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The long road to outdoor stability: real-world challenges for controlling perovskite materials for solar cells

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Perovskite photovoltaics have demonstrated impressive performance in controlled laboratory tests, with power conversion efficiencies as high as 27%, positioning them as promising alternatives to conventional silicon-based solar cells for various applications. However, despite these major advances, operational stability remains a limiting factor for commercial applications. Large-scale implementation requires addressing challenges such as susceptibility to environmental stressors and the need for long-term outdoor stability, which few standardized outdoor testing studies and publications have accurately assessed. In this review, it is suggested that several stressors, such as temperature and irradiance fluctuations, UV-light, humidity, and precipitation have a significant relevance on the long-term stability. The main degradation reactions initiated by these stressors are reviewed for different compositions of perovskite absorbers and charge transport layers. Furthermore, we reviewed recent *in situ* and/or *in operando* studies of perovskite materials and devices under controlled conditions, which intend to elucidate reaction mechanisms and/or interactions among the device's layers in state-of-the-art perovskite compositions, which pave the way for a more rational design of stable devices and highlight the importance of developing *in situ* or *in operando* studies for each perovskite composition interacting with other materials within the cell stack. For controlling perovskite solar cells (PSCs) and improving outdoor stability, we emphasize strategies such as bulk modifications, interface engineering, and back electrode design, and discuss each specific strategy used in the literature that has been proved in outdoor conditions and its direct effect on device real-world stability. Moreover, additional strategies such as optimized interconnection layouts, protective functional layers, and advanced encapsulation materials are discussed. Additionally, in this work, an assessment of different measurement approaches aligned with international standards such as IEC 61215 and ISOS was carried out across different climate zones. Our analysis reveals a predominant focus on temperate climates in outdoor testing, along with growing interest in correlating indoor accelerated aging data with measured outdoor performance. In particular, tropical climates, with consistently high humidity, temperature, and solar radiation, provide an ideal setting for exhaustive and accelerated stability tests to evaluate strategies for improving perovskite devices. This review also emphasizes the need for expanding outdoor testing in diverse climates, particularly those enabling rapid feedback and decision-making, as a critical step towards ensuring PSC stability and commercial viability.

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1. Introduction

The exceptional properties of metal-halide perovskite (MHP) semiconductors, including high charge mobility,^{1–3} tunable bandgap,^{4–8} long diffusion lengths,^{5,9} and the ability to be processed at relatively low temperatures (<150 °C),² have enabled their use in the development of one of the most promising emerging photovoltaic solar technologies: PSCs. Although, during the initial years, the devices exhibited



relatively low stability and power conversion efficiency (PCE), the effort of the scientific community led to the fabrication of devices with current record PCEs of 26.95% for single-junction cells and 30.1% for full-perovskite tandem devices,¹⁰ matching and even surpassing in some cases the performance of commercial technologies. For context, within these, top performing silicon-based panels using n-type substrates for household applications can deliver PCEs of 24–25% under AM1.5,^{11,12} triple-junction GaInP/GaAs/Ge and quadruple-junction AlInGaP/AlInGaAs/InGaAs/Ge modules for aerospace applications display PCEs approaching 30 and 32% respectively under AM0,¹³ light-weight CdTe panels show PCEs exceeding 19%,¹⁴ and flexible CIGS modules can deliver 17% PCE under AM1.5.¹⁵

The progress of PSCs has been made possible through continuous improvements in manufacturing processes,¹⁶ compositional optimization,¹⁷ development of passivation of the perovskite absorber,^{18–20} interface engineering,^{21–24} and encapsulation strategies.^{25–28} Particularly, encapsulation of the

devices remains a key area of study for PSCs, as silicon (the market leader) has demonstrated operational stability of up to 25 years under real-world operating conditions, due to better intrinsic stability, but mainly to the use of proper encapsulation materials.^{29–32}

The performance and stability of PSCs can be evaluated under controlled laboratory conditions (Indoor) or outdoor (uncontrolled) conditions. The second one is difficult to replicate in the laboratory because of the variability and unpredictable nature of climatic conditions worldwide,^{33,34} however, both approaches are essential and have provided valuable knowledge for developing this technology. Indoor stability evaluations are intended for accelerated testing and the emulation of certain outdoor conditions,^{35,36} while outdoor stability testing demonstrates device performance under real-world operating conditions; which is crucial for deploying this technology in different countries and climates, as performance may vary significantly in tropical, desert, or temperate regions,



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for meniscus-guided coating techniques. He develops flexible p-i-n devices and addresses the key challenges associated with large-area and scalable processing.

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energy applications, with particular emphasis on renewable-energy technologies.

Esteban Velilla earned his MSc in Engineering from the University of Antioquia (UdeA) in 2007 and his PhD in Materials Engineering in 2021, where his research addressed the fabrication, characterization, and modeling of large-area perovskite solar cells (PSCs). He currently serves as a full professor at UdeA and his research focuses on energy management, artificial intelligence, emulation, simulation, and the characterization of devices for



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perovskite solar cells, and perovskite-based photoanodes for water splitting.

Juan Felipe Montoya obtained his BSc in Chemical Engineering from the University of Antioquia (UdeA) and PhD in Chemistry from the Autonomous University of Barcelona (Spain). Since 2023, he has been an Associate Professor at the Institute of Chemistry (UdeA) and conducts his research in the CIDEMAT and CATALAD research groups. His research mainly focuses on the development of materials and precursors for scalable



where factors such as device and environment temperature, humidity, solar irradiance, and spectral distribution fluctuate continuously and interact in complex ways.³⁷

As a result of the rapid increase in efficiency and the need of having stable PSCs, nowadays more effort has been placed on stability evaluations, both indoor and outdoor, which motivated the scientific community to rapidly adopt consensus methodologies for assessing the stability of emerging photovoltaic technologies, where the proposed protocol during the International Summit on Organic Photovoltaic Stability (ISOS) in 2011,^{33,38} served as a base for a broader consensus for PSCs in 2023.³⁹ These protocols encourage the comparison of results across laboratories, to promote a deeper understanding of degradation processes, and ultimately extend the device lifetime. ISOS protocols are primarily designed for testing small cells or minimodules in laboratory environments and do not replace industry testing standards. For modules, stability assessments must adhere to the International Electrotechnical Commission (IEC) standard 61215.³⁵ A significant difference is that IEC standardized testing involves more rigorous assessments of device stability compared to individual laboratory tests. In fact, in the case of PSCs, according to the Perovskite Database, more than 43 000 devices were reported and published between 2015 and 2025, most of which do not include associated stability data. Even so, more than 7000 devices have been identified with stability information referenced to ISOS protocols.^{40,41} Of these, only 5 reports use the ISOS-O protocol, 3 reports use the ISOS-LC-1 protocol, and 2 reports use the ISOS-T-3 protocol (Table S1). These ISOS tests are particularly relevant for any photovoltaic technology intended for outdoor use, as they reproduce realistic conditions and trigger failure mechanisms related to layer or contact delamination.⁴² Furthermore, these tests are included in the international qualification standards for photovoltaic technologies.⁴³ To

determine how many studies assessed stability under real-world operating conditions, we performed a bibliographic search using the Scopus database, complemented by articles already registered in the Perovskite database. This search returned 248 documents. A systematic review of the title, abstract, and keywords of each article was subsequently performed, with the aim of identifying those that explicitly reported outdoor stability tests. As a result, it was found that less than 30% of the articles (around 60 relevant publications) met this criterion. The complete procedure is described in greater detail in Note S1 and Table S2.

Meanwhile, as illustrated in Fig. 1a, evaluations under outdoor conditions are relatively scarce, but have demonstrated that PSCs are steadily moving closer to achieve operational stability within the several years range. The first report on outdoor stability evaluation of PSCs was published in 2015 for a small area mesoscopic carbon electrode (printable) PSCs, which remained stable during 1 week of outdoor exposure,⁴⁴ with few reports or remarkable improvements until 2022. However, after 2023 more reports have been published for small labcells with sizes (0.05–10 cm²), minimodules (10–200 cm²), sub-modules (200–800 cm²) and even modules (> 800 cm²)⁴⁵ with total evaluation times for several thousand hours, reaching a maximum of 4 years of outdoor evaluation, as reported by Remec *et al.*⁴⁶ This is the longest and most comprehensive stability result reported for PSCs to date. In this work, the authors monitored encapsulated p–i–n devices for four years under real-world operating conditions in Berlin, Germany. The results showed exceptional stability, with an efficiency loss of only ~2% during the first two summers and 15% over the entire evaluation time. However, changes in device behavior were observed. During the summer months, efficiency levels dropped by as much as 30% compared to winter. These decreases are attributed to changes in the solar



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nanostructures with a focus on advancing energy and optoelectronic applications through the development of photoelectrochemical (PEC) devices and perovskite solar cells (PSCs). His research specifically addresses achieving high stability and large-area scalability by using full-solution processing techniques.



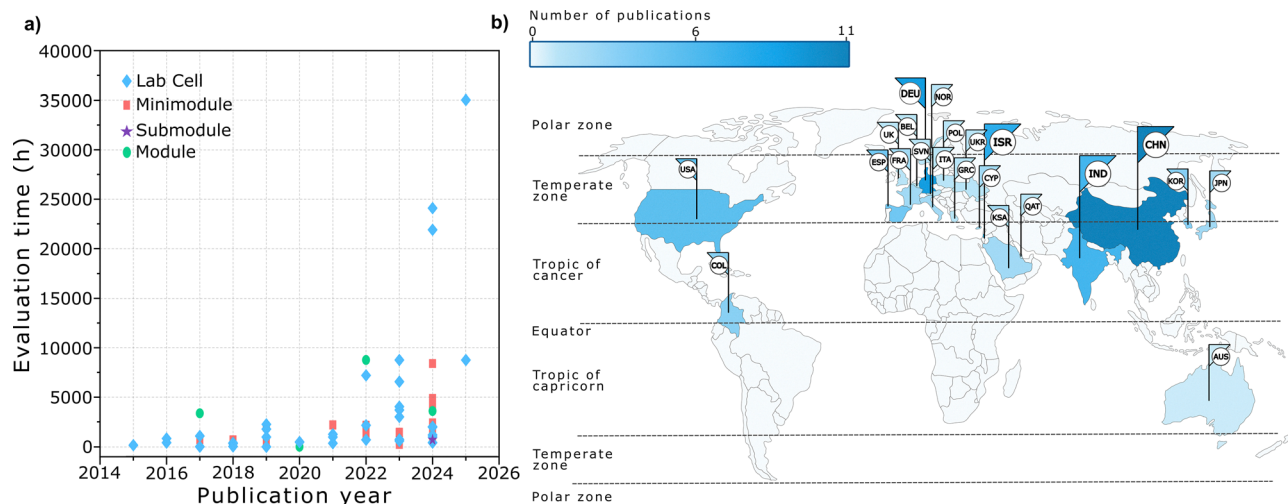


Fig. 1 Overview of outdoor stability evaluations for perovskite solar cells. (a) Evolution of the total outdoor evaluation time per device as a function of its size and the publication year. (b) Geographical distribution of reported studies on the stability of PSCs under outdoor conditions. Each country is shaded according to the number of reported evaluations. Country abbreviations: USA (United States), COL (Colombia), ESP (Spain), ITA (Italy), KSA (Saudi Arabia), IND (India), CHN (China), JPN (Japan), KOR (South Korea), AUS (Australia), NOR (Norway), POL (Poland), UKR (Ukraine), UK (United Kingdom), GRC (Greece), FRA (France), SVN (Slovenia), BEL (Belgium), ISR (Israel), QAT (Qatar), DEU (Germany), CYP (Cyprus).

spectrum, operating temperature, losses in maximum power point (MPP) tracking, as well as metastability effects. In contrast, in the winter, the devices experienced efficiency recovery cycles demonstrating that some efficiency drops were reversible. Thus, recovery and degradation cycles were evident for the winter and summer seasons, respectively, evidencing the particular behavior of perovskite materials degradation. These results confirm the potential of long-term stability of PSCs and highlight the need for extended outdoor testing in diverse climate zones to accurately assess their real-world performance and potential for commercial applications.

As real-world operating conditions are dependent on the climate conditions (the country of evaluation), it becomes essential to understand where the devices are tested as this will lead to different performances according to the geographical location. Fig. 1b shows a world map of stability evaluations of PSCs under real-world operating conditions. Few reports are available in the Equator zone and Tropics, while most evaluations have been concentrated in Europe (Germany, France, Spain, Italy, Cyprus), some countries in Asia (China, India, Saudi Arabia, Japan) and Australia. These regions cover a variety of climates, from temperate (Europe) to more extreme and hotter climates (Mediterranean in Cyprus and Israel, desert in Saudi Arabia, tropical and monsoon in India, and subtropical zones in China). Different climatic conditions offer relevant information on various degradation mechanisms, as evidenced in the comparative study conducted in Cyprus, Germany, and Israel⁴⁷ that highlights the importance of multi-climatic testing, as delamination of encapsulated devices occurred in Cyprus and Israel within weeks, while in Germany it was not detected after 2.5 years of exposure. This suggests that extreme climatic conditions, such as high temperatures and intense radiation, accelerate some degradation mechanisms, such as encapsulant-induced delamination. Therefore, to understand

the stability of PSCs, the results indicate the need to conduct evaluations under real-world operating conditions in more locations around the world, particularly in tropical environments. Collecting data under different environmental conditions can help identify specific degradation patterns enabling the development of more robust encapsulation strategies and materials, a key approach for the successful commercialization of this technology. This reinforces the need to promote research in understudied locations to better understand the stability of PSCs under the full range of possible environmental conditions.

The intrinsic instability of perovskite devices has been identified as an issue since the early stages of research in the field.^{48–57} Correspondingly, it has been tackled by numerous investigations for over 10 years now, and excellent reviews on the topic can be found in the literature.^{58–62} Most of them correspond to strategies evaluated on controlled laboratory conditions, which have significantly contributed to making perovskite materials more stable. For bringing PSCs closer to commercialization, it is required to continue research studies on long-term evaluation under real-world operating conditions as it provides data from a complex environment that is difficult to replicate under laboratory conditions. On the other hand, the interaction of each cell component, according to the nature of the materials and their interfaces, also needs to be studied. Consequently, this work provides a comprehensive overview of the stability of PSCs under real-world operating conditions. We discuss the primary factors contributing to device degradation, including exposure to variable environmental stressors such as moisture, light, and heat. Furthermore, we collect recent advancements in fabrication strategies aimed at enhancing the intrinsic and extrinsic stability of PSCs. We highlight methods such as bulk compositional engineering, interface engineering, and optimized electrode architecture that can be



implemented during device fabrication. Other strategies include the application of functional barrier layers, climate-adapted encapsulation techniques, and robust interconnect designs. Additionally, we summarized different parameters and protocols to be considered when performing outdoor evaluations. By compiling this information, this review aims to guide future research toward the development of durable and efficient PSCs suitable for widespread deployment.

2. Factors that trigger device degradation under real-world operation

The issue of PSC stability is a complex matter that depends on numerous variables that are typically categorized as intrinsic or extrinsic. The following section will examine the extrinsic degradation factors that impact the stability of PSC devices during outdoor testing. The fluctuating environmental conditions to which devices are exposed during the ageing process when subjected to outdoor conditions are considered here as extrinsic stressors. The intrinsic characteristics and thermodynamic stability of perovskite materials must be briefly discussed before examining how external stressors affect PSCs. The ABX₃ structure consists of a monovalent cation (usually cesium (Cs), methylammonium (MA), formamidinium (FA) or mixtures), a divalent metal (usually Pb or Sn), and pure or mixed halides (I, Br, Cl).⁴⁰ This family of materials has been considered “soft crystalline materials” due to their low formation energy and stability dictated by a delicate thermodynamic equilibrium.⁶³

Even under dry, solvent-free conditions, methylammonium lead iodide (MAPbI₃) and other organic–inorganic hybrid perovskites exhibit thermodynamically favorable synthesis and rapid crystallization.⁶⁴ This spontaneous formation is because MAPbI₃ has a lower Gibbs free energy than the precursor materials because it is more stable under synthesis conditions than its constituents (methylammonium iodide (MAI) and lead iodide (PbI₂)).⁶⁵ Though, this favorable formation energy should not be mistaken for long-term thermodynamic stability. After formation, MAPbI₃ is in the Gibbs free energy landscape in a shallow local minimum.⁶⁶ Despite being more thermodynamically stable than the precursors, this state is not always the global thermodynamic minimum. Under certain degradation conditions, MAPbI₃ can decompose into PbI₂, water, methyl amine (CH₃NH₂), and molecular iodine (I₂), which exhibits the lowest total Gibbs free energy.⁶⁷ The volatility of CH₃NH₂ and I₂ makes the process irreversible in the presence of open or semi-open conditions because it facilitates their removal from the system and shifts the equilibrium in favor of decomposition. Thus, the degradation processes affecting MAPbI₃, should be understood in the context of its thermodynamically favored formation and inherent metastability, as discussed above. As a result, the concept of intrinsic instability addresses fundamental instability of the perovskite itself without the influence or contributions of the external environment.

Examples of these include ion migration, vacancy diffusion, and defect assisted non-radiative recombination.^{68,69} These processes occur mostly due to physical, chemical, and thermodynamic reasons inherent to the properties of perovskites, such as soft ionic lattice, dynamically evolving structure, and low defect formation energy.^{70–72} It has been shown that these events shorten device lifespans and performance due to internal reorganization, phase segregation, and structural instability.⁷³ Environmental isolation alone cannot stop intrinsic degradation since it is governed by the material's basic thermodynamic and kinetic landscape, in contrast to extrinsic degradation.⁷⁴ Otherwise, environmental stressors lead to chemical reactions of the perovskite phase that can accelerate degradation. The conversion of iodide (I[−]) into molecular iodine (I₂) upon exposure of the perovskite to light and oxygen, which results in the collapse of the active phase structure, is an example of these processes.^{75–77}

Since PSCs frequently function at high temperatures in real-world settings, thermal instability is the main extrinsic degradation factor for most compositions, especially MAPbI₃. Outdoors, device temperatures can average between 70 °C and 75 °C during the summer in warm climates or under high irradiance. These are already considerably high values for MAPbI₃-based perovskites. This material degrades slowly at operating temperatures between 65 and 85 °C and decomposes rapidly at the temperatures normally used to encapsulate solar cells (135–150 °C).^{50,78,79} These results suggest that MAPbI₃ is not suitable for long-term outdoor stability. In such demanding applications, stabilization by device-level mitigation techniques or compositional engineering strategies may be necessary.

For instance, formamidinium lead iodide (FAPbI₃), which is more thermally stable and does not permanently disintegrate below 95 °C, is formed when methylammonium is substituted with formamidinium.^{80,81} Thermodynamically, the black α -phase of FAPbI₃ requires 150 °C to form.⁸² However, despite its good optoelectronic properties at room temperature, over time it spontaneously transforms (due to its high formation temperature) into the yellow δ -phase.⁸² This phase transition is driven solely by enthalpy and can occur without environmental stress. To preserve the photoactive phase, FAPbI₃ must be both kinetically trapped and thermodynamically stabilized. Effective strategies include A-site alloying with MA⁺ or Cs⁺,⁸³ halide mixing, and dimensional or interfacial confinement.^{84,85} Reports from specific studies indicate that FAPbI₃ can undergo reversible gaseous degradation at moderate temperatures, remaining reversible up to about 100 °C; above this temperature, some studies have observed polymerization of the organic cation, which may lead to the formation of carbon nitride species.^{86–88} Consequently, FAPbI₃ exhibits potential for practical use as a stable PSC material in the range of −40 to 75 °C, suitable for PSC operating outdoors. Other compositions such as cesium lead iodide (CsPbI₃), a wholly inorganic perovskite, have shown promising stability but less PCE. In general, one of the most promising methods for enhancing the intrinsic thermodynamic stability of perovskite absorbers is compositional engineering at either the A or X site.^{89,90} Apart from



compositional approaches, other strategies have been suggested and evaluated, such as the use of stable support materials, additive engineering, and the introduction of barrier layers at interfaces that block moisture and protect perovskite.^{91–94}

Most of these strategies have been indoor tested, which does not accurately represent the overall degradation behavior of the devices.⁴⁰ In the real environment, there are several simultaneous factors such as temperature, humidity, light, and oxygen that place considerable demands on the integrity of the PSCs. These factors may operate jointly or independently. In the following, we examine the effects of these factors on stability and strategies that can be employed to mitigate them. First, we discuss the effect on PSCs stability of each environmental stressor separately to understand each correspondent degradation pathway. Then, we discuss the synergic effect on device stability of combined environmental stressors, focusing on the combination of the most critical for PSCs stability. Note that outdoor operating PSCs are simultaneously submitted to a combination of environmental stressors depending on the quality of device encapsulation. Moreover, according to our literature survey on stability testing some environmental factors contribute more than others to the irreversible degradation of PSCs. Below we will discuss environmental factors in order of importance.

2.1. Temperature

PSCs are submitted to thermal stress in different stages of their fabrication and operation. First, an annealing step is required to form the perovskite film and other transport layers, then typical encapsulation process requires temperatures higher than 140 °C.⁹⁵ During outdoor operation, the high temperatures can activate degradation processes on the active absorber or transport layers.⁹⁶ Unlike other environmental stressors such as humidity or oxygen, thermal degradation cannot be avoided with high quality encapsulation. Thus, it is important to understand the timescale of stability of PSCs under periods of thermal stress experimented during their fabrication and operation. It is of particular importance to understand device degradation upon exposure to seasonal and diurnal temperature cycles. Recently, Abate *et al.* reviewed effective strategies for improving PSCs stability under temperature cycling, a key subject for the development of PSCs with exceptional real-world outdoor operating stability.⁹⁷ Since most of the strategies rely on modifications to the perovskite absorber layer and controlling the interaction and/or reactions among the stacked layers, we briefly discuss what are the main effects of thermal stress on these layers. One of the main concerns regarding the perovskite absorber is the phase stability which is the ability to maintain the desired photoactive structure under thermal stress without transforming into a non-photoactive phase or segregating into several phases. Given the size of Cs and FA cations they tend to form non-photoactive yellow delta phase CsPbI₃ and FAPbI₃ perovskites at room temperature.⁹⁰ Thus, quenching a metastable black perovskite phase at room temperature is the major challenge for using CsPbI₃ and FAPbI₃ as absorber layers despite they are more stable than MAPbI₃ to decomposition

reactions at outdoor operational temperatures as discussed above. Hence, for the successful implementation of compositional tuning, it is necessary to consider the phase stability of the perovskite compound according to the degree of A-site or X-site alloying. The temperature range of phase stability for several perovskite compounds have been summarized elsewhere.⁵⁹ In consequence, studies examining long-term structural stability against phase segregation or phase transition across operational temperatures must be developed to find highly stable perovskite absorbers under real world operating conditions. Another critical point to reach higher thermally stable PSCs is to understand degradation reactions triggered by interactions among the stacked layers. Some common organic hole transport materials such as PEDOT⁹⁸ or Spiro-OMeTAD⁹⁹ are known to have thermal stability problems. Thus, inorganic charge transport layers, mainly metal oxides have been chosen because of their higher thermal stability. However, although some oxides are thermally stable in isolation, they can react with the perovskite or other layers. Therefore, the effect of metal oxide layers on the thermal stability of the device must be carefully analyzed. Some oxide layers such as SnO₂¹⁰⁰ or NiO_x¹⁰¹ have been reported as highly stable and efficient charge transport layers for PSCs. Another degradation pathway of PSCs caused by thermal stress is the diffusion of mobile halide ions of the perovskite layer into the contact layers.¹⁰² This could be a major issue for long-term stability of PSCs under outdoor operation conditions that can reach temperatures as high as 75 °C. Moreover, devices operate under electric bias and light, both factors that accelerate ion migration and could affect hysteresis of PSCs.¹⁰³ On the other hand, ion migration could lead to reactions between halides and metal contacts accelerating degradation of the full device.¹⁰⁴ Therefore, it is necessary to use stable contact layers or barrier layers on top of the perovskite to suppress the reaction of halides with metal electrodes. Another strategy to avoid metal induced degradation is the substitution of metal by carbon electrodes. A big data driven PSCs stability analysis has shown higher device stability for PSCs with carbon electrodes.¹⁰⁵ In summary, there are several degradation pathways for PSCs under thermal stress which must be controlled to achieve long-term device stability in outdoor operation conditions. Since this stressor cannot be avoided through high quality encapsulation, it is necessary to increase intrinsic stability of the full device stack by implementing strategies that improve the stability of the perovskite absorber itself and its reactions with the stacked layers. These strategies will be examined in Section 3. Their effective design depends on better understanding of temperature effect on PSCs through *in situ* studies employing advance characterization techniques under thermal stress. Recently, Ruellou *et al.*¹⁰⁶ studied structural changes of the FAPbI₃ perovskite by *in situ* X-ray diffraction (XRD) under controlled temperature. They found that pure α -FAPbI₃ perovskite is stable in air up to 145 °C, confirming its high intrinsic stability. The thermal degradation of α -FAPbI₃ perovskite was monitored by the decrease in the normalized intensity of the (100) diffraction peak which started at 145 °C. This degradation correlates with



the formation of PbI_2 which started to form at 150°C as evidenced by the increase in the normalized intensity of the PbI_2 (100) diffraction peak. This result supports a direct degradation pathway of $\alpha\text{-FAPbI}_3$ to PbI_2 . In contrast to the dark conditions, the formation temperature of PbI_2 is reduced to 130°C and does not correlate with the decomposition temperature of perovskite (110°C) for the same experiment carried out under illumination. This suggests a different degradation pathway which involves the formation of an amorphous or poorly ordered phase not detectable by XRD. This finding is relevant for PSCs in real operation conditions because heat is always combined with light. In this scenario thermally activated photodecomposition pathways can emerge yielding $\alpha\text{-FAPbI}_3$ perovskite less stable than expected. Moreover, compositional tuning of the FAPbI_3 perovskite can also decrease the thermal stability. For instance, the introduction of methyl ammonium bromide (MABr) in the precursor solution can decrease the onset temperature of perovskite decomposition up to 50°C compared to pure $\alpha\text{-FAPbI}_3$.¹⁰⁶ Thus, the compositional tuning usually used for the stabilization of the photoactive $\alpha\text{-FAPbI}_3$ polymorph can substantially decrease the thermal stability of the perovskite absorber. This finding demonstrates that *in situ* characterization studies must be conducted for each specific formulation of the perovskite precursor rather than the common assumption about the high thermal stability of the $\alpha\text{-FAPbI}_3$ perovskite. Furthermore, the charge extraction layers can change the degradation dynamics of the $\alpha\text{-FAPbI}_3$ perovskite as demonstrated for titanium dioxide (TiO_2) and spiro-OMeTAD interacting with the FAPbI_3 perovskite under controlled temperature.¹⁰⁶ This highlights the necessity of evaluating in real operation conditions the stability of the whole PSCs stack, rather than focusing solely on the perovskite absorber layer.

2.2. Light

During outdoor operation, PSCs are exposed to changes in sunlight's radiation intensity and spectral distribution caused by weather variability and geographical location. Ultraviolet (UV) and blue wavelengths of sunlight are of main concern because of their impact on PSCs stability.¹⁰⁷ Unlike atmospheric stressors such as oxygen or humidity, light-induced degradation cannot be prevented by encapsulation; therefore, the semiconductor must be intrinsically stable or externally protected by optical filters against light exposure. Some reviews have discussed photoinduced phenomena in perovskite materials such as ion migration, halide segregation, and phase segregation in bromine rich ($>20\%$ Br/I ratio) mixed halide perovskites.^{108,109} These photo-induced changes are not necessarily detrimental to the performance of solar cells. Sometimes these processes in perovskite materials can be reversible or even increase solar cell PCE as observed for PSCs exposed to 1 sun illumination for more than 1500 hours.¹¹⁰ Indeed, big data analysis reports stable PSCs in timeframes higher than 10 000 hours under simulated sunlight illumination.¹⁰⁵ Our literature revision (Fig. 1a) shows recent reports of outdoor operation of up to 35 000 hours demonstrating outstanding stability of PSCs to prolonged cycles of illumination under variable weather

conditions. These highly stable PSCs have been achieved by applying strategies such as implementing stable contact layers on the stacked device, structural modification of the perovskite absorber layer, and choosing encapsulants that filter UV light,¹⁰⁹ as will be discussed in Section 3. The high stability achieved up to date implies successful hindering of irreversible photodegradation processes by applying some of the mentioned strategies. Among the irreversible photo-induced degradation processes are the oxidation of iodide ions in the lattice by photogenerated holes, increasing halide vacancy concentration and forming neutral iodine interstitials. Eventually this process could lead to the reduction of Pb^{2+} to Pb^0 which is an irreversible decomposition pathway of the perovskite absorber.^{35,111} New insights into light induced degradations mechanism have been achieved by *in situ* photoluminescence (PL) studies of PSCs under blue LED illumination.¹¹² PL imaging was recorded along with periodic *I-V* curves and transient photovoltage measurements. The PL maps revealed increased spatial non-uniformity of PL intensity as device degradation takes place. This is evidenced by the emergence of low intensity PL regions associated with localized failure sites. In this way, the diminution in macroscopic performance of the device was correlated with the observed spatial heterogeneous degradation implying that degradation takes place faster in certain areas such as defect clusters or grain boundaries. These results highlight the importance of spatial and *in situ* characterization techniques for gaining insights into light induced degradation mechanisms.¹¹² Following this approach, Frohna *et al.* developed a multimodal *operando* microscopy toolkit to spatially assess charge transport, recombination losses and compositional changes of operating PSC devices.¹¹³ Devices under one sun illumination (at variable bias) were characterized by Hyperspectral *operando* luminescence microscopy and Nanoprobe synchrotron X-ray fluorescence to measure the spatial variation of charge transport losses, chemical composition and recombination losses of PSC devices before and after extended operation. Results demonstrate that devices with the highest macroscopic performance show the lowest initial PCE spatial heterogeneity. Hence, authors demonstrate the crucial role of interface and compositional engineering to homogenize charge extraction in the device which. Although previous studies have demonstrated that hybrid lead-halide perovskites can tolerate spatial disorder in chemistry,¹¹⁴ the most recent research demonstrates that PSC devices cannot tolerate spatial heterogeneity in charge extraction originating from interfacial defects or local composition heterogeneity.¹¹³ Another recent study decouples the light induced degradation from thermal decomposition processes.¹¹⁵ This is achieved by characterizing a mixed cation $(\text{FA}_{0.73}\text{MA}_{0.27})\text{Pb}(\text{I}_{0.945}\text{Br}_{0.055})_3$ perovskite by means of *in situ* electron paramagnetic resonance (EPR) spectroscopy, a technique highly sensitive to the formation of free carriers. It was demonstrated that photogenerated charge carriers trigger a decomposition pathway that is mediated by radicals localized in the carbon of formamidinium leading to a higher degradation rate compared to that in darkness. In contrast, temperature induced decomposition leads to



methylammonium and iodide release without involving radicals as demonstrated by the combination of *in situ* EPR with *in situ* XRD, as well as thermogravimetric and calorimetric analysis. This demonstrates that illumination produces reactive species such as radicals which react with the organic cation increasing degradation rate compared to that of PSC submitted to thermal stress alone.¹¹⁵

2.3. Moisture

Metal halide perovskites suffer several degradation reactions with water which form monohydrated or di-hydrated perovskites when water molecules penetrate the perovskite structure as summarized in some review articles.^{116,117} These reactions induce reversible and irreversible degradation pathways of perovskite films. The irreversible decomposition processes led to the formation of PbI_2 which implies the failure of the PSCs device. Indeed, several studies reported degradation of unencapsulated PSCs after a few hundred hours of exposure to air with relative humidity higher than 50%.^{118,119} The intrinsic instability of perovskite materials under moisture points out the necessity of developing strategies to prevent degradation by contact with water. Some approaches such as compositional tuning of the perovskite absorber, and modification of the PSC stack by introducing surface treatments, passivation, and hydrophobic layers have been reviewed elsewhere.⁵⁹ These strategies focus on increasing the intrinsic resistance of the solar cell stack to moisture-induced degradation. In Section 3, we will review some of these strategies that have been outdoor tested. The degree of success of these strategies determines the requirements for the maximum water vapor transmission rate achieved by encapsulation. This is relevant for flexible PSCs because their encapsulants have not reached the same barrier quality as their rigid counterparts.^{95,120,121} Hence, increasing the intrinsic stability of perovskite to moisture induced degradation will allow the simplification of encapsulation processes and architectures which are expensive and complex, as it has been reviewed elsewhere.⁹⁵ On the other hand, rigid glass encapsulated PSCs have demonstrated encapsulations that prevent moisture degradation on timescales up to one year.⁸⁴ Although moisture is considered up to date detrimental for PSC performance, a recent study demonstrates new insights into the self-healing or self-passivation induced by humid air on methylammonium (MA) free compositions.¹²² Since formamidinium (FA) rich compositions have been used recently in the highest performance devices,⁸⁰ a deeper understanding of the role of oxygen and water in the reactions of FA based perovskites is required. For instance, in a $\text{FA}_{0.7}\text{Cs}_{0.3}$ mixed Br/I wide bandgap perovskite in the presence of humid air and ambient illumination, the formation of a surface layer that contains O, OH and N-based anions was demonstrated. These anions promote the formation of cyanide and/or formamidinate that bind to Pb. The role of these ligands on defect passivation was demonstrated by enhanced photoluminescence quantum yield as well as improved performance of PSCs attributed to both reduced surface recombination and increased bulk carrier lifetime.¹²² Based on these conclusions, we suggest the necessity of deeper

studies into the FA related surface chemistry for a broad set of perovskite compositions because they can lead to highly stable perovskite films as well as to elucidate the role of environmental factors, such as moisture or oxygen, in the surface reactions of perovskites. As discussed, these environmental factors do not always act as “stressors”; in fact, under certain conditions they may contribute to the stability of perovskites. In a recent study, Kelly *et al.* correlated the changes in the perovskite lattice with device performance by combining *operando* grazing-incidence wide-angle X-ray scattering (GIWAXS) with periodical measurements of *I-V* curves.¹²³ The data demonstrated the high defect tolerance of PSCs. Despite the formation of hydrated perovskite phases, they did not correlate with changes in PSCs performance. This highlights the importance of better understanding of the role of humidity in device stability. Most of the scientific literature have established that water led to perovskite degradation based on studies of MAPbI_3 aging under humid conditions. However, a detailed comparative study of triple cation and MAPbI_3 aging under humid conditions by means of *operando* GIWAXS demonstrates the central role of perovskite composition on humidity resistance. This contrasting behavior was attributed to reduced ion migration in triple cation perovskite compared to MAPbI_3 which showed higher rate of iodide migration caused by exposure to moisture.¹²³ Thus, we suggest studying the effect of humidity on PSC stability for each composition of the perovskite absorber, preferably combining periodic PSC performance measurements with *operando* material characterization techniques. A prominent example is the study by Mejaouri *et al.* which investigates the chemical and structural evolution of the hybrid perovskite film $\text{Cs}_{0.05}(\text{MA}_{0.15}\text{FA}_{0.85})_{0.95}\text{Pb}(\text{I}_{0.84}\text{Br}_{0.16})_3$ (CsMAFA) after aging under humidity-controlled conditions.¹²⁴ By analyzing perovskite films at different scales through photoluminescence, X-ray diffraction spectroscopy, cathodoluminescence, selected area diffraction, and energy dispersive X-ray spectroscopy, it was identified several degradation products and their optical and chemical properties at the microscopic level. Upon degradation induced by water different phases such as lead iodide (PbI_2), inorganic mixed halide $\text{CsPb}(\text{I}_{0.9}\text{Br}_{0.1})_3$ and lead-rich $\text{CsPb}_2(\text{I}_{0.74}\text{Br}_{0.26})_5$ were identified.¹²⁴ This demonstrates that humidity induces complex phase-segregation and crystallization processes. Elucidation of degradation mechanisms that describe the formation sequence of these byproducts is crucial for designing stable perovskite compositions or barrier layers to suppress such reaction pathways.

2.4. Oxygen

As PSCs are exposed to air under outdoor conditions, it is necessary to understand the stability of the materials comprising the solar cell stack under oxygen exposure. It is of particular importance to understand the interaction of the charge transport and active absorber layers with oxygen because they can induce degradation. Although metal halide perovskites have demonstrated to be stable to oxygen in dark and dry conditions, they easily decompose in the presence of both oxygen and light.^{75,125} Photodecomposition reactions are initiated by the



diffusion and adsorption of oxygen at iodide vacancies, occurring both at the surface and within the bulk of the perovskite crystallites.⁶⁷ Thus, it is critical to control the density of iodide vacancies in perovskite films to avoid oxygen-induced degradation.⁶⁷ On the other hand, less acidic A-site cations such as cesium or formamidinium could be more stable to photooxidation because the degradation mechanism involves acid–base reaction between the A-site cation and superoxide. Another concern for PSCs stability is the possible degradation caused by the contact between charge transport layers and oxygen. Typically, these layers are made of small organic molecules, polymers or inorganic materials, mainly oxides. Most organic materials are particularly susceptible to oxidation as has been demonstrated when they are employed in organic photovoltaics.¹²⁶ Some reviews give detailed information about the interaction of organic semiconductors and oxygen.^{126–128} Inorganic charge transport layers, mainly based on oxides such as titanium dioxide (TiO₂), tin oxide (SnO₂), and nickel oxide (NiO_x) have been used as alternatives to organic materials. However, some metal oxides interact with light and oxygen to form superoxide which can decompose materials in contact with it. For instance, TiO₂ promotes photooxidation reactions of organic molecules.¹²⁹ Some studies have demonstrated that PSCs that use TiO₂ as an electron transport layer show instability due to photocatalytic reactions. Thus, strategies such as TiO₂/perovskite interface modification or replacing TiO₂ by SnO₂ must be implemented to increase PSCs stability.^{130,131} As mentioned in the case of moisture, it is necessary to continue developing strategies for increasing the intrinsic stability of PSCs to oxygen as well as to improve encapsulants to avoid oxygen ingress to devices during outdoor testing.

2.5. Challenges for controlling degradation factors in outdoor exposed PSCs

For controlling perovskite materials and solar cells exposed to outdoor conditions, the mentioned degradation factors need to be completely understood, which is quite complicated, due to the large variability they have depending on diurnal and seasonal variations. Laboratory-controlled exposure to these stressors has been shown to induce significant degradation in devices, and some studies suggest a correlation with outdoor performance.^{132,133} However, they differ from real-world conditions which can be both harsher and more irregular, namely due to unexpected variations of the spectrum depending on the location (*i.e.* tropical zone) which modifies the operating point and thus causes varying stress conditions¹³⁴ as well as natural cycling of the devices between day and night.

Particularly, the range of values of environmental variables such as temperature, humidity, irradiance, and ultraviolet radiation are highly dependent on the geographical location where the PSCs are tested. This points out the necessity to collect a large set of data on outdoor testing of PSCs in a wide range of geographical locations. As analyzed in section one, most of the research articles report outdoor testing of PSCs in countries located in the temperate zone (35°–66.5° latitude). These studies offer a set of data of PCE, and lifetime of PSCs

evaluated in a particular climate zone characterized by four distinct seasons (spring, summer, autumn, and winter), moderate temperature and irradiation variations during diurnal cycles, and particular types of precipitation.

However, a comprehensive understanding of outdoor performance and stability of PSCs requires collecting data of PSCs outdoor operating under well differentiated climatic zones which are classified based on geographical latitude. As schematically illustrated in Fig. 2 the main environmental factors that trigger PSCs degradation are the temperature variations, the solar irradiance, the humidity levels, the UV index, and the precipitations. All these factors are highly dependent on geographical latitude. Three main climatic zones classified according to the geographical latitude are denoted as tropical (0°–35° latitude), temperate (35°–66.5° latitude), and polar (66.5°–90° latitude). For each geographical zone the minimum, maximum, and range of variation of some climatic parameters differ significantly. For instance, the tropical zone is characterized by high temperatures throughout the year (25–37.75 °C) with little seasonal variations while in the temperate zone the temperature presents higher seasonal variations and lower range values (0–25 °C). These temperature differences between geographical zones significantly affect the commonly used lifetime indicators of PSCs such as T_{80} or T_{50} . A recent report of PSCs outdoor testing in the temperate zone found that the day/light time maximum temperatures have a more significant effect on the long-term degradation than the minimum temperatures during the dark/night cycles.³⁷ Therefore, a higher long-term degradation could be expected for PSCs outdoor tested in the tropical zone compared to devices evaluated in temperate or polar zones where the maximum diurnal temperatures are lower. Moreover, the annual average range of values for environmental stressors such as irradiance, moisture, and UV index are higher in the tropical zone as shown in Fig. 2. These stressors affect PSCs stability as already discussed above; therefore, higher degradation rates and more demanding conditions on encapsulation materials could be expected for outdoor evaluation in tropical zones. In this literature revision we only found 21 research articles reporting the lifetime of outdoor PSCs tested in 7 countries located in the tropical zone, with only 3 reports near to the equator line and none in the polar zone (Table S2). Since the lifetime indicators of PSCs (T_{80} , T_{50} , *etc.*) are climate-dependent, there is a need in the field of PSCs to collect more data of outdoor performance and stability of devices evaluated in countries located out of the temperate zone. This could allow the scientific community to develop models and testing protocols for predicting the long-term performance of PSCs operating in any climatic zone, while evaluating the strategies for device stability under these realistic conditions, as discussed in Section 3, is of paramount importance for further insertion of perovskite photovoltaics in the market.

In the next two Sections (2.6 and 2.7), the variables with the strongest influence on PCE reduction during outdoor and indoor accelerated tests will be discussed, focusing on their coupled or synergistic effects. In particular, the variables that



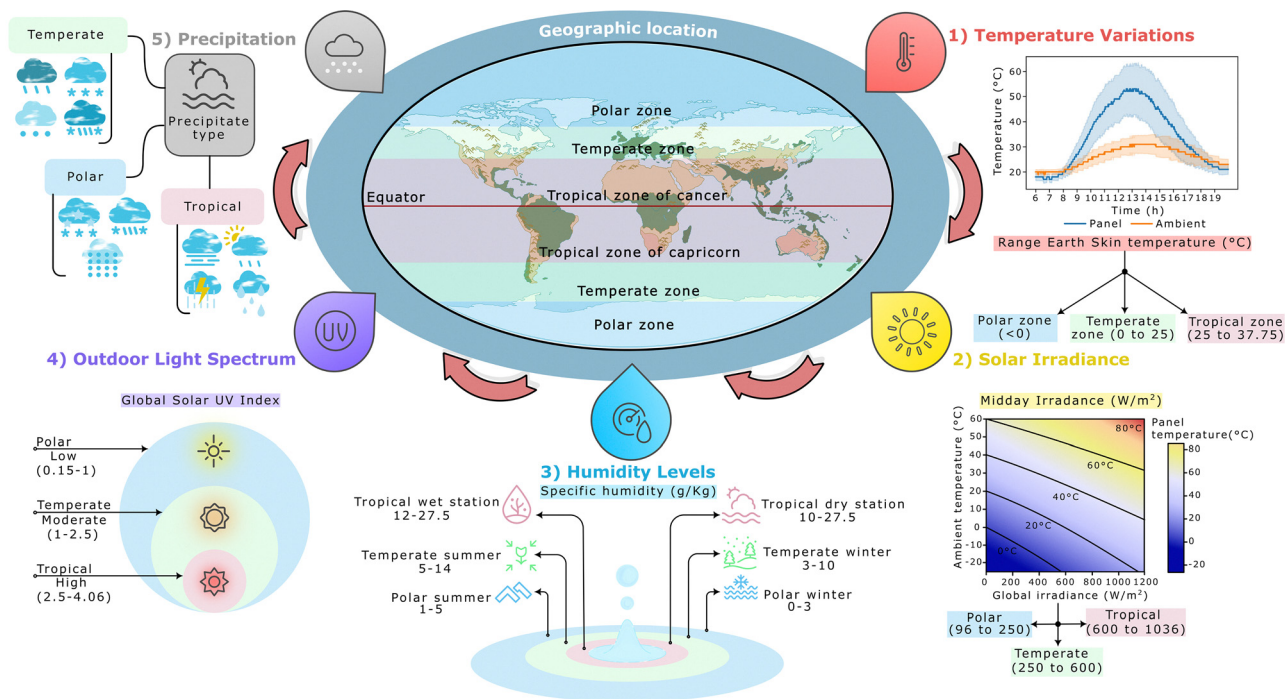


Fig. 2 Environmental stressors affecting perovskite solar cell stability during outdoor operation. Center: world map highlighting three main climate zones classified according to geographical latitude. Tropical (0° – 35° latitude), temperate (35° – 66.5° latitude), and polar (66.5° – 90° latitude) zones are colored in purple, light green, and light blue, respectively. Around the world map, the main environmental parameters that trigger PSCs degradation are depicted. The range of variations of the average annual value (year 2023¹⁴⁵) of each parameter depending on the geographical location is shown for each climate zone. Environmental parameters are shown from greatest to least importance in clockwise direction. (1) Panel temperature variation in tropical zone (Colombia, data by the authors), below this graph, the ranges of earth skin temperature (annual average for year 2023¹⁴⁵) are shown in each main climate zone. (2) Simulation of the steady-state panel temperature depending on solar irradiance and ambient temperature, below this graph are shown the ranges of solar irradiance for each climate zone (annual average for year 2023¹⁴⁵), (adapted from ref. 146, CC BY 4.0, 2018). (3) Range of specific humidity for each climate zone (annual average for year 2023¹⁴⁵) showing also the seasonal variations within each one (from left to right). (4) Ranges of global solar ultraviolet (UV) index according to the climate zone (annual average for year 2023¹⁴⁵). (5) Scheme of precipitate types for each climate zone. The icons represent snow, sleet, and hail for the polar zone; rain, snow, sleet, and hail for the temperate zone; and rain, drizzle, downpour, and hail for the tropical zone.

can be mitigated through mechanical or physical barrier materials, such as moisture, oxygen, and precipitations, will not be considered in the context of this synergistic degradation. However, variables that cannot be eliminated, such as temperature, light, and bias, will be studied in pairs; temperature with light/UV, and temperature with electrical bias. These combinations have shown the greatest influence on the degradation of encapsulated PSCs.

2.6. Temperature and light/UV: coupled factors effects

To investigate the impact of external factors on the stability of perovskite devices, each component must be evaluated separately. However, light and temperature are difficult to isolate because they are intrinsically related, as each absorbed photon generates either charge carriers or heat, leading to photothermal heating and subsequent degradation. Kamppinen *et al.* address this challenge with a fully coupled opto-electro-thermal model that quantifies parasitic absorption, recombination and Joule heating separately before feeding these heat sources into a steady-state heat equation to predict device temperatures under defined illumination and ambient conditions.¹³⁵ Under one sun (1000 W m^{-2}), 20°C ambient temperature and

about 1 m s^{-1} wind, large-bandgap (2.2 eV) devices stabilize at around 30°C , whereas small-gap (1.2 eV) devices reach approximately 44°C . These values correspond to the equilibrium temperatures reached under the stated ambient and wind conditions. This temperature rise deviates power production predictions by 1–6%, highlighting that thermalization losses, while accurately quantified, cannot be eliminated merely by optical design. Their specific parameters guide absorber and device engineering strategies to minimize photothermal heating and improve MPPT power predictions across perovskite compositions. Complementing this work, Cui *et al.*¹³⁶ categorize photothermal conversion into plasmonic localized heating, nonradiative semiconductor relaxation, and molecular vibrational heating, offering mathematical models and experimental protocols such as transfer-matrix, Beer–Lambert analyses, and temperature-decay calorimetry to isolate the thermalization term from pure optical effects. Integrating these methods with Kamppinen *et al.*'s model enables empirical validation of simulated heating pathways, closing the loop between theory and experiment.

Since UV exposure can be largely suppressed and illumination primarily induces photothermal heating, temperature



emerges as the dominant stressor under outdoor MPPT conditions for most of the encapsulated device architectures and climates studied. In both n-i-p and inverted p-i-n architectures, the front-end layers inherently block the majority of UV radiation. For example, in n-i-p configurations, ITO and compact TiO₂ layers, commonly used as transparent conductive electrode and ETL respectively, effectively filter out high-energy UV photons, enhancing device stability under illumination.^{137–139} Likewise, in inverted p-i-n PSCs, PEDOT:PSS layers exhibit notable optical absorption in the UV range (approximately 300–400 nm), while remaining transparent in the visible spectrum, which enables them to serve as effective UV attenuation front layers.^{140–142} In devices lacking these layers or using different encapsulations, UV degradation may be significantly more pronounced. However, in most outdoor-tested PSC devices, UV-induced degradation plays a minor role compared to thermally driven failure modes.

Building on these findings, Islam *et al.* conducted complementary stability tests on fully encapsulated semi-transparent MAPbI₃ cells: dark annealing at 85 °C for 1000 h to isolate pure thermal aging and continuous one-sun MPPT at a fixed ~30 °C for nearly 4000 h to capture light-induced photothermal heating under controlled temperature.¹⁴³ By comparing performance loss, they conclusively show that under their experimental setup thermal stress alone drives most of the stability decay. This conclusion is further supported by later research by Lee *et al.*¹⁰⁷ and Muhammad *et al.*,³³ which discovered that light/UV exposure results in mostly reversible photocatalytic damage with minimal irreversible loss. Taken together, these results indicate that temperature exposure (especially when amplified by light-induced photothermal heating)¹⁴⁴ is a major driver of long-term performance decline under the studied outdoor-like conditions, though other stressors may dominate in different environments without a proper encapsulation.

2.7. Temperature and electrical bias: coupled factors effects

In real operating conditions, PSCs are continuously exposed to high temperatures due to ambient heat and self-heating induced by sunlight (as discussed in Section 2.6), as well as electrical bias. Typically, this is forward bias during power generation, but occasionally it is a reverse bias caused by shading or mismatch conditions.¹⁴⁷ The combined effect of these two stress factors, temperature and electrical bias, accelerates degradation to a greater extent than either factor alone.¹⁴⁸

In accelerated indoor testing, researchers have observed that degradation occurs under combined thermal and electrical stress, even when PSCs are encapsulated to exclude oxygen and moisture.¹⁴⁹ Some studies have reported that PSCs exposed to high temperature and bias have a rapid initial decrease in power conversion efficiency (PCE), attributed mainly to the formation of additional recombination centers, interface charge accumulation and ion migration (*e.g.* halide or cation redistribution) within the electric field during the initial hours of operation.¹⁵⁰ However, this performance loss is often

partially reversible when the device is left in the dark for a comparable period.^{149,151} In contrast, slower degradation over longer periods is driven by irreversible processes such as the decomposition of the active layer and the chemical degradation of the films and interfaces.

Erdil *et al.*¹⁵² simulated field-like stress using bias/rest cycling. They demonstrated that, over the course of months, cumulative ion migration and interfacial recombination cause a persistent VOC loss, despite partial recovery occurring overnight. The authors emphasized a seasonal analogy; winter conditions are like rest phases (lower temperature and irradiance), while summer conditions are similar to bias phases (higher temperature and irradiance). Remec *et al.* provided additional validation of this phenomenon,⁴⁶ using extended outdoor datasets to confirm the trends previously reported by Erdil *et al.* over shorter timescales. According to their results, high irradiance and prolonged MPPT bias at elevated temperatures could contribute to substantial degradation in the summertime driven by increased ionic redistribution and defect formation.

Rapid ionic migration and interfacial degradation are hallmarks of bias-induced degradation (PID) in perovskite solar cells, especially when exposed to high bias and high temperatures. Studies such as that of Nakka *et al.*¹⁵³ reported a 59% PCE loss after 55 h at –1000 V and 25 °C, escalating to over 90% PCE loss under the same bias at 60 °C and 60% RH. Although real-world PSC voltage is much lower, continuous forward bias at MPP (about 1 V) combined with elevated temperatures (~60–75 °C from photothermal self-heating) could trigger analogous, but slower degradation pathways.^{46,154,155}

In particular, the electric field facilitates the migration of ions, notably I[–], MA⁺ and FA⁺, which accumulate at interfaces. This distorts the local band alignment and promotes non-radiative recombination driven by ionic redistribution, lattice distortion and field-screening effects.^{156–158} Electrical bias and elevated temperatures simultaneously accelerate the irreversible decomposition of the perovskite absorber^{154,159} particularly the volatilization of organic cations and trigger chemical reactions at interfaces with charge transport layers. Ruellou *et al.*¹¹⁵ confirmed this by observing radical generation and Spiro-OMeTAD de-doping under heat and illumination *via in situ* EPR and XRD.

Interestingly, degradation driven by electrical bias in PSCs doesn't appear to progress in a gradual or linear manner. Instead, it tends to exhibit a threshold-like response.¹⁶⁰ For this reason, the researchers suggest maintaining MPP bias conditions for the PSCs during outdoor evaluation.³⁹ Devices disconnected from an electrical circuit and exposed to illumination are biased in V_{oc} conditions, which has been demonstrated to cause faster degradation than MPP.¹⁶¹ MPPT operation better simulates field conditions and leads to more representative aging behavior. Partial shading events introduce degradation mechanisms like ion accumulation, hot spot formation, and electrochemical damage.^{147,161,162} A shaded cell in a PSC module, especially one with series-connected cells, may be reverse-biased by its illuminated neighbors, causing a



voltage stress that is significantly higher than the normal forward-operation levels (approximately 1 V).¹⁶³ Even modest reverse voltages can induce hot-spot formation, permanent shunting, electrode delamination and ion migration, particularly at elevated temperatures.^{164,165} These effects are not observed during typical MPP operation⁵⁸ and are described as a phenomenon known as reverse-bias pinning;¹⁶⁶ whereby shaded perovskite cells remain stuck in reverse even after the shading event ends. This causes irreversible drops in current output and efficiency.¹⁶⁶ Module designs should include bypass diodes, ideally one for every two or fewer cells, to lessen stress-induced degradation. This ensures that shaded cells are not exposed to detrimental stress under reverse bias.¹⁶⁷

To compare the effects of the environmental stressors discussed in this section with varying intensities across different climatic zones, a temporal evaluation of climatic variables and their impact on device performance was conducted at different time scales. The first case study was conducted in a temperate climate at the HZB (Helmholtz-Zentrum Berlin, Germany), where a one-year outdoor evaluation was carried out by Jinzhao Li *et al.*¹⁶⁸ This included both the photovoltaic performance and stability of PSCs as illustrated in Fig. 3a. In contrast, evaluations in a tropical climate, (University of Antioquia, Medellin, Colombia) were also conducted (see Fig. S1). Here, climatic variables were monitored over 18 months, with a focused analysis of their effect on device performance over a 5-month period (Fig. 3b).

Fig. 3a(1) shows that the devices depicted in the inset remained stable for most of the testing period (from October to June). However, starting in June, both the performance ratio and the power output at an irradiance of 500 W m^{-2} began to decline, reaching the T_{80} threshold at nearly one year of operation. By correlating the degradation rate with the climatic variables presented in Fig. 3a(2), it becomes evident that during the months of lower degradation rates, both irradiance and device temperature were significantly lower compared to the summer months highlighted in red (irradiance $> 1000 \text{ W m}^{-2}$ and temperature $> 30 \text{ }^\circ\text{C}$). This clearly demonstrates that increased temperature and irradiance accelerate the degradation processes in perovskite solar cells, as discussed earlier.

When comparing this behavior to the tropical zone, Fig. 3b(1) shows a much faster degradation. From the very beginning of the evaluation, power output at both 500 W m^{-2} and 800 W m^{-2} declined at a faster rate than in the temperate climate, reaching the T_{80} point in less than 5 months. Correlating this with the climatic variables in Fig. 3b(2), we observe that throughout the 18-month evaluation, both irradiance and device temperature consistently matched or exceeded the levels seen during summer months in the temperate zone. Device temperatures frequently surpassed $60 \text{ }^\circ\text{C}$, with irradiance often exceeding 1200 W m^{-2} . As shown in Fig. 2, UV exposure is higher for tropical than for temperate zones (depending on device architecture and encapsulation). Irradiance levels in temperate climates are nearly half those in tropical regions.

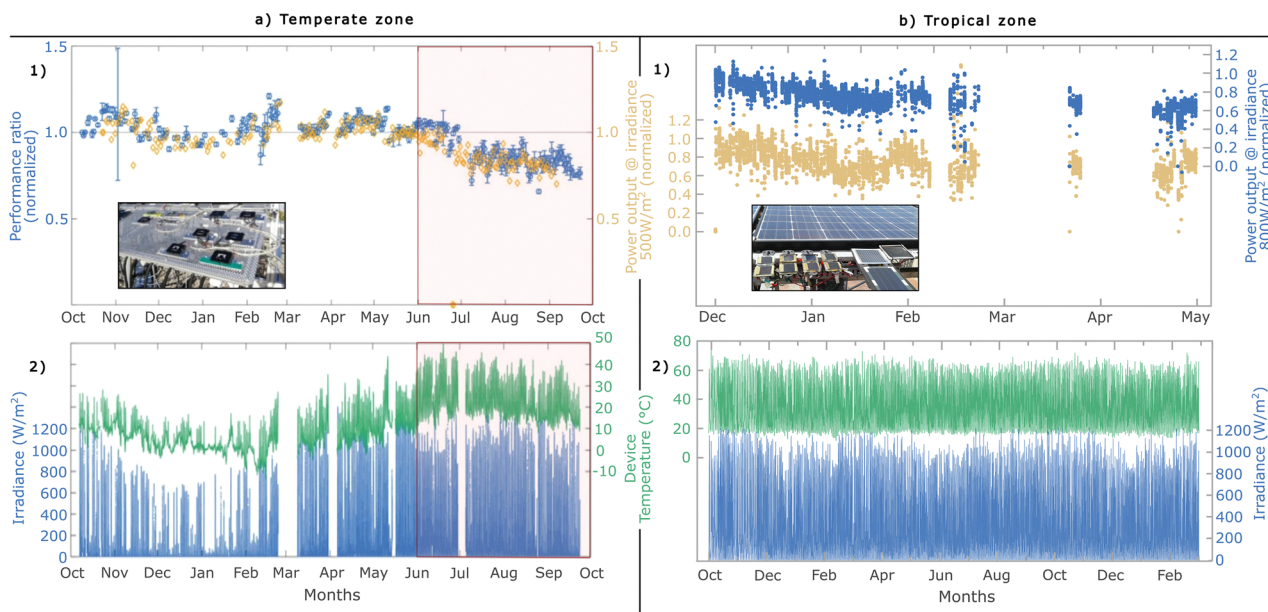


Fig. 3 Comparative analysis of perovskite solar cell performance and environmental conditions in temperate (Berlin, Germany) and tropical zones (Medellin, Colombia in the set-up showed in the Fig. S2). (a)(1) Evolution of the performance ratio (blue points) and power output at 500 W m^{-2} irradiance (gold points) over one year in the temperate zone, where a decline in performance is observed during the summer months (highlighted in red). The inset shows the tested devices under outdoor conditions. (a)(2) Corresponding irradiance (blue line) and device temperature (green line) trends over the same period, illustrating seasonal variations that influence device performance. (b)(1) Power output at irradiances of 500 W m^{-2} (gold points) and 800 W m^{-2} (blue points) over five months in the tropical zone, showing the stability of device performance under continuous high temperature and irradiance. The inset depicts the photovoltaic installation under evaluation. (b)(2) Continuous monitoring of irradiance (blue line) and device temperature (green line) in the tropical zone over eighteen months, highlighting consistently high temperatures and stable irradiance throughout the exposure time. Figures (a)(1) and (a)(2) were adapted from ref. 168, CC BY 4.0, 2023. Figures (b)(1) and (b)(2) are data from the authors.



This increases the risk of UV-induced degradation. These high UV levels affect not only UV-sensitive layers, such as the inorganic oxides¹⁶⁹ and perovskite materials (where it can accelerate ion migration¹⁷⁰), but also the encapsulation materials used in outdoor testing.¹⁷¹ Therefore, robust encapsulation materials are required to improve stability and standardize stable device structures for those climate zones.

Finally, we want to emphasize that tropical climates near the equator offer stable and consistent environmental conditions throughout the year, as shown in Fig. 3. Fig. S1a provides a more detailed explanation of this behavior, attributing the stability of the device temperature (observed in Fig. 3b(2)) to the uniformity of both irradiance and ambient temperature. Furthermore, Fig. S1b demonstrates that the peak sun hours (PSH) remained statistically stable over the course of almost a year, indicating a consistent solar exposure pattern. The absence of distinct seasons in tropical climates creates stringent real-world testing conditions with consistent day/night cycles. As a result, tropical regions serve as reference platforms for accelerated stability testing, material benchmarking and the development of robust photovoltaic technologies.

3. Fabrication strategies for having stable PSCs operating under real-world conditions

The inherent instabilities in perovskite devices arising from susceptibilities of the perovskite material itself, the stacking of materials of different nature to fabricate devices, extrinsic factors and eventually the introduction of additional elements in the devices to scale them up⁵⁸ are of critical importance when operating devices under real-world conditions and adopting appropriate strategies to deal with them is key for successful operation. Encapsulation and other approaches involving some sort of control of the device once it has been fabricated^{28,172} are widely adopted in the field and analyzed in a later section of this review. Despite this, tackling the instability issue from the fabrication of the devices themselves is naturally of great importance and correspondingly many efforts have been made towards this in the literature, resulting in a wide gamut of strategies to enhance stability, namely the stabilization of photoactive phases *via* composition engineering,^{80,81,83,173} using 2D/3D perovskite heterojunctions to reduce interface-related instabilities^{84,174–176} and modulation of perovskite/charge-transport-layer interface employing diverse additives.^{20,177–179} In this section, an overview of the main strategies for enhancing device stability tested under real-world conditions is presented, and the problems discussed in Section 2 are directly addressed here through the actual solutions implemented. The strategies are focused on the device fabrication and presented in 3 groups depending on the aspect tackled by them: bulk modifications, interface modulation and back electrode design, schematically illustrated in Fig. 4a.

Bulk modifications collect those strategies targeting a change in the bulk of the materials used in devices (perovskite

absorber, HTL, ETL), namely by introducing new materials with enhanced performance or by modifying existing ones using additives. Interface modulation gathers strategies aiming to improve the interface between the perovskite and the charge transporting materials and/or between these materials and the corresponding electrodes. Back electrode design groups the strategies dealing with the replacement of the common metal back electrodes employed in devices. Carbon electrodes shine in this aspect, as its success in enhancing stability under simulated conditions is well known in the literature.¹⁸³ The papers mentioned here are only those corresponding to works in which the focus was the strategy in question, since other approaches are discussed in other sections. Finally, a comparison between the outdoor stability results of the typical device architectures (n-i-p, p-i-n and mesoscopic carbon) is done to benchmark the overall stability of the architectures. It is important to note that the discussion on the mesoscopic cells employing carbon is treated separately from the back-electrode design, because the approach itself goes beyond simply replacing the electrode involving a significantly different approach for the fabrication of the device itself.

3.1. Tuning device fabrication to enhance stability

3.1.1. Bulk modifications. One of the most widely adopted bulk modifications is tuning the perovskite absorber layer composition, which has proven to be a powerful approach to tackle both performance and stability of devices since it has a direct effect on properties such as bandgap and stability.⁸¹ Performance is directly influenced since the composition sets the bandgap, which defines the maximum attainable PCE according to the Shockley–Queisser limit,¹⁸⁴ and impacts charge-carrier lifetime, *e.g.*, FAPbI₃ has a higher charge-carrier lifetime compared to MAPbI₃ thanks to the faster reorientation motion of the FA⁺ upon photogeneration of carriers.^{185–187} Furthermore, solvent compatibility depends on the composition, namely Cs-rich phases have poor solubility in organic solvents¹⁸⁸ and require high-coordination solvents like DMSO,¹⁸⁹ which heavily influences the quality of the absorbing films. High-coordination solvents usually have higher boiling points, hence making the crystallization step upon annealing slower. This could yield variable surface roughness which must be accounted for since it will directly impact the interface with the next layer and thus charge transport between them and charge recombination.¹⁹⁰ Stability is inherently affected because the chemical bonding in the material depends on the perovskite absorber composition. Inclusion of FA usually leads to enhanced thermal and light stability owing to the strength of the C=N double bond in FA⁺ due to the contribution of the conjugated π bond and protonated FA⁺ that hampers deprotonation.¹⁹¹ Additionally, FA⁺ displays low polarity and orientational mobility, which can increase the activation energy of ion migration for enhanced stability.¹⁹²

In this context, mixed- and triple-cation perovskite compositions have been tested extensively, aiming to synergize the benefits from different cations in the material and displaying promising outdoor stability from the early days.^{83,193–195}



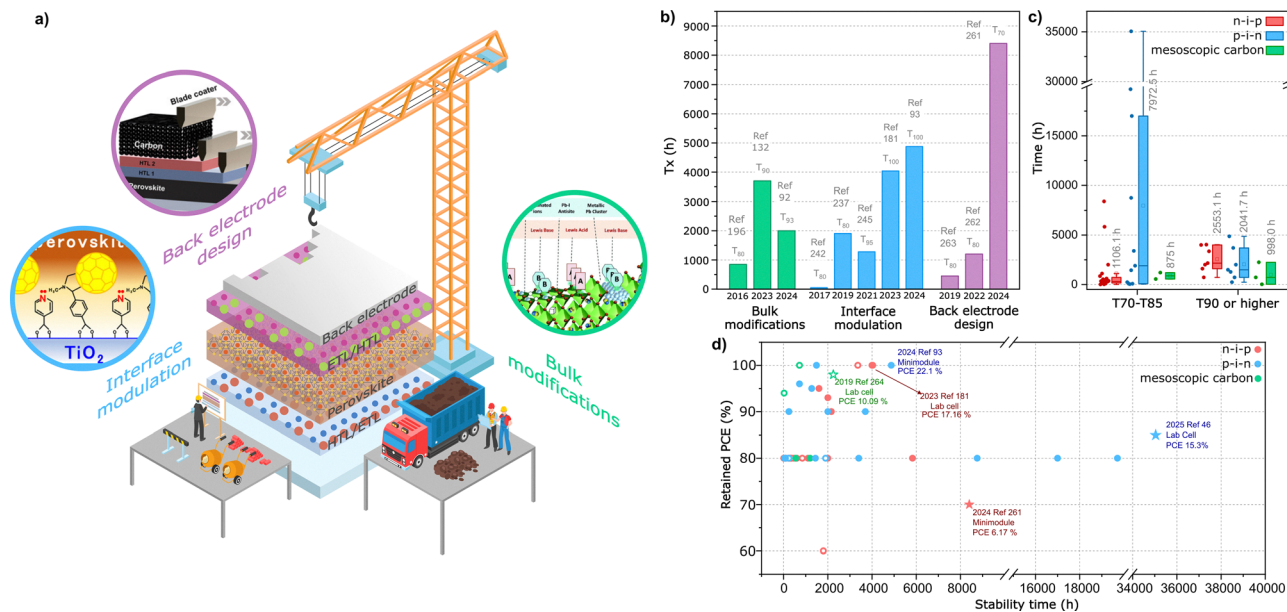


Fig. 4 (a) Schematic illustration of the three main device fabrication strategies aimed at improving the real-world operational stability of perovskite solar cells. Inset images represent: bulk modifications (adapted from ref. 180, CC BY 4.0, 2024); interface modulation (adapted with permission from ref. 181, Copyright © 2023, American Chemical Society); and back electrode design (adapted with permission from ref. 182, Copyright © 2023, Elsevier). (b) Representative outdoor stability results grouped by year and fabrication strategy used to enhance device stability. The y -axis data (T_{80} , T_{90} , T_{95} and T_{100}) differ depending on the stability result available from the reference in question. (c) Box plot showing a comparison between outdoor stability results between n-i-p, p-i-n and mesoscopic carbon device architectures grouped as T_{70} – T_{85} and T_{90} or higher. The inset text represents the corresponding average values of stability time by architecture (Axis break was applied between 20 000 and 30 000 for better readability). (d) Percentage of retained PCE as a function of test time based on reviewed papers displaying explicit outdoor evaluation. The hollow points correspond to results which have entries in the Perovskite Database.⁴¹ The star points mark the longest outdoor stability time for each architecture, and the corresponding efficiency (PCE) for these cases is also included (Axis breaks were applied between 10 000–15 000 and 20 000–34 000 for better readability).

Compositions with mixed A-cations, offer a straight-forward way to achieve this synergy and tune the Goldschmidt tolerance factor, an empirical index used to predict the preferentially formed structure which is important to promote the formation of photoactive α -perovskite phases.⁹⁰ In this sense, combining FA with a smaller cation, namely MA or Cs, allowed the benefits of FA while bringing down the tolerance factor into a suitable range to promote the formation of α -phase. Indeed, already in 2015 a mixed composition of $(\text{FAPbI}_3)_{0.85}(\text{MAPbI}_3)_{0.15}$ enabled an optimized PCE going from 13.5 to 17.3% in a FTO/TiO₂/perovskite/Spiro-OMeTAD/Au configuration.¹⁹³ The same perovskite composition enabled cells showing a T_{80} of 846 h upon exposure in Spain following the ISOS-O-2 protocol.¹⁹⁶ This approach is highly compatible with scalable deposition methods like blade coating and slot-die coating, since perovskite films fabrication relies on solution-based precursors, which can be adapted to large-area deposition systems.¹⁹⁷ However, scalability challenges include ensuring uniformity of cation distribution over large areas, as variations in precursor stoichiometry can lead to phase segregation or inconsistent film quality.¹⁹⁸ Precise control of precursor mixing and environmental deposition parameters, such as temperature and humidity, is critical to maintain reproducibility in high-throughput settings. It is important to note that, given the pioneering nature of the work, although the A-site alloying directly addressed the intrinsic thermodynamic instability of

FAPbI₃, intrinsic temperature instability during operation remained an issue. In several of the devices, following evaluation, the presence of iodine (I) in the Spiro-OMeTAD layer was noted, clearly indicating ion migration. This is expected as no direct attempt of “ion fixation” was pursued, which could be aggravating for stability given that the authors identified I[−] ions as the most likely mobile ones. This highlighted already in the early years the need to devise clever strategies to tackle intrinsic instabilities.

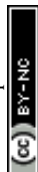
Accordingly, in the following years this A-site cation tuning has been complemented with several other strategies. Finely tuning the solvent system employed in the perovskite precursors along with the deposition technique has proven to be a suitable approach to modify the bulk of the layer and improve performance in outdoor conditions. Both aspects go hand in hand since they determine the crystallization kinetics of the perovskite film. The solvents employed in perovskite precursor solutions often have either a low boiling point and high saturated vapor pressures, *e.g.*, 2-methoxyethanol (2-ME), or a high boiling point and low saturated vapor pressures, *e.g.* *N,N*-dimethylformamide (DMF) and *N*-methyl-2-pyrrolidone (NMP). The former displays a weaker lead Pb²⁺ coordination, in agreement with its low donor number and low polarity,^{189,199} and hence can promote rapid nucleation as solute precipitation rapidly becomes more energetically favored upon solvent removal. When uncontrolled, this could restrict grain growth



and crystallinity,⁹² which is deleterious owing to the appearance of more grain boundaries that are known to have negative impacts on ion transport and perovskite decomposition over time, as has been reviewed in several works available in the literature.^{200–202} The latter, typically polar aprotic in nature,²⁰³ instead shows a high Pb²⁺ coordination ability, *e.g.*, DMF coordinates Pb²⁺ *via* the Pb–O bond,^{204,205} thus exhibiting slow supersaturation due their low evaporation rates. This eventually limits the rate of nucleation, which can be exploited to compensate for the potentially harmful fast nucleation when using the previous solvents, but could compromise the control of the film uniformity, coverage and thickness over large areas.²⁰³ When processing *via* blade coating, the drawbacks of high boiling point-solvents have been overcome by meticulous meniscus control during coating. When aiming for high throughput in blade coating, the deposition falls in the Landau–Levich regime, where viscous forces and Marangoni flow of the solute influence the amount of solution transferred from the front meniscus to the substrate. This makes the uniformity of the coating highly susceptible to uncontrolled airflow, *e.g.*, from the environment at the vicinity of the coating tool. This could stochastically alter the front meniscus at different positions, yielding inhomogeneous film thickness which has a direct effect in performance and stability. In view of this, and by using an NMP/DMF precursor whose high affinity towards lead allows slow nucleation leading to excellent crystallinity and grain size, Yoo *et al.* employed a high-velocity nitrogen stream directed towards the front of the meniscus during coating to induce an inwards contraction of it. This lowered the amount of precursor solution advancing towards the substrate homogeneously throughout the deposition front and accelerated drying of the wet film, enabling film thinning owing to the reduced duration of wet films. Consequently, a uniform and smooth film was achieved despite the high boiling point of the solvent system.⁹² In this way, encapsulated n–i–p min-modules with a FTO/SnO₂/KCl/FAPbI₃/PTABr/Spiro-OMeTAD/Au configuration showed an approximate T_{93} of 2000 h under outdoor operation conditions in China. As this approach focused on increasing the crystal quality of the perovskite film, the most important factor contributing to stable outdoor behavior was the reduction in grain boundaries. This directly tackles the resilience of the device against the combined effect of temperature and bias. Namely grain boundaries in perovskites are commonly rich in point defects, namely Γ^- ions,²⁰⁶ which are highly mobile upon biasing or heating. Given the downward band bending in the device during operation, such accumulation is aggravated by further migration of ions in the vicinity of the boundary towards it, which can result in decomposition into PbI₂ in presence of humidity^{118,207} or formation of non-photoactive δ -FAPbI₃.²⁰⁸ Although not the main focus of the work, the surface treatment of the perovskite with PTABr also plays a role in stability, as it tackles the negative effect of possible humidity leaking into the encapsulated devices. PTA⁺ can be readily adsorbed *via* hydrogen or ionic bonds between its ammonium groups and the perovskite cages, while its hydrophobic phenyl groups prevents water-

induced reactions at the vulnerable interface.²⁰⁹ Alternatively, the drawbacks of low boiling point solvents can be bridged by including small amounts of high boiling point lead-coordinating ones and smart ink design. Dagar *et al.*¹⁶⁸ employed a combination of 2-ME with small amounts of NMP that compensates for the otherwise restricted grain growth promoted by the sole use of 2-ME by forming an intermediate adduct with the perovskite precursor materials (FAI-PbI₂-NMP) that aids crystallization prior to the formation of the final FAPbI₃ α -phase.²¹⁰ Effectively, such adduct slows the overall crystallization, which in turn enhances grain growth and crystallinity. Upon deposition *via* slot-die coating, despite good crystal quality, ribbing effects were present, for which further addition of acetonitrile (ACN) as a low-lead-coordinating co-solvent²¹¹ was used. This caused the ink viscosity to decrease and led to an earlier onset of the formation of intermediate solvate phase owing to the low boiling point of ACN, which in turn removed the ribbing issues. This increased homogeneity translates to better phase purity of α -FAPbI₃ perovskite and improved crystallinity, ultimately leading to encapsulated p–i–n lab cells with an ITO/Methoxy-substituted [2-(9*H*-carbazol-9-yl)ethyl]phosphonic acid (MeO-2PACz)/FAPbI₃/LiF/C₆₀/BCP/Cu configuration displaying a remarkable T_{80} of approximately 8760 h in Germany.¹⁶⁸ Similarly to the strategy followed by Yoo *et al.*, the mixed use of solvents here tackled the combined effect of temperature and bias given its focus on increasing crystallinity and grain size. In this context, techniques like blade coating and slot-die coating benefit from tailored solvent systems to control nucleation and crystallization, hence making these approaches compatible with such techniques. For instance, the use of nitrogen-stream-assisted meniscus control in blade coating is a scalable approach compatible with roll-to-roll processing.⁹² Still, challenges remain, namely the need for precise control of solvent ratios and drying conditions, which may require advanced equipment for gas flow or temperature regulation in industrial settings. Furthermore, the formation of intermediate adducts (*e.g.*, FAI-PbI₂-NMP) to enhance crystallinity is scalable but requires careful precursor formulation to avoid batch-to-batch variations, which could impact production consistency.

Bulk modifications of charge transporting layers also allow improvements in outdoor stability. In state-of-the-art p–i–n devices a common practice is to employ a wide gamut of self-assembling monolayers (SAMs) as HTL, namely [2-(9*H*-carbazol-9-yl)ethyl]phosphonic acid (2PACz), MeO-2PACz, Methyl-substituted [2-(9*H*-carbazol-9-yl)ethyl]phosphonic acid (Me-4PACz) and Methoxy-substituted [2-(9*H*-carbazol-9-yl)ethyl]phosphonic acid (MeO-4PACz).^{212,213} Due to their molecularly thin nature, they define the quality of the transparent conductive electrode (TCO)/HTL/perovskite interface, namely by guaranteeing good adhesion of the perovskite layer to the TCO substrate and by protecting the latter from ion-induced indium reduction (typical TCO is ITO). The former aspect is dealt with molecular design, where typically moieties of –PO(OH)₂ bind to ITO,²¹⁴ while the latter is dealt with by the ion blocking capability of the SAM molecules. Jiang *et al.* employed DFT calculations to show that,



for a FA-rich perovskite composition, modifying the SAM composition by mixing MeO-2PACz and Me-4PACz resulted in decreased $\text{In}^{3+} \rightarrow \text{In}^0$ reduction stemming from electrochemical reactions at the buried interface with the perovskite organic cation. Such reactions are especially promoted under operation by the electric field that can transport FA cations closer to the interface, where if they are able to penetrate through the SAM the reduction reaction will occur. Furthermore, light exposure can aggravate this, as photoinduced production of HI near the interface yields I^- ions that can react with In^0 to form InI_3 , effectively etching the electrode, hence harming performance over time.²¹⁵ The main difference between MeO-2PACz and Me-4PACz lies in their hydrophilicity, given that the MeO group in the former has a stronger interaction with ions compared to the latter, non-polar in nature. This causes FA ions to display an increased ability to reach the ITO interface in the case of sole use of MeO-2PACz, in which case the aforementioned reactions can take place. Including instead a modest amount of Me-4PACz can alleviate this while maintaining a good adhesion of the perovskite layer upon coating.¹³² Consequently, encapsulated lab cells fabricated with such HTL in an ITO/MeO-2PACz+Me-4PACz/Rb_{0.05}Cs_{0.05}MA_{0.05}FA_{0.85}Pb(I_{0.95}Br_{0.05})₃/C₆₀/SnO₂/Ag configuration displayed a T_{90} of 3700 h of outdoor exposure in the USA. This approach focused mainly on addressing the instability against the combined effect of electrical bias and light exposure. In view of high-throughput fabrication for market insertion, these SAMs are typically applied *via* solution processing, which is compatible with high-throughput techniques like slot-die coating or spray coating.²¹⁶ Their molecularly thin nature minimizes material consumption, reducing costs for large-scale production.²¹⁶ However, achieving uniform SAM deposition over large areas requires precise control of surface preparation and coating conditions to ensure consistent adhesion and ion-blocking properties, which could be challenging on an industrial scale.²¹⁷ Such difficulties arise from inhomogeneous coating namely owing to the tendency of phosphonic acid groups to aggregate in solution^{217,218} and partial SAM desorption from the substrates during perovskite precursor coating.^{219,220}

In the case of n-i-p devices, a typical ETL is SnO₂, which upon annealing during processing displays surface protonation resulting in the formation of -OH groups that act as interfacial electronic defects yielding non-radiative recombination-related losses.²²¹ The formation is driven by the presence of two active sites on the surface of the as-deposited SnO₂ films: a bridge oxygen and a 5-fold (under)coordinated Sn. The latter acts as a Lewis acid attracting water molecules from the air in the vicinity of the film, which readily dissociate and transfer a proton to the bridge oxygen.²²² A clever strategy to mitigate this is using pH-induced agglomeration of SnO₂ nanoparticles in the corresponding precursor used for deposition.⁹¹ Fine control to reduce the pH of the precursor by addition of HNO₃ acid allows a controlled agglomeration of nanoparticles thanks to the appearance of oxo groups linking them guided by the equilibrium given by $\text{Sn-OH} + \text{OH}^- \leftrightarrow \text{Sn-O}^- + \text{H}_2\text{O}$. The addition of HNO₃ shifts the equilibrium in solution to the left,

which owing to the reduced charge of colloidal nanoparticles allows them to come in close contact, hence aggregating by crosslinking between them upon the formation of oxo groups (Sn-O-Sn). This prevents protonation of oxygen during later annealing of the as-deposited SnO₂ films, significantly lowering the amount of defective -OH groups appearing in the final film and hence decreasing the losses due to undesired recombination. The main consequence of this is an enhanced performance, which to some extent aids long term performance because larger losses can be somehow compensated by the higher initial performance. Additionally, oxo group crosslinking also allows the final SnO₂ film to be more compact and display less pinholes, key to prevent defect-initialized permeation of moisture or oxygen.⁹¹ In this way, encapsulated n-i-p lab cells with an ITO/SnO₂/perovskite/Spiro-OMeTAD/Au configuration maintained unchanged power output during 720 h hours of outdoor operation in South Korea. The pH-induced agglomeration strategy addresses the effect of both potential oxygen and moisture that may enter the device when encapsulation is not perfect, key in outdoor operation regions with tropical climate with a high humidity throughout the year that could trigger such permeation. In addition, it could potentially be integrated into existing coating workflows, since ultimately the key upgrade would be the addition of HNO₃. However, the challenge lies in maintaining consistent pH control and nanoparticle dispersion in large-scale precursor solutions, as variations could lead to defects or reduce film quality.²²³

3.1.2. Interface modulation. This is one of the key aspects of improving the overall performance and stability of PSCs. This strategy can influence the charge transfer, recombination and the operational lifetime. Interface modulation refers to strategies like adding buffer layers at the perovskite/charge transport material interface, which act as defect passivation materials that reduce non-radiative recombination and enhance stability.^{224,225} In this context, adding adducts onto the surface of perovskite films can patch grain boundary cracks and passivate interfacial defects.²²⁶ Lewis base passivation can reduce nonradiative electron-hole recombination and enhance photoluminescence lifetimes by nearly up to 2 μs .²²⁷ Interfacial recombination occurs due to poor contact or defect-rich interfaces. Strategies like employing self-assembled monolayers (SAMs) with high polarity can improve the band alignment and minimize the energy loss suppressing recombination and improving open-circuit voltage (V_{oc})²²⁸ and stability. Chemical surface treatments and interface modulation using hydrophobic or encapsulating materials have been shown to enhance the long-term stability of perovskite solar cells. For example,²²⁹ incorporating 2D perovskite materials into the perovskite layer improved moisture tolerance and stability without significantly compromising efficiency. Unfortunately, not all devices made using the previously mentioned strategies have been tested under real-world operating conditions, and most of them have been evaluated under simulated-laboratory conditions; additionally, most of the devices are less than 10 cm² of area (lab cells).



Recent studies have used self-assembled monolayers (SAMs) to enhance light stability and outdoor performance. Sekimoto *et al.*¹⁸¹ proposed the use of isonicotinic acid/C₆₀ SAM between the electron transport layer and the perovskite. Long-term outdoor stability was tested in Osaka, Japan, on encapsulated cells under open-circuit conditions for six months, with a cumulative light exposure of 774 kWh m⁻². The results showed that the unpassivated TiO₂ surface reduced thermal stability, whereas the cell with the modified interlayer, with an ITO/ATO/c-TiO₂/mp-TiO₂/C₆₀SAM + isonicotinic acid/perovskite/BABr/PTAA/Au configuration, exhibited an almost unchanged PCE and improved durability against light and heat, with an estimated T_{100} of approximately 4032 hours. The origin of such improvement lies in the interlayer suppressing the deleterious cathode reaction, $\text{Pb}^{2+} + 2\text{e}^{-} \rightarrow \text{Pb}^0$, at the ETL/perovskite interface, which in turn also suppresses the detrimental anode reaction, $\text{I}^{-} + \text{h}^{+} \rightarrow \text{I}^0$, at the opposite HTL/perovskite interface. Such reactions ultimately originate from the volatility of the MA cation used in the triple-cation composition studied in this work. Loss of organic cations and iodine upon exposure to white light is known to take place in MA-rich compositions, being less of an issue when MA is replaced by FA or I by Br.²³⁰ This leaves PbI₂ as a byproduct that can further undergo photolysis, yielding Pb⁰ by the previously shown reaction thanks to the availability of photogenerated electrons.²³¹ The interface treatment with isonicotinic acid improved the electron extraction at the ETL/perovskite interface, both favoring performance and stability by effectively reducing the availability of “free” photogenerated electrons to reduce Pb²⁺. This approach hence directly confronts the issue of light instability, thus yielding resilient outdoor performance despite light-induced permeation of iodine into the HTL taking place. Interlayers of potassium pyrophosphates have also been used to improve the performance of semi-transparent perovskite solar cells. In a recent study,²³² interface passivation (SnO₂/perovskite) was performed by bonding potassium pyrophosphate (KPP) to the SnO₂ surface, enabled by the coordination of Sn *via* Sn–O linkages with the pyrophosphate group (P₂O₇⁴⁻).²³³ The compound was able to passivate undercoordinated atom sites, allowing a higher proportion of Sn⁴⁺ which is beneficial for the optoelectronic properties.²³⁴ The Sn²⁺ counterpart instead corresponds to Sn dangling bonds due to oxygen vacancies and produces non radiative recombination at the SnO₂/perovskite interface given that a trap state within the bandgap is formed.²³⁴ The proposed strategy in turn yielded perovskite films with enhanced crystallinity, which as mentioned before can tackle the instability caused by electrical bias due to reduced grain boundaries. The authors also claimed enhanced thermal stability since they found less PbI₂ in their perovskite films following thermal annealing. However, since they employed a double-cation composition (mostly FA with some MA) and PbI₂ is not amongst the decomposition products of FAPbI₃ in the temperature range of outdoor operation (*e.g.*, 70–75 °C in summer), the lower presence of PbI₂ could be ascribed to the same phenomenon highlighted in the previous approach, where in this case the prevention of charge trapping

at the relevant interface hinders photolysis of photogenerated PbI₂ by the light-induced loss of MA cations. Ultimately this enabled an estimated T_{80} of 1104 hours of outdoor exposition in South Korea for devices with an ITO/SnO₂/perovskite/Spiro-OMeTAD/Au configuration. The perovskite/HTL interface in more mainstream n–i–p configuration has also been modified. For instance, deposition of a delaminated 2D Ti₃C₂ MXene layer intercalated with 3-phosphonopropionic acid (H3pp) onto a quadruple-cation perovskite film has demonstrated to down-shift both surface work function and valence band of the perovskite film, allowing enhanced hole-extraction at the interface. Furthermore, it enabled a 10-fold and 1000-fold decrease in the density of shallow and deep interface trap states, respectively.²³⁵ Ti₃C₂ typically displays Ti outer layers functionalized with O and OH groups that allow an electrostatic interaction with the perovskite, leading to a transferred charge density highly localized at the first Pb–I layer of the perovskite material. This forms an interface dipole whose magnitude depends on the groups involved (*e.g.*, O or OH), being directly responsible for the surface work function shift.²³⁶ This effect is further complemented by the presence of H3pp, resulting in decreased trap density at the interface. Karimipour *et al.* evaluated the lab cell devices with a FTO/c-TiO₂/m-TiO₂/Perovskite/MXene:H3pp/Spiro-OMeTAD/Au configuration under outdoor exposure in Spain and obtained a T_{80} of 600 hours. The focus on diminished charge trapping at the interface points to enhanced stability against light, as previously explained, given that effectively less photogenerated charges will be available at the interface to undergo deleterious redox reactions. Inverted p–i–n configuration has also been the subject of different outdoor testing, particularly by modifying the HTL/perovskite interface. An important aspect that can be enhanced with such an interface is reproducibility, which is particularly relevant for industrial-scale production. The solution-processing required for device fabrication yields inherent pinholes and non-uniformities in the films that can lead to device failure due to shunt paths. Back in 2019, Ramirez *et al.*²³⁷ tackled this issue by inserting a 100 nm mesoporous layer of Al₂O₃ at the HTL/perovskite interface, which homogenized the perovskite film coverage to prevent the formation of pinholes and thus shunting paths through the layer upon deposition of further charge carrier films and metal electrode. This led to enhanced reproducibility evident in less steep decline in PCE when increasing device size and more consistent V_{oc} values. Pinholes yield a reduced distance between opposite electrodes, which in turn allows an increased electric field upon operation.²³⁸ Hence, in view of stability, a reduction in pinhole formation enables better stability against bias. Complementarily, the mesoporous Al₂O₃ layer also reduced the trap states at the HTL/perovskite interface,²³⁹ further enhancing bias resilience and ultimately leading to minimodules with an ITO/NiO_x/m-Al₂O₃/MAPbI₃/PCBM/Rhodamine/Ag configuration displaying a T_{80} of around 1900 h under outdoor conditions in Colombia, an outstanding stability achieved under harsh tropical climatic conditions (Fig. 3b). In an industrial environment, deposition of Al₂O₃ nanoparticles is well-suited for high-throughput processing.



Screen printing, in particular, is a mature industrial technique that can facilitate large-scale production of mesoporous layers, provided uniformity is maintained.²⁴⁰ However, challenges include controlling the porosity and thickness of the mesoporous layer over large areas, which is especially critical for such thin layers. Additionally, variations in nanoparticle dispersion or coating conditions could lead to inconsistent film quality, requiring optimization of precursor inks and deposition parameters.

Fei *et al.*⁹³ modified the buried ITO/PTAA/perovskite interface by adding an aromatic phosphonic acid, [2-(9-ethyl-9H-carbazol-3-yl)ethyl]phosphonic acid (EtCz3EPA). At such interface, mainly three types of defects are present following fabrication: I_i^- (type I), I_i^+ (type II) and a defect related to the formation of an amorphous perovskite region (type III) stemming from phase segregation.⁵² The density of all such defects increases following illumination during operation due to the UV region of the spectrum. Type II defects increased due to accelerated I-migration upon illumination, and since trapping of photogenerated electrons is possible by such defects excess holes are created which trigger A-cation migration and phase segregation.²⁴¹ A weak anchoring between the perovskite and the ITO/PTAA substrate was identified as the main cause for this inadequate evolution under illumination, and hence EtCz3EPA was introduced as an interface layer capable of anchoring both ITO and perovskite at opposite ends. The $-COOH$ and $-PO(OH)_2$ groups of the molecule could link to the $-OH$ groups of ITO, while the $-NH-$, $=O$, halide ($-X$) and carbazole groups could interact with the Pb^{2+} in the perovskite. Furthermore, the preferred in-plane orientation of the carbazole group allows better interaction of N with Pb^{2+} in the perovskite, indeed resulting in better anchoring compared to sole PTAA. Consequently, type-III defects (amorphous perovskite phase) were significantly suppressed and the increase of all defect types upon illumination was lowered, thus allowing reduced cation migration and phase segregation. This strategy hence directly bridged issues stemming from the coupled effects of temperature, electrical bias, and light, given the reduction of defects capable of trapping photogenerated carriers and whose migration is triggered during operation by heat and electric fields. The authors obtained outstanding outdoor stability results in Colorado (USA) with a T_{100} of 4872 hours for minimodules with an ITO/PTAA/ $FA_{0.9}CS_{0.1}PbI_3/C_{60}/BCP/Cu$ configuration displaying an aperture area of 17.88 cm^2 . In view of industrial fabrication, this approach displays the same benefits and drawbacks as typical SAMS, namely compatibility with slot-die and spray coating²¹⁶ and challenging uniformity over large areas.²¹⁷ Alternatively, an ETL based on perylendiimide derivatives compared to Phenyl C61 Butyric Acid Methyl ester (PCBM) has proven to improve the stability of (ITO/PEDOT:PSS/MAPbI₃/ETL/Ag) solar cell.²⁴² In the study by Akbulatov *et al.*, TOF-SIMS depth profiling revealed that, after device operation, a layer of AgI is formed at the ETL/Ag interface regardless of the type of ETL employed, although when using PCBM the proportion of such layer was higher. This is a result of MAPbI₃ splitting into PbI₂ and MAI (*e.g.*, owing to photothermal

decomposition upon operation),⁹³ the latter one being a volatile compound capable of diffusing through the ETL, reach the back Ag electrode and corrode it to AgI.^{104,243,244} Although such process takes place for both types of ETL, in the case of the perylendiimide derivative, such diffusion only occurs through grain boundaries, which has a less negative effect on device performance over time. Additionally, MAI decomposition product was found to dimerize to (MAI)₂, which should be intercalated in the ETL for diffusion to take place. DFT calculations showed this to be the main reason for increase stability, since such process can readily take place in PCBM while being heavily unfavorable for the perylendiimide derivative, which forms a very compact 3D network *via* van der Waals interactions. In this way, the proposed strategy bridged instabilities arising from the coupled effect of temperature and light, as harmful effects stemming from photothermal decomposition were mitigated. The fabricated lab cell scale devices were exposed to natural outdoor conditions in the Negev desert (Israel) to obtain a T_{80} of about 41 hours when using the perylendiimide derivatives compared to less than 5 hours for PCBM.²⁴² Boron chloride subphthalocyanine (Cl₆SubPc) has also been used to passivate the perovskite/fullerene ETL interface in inverted PSCs.²⁴⁵ In an ITO/Cu:NiO_x/perovskite/ETL/C60:BCP/Ag device, Chen *et al.* found that N and B atoms from Cl₆SubPc underwent strong chemical bonding with migrating ions from the perovskite, N allowing I^- adsorption (more than 20 times stronger compared to bare C₆₀ based on DFT and molecular dynamics simulations), key to prevent metal electrode corrosion upon operation, and B enabling electron transfer with both perovskite and C₆₀.²⁴⁶ Furthermore, Cl ions from Cl₆SubPc display higher binding energy towards Pb compared to bare C₆₀, hence allowing stronger linking between perovskite and ETL. This resulted in effective inhibition of the halide ion migration at the perovskite/ETL interface, directly mitigating temperature and bias effects, *e.g.*, suppression of electrode corrosion as the rate of corroding migrating ions reaching it is lower. Synergistically, the enhanced interface linkage favors charge transport, *i.e.*, aiding the issue of charge accumulation at the interfaces whose deleterious effect has been previously described and related to light exposure. Consequently, the outdoor testing results in China showed a T_{87} of 1728 hours under ISOS-O-1 for lab cells. The use of specific materials like perylendiimide derivatives and Cl₆SubPc would still be highly compatible with solution processing at large scale, however, their scalability is limited by their synthesis complexity and cost, which may require highly specialized production facilities.²⁴⁷

The paper presented by Cao *et al.*²⁴⁸ used NiO_x not as a direct HTL but as a redox-sensitive metal oxide into a perovskite to obtain an active composite film. NiO_x nanoparticles (NP) were incorporated into the perovskite layer using the antisolvent process to prevent possible surface $-OH$ groups at the NPs surface from triggering proton removal of the A-site organic cations.²⁴⁹ With the aim of chemically passivating Pb⁰ and I⁰ defects, the authors successfully employed the Ni²⁺/Ni³⁺ ion pair, displaying a suitable redox potential and cost-



effectiveness, to continuously shuttle electrons leading to oxidation ($2\text{Ni}^{3+} + \text{Pb}^0 \rightarrow 2\text{Ni}^{2+} + \text{Pb}^{2+}$) and reduction ($\text{Ni}^{2+} + \text{I}^0 \rightarrow \text{Ni}^{3+} + \text{I}^-$) of such defects, respectively. Additionally, NiO_x causes a slight increase in the oxidation energy of I^-/I_3^+ and I_3^-/I^0 , offering energetic passivation by increasing the energy required for iodine oxidation.²⁵⁰ X-ray Photoelectron Spectroscopy (XPS) and Raman analysis demonstrated the formation of strong Ni–I and Pb–O bonds, which hampers loss of volatile iodide due to thermal stimuli, whilst DFT calculations showed an increase of the formation energy of iodide vacancies in PbI_6 octahedral frameworks from 0.45 to 0.97 eV, hence impeding such vacancies and ion migration. Furthermore, the p-type doping and barrier effect of NiO_x shifted the electric field at the HTL/perovskite interface from positive to slightly negative, which is highly beneficial for charge extraction at such interface.^{250,251} The combined effect of these phenomena offers a thorough approach to mainly tackle coupled temperature and bias instabilities by fixation of volatile species, compensation of charges losses that produce defects and reduction in charge accumulation at interfaces. In this way min-modules were fabricated, consisting of an effective area of 18 cm² and 8 series-connected perovskite sub-cells fabricated on a FTO substrate (6×6 cm²). They were subjected to outdoor exposure in China and displayed a T_{95} of 1600 hours for devices with a FTO/ZnO–TiO₂/perovskite/ NiO_x –NiPc/Au configuration. When considering high-throughput fabrication, some potential challenges could be ensuring uniform NiO_x nanoparticle distribution and preventing aggregation, which could introduce defects. The need for precise control of antisolvent timing and nanoparticle concentration may complicate high-throughput processing, requiring automated systems to maintain consistency.

3.1.3. Back electrode design. Gold and silver electrodes, despite their excellent conductivity and energy alignment, suffer from interfacial degradation that limits PSC lifetimes.^{104,244,252,253} Halide ions (I^- , Br^-) released under the action of light, temperature, or humidity form insoluble metal halides with the electrodes (AuI₃, AgI), which increases contact resistance.²⁵⁴ At the same time, the bidirectional migration of Ag^+ and I^- induces dendrites that create bypasses,^{252,255,256} while Pb^{2+} –Au galvanic reactions and photocorrosion further deteriorate the integrity of the electrode.^{253,257} Ultrathin ion-blocking interlayers (*e.g.*, NiO_x , Copper(I) thiocyanate (CuSCN)), have proven effective at mitigating these pathways and extending device stability²⁵⁸ although further outdoor testing of such strategies is lacking.

Accordingly, some efforts have been made to find alternatives to replace these metals as counter electrodes. Notably carbon-based materials thanks to their outstanding conductivity and inertness,¹⁸³ specially against ion migration between perovskites and metal electrodes,²⁵⁹ not requiring heating of the device during deposition which is itself an intrinsic advantage for device stability, alongside low-cost fabrication and mechanical flexibility.²⁶⁰ A big-data-driven analysis of overall strategies to enhance the stability of PSCs even demonstrated that employing carbon-based electrodes can outperform any

other strategy available at the time of publication,¹⁰⁵ thus highlighting the potential of this carbon-based-electrode approach. However, examples of this approach being tested outdoors are rather limited.

Carbon-based materials have been implemented in back electrodes of perovskite solar devices in two main ways: as thin films replacing the typical metal electrode^{261–263} and as a mesoscopic thick films (thicknesses of 10–12 μm are common) that serve as back electrode and simultaneously pose a template onto which the perovskite absorber material can be infiltrated,^{264–266} implying a fabrication approach different from the typical layer by layer deposition strategy. Herein, only the first group is discussed, and the second one is treated in the following section.

The carbon-based back electrodes of outdoor evaluated perovskite devices have been manufactured using different techniques, namely film transfer and typical thin film deposition, *e.g.*, blade coating and screen printing. Using an HTL-free configuration, a film-transfer approach has demonstrated good results.²⁶³ For this, a hybrid ONC/MWCNT (Organic Nanocrystal/Multi-Walled Carbon Nanotubes) film was prepared by self-assembly, with an overnight vacuum drying process that yielded free standing films. Subsequently, the films were transferred onto devices to work as back electrode by performing a pre-wetting in acetonitrile followed by a gentle stamping onto the device. Conventional devices using Au as back electrode can readily undergo photo-induced Au diffusion into the perovskite layer following operation, which has been demonstrated to have detrimental effects on performance and in this case is highly likely due to the absence of an HTL that could act as a diffusion barrier.²⁶⁷ Hence, the replacement of Au for the hybrid ONC/MWCNT layer is key to coping with this issue. Upon outdoor exposure, no encapsulation was employed thanks to the resilience of the hybrid electrode to air and moisture, which coupled with an inorganic perovskite absorber within a device with a glass/FTO/c-TiO₂/mp-TiO₂/CsPbBr₃/carbon-based electrode resulting in an estimate T_{80} of 450 h in Israel. This strategy is particularly clever since it removes the complication of the deposition of the electrode (usually from a slurry-like solution) onto the perovskite material which can be experimentally difficult and severely damage the substrate material. Furthermore, it showcases a smart way to tackle thermal and light instability originated at a device level, not directly from the perovskite itself or the adjacent charge transport layers, as the main addressed issue stemmed from metal electrode migration. In addition, the hydrophobic nature of the carbon electrode offers an additional barrier against moisture induced degradation, aiding device stability in view of possible leakages through the encapsulation system. Another study²⁶² reported a similar HTL-free device with a “bio-inspired” carbon-based back electrode fabricated with graphitic carbon from an invasive plant species. In contrast to the previous work, a full solution-processed device was studied in which an inter-layer of graphitic carbon and $\text{MAPbI}_{3-x}\text{Cl}_x$ perovskite was inserted between the absorber and the back-carbon electrode. Furthermore, the deposition of the electrode was performed by



brush-painting, reducing damage to the underlying perovskite layer, which is important to reduce pinholes that produce locally increased electric fields with negative effects in terms of ion migration. In this way, encapsulated n-i-p lab cells with a FTO/TiO₂/Al₂O₃/perovskite/carbon-based electrode architecture displayed an approximate T_{80} of 1200 h under outdoor exposure in Norway. More recently, an encapsulated n-i-p flexible minimodule (ITO/SnO₂/Cs_{0.17}FA_{0.83}PbI_{2.7}Br_{0.3}/PTAA/Carbon) with a screen-printed carbon back electrode with remarkable stability has been achieved, directed towards application in a perovskite-powered global positioning system (GPS) for an internet of things (IoT) device that monitors wildlife bison.²⁶¹ Such device achieved a T_{70} of 8400 h under outdoor exposure in Poland and exploited the same inherent benefits of carbon materials as previously stated. Overall, the carbon-based electrode strategy is highly compatible with high-throughput fabrication. Screen printing, as demonstrated in Poland,²⁶¹ aligns well with roll-to-roll processing and is cost-effective due to the low cost and inertness of carbon materials. The film-transfer approach,²⁶³ while innovative, is less scalable due to its time-intensive preparation and manual steps but could be improved with automated lamination systems. Challenges for solution-processed electrodes include ensuring uniform conductivity and adhesion over large areas, which can be addressed through optimized ink formulations and automated deposition systems. The absence of high-temperature processing and the mechanical flexibility of carbon electrodes further enhance their suitability for industrial-scale production.

A summary of the outdoor stability results discussed above is shown in Table 1, analyzing the key concept strategy used in the cited papers and its effect on device stability. It is important to note that this represents only a minor subset of the comprehensive body of work on perovskite solar cells. For instance, the Perovskite Database^{40,41,268} includes over 43 000 entries of published device data. This contrast underscores the limited experimental efforts devoted to testing devices under real-world conditions. Given that perovskite solar cell research has accelerated within the past 15 years, this limited number of outdoor studies is understandable. It also highlights the need to bridge the gap between advanced device architectures and their real-world performance.

Despite this, there is a clear trend: more outdoor stability testing, with more robust devices and longer lifetimes. Fig. 4b shows how manufacturing strategies for improving outdoor stability have evolved over time. It presents representative results from different approaches, including T_{80} , T_{90} , T_{93} , T_{95} and T_{100} values, tracked over several years. Among these, interface modulation has been the most studied strategy, achieving notable improvements in stability. For example, devices advanced from a T_{80} of just 41 hours in 2017²⁴² to minimodules with no observable performance loss after 4872 hours of outdoor exposure.⁹³ The bulk modification approach ranks second in terms of the number of studies, with results such as a T_{93} of 2000 hours in minimodules.⁹²

Notably, while the use of carbon back electrodes has demonstrated outstanding stability potential, such as a reported T_{70} of

8400 hours,²⁶¹ this strategy has received relatively little attention. Only three outdoor studies on carbon-based devices are included in Fig. 4b. In general, outdoor stability results in this approach trail behind those interface and bulk modifications. This issue may be attributed to the intrinsic challenges associated with exposing devices into the environment, including inadequate interfacial adhesion, suboptimal surface wettability, and insufficient ohmic contact between the carbon electrode and underlying layers.¹⁸³ These issues can be further compounded in outdoor environments. Nevertheless, one of the most promising approaches to producing stable, cost-effective perovskite modules is still the use of HTL-free designs with carbon electrodes (discussed in the next subsection). There is considerable potential for large-scale deployment, particularly in situations where longevity exceeds peak efficiency. This is due to their chemical robustness, moisture resistance, and compliance with ambient, low temperature processing.

Unlike the more chemistry-oriented strategies of interface and bulk modification, the carbon back-electrode approach requires substantial engineering effort. The materials and methods for carbon electrode integration into the stack are important to achieving stable performance outdoors. The mentioned strategies can be used combined in high-efficiency devices, especially in tandem structures which require advanced techniques to improve performance.^{175,269–274} This is particularly relevant for tandem devices, which often require complex architectural strategies to optimize both performance and stability.^{275–280} While these approaches have yielded some of the highest efficiencies reported to date, they too lack systematic benchmarking under outdoor conditions. Expanding real-world validation across both single junction and tandem architectures will be essential to ensure perovskite photovoltaics meet commercial reliability standards. Some studies display devices with promising stability results under stringent conditions, such as T_{85} of 1250 hours,²⁶⁹ T_{95} of 2000 hours,²⁷³ and T_{96} of 2000 hours²⁷⁴ under ISOSL3 protocols. Though developments seem promising, few are reported in the literature specifically showing outdoor testing.

As demonstrated in Table 1, the fabrication approaches exhibited favorable results in outdoor conditions. All studies incorporated at least one complementary indoor test, thus enabling the correlation of degradation mechanisms observed under controlled laboratory conditions with real-world operating behavior. It was demonstrated that strategies based on bulk modification (B.M.) exhibited the longest stability times, reaching $T_{80} > 5000$ h under ISOS-L1I, which an inherent improvement in perovskite stability. On the other hand, more stringent protocols (ISOS-D2, ISOS-L3) have been used to evaluate Interface Modulation (I.M.) approaches, with T_{90} values approaching 2000 hours and T_{100} values around 1100 hours. These findings highlight the efficacy of interfacial strategies even under more demanding ISOS protocols. Finally, even without encapsulation, back electrode design (B.E.D.) techniques produced T_{90}/T_{100} values greater than 1200 hours, highlighting their role in mitigating degradation pathways.



Table 1 Summary of relevant data from papers displaying outdoor results of perovskite solar devices using several fabrication approaches. Device sizes are presented as defined in ref. 45. Figures were adapted or reprinted with permission from ref. 168 (CC BY 4.0, 2023),⁹¹ (CC BY-NC 4.0, 2024),⁹² (Copyright © 2024, Elsevier),²³⁷ (Copyright © 2019, Elsevier),²⁴⁵ (Copyright © 2021, Elsevier),¹⁸¹ (Copyright © 2023, American Chemical Society),²⁴⁸ (Copyright © 2024, American Chemical Society),⁹³ (CC BY-NC 4.0, 2024)

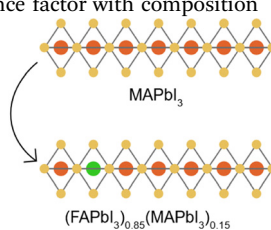
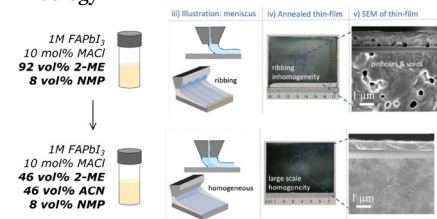
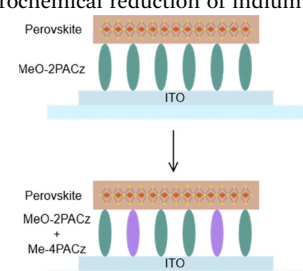
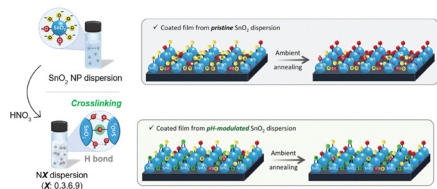
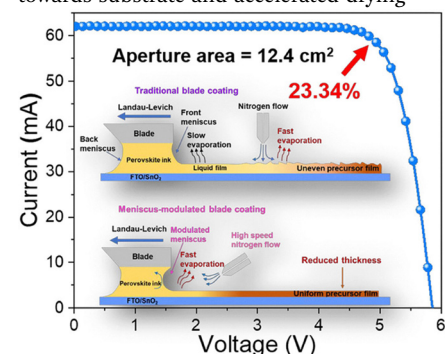
Approach	Year	Device size	Stability parameter	Stability time (h)	Country	Key concept	Ref.
B.M.	2016	Lab cell	T_{80}	846	Spain	Stabilization of α -phase by modifying the tolerance factor with composition 	196
	2023	Lab cell	T_{80}	8760	Germany	Fine tuning of adduct formation to template crystallization and modification of solution rheology 	168
	2023	Lab cell	T_{90}	3700	USA	Mixing of SAMs to enhance perovskite adhesion to TCO substrate and suppress electrochemical reduction of indium in ITO 	132
	2024	Submodule	T_{100}	720	South Korea	Link of SnO ₂ nanoparticles with oxo groups to prevent the formation of -OH electronic defects 	91
	2024	Minimodule	T_{93}	2000	China	Reduction of precursor solution advancing towards substrate and accelerated drying 	92



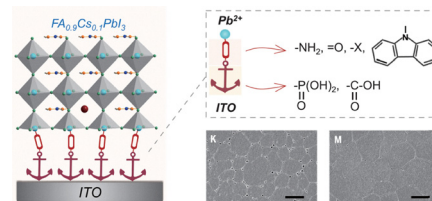
Table 1 (continued)

Approach	Year	Device size	Stability parameter	Stability time (h)	Country	Key concept	Ref.
I.M.	2017	Lab cell	T_{80}	41	Israel	Replacement of [60]PCBM with ETL showing lower diffusion rate of volatile perovskite decomposition product	242
	2019	Minimodule	T_{80}	1900	Colombia	Perovskite morphology homogenization using mesoporous underlayer	237
	2021	Lab cell	T_{95}	1272	China	Chemical bonding of migrating ions to prevent electrode corrosion and enhance electronic between perovskite and ETL	245
	2023	Lab cell	T_{100}	4032	Japan	Suppression of electrochemical lead reduction and iodine oxidation	181
	2023	Lab cell	T_{80}	600	Spain	Decrease of interfacial deep and shallow traps	235
	2024	Minimodule	T_{80}	1104	South Korea	Passivation of undercoordinated tin atoms in HTL via phosphate bonding	232
	2024	Minimodule	T_{95}	1600	China	Suppression of lead reduction and iodine oxidation via Ni ²⁺ /Ni ³⁺ ion pair	248



Table 1 (continued)

Approach	Year	Device size	Stability parameter	Stability time (h)	Country	Key concept	Ref.
2024		Minimodule	T_{100}	4872	USA	Chemical engineering of SAMs to enhance anchoring of perovskite to ITO	93
B.E.D	2019	Lab cell	T_{80}	450	Israel	Fabrication of free standing back electrode with subsequent transfer onto the device	263
	2022	Minimodule	T_{80}	1200	Norway	Coupling of paint-brushed carbon back-electrode <i>via</i> graphitic carbon-MAPbI ₃ -xCl _x interlayer	262
	2024	Minimodule	T_{70}	8400	Poland	Use of screen-printed highly stable carbon back-electrode	261



In the ref. 91, which employs the bulk modification (B.M.) strategy, the ISOS-D3 protocol is used under conditions of 85 °C and 85% relative humidity. The device exhibits greater stability in indoor-D3 ($T_{80} = 1080$ h) than in outdoor tests ($T_{80} = 720$ h). This occurs because, although ISOS-D3 imposes temperature and humidity stress, this stress is constant, controlled, and static. Under such conditions, uniform diffusive and thermal processes dominate degradation, allowing both the encapsulant and the perovskite to maintain stable behavior. In contrast, the encapsulated device is subjected to a highly dynamic and multifactorial environment during outdoor tests. This environment includes real solar irradiance (including UV), day-night thermal cycles, humidity variations, atmospheric oxygen, and climatic agents like wind gusts, dew, and intermittent shading. Together, these elements speed up the encapsulant's and interfaces' deterioration and cause failure pathways that do not appear under controlled damp-heat. As a result, even though B.M. greatly increases stability indoors, its outdoor performance shows that the encapsulation used is insufficient to mitigate the entire range of degradation mechanisms, which explains the lifetime reduction from 1080 to 720 hours.

Regarding the ref. 237 and the interface modulation (I.M.) strategy, the ISOS-D1 protocol is used with a relative humidity of 60%. Here, the absence of encapsulation is crucial in explaining the marked difference between indoor ($T_{72} = 61$ h) and outdoor ($T_{80} = 1900$ h) stability. In indoor environments without encapsulation, degradation is driven by direct exposure to ambient humidity, immediately compromising the functional interfaces. In contrast, for outdoor testing, these devices were encapsulated, preventing direct water permeation, and allowing the interfacial strategy to demonstrate its effectiveness under more complex climatic conditions. This shows that, for humidity-sensitive devices, encapsulation can become the dominant factor determining overall stability.

Finally, in the article²⁶³ evaluates the back electrode design (BED) strategy using the ISOS-D1 protocol under ambient

temperature and relative humidity conditions between 25–30%. In this case, the unencapsulated device exhibits a T_{100} of 1440 h, while outdoors, even when encapsulated, the T_{80} decreases to 450 h. This demonstrates that high indoor stability does not necessarily translate to real-world conditions. Outdoors, the simultaneous presence of UV irradiance, fluctuating humidity, and thermal cycles induce degradation mechanisms, especially in the encapsulant and back contact, that remain inactive under a stable indoor environment. Consequently, indoor tests are useful for preliminary diagnosis but mandate respective outdoor validation.

3.1.4. Indoor-tested strategies and their transferability to real-world operation. The strategies discussed in the previous sections (Table 1) have been tested under outdoor conditions, validating that they are adequate to integrate into devices. However, many strategies developed for PSCs have still not been proven under real-world operation. In the state of the art of stability evaluation of PSCs, the majority of tests correspond to ISOS indoor protocols, which enable the assessment of different strategies for improving device durability under standardized conditions. Indoor protocols are indispensable because they isolate mechanisms (single-stress effects), accelerate degradation (rapid evaluation), provide reproducible conditions, enable fast screening, and allow global standardization, while outdoor tests provide the ultimate validation. This verification is essential to confirm that, under real-world operating conditions, the strategy evaluated indoors can function effectively. For this reason, it is necessary to ask whether a strategy that performed well in indoor tests can be directly assumed to work outdoors. The answer is negative. The two approaches are complementary, not interchangeable. To address this gap, different methods exist today to attempt to predict the fingerprint of outdoor performance from indoor tests, which are discussed in Section 5.3.

To illustrate this point, we discuss representative examples in which a strategy that performs well under indoor conditions requires additional considerations to succeed outdoors, highlighting that indoor performance alone is not sufficient for real-



world deployment. A prime example of this is shown by Fei *et al.*, who demonstrate that devices using PTAA operate well and remain stable under LED-based indoor protocols but degrade rapidly outdoors.⁹³ As the chemical mechanism behind this instability has already been described in Section 3.1.2, here we emphasize that this degradation pathway is activated primarily by UV exposure, an effect absent in conventional ISOS indoor tests. Consequently, strategies that appear robust under indoor conditions may not translate to outdoor stability unless they are specifically designed to withstand UV-induced stress. In this context, Fei *et al.* show that replacing PTAA with the phosphonic-acid carbazole molecule EtCz3EPA, which provides stronger anchoring at the buried interface, is necessary to preserve device performance under real sunlight. This demonstrates that although PTAA can perform well under ISOS protocols, including ISOS-LC or indoor temperatures even higher than those experienced outdoors in the study by Fei *et al.*, while humidity-related effects are effectively suppressed through encapsulation, factors unique to outdoor environments, such as UV exposure, can activate degradation pathways not triggered indoors. This makes it necessary to adjust or replace the HTL with another material that preserves efficient hole transport while avoiding UV-induced instability. Taken together, this case shows how indoor robustness does not guarantee outdoor viability when specific spectral stressors are missing from indoor protocols.

Similarly, a widely used indoor passivation strategy involves the incorporation of PEAI to form a 3D/2D heterojunction, which effectively suppresses surface recombination under mild ISOS conditions. However, the work by Wang *et al.* shows that PEAI is not transferable to outdoor-like stressors because its mono-ammonium nature makes it highly susceptible to photo-thermal degradation.²⁸¹ Under simultaneous light and heat, PEAI deprotonates and reacts with PbI_2 , producing Pb^0 and triggering rapid iodide migration, processes that do not occur under standard indoor protocols. To overcome this limitation, the authors replace PEAI with the di-ammonium spacer BDAI₂, which forms the photothermally robust Dion–Jacobson phase BDAPbI₄. The strong bidentate hydrogen bonding of the di-ammonium cation stabilizes the PbI_6 network, suppresses deprotonation, and prevents structural collapse under harsh ageing. This finding underscores that certain molecules performing excellently indoors may still require structural redesign to withstand full-spectrum outdoor stressors.

In the context of carbon electrodes, Lukas *et al.* provide another example of an indoor strategy that cannot be directly transferred outdoors.²⁸² Although PCBM is widely used as the electron-extraction layer in indoor PIN devices, the authors show that it is incompatible with carbon pastes, since it dissolves and intermixes during contact formation. This prevents the fabrication of stable or reproducible devices, and previous work showed that only an evaporated chromium layer could protect PCBM, which increases cost and defeats the purpose of low-cost carbon contacts.²⁸³ To create a fully low-cost PIN/carbon architecture, the authors replace PCBM with ALD-SnO₂. However, SnO₂ alone gives limited performance and

poor stability because its rigid inorganic surface forms a weak electronic interface with the carbon electrode, leading to contact resistance increases and interfacial degradation. Adding a thin PH1000 layer on top of SnO₂ overcomes these limitations by improving energy alignment, charge transport, and interface quality. As a result, the SnO₂/PH1000 and carbon configuration shows significantly improved stability and functionality, demonstrating a viable pathway toward low-cost PIN devices with carbon top electrodes. This illustrates a different type of indoor–outdoor mismatch rooted not in photochemistry but in interface compatibility and mechanical robustness.

Furthermore, encapsulation provides another example of a strategy that performs well under indoor ISOS tests but fails outdoors. Emery *et al.* show that UV-glue “LAB” encapsulation, commonly used in perovskite research, preserves more than 93% of the initial efficiency for over 1500 h under ISOS-D1 storage, giving the impression of strong protection.²⁸⁴ However, under damp-heat conditions (85 °C/85% RH) or real outdoor exposure, LAB-encapsulated devices degrade rapidly, often failing within days. This poor transferability arises from the inherent limitations of UV-cured epoxies, including micro-cracking, high WVTR, and interfacial damage introduced during curing. Outdoors, these weaknesses permit moisture ingress and edge-initiated decomposition, effects not revealed by indoor tests. To address this issue, the authors implemented a commercial lamination stack consisting of a polyolefin encapsulant and a butyl rubber edge seal. This provides low permeation rates and higher mechanical stability. Devices encapsulated using this method pass IEC 61215 damp-heat testing and retain almost full efficiency after spending more than ten months outdoors. This demonstrates that, like interfaces and contact layers, encapsulation must be validated directly under outdoor conditions to ensure real-world durability.

Collectively, these examples show different strategies for improving device stability and highlight the gap between good development in indoor tests and outdoor tests, spanning from interface modifications to back electrode design and encapsulation. This message makes clear that the strategies at the edge of the state of the art tested only indoors should be tested outdoors to analyze their transferability and ensure adequate development or the needed optimizations. Therefore, we propose testing some recent strategies in outdoor conditions that work very well indoors and could have promising behavior outdoors. Two high-efficiency interface modulation strategies recently reported under indoor ISOS testing highlight the need for outdoor validation. Liu *et al.* introduce a bimolecular treatment combining PDAI₂, which induces a field-effect at the perovskite/C₆₀ interface to block holes, with 3MTPAI, which passivates iodide vacancies through S–Pb interactions;²⁷⁴ this synergistic design suppresses interfacial recombination and enables certified PCEs above 25% with >1500 h indoor stability (>T₉₀). Chen *et al.* develop dual-site aromatic sulfonate ligands such as 4Cl-BZS that bind two adjacent Pb²⁺ sites and improve the energetic and mechanical coupling with C₆₀, producing >26% certified efficiencies and strong stability



under ISOS-L3 protocols during 1200 h (T_{95}).²⁷² Both treatments rely on ultrathin molecular layers that are solution-processable and compatible with scalable coating, but neither was evaluated under sunlight containing UV, humidity variations, or thermal cycling. Their excellent indoor performance therefore makes them compelling candidates for outdoor testing to assess the robustness and transferability of these interface-engineering concepts under real-world conditions.

In addition to interface engineering, several recent indoor-only stability advances based on molecular and polymer additives for bulk modification highlight promising yet unvalidated pathways for real-world operation. Sun *et al.* introduce a step-wise melting-polymerizing molecule (SMPM) that melts, infiltrates grain boundaries, and polymerizes *in situ*, forming a hydrophobic grain-scale encapsulation that strongly suppresses humidity-driven degradation (T_{95} at 2000 h at 85% RH) and stabilizes FAPbI₃ through hydrogen bonding and Pb²⁺ coordination.²⁸⁵ Simultaneously, Shi *et al.* develop mechanically interlocked networks (MINs) that regulate colloidal particle distribution during printing, yielding highly uniform large-area films with enhanced crystallinity and mechanical robustness, an approach fundamentally compatible with scalable coating methods such as blade or slot-die processing ($>T_{90}$ after 1000 h at 65 °C).²⁸⁶ Finally, Xiong *et al.* propose an additive strategy using alkali metal oxalates to homogenize Cl distribution, suppress surface aggregation, and improve interfacial coupling, enabling $>27\%$ indoor PCE and strong thermally driven MPPT stability ($>T_{80}$ after 1000 h at 85 °C).²⁸⁷ None of these three approaches have been tested in outdoor conditions with fluctuations in humidity, temperature, or light with UV components. However, all are based on ultra-thin molecular layers or solution-processable additives that are in principle compatible with large-scale manufacturing techniques. These strategies are therefore excellent candidates for systematic outdoor validation to determine their actual transferability to real-world conditions due to their outstanding indoor performance.

Ultimately, these examples show that potentially viable strategies for outdoor performance have not yet been evaluated under real operating conditions, and they corroborate whether they are truly viable or if some adjustment or optimization of the molecule or strategy is necessary. These are directly applicable improvements for devices exposed to actual outdoor conditions, and they could accelerate the stability of solar cell devices and bring their commercialization closer.

3.1.5. HTL-Free carbon devices. Analyzing the effect of different strategies on device stability requires a layer by layer investigation of how each component contributes to the degradation of the device's PCE. Any layer that degrades contributes to aggregate performance loss. The layers with the largest impact on stability are the light absorber material, the charge transport layers, and the counter electrode. Section 2 covered the degradation of the light absorber material, and Section 3.1.3 examined the counter electrode. There, it was determined that carbon-based counter electrodes exhibit enhanced long-term stability in outdoor environments and offer a scalable, low

temperature processable substitute for noble metal contacts. Nevertheless, in PSCs, the progressive reduction in PCE is also due to the degradation of the other layers of the device, especially the HTLs.

HTLs used in n-i-p devices (such as Spiro-OMeTAD and PTAA) show thermal and photo-induced degradation in the operational range ($\approx 70\text{--}85$ °C).²⁸⁸ In doped Spiro-OMeTAD, device performance drops rapidly at ≈ 85 °C while other layers remain comparatively stable; part of this decay is irreversible due to dopant loss. In particular, tBP evaporates at ≈ 85 °C, causing voids, pinhole formation, and morphological failure,^{289,290} effects that can be partially mitigated by the top metal electrode.^{288,291} Also, under humid conditions, the hygroscopic Li-TFSI dopant accelerates underlying perovskite degradation; higher Li-TFSI content correlates with faster moisture-driven decay, and at high RH Spiro-OMeTAD or PTAA films crack and fragment on the perovskite, enabling water and ion ingress.^{119,288,292} Meanwhile, Au electrode migration through Spiro-OMeTAD occurs under illumination even at 70 °C, causing shunts and trap formation.^{288,293} For PTAA, doping with Li-TFSI + tBP increases conductivity but also introduces light-assisted oxidation features (*e.g.*, a new absorption band at 440–550 nm assigned to PTAA⁺ under illumination), while shifting the energy levels and affecting morphology; optimized films still benefit from thin, pinhole-free coverage to minimize transport losses.²⁹⁴ A remarkable exception of this degradation pathways is the work by Babu *et al.*, who reported a PTAA based carbon electrode perovskite module with operational stability lasting one year.²⁶¹ Maybe, this result was made possible using a highly durable encapsulant that mitigated moisture and HTL degradation effects. Nevertheless, the requirement for multi-layer encapsulation and the high cost of the HTL material also resulted in extra expenses.

Although the device is protected from moisture and light by carbon electrodes, organic HTLs still degrade significantly, and the most stable carbon based devices reported in the literature are typically HTL-free. This is due to the intrinsic stability issues associated with organic HTLs. Passatorntaschakorn *et al.* fabricated HTL free, carbon-electrode PSCs that achieved a maximum PCE of 11.09% and retained 73% of their initial efficiency after about 20232 h under ambient ISOS-D1 (~ 25 °C, 35–45% RH).²⁹⁵ In comparison, Spiro-OMeTAD based-devices with 13.95% of initial PCE retained only 40% PCE over the same period in the same test.

To overcome the hole extraction barrier associated with HTL free devices, Lin *et al.* developed a 1D/3D heterojunction by incorporating thiocholine iodide derived TcHPbI₃ at the surface of CsPbI₃.²⁹⁶ Their optimized devices achieved a certified PCE of 18.7% and retained over 90% of their initial efficiency after 300 hours of continuous maximum power point tracking (1 sun exposure) under inert conditions. This strategy also shows potential for outdoor deployment. Selvakumar Pitchaiya *et al.* demonstrated that HTL-free, carbon electrode PSCs retained more than 80% of their initial PCE after 50 days (approximately 1200 hours) of outdoor exposure.²⁹⁷ Likewise, Siram *et al.* reported that fully inorganic CsPbBr₃ unencapsulated devices



using a carbon nanotubes (CNT)/perylene diimide (PDI) back contact, without any HTL, lost less than 30% of their initial performance after approximately 6 weeks of outdoor testing in desert weather conditions.²⁶³

Despite their excellent stability, HTL free devices usually have lower efficiency because of less hole extraction. Alternatively, inorganic HTLs like CuSCN and NiO_x can be used. Unlike Spiro-OMeTAD and PTAA, which degrade at about 85 °C, these materials can withstand thermal and photo-oxidative degradation up to 100 °C.^{298–301} Long-term stability is enhanced by their nonhygroscopic nature, which inhibits ion migration and moisture-induced film swelling.^{263,302,303} Zhang *et al.* carried out a large-scale study, which included over 7000 PSCs, demonstrating that simply swapping out organic HTLs for carbon electrodes increased normalized device lifetime by 4.2 times.¹⁰⁵ Stability was increased around 7.2 times by adding an inorganic HTL below the carbon contact. When compared to organic HTLs, inorganic HTLs alone (without carbon electrode) provided a 1.8-fold increase in T_{S80m} values, indicating that they may be able to strike a balance between stability and hole extraction. Wu *et al.* demonstrated that a CuSCN and carbon nanotube bilayer improved the open-circuit-voltage from 0.85 V to 1.01 V and fill factor from 0.60 to 0.73, resulting in a champion PCE of 17.58%. These devices maintained approximately 80% of their initial performance after 1000 hours of continuous one-sun illumination under an inert atmosphere.³⁰⁴

Unlike organic HTLs, inorganic HTLs such as NiO_x and CuSCN usually offer higher stability but often suffer from incomplete coverage, high surface roughness or low hole mobility, even with careful synthesis and interface engineering.^{305–307} These morphological defects are conducive to interfacial recombination, reducing the PCE compared to organic HTLs, which spontaneously form a thin, smooth, high-hole mobility film. For instance, mesoporous or nanoparticle-based NiO_x films require optimization,³⁰⁸ particularly in terms of thickness, doping, and nanoparticle dispersion, to minimize voids and achieve the improved efficiencies reported for electrochemically deposited NiO_x and dopant or additives modified NiO_x.^{309,310} Likewise, CuSCN-based devices typically achieve efficiencies of ~15–16% PCE but with interface passivation (*i.e.*, (3-Mercaptopropyl)trimethoxysilane (MPTMS)), improved efficiencies can surpass 20%.^{311,312} Thus, while inorganic HTLs can be more stable, the quality of the organic HTLs in as-fabricated films is better, unless further engineering on the inorganic HTLs is undertaken.

Despite the mentioned advances in HTLs, eliminating the HTL layer offers several advantages. It enables ambient, low-temperature processing, removes the need for controlled-atmosphere HTL deposition and dopant incorporation, and avoids high-temperature annealing steps. Passatorntaschakorn *et al.* reported that this simplification improved reproducibility and reduced module material costs by approximately 97% compared to HTL based workflows.²⁹⁵ Carbon paste electrodes cost only \$0.01 per gram, about 1.4% the price of silver, leading to a 69.7% reduction in perovskite module material costs compared to noble metal based devices.³¹³ Projections from techno-

economic modeling indicate that HTL-free, carbon electrode PSCs could achieve manufacturing costs as low as \$22.70 per m² in large-scale production, highlighting their potential for cost effective, high throughput commercial production.³¹⁴

3.2. Influence of device architecture on stability

The mainstream device architecture used in the field of perovskite solar cells are n–i–p and p–i–n, which differ in the order of stacking the layers: in the former, sunlight first goes through the ETL before reaching the perovskite absorber while in the latter it first goes through the HTL. In the n–i–p architecture, instability issues related to the use of Spiro-OMeTAD have been frequently mentioned, although big-data driven investigation have shown that in general, and not being limited to outdoor stability results, n–i–p devices have shown slightly higher overall stability.¹⁰⁵ In this sense, it is interesting to compare these tendencies exclusively in the outdoor stability landscape. Additionally, the mesoscopic carbon architecture is considered, due to its considerably different approach to fabrication: the transport layers and/or the carbon back electrode are deposited first followed by an infiltration of the perovskite absorber. This architecture is particularly remarkable since the earliest results of relatively high outdoor stability were attained following this configuration.^{44,266}

Fig. 4c shows a box plot comparing the outdoor stability results of the mentioned architectures across all the papers reviewed in this work which explicitly showed outdoor evaluation, in which the insets correspond to the average stability time. It is important to note that more papers were considered here than the ones previously shown in this section, since the discussion was limited to works in which the fabrication strategy was the focus for a richer overview of such strategies. The distinction between T_{70-T85} and T_{90} or higher is made to facilitate the comparison between the results, as the reported values were not standardized to a benchmark parameter. A total of 50 reports were considered, of which 27 (54%), 18 (36%) and 5 (10%) employ the n–i–p, p–i–n and mesoscopic carbon architectures, respectively. From those works using the n–i–p architecture, 20 (74.07%) reported a T_{70-T85} while 7 (25.93%) reported a T_{90} or higher. For the p–i–n and mesoscopic carbon architectures these numbers correspond to 11 (61.11%) and 7 (38.89%), and 2 (40%) and 3 (60%), respectively. In agreement with the general tendencies in perovskite photovoltaic research, most outdoor stability results correspond to n–i–p devices, followed by p–i–n and mesoscopic carbon. Except for the mesoscopic carbon architecture, in most works an outdoor stability corresponding to T_{70-T85} has been reported, although in recent years improvements in stability have led to a higher ratio of reports of T_{90} or higher, *e.g.*, in Fig. 4b most of the reported stability times in 2023 and 2025 are T_{90} or higher. In terms of T_{70-T85} , a higher average of 7972.5 h has been achieved in p–i–n devices, followed by n–i–p (1106.1 h) and mesoscopic carbon (875 h). In the case of T_{90} or higher, the highest average of 2553.14 h was achieved with n–i–p architecture, followed by p–i–n (2041.71 h) and mesoscopic carbon (998 h). Nonetheless, the discrepancy in number of reports and



time of evaluation makes the comparison of averages elusive and instead looking at the best result from each architecture shines light onto their viability for stable real-world operation. In terms of T_{70} – T_{85} , comparable outdoor stability times of 8400 h²⁶¹ and 35040 h⁴⁶ have been achieved in n–i–p and p–i–n architectures, respectively, while mesoscopic carbon lags with a maximum value of 1200 h. As for T_{90} or higher, the p–i–n architectures comes on top with a maximum outdoor stability time of 4870 h,⁹³ followed by n–i–p with 4032 h¹⁸¹ and mesoscopic carbon with 2250 h.²⁶⁴ In general, this hints to higher attained outdoor stability for the p–i–n architecture, although in general this shows a promising landscape of durable devices regardless of architecture.

Fig. 4d shows an overview of the retained PCE as a function of the stability time of all the reviewed reports that explicitly showed outdoor evaluation, in which the bulk of results are concentrated in times below 4000 h and those displaying the longest time for each architecture (highlighted as a star) follow the same trend as the T_{90} or higher discussed above: p–i–n > n–i–p > mesoscopic carbon. Furthermore, for both n–i–p and p–i–n architectures, the longest outdoor stability periods without any performance loss were achieved in 2023 and 2024, further corroborating the recent trend. Interestingly, only 10 of the reports considered here (shown as hollow data points in Fig. 4d) are found as entries in the Perovskite Database, pointing to a lack of willingness in the community to update this valuable resource. Having centralized access to up-to-date outdoor stability data is key to keeping track of progress in the field and possibly facilitate collaboration between different research groups in the world, and thus it is highly encouraged for following investigations to update the database.

4. Additional strategies for stable device operation under real world operating conditions

When discussing the stable operation of both, PSCs and perovskite solar modules (PSMs) under real-world conditions, it refers to maintaining the device's stability while operating under illumination, with a working load/bias, and within the specific environmental conditions of its deployment.⁶⁰ As highlighted in previous sections, various strategies exist to enhance the intrinsic stability of PSCs. However, other processes are essential to further improve device performance and durability. Both small-scale lab cells and large-area PSMs are vulnerable to harsh outdoor conditions without proper encapsulation, a critical requirement for any commercial technology.^{94,315,316} Moreover, several strategies can be employed to optimize device operation, as indicated in Fig. 5, and discussed below.

4.1. Device interconnection and design

The first strategy includes device interconnection and design, which is crucial for controlling current and voltage output of large-area devices,^{315,317} which can be done by modulating series and parallel connection modes. Gao *et al.*³¹⁸

demonstrated that, for four PSCs with an active area of 6.12 cm², the most effective way to maximize output power in PSM was to first connect two pairs of cells in series and then interconnecting them in parallel. This type of design is especially relevant for scaling up, as electrical interconnection becomes mandatory in large-area modules. In this context, the PSM interconnection design is based on three steps known as P1, P2 and P3, which are essential to ensure efficient and stable operation; these steps are typically carried out by laser ablation.^{319,320} P1 consists of the removal of the transparent conducting oxide (TCO), such as indium tin oxide (ITO) or fluorine-doped tin oxide (FTO), directly on the substrate, in order to electrically isolate the subcells and avoid lateral short circuits. P2 is the most critical process, as it allows the electrical series connection between subcells by carefully removing the top layers such as HTL, PVKT and ETL without damaging the TCO, that is, this connection allows current to flow from the back contact of one cell to the TCO of the next, achieving a series interconnection that sums the voltage of each subcell. By removing the top layers of the stack, the TCO is exposed, allowing the back contact of the adjacent cell to be deposited directly on it, forming an ohmic connection that ensures efficient charge transfer. However, poor execution of P2 can cause damage to the TCO or incomplete contacts, increasing the series resistance or inducing the appearance of short circuits. Finally, P3 is responsible for isolating the subcells present in the device.^{319–322}

Now, under real-world operating conditions, one of the main limitations faced by PSMs is the phenomenon of reverse bias.³²³ This occurs when one or more series-connected cells are partially or fully shaded, generating an inverse voltage that intensifies ionic migration processes. That is, when a configuration is in series, all cells share the same current and when one of them is shaded, the others force the current through it, generating a reverse potential. This charge redistribution generates an electric field that favors the movement of mobile ions such as I[−], MA⁺, or FA⁺ from the bulk to the interfaces, promoting undesirable reactions, vacancy formation, tunneling phenomena, and delamination, ultimately leading to partial or irreversible degradation of the device.³²⁴ With a short polarization duration, reverse polarization degradation is partially recoverable through the redistribution of migrated ions. When polarization is prolonged or repeated, the degradation gradually becomes irreversible, involving I[−] oxidation by injected holes.⁵² In summary, the combination of localized charges, reverse voltages, and accelerated ionic transport in series/parallel configurations under realistic conditions can trigger chemical degradation processes both at the interfaces and in the active volume of the cell.¹⁶² This phenomenon is frequently observed in series-connected configurations, especially in outdoor-tested minimodules, where variable environmental conditions such as shading, dust, or intrinsic defects cause cells to operate asynchronously, resulting in hot-spot formation, localized overheating, and energy loss.^{237,325–328} A representative example of this configuration has been reported in large area perovskite solar panels (PSPs)



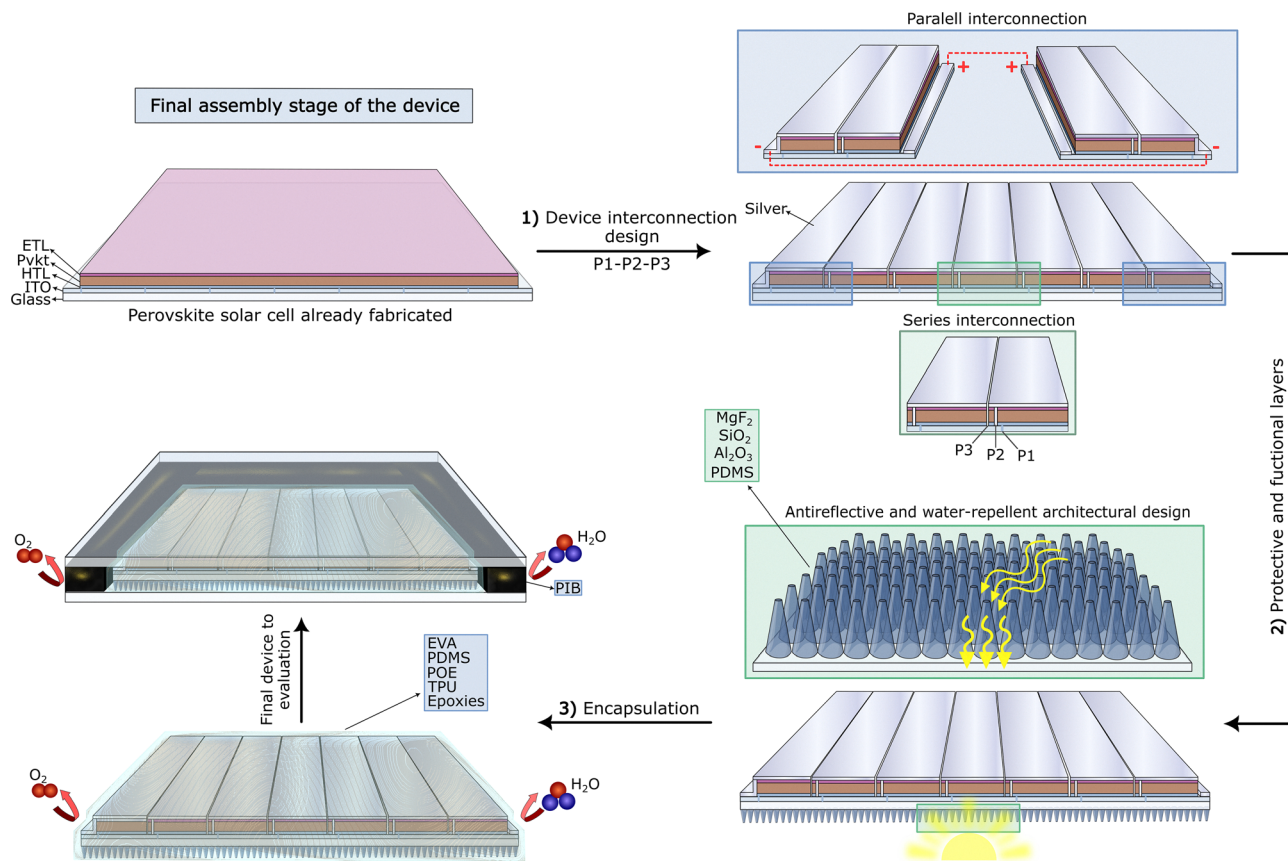


Fig. 5 Schematic process line illustrating other strategies to enhance the long-term stability and performance of PSCs under real-world operating conditions. These strategies include: (1) optimized device interconnection designs, such as parallel or series configurations enabled by P1–P2–P3 laser patterning and contacts (e.g., Silver); (2) functional surface modifications, including self-cleaning and antireflective coatings composed of materials (e.g., MgF₂, SnO₂, Al₂O₃, or PDMS) that also improve light harvesting; and (3) encapsulation using barrier materials (e.g., PIB, EVA, PDMS, POE, TPU, or epoxy resins) to protect the device from moisture (H₂O) and oxygen (O₂) access.

with an area of 0.5 m², composed of minimodules with 10 series-connected cells, arranged in 5 parallel strings, each consisting of 8 modules in series, totaling 40 PSMs per panel.³²⁵

To address these challenges, various mitigation strategies have been proposed, such as the use of ion-blocking layers like doped NiO_x, Al₂O₃, or self-assembled monolayers (SAMs), which limit ionic movement toward sensitive interfaces,^{329–332} reducing mobile ionic defects through compositional engineering; and incorporating bypass diodes to protect shaded cells. However, the implementation of bypass diodes can increase system costs, particularly in PSCs, which have lower breakdown voltages compared to conventional photovoltaic technologies. In this context, carbon-based PSMs have shown promise as a robust solution. Recent studies report that these modules can withstand reverse voltages of up to –9 V, passing the hot-spot test defined by the IEC 61215-2: 2016 standard without efficiency loss, highlighting their potential for industrialization and suitability for large-scale applications.³³³ Table 2 below summarizes the main structural and design strategies employed to mitigate ionic migration induced by reverse bias.

4.2. Protective and functional layers

To maximize power generation, it is important both to maintain stable performance and to have the maximum PCE possible. As previously discussed, it can be achieved by different strategies for the cell stack, but it can also be done after the PSM is fabricated. Specifically, it is possible to incorporate protective and functional layers such as anti-reflective,^{335–338} self-cleaning^{338,339} and UV-absorbing coatings.^{340,341} For instance, super-hydrophilic antireflection (AR) films coated on photovoltaic glass have been demonstrated to significantly improve solar transmittance across broadband wavelengths (380–1100 nm), thereby boosting efficiency.³⁴² However, while AR films enhance optical performance,^{335,337} they do not inherently improve device stability.^{341,343–345} Currently, anti-reflective films have been applied to 90% of commercial solar panels.^{346,347} Commonly used materials to fabricate antireflection films include magnesium fluoride (MgF₂),^{348,349} zinc oxide (ZnO),^{350,351} silicon oxide (SiO₂),^{352–355} titanium oxide (TiO₂),^{356,357} aluminum oxide (Al₂O₃)³⁵⁸ and polydimethylsiloxane (PDMS),³³⁷ *etc.* Among these, the most widely used is SiO₂, due to its low price and good chemical stability. It is often used for the preparation of anti-reflection films



Table 2 Different strategies employed to mitigate reverse polarization-induced ion migration

Configuration/modification	Blocking material	Evaluated conditions	Results	Ref. (DOI)
PSCs with series inter-connection and bypass diode	Bypass diode + carbon	−9 V, 1 week (IEC 61215–2)	Without loss of efficiency	333
PSCs with doped NiO _x layer	NiO _x (e.g. Cu:NiO _x , Ag:NiO _x)	Reverse bias, thermal stress	Reduction of ionic accumulation, enhanced thermal stability	331
PSCs with SAM on SnO ₂	Self-assembled monolayer	Leakage current and <i>J</i> – <i>V</i> measurement	Partial suppression of ion migration	334
PSCs with ultrathin Al ₂ O ₃ ALD	Al ₂ O ₃	Thermal + electrical stress (MPPT, 85 °C, 85%)	Hindered I [−] and MA ⁺ migration, improved interfacial stability	332
PSCs with secure inter-connection and appropriate encapsulation	Encapsulated + POE + PIB	Outdoor test, simulated partial shade	Sustained stability for 10 months	284

by a simple and controllable sol–gel method. In this same context, to further improve optical efficiency, researchers have explored the use of nanostructures, such as nanocones^{338,359} made from the materials like MgF₂, ZnO, SiO₂, TiO₂, Al₂O₃ or PDMS, which represent a promising solution for reducing reflection losses. These structures allow the generation of a refractive index (RI) gradient at the air–solid interface, effectively reducing surface reflection.^{359,360}

Looking ahead, future challenges of these protective and functional layers will involve optimizing integration during lab-scale production, while ensuring cost-effectiveness. The development of new materials or hybrid coatings with multifunctional properties, such as combining anti-reflective, self-cleaning, and UV-absorbing capabilities, could provide more holistic solutions. Overcoming these challenges will be crucial for the continued advancement of PSM and its successful integration into large-scale renewable energy systems.

4.3. Encapsulation

Encapsulation is essential for protecting PSCs from environmental stressors such as moisture, oxygen, and UV radiation. Materials used for encapsulation must meet several criteria: chemical inertness, high light transmittance (>90%), low water vapor transmission rate (WVTR) between 10^{−3} and 10^{−6} gm^{−2} day, low oxygen transmission rate (OTR) between 10^{−3} and 10^{−5} g m^{−2} day, high electrical resistivity, and mechanical durability.^{31,95,172,284,361,362} In addition, functional carbon electrodes or carbon-based layers can serve not only as conductive elements but also as protective barriers or encapsulants, resisting moisture and oxygen ingress and thus enhancing device stability.^{363–365} These carbon layers can reduce the influence of the external factors and degradation pathways, further supporting the long-term operation of perovskite solar cells in real-world conditions. On the other hand, traditional encapsulants like ethylene vinyl acetate (EVA) for glass-to-glass encapsulation, commonly used in silicon solar cells, require lamination temperatures around 150 °C and are prone to optical degradation, delamination and acetic acid formation, making them unsuitable for PSCs.^{94,361,366} Therefore, alternatives to EVA exist, such as ionomers,^{42,367} polyurethane²⁶⁴ or polyolefins (POE).^{94,264} Even though EVA has been used for encapsulation

of PSM and panels,^{29,368} the reduction of the processing temperature to values around 85 °C does not allow proper cross linking, leading to premature moisture and oxygen interaction with the devices.³²⁵ Also, for rigid devices, glass-to-glass encapsulation with UV-curable polymers has proven effective. Specifically epoxies, widely used in the industry, exhibit good moisture resistance and studies have demonstrated the stability of UV-epoxy-cured devices in the range of 500 to 2000 h of outdoor exposure.^{237,369} However, it has been determined that the exothermic curing process of UV-curable epoxy resins can damage cells due to their mechanical rigidity and thermal stress,³⁶¹ they are also prone to aging and yellowing³⁷⁰ and they may be also susceptible to delamination at elevated temperatures.³³ To mitigate these effects, polyvinylpyrrolidone (PVP) interlayers have been implemented.³¹⁵ For the case of flexible devices, using polymeric substrates such as polyethylene terephthalate (PET), polyethylene naphthalate (PEN) and polyimide (PI), the encapsulation materials require high ductility and compatibility with roll-to-roll fabrication processes. Thermal load during encapsulation must also be carefully managed to avoid damaging organic substrates.³⁶¹ Promising alternatives include solution-based poly(methyl methacrylate)/styrene-butadiene (PMMA/SB) bilayer structures, which are cost-effective and scalable for both rigid and flexible devices.²⁶

Other materials, such as thermally evaporated MgF₂ or MoO_{3−x} capping layers, combined with glass and UV-curable polymers, have shown excellent encapsulation performance.³⁷¹ Recent advancements in encapsulation materials include the use of thermoplastic polyurethanes (TPUs), which stand out for their high thermal stability, scalability, and flexibility.³⁷² In addition, TPUs have good chemical inertia against environmental agents such as moisture, oxygen and UV radiation, preventing the degradation of perovskites to PbI₂ and thus improving device stability.³⁷⁰ For example, polyether based TPUs are preferred for moisture exposure applications, since they offer greater hydrolysis resistance. However, at high temperatures, these materials may suffer oxidation, which causes yellowing and loss of mechanical properties. Similarly, prolonged exposure to UV radiation degrades the material and



hinders encapsulation.³⁷⁰ TPUs are formed by cross-linking polyether polyols (e.g., polypropylene glycol (PPG) and polyethylene glycol (PEG)) with isocyanates (e.g., isophorone diisocyanate (IPDI) and toluene diisocyanate (TDI)). These materials are highly tunable, making them suitable for diverse applications, including PSCs encapsulation.³⁷³ Butyl rubber-based polymers, such as polyisobutylene (PIB) has been validated as an effective material for both primary encapsulation and edge sealing because of its efficient moisture barrier properties³³ achieved by its low water permeability and stability in accelerated aging tests.³³ In this context, edge sealants play a crucial role in preventing delamination and cracking at the encapsulant edges, ensuring the long-term structural integrity of the device. The use of PIB as edge sealant with desiccant materials to prevent moisture and oxygen ingress, combined with polyolefins (POE) films for mechanical stability has demonstrated to be a good strategy, inspired by commercial CIGS module encapsulation, as it has demonstrated excellent outdoor stability (ISOS-O3) when compared to typical encapsulation using UV curable epoxies, as they were able remaining stable for over 10 months.²⁸⁴ This effectiveness is largely due to the careful selection of materials, where the selection of the edge sealant is due to its low water vapor transmission rate (WVTR) and low glass transition temperature (T_g),⁹⁴ making it compatible with mechanical stresses induced by extreme weather conditions.³⁷⁴ Additionally, POE films have excellent adhesion, water resistance and Young's modulus of 8 MPa (mechanical property that measures the stiffness of a material, that is, its ability to resist elastic deformation under an applied load), contributing to mechanical stability.⁴² However, their application requires temperatures above 80 °C, which could compromise the structural integrity of the device.³¹⁵ Furthermore, POE's high volumetric resistivity ($\geq 10^{17} \Omega \text{ cm}$ at 23 °C) minimizes the risk of potential-induced degradation, thus prolonging the device lifetime.³⁶¹ In general, materials such as polyisobutylene (PIB) and polyolefins (POE) not only offer a high barrier to the permeation of water and oxygen vapor, but also have chemical inertia, which allows them to avoid reactions with volatile species such as HI or MAI, which can be released from bulk in conditions of thermal stress. This inertia makes them excellent candidates for sealing edges and encapsulant. In addition, their high volumetric resistivity and low vitreous transition temperature allows them to mechanically adapt to real environmental conditions without compromising structural integrity.

Other materials have been extensively studied and employed in the encapsulation of PSCs, standing out for their effectiveness as moisture barriers, compatibility with devices, and ease of processing. Among them, polydimethylsiloxane (PDMS) has demonstrated high hydrophobicity and flexibility, contributing to extending PSCs stability up to 3000 hours.³¹⁵ In general terms, hydrophobic materials such as PDMS and TPU are advantageous in applications where environmental exposure to moisture is critical. Its repellent water surfaces limit the adsorption and dissemination of H_2O , which reduces the probability of decomposition reactions driven by hydrolysis.

These properties, combined with their mechanical adaptability, make them suitable candidates for flexible substrates in outdoor environments. For applications requiring superior thermal and chemical stability, silicones and glass frits have been used as alternative encapsulation solutions.³¹⁶ Overall, the combination of encapsulants such as POE or PU with edge sealants like PIB or epoxies has proven to be an effective strategy to enhance the durability and reliability of PSCs, providing increased protection against environmental factors and ensuring long-term stability.³³ In fact, recent studies have reported operating times exceeding one year under real-world conditions, using POE as the encapsulant and PIB as the edge sealant, confirming the technical feasibility of this combination for long-lasting photovoltaic applications.³⁷⁵

The encapsulation process itself is critical to ensure their stability and durability under real operating conditions, as can be done with and without vacuum assistance. It is commonly carried out in a vacuum laminator, which removes the air and prevents the formation of air bubbles inside the device. This approach is compatible with industrial processes and allows for homogeneous fusion and curing of both the barrier films, *i.e.* POE, PIB and ionomers,^{94,284,376} ensuring uniform and effective protection barrier against moisture and oxygen.³⁷⁷ As POE films are polymers that require a high-temperature cross-linking process, depending on the vacuum used, they can be incompatible with the thermal sensitivity of MAPI-based PSCs. In contrast, ionomers can be processed at lower temperatures, as their cross-links are formed through ionic interactions.³⁷⁸ However, these ionomers present additional challenges, such as poor adhesion to glass surfaces. To reduce water permeability and improve adhesion in the encapsulation, the use of the previously mentioned edge sealants based on PIB filled with edge desiccant is recommended,¹²¹ which offer a low water vapor transmission rate (WVTR) and a low glass transition temperature.³⁷⁴ PIB can be processed at a lower temperature than ionomers or POE. However, it has been observed that configurations combining POE with edge sealants can suffer from delamination under thermal cycling, highlighting the importance of optimizing processing conditions.³⁷⁷ In conclusion, the selection of the encapsulant should consider a combination of moisture and oxygen barrier, mechanical strength, and thermal compatibility, avoiding processing temperatures that may compromise the integrity of the perovskite material and is still under continuous investigations. To provide a clearer comparison of the encapsulation strategies discussed, Table 3 summarizes key materials used for both edge sealing and primary encapsulation in perovskite solar cells. It highlights their main advantages, limitations, type of solar cell substrates tested, and suitability for real-world operating conditions.

In conclusion, encapsulation remains one of the most critical steps to achieve long-term stability of PSCs outdoors. Among the strategies evaluated, the combination of polyolefins (POE) as the primary encapsulant and polyisobutylene (PIB) as the edge sealant has proven to be one of the most efficient solutions. This is attributed to its excellent moisture barrier



properties, mechanical flexibility, and chemical stability, which together help mitigate environmental degradation mechanisms such as delamination, oxygen ingress, and moisture diffusion. However, challenges such as high processing temperatures (in the case of POE) and poor adhesion of certain ionomers to glass remain obstacles to its large-scale implementation. Future efforts should focus on improving material compatibility, reducing curing temperatures, and developing industrial-scale encapsulation techniques (e.g., vacuum lamination) that preserve device integrity.

Due to the absence of standardized test conditions, comparing the performance of the various encapsulation materials reported in literature remains a challenge. To address this, it is essential to establish unified evaluation protocols that reflect real-world (outdoor) operating environments and consider key variables such as device configuration, climate exposure, and mechanical stress. Advancing in this direction will not only enable more meaningful comparisons but also accelerate the development of durable and commercially viable PSCs. In this context, the Photovoltaic Accelerator for Commercializing Technologies (PACT) at Sandia National Laboratories³⁷⁹ plays a pivotal role by conducting outdoor field testing, validating performance metrics, and promoting the creation of test standards aimed at accurately assessing field behavior and predicting long-term degradation. In the following section, we review the main advances in characterization of PSCs under real-world operating conditions.

5. Comprehensive characterization of PSCs under real-world operating conditions

5.1. Standard protocols and key parameters for outdoor testing

Device performance can be evaluated in a controlled laboratory (indoor) or under natural sunlight (outdoors) conditions, allowing for direct comparisons of device behavior under specific conditions like Standard Test Conditions (STC, irradiance of 1000 W m^{-2} and cell temperature of $25 \text{ }^\circ\text{C}$). However, to better assess the impact of real-world environmental factors on performance, additional power rating conditions (PRCs) are recommended (IEC 61215-1-1, 2016; IEC 61215-2, 2016; IEC 61853-1, 2011). These include Nominal Operating Cell Temperature (NOCT: 800 W m^{-2} , $20 \text{ }^\circ\text{C}$ ambient temperature), Low-Irradiance Condition (LIC: 200 W m^{-2} , $25 \text{ }^\circ\text{C}$ cell temperature), High-Temperature Condition (HTC: 1000 W m^{-2} , $75 \text{ }^\circ\text{C}$ cell temperature), and Low-Temperature Condition (LTC: 500 W m^{-2} , $15 \text{ }^\circ\text{C}$ cell temperature). Additionally, thermal performance tests are conducted beyond electrical measurements to comprehensively characterize photovoltaic devices, including solar cells, sub-assembly of cells, and PV modules. Therefore, the instrumentation, infrastructure, test duration, and estimation procedures may vary significantly between indoor and outdoor testing settings.

For indoor testing, the primary light source is the solar simulator, allowing control over the irradiance level. In contrast, the device's temperature can be externally controlled to reach the target temperature. As a result, a complete characterization is carried out, including performance and temperature coefficients.³⁸¹ On the other hand, achieving performance reproducibility in natural sunlight is challenging. First, there is no control over the irradiance level; it varies depending on the presence of clouds, haze, smoke, and seasonal changes. Second, keeping consistent operating conditions as PRCs to ensure similar temperature and irradiance levels during exposure is difficult. Third, depending on the location, different operating conditions may arise throughout the year due to seasonal variations.³⁸² Thus, the PRCs cannot be measured during exposure or may not be the most representative conditions concerning location. This leads to uncertainty in the exposure duration because there is no solid knowledge about the time necessary to measure data that is considered statistically reliable. Moreover, additional uncertainty could be derived from additional elements used for outdoor testing such as tracker mechanisms, the equipment used to measure the performance (accuracy, sampling time, etc.), and data processing to estimate performance.^{383,384} Therefore, the instrumentation, equipment, and data processing for outdoor tests may differ from those used indoors to ensure reproducible tests.

To fully characterize PV devices, it is necessary to evaluate the impact of temperature variations throughout the months (seasonal behavior) on their performance. The effect of temperature on PV devices is mainly considered through the thermal coefficients (TCs) for maximum power (P_{max}), open circuit voltage (V_{oc}), and short circuit current (I_{sc}). These are typically estimated as the slope of the linear regression of the processed I - V curve data as a function of device temperature ranging between 25 and 65 Celsius degrees ($^\circ\text{C}$). TCs could be sensitive to solar simulator spectral distribution depending on PV technology (temperature-dependent spectral mismatch), requiring a spectral correction. In contrast, under natural sunlight, spectral correction is unnecessary³⁸⁵ (IEC 60891 Ed. 3.0 b:2021, 2021). Accordingly, the impact of irradiance and temperature on the performance of the PV device under test must be measured during the evaluation time or extrapolated to complete the characterization of any PV technology. Therefore, the estimation of TCs and the I - V curve adjustment procedures play a prominent role in significantly reducing the exposure time of outdoor tests by correcting the performance at PRCs or target conditions.³⁸⁶

It is noted that TCs show an irradiance dependence in the case of inefficient or older PV silicon devices. In contrast, efficient and stable devices show an almost constant behavior with irradiance.³⁸⁷ In the case of PSCs, the influence of temperature on the performance of perovskite minimodules have exhibited nonlinear behavior, improving performance at high irradiance levels, highlighting a remarkable competitive aspect in contrast to silicon technology.³⁶⁹ This fascinating and positive effect of temperature on performance was also observed in modules, highlighting an inherent property of the technology



Table 3 Summary of encapsulant materials for PSCs, their performance under real-world conditions, and recommended applications

Material	Compatibility with perovskite	PSCs tested under real conditions	Advantages	Limitations	Type of substrate	Ref.
Ethylene Vinyl Acetate (EVA)	No	No	Economical and widely used for traditional technology	Prone to optical degradation, delamination, and acetic acid formation. Requires high temperature lamination that degrades the perovskite	Rigid	94 and 361
Thermoplastic Polyurethane (TPU)	Yes	Yes	High thermal stability, lightweight, processable	Still in large-scale validation	Rigid and flexible	264, 370 and 373
Polyolefins (POE)	Yes	Yes	Good adhesion, high resistivity, flexible	Curing temperatures > 80 °C	Rigid and flexible	94, 264 and 373
Polyisobutylene (PIB)	Yes	Yes	Low WVTR, good adhesion, flexible	May delaminate under thermal cycling if not optimized	Rigid and flexible	33 and 370
Ionomers	No	Partial ^a	Processable at low temperatures, moderate performance	Poor adhesion to glass	Flexible	42 and 367
Epoxy resins UV curable	No	Partial ^a	Good resistance to humidity	Susceptible to damage from exothermic curing (thermal stress and stiffening), release of volatile compounds and delamination at high temperatures	Rigid	26, 370 and 380
Bilayer (PMMA/SB)	Yes	Yes	High stability under real-world conditions, corrosion and water resistance, good adhesion to substrates, and can be applied <i>via</i> a simple and cost-effective solution-based process, without the need for complex techniques such as ALD or UV/thermal curing	It requires considerable thickness (> 10 μm), which can affect flexibility; in addition, the solvent in the SB layer can damage the Transport Layers	Rigid and flexible	26
Combined (PIB + POE)	Yes	Yes	High barrier against moisture and oxygen, excellent adhesion, good mechanical stability, high electrical resistivity, and outstanding durability under real-world operating conditions	Use of processing temperatures above 80 °C for POE may compromise the integrity of some perovskites and has a risk of delamination during prolonged thermal cycling	Rigid and possibility flexible	372
Multilayer (Polyvinyl chloride belt/silicone/barrier foil/UV adhesive/perovskite carbon module/UV adhesive/barrier foil/double sided tape/polycarbonate sheet)	Yes	Yes	Flexibility for curved surfaces and provides mechanical robustness against harsh environmental conditions	Initial efficiency drops due to repeated bending and susceptible to surface soiling	Flexible	261

^a Partial: These materials have been implemented or evaluated under real-life operating conditions; however, they have limitations in terms of adhesion, chemical stability, and resistance to moisture and oxygen. Therefore, they require improvements to be implemented under outdoor operating conditions.

that could be exploited for energy production.³²⁵ However, a strong dependence of TCs on the architecture and quality of devices was observed and interpreted due to ion migration and accumulation at interfaces.³⁸⁸

Accordingly, an outdoor performance characterization test requires an encapsulated device to withstand harsh conditions. Encapsulation protects the devices by improving their thermal and mechanical properties under IEC 61215 tests,³⁸⁹ such as thermal cycling and/or damp heat, while also enhancing electrical insulation, reducing extrinsic degradation processes, and extending their lifetime.⁹⁵ Few studies of encapsulated modules or devices with interconnected cells have been reported for perovskite solar cells. In contrast, most studies

focus on individual solar cells, even performing tests to comply with international standards such as IEC 61215.³⁵ In this context, the fact that small devices can pass some of these tests is insufficient to guarantee successful performance in a large module or interconnected device, since the fabrication and encapsulation processes demonstrated at the cell level cannot necessarily be extrapolated to modules, as scaling introduces additional challenges.³¹⁶

It is worth noting that the standard IEC 61215 is intended to qualify PV modules, considering several tests to detect the most common defects (cell cracks and hotspots) and sources of degradation (discoloration, delamination, corrosion, breakage, *etc.*) observed in commercial PV technologies.³⁹⁰ Moreover,



because the qualification testing of the standard does not consider all failure mechanisms that can occur outdoors, it is impossible to predict or ensure the lifetime of devices from these tests.^{33,391}

In this regard, PV devices' performance evolution over time is characterized by the well-known degradation rate parameter, providing information related to the performance losses per year, allowing estimation of the energy production and leveled cost of energy (LCOE). A degradation rate of 0.55%/year is associated with a 25-year lifetime of mature PV technologies such as silicon, which are optimized to withstand outdoor conditions. Nevertheless, the degradation rate depends on various factors, including weather conditions, installation processes, mounting configuration, maintenance guidelines, system operation (inverter operation, clipping, sizing, *etc.*), and initial PV device performance stabilization (especially for thin-film modules). Therefore, at least a few years of performance data are required to observe longer-term trends and estimate accurate values.^{392,393} This fact is a critical issue for perovskite photovoltaic technology because most outdoor reports consider months of data, as previously shown in Fig. 1. Therefore, it is not easy to decouple outdoor performance from seasonal weather conditions. However, the degradation patterns or degradation shapes of performance curves observed for silicon technology³⁹⁴ have been also observed in long-term outdoor tests (months) of PSCs, suggesting that similar degradation models such as linear, concave, and convex can be used to estimate lifetime and correlated with important parameters such as the ideality factor to elucidate the degradation or failure modes.^{69,395}

Regarding long-term tests, the perovskite community has adopted the International Summit on Organic Photovoltaic Stability (ISOS) protocols, which suggest three procedures for outdoor testing.³⁴ ISOS-O1 suggests keeping the device at V_{oc} or maximum power point (MPP). ISOS-O-2 suggests additionally measuring I - V curves under natural sunlight. ISOS-O-3 suggests keeping the device under MPP and measuring the I - V curve outdoors and indoors. Nevertheless, only one operative point is observed during outdoor exposure (V_{oc} or P_{max}). Keeping the device at I_{sc} or V_{oc} is known to negatively impact the lifetime of PV devices,⁷¹ leading to a misinterpretation of degradation mechanisms and, consequently, to unreliable stability assessments.^{168,395,396,403} In contrast, P_{max} is the most realistic operating condition for a PV device. However, this operative point is hardly dependent on the implemented maximum power point tracking technique (hardware) and algorithms (numerical method). Hence, various factors must be optimized to ensure the accurate tracking of PV devices, including hardware implementation, required parameters, convergence, tracking speed, implementation complexity, and efficiency, among others.^{99,404} Thus, differences in the behavior and values of MPP are expected to be observed, depending on the device used for tracking and the type of test (indoor or outdoor).

Accordingly, various algorithms or techniques for tracking the MPP have been proposed, with the most commonly used,

being Constant Voltage (CV), Perturb & Observe (P&O), and Incremental Conductance (IC).³⁹⁶⁻³⁹⁸ CV typically sets the MPP tracking (MPPT) value at a fixed percentage of the V_{oc} values, ranging between 65% and 85%. However, this value depends on the weather conditions and the device under test. IC is one of the highest-accuracy algorithms, relying on the real-time estimation of derivatives (dI/dV) and step size, demanding higher hardware resources. P&O is a straightforward algorithm to implement, but it depends mainly on the perturbation step size (large step sizes can reduce accuracy, and small step sizes imply low speed), resulting commonly in drawbacks such as a slow response under rapidly changing weather conditions and oscillation problems around the MPP.³⁹⁹ This numerical stability issue has also been observed in perovskite devices exhibiting hysteresis.^{396,403}

While conventional math-based MPPT algorithms perform adequately under uniform irradiance conditions and can be readily implemented in physical controllers due to their relatively low complexity, their effectiveness could significantly diminish under rapidly changing, non-uniform irradiation or shading effects.^{390,400} Hence, it is expected that some algorithms and hardware strategies implemented in MPPT controllers may perform properly for cell devices but improperly for module devices, considering that partial shading on modules is more common, producing multiple local MPPs on the power-voltage curve, making it a challenging task that has been overcome using algorithms based on Artificial Intelligence (AI), soft computing, and data-driven algorithms for tracking the true global MPP.^{398,410} Alternatively, strategies such as optimizing the duty cycle perturbation and sampling rate,⁴⁰¹ or employing galvanostatic approaches⁴⁰² can be explored to improve the effectiveness of MPPT, for instance, using the P&O algorithm.

It is important to note that the ability of MPPT systems to track the MPP of PV devices accurately depends not only on the algorithm implemented but also on the design of the buck converter (hardware). The converter must be appropriately sized to enhance efficiency, considering the operating mode (such as continuous conduction mode), the duty cycle (the efficiency tends to decrease when the duty cycle is reduced), *etc.* These parameters are linked to the quality and cost of the MPPT converter.⁴⁰³ Herein, the application of real-time simulation technology, such as Hardware-in-the-Loop (HIL),⁴⁰⁴ has overcome the limitations of hardware and simulation techniques as it allows for testing MPPT controllers as well as adjusting model parameters.⁷⁸ This results in a powerful tool to design a MPPT controller considering the particularities of PSC, such as hysteresis behavior, and evaluating the performance stabilization behavior observed in perovskite devices depending on the MPPT technique used to extract P_{max} .⁴⁰⁵

Although several alternative strategies can be used or adapted to improve MPPT efficacy, only a single operating point can be monitored during the outdoor test. In contrast, frequently tracing I - V curves of devices under outdoor test enables the estimation of key performance parameters (P_{max} , I_{sc} , V_{oc} and FF) which can be correlated with weather variables



(irradiance and temperatures) as suggested by IEC 61853. This approach enables the consideration of seasonal effects, offering valuable insights into performance degradation, estimating a reliable degradation rate, and predicting device lifetime based on statistically robust data to ensure test reproducibility.⁶⁸ Furthermore, the outdoor test data can be processed to estimate thermal characteristics (TCs),⁵⁸ the impact of weather conditions on performance based on the PRC (STC, NOCT, *etc.*), to perform a proper comparison with other PSC architectures or PV technologies,³⁹⁴ as well as for the diagnosis of faults. In addition, due to the significant variability of hysteresis exhibited by perovskite devices under different conditions of irradiance and temperature,⁵⁹ recording forward and reverse curves during stability tests helps to study the hysteresis behavior and gain insights into performance degradation.⁴⁰¹

5.2. Infrastructure requirements for outdoor testing

The type of outdoor test determines the variables to be monitored for a given PV technology. For instance, the energy rating of PV modules (IEC 61853-3⁴⁰⁶), which is the energy output (watt-hours) produced over a period of time as a function of the climate profile (incident irradiance and device temperature) under STC. However, this methodology does not consider degradation mechanisms or transient behaviors. Additionally, the accuracy of energy production estimation depends on the sampling time of measurements and the procedure used to track the MPP, as was mentioned. On the other hand, evaluating the impact of weather variables on PV performance (IEC 61853-1⁴⁰⁷) requires characterization under various ambient conditions.

For a comprehensive evaluation of outdoor performance, a dedicated infrastructure is required to monitor electrical and environmental parameters. The essential components for accurate and reliable measurements over time are outlined in Fig. 6. A well-designed outdoor solar station allows correlation of environmental variables and performance stability over time under real stress conditions. This approach is essential for assessing long-term stability and optimizing solar energy systems. Fig. S2 shows a real solar station designed in accordance with the IEC recommendations presented in Fig. 6. This setup is intended for evaluating and characterizing the performance of PV devices and monitoring their evolution, including stability studies, under natural sunlight without a solar tracker. To ensure the reliability of the measurements, the components must be calibrated and installed in accordance with regulations (IEC 61215⁴⁰⁸). With respect to the test module, it must be mounted in a coplanar arrangement with the irradiance sensor (pyranometer or reference cell) at a distance of ≤ 0.3 m away to minimize measurement errors. Moreover, panel temperature sensors must be positioned to various locations on the back of the test module to allow for better measurement statistics and to obtain the panel temperature measurement from multiple regions. On the other hand, to prevent panel shading, wind speed sensors should be placed at 0.7 m. This also ensures reproducibility in the measurement of convective cooling by wind. The inclination of the test modules should be optimized

to obtain maximum energy production. However, the inclination of the modules can be adjusted to facilitate maintenance or to install solar tracking systems.

Fig. 6 illustrates a full outdoor test setup compliant with IEC standards that include additional elements for the purpose of monitoring both the weather parameters and device performance. While advanced meteorological instruments, such as anemometers, rainfall sensors, and snow sensors, may provide valuable data for targeted analyses and fair comparisons between different climate regimes, in most cases, the observation of a subset of the most critical environmental parameters is sufficient. That is, solar irradiance, ambient and device temperatures, and relative humidity afford the necessary information for the analysis of degradation phenomena and the association of performance loss with the corresponding environmental stress.³⁷ These parameters are directly related to dominant PSC degradation phenomena such as thermally induced phase transitions, moisture-induced decomposition, and light-activated ionic transport.¹³⁹ Accordingly, for benchmarking long-term outdoor stability, for example, a basic setup focusing on the most critical parameters is most effective and enables broader implementation of outdoor test protocols. Advanced sensors may still be included in studies aiming to investigate specific aspects of environmental influence or to generate high-resolution performance prediction models. A simplified setup is shown in Fig. S3 and described more extensively in Note S2, which includes recommendations for cost-effective choices in the selection of device components.

In addition, another important component in solar installations is the use of systems that track the sun's path continuously, optimizing incident radiation. This increases energy production between 22% and 56% compared to fixed installations.⁴⁰⁹ There are two main classifications for solar tracking systems. Single-axis trackers are devices that enable the tracking of the sun by rotating on a single axis, typically in the east-west direction. Dual-axis trackers adjust in two directions to account for seasonal variations.

Energy efficiency depends not only on these systems, but also on geographical location, weather conditions, and the configuration of mechanical devices. However, solar tracking systems have progressed significantly with the implementation of artificial intelligence-based systems (AI). The use of real-time weather data has helped to improve the accuracy of solar trajectory prediction.⁴¹⁰ The use of AI for monitoring purposes has not yet been widely adopted for PSCs. The integration of this technology has the potential to improve energy production by increasing total incident radiation and influencing operational stability.

5.3. Bridging indoor testing and outdoor performance prediction

For commercial solar technologies, accurate performance prediction in photovoltaic (PV) devices under standard indoor measuring conditions and translation to outdoor operation is imperative. With this approach, there can be effective performance assessment across diverse climatic environments and



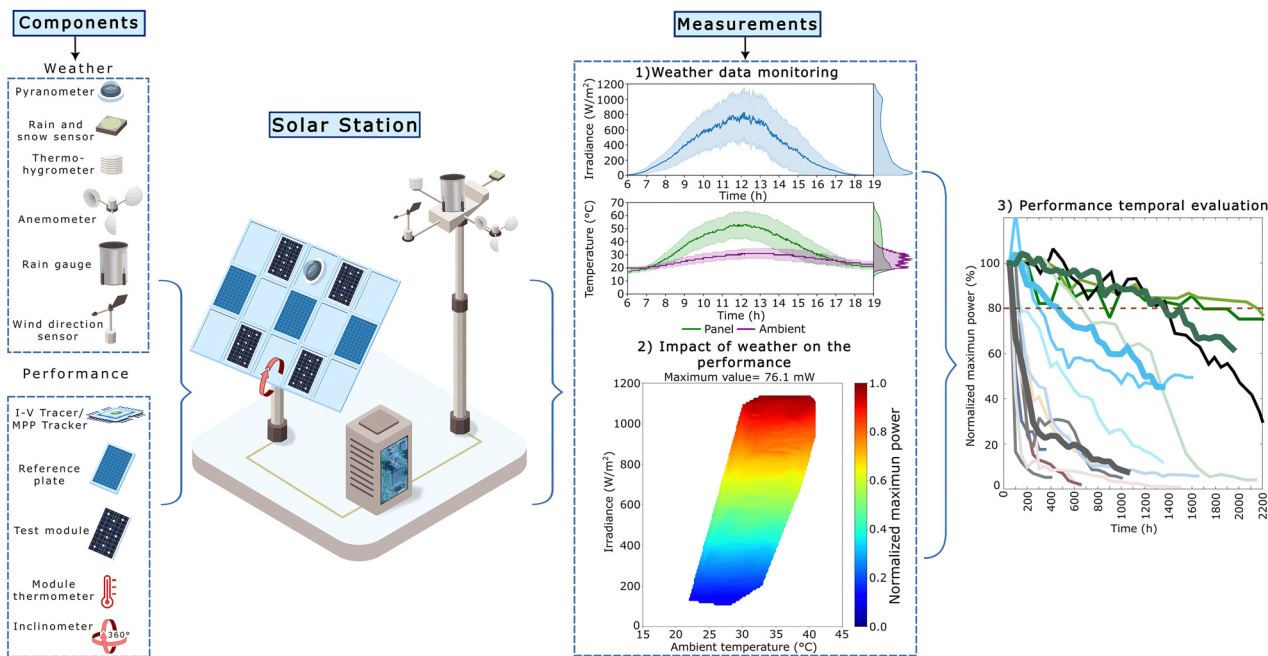


Fig. 6 Solar station components and environmental monitoring setup for evaluating PSCs performance under real-world conditions. The left section illustrates the key components of the solar station, including meteorological sensors (e.g., pyranometer, anemometer, and rain gauge) for weather data acquisition, as well as performance monitoring devices (e.g., *I*-*V* tracer, reference plate, and module thermometer) used to assess PSCs behavior. These components enable the collection of both operational and environmental data. The center section presents a schematic representation of the solar station, highlighting its integrated monitoring capabilities. The right section displays key measurement outputs. This includes (1) time-series plots of meteorological parameters such as irradiance and panel temperature, providing insight into daily environmental variations, (2) a performance map that correlates irradiance and ambient temperature with normalized power output (adapted with permission from ref. 395, Copyright © 2021, Springer Nature) and (3) A long-term performance evaluation graph showing the temporal degradation of PSCs efficiency under outdoor conditions (adapted with permission from ref. 395, Copyright © 2021, Springer Nature). This comprehensive monitoring approach allows for an in-depth analysis of how environmental stressors impact PSCs stability and efficiency.

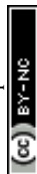
provides a sound basis for estimating operational reliability and service lifetime. Establishing robust indoor–outdoor correlations further streamline certification procedures, informs warranty and financing frameworks, and enhances manufacturers' confidence in the long-term field performance of photovoltaic modules.

5.3.1. Established modelling strategies in silicon photovoltaics and their transferability to PSCs. For silicon PV technology, numerous studies have been successful in developing models for the estimation of long-term outdoor performance. Aissa Meflah *et al.*⁴¹¹ for instance, proposed a procedure for the estimation of the stability of outdoor performance by the two-diode model and method of correction by irradiance (eqn (1)). The model in calculating the panel output current uses eqn (1) where I_{ph} is the photocurrent; I_{rs1} and I_{rs2} are the two diodes' reverse saturation currents, respectively. The contact loss is included in I_{rs1} , whereas recombination loss occurring at the depletion region is denoted by I_{rs2} . V_{t1} and V_{t2} are the thermal voltages in the two diodes; R_s is the series resistance in representation of the resistive loss occurring in the flow of current through the material within the semiconductor and the contact interconnections; R_{sh} is the shunt resistance accounting for leakage paths through the p–n junction describing the presence of alternate unintended path of current; and n_1 and n_2

are the diodes' ideality factors.

$$I = I_{\text{ph}} - I_{\text{rs1}} \left(\exp \left(\frac{V + IR_s}{n_1 V_{t1}} \right) - 1 \right) - I_{\text{rs2}} \left(\exp \left(\frac{V + IR_s}{n_2 V_{t2}} \right) - 1 \right) - \frac{V + IR_s}{R_{\text{sh}}} \quad (1)$$

Performance measurements are time-corrected in response to variations in irradiance, temperatures, and electrical parameters throughout the durations of sunny and cloudy days to compensate for variations in real-world operating conditions. The model iterates the absorbed irradiance (G), implicitly included in the photocurrent (I_{ph}), at different times throughout the day to match the measured short-circuit current (I_{sc}). A linear correction coefficient is implemented to compensate for module-specific and spectrally differing absorptions. In addition, the thermal voltage and the diode saturation currents are re-evaluated at all time steps to correspondingly model time-dependent variations in the environment including changes in temperatures and irradiances on a daily cycle basis. The approach enables long-term stability modelling with minimal loss in accuracy (approx. 5%),⁴¹¹ providing a valid estimate of silicon PV module performance on a long timescale for longer operating periods.



In contrast, applying these established strategies directly to perovskite solar cells has proven significantly more challenging. Primarily due to the inherently dynamic and transient nature of the PSCs degradation. Although two-diode equivalent circuit plus irradiance adjustment method-type model applications have been effective in silicon-module outdoor performance prediction, the respective applications on PSCs show markedly smaller success rates. Such lack of efficacy is a direct outcome stemming from the crucial difference in the mechanism of degradation. Silicon-based PV technology is predominantly affected by physical, electrochemical, and thermal degradation stemming from well-known mechanisms such as corrosion, microcrack formation, and potential-induced degradation (PID), with typical yearly performance losses of $\sim 0.5\text{--}0.6\%$ under temperate climates.^{392,412} PSCs exhibit degradation driven by humidity and temperature dependent ionic migration, interfacial reactions, volatile species transport, and light-induced phase segregation.⁴¹³ These processes introduce hysteresis, transient behavior, and path dependent degradation trajectories that violate the core assumptions of equivalent circuit-based models.

Another dimension to consider is the relative susceptibility of PSCs towards environmental degradation. Silicon modules benefit from decade-long advancements in the field of encapsulation technology capable of slowing moisture as well as oxygen penetration, which has enabled lifespans exceeding 25 years in modules. In contrast, PSCs are significantly more susceptible to minor defects in the encapsulation domain. Even in optimal conditions of encapsulation, native processes of degradation such as ionic migration may still pose a threat towards stability.⁴¹⁴ Dynamic behaviors, partial reversible defect passivation under light, nocturnal relaxation, light soaking effects, and daily thermal/photofatigue cycling,^{415,416} complicate any attempt to map short-term indoor measurements to long-term field performance. These reversible-irreversible interactions challenge silicon derived models that assume monotonic and mechanism consistent degradation pathways.

These discrepancies explain why silicon PV modelling approaches must be modified for PSCs and why new strategies should account for PSC-specific transient and environmental degradation phenomena.

5.3.2. Assessment of indoor accelerated aging protocols for PSCs degradation modelling. The PSC field is developing indoor test protocols to reproduce long-term outdoor degradation. This problem has been investigated by means of a comparison between indoor accelerated aging and outdoor conditions. For instance, Zhao *et al.* used the ISOS-L3 protocol at different temperatures and 65% relative humidity and found that the transient degradation behavior of PSCs showed an Arrhenius-type temperature dependence.¹⁵⁵ After conducting a 1200-hour ISOS-L1I stability test on semitransparent PSCs, Liu *et al.* found that continuous 1-sun illumination only caused a 3.5% degradation.⁴¹⁷ A linear extrapolation of the degradation rate led them to predict a T_{80} lifetime of more than 6500 hours. This is equivalent to over 20 years of outdoor operation, when the average daily irradiance in Princeton ($\sim 850 \text{ Wh m}^{-2} \text{ day}^{-1}$)

is included, although this assumes constant degradation rates and similar failure modes between indoor and outdoor environments, which may not reflect real-world PSC behavior.⁴¹⁷ For this reason, this remains an indoor-only validation; the extrapolated lifetime is derived under fixed illumination and static environmental stresses, without incorporating factors that are known to vary outdoors, such as spectral irradiance shifts, daily or seasonal humidity and temperature ramp rates, or module scale thermal gradients. Therefore, the equivalence to a 20-year outdoor duration should be interpreted as a benchmark estimate rather than a quantitative field prediction.

In addition to following the ISOS-D3 procedure for accelerated aging and extending IEC 61215: 2021, Kobayashi *et al.* performed 3000 hours of moist heat testing at 85 °C and 85% relative humidity.⁴¹⁸ The T_{90} data showed an approximate T_{90} value of 3260 hours, which was then converted to a projected outdoor durability estimate of 20 years. This projection was based on benchmarks for silicon PV panel aging and therefore carries uncertainty when applied to PSCs, as the degradation mechanisms and environmental sensitivities differ substantially between the two technologies. Although this study demonstrates remarkable indoor durability and parallelism with silicon-based predictive models, the inherently nonlinear and transient degradation mechanisms of PSCs underscore the need for complementary testing approaches that account for the specific failure modes and sensitivities inherent to perovskite photovoltaics. In particular, the acceleration factors are inherited from silicon IEC protocols and are not calibrated against PSCs specific degradation kinetics, such as ion migration, bias induced interface reactions, or the sensitivity to humidity and temperature ramp rates. As a result, the 20-year estimate reflects equivalent stress severity, but not necessarily the same failure modes as those active under outdoor operation.

To achieve this objective, Hans Köbler and his team conducted a comparative analysis of constant illumination, light-cycle (LC), and outdoor testing methodologies with the aim of investigating the transient behaviors of PSCs.⁴¹⁹ The study revealed significant discrepancies between indoor and outdoor test results attributable to transient effects. A correlation between indoor and outdoor performance was observed; however, the different behaviors could be attributed to variations in hole transport materials and their effect on device stability. Therefore, LC tests are not universally predictive, the development of improved testing protocols and robust encapsulation strategies is necessary to address correlation errors between indoor stability predictions and actual outdoor performance of PSCs. Mark Khenkin *et al.* reported a two-year outdoor stability study,¹³³ demonstrating a strong correlation between LC tests and real-world outdoor performance. The research findings indicate that different PSC architectures manifest distinct behavioral patterns when subjected to stability testing. The triple cation/ NiO_x perovskite cells demonstrated a uniform decline in performance under both LC and outdoor conditions, supporting the suggestion that LC tests can reliably indicate long-term trends. The most stable dual-cation/SAM perovskite



cells also performed well outdoors, with no significant degradation over a year. These results suggest that LC tests may reflect degradation caused by ion migration during night/day cycles better than traditional constant illumination experiments. Ulas Erdil *et al.* proposed an innovative accelerated aging protocol that applies a continuous forward bias in the dark, followed by a post-bias rest phase, to simulate migration-driven degradation typically observed under outdoor exposure conditions.¹⁵² Unlike conventional tests, this methodology reflects natural cycles and variations better, without needing illumination. It shows that forward polarization impairs charge transport by increasing recombination losses. There is some recovery during rest, but it also causes a decline. The degradation patterns in this indoor study are like those in an outdoor 20-month study in Berlin, showing that the method is relevant. This low-cost, non-illuminating strategy has the potential to be a promising alternative to conventional stress tests for assessing PSC stability. Together, these studies show that LC and bias-driven indoor tests can reproduce specific outdoor degradation signatures, mainly those linked to ion migration and transient interfacial processes, but their predictive value remains limited. The correlations observed are architecture dependent, and each protocol isolates only a subset of relevant stressors. None of the approaches incorporate real spectral variation, diurnal MPP transients, humidity or temperature ramping, or module-scale gradients. Thus, while mechanistically informative, these tests do not yet provide a complete basis for quantitative outdoor lifetime prediction, underscoring the need for multi-stressor, climate-relevant stability frameworks.

Simultaneously, Jiang *et al.* developed a temperature-dependent degradation model based on accelerated indoor testing under 1.2-sun illumination at temperatures ranging from 25 °C to 85 °C.¹³² Arrhenius analysis was applied to the data, resulting in an apparent activation energy of approximately 0.59 eV. Subsequently, the model was combined with recorded outdoor ambient temperature data, incorporating an estimated compensation for the internal temperature of the device of 40–50 °C, in order to calculate weekly degradation rates and predict cumulative performance losses under outdoor conditions. Over a 26-week period, the degradation trends showed a high degree of agreement with actual measurements, validating the use of indoor thermal stress testing to predict real-world behavior, provided the dominant degradation mechanism remains the same. To make standard accelerated stress tests more reflective of behavior in the real world, Song *et al.* highlights the importance of including the actual conditions of devices exposed to outdoor environments in indoor testing.³⁶ The proposed method incorporates light and dark cycles, temperature variations, and other environmental elements relevant to outdoor operation. Additionally, the influence of perovskite composition on degradation was investigated, showing that during the night, FAMA-based solar cells self-recovered, while ion migration caused deterioration in FA/Cs-based devices during thermal cycles. This study proposes the development of real-world-like tests to increase the

accuracy of degradation behavior across different perovskite compositions. These results show that temperature-activated indoor models can reproduce outdoor behavior only when the same dominant mechanism is active. However, as demonstrated by real world emulation protocols that include light/dark cycling, temperature variations, and composition dependent ion migration, can trigger mechanism changes that are not captured by purely temperature-based Arrhenius extrapolations. This confirms the need for accelerated indoor tests that integrate the full transient coupling of illumination, temperature, bias, and environmental factors to achieve robust outdoor predictability.

5.3.3. Advanced data-analytic and machine learning tools for lifetime prediction. In order to capture couple predictive models. Tress *et al.* introduced a predictive framework based on mapping PCE as a function of temperature and light intensity, $PCE(T,I)$, through stabilized MPP tracking.⁴²⁰ Using real irradiance and temperature data from Lausanne, they recreated 24 representative daily cycles in the laboratory and integrated the $PCE(T,I)$ function over time to estimate outdoor energy yield.⁴²⁰ By comparing the predicted and actual energy harvested during emulated operation, it was possible to distinguish between reversible degradation (recovering overnight) and irreversible degradation (accumulating over the test period). The proposed approach constitutes a pragmatic methodology for the emulation of realistic outdoor conditions, thereby facilitating the isolation of the respective contributions of distinct degradation mechanisms to the overall performance loss.

To improve the accuracy of indoor stability tests, Noor Titan *et al.* proposed a new indoor aging protocol to more accurately predict the long-term outdoor stability of PSCs.⁴²¹ The study showed that traditional continuous illumination tests do not replicate real-world day–night cycling, so an accelerated aging method was introduced that combines elevated temperatures with shortened light/dark cycles. The improvement in the correlation between indoor and outdoor degradation behaviors is significant. An Arrhenius-based equation was presented for calculating cycle duration. This method provides a solid framework for simulating the degradation of PSCs. For example, the test has different cycle durations depending on the temperature: 12 hours of light/12 hours of darkness at 25 °C, but at 55 °C, the cycle is reduced to 1.5 hours of light/1.5 hours of darkness. The acceleration factor, which defines the rate at which the degradation cycle occurs relative to real-world conditions, ranged from 5× to 46×. This acceleration factor was obtained by increasing the temperature and reducing the cycle duration. Among the PSC architectures tested, those based on SAM showed the fastest degradation rate (46×), while NiO_x-based PSCs and NiO_x/SAM hybrids demonstrated significantly greater stability (~5×). The model was validated on different hole transport layers in PIN architectures, demonstrating that degradation trends at higher temperatures with shorter cycles are like those at lower temperatures with longer cycles. This confirms the reliability of the estimation model (see Note S3).



The integration of illumination and temperature in these indoor protocols has been shown to enhance the predictive accuracy of the models. Furthermore, the incorporation of environment-dependent acceleration factors has been demonstrated to further improve the correspondence between the indoor and outdoor degradation trends. However, variable humidity and differences in perovskite composition can alter the dominant degradation mechanisms, especially in extreme climatic conditions where encapsulants may fail. This makes it necessary to validate data collected outdoors under variable climatic conditions, mainly in tropical climates with high UV, humidity changes, and constant high irradiance. This ensures the opportunity to adjust acceleration models to generalize and strengthen different environmental stress regimes.

Machine learning (ML) has been used to predict long-term PSC performance under real conditions. These methods use short-term testing to generate fast and accurate predictions. Unlike conventional models, these models could interpret the nonlinear and combined effects of stress factors present in outdoor conditions. Consequently, they facilitate the reduction of testing time, provide guidance on material and architecture optimization, and support the transferability of lifetime predictions across various climates and device configurations.

Graniero *et al.* conducted a comprehensive ML study using both the Perovskite Database Project and a high-resolution in-house aging dataset.⁴²² The demonstration showed that nonlinear models, *e.g.*, the Random Forest algorithm, can identify key elements that affect stability, such as perovskite composition, charge transport layers, and environmental exposure. Researchers emphasized the fundamental importance of a high-quality dataset for model accuracy. The study found no significant improvement with larger datasets if the data quality was not improved because there were inconsistencies among common aging data metrics. Standardized and high-quality datasets are needed to improve machine learning reliability and ML-based predictions should be validated against long-term outdoor datasets before being used as definitive lifetime estimations.

In a similar vein, Odabaşı and Yıldırım employed ML-based meta-analysis to analyze a curated dataset of 404 PSCs, which had been compiled from 181 publications.⁴²³ Subsequently, the researchers employed association rule mining and decision tree modeling, thereby identifying strategies such as mixed cations, multi-spin deposition, SnO₂ as ETL, and storage at low humidity. This process yielded models with interpretable rules, such as high stability at <30% humidity. The study demonstrates the value of statistical learning in deriving practical design guidelines, although such guidelines are not yet developed to directly extrapolate outdoor lifetimes, and their applicability remains limited without controlled real-world validation.

To enhance the accuracy of ML models, Kouroudis *et al.* advanced the field by introducing a wavelet-assisted ML framework to quantitatively predict outdoor PSC degradation from short-term indoor aging tests.³³⁴ The methodology combines signal preprocessing, utilizing IIR filtering and Daubechies-2

wavelet transforms, with a Kernel Ridge Regression (KRR) model trained on time-series degradation data collected under controlled indoor conditions (varied light intensity, atmosphere, and temperature). To avoid overfitting, indoor tests were selected using the Fréchet distance metric. The model's robustness was enhanced using Bayesian optimization and six-fold cross-validation. The model predicted outdoor degradation behavior effectively (MSE < 0.24) across various climates, *e.g.*, Barcelona and the Negev desert, but the predictions remain dependent on the representativeness of the indoor stress profiles used for training.

5.3.4. Perspectives to bridging indoor testing and outdoor performance prediction. Indoor evaluation protocols have enabled important progress in predicting outdoor performance. However, perovskite solar cells have transient degradation mechanisms that cause them to diverge from the patterns observed in other technologies. Therefore, simple extrapolations from previously used models are unreliable. Additionally, models that consider only a single stress factor can reproduce degradation trends only under specific conditions and typically fail when conditions change. When models couple multiple variables, their predictive accuracy improves. However, their validity varies across climatic zones, since factors such as humidity and daily thermal cycles can shift the dominant degradation mechanism.

Consistent with this, a recent review of accelerated degradation models⁴²⁴ concluded that traditional single-stressor approaches such as Arrhenius or Eyring capture temperature-driven degradation but break down when humidity, light, or electrical bias introduce additional pathways. Humidity-driven degradation requires dual-stressor formulations like Hallberg-Peck, while PID follows inverse-power-law kinetics, and no single model can represent all coupled effects observed in real devices. The review therefore recommends the development of multi-variable degradation models, such as Arrhenius-light, Arrhenius-RH, or Subramaniyan-type conjunction models, that explicitly integrate temperature, humidity, illumination, and electrical bias within a unified acceleration framework to enable more realistic stability predictions under complex stress conditions.

ML also shows strong potential for improving outdoor prediction when supported by high-quality datasets. Current studies consistently demonstrate that accurate outdoor predictions are only possible when indoor training data capture the same dominant stress factors operating in the field. For example, Kouroudis *et al.*³³⁴ showed that 1-sun nitrogen tests yielded the lowest prediction error because the outdoor devices used for validation were encapsulated and therefore degraded mainly through thermal-ionic pathways; in contrast, indoor air tests performed more than twice as poorly, indicating that models trained under mismatched stress profiles fail to reproduce outdoor behavior. They also demonstrated that prediction of accuracy declines when indoor and outdoor protocols diverge, such as differences in UV shielding, highlighting the sensitivity of ML models to inconsistent stress environments. In practice, this means that ML models should be trained



within the same degradation regime they are expected to predict, encapsulated outdoor devices require encapsulation-relevant indoor stress profiles, whereas unencapsulated devices demand training data that explicitly include air exposure, humidity, and UV.

In this context, a simplified but practical approach is for each laboratory to train ML models using its own indoor protocols, device architectures, and encapsulation strategies. Such lab-specific training can yield accurate predictions for that laboratory device, allowing ML to serve as a tool for forecasting outdoor performance under the specific conditions relevant to that fabrication line. In contrast, building a generalized, climate agnostic outdoor-prediction model requires structured dataset expansion. High-quality indoor measurements under controlled 1-sun illumination, varied temperatures, and well-defined atmospheres (N_2 vs. air) should form the core of the training set, recorded through stabilized MPP or PCE time series. Additional stressors should be included only when they correspond to realistic degradation pathways for the devices being modeled, such as humidity-driven changes, light/dark transients, seasonal effects (self-recovery), solar cells architectures, or composition-dependent ion migration. Additionally, given that machine learning predictions exhibited reduced accuracy when indoor and outdoor stress profiles diverged (e.g., UV exposure, imperfect encapsulation, moisture ingress), a subset of device data could comprise measurements that integrate indoor and outdoor measurements across disparate climate zones. The expansion of the diversity of datasets under controlled and well-adapted protocols is of greater value than the mere increase in size of the dataset. This is a prerequisite for machine learning models that aim to generalize across different laboratories and climate zones.

6. Conclusions and perspectives

Recent advancements in PSCs have shown encouraging results, with some studies reporting multi-year operational lifetimes. However, transitioning from laboratory success to large-scale and reliable outdoor deployment requires directly confronting several critical bottlenecks. While traditional performance benchmarks and indoor evaluation tests are useful for initial assessments, they still fail to capture the complexities of real-world conditions, such as fluctuating irradiance and temperature variations. Environmental and operational coupled factors, such as high humidity, extreme temperatures, UV exposure and bias play a significant role in different PSCs degradation mechanisms, and become very dependent on the specific location, as each climate presents unique fluctuations. This can be significantly understood by adequate modeling of coupled factors (e.g., simultaneous UV exposure, humidity, high temperature, and electrical bias), as recent advances in *in situ* and *in operando* studies provide critical insights into these mechanisms, enabling a deeper understanding of interfacial reactions and paving the way for more rational design principles that enhance device robustness. While improvements in PSCs

stability, such as those achieved through interface engineering and carbon-based back electrodes, show promise, their long-term viability under outdoor exposure conditions has not been fully validated yet, as required by industry. Therefore, research must pivot from laboratory tuning to a rational, physics-based approach for degradation mitigation. This involves leveraging these advanced *in situ* and *in operando* diagnostic tools to monitor interfacial reactions in real-time, enabling the design of intrinsically more robust bulk materials, highly stable interface layers, and self-healing or passivating encapsulation strategies. On the other hand, the diagnostic tools used in outdoor evaluations, such as thermal coefficients and *I-V* curve tracing, help to identify key degradation pathways, but they alone are not enough to fully predict long-term performance. Issues like insulation, leakage currents, and the formation of hot spots must also be addressed to ensure the durability and stability of PSCs modules. As most stability evaluations are still conducted under controlled environments, there is still a need for real-world testing. This gap in outdoor testing is especially prominent in regions like Latin America, Africa, and Southeast Asia, where climate-specific degradation mechanisms remain insufficiently understood. In general, more outdoor testing in tropical climates is required to collect data of PSC degradation under stringent climate conditions with consistent day/night cycles, serving as reference platforms for accelerated stability testing, material benchmarking and the development of robust commercial photovoltaic technologies, where other engineering issues (insulation breakdown, localized hot spots, and leakage currents) come out as area increases, limiting long-term durability and making more evident the performance and stability gap between small-area laboratory devices and commercial-sized modules. In this sense, more systematic outdoor evaluations of mini-modules, sub-modules, and full modules are needed to address scaling challenges and ensure reliable real-world performance characterization. However, even if these evaluations are currently being conducted, it takes several years to be reported to the community, which combined with a lack of clarity about which standard is most appropriate to report for specific cases, makes it difficult to compare materials and device architectures under different climatic conditions, reporting data across the community using agreed-upon testing protocols. Finally, by focusing research efforts from rational materials and devices design to standardized, geographically diverse outdoor validation, the field can transition from reporting promising laboratory results to delivering a truly viable, commercially scalable, and globally stable perovskite photovoltaic technology. The long road to outdoor stability demands a fundamental shift toward real-world testing and predictive engineering to fully unlock the transformative potential of PSCs.

Disclaimer

We have used AI assistance to help with the English language formulation and some grammatical aspects. All technical points, Ideas, original text, and results in this review are our



own original contributions provided in only some parts to the AI system. We have thoroughly written the entire review and verified that no information has been fabricated or misrepresented in the grammatical improvement process.

Author contributions

J. J. P. L. and K. G. R. B. contributed equally to this work. All the authors wrote, analyzed and reviewed the manuscript.

Conflicts of interest

The authors declare no competing financial interest.

Data availability

All data supporting this study is included in the manuscript. All figures are original, created by the authors, and appropriately cited where external sources were used to support their illustrations.

The data supporting this article have been included as part of the supplementary information (SI). Supplementary information is available, this document provides the Scopus-based literature review methodology and datasets (Tables S1–S3) for outdoor stability reports and manufacturing strategies. It includes technical specifications and 18-month environmental data from the Medellín monitoring station. Additionally, it offers practical guidelines for developing low-cost outdoor testing infrastructure. See DOI: <https://doi.org/10.1039/d5cs01085c>.

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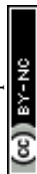
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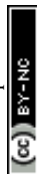
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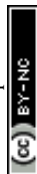
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