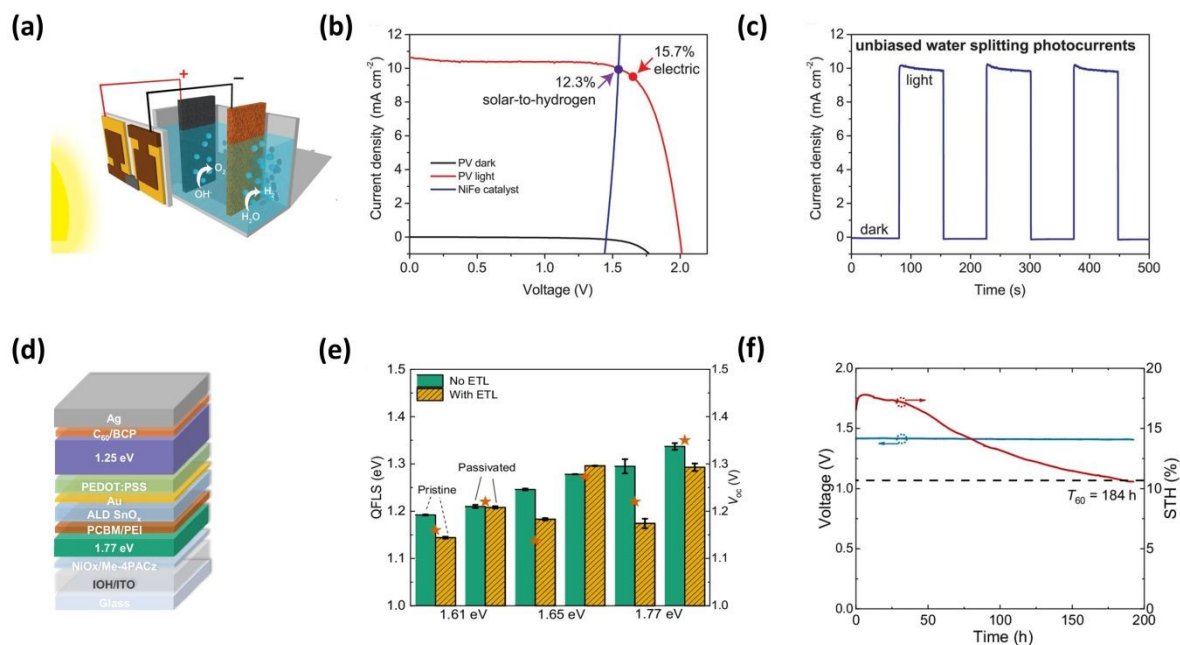
Type	Tandem PV subcells	PV area (cm ²)	Catalyst (HEC OEC)	Electrolyte	Stability	Photocurrent (mA cm ⁻²)	V _{OP} (V)	STH (%)	Year	Ref.
2T	Cs _{0.2} FA _{0.8} Pb(Br _{0.3} I _{0.7} /FA _{0.7} MA _{0.3} Pb _{0.5} Sn _{0.5} I ₃)	0.1	Pt IrO _x	0.5 M H ₂ SO ₄	120 h (95%)	14.1	-	17	2023	[38]
2T	FA _{0.8} Cs _{0.2} PbBrI ₂ /FA _{0.7} MA _{0.3} PbI ₃	0.12	Pt IrO _x	0.5 M H ₂ SO ₄	-	~7	-	8.5	2021	[67]
2T	Cs _{0.2} FA _{0.8} Pb(I _{0.6} Br _{0.4} /Cs _{0.05} FA _{0.7} MA _{0.25} Pb _{0.5} Sn _{0.5} I ₃)	1	Pt/C RuO ₂	H ₂ O	184 h (over 60%)	14.7	1.42	17.8	2025	[39]
4T	CsPbI _{1.5} Br _{1.5} /CsPb _{0.5} Sn _{0.5} I _{2.7} Br _{0.3}	0.1	Pt/C Ni/Fe	1 M KOH	2 h (97.2%)	10.71	1.62	13.1	2022	[43]
4T	CH ₃ NH ₃ PbI ₃ /CH ₃ NH ₃ PbI ₃	0.318	NiFe-LDH NiFe-LDH	1 M NaOH	2 h (~80%)	9.61	1.63	12.3	2014	[68]
4T	(FA _{0.7} MA _{0.3}) _{0.935} Cs _{0.065} Pb(I _{0.89} Br _{0.11}) ₃ / (FA _{0.7} MA _{0.3}) _{0.935} Cs _{0.065} Pb(I _{0.89} Br _{0.11}) ₃	1 and 1 (parallel illumination)	Pt CoFe	1 M KOH + 0.5 M NaCl	10 h (90%)	11.53	1.56	14.18	2025	[69]
4T	(FA _{0.7} MA _{0.3}) _{0.935} Cs _{0.065} Pb(I _{0.89} Br _{0.11}) ₃ / (FA _{0.7} MA _{0.3}) _{0.935} Cs _{0.065} Pb(I _{0.89} Br _{0.11}) ₃	2 and 2 (parallel illumination)	Pt NiFe	1 M KOH	-	11.18	-	13.75	2025	[70]

3

4 To address the limitations of organic tandem PVs, halide perovskite tandem PVs have emerged
 5 as a promising alternative. Perovskite materials offer superior optoelectronic properties — such as high
 6 absorption coefficients, long carrier diffusion lengths, and tunable bandgaps — that make them
 7 particularly advantageous for tandem solar cell integration.^{71, 72} In particular, the ability to engineer both
 8 wide- and narrow-bandgap perovskite absorbers has enabled the development of monolithic and four-
 9 terminal devices capable of delivering high photovoltages and stable operation.⁷³⁻⁸¹ Combining these
 10 strengths, perovskite tandems features have led to significant improvements in STH efficiency, with
 11 recent reports exceeding 17%.^{39, 82} Consequently, all-halide perovskite tandem architectures
 12 theoretically allow for optimal efficiency due to the tunability of the band gaps in both subcells and the
 13 specialized function of each absorber.⁸³



1

2 **3.2.1 Performance optimization and stability challenges**

3

4 **Fig. 5** (a) Schematic diagram of the water-splitting driven by the four-terminal perovskite tandem PV.
 5 (b) $J-V$ curves of the perovskite tandem cell under dark and simulated AM 1.5G illumination, and the
 6 NiFe/Ni foam electrodes in a two-electrode configuration. The illuminated surface area of the
 7 perovskite cell was 0.318 cm^2 , and the catalyst electrode areas (geometric) were $\sim 5 \text{ cm}^2$ each. (c)
 8 Current density–time curve of the integrated water-splitting device without external bias under chopped
 9 simulated AM 1.5G illumination. Adapted with permission.⁶⁸ Copyright 2014, American Association
 10 for the Advancement of Science. (d) Inverted (p-i-n) perovskite tandem solar cell layout. (e) Quasi-
 11 Fermi level splitting of 1.61, 1.65, and 1.77 eV perovskite films. The stars represent the maximum V_{OC}
 12 obtained using each perovskite absorber. (f) Voltage (blue) and STH conversion efficiency (red) vs.
 13 time using the integrated PV-EC system during 192 h continuous operation at approximated 1-sun
 14 equivalent light intensity. Adapted with permission.³⁹ Copyright 2025, Springer Nature.

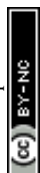
15

16 To address the economic barriers of solar hydrogen production, Luo et al. proposed a high-
 17 efficiency water splitting device that eliminates expensive noble metal catalysts.⁶⁸ **Fig. 5a** depicts the
 18 schematic of this integrated system, where two $\text{CH}_3\text{NH}_3\text{PbI}_3$ perovskite solar cells are connected in
 19 series to form a tandem configuration. This photovoltaic source is wired to Earth-abundant nickel-iron
 20 layered double hydroxide (NiFe-LDH) electrodes, which serve as bifunctional catalysts for both the
 21 hydrogen and oxygen evolution reactions. The tandem architecture allows the device to generate a high
 22 open-circuit voltage of 2.00 V, providing the necessary thermodynamic potential and overpotential to

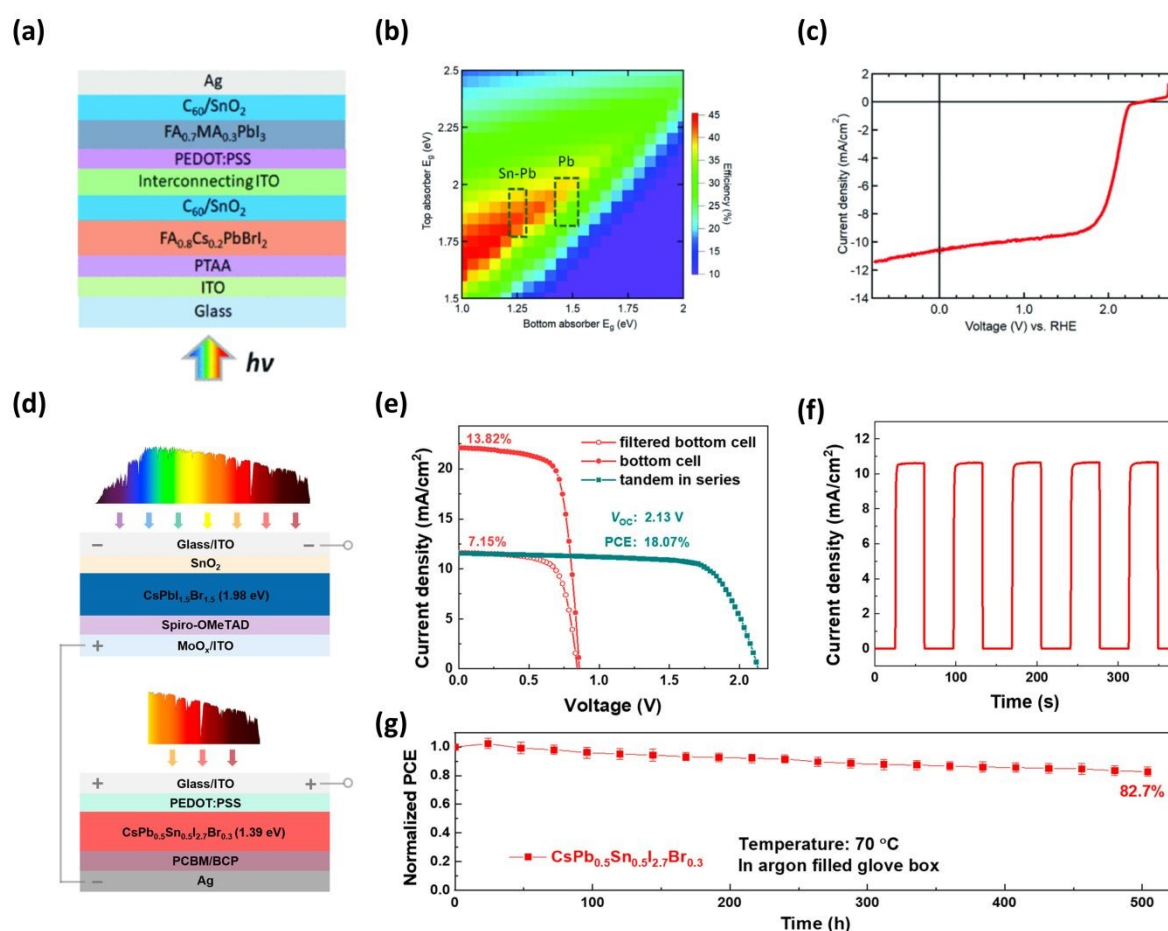


1 drive water electrolysis without external bias. The efficiency of this system is defined by the coupling
2 between the photovoltaic power output and the electrochemical load. **Fig. 5b** displays the J - V
3 characteristics of the perovskite tandem cell alongside the polarization curve of the NiFe-LDH
4 electrolyzer. The intersection of these two curves represents the operating point of the integrated device.
5 Notably, this operating point occurs at a current density of approximately 10 mA cm^{-2} and a voltage of
6 1.63 V , which is remarkably close to the maximum power point of the solar cell. This optimal
7 impedance matching minimizes electrical losses and results in a STH conversion efficiency of 12.3%.
8 The capability of the device to perform unassisted water splitting is further confirmed by the
9 photocurrent measurements over time. **Fig. 5c** shows the current density profile of the integrated system
10 under chopped simulated sunlight with no external bias applied. The device rapidly reaches a stable
11 photocurrent of approximately 10 mA cm^{-2} upon illumination, maintaining this performance during the
12 initial testing period. This demonstration confirms that solution-processed perovskite photovoltaics
13 combined with low-cost, Earth-abundant catalysts can achieve STH efficiencies exceeding the 10%
14 benchmark required for practical viability.

15 Building on the potential of perovskite technologies, Wang et al. developed a monolithic all-
16 perovskite tandem solar cell specifically optimized for high-voltage water splitting applications.³⁹ **Fig.**
17 **5d** illustrates the device architecture, which consists of a wide bandgap ($\text{Cs}_{0.2}\text{FA}_{0.8}\text{Pb}(\text{I}_{0.6}\text{Br}_{0.4})_3$, 1.77
18 eV) top cell and a narrow bandgap ($\text{Cs}_{0.05}\text{FA}_{0.7}\text{MA}_{0.25}\text{Pb}_{0.5}\text{Sn}_{0.5}\text{I}_3$, 1.25 eV) bottom cell connected by a
19 recombination layer. A critical challenge in such wide-bandgap perovskites is the voltage loss due to
20 non-radiative recombination at the electron transport layer interface. To mitigate this, the authors
21 applied a propane-1,3-diammonium iodide (PDAI_2) surface treatment. As shown in **Fig. 5e**, the quasi-
22 Fermi level splitting measurements reveal that the PDAI_2 treatment significantly reduces energetic
23 losses, enabling the 1.77 eV perovskite to achieve an open-circuit voltage closer to its radiative limit
24 compared to the pristine control. This voltage-optimized tandem cell was integrated into a PV-EC flow
25 cell system to evaluate its performance and stability. **Fig. 5f** presents the long-term operational data of
26 this system under continuous illumination. The device achieved a record-high STH efficiency of 17.8%



1 at the start of the operation. More importantly, the system demonstrated substantial durability,
 2 maintaining stable voltage output and retaining over 60% of its initial efficiency after more than 180
 3 hours of continuous operation. While the voltage remained relatively constant, the gradual decline in
 4 STH efficiency was attributed to charge collection losses in the narrow bandgap subcell, highlighting
 5 that while interface engineering can maximize initial performance, intrinsic stability of the narrow
 6 bandgap perovskite absorber remains a key focus for future improvements.



7
 8 **Fig. 6** (a) Schematic illustrations of device structure of a FA_{0.8}Cs_{0.2}PbBr₂/FA_{0.7}MA_{0.3}PbI₃ tandem solar
 9 cell. (b) Theoretical efficiency of two-terminal tandem solar cells as a function of top and bottom
 10 absorber bandgaps. (c) Linear sweep voltammetry of the perovskite tandem PV driven EC system was
 11 performed in a three-electrode setup featuring a Pt cathode and an IrO_x anode.⁶⁷ Copyright 2025, IEEE.
 12 (d) Schematic diagram of the four-terminal tandem solar cell made by combining a CsPb_{0.5}Sn_{0.5}I_{2.7}Br_{0.3}
 13 narrow bandgap perovskite as the bottom subcell and a semi-transparent CsPbI_{1.5}Br_{1.5} wide bandgap
 14 perovskite as the top subcell. (e) $J-V$ curves of the integrated four-terminal all-inorganic perovskite
 15 tandem solar cell measured using a metallic mask with an aperture area of 0.1 cm² under AM 1.5 G
 16 illumination with a scan rate of 50 mV s⁻¹ in the reverse scan direction. (f) Practical operating current
 17 density of the assembled solar water splitting system under chopped AM 1.5G illumination without
 18 external bias. (g) Thermal stability test of un-encapsulated CsPb_{0.5}Sn_{0.5}I_{2.7}Br_{0.3} solar cells measured

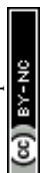


1 under 70°C aging condition in an argon-filled glovebox. Adapted with permission.⁴³ Copyright 2025,
2 American Chemical Society.

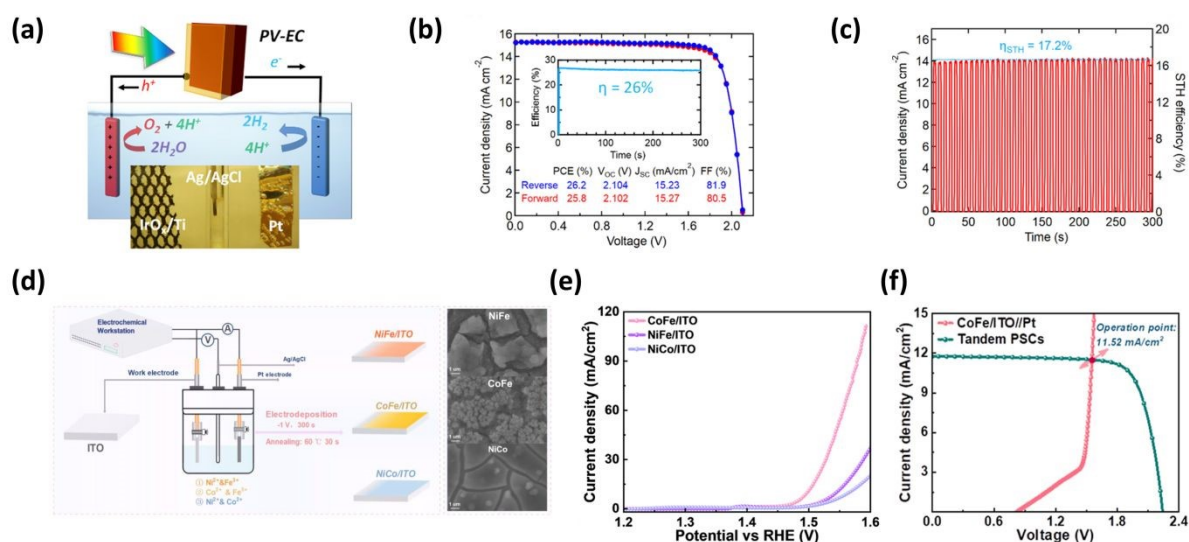
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4 Stability under heat, moisture, and UV exposure has remained a major challenge for all-
5 perovskite tandem solar cells, prompting extensive efforts to improve both the intrinsic and extrinsic
6 durability of their components. Song et al. addressed this by introducing a monolithic all-perovskite
7 tandem configuration that completely eliminates chemically unstable Sn-containing perovskites (**Fig.**
8 **6a**).⁶⁷ They utilized a wide bandgap $\text{FA}_{0.8}\text{Cs}_{0.2}\text{PbBrI}_2$ top cell and a narrower bandgap $\text{FA}_{0.7}\text{MA}_{0.3}\text{PbI}_3$
9 bottom cell, employing fully solution-processed, Pb-only materials. As shown in the theoretical
10 efficiency simulation (**Fig. 6b**), shifting from the conventional Sn-Pb based bottom cell (left box) to a
11 pure Pb-based one (right box) involves a trade-off in theoretical efficiency limits. The practical viability
12 of this approach for unassisted water splitting was evaluated using a PV-EC system. **Fig. 6c** presents
13 LSV of the tandem device in a three-electrode configuration. The device exhibits a high onset potential
14 of approximately 2.1 V vs. the reversible hydrogen electrode, which provides ample voltage to drive
15 the water splitting reaction. Consequently, the system achieves a photocurrent density of 10.5 mA cm⁻²
16 at zero bias, confirming that pure Pb-based all-perovskite tandem cells can effectively power
17 unassisted water electrolysis while offering enhanced material stability.

18 Sun et al. demonstrated the first integrated four-terminal tandem solar cell composed entirely
19 of all-inorganic perovskite absorbers to achieve both high efficiency and superior thermal stability.⁴³
20 **Fig. 6d** depicts the schematic of this four-terminal architecture, featuring a semi-transparent inorganic
21 wide-bandgap $\text{CsPbI}_{1.5}\text{Br}_{1.5}$ top cell and a narrow-bandgap $\text{CsPb}_{0.5}\text{Sn}_{0.5}\text{I}_{2.7}\text{Br}_{0.3}$ bottom cell. This all-
22 inorganic design effectively mitigates the volatility issues of organic cations. The photovoltaic
23 performance of this integrated device is shown in **Fig. 6e**, where the tandem cell achieves a remarkable
24 PCE of 18.07% with a record-high V_{OC} of 2.13 V, making it highly suitable for driving electrochemical
25 loads. **Fig. 6f** displays the operational stability of this system under chopped illumination, delivering a
26 water-splitting current density of ~ 10.65 mA cm⁻² without external bias. This corresponds to a STH
27 efficiency of 13.10%, establishing a new benchmark for all-inorganic perovskite-based solar fuel



1 production. The stability of the inorganic narrow-bandgap absorber, a critical component of this system,
 2 is highlighted in **Fig. 6g**. The unencapsulated device retained over 82% of its initial efficiency after 504
 3 hours of thermal aging at 70°C, demonstrating exceptional robustness compared to hybrid counterparts.



5
 6 **Fig. 7** (a) Schematic diagram of perovskite tandem PV-driven water splitting. The inset is a photo of
 7 the IrO_x anode, Pt cathode, and Ag/AgCl reference electrode in an electrolyzer. (b) J - V curves of an
 8 all-perovskite tandem solar cell under simulated 1 sun illumination measured from reverse and forward
 9 scans. The inset is the stabilized efficiency measurement of the tandem cell for 300 s. (c) Current density
 10 and corresponding STH conversion efficiency of the tandem PV-driven EC device at zero external bias
 11 under chopped 1 sun illumination. Adapted with permission.³⁸ Copyright 2023, American Chemical
 12 Society. (d) Schematic diagram of preparation of nonprecious metal catalysts by electrodeposition and
 13 corresponding SEM images of NiFe, CoFe, and NiCo on ITO substrates. (e) J - V characteristics of NiFe,
 14 CoFe, and NiCo on ITO substrates, corrected by iR drop. (f) J - V characteristics of the two-electrode
 15 electrochemical cell and the all-perovskite tandem solar cell. Adapted with permission.⁶⁹ Copyright
 16 2025, American Chemical Society.

17

18 To harness the high photovoltage potential of perovskite multi-junction cells for practical fuel
 19 production, Song et al. constructed a PV-EC system driven by a monolithic all-perovskite tandem solar
 20 cell.³⁸ **Fig. 7a** illustrates the system configuration, where the tandem solar cell is wired to a water
 21 electrolyzer comprising a Pt cathode and an IrO_x anode in an acidic electrolyte. This physical separation
 22 allows for the optimization of the photovoltaic and electrochemical components independently. The
 23 driving force of this system is the high-performance all-perovskite tandem solar cell; as shown in the J -



1 V characteristics in **Fig. 7b**, the device delivers a remarkable V_{OC} exceeding 2.1 V and a fill factor over
2 80%. This high voltage is critical for driving the water splitting reaction without external bias.
3 Consequently, the integrated PV-EC system demonstrates exceptional performance. **Fig. 7c** presents
4 the operation of the device under chopped 1-sun illumination at zero external bias, where it maintains
5 a steady photocurrent density of approximately 14.1 mA cm⁻². This performance translates to a
6 benchmark STH conversion efficiency of 17.2%, confirming that all-perovskite tandem solar cells can
7 effectively power high-efficiency unassisted water splitting.

9 3.2.2 Perovskite tandem PV driven seawater electrolysis

10 Toward practical water electrolysis, it is necessary to check the operation in the seawater
11 environment. While seawater represents 96.5% of water resources from the Earth, its direct utilization
12 in electrolysis is severely hampered by the presence of chloride ions, which trigger the competitive
13 chlorine evolution reaction and accelerate anode corrosion. Addressing these challenges requires
14 electrocatalysts with exceptional selectivity for the OER over chlorine evolution. Li et al. proposed a
15 solution by synthesizing a cobalt-iron (CoFe) catalyst on indium tin oxide (ITO) *via* a rapid in situ
16 electrodeposition technique (**Fig. 7d**).⁶⁹ This catalyst features a nanoporous architecture that maximizes
17 the active surface area and exhibits superhydrophilicity, facilitating the rapid detachment of gas bubbles
18 during electrolysis. In electrochemical characterization within simulated alkaline seawater, the CoFe
19 electrode demonstrated superior catalytic activity, achieving a low overpotential of 268 mV at 10 mA
20 cm⁻² (**Fig. 7e**). This performance notably surpassed that of other bimetallic catalysts such as NiFe (315
21 mV) and NiCo (330 mV), confirming its high selectivity for OER. To realize unassisted solar hydrogen
22 production, the authors integrated this robust catalyst with perovskite tandem solar cells in a PV-EC
23 configuration. The integrated system operated at a voltage of ~1.56 V with a current density of 11.52
24 mA cm⁻², yielding a remarkable STH efficiency of 14.18% (**Fig. 7f**). The system demonstrated stable
25 operation for over 10 hours with approximately 90% retention of its initial current density. Furthermore,



1 a comprehensive techno-economic analysis projected a levelized cost of hydrogen of 7.17 \$ kg⁻¹,
2 suggesting a promising pathway toward cost-competitive green hydrogen generation.

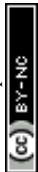
3 All-perovskite tandem structures have rapidly evolved from proof-of-concept devices to
4 highly efficient and stable systems. Recent progress includes the development of Sn-free narrow
5 bandgap materials to enhance stability, the adoption of all-inorganic compositions for improved
6 environmental resistance, and detailed investigations of interfacial degradation mechanisms.
7 Nevertheless, the narrow bandgap perovskite subcell often exhibits poor chemical and thermal stability,
8 posing a critical limitation to long-term device reliability.⁸⁴ Additionally, the use of similar polar
9 solvents for both subcells often causes solvent-induced damage to the bottom perovskite layer during
10 top-cell deposition, complicating tandem device fabrication.⁸⁵

12 3.3 Perovskite-organic tandem PV driven water electrolysis systems

13 **Table 3** Perovskite-organic hybrid tandem PV driven water electrolysis systems

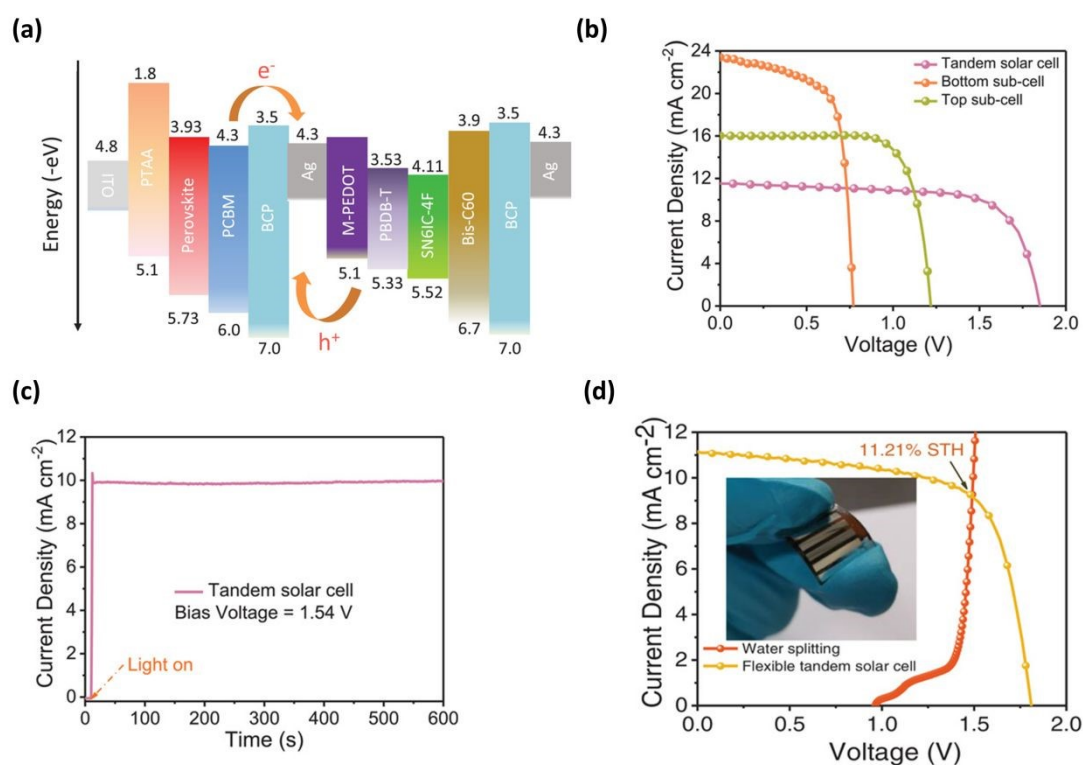
Type	Tandem PV subcells	PV area (cm ²)	Catalyst (HEC OEC)	Electrolyte	Stability	Photocurrent (mA cm ⁻²)	V _{OP} (V)	STH (%)	Year	Ref.
2T	MAPbI _{2.95} Cl _{0.05} /PM6:Y6	1	Pt Ni foam	1 M NaOH	2000h (91% in N ₂)	9.1	1.68	11.2	2022	[82]
2T	Cs _{0.1} (FA _{0.6} MA _{0.4}) _{0.9} Pb(I _{0.6} Br _{0.4}) ₃ /PBDB-T:SN61C-4F	0.13	Pt/C NiFe/C NT	1 M KOH	-	10	1.5	12.3	2020	[86]
2T	FA _{0.8} Cs _{0.2} Pb(I _{0.5} Br _{0.5}) ₃ /PM6:PM7:Y6:PC ₇₁ B M	1	Pt/C/CFP Ni-TFBDC/CF P	1 M KOH	-	13.62	-	16.75	2023	[87]

14
15 Perovskite-organic hybrid tandem structures integrate the complementary strengths of
16 perovskite and organic light absorbers within a single architecture. By combining a wide bandgap
17 perovskite top absorber with a narrow bandgap organic bottom absorber, the hybrid configuration
18 converts high-energy photons efficiently while allowing the organic layer to harvest longer wavelengths
19 that pass through the upper cell. All-organic tandems provide mechanical flexibility and can be



1 fabricated at low temperatures, yet their intrinsically small photovoltage and limited operational
 2 stability constrain practical PV-EC use. Conversely, all-perovskite tandems deliver larger voltages and
 3 superior charge transport but remain vulnerable to moisture ingress and interfacial degradation,
 4 especially in the narrow bandgap perovskites. This spectral splitting not only raises the overall
 5 photovoltage to levels sufficient for unbiased water electrolysis but also enhances durability: the
 6 inorganic top cell blocks ultraviolet and high-energy radiation, mitigating photochemical stress in the
 7 underlying organic layer.^{88, 89}

9 3.3.1 Strategies for high-voltage and flexible tandem PVs in water electrolysis applications



10

11 **Fig. 8** (a) Energy level diagram of a perovskite-organic tandem solar cell. (b) The *J-V* curves of a wide
 12 bandgap perovskite Cs_{0.1}(FA_{0.6}MA_{0.4})_{0.9}Pb(I_{0.6}Br_{0.4})₃ top subcell, a narrow bandgap PBDB-T:SN61C-
 13 4F organic bottom subcell, and the perovskite-organic tandem solar cell under AM 1.5G illumination.
 14 (c) The stabilized photocurrent of the perovskite-organic tandem solar cell was plotted as a function of
 15 continuous illumination time at the MPP condition with a steady bias voltage of 1.54 V. (d) The *J-V*
 16 curves of a flexible perovskite-organic tandem solar cell under simulated AM 1.5G illumination and
 17 NiFe LDH electrodes in a two-electrode system used for water splitting (inset is the photograph of the
 18 flexible tandem solar cell). Adapted with permission.⁸⁶ Copyright 2020, WILEY-VCH Verlag GmbH
 19 & Co. KGaA, Weinheim.

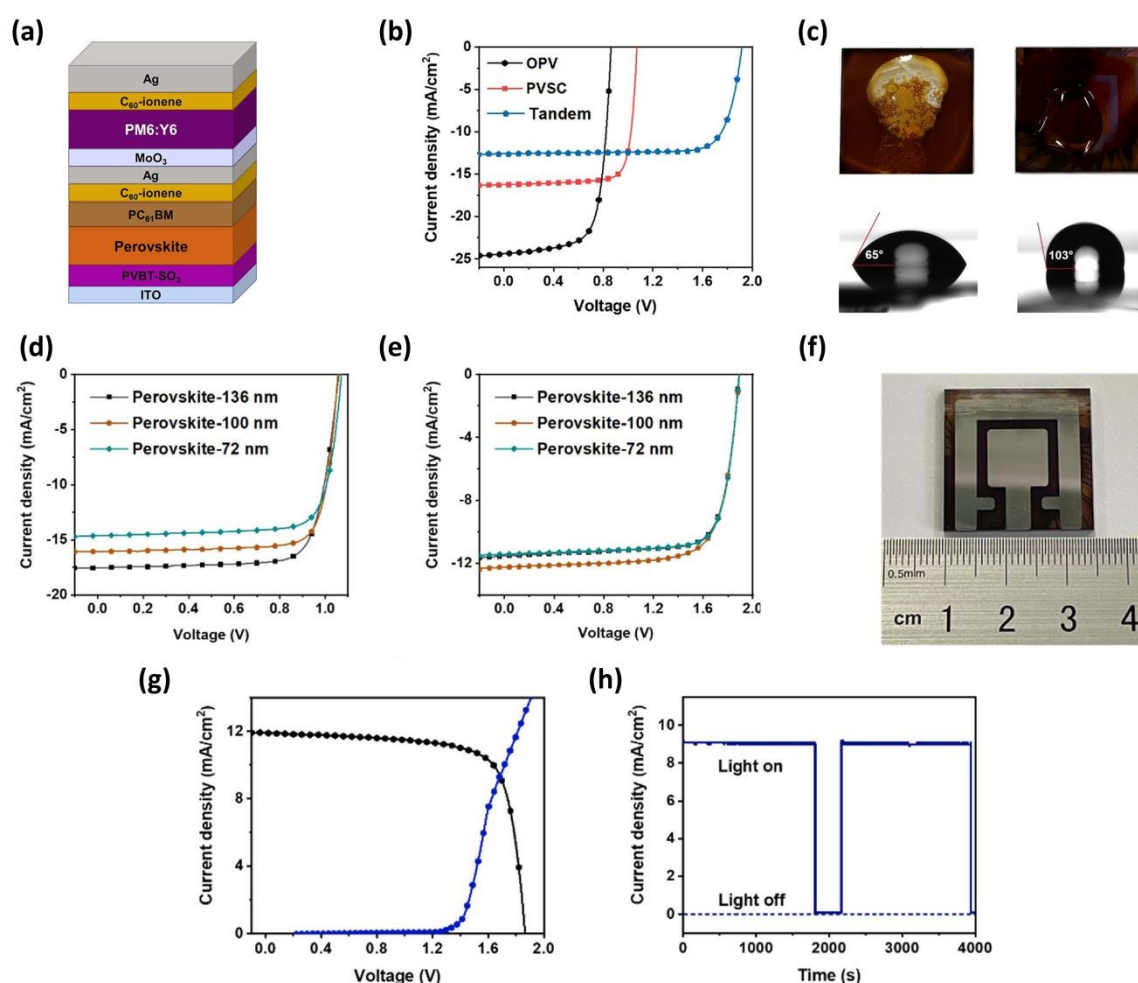


1
2 Li et al. designed a hybrid tandem solar cell combining a wide bandgap perovskite top cell and
3 a narrow bandgap organic bottom cell to secure the high V_{OC} essential for efficient water electrolysis
4 driving.⁸⁶ To minimize voltage loss in the 1.74 eV bandgap $Cs_{0.1}(FA_{0.6}MA_{0.4})_{0.9}Pb(I_{0.6}Br_{0.4})_3$ perovskite
5 film, they employed an interfacial passivation strategy using phenmethylammonium bromide (PMABr).
6 As illustrated in the energy level diagram in **Fig. 8a**, a PMABr layer was inserted between the perovskite
7 and the electron transport layer (PCBM) within the p-i-n configuration
8 (ITO/PTAA/perovskite/PCBM/BCP/Ag). This passivation treatment effectively passivated defects at
9 grain boundaries and surfaces, reducing trap density and suppressing non-radiative recombination,
10 which contributed to achieving an enhanced V_{OC} of 1.22 V in the single-junction device. By
11 monolithically integrating the optimized wide bandgap perovskite top cell with a PBDB-T:SN6IC-4F
12 based organic bottom cell (1.30 eV), a two-terminal perovskite-organic tandem solar cell was
13 fabricated. As evidenced by the $J-V$ characteristics in **Fig. 8b**, the rigid substrate-based tandem device
14 achieved a PCE of 15.13%, with a remarkable V_{OC} of 1.85 V, a short-circuit current density (J_{SC}) of
15 11.52 mA cm⁻², and a FF of 70.98%. Notably, the high V_{OC} of 1.85 V sufficiently exceeds the
16 thermodynamic potential required for water splitting (1.23 V), demonstrating the potential to drive an
17 electrolysis system without external bias. To evaluate device stability, the stabilized power output was
18 measured at the maximum power point (1.54 V) as shown in **Fig. 8c**. The device maintained a stable
19 efficiency of approximately 14.98% for 600 seconds without initial degradation, suggesting that
20 PMABr passivation effectively suppressed light-induced phase segregation common in mixed-halide
21 perovskites, thereby enhancing operational stability. Leveraging the low-temperature processing
22 advantages of the materials, the study also successfully implemented the tandem architecture on a
23 flexible substrate (PET/ITO). As presented in **Fig. 8d**, the flexible tandem solar cell exhibited excellent
24 performance comparable to its rigid counterpart, recording a PCE of 13.61% and a V_{OC} of 1.80 V.
25 Furthermore, when this flexible tandem device was coupled with a water splitting electrolyzer
26 composed of a NiFe/CNT oxygen evolution catalyst and a Pt/C hydrogen evolution catalyst, it achieved



1 a STH efficiency of 11.21%, as indicated by the operating point in **Fig. 8d**. These results demonstrate
 2 that solution-processed perovskite-organic hybrid tandem technology is a promising route not only for
 3 high-efficiency, but also for light-weight hydrogen production systems such as floating solar fuel
 4 production on water as discussed in Section 2.4.

6 3.3.2 Scalability strategies in monolithic perovskite-organic tandem structure



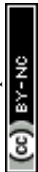
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8 **Fig. 9** (a) Device structure of a $\text{MAPbI}_{2.95}\text{Cl}_{0.05}/\text{PM6:Y6}$ tandem solar cell. (b) J - V curves of
 9 $\text{MAPbI}_{2.95}\text{Cl}_{0.05}$ (denoted as PVSC) and PM6:Y6 (denoted as OPV) subcells, and the tandem solar cell.
 10 (c) Photographs of water droplets on the surface of the $\text{MAPbI}_{2.95}\text{Cl}_{0.05}$ solar cell (left) and the tandem
 11 solar cell (right) and corresponding contact angles. (d) J - V curves of representative $\text{MAPbI}_{2.95}\text{Cl}_{0.05}$
 12 solar cells and (e) corresponding tandem devices with different perovskite thicknesses. (f) Photograph
 13 of a 1 cm^2 tandem solar cell. (g) J - V curves of the tandem solar cell under simulated AM 1.5G
 14 illumination and the electrochemical cell (Pt||Ni foam) for water splitting. (h) Current density-time
 15 curve of the water splitting device under chopped AM 1.5G illumination. Adapted with permission.⁸²
 16 Copyright 2022, American Chemical Society.



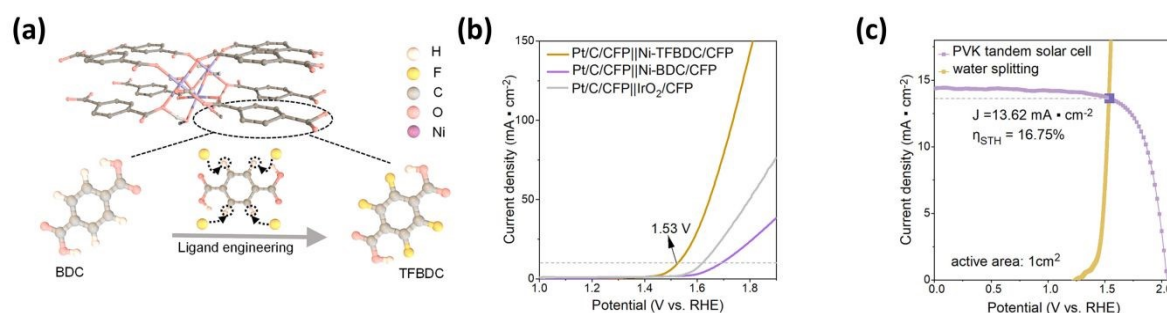
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Zhang et al. developed a monolithic perovskite-organic tandem solar cell structure that integrates a methylammonium lead perovskite front cell with a PM6:Y6 organic back cell to achieve high performance through solution processing.⁸² As illustrated in the device architecture (**Fig. 9a**), the two subcells are electrically connected *via* a robust, solvent-resistant interconnecting layer consisting of C₆₀-ionene/Ag/MoO₃, which is designed to form ohmic contacts with low contact resistance. This configuration enables the use of an ultrathin perovskite layer (~100 nm), minimizing lead content while maintaining high efficiency. The photovoltaic performance of the tandem device significantly surpasses that of the individual subcells, as evidenced by the *J-V* curves shown in **Fig. 9b**. The champion tandem device achieved a PCE of 19.2%, with a V_{OC} of 1.92 V, a J_{SC} of 12.6 mA cm⁻², and a FF of 79%. The high V_{OC} , which nearly equals the sum of the subcell voltages, confirms the effectiveness of the C₆₀-ionene-based interconnecting layer in facilitating charge recombination. Beyond efficiency, the tandem design offers superior stability against moisture compared to single-junction perovskite cells. When exposed to water droplets, the hydrophobic organic rear cell acts as a protective barrier, preventing the rapid decomposition observed in bare perovskite films. As quantified in **Fig. 9c**, the tandem device exhibited a water contact angle of 103°, significantly higher than the 65° measured for the perovskite single-junction device, attributing the enhanced environmental stability to the hydrophobic nature of the polymer active layer. To optimize the tandem performance, the thickness of the perovskite front cell was carefully tuned to ensure current matching. While reducing the perovskite thickness from approximately 136 nm to 72 nm resulted in a decrease in current for single-junction cells (**Fig. 9d**), the tandem devices incorporating an ~100 nm perovskite layer yielded the highest efficiencies (**Fig. 9e**). This thickness provides an optimal balance, allowing sufficient transparency for the bottom organic cell while maintaining adequate absorption for the top cell, whereas thicker films (~450 nm) were shown to block a significant portion of the visible light required by the rear cell. Demonstrating the scalability of this approach, the authors successfully fabricated large-area tandem devices (1 cm²) using the thickness-insensitive interconnecting layer (**Fig. 9f**), achieving a PCE of 17.8% with high reproducibility. These



1 1 cm² tandem solar cells were further utilized to drive solar-to-hydrogen conversion. The *J-V*
 2 characteristics in **Fig. 9g** show the intersection of the tandem cell and the electrolyzer curves, resulting
 3 in an operating current density of 9.1 mA cm⁻² and a STH conversion efficiency of 11.2%. The system
 4 demonstrated stable operation under chopped illumination (**Fig. 9h**), highlighting the potential of these
 5 solution-processed perovskite-organic tandem solar cells for efficient and practical green hydrogen
 6 production.

8 3.3.3 Achieving high STH efficiency via PV-catalyst optimization



9
 10 **Fig. 10** (a) Illustration of the ligand engineering process used to prepare Ni-TFBDC. (b)
 11 Electrochemical water splitting curves of Pt/C/CPF||Ni-TFBDC/CFP, Pt/C/CPF||Ni-BDC/CFP, and
 12 Pt/C/CPF||IrO₂/CFP cells. (c) *J-V* curve of the FA_{0.8}Cs_{0.2}Pb(I_{0.5}Br_{0.5})₃/PM6:PM7:Y6:PC₇₁BM
 13 tandem solar cell and the Pt/C/CPF||Ni-TFBDC/CFP cell. Adapted with permission.⁸⁷ Copyright 2023,
 14 American Chemical Society.

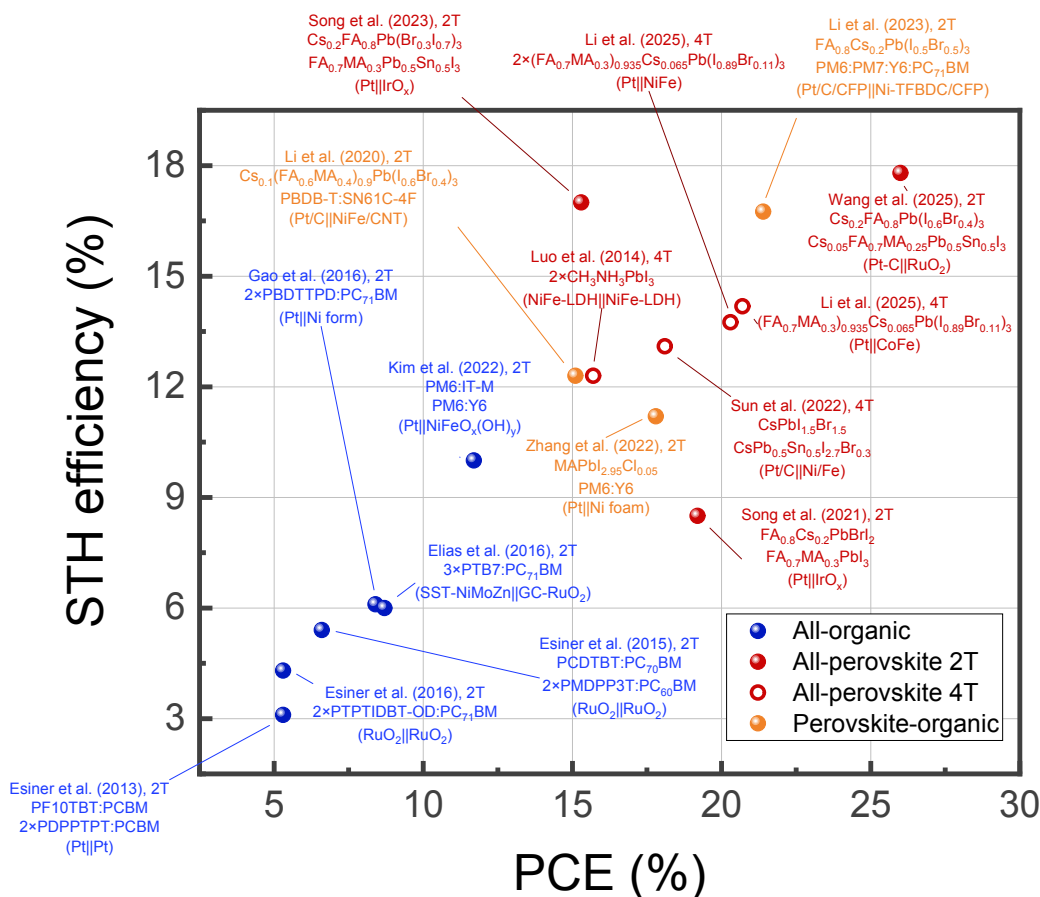
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 16 Li et al. demonstrated a strategy to enhance the intrinsic activity of metal-organic framework
 17 (MOF) electrocatalysts for oxygen evolution reaction through ligand engineering, which was
 18 subsequently integrated with perovskite photovoltaics for efficient solar-to-hydrogen conversion.⁸⁷ As
 19 illustrated in **Fig. 10a**, the authors synthesized a fluorinated MOF, denoted as Ni-TFBDC, by
 20 substituting the terephthalic acid (BDC) ligands in the nickel-coordinated framework with
 21 tetrafluoroterephthalic acid (TFBDC). The substitution of BDC with TFBDC significantly boosts
 22 oxygen evolution reaction performance by synergistically modulating the electronic structure,
 23 conductivity, and surface properties of the MOF. The introduction of highly electronegative fluorine



1 atoms lowers the oxidation state of nickel centers, as confirmed by X-ray absorption spectroscopy and
2 DFT calculations, which facilitates the reconstruction into highly active γ -NiOOH species during
3 catalysis. This ligand engineering also narrows the bandgap from 0.431 eV to 0.235 eV and drastically
4 reduces charge transfer resistance, thereby accelerating electron transport. Furthermore, steric repulsion
5 between fluorine atoms expands the lattice and reduces crystallinity, leading to a substantial increase in
6 specific surface area and hydrophilicity, while simultaneously shifting the reaction pathway toward the
7 kinetically favorable lattice oxygen mechanism. The electrocatalytic performance of the resulting
8 material was evaluated in a two-electrode water splitting configuration using a Pt/C cathode, as shown
9 in **Fig. 10b**. The Ni-TFBDC-based cell exhibited superior performance, requiring a voltage of only 1.53
10 V to drive a current density of 10 mA cm⁻², which notably outperformed the control device based on
11 Ni-BDC (1.69 V) and the commercial IrO₂-based system (1.61 V). To achieve unbiased solar water
12 splitting, the Ni-TFBDC electrolyzer was coupled with a high-voltage perovskite-organic tandem solar
13 cell. **Fig. 10c** displays the intersection of the photovoltaic J - V curve and the electrolyzer polarization
14 curve, indicating an operating current density of 13.62 mA cm⁻². This configuration resulted in a
15 remarkable STH efficiency of 16.75% under AM 1.5G illumination.

16



1 **4 Conclusion and prospects**

2
3 **Fig. 11** Summary of the relationship between the PCE of the tandem PVs and STH efficiency of solar
4 water splitting driven by corresponding tandem PVs. The data points differ by the type of tandem PV
5 configurations: all-organic tandem (blue), all-perovskite tandem (red), and perovskite-organic tandem
6 (orange). Annotations indicate the tandem PV subcells, two- or four-terminal (2T or 4T) PV
7 configuration, and catalysts used for the HER and the OER (HEC||OEC).

8
9 In this review, we have examined three types of tandem solar cell architectures that are
10 compatible with solution-based fabrication processes: organic tandem, halide perovskite tandem, and
11 hybrid tandem structures. For each configuration, we investigated how they have been implemented in
12 PV-EC water electrolysis systems, highlighting the specific design strategies and material optimizations
13 employed to enhance device performance and achieve sufficient photovoltage for overall water
14 splitting. Through this analysis, we aimed to elucidate the potential and limitations of each tandem
15 structure in the context of bias-free, efficient solar hydrogen production.



1 Organic tandem PV structures offer advantages in terms of solution processability and
2 mechanical flexibility. However, due to their intrinsically low charge carrier mobility and limited
3 photovoltage, they face challenges in supplying sufficient voltage for water electrolysis.^{90,91} In addition,
4 poor ultraviolet stability and degradation under operational conditions hinder their long-term durability.
5 The maximum STH efficiency stands at 10% driven by the organic-organic tandem PV, representing a
6 relatively modest performance.⁶³

7 Leveraging the superior optoelectronic properties of perovskites, perovskite-perovskite
8 tandem PV configurations have emerged as the most promising candidates for water electrolysis,
9 achieving STH efficiencies approaching 17.8%. However, challenges regarding the difficulty of large-
10 area fabrication remain significant hurdles. Also, the chemical and thermal instability of narrow
11 bandgap perovskites — particularly their sensitivity to moisture and oxygen — negatively impacts
12 device stability and longevity.^{92, 93}

13 The perovskite-organic tandem PV structure combines a wide bandgap perovskite top cell with
14 a narrow bandgap organic bottom cell, enabling high operating voltage and broad spectral absorption.
15 Perovskites provide strong visible absorption, while organics enhance near-infrared response. Their
16 compatibility with solution processing and flexible substrates makes this architecture attractive for
17 lightweight, large-area, and bias-free water electrolysis applications. This configuration has already
18 demonstrated sufficient photovoltage for overall water splitting.⁹⁴⁻¹⁰¹ Perovskite-organic tandem PVs
19 achieved a maximum STH efficiency of 16.38%.⁸⁷ However, stability remains a challenge due to halide
20 segregation in wide-bandgap perovskites, photodegradation of narrow bandgap polymers, and general
21 vulnerability of the absorber layers to moisture, oxygen, and thermal stress.¹⁰²⁻¹⁰⁸ Recent approaches to
22 mitigate these issues include the use of fully inorganic perovskites.^{95, 100, 109-112}



1 **Table 4** Performance of 2T tandem PV driven water electrolysis systems

Year	PV						EC							Ref.
	Tandem PV subcells	PV area (cm ²)	J_{sc} (mA cm ⁻²)	V_{oc} (V)	FF (%)	PCE (%)	Catalyst (HEC OEC)	Catalyst area (cm ²)	Electrolyte	Stability	Photocurrent (mA cm ⁻²)	V_{OP} (V)	STH (%)	
2013	PF10TBT:PCBM / PDPPTPT:PCBM / PDPPTPT:PCBM	0.09 ~ 0.16	4.42	2.23	51	5.3	Pt Pt	5 cm and 0.35 mm in diameter	1 M KOH	-	-	1.70	3.1	[58]
2015	PCDTBT:PC ₇₀ BM / PMDPP3T:PC ₆₀ BM / PMDPP3T:PC ₆₀ BM	0.0676	5.81	2.03	57	6.7	RuO ₂ RuO ₂	1.1 and 1.3	1 M KOH	-	4.4	1.49	5.41	[59]
2015	PCDTBT:PC ₇₀ BM / PMDPP3T:PC ₆₀ BM / PMDPP3T:PC ₆₀ BM	0.0676	5.78	2.00	56	6.5	Co ₃ O ₄ /NiMoZn	1	0.1 M KBi	-	3.98	1.56	4.89	[59]
2015	PCDTBT:PC ₇₀ BM / PMDPP3T:PC ₆₀ BM / PMDPP3T:PC ₆₀ BM	1.7	5.55	2.03	54	6.1	RuO ₂ RuO ₂	1.2	1 M KOH	-	2.94	1.67	3.61	[59]
2016	PTPTIDBT-OD:PC ₇₁ BM / PTPTIDBT-OD:PC ₇₁ BM	0.0676	4.14	1.76	73	5.3	RuO ₂ RuO ₂	1.1 and 1.3	1 M KOH	-	3.46	1.50	4.3	[60]
2016	PTB7:PC ₇₁ BM / PTB7:PC ₇₁ BM / PTB7:PC ₇₁ BM	0.09	5.4	2.13	76	8.7	SST-NiMoZn GC-RuO ₂	-	0.1 M KPi	50 h (79%) ^a	4.53	1.75	6.0	[61]
2016	PBDTTPD:PC ₇₁ BM / PBDTTPD:PC ₇₁ BM / PBDTTPD:PC ₇₁ BM	0.1	3.95	2.75	68.15	7.42	Pt Ni foam	-	1 M NaOH	-	5.4	1.5	6.1	[62]
2020	Cs _{0.1} (FA _{0.6} MA _{0.4}) _{0.9} Pb(I _{0.6} Br _{0.4}) ₃ / PBDB-T:SN61C-4F	0.13	11.52	1.85	70.98	15.13	Pt/C NiFe/CNT	-	1 M KOH	-	10	1.5	12.3	[86]
2021	FA _{0.8} Cs _{0.2} PbBrI ₂ / FA _{0.7} MA _{0.3} PbI ₃	0.12	11.13	2.153	80	19.2	Pt IrO _x	-	0.5 M H ₂ SO ₄	-	~7	-	8.5	[67]
2022	PM6:IT-M / PM6:Y6	0.56	9.81	1.84	65	11.7	Pt NiFeO _x (OH) _y	2	1 M KOH	2 h (80.8%)	8.2	1.46	10.0	[63]
2022	MAPbI _{2.95} Cl _{0.05} / PM6:Y6	1	12.6	1.92	79	15.6	Pt Ni foam	-	1 M NaOH	2000h (91% in N ₂)	9.1	1.68	11.2	[82]
2023	Cs _{0.2} FA _{0.8} Pb(Br _{0.3} I _{0.7}) / FA _{0.7} MA _{0.3} Pb _{0.5} Sn _{0.5} I ₃	0.1	15.23	2.104	81.9	26.2	Pt IrO _x	-	0.5 M H ₂ SO ₄	120 h (95%)	14.1	-	17	[38]
2023	FA _{0.8} Cs _{0.2} Pb(I _{0.5} Br _{0.5}) / PM6:PM7:Y6:PC ₇₁ BM	1	14.23	2.05	73.9	21.38	Pt/C/CFP Ni-TFBDC/CFP	-	1 M KOH	-	13.62	-	16.75	[87]
2025	Cs _{0.2} FA _{0.8} Pb(I _{0.6} Br _{0.4}) / Cs _{0.05} FA _{0.7} MA _{0.25} Pb _{0.5} Sn _{0.5} I ₃	1	16.0	2.08	78	26.0	Pt/C RuO ₂	1	H ₂ O	184 h (over 60%)	14.7	1.42	17.8	[39]

^a Tested under day/night cycles (16 h light/8 h dark) with UV filter.

2
3

1 **Table 5** Performance of 4T tandem PV driven water electrolysis systems

Year	PV						EC						Ref.	
	Tandem PV subcells	PV area (cm ²)	J_{sc} (mA cm ⁻²)	V_{oc} (V)	FF (%)	PCE (%)	Catalyst (HEC OEC)	Catalyst area (cm ²)	Electrolyte	Stability	Photocurrent (mA cm ⁻²)	V_{OP} (V)		STH (%)
2014	CH ₃ NH ₃ PbI ₃ / CH ₃ NH ₃ PbI ₃	0.318	10.0	2.00	-	15.7	NiFe-LDH NiFe-LDH	5	1 M NaOH	2 h (~80%)	9.61	1.63	12.3	[68]
2022	CsPbI _{1.5} Br _{1.5} / CsPb _{0.5} Sn _{0.5} I _{2.7} Br _{0.3}	0.1	11.56	2.13	73.4	18.07	Pt/C Ni/Fe	-	1 M KOH	2 h (97.2%)	10.71	1.62	13.1	[43]
2025	PBQx-TF:FPCC-Br / PBQx-TF:FPCC-Br	1 and 1 (parallel illumination)	7.90	2.16	69.7	11.9	Pt Pt	-	1M KOH	600 s (72%)	5.62	1.93	6.91	[64]
2025	(FA _{0.7} MA _{0.3}) _{0.935} Cs _{0.065} Pb(I _{0.89} Br _{0.11}) ₃ / (FA _{0.7} MA _{0.3}) _{0.935} Cs _{0.065} Pb(I _{0.89} Br _{0.11}) ₃	1 and 1 (parallel illumination)	11.79	2.25	78	20.65	Pt CoFe	1	1 M KOH + 0.5 M NaCl	10 h (90%)	11.53	1.56	14.18	[69]
2025	(FA _{0.7} MA _{0.3}) _{0.935} Cs _{0.065} Pb(I _{0.89} Br _{0.11}) ₃ / (FA _{0.7} MA _{0.3}) _{0.935} Cs _{0.065} Pb(I _{0.89} Br _{0.11}) ₃	1 and 1 (parallel illumination)	11.62	2.28	-	-	Pt NiFe	2	1 M KOH	-	11.18	-	13.75	[70]

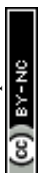
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1 To provide a comprehensive overview of the recent progress, **Fig. 11** summarizes the
2 relationship between the PCEs of tandem PVs discussed in this review and the STH efficiencies
3 obtained from water electrolysis systems driven by corresponding tandem PV devices, to enable a more
4 in-depth comparison between PV and EC studies, we include additional extended parameters and
5 present them in chronological order in the newly introduced **Tables 4 and 5**. The plot reveals a distinct
6 technological evolution: early research was dominated by all-organic tandems with modest efficiencies,
7 whereas recent advancements have been driven by all-perovskite tandems and perovskite-organic
8 tandems, which now occupy the high-efficiency regime. This trend illustrates a clear positive
9 correlation between PV performance and hydrogen production, confirming that high-performance
10 photovoltaic absorbers are a prerequisite for achieving STH values exceeding 15%. Furthermore, as
11 annotated in **Fig. 11**, these record-breaking efficiencies are being realized using a diverse range of
12 electrocatalysts — from noble metals (e.g., Pt, IrO_x) to earth-abundant transition metal compounds (e.g.,
13 NiFe-LDH) — marking a significant step toward practical and cost-effective solar fuel production.

14 To transition from laboratory-scale proof-of-concept devices to practical, scalable hydrogen
15 production systems, future research must converge on the following critical areas:

16 *1. Intrinsic stability of solution processable PV materials:* The stability demands placed on solution-
17 processable PV materials in integrated PV-EC systems are fundamentally distinct from those
18 encountered in grid-connected operation. For perovskite light absorbers, the central concern is the
19 diurnal cycling through multiple voltage states inherent to direct-coupled operation: at dawn and dusk,
20 when the electrolyser has not yet reached its onset threshold, the perovskite cell dwells near V_{OC} ,
21 precisely the condition shown to maximise charge accumulation, accelerate ion migration, and produce
22 the lowest operational stability, meaning the device experiences its most damaging electrical state
23 exactly when it is performing no useful water electrolysis. For organic light absorbers, the critical threats
24 are thermally driven morphological coarsening of the BHJ under outdoor temperature cycling, which
25 simultaneously degrades FF and J_{SC} , and the well-documented burn-in efficiency loss in the first tens
26 to hundreds of hours of operation. They carry a uniquely severe consequence in PV-EC: if efficiency



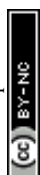
1 loss is sufficient to shift the operating voltage below the electrolyser onset threshold, hydrogen
2 production ceases entirely rather than merely declining. Compounding both material classes is the
3 proximity of the liquid electrolyte environment, which imposes moisture ingress and chemical
4 compatibility requirements on encapsulation that far exceed those of standard photovoltaic deployment,
5 alongside the toxicity constraint of preventing lead or organic component leaching into the electrolyte.
6 Critically, no current stability standard captures these coupled stressors, as none incorporates variable-
7 voltage operation along an electrolyser load line, start-stop cycling with open-circuit dwell, or
8 interfacial chemical exposure; prospective work must therefore develop PV-EC specific accelerated
9 stress test protocols that treat the photovoltaic and electrochemical subsystems as a coupled degrading
10 system rather than independently qualified components.

11 *2. System-level integration and geometric optimization: Achieving high-performance integrated PV-*
12 *EC systems based on solution-processable perovskite and organic absorbers demands a co-design*
13 *philosophy in which geometric, catalytic, and system-level parameters are treated as primary*
14 *optimisation variables from the outset, rather than as secondary engineering adjustments applied to*
15 *independently optimised components. The most critical geometric variable is the PV-to-electrolyser*
16 *area ratio, which determines the current density at which the electrolyser operates. For tandem*
17 *architectures combining perovskite front cells with organic rear cells, photocurrent matching between*
18 *subcells introduces an additional geometric constraint: optical and parasitic losses at the interconnection*
19 *layer directly limit the current delivered to the electrolyser, and sub-optimal matching carries a threshold*
20 *consequence in PV-EC operation whereby the system ceases hydrogen production entirely rather than*
21 *operating at reduced efficiency. Crucially, the cost reduction potential uniquely offered by solution-*
22 *processable PV materials can only be fully realised at the system level if the electrocatalyst cost is*
23 *addressed in parallel, the replacement of precious metal catalysts such as platinum and iridium oxide*
24 *with earth-abundant alternatives, including transition metal phosphides, sulfides, and layered double*
25 *hydroxides for the hydrogen and oxygen evolution reactions respectively, is therefore not merely a*
26 *materials science objective but a system-level economic necessity, since retaining rare-metal catalysts*



1 in a system built around low-cost solution-processed absorbers creates a cost asymmetry that
2 undermines the economic case for the entire integration strategy. Prospectively, realising the full
3 optimisation potential of these materials will require integrated multiphysics modelling frameworks that
4 simultaneously treat optical design, electrochemical load matching, area ratio, earth-abundant catalyst
5 selection, encapsulation geometry, and deployment configuration as a coupled optimisation problem: a
6 level of co-design sophistication that the field has not yet systematically achieved.

7 *3. Scalability and technical challenges:* Solution-processed 2T tandem PV devices have recently
8 emerged as a promising platform for highly efficient solar-driven hydrogen production, demonstrating
9 sufficiently high photovoltage and excellent fill factor for unbiased water splitting at the laboratory cell
10 level. Representative devices have reported V_{OC} exceeding 1.8 V, in some cases surpassing 2 V, together
11 with FF over 80%, clearly indicating the potential for self-sustained water electrolysis without external
12 bias. However, extending such performance to large-area modules requires precise control of film
13 formation uniformity, which is an inherent limitation of solution-based processes. It also demands
14 careful optimization of the resistance–transmittance characteristics of transparent electrodes, interfacial
15 quality, and current matching between serially connected subcells. During scaling, factors such as
16 increased sheet resistance of transparent electrodes, coating non-uniformity, crystallization variability,
17 and interfacial defects tend to accumulate, resulting in higher series resistance and non-ideal
18 recombination losses that degrade FF and sometimes V_{OC} .³⁸ Moreover, in the 2T tandem structure,
19 current mismatch between subcells constrains overall operating current, so even minor deviations in
20 optical or electrical matching can directly lead to significant module performance losses. While such
21 losses primarily appear as reduced power output in standalone PV systems, they become far more
22 detrimental in directly coupled PV-EC systems, where the operating voltage must remain above the
23 practical water electrolysis threshold to maintain unassisted operation and steady hydrogen
24 generation.¹¹³ Therefore, maintaining the high photovoltage and fill factor achieved at small scales in
25 module-level devices should be regarded not merely as a goal for efficiency enhancement but as a
26 prerequisite for stable, system-level hydrogen production.¹¹⁴ In particular, for PV-EC systems based on



1 solution-processed tandem devices, performance losses caused by film non-uniformity, current
2 mismatch, and interfacial defects during large-area fabrication are critical. Consequently, research on
3 large-area implementation remains markedly insufficient. This limitation acts as a key bottleneck,
4 hindering the translation of small-area high performance to module-scale devices and thereby delaying
5 stable hydrogen production at the system level. To achieve this, process stabilization strategies that can
6 precisely control film uniformity and interfacial quality in large-area solution processing are essential.
7 In particular, (i) controlling the fluid dynamics during coating to ensure meniscus stability, (ii) securing
8 interlayer process compatibility through immiscible or carefully designed orthogonal solvent
9 combinations, and (iii) employing flash drying, infrared, or thermal-assisted drying to regulate the
10 crystallization of the top layer while mitigating thermal stress on the underlying layers are required as
11 part of scalable and refined solution-processing techniques.^{115, 116} Meanwhile, (iv) the development of
12 low-resistance transparent electrodes and highly efficient interconnecting layer/recombination
13 junctions, (v) the use of bandgap combinations and optical–electrical designs less sensitive to spectral
14 variations, and (vi) the optimization of laser scribing-based monolithic interconnections serves as
15 critical factors in minimizing resistive and interconnection losses during large-area module
16 fabrication.^{117, 118} Module design must further ensure voltage retention and current scalability through
17 serial stack matching and serial expansion. In this architecture, scalability should not rely solely on
18 increasing the PV voltage itself but rather on proportional co-scaling of PV module voltage and
19 electrolyser stack voltage through serial interconnection.¹¹⁹ Furthermore, achieving long-term durability
20 requires encapsulation and interfacial stabilization techniques that suppress ion migration and
21 interfacial degradation, together with the establishment of accelerated lifetime evaluation protocols
22 under combined thermal, light, and electrical stresses.¹²⁰ Beyond efficiency enhancement, process
23 development should also account for manufacturing yield, reproducibility, PV-EC integration, and
24 techno-economic metrics based on the levelized cost of hydrogen.¹²¹ With these technological advances
25 and systematic process control, solution-processed 2T perovskite tandem PV-EC systems are expected
26 to realize both high-voltage operation and efficient unassisted water electrolysis, paving the way for
27 practical large-area manufacturing and sustainable STH production.



1 In conclusion, while solution-processed tandem PV-EC systems have made remarkable strides
2 in efficiency, the focus must now shift toward holistic system engineering that simultaneously addresses
3 stability, scalability, and cost. By converging material innovations with rational system design, these
4 technologies hold the potential to become a cornerstone of the future green hydrogen economy.
5

6 **Author contributions**

7 H.-B. K. led the review and carried out the investigation and visualization. W. C. surveyed the relevant
8 literatures. T. H. L. contributed to conceptualization and edited the manuscript.
9

10 **Conflicts of interest**

11 There are no conflicts to declare.
12

13 **Data availability**

14 No primary research results, software or code have been included and no new data were generated or
15 analysed as part of this review.
16

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20 **Notes and references**

21 1. A. M. Oliveira, R. R. Beswick and Y. Yan, *Curr. Opin. Chem. Eng*, 2021, **33**, 100701.



- 1 2. A. G. Olabi and M. A. Abdelkareem, *Renew. Sustain. Energy Rev.*, 2022, **158**, 112111.
- 2 3. B. C. Tashie-Lewis and S. G. Nnabuiife, *Chem. Eng. J. Adv.*, 2021, **8**, 100172.
- 3 4. H. H. Cho, V. Strezov and T. J. Evans, *Sustain. Mater. Technol.*, 2023, **35**, e00567.
- 4 5. N. Ma, W. Zhao, W. Wang, X. Li and H. Zhou, *Int. J. Hydrogen Energy*, 2024, **50**, 379-
5 396.
- 6 6. G. Squadrito, G. Maggio and A. Nicita, *Renew. Energy*, 2023, **216**, 119041.
- 7 7. F. Urbain, V. Smirnov, J.-P. Becker, U. Rau, J. Ziegler, B. Kaiser, W. Jaegermann and
8 F. Finger, *Sol. Energy Mater. Sol. Cells*, 2015, **140**, 275-280.
- 9 8. S. Hu, C. Xiang, S. Haussener, A. D. Berger and N. S. Lewis, *Energy Environ. Sci.*,
10 2013, **6**, 2984.
- 11 9. K. Zhang, M. Ma, P. Li, D. H. Wang and J. H. Park, *Adv. Energy Mater.*, 2016, **6**,
12 1600602.
- 13 10. J. Jia, L. C. Seitz, J. D. Benck, Y. Huo, Y. Chen, J. W. Ng, T. Bilir, J. S. Harris and T.
14 F. Jaramillo, *Nat Commun*, 2016, **7**, 13237.
- 15 11. C. Jiang, S. J. A. Moniz, A. Wang, T. Zhang and J. Tang, *Chem. Soc. Rev.*, 2017, **46**,
16 4645-4660.
- 17 12. F. Finger, K. Welter, F. Urbain, V. Smirnov, B. Kaiser and W. Jaegermann, *Z. Phys.*
18 *Chem.*, 2020, **234**, 1055-1095.
- 19 13. A. Vilanova, P. Dias, T. Lopes and A. Mendes, *Chem. Soc. Rev.*, 2024, **53**, 2388-2434.
- 20 14. F. Qureshi and M. Tahir, *Int. J. Hydrogen Energy*, 2024, **69**, 760-776.
- 21 15. A. F. Palmstrom, G. E. Eperon, T. Leijtens, R. Prasanna, S. N. Habisreutinger, W.
22 Nemeth, E. A. Gaulding, S. P. Dunfield, M. Reese, S. Nanayakkara, T. Moot, J. Werner,
23 J. Liu, B. To, S. T. Christensen, M. D. McGehee, M. F. A. M. van Hest, J. M. Luther, J.
24 J. Berry and D. T. Moore, *Joule*, 2019, **3**, 2193-2204.
- 25 16. Y. Jiang and Y. Qi, *Mater. Chem. Front.*, 2021, **5**, 4833-4850.



- 1 17. L. Mao, J. Tong, S. Xiong, F. Jiang, F. Qin, W. Meng, B. Luo, Y. Liu, Z. Li, Y. Jiang,
2 C. Fuentes-Hernandez, B. Kippelen and Y. Zhou, *J. Mater. Chem. A*, 2017, **5**, 3186-
3 3192.
- 4 18. H. Lai, J. Luo, Y. Zwirner, S. Olthof, A. Wiczorek, F. Ye, Q. Jeangros, X. Yin, F.
5 Akhundova, T. Ma, R. He, R. K. Kothandaraman, X. Chin, E. Gilshtein, A. Müller, C.
6 Wang, J. Thiesbrummel, S. Siol, J. M. Prieto, T. Unold, M. Stolterfoht, C. Chen, A. N.
7 Tiwari, D. Zhao and F. Fu, *Adv. Energy Mater.*, 2022, **12**, 2202438.
- 8 19. M. Raïssi, S. Wageh, A. A. Al-Ghamdi and D. Rousseau, *J. Mater. Chem. A*, 2023, **11**,
9 25578-25594.
- 10 20. H. Liu, M. H. Yu, C. C. Lee, X. Yu, Y. Li, Z. Zhu, C. C. Chueh, Z. a. Li and A. K. Y.
11 Jen, *Adv. Mater. Technol.*, 2021, **6**, 2000960.
- 12 21. P. Subudhi and D. Punetha, *Prog. Photovolt. Res. Appl.*, 2023, **31**, 753-789.
- 13 22. Y. Sun, T. Liu, Y. Kan, K. Gao, B. Tang and Y. Li, *Small Sci.*, 2021, **1**, 2100001.
- 14 23. N. Ahn and M. Choi, *Adv. Sci.*, 2024, **11**, e2306110.
- 15 24. D. Zhang, D. Li, Y. Hu, A. Mei and H. Han, *Commun. Mater.*, 2022, **3**, 58.
- 16 25. Z. Ni, H. Jiao, C. Fei, H. Gu, S. Xu, Z. Yu, G. Yang, Y. Deng, Q. Jiang, Y. Liu, Y. Yan
17 and J. Huang, *Nat. Energy*, 2021, **7**, 65-73.
- 18 26. Y. Zhao, Z. Wu, X. Liu, Z. Zhong, R. Zhu and J. Yu, *J. Mater. Chem. C*, 2021, **9**, 13972-
19 13980.
- 20 27. A. J. Clarke, J. Luke, R. Meitzner, J. Wu, Y. Wang, H. K. H. Lee, E. M. Speller, H.
21 Bristow, H. Cha, M. J. Newman, K. Hooper, A. Evans, F. Gao, H. Hoppe, I. McCulloch,
22 U. S. Schubert, T. M. Watson, J. R. Durrant, W. C. Tsoi, J.-S. Kim and Z. Li, *Cell Rep.*
23 *Phys. Sci.*, 2021, **2**, 100498.
- 24 28. J. K. Nørskov, J. Rossmeisl, A. Logadottir, L. Lindqvist, J. R. Kitchin, T. Bligaard and
25 H. Jonsson, *J. Phys. Chem. B*, 2004, **108**, 17886-17892.
- 26 29. J. Rossmeisl, Z. W. Qu, H. Zhu, G. J. Kroes and J. K. Nørskov, *J. Electroanal. Chem.*,
27 2007, **607**, 83-89.



- 1 30. S. Zhou, N. Liu, Z. Wang and J. Zhao, *ACS Appl. Mater. Interfaces*, 2017, **9**, 22578-
2 22587.
- 3 31. K. Zeng and D. Zhang, *Prog. Energy Combust. Sci.*, 2010, **36**, 307-326.
- 4 32. C. Xiang, K. M. Papadantonakis and N. S. Lewis, *Mater. Horiz.*, 2016, **3**, 169-173.
- 5 33. Y. Wang, Y. Wu, J. Schwartz, S. H. Sung, R. Hovden and Z. Mi, *Joule*, 2019, **3**, 2444-
6 2456.
- 7 34. T. Baumeler, N. Arora, A. Hinderhofer, S. Akin, A. Greco, M. Abdi-Jalebi, R. Shivanna,
8 R. Uchida, Y. Liu, F. Schreiber, S. M. Zakeeruddin, R. H. Friend, M. Graetzel and M. I.
9 Dar, *J. Phys. Chem. Lett.*, 2020, **11**, 10188-10195.
- 10 35. K. T. Fountaine, H. J. Lewerenz and H. A. Atwater, *Nat. Commun.*, 2016, **7**, 13706.
- 11 36. J.-F. Guillemoles, T. Kirchartz, D. Cahen and U. Rau, *Nat. Photonics*, 2019, **13**, 501-
12 505.
- 13 37. S. Rühle, *Sol. Energy*, 2016, **130**, 139-147.
- 14 38. Z. Song, C. Li, L. Chen, K. Dolia, S. Fu, N. Sun, Y. Li, K. Wyatt, J. L. Young, T. G.
15 Deutsch and Y. Yan, *ACS Energy Lett.*, 2023, **8**, 2611-2619.
- 16 39. J. Wang, B. Branco, W. H. M. Remmerswaal, S. Hu, N. R. M. Schipper, V. Zardetto, L.
17 Bellini, N. Daub, M. M. Wienk, A. Wakamiya, H. J. Snaith and R. A. J. Janssen, *Nat.*
18 *Commun.*, 2025, **16**, 174.
- 19 40. W. E. McMahon, J. F. Geisz, J. Buencuerpo and E. L. Warren, *Sustain. Energy Fuels*,
20 2023, **7**, 461-470.
- 21 41. R. Fan, S. Cheng, G. Huang, Y. Wang, Y. Zhang, S. Vanka, G. A. Botton, Z. Mi and M.
22 Shen, *J. Mater. Chem. A*, 2019, **7**, 2200-2209.
- 23 42. R. Witteck, J. F. Geisz, E. L. Warren and W. E. McMahon, *Sol. RRL*, 2023, **8**, 2300782.
- 24 43. Q. Sun, Z. Zhang, T. Zhang, Y. Feng, A. Gu, H. Yu, M. Zhang, X. L. Zhang, J. Zhu, Y.
25 Shen and M. Wang, *ACS Energy Lett.*, 2022, **7**, 4215-4223.
- 26 44. Z. Shokrollahi, M. Piralaee and A. Asgari, *Sci. Rep.*, 2024, **14**, 11515.



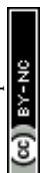
- 1 45. B. Abdollahi Nejand, D. B. Ritzer, H. Hu, F. Schackmar, S. Moghadamzadeh, T. Feeney,
2 R. Singh, F. Laufer, R. Schmager, R. Azmi, M. Kaiser, T. Abzieher, S. Gharibzadeh, E.
3 Ahlswede, U. Lemmer, B. S. Richards and U. W. Paetzold, *Nat. Energy*, 2022, **7**, 620-
4 630.
- 5 46. E. Raza and Z. Ahmad, *Energy Rep.*, 2022, **8**, 5820-5851.
- 6 47. Gurudayal, D. Sabba, M. H. Kumar, L. H. Wong, J. Barber, M. Gratzel and N. Mathews,
7 *Nano Lett.*, 2015, **15**, 3833-3839.
- 8 48. V. A. Martinez Lopez, H. Ziar, J. W. Haverkort, M. Zeman and O. Isabella, *Renew.*
9 *Sustain. Energy Rev.*, 2023, **182**, 113407.
- 10 49. C. Chen, L. Wang, W. Xia, K. Qiu, C. Guo, Z. Gan, J. Zhou, Y. Sun, D. Liu, W. Li and
11 T. Wang, *Nat. Commun.*, 2024, **15**, 6865.
- 12 50. Martin A. Green, Ewan D. Dunlop, M. Yoshita, N. Kopidakis, K. Bothe, G. Siefer, X.
13 Hao and Jessica Y. Jiang, *Prog. Photovolt: Res. Appl.*, 2025, **33**, 795-810.
- 14 51. J. Wang, J. Li, Y. Wang, J. Ren, P. Bi, H. Li, J. Dai, S. Zhang and J. Hou, *Adv. Mater.*,
15 2025, **37**, e10378.
- 16 52. Z. Liu, R. Lin, M. Wei, M. Yin, P. Wu, M. Li, L. Li, Y. Wang, G. Chen, V. Carnevali,
17 L. Agosta, V. Slama, N. Lempesis, Z. Wang, M. Wang, Y. Deng, H. Luo, H. Gao, U.
18 Rothlisberger, S. M. Zakeeruddin, X. Luo, Y. Liu, M. Gratzel and H. Tan, *Nat. Mater.*,
19 2025, **24**, 252-259.
- 20 53. X. Jiang, S. Qin, L. Meng, G. He, J. Zhang, Y. Wang, Y. Zhu, T. Zou, Y. Gong, Z. Chen,
21 G. Sun, M. Liu, X. Li, F. Lang and Y. Li, *Nature*, 2024, **635**, 860-866.
- 22 54. T. H. Lee, Y. Dong, R. A. Pacalaj, S. Y. Park, W. Xu, J. S. Kim and J. R. Durrant, *Adv.*
23 *Funct. Mater.*, 2022, **32**, 2208001.
- 24 55. M. V. Khenkin, E. A. Katz, A. Abate, G. Bardizza, J. J. Berry, C. Brabec, F. Brunetti,
25 V. Bulović, Q. Burlingame, A. Di Carlo, R. Cheacharoen, Y.-B. Cheng, A. Colmann,
26 S. Cros, K. Domanski, M. Dusza, C. J. Fell, S. R. Forrest, Y. Galagan, D. Di Girolamo,
27 M. Grätzel, A. Hagfeldt, E. von Hauff, H. Hoppe, J. Kettle, H. Köbler, M. S. Leite, S.



- 1 Liu, Y.-L. Loo, J. M. Luther, C.-Q. Ma, M. Madsen, M. Manceau, M. Matheron, M.
2 McGehee, R. Meitzner, M. K. Nazeeruddin, A. F. Nogueira, Ç. Odabaşı, A. Osherov,
3 N.-G. Park, M. O. Reese, F. De Rossi, M. Saliba, U. S. Schubert, H. J. Snaith, S. D.
4 Stranks, W. Tress, P. A. Troshin, V. Turkovic, S. Veenstra, I. Visoly-Fisher, A. Walsh,
5 T. Watson, H. Xie, R. Yıldırım, S. M. Zakeeruddin, K. Zhu and M. Lira-Cantu, *Nat.*
6 *Energy*, 2020, **5**, 35-49.
- 7 56. M. Cai, Y. Wu, H. Chen, X. Yang, Y. Qiang and L. Han, *Adv. Sci. (Weinh.)*, 2017, **4**,
8 1600269.
- 9 57. V. Andrei, G. M. Ucoski, C. Pornrunroj, C. Uswachoke, Q. Wang, D. S. Achilleos, H.
10 Kasap, K. P. Sokol, R. A. Jagt, H. Lu, T. Lawson, A. Wagner, S. D. Pike, D. S. Wright,
11 R. L. Z. Hoye, J. L. MacManus-Driscoll, H. J. Joyce, R. H. Friend and E. Reisner,
12 *Nature*, 2022, **608**, 518-522.
- 13 58. S. Esiner, H. van Eersel, M. M. Wienk and R. A. Janssen, *Adv. Mater.*, 2013, **25**, 2932-
14 2936.
- 15 59. S. Esiner, R. E. M. Willems, A. Furlan, W. Li, M. M. Wienk and R. A. J. Janssen, *J.*
16 *Mater. Chem. A*, 2015, **3**, 23936-23945.
- 17 60. S. Esiner, G. W. P. van Pruissen, M. M. Wienk and R. A. J. Janssen, *J. Mater. Chem. A*,
18 2016, **4**, 5107-5114.
- 19 61. X. Elias, Q. Liu, C. Gimbert-Suriñach, R. Matheu, P. Mantilla-Perez, A. Martinez-Otero,
20 X. Sala, J. Martorell and A. Llobet, *ACS Catalysis*, 2016, **6**, 3310-3316.
- 21 62. Y. Gao, V. M. Le Corre, A. Gaitis, M. Neophytou, M. A. Hamid, K. Takanabe and P.
22 M. Beaujuge, *Adv. Mater.*, 2016, **28**, 3366-3373.
- 23 63. Y. K. Kim, T. H. Lee, J. Yeop, W. J. Byun, J. H. Kim, J. Y. Kim and J. S. Lee, *Appl.*
24 *Catal. B: Environ.*, 2022, **309**, 121237.
- 25 64. Y. Xiao, J. Wang, Y. Cui, Y. Wang, Z. Chen, S. Cheng, H. Yuan, J. Qiao, Y. Yang, W.
26 Wang, N. Yang, Y. Yu, R. Yu, X. Hao and J. Hou, *Energy Environ. Sci.*, 2025, **18**, 3259-
27 3268.



- 1 65. Y. Fu, T. H. Lee, Y. C. Chin, R. A. Pacalaj, C. Labanti, S. Y. Park, Y. Dong, H. W. Cho,
2 J. Y. Kim, D. Minami, J. R. Durrant and J. S. Kim, *Nat. Commun.*, 2023, **14**, 1870.
- 3 66. Z. Li, S. Fang, H. Sun, R. J. Chung, X. Fang and J. H. He, *Adv. Energy Mater.*, 2023,
4 **13**, 2203019.
- 5 67. Z. N. Song, C. W. Li, L. Chen, S. Rijal, J. L. Young, T. G. Deutsch and Y. F. Yan, *IEEE*
6 *Photovolt. Spec. Conf.*, 2021, **20-25**, 2222-2225.
- 7 68. J. Luo, J. H. Im, M. T. Mayer, M. Schreier, M. K. Nazeeruddin, N. G. Park, S. D. Tilley,
8 H. J. Fan and M. Gratzel, *Science*, 2014, **345**, 1593-1596.
- 9 69. Y. Li, Y. Li, Z. Ma, K. Yue, Q. Yang, X. Li, Q. Zhang, F. Gou, H. Du, C. Cheng, M.
10 Mao, D. Xiang, Z. Lv, K. Liu, B. Chen, R. Xu, Q. Yin, B. Luo, J. Zhan, K. Sun, C. Tang
11 and Z. Pan, *ACS Appl. Mater. Interfaces*, 2025, **17**, 32530-32543.
- 12 70. Y. Li, Z. Ma, S. Hou, X. Li, S. Wang, Z. Du, Y. Chen, Q. Zhang, Y. Li, Q. Yang, Z.
13 Huang, L. Bai, H. Yu, Q. Liu, Y. Xiang, M. Zhang, J. Yu, J. Xie, Y. Zhou, C. Tang, K.
14 Sun and L. Ding, *J. Colloid Interface Sci.*, 2025, **677**, 599-609.
- 15 71. S. D. Stranks, G. E. Eperon, G. Grancini, C. Menelaou, M. J. Alcocer, T. Leijtens, L. M.
16 Herz, A. Petrozza and H. J. Snaith, *Science*, 2013, **342**, 341-344.
- 17 72. C. C. Stoumpos, C. D. Malliakas and M. G. Kanatzidis, *Inorg. Chem.*, 2013, **52**, 9019-
18 9038.
- 19 73. S. Tan, C. Li, C. Peng, W. Yan, H. Bu, H. Jiang, F. Yue, L. Zhang, H. Gao and Z. Zhou,
20 *Nat. Commun.*, 2024, **15**, 4136.
- 21 74. J. Zhou, H. Qiu, T. Wen, Z. He, C. Zou, Y. Shi, L. Zhu, C. C. Chen, G. Liu, S. Yang, F.
22 Liu and Z. Yang, *Adv. Energy Mater.*, 2023, **13**, 2300968.
- 23 75. C. Li, L. Chen, F. Jiang, Z. Song, X. Wang, A. Balvanz, E. Ugur, Y. Liu, C. Liu, A.
24 Maxwell, H. Chen, Y. Liu, Z. Wang, P. Xia, Y. Li, S. Fu, N. Sun, C. R. Grice, X. Wu,
25 Z. Fink, Q. Hu, L. Zeng, E. Jung, J. Wang, S. M. Park, D. Luo, C. Chen, J. Shen, Y.
26 Han, C. A. R. Perini, J.-P. Correa-Baena, Z.-H. Lu, T. P. Russell, S. De Wolf, M. G.



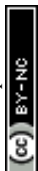
- 1 Kanatzidis, D. S. Ginger, B. Chen, Y. Yan and E. H. Sargent, *Nat. Energy*, 2024, **9**,
2 1388-1396.
- 3 76. T. Leijtens, R. Prasanna, K. A. Bush, G. E. Eperon, J. A. Raiford, A. Gold-Parker, E. J.
4 Wolf, S. A. Swifter, C. C. Boyd, H.-P. Wang, M. F. Toney, S. F. Bent and M. D.
5 McGehee, *Sustain. Energy Fuels*, 2018, **2**, 2450-2459.
- 6 77. J. Kim, S. H. Lee, C. H. Chung and K. H. Hong, *Phys. Chem. Chem. Phys.*, 2016, **18**,
7 4423-4428.
- 8 78. G. E. Eperon, S. D. Stranks, C. Menelaou, M. B. Johnston, L. M. Herz and H. J. Snaith,
9 *Energy Environ. Sci.*, 2014, **7**, 982.
- 10 79. D.-Y. Son, J.-W. Lee, Y. J. Choi, I.-H. Jang, S. Lee, P. J. Yoo, H. Shin, N. Ahn, M.
11 Choi, D. Kim and N.-G. Park, *Nat. Energy*, 2016, **1**, 16081.
- 12 80. M. Saliba, T. Matsui, J. Y. Seo, K. Domanski, J. P. Correa-Baena, M. K. Nazeeruddin,
13 S. M. Zakeeruddin, W. Tress, A. Abate, A. Hagfeldt and M. Gratzel, *Energy Environ.*
14 *Sci.*, 2016, **9**, 1989-1997.
- 15 81. J. H. Noh, S. H. Im, J. H. Heo, T. N. Mandal and S. I. Seok, *Nano Lett.*, 2013, **13**, 1764-
16 1769.
- 17 82. Z. Zhang, C. Cueto, Y. Ding, L. Yu, T. P. Russell, T. Emrick and Y. Liu, *ACS Appl.*
18 *Mater. Interfaces*, 2022, **14**, 29896-29904.
- 19 83. G. E. Eperon, T. Leijtens, K. A. Bush, R. Prasanna, T. Green, J. T. Wang, D. P.
20 McMeekin, G. Volonakis, R. L. Milot, R. May, A. Palmstrom, D. J. Slotcavage, R. A.
21 Belisle, J. B. Patel, E. S. Parrott, R. J. Sutton, W. Ma, F. Moghadam, B. Conings, A.
22 Babayigit, H. G. Boyen, S. Bent, F. Giustino, L. M. Herz, M. B. Johnston, M. D.
23 McGehee and H. J. Snaith, *Science*, 2016, **354**, 861-865.
- 24 84. R. Wang, M. Mujahid, Y. Duan, Z. K. Wang, J. Xue and Y. Yang, *Adv. Funct. Mater.*,
25 2019, **29**, 1808843.
- 26 85. D. P. McMeekin, S. Mahesh, N. K. Noel, M. T. Klug, J. Lim, J. H. Warby, J. M. Ball,
27 L. M. Herz, M. B. Johnston and H. J. Snaith, *Joule*, 2019, **3**, 387-401.



- 1 86. Z. Li, S. Wu, J. Zhang, K. C. Lee, H. Lei, F. Lin, Z. Wang, Z. Zhu and A. K. Y. Jen,
2 *Adv. Energy Mater.*, 2020, **10**, 2000361.
- 3 87. X. Li, X. Wu, B. Li, S. Zhang, Y. Liu, Z. Li, D. Zhang, X. Wang, Q. Sun, D. Gao, C.
4 Zhang, W. H. Huang, C. C. Chueh, C. L. Chen, S. Yang, S. Xiao, Z. Wang and Z. Zhu,
5 *ACS Nano*, 2023, **17**, 23478-23487.
- 6 88. K. O. Brinkmann, T. Becker, F. Zimmermann, C. Kreusel, T. Gahlmann, M. Theisen,
7 T. Haeger, S. Olthof, C. Tuckmantel, M. Gunster, T. Maschwitz, F. Gobelsmann, C.
8 Koch, D. Hertel, P. Caprioglio, F. Pena-Camargo, L. Perdigon-Toro, A. Al-Ashouri, L.
9 Merten, A. Hinderhofer, L. Gomell, S. Zhang, F. Schreiber, S. Albrecht, K. Meerholz,
10 D. Neher, M. Stolterfoht and T. Riedl, *Nature*, 2022, **604**, 280-286.
- 11 89. X. Wu, Y. Liu, F. Qi, F. Lin, H. Fu, K. Jiang, S. Wu, L. Bi, D. Wang, F. Xu, A. K. Y.
12 Jen and Z. Zhu, *J. Mater. Chem. A*, 2021, **9**, 19778-19787.
- 13 90. B. Qi and J. Wang, *J. Mater. Chem.*, 2012, **22**, 24315.
- 14 91. N. K. Elumalai and A. Uddin, *Energy Environ. Sci.*, 2016, **9**, 391-410.
- 15 92. C. Li, Z. Song, C. Chen, C. Xiao, B. Subedi, S. P. Harvey, N. Shrestha, K. K. Subedi,
16 L. Chen, D. Liu, Y. Li, Y.-W. Kim, C.-s. Jiang, M. J. Heben, D. Zhao, R. J. Ellingson,
17 N. J. Podraza, M. Al-Jassim and Y. Yan, *Nat. Energy*, 2020, **5**, 768-776.
- 18 93. S. H. Turren-Cruz, A. Hagfeldt and M. Saliba, *Science*, 2018, **362**, 449-453.
- 19 94. Z. Jia, X. Guo, X. Yin, M. Sun, J. Qiao, X. Jiang, X. Wang, Y. Wang, Z. Dong, Z. Shi,
20 C. H. Kuan, J. Hu, Q. Zhou, X. Jia, J. Chen, Z. Wei, S. Liu, H. Liang, N. Li, L. K. Lee,
21 R. Guo, S. V. Roth, P. Muller-Buschbaum, X. Hao, X. Du and Y. Hou, *Nature*, 2025,
22 **643**, 104-110.
- 23 95. Y. Han, J. Fu, Z. Ren, J. Yu, Q. Liang, Z. Xu, X. Xie, D. Li, R. Ma, M. Cao, Y. Sun, C.
24 Yang, J. He, X. Chang, K. Liu, P. W. K. Fong, J. Huang, H. Liu, Z. Liu, D. Xu, L. Cheng,
25 J. Zhang, G. Yang, X. Lu, Y. Zhu, Q. Tai, Q. Lin, H. Hu, Y. Yang and G. Li, *Nat.*
26 *Energy*, 2025, **10**, 513-525.



- 1 96. S. Jiang, R. Wang, M. Li, R. Yu, F. Wang and Z. a. Tan, *Energy Environ. Sci.*, 2024, **17**,
2 219-226.
- 3 97. Z. He, R. Yu, Y. Dong, R. Wang, Y. Zhang and Z. Tan, *Nat. Commun.*, 2025, **16**, 1773.
- 4 98. K. Lang, J. Xu, H. Han, H. Liu, Y. Fu, X. Zhang, Z. Sun, Q. Shi, Z. a. Tan and J. Yao,
5 *Adv. Funct. Mater.*, 2025, **35**, 2502966.
- 6 99. Z. Song, J. Wang, Y. Bao, J. Zeng, D. Wang, J. He, P. Zhu, B. Jiang, Z. Liu, S. He, Y.
7 Hou, Z. Hu, C. Xie, Y. Chen, Y. Liu, X. Wang and B. Xu, *Energy Environ. Sci.*, 2025,
8 **18**, 4883-4892.
- 9 100. S. Liu, L. Hao, J. Yu, Y. Xu, Y. Dou, J. Xie, Y. Wang, K. Zhang, F. Huang and Y. Cao,
10 *ACS Nano*, 2025, **19**, 748-759.
- 11 101. J. G. Son, S. Ameen, J. Roe, S. Park, J. Seo, J. Kim, A. B. Faheem, H. e. Koo, S. O. Oh,
12 Y. Jo, J. W. Kim, Y. Lee, Y. S. Shin, H. Jang, D. Lee, S. Hur, K. K. Lee, S. Cho, D. S.
13 Kim, J. Y. Kim and B. Kim, *Adv. Energy Mater.*, 2025, **15**, 2404092.
- 14 102. E. T. Hoke, D. J. Slotcavage, E. R. Dohner, A. R. Bowring, H. I. Karunadasa and M. D.
15 McGehee, *Chem. Sci.*, 2015, **6**, 613-617.
- 16 103. S. Wu, M. Liu and A. K. Y. Jen, *Joule*, 2023, **7**, 484-502.
- 17 104. X. Meng, X. Liu, Q. Zhou, Z. Liu and W. Chen, *Nano Energy*, 2024, **128**, 109984.
- 18 105. H. Hoppe and N. S. Sariciftci, *J. Mater. Res.*, 2011, **19**, 1924-1945.
- 19 106. H. Neugebauer, C. Brabec, J. C. Hummelen and N. S. Sariciftci, *Sol. Energy Mater. Sol.*
20 *Cells*, 2000, **61**, 35-42.
- 21 107. S. Gunes, H. Neugebauer and N. S. Sariciftci, *Chem. Rev.*, 2007, **107**, 1324-1338.
- 22 108. T. Liu, Q. C. Burlingame, M. R. Ivancevic, X. Liu, J. Hu, B. P. Rand and Y. L. Loo,
23 *Adv. Energy Mater.*, 2023, **13**, 2300046.
- 24 109. W. Chen, D. Li, X. Chen, H. Chen, S. Liu, H. Yang, X. Li, Y. Shen, X. Ou, Y. Yang, L.
25 Jiang, Y. Li and Y. Li, *Adv. Funct. Mater.*, 2021, **32**, 2109321.
- 26 110. Y. Wang, B. Liu, D. Zhang, H. Yu, X. Wu, D. Gao, B. Li, C. Zhang, W. Liu, Z. Yu, N.
27 Wang, L. Wang, X. Li, H. Yan and Z. Zhu, *Small*, 2025, **21**, e2411031.



- 1 111. H. Zhang, Y. Luo, T. A. Dela Peña, R. Ma, H. Yan, M. Li, M. Suryawanshi, J. Wu and
2 A. Uddin, *Adv. Mater. Interfaces*, 2025, **12**, 2500204.
- 3 112. Y. Li, Y. Yan, Y. Fu, W. Jiang, M. Liu, M. Chen, X. Huang, G. Lu, X. Lu, J. Yin, S.
4 Wu and A. K. Jen, *Angew. Chem. Int. Ed.*, 2024, **63**, e202412515.
- 5 113. J. G. Son, H.-e. Koo, W. Lee, D. Kim, S. Park, J. Roe, J. Seo, J. M. Ha, H. Lee, W. Lee,
6 H. Y. Woo, S. Cho, D. S. Kim, S.-J. Shin and J. Y. Kim, *Energy Environ. Sci.*, 2026, **19**,
7 1540-1550.
- 8 114. X. Chen, Z. Jia, Z. Chen, C. Zhou, S. Huang, X. Xia, S. Liang, P. Wang, T. Jiang, T.
9 Liu, X. Xu, B. Yan, J. Yao, X. Lu, W. Shen, H. Zhu and Y. M. Yang, *Adv. Mater.*, 2025,
10 **37**, e2500190.
- 11 115. C. Kan, C. Luo and Y. Hou, *Energy Environ. Sci.*, 2026, **19**, 1101-1123.
- 12 116. P. Wu, D. Thrithamarassery Gangadharan, M. I. Saidaminov and H. Tan, *ACS Cent.*
13 *Sci.*, 2023, **9**, 14-26.
- 14 117. B. Taheri, F. De Rossi, G. Lucarelli, L. A. Castriotta, A. Di Carlo, T. M. Brown and F.
15 Brunetti, *ACS Appl. Energy Mater.*, 2021, **4**, 4507-4518.
- 16 118. M. Zhang and Z. Lin, *Energy Environ. Sci.*, 2022, **15**, 3152-3170.
- 17 119. O. Astakhov, T. Cibaka, L. Wieprecht, U. Rau and T. Merdzhanova, *ChemSusChem*,
18 2025, **18**, e202402027.
- 19 120. S. Baumann, G. E. Eperon, A. Virtuani, Q. Jeangros, D. B. Kern, D. Barrit, J. Schall, W.
20 Nie, G. Oreski, M. Khenkin, C. Ulbrich, R. Peibst, J. S. Stein and M. Köntges, *Energy*
21 *Environ. Sci.*, 2024, **17**, 7566-7599.
- 22 121. D. Kumar, C. Zhang, E. Holubnyak and S. Demirkesen, *Int. J. Hydrogen Energy*, 2024,
23 **95**, 389-401.
- 24



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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.

