Sub-millimetre light detection and ranging using perovskites

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Light detection and ranging (LiDAR) technology measures the time for a short-pulsed light to scatter back from a target object and return to a detector. The measured time is converted to distance using the speed of light. By scanning illumination over a target scene, the system reconstructs a three-dimensional (3D) distance using the speed of light. By scanning illumination over a target scene, the system reconstructs a three-dimensional (3D) distance using the speed of light. Silicon electronics have been the principal option for LiDAR photodetectors in the range of 850–950 nm. However, its indirect bandgap leads to a low absorption coefficient in the near-infrared region, as well as a consequent trade-off between speed and efficiency. Here we report solution-processed lead–tin binary perovskite photodetectors that have an external quantum efficiency of 85% at 850 nm, dark current below $10^{-8}$ A cm$^{-2}$ and response time faster than 100 ps. The devices are fabricated using self-limiting and self-reduced tin precursors that enable perovskite crystallization at the desired stoichiometry and formation of the interface defects with the hole transport layer. The approach removes oxygen from the solution, converts Sn$^{4+}$ to Sn$^{2+}$ through comproportionation, and leaves neither metallic tin nor SnO$_2$ residues. To illustrate the potential of these solution-processed perovskite photodetectors in LiDAR, we show that they can resolve sub-millimetre distances with a typical 50 μm standard deviation.
and it has been shown that oxygen vacancies in metal oxide may contribute to tin oxidation in Pb–Sn perovskites. In this Article, we report a strategy to overcome the incompatibility of NiO, and Pb–Sn perovskites, and create fast and efficient solution-processed photodetectors. We use self-limiting and self-reduced tin precursors. This approach removes oxygen from the solution, converts Sn⁴⁺ to Sn⁺⁺ through a comproportionation reaction and leaves no tin residue after filtration. This allows us to incorporate crystalline NiOₓ as the HTL in Pb–Sn perovskite devices. The resulting devices exhibit an EQE of 85% at 850 nm, dark current below 10⁻¹² A cm⁻² and response time faster than 100 ps. The combined efficiency and speed is more than two times higher than fast Si photodetectors and more than 100 times higher than previously reported solution-processed photodetectors. To illustrate the potential of this approach, we show that the solution-processed perovskite photodetectors can be used to resolve sub-millimetre distances with typical 50 µm standard deviations.

**Pb–Sn photodetector with NiOₓ HTL**

We first fabricated photodetectors with an inverted planar architecture (indium tin oxide (ITO)/HTL/Pb–Sn/C60/bathocuproine (BCP)/Ag), where Pb–Sn is MA₃FA₃PbₓSnₓI₃, and the HTL is NiOₓ or PEDOT:PSS as the control (Fig. 1d, inset). To limit the formation of Sn⁺⁺ vacancies in the perovskite film, we began by using a previously reported metallic tin precursor strategy. We added 0.5 mg ml⁻¹ Sn metallic microparticles (diameter, <150 µm) to the precursor solution, which we vigorously stirred for 10 min. We observed that the addition of metallic tin powder to the precursor solution decreased the EQE of the devices fabricated with NiOₓ as the HTL (Fig. 1d). This finding is in contrast to the improvement we saw in control PEDOT:PSS devices on the addition of Sn powder (Extended Data Fig. 3).

To explore the origins of the drop in EQE observed in NiOₓ devices on incorporating metallic Sn powder into the precursor solution, we considered the possibility that residual Sn species may remain in the precursor solution following filtration—something that could arise due to an inhomogeneous particle distribution (Fig. 2a). To test this hypothesis, we dispersed metallic Sn powder (<150 µm) in a dimethylformamide:dimethyl sulfoxide (DMF:DMSO) (3:1 v/v) solution, stirred vigorously and analysed following filtration (0.22 µm filter). Inductively coupled plasma elemental analysis revealed the presence of Sn species in the solution after filtration (Supplementary Fig. 1). We then treated the NiOₓ HTL with this solution and analysed its surface. Scanning electron microscopy (SEM) images of the NiOₓ surface show a distribution of microparticles (<200 nm) on the surface of NiOₓ. Energy-dispersive X-ray (EDX) mapping demonstrates that these microparticles are Sn (Fig. 2b). Atomic force microscopy (AFM) imaging and X-ray photoelectron spectroscopy (XPS) analysis further support the presence of residual Sn on the surface of NiOₓ (Supplementary Notes 2–4).

**Compatible reducing strategy with NiOₓ**

The ideal reducing agent should provide a favourable balance among the surface area, reaction time and ease of removal, all with minimum probability of passing through the filtration step. The surface of the desired reducing agent should be of a form that exposes a
Fig. 2 | Reducing strategy for Sn precursor. a, Schematic of the Sn metallic powder reducing precursor process. b, SEM images and EDX Sn elemental mapping from the Si/NiO, surface treated with Sn powder solution before filtration (i). The blue colour in the elemental map is assigned to elemental Sn (Supplementary Fig. 2 shows the EDX spectrum and analysis details). SEM images and EDX Sn elemental mapping from the Si/NiO, surface treated with filtered Sn powder solution (ii). The blue colour in the elemental map is assigned to elemental Sn (Supplementary Fig. 3 shows the EDX spectrum and analysis details). c, Schematic showing the Sn metallic wire reducing precursor process. d, Surface SEM images of Pb–Sn perovskite spin coated on NiO, with different reducing strategies: NiO,–pristine precursor (i), NiO,–TPRP (ii) and NiO,–TWRP (iii). e, Histogram of the perovskite grain size for NiO,/Pb–Sn perovskites in previously shown SEM images (Supplementary Fig. 5 shows a comparison with PEDOT:PSS). f, EQE of the devices using the TWRP strategy.

high surface area to the solution and does not disintegrate into smaller pieces during long-duration reactions with oxygen. We posited that if we could simultaneously satisfy these requirements, the Sn surface would function as a self-limiting reducing agent for Sn⁴⁺, and that this would ensure that the time of reaction is not a critical process parameter: similarly, it would provide a wide process window in which high performance was achieved. Experimentally, we found that tin wire satisfied these requirements (Fig. 2c): the reactive macroscopic surface of Sn wire acts as a self-limiting reducing agent to convert unwanted Sn⁴⁺ to Sn²⁺, allowing Sn wire to react with oxygen in the precursor without leaving metallic Sn or SnO₂ residue after filtration.

To explore the effect of reducing strategy (tin-wire-reduced precursor (TWRP) versus TPRP) on perovskite film growth, we compared the surface morphology of perovskite films on different HTLs. The SEM images of the surface show that the grain size of perovskite films grown on NiO, is sensitive to the reducing strategy employed—in contrast to the case of PEDOT:PSS (Supplementary Note 1). This agrees with the following picture of the surface reactivity of NiO₂: unlike organic HTLs such as PEDOT:PSS, NiO₂—with its surface that includes exposed oxygen—is prone to react with residual metallic Sn, leading to the formation of an interfacial non-uniform and non-conformal carrier extraction barrier (Extended Data Fig. 4). The residual Sn species—whether metallic or oxide—present at the interface with NiO₂ impede the nucleation and crystallization of perovskite and lead to a smaller grain size (Fig. 2d,e).

A Foster diagram shows that different forms of SnO₂ have lower formation energies compared with NiO, (Extended Data Fig. 4a). Defects derived from this interface (NiO₂–perovskite) can be asymptomatic for d.c. photovoltaic operation under solar irradiance and yet be materially detrimental to the rapid charge extraction needed for time-of-flight (ToF) measurements. Density functional theory (DFT) simulations (based on the Perdew–Burke–Ernzerhof (PBE) functional within spin-polarized treatments) further support the negative impact of SnO₂ formation on the performance of devices and the negative effect on energy-level alignment in devices (Extended Data Fig. 4b–g). This points to the importance of avoiding the presence of Sn residual species on NiO₂ as they are deleterious to the crystallization kinetics of Pb–Sn perovskites and work against the energy-level alignment needed within p–i–n (PIN) devices (Supplementary Note 2).

When we used the TWRP strategy, we no longer saw negative impacts akin to those observed when extra clusters of Sn species were present in the perovskite precursor mix (Fig. 2d,e). Figure 2f shows that the NiO₂ HTL with the TWRP results yields an EQE of ≈ 85%, which is higher than the control PEDOT:PSS devices. NiO₂ devices show a lower dark current compared with PEDOT:PSS, achieving values below 10⁻⁴ A cm⁻² (Extended Data Fig. 3c). The representative dark-current curves for PEDOT:PSS and NiO₂ are shown in Extended Data Fig. 3b.

Pb–Sn photodetector performance
We compare these results with those of the best near-infrared solution-processed photodetectors across organics, quantum dots and perovskites. Figure 3a shows the EQE versus dark current in the highest-performing previously reported detectors having a detection
responsivity reported in Fig. 3b, we obtain a specific detectivity of
Figure 3b shows the overlay of the Pb–Sn photodetector responsiv-
layers, which results in higher dark currents and lower EQE values.
(Fig. 3a). For Si, high speed can only be attained with thin active
solution-processed photodetectors at LiDAR-relevant wavelengths
points higher than the highest-performing previously reported
514
ing at LiDAR-relevant wavelengths (Supplementary Table 4).
Fig. 3 | Performance of Pb–Sn photodetectors. a, Plot of the maximum EQE for wavelengths >850 nm versus the minimum dark current at the operating
bias. For devices operating without external bias, the dark current at 50 mV is plotted (Extended Data Fig. 5a and Extended Data Table 2). b, Responsivity
of our Pb–Sn photodetector compared with commercial Si fast and slow photodetectors. Details of commercial fast and slow photodetectors from different
companies are given in Extended Data Fig. 5b and Extended Data Table 1.

Fig. 4 | Speed measurements for Pb–Sn devices. a, Schematic illustrating the TPC measurements. An RF probe with a two-point contact (signal–ground)
is used to make a connection with the bottom ITO electrode and top Ag electrode with different pixel areas. Direct contact of the RF probe with the surface
of the ETL (without an Ag electrode) leads to a working device with a small area (<10 × 10 μm²). b, A 3D plot showing the normalized response versus
time for different pixel sizes. The Y axis is the diameter of the pixel in micrometres. c, Comparison of Pb–Sn response time (with direct probe contact) with
commercialized 2.0 GHz Si and 5.0 GHz GaAs photodetector. Extended Data Fig. 5c and Extended Data Table 1 show the detailed characteristics of the
shown commercial photodetector.

range of >850 nm. Extended Data Table 1 is limited to reports that
include the measured speed of photodetection.

Integrating the TWRP strategy for Pb–Sn perovskites with the
NiOx HTL offers values of EQE that are 20 absolute percentage
points higher than the highest-performing previously reported
solution-processed photodetectors at LiDAR-relevant wavelengths
(Fig. 3a). For Si, high speed can only be attained with thin active
layers, which results in higher dark currents and lower EQE values.
Figure 3b shows the overlay of the Pb–Sn photodetector responsivity
spectra with high- and low-speed Si photodetectors. The quantum
efficiency of slow commercial Si photodetectors approaches
≈ 93% at 905 nm, but these values are reduced in fast Si photodetectors (EQE <50%). Direct noise measurements in a wide fre-
quency range using a signal analyser show the equivalent noise
level of 100 fA Hz−½ for Pb–Sn perovskite photodetectors. Using the
responsivity reported in Fig. 3b, we obtain a specific detectivity of ≈
1.5 × 10¹ⁱ cm Hz¹/₂ W⁻¹ at 905 nm (Supplementary Fig. 8).

We carried out stability measurements, under ambient condi-
tions, on encapsulated Pb–Sn photodetectors that use NiOx, as the
HTL. Devices with the NiOx HTL retained 95% of their photores-
ponse after 10 h of continuous illumination (Supplementary Fig. 9).
We compare the best-reported stability for photodetectors operat-
ing at LiDAR-relevant wavelengths (Supplementary Table 4).

We used transient photocurrent (TPC; Fig. 4a) to characterize
the response time of Pb–Sn perovskite photodetectors. A two-point
contact (signal–ground) radio-frequency (RF) probe with a pitch of
1.000 μm was used to contact the bottom ITO electrode and top Ag
electrode, whose size defines the detection area. Figure 4b shows the
TPC results for different pixel sizes. An analysis of the response time
versus pixel area (Extended Data Fig. 6) reveals that the response
time decreases with the pixel area and does not plateau even at the
smallest pixel area of ≈0.005 mm². Considering a dielectric constant of ε ≈ 51, estimated via impedance spectroscopy (Extended Data
Fig. 7), we conclude that the geometrical capacitance of the Pb–Sn
perovskite plays an important role in determining speed (Extended
Data Fig. 6); even at the smallest pixel area reported here, we had
not yet reached transport-limited performance, and thus, we were
motivated to develop a strategy to circumvent the geometrical
capacitance limitation.

Direct contact of the signal side of the RF probe with the ETL
surface of the device resulted in a contact area of approximately
10 × 10 μm². This strategy enabled us to reach a full-width at
half-maximum response time of 100 ps. Figure 4c shows a compar-
ison between the response time of the Pb–Sn photodetector with the
direct probe contact and a commercial fast Si and GaAs photodi-
oxide (Extended Data Table 2 provides the details of the commercial

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photodetector). We note that the data shown in Fig. 4c are not for the same detector area; however, a Si photodetector operates in a transport-limited regime and is therefore not limited by geometrical capacitance. Consequently, a decrease in the Si detector size does not lead to a faster response time (Extended Data Fig. 6). As evident from Fig. 4c, the Pb–Sn photodetector offers a narrower full-width at half-maximum value and a comparable fall time (90% to 10%) to 5.0 GHz GaAs photodetector.

**LiDAR with Pb–Sn perovskites**

We developed a system to apply the photodetectors developed here in the LiDAR-based estimation of the distance of objects. We used a Michelson interferometer configuration in a ToF setup (Fig. 5a). Using a beamsplitter, we referenced the pulsed light to a commercial InGaAs at a fixed position. Figure 5b shows the response of a Pb–Sn photodetector at different positions of the moving mirror of the interferometer. The contour graph (Fig. 5c) shows the light travel distance versus time, and the slope of the line gives the speed of light at 299 mm ns⁻¹. To visualize the accuracy of the depth estimation using the Pb–Sn perovskite photodetector, we moved the mirror based on a predefined map and estimated the distance by the same ToF setup shown in Fig. 5a (Extended Data Fig. 8). We then reconstructed a 3D image using these individual measurements (Fig. 5d).

**Conclusions**

PIN photodetectors are used in short-range LiDAR applications. Our Pb–Sn devices meet the specifications in terms of EQE, response time and low dark current that make them competitive in these short-range high-accuracy LiDAR applications. However, to expand its applicability to long-range detection, multiplication will be needed to detect small signals (Supplementary Table 2). The flexibility of solution-processed perovskites may also allow incorporation into an avalanche amplification structure, and will be an important subject of further investigation.

**Methods**

**Materials and preparation.** Unless indicated otherwise, all the materials were purchased from Sigma-Aldrich and used as received. Formamidinium iodide (FAI), methylammonium iodid (MAI), PbI₂, and SnI₂ were acquired from Great Cell and Alfa Aesar. A perovskite precursor solution (1.7 M) was prepared by mixing the precursors based on the molar ratio of FA₀.₇MA₀.₃₀Pb₀.₅Sn₀.₅I₀.₃ in the mixture of DMF-DMSO (3:1 v/v). The SnF₂ precursor was added by 10 mol% relative to the SnI₂ amount. We noticed that it is important to preheat the precursor glass vial at 100 °C for 15 min. The obtained mixture was prepared inside the
nitrogen glovebox. After stirring at room temperature for 2 h, the solution was filtered through a 0.45 μm polytetrafluorethylene (PTFE) syringe filter. For the TPRP samples, tin powders (5 mg mL⁻¹) were added, stirred at room temperature for 10 min and filtered using a 0.45 μm PTFE syringe filter before use. For the TWP samples, a few-centimetre-long Sn wire (99.999%, MilliporeSigma; 0.5 mm diameter) was dropped inside the solution, which was stirred for an additional hour before filtration. We noticed that expanding the time of this reaction to more than one hour does not affect the performance. A 1.0 M NiO₃ solution was prepared by filtering the intrinsically mixed Ni(NO₃)₂·6H₂O with ethylene glycol (anhydrous, 99.8%) followed by the addition of ethyleneamine (≥99.0%) in a molar ratio of 1:1.1:0.8 with respect to the Ni precursor. The solution was stirred overnight, followed by its dilution to the favourable concentration of 0.23 M. The solution was filtered using a 0.45 μm polyvinylidene fluoride (PVDF) syringe filter before deposition.

Device fabrication. For NiO₃ films, the as-prepared NiO₃ solution was spin coated on a precleaned ITO substrate at 4,000 rpm for 90 s and annealed at 300 °C on a hotplate for 1 h in ambient air. We tested the ultraviolet illumination post-treatment, but it had a negative impact on the device performance. For PEDOT/PSS films, the solution (Xian PLT) was spin coated on the substrates at 4,000 rpm for 30 s followed by annealing at 150 °C on a hotplate for 30 min in ambient air. The substrates were transferred to a nitrogen glovebox after annealing. We also diluted the PEDOT:PSS with methanol in a different ratio, but we did not observe any statistical and reproducible difference on the EQE of the devices. For the device fabrication, the spin coated substrates were deposited in a nitrogen glovebox. After stirring at room temperature for 2 h, the solution was filtered using a 0.45 μm polytetrafluoroethylene (PTFE) syringe filter before deposition.

Device characterization. Dark current–voltage (J–V) characteristics were measured using a Keithley 2400 sourcemeter by scanning the bias between −0.3 and 0.3 V for pixels with an active area of 0.1 mm². The EQE spectra were measured using a Newport QUANTX-300 system without applying any external bias, with J–V and EQE also measured at 1 V and 10 V, respectively. The EQE spectra were recorded under a constant light intensity of 1,000 μW cm⁻² at a detection wavelength of 450 nm. The device was kept in the dark and short-circuit conditions. The impedance spectra were collected as the frequency scans from 1 Hz to 1 MHz (Extended Data Fig. 7a). The capacitance component C is calculated based on the relationship with impedance Z and angular frequency ω (ref. 43).

$$C = \frac{1}{(ωZ)}$$

where ω is the permittivity of free space, ω is the dielectric constant of the material, d is the thickness of the light-absorbing perovskite layer and A is the overlapping area between the two electrodes (d = 500 nm and A = 0.1 cm²). To calculate the accurate value of C, a proper equivalent-circuit representative of the resistive and capacitive responses of the system should be fitted (Extended Data Fig. 7a,b, solid lines). Based on previous studies on perovskite solar cells, the equivalent circuit of a modified version of the standard Randles circuit (Extended Data Fig. 7c), which delivers the best fit to the measured data. Fitting results, as well as the calculated dielectric constant, are tabulated in Extended Data Fig. 7d. The obtained value for the dielectric constant is in good agreement with the values reported for metal halide perovskites.

Noise measurement. Noise spectra have been acquired using a signal analyser (N9010A, Agilent) combined with a low-noise transimpedance preamplifier at 10 V A⁻¹ gain (FEMTO, DHPCA-100). The devices (area, 0.1 cm²) were operated under short-circuit and dark conditions.

LiDAR demo. For distance estimation, we used the same equipment and arrangement as the TPC setup to measure the time response, but we split the femtosecond laser using a 50:50 beamsplitter before using one-half of the split beam to illuminate the sample. The other part was used as a reference and was directed to a 30 GHz InGaAs photodetector (Thorlabs, DXM30BF). The oscilloscope was triggered by the photodetector signal collected as the frequency scans from 1 Hz to 1 MHz (Extended Data Fig. 7c), which delivers the best fit to the measured data. Fitting results, as well as the calculated dielectric constant, are tabulated in Extended Data Fig. 7d. The obtained value for the dielectric constant is in good agreement with the values reported for metal halide perovskites.

SEM/EDX mapping measurement. High-resolution SEM images were acquired using a Hitachi SU5000 instrument. The samples were prepared by depositing Pb–Sn PVK on top of NiO₃/ITO and PEDOT/ITO substrates using the same condition as device fabrication. To map the elemental composition of certain SEM samples, EDX analysis was carried out using a Bruker EDX detector. The EDX measurement conditions were set in the range of 5–15kV.

AFM. The film morphology was characterized using an AFM instrument (Asylum Research Cypher) operated in the a.c. tapping mode equipped with AC240TM-R3 probes (resonant frequency, 70 kHz; spring constant, 2 N m⁻¹). To obtain high-quality images, areas of 5 × 5 μm² were probed using 1 Hz scanning rate and 1.024 lines per point. XPS. XPS analysis was performed using an ESCALAB 250Xi system (Thermo Fisher Scientific) with an Al Kα source and a 500 μm X-ray spot size. A beam of low-energy argon-ion clusters (3 keV) was vertically employed on a raster area of 1 mm².

EQE measurements. The responsivity and quantum efficiency of the devices were measured using a QUANTX-300 Newport measurement system. No external bias was applied, and the measurements were taken under a constant nitrogen flow. A standard calibrated Si/Ge reference detector (BD005, Newport) was used to provide values of quantum efficiency/responsivity over the wavelength range from 350 to 1,800 nm. Calibration was conducted before each set of measurements to create the reference values. A Si photovoltaic test cell was used to further validate the calibration procedure.

Impedance spectroscopy. Impedance spectroscopy is employed to estimate the dielectric constant of the Pb–Sn perovskite film. For this purpose, the fabricated device with PEDOT:PSS as the HTL is connected to an Autolab potentiostat/galvanostat module. An a.c. voltage perturbation of 20 mV is applied as the device is kept in the dark and short-circuit conditions. The impedance spectra were collected as the frequency scans from 1 Hz to 1 MHz (Extended Data Fig. 7a). The capacitance component C is calculated based on the relationship with impedance Z and angular frequency ω (ref. 43).
was reconstructed using the estimated travel distance by considering the peak position of the response and speed of light (Extended Data Fig. 8).

DFT calculation. The DFT calculations were performed using the FHI-aims all-electron code. The default numerical settings, referred to as ‘tight’ in FHI-aims, were used. Spin-polarized treatment for Ni atom was incorporated into all the calculated electronic properties and geometry relaxation. More specifically, NiO in all the structures (that is, bulk and superlattice calculations) adopted a type-II antiferromagnetic phase (that is, the spin-ordered vector along the [111] direction). Local minimum-energy geometries of the Bors–Oppenheimer surface were obtained with residual total energy gradients below $1 \times 10^{-4}$ eV Å$^{-1}$ for atomic positions by using the PBE generalized gradient approximation functional. To analyse the band alignment of NiO/SnO and NiO/SnO$_2$, we adopted the method described earlier, in which the valence-band offset, $\Delta E_{\text{val}}$, is defined as below.

$$\Delta E_{\text{val}}(\text{NiO}/\text{SnO}, \text{SnO}_2) = (E_{\text{val}}(\text{SnO}) - E_{\text{val}}(\text{NiO})) + \Delta V,$$

where $E_{\text{val}}(\text{NiO})$, $E_{\text{val}}(\text{SnO})$, and $E_{\text{val}}(\text{SnO}_2)$ represents the position of the valence-band maximum (VBM) with respect to the average electrostatic potential in materials SnO/ SnO$_2$ (NiO), and $\Delta V$ represents the potential alignment term obtained from the superlattice calculations (Extended Data Fig. 4). According to another study, the $E_{\text{val}}(\text{NiO})$, $E_{\text{val}}(\text{SnO})$ value is obtained by using a hybrid functional (HSE06), and the potential alignment term $\Delta V$ is calculated based on a fully relaxed superlattice by the PBE generalized gradient approximation functional (which is proved to give similar accuracy as HSE06, but within less computational cost). More specifically, instead of using the plane-average electrostatic potential to calculate $\Delta V$, the electronic potential at the cores of oxygen atoms is used to eliminate the oscillations caused by the ionic cores (Extended Data Fig. 4b,c). The final band-alignment schematic plots are shown in Extended Data Fig. 4d,e.

Photodetector performance modelling. The temporal response ($\tau_t$) of a photodetector is modelled based on its dependence on the carrier transient time ($\tau_c$), RC-limited time response ($\tau_{\text{RC}}$) and diffusion time ($\tau_{\text{dif}}$) as follows.

$$\tau_t = \sqrt{\tau_c^2 + (\tau_{\text{RC}})^2 + (\tau_{\text{dif}})^2}$$

The transient time is predicted based on the bias across the sample ($V$), mobility ($\mu$) and film thickness ($d$).

$$\tau_c = \frac{d^2}{\mu V}$$

The RC constant is another limitation to the photodetector speed as it defines the time it takes to charge/discharge the capacitors in the electrical circuit. Assuming a simple parallel-plate capacitance model, the equation is derived as below.

$$\tau_{\text{RC}} = 2.2RC = 2.2\frac{AR\tau_0}{d}$$

In the calculations, we have assumed that the diffusion time is negligible when the drift and RC time constant, and the field is linear across the sample thickness.

Finally, the −3 dB bandwidth of a photodetector relates to the temporal response according to the following equation.

$$\text{BW} = 0.35\sqrt{\frac{2.2AR\tau_0}{d}^2 + \frac{d^2}{\mu^2 V^2}}$$

Using the Beer–Lambert law and assuming that the reflectance is negligible, the absorbance of a material relates to its transmittance ($T$), absorption coefficient ($a$) and thickness ($d$) as below.

$$A = 1 - T = 1 - e^{-ad}$$

The EQE is defined as the number of photogenerated carriers divided by the number of incident photons. The internal quantum efficiency (IQE) is defined as the ratio of the photogenerated carriers to the number of absorbed photons. Therefore, the division of these two quantities gives the percentage of absorbed photons (A). As a result, the following equation describes the relationship between the EQE and thickness of the photovoltaic layer.

$$\text{EQE} = \text{IQE} \times (1 - e^{-ad})$$

In this paper, for comparison between Si and Pb–Sn, it is assumed that IQE at 905 nm is equal to unity for both architectures (IQE=1.0). We note that assuming IQE of 0.7 to 1.0 does not change the conclusions of the model regarding the performance advance of Pb–Sn-perovskites over Si (Supplementary Fig. 10). To improve readability, a table containing the raw data obtained from device modelling is provided in Supplementary Table 3. In each column, we have referred to the equation listed in Methods.

Data availability

The datasets generated during and/or analysed during the current study are available from the corresponding author on reasonable request.

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Extended Data Fig. 1 | Performance modeling of the photodetector. Equations are provided in Methods section. **a**, Percentage of absorbed light vs. thickness at indicated absorption coefficients. EQE and bandwidth of **b**, Silicon and **c**, PbSn perovskite vs. film thickness. **d**, Comparison of silicon and PbSn perovskite from the last two panels. Parameters used for the modeling are given in Extended Data Fig. 2a.
Extended Data Fig. 2 | Combined speed × efficiency characteristic of the photodetector. a, Parameters used in the modeling of PbSn and Silicon photodetectors. b, Combined speed × efficiency modeling of silicon and PbSn PVK. c, Comparison between estimated performance and actual devices.
Extended Data Fig. 3 | Performance of NiOx vs PEDOT:PSS HTL. 

a, EQE comparison between PEDOT:PSS and NiOx using tin-powder reduced precursor (TPRP) strategy. 
b, Representative current density vs. voltage (JV curves) under the dark condition for the mentioned HTLs. Tin wire-reduced precursor strategy is used for all three shown cases. 
c, Comparison of dark current density at 50 mV for devices fabricated with PEDOT:PSS, NiOx, and without HTL with different Sn reducing strategy.
Extended Data Fig. 4 | Possibility of interfacial SnOx formation. a, Diagram comparing the formation energy of different form of oxide for Ni and Sn. 
Electrostatic potential (in eV) at the cores of oxygen atoms along the longitudinal direction of the b, DFT simulated structure of formed SnO layer on top of NiO HTL. c, NiO/SnO (001) and d, NiO/SnO2 (001) longitudinally lattice-matched supercell calculated by DFT-PBE within spin-polarized treatment for Ni.
In each subplot, the top are the atomic structures of a (001)-oriented NiO/(SnO/SnO2) superlattice; Ni atoms are shown in grey, Sn atoms are purple, and O atoms are red. Schematic plots of the band alignment for e, NiO/SnO and f, NiO/SnO2 interface. The bandgap values are obtained according to a bulk calculation with HSE06 functional. g, Energy level diagram for different layers of fabricated devices, including SnO.
Extended Data Fig. 5 | Citation for the data used in the manuscript. a, Fig. 3a; b, Fig. 3b; c, Fig. 4c. The numbers shown in the Figure, refer to the raw number in Extended Data Tables 1 and 2.
Extended Data Fig. 6 | Geometrical capacitance effect in photodetectors time response. Measured fall time of the pixels is plotted vs. pixel area (mm²). The area for direct probe connection is estimated with the extension of the capacitance-limited regime. The area could be smaller than the estimated one. Area effect on response time of silicon photodetector is also compared with PbSn. Silicon below 0.1 mm² reaches the plateau, and further decrease in the area does not decrease the response time. This behaviour indicates that the relatively poor carrier mobility limits the speed of Si PIN PDs.
Extended Data Fig. 7 | Impedance measurement of the PbSn device. a, Impedance plot measured in the dark under short-circuit conditions. b, Capacitance spectra obtained under the same condition as panel a. Black lines correspond to fits. c, Modified Randles equivalent circuit used for the fitting with Rs, Cg, and Cs, related to series resistance, geometrical capacitance, and interface accumulation capacitance and R2 and R1 as surface recombination resistance. d, Parameters obtained by fitting the impedance data with Randles equivalent circuit.

| Parameters | Fitted value |
|------------|--------------|
| Rs (Ω)     | 10.24        |
| R2 (Ω)     | 8.07 × 10⁻⁶  |
| R3 (Ω)     | 7.73 × 10⁻⁷  |
| Cg (F)     | 6.88 × 10⁻⁹  |
| Cs (F)     | 8.99 × 10⁻⁹  |
| ε          | 51           |
Extended Data Fig. 8 | Mapping of the distance using PbSn photodetector. a, The input for the mirror's position in the ToF setup shown in Fig. 5a for 105 individual measurements. The motorized stage controls the position of the mirror (distance in respect to the light source). The distance of zero is arbitrary, and it means that light is traveling the same distance as the reference pathway toward the InGaAs detector in the interferometer. b, Graph showing the response when the mirror has moved by 1.0, 3.0, 5.0, and 7.0 mm. For better visualization of peak positions, the response magnitude has multiplied by a power of 400. Averages and standard deviations of estimated depths based on the response time are shown on right side c, The output of the estimated distances using the peak position of PbSn photodetector response. The 3D visualization of these results is shown in Fig. 5d.
Extended Data Table 1 | Comparison of the solution-processed photodetectors

| # code | HTL/Absorber/ETL | Max EQE% @ λ > 850 nm | Min dark current (A/cm²) @ 50 mV | Fall time (ns) | Area (mm²) | Bias (V) | Stability | Ref. |
|-------|-----------------|------------------------|----------------------------------|---------------|------------|---------|-----------|------|
| 1     | PTAA/ MAPbI₃/ F8C:PTB7-Th/ C₆₀ | 54                      | 1×10⁻⁶                           | 5.6           | 0.1        | 0       | 50% (storage, 120h) | 50   |
| 2     | MoO₃/ PTBTT:PCBM/PEIE | 23                      | 8×10⁻⁵                           | 230           | 0.5        | 0       | -         | 51   |
| 3     | PEDOT:PSS/ PTT:PCBM | 17                      | 1×10⁻⁴                           | 62            | 1          | 0       | -         | 52   |
| 4     | MoO₃/ PbS-QD/ TiO₂ | 30 @ 0.5 V               | 1×10⁻⁴                           | 292           | -          | 0       | -         | 53   |
| 5     | PEDOT:PSS/ (FA₆Sh)₅b,MAPbI₃,₅/ C₆₀ | 64 @ 0.2 V              | 8×10⁻⁵                           | 9100          | 3.8        | 0.2     | 96% (storage, 90days) | 54   |
| 6     | PTAA/ MAPbI₃/ PDPPTDTPT:PCBM | 20 @ 0.2 V              | 4×10⁻⁴                           | 5             | 0.1        | 0.2     | -         | 55   |
| 7     | PbS-QD | 33                      | 1×10⁻⁷                           | 70            | 1.96       | 5       | -         | 56   |
| 8     | NiO₃/ PbS/ C69/ BCP/ Ag | 85⁺                    | 5×10⁻⁵                           | <0.2          | <0.001     | 0       | 96% (continuous illumination, 10 hours) | This work |

*EQE and speed was measured on the same device. EQE was measured on the pixels with a area of 0.049 cm². Time response was measured for a range of pixel areas as it is shown in Extended Data Fig. 6. Fastest measured response times are listed. The effect of device area on response time is shown in extended data Fig. 6. Response times in the capacitance-limited region (larger pixel area) do not reflect the charge carrier mobility or speed of carrier transfer rate across the device, as these response times are governed by the capacitance and resistance of devices. Details of measurement conditions and more references for stability are listed in Supplementary Table 4.
### Extended Data Table 2 | Comparison of commercial silicon photodetector

| code No. | Company   | Model     | R (A/V) @ 850 nm | EQE @ 850 nm | Speed (GHz) | Rise time (ps) | Dark (A) | Diameter (mm) | Area (cm²) | Current Density |
|----------|-----------|-----------|------------------|--------------|-------------|----------------|----------|---------------|------------|----------------|
| a1       | Thorlabs  | DET026A   | 0.38             | 0.55         | 2.0         | 230            | 3.5×10⁻¹¹ | 2.5×10⁻⁶     | 4.9×10⁻⁶   | 1.4×10⁻⁷       |
| a2       | Hamamatsu | S9055-01  | 0.25             | 0.36         | 2.0         | 250            | 2.0×10⁻¹² | 1.0×10⁻⁶     | 7.8×10⁻⁶   | 2.5×10⁻⁶       |
| a3       | Newport   | 818-BB-20 | 0.14             | 0.20         | 1.0         | 350            | 1.1×10⁻¹⁰ | 8.8×10⁻⁶     | 6.1×10⁻⁶   | 1.8×10⁻⁶       |
| a4       | Newport   | 818-BB-21 | 0.47             | 0.69         | 1.2         | 300            | 1.0×10⁻¹⁰ | 4.0×10⁻⁶     | 1.3×10⁻⁶   | 7.9×10⁻⁶       |
| a5       | Thorlabs  | PDA100A2  | 0.64             | 0.94         | <0.01       | <30000>        | <1×10⁻⁶  | 9.8           | 0.75       | <1.3×10⁻⁶      |