

## RESEARCH ARTICLE

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## Durable dielectric switching and photo-responsivity in a Dion–Jacobson hybrid perovskite semiconductor†

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Two-dimensional hybrid perovskite (2DHPs) semiconductors have shown great potential in multifunctional applications, especially dielectric switches and photoelectric detection owing to their diverse structural feasibility and excellent semiconducting features. However, most of the reported 2DHPs belong to the Ruddlesden–Popper (R–P) type, which unavoidably suffers from weak van der Waals interactions, degrading their reliability for long-term operation. The recently developed environmentally stable Dion–Jacobson (D–J) phase perovskites provide us with an opportunity to design 2DHPs with durable working performances. Here, we report a D–J phase perovskite material (BDA)MA<sub>2</sub>Pb<sub>3</sub>Br<sub>10</sub> (**1**, where BDA<sup>2+</sup> is 1,4-butanediammonium and MA<sup>+</sup> is methylammonium) that displays a remarkable switchable dielectric anomaly at 322 K. Meanwhile, the single crystal devices of **1** exhibit impressive photoelectric detection performances, featuring a large on/off current ratio ( $\sim 6.19 \times 10^3$ ), remarkable responsivity ( $\sim 11.79 \text{ mA W}^{-1}$ ), and high detectivity ( $\sim 2.9 \times 10^{11}$  Jones). Additionally, due to the phase stability conferred by the D–J structure, these devices exhibit excellent photoresponsivity and resistance to fatigue, as evidenced by over 1200 photoresponse cycles, along with noteworthy stability in dielectric switching (maintained for two months). This study presents a comprehensive methodology for designing highly stable D–J phase perovskites with exceptional performance in dielectric switching and photoelectric detection, thereby expanding the repertoire of 2D D–J multifunctional hybrid perovskites.

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## Introduction

Two-dimensional hybrid perovskites (2DHPs), with their structural diversity and composition tunability, have emerged as a hotspot of research in numerous fields, including solar harvesting, photoelectric diodes, ferroelectrics, and nonlinear optics.<sup>1–4</sup> In particular, in the region of phase transition materials, the dynamic synergistic effect between organic and inorganic components in 2DHPs significantly facilitates the

introduction of thermal-triggered ordered–disordered transitions.<sup>5–8</sup> This characteristic results in notable dielectric anomalies, making 2DHPs suitable for applications as thermal stimuli dielectric switching materials, offering novel capabilities such as temperature sensing, smart switching, and information processing.<sup>9–12</sup> Moreover, benefiting from the excellent semiconductor properties brought by the inorganic framework, including high optical absorption, defect tolerance, and long carrier lifetime, 2DHPs have also sparked great research enthusiasm in the field of photoelectric detection.<sup>13–18</sup> Therefore, 2DHPs are considered an ideal platform for integrating dielectric switching and photo-responsivity features, thereby promoting the development of multifunctional materials. For example, Sun *et al.* reported a series of 2DHPs that synchronously exhibited excellent photosensitivity and giant dielectric anomalies, showcasing their significant application value.<sup>19–21</sup> Despite these significant achievements, most reported dielectric switching 2DHPs are based on the Ruddlesden–Popper (R–P) type.<sup>22–25</sup> Structurally, the interlayer cations in R–P 2DHPs are loosely connected through weak van der Waals interactions, rendering them vulnerable to

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long-term working conditions and limiting their practical applications.<sup>26–28</sup> Thus, it is of strategic and scientific significance to explore durable dielectric switching 2DHPs.

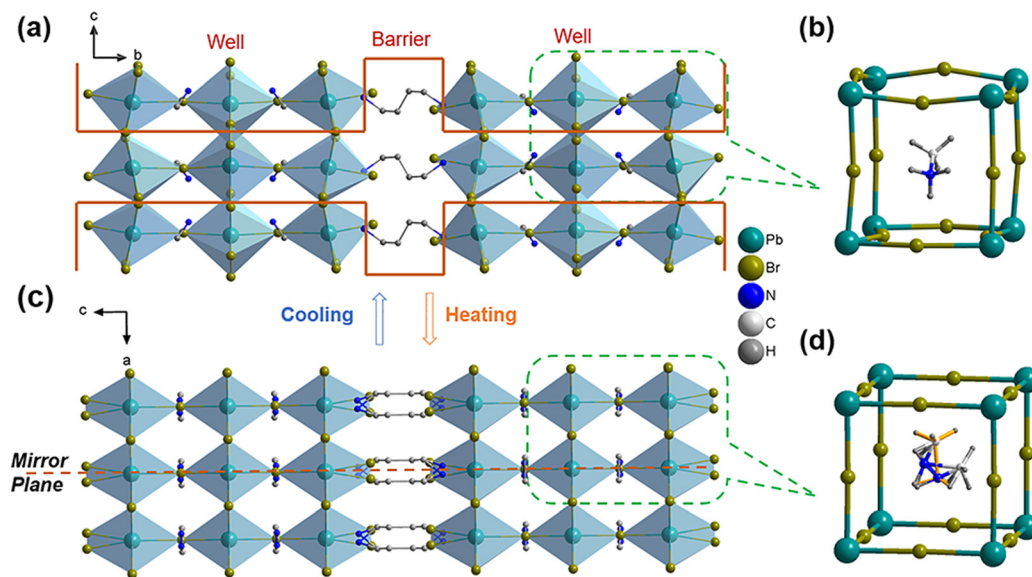
The recently developed Dion–Jacobson (D–J) hybrid perovskites provide us with an opportunity to obtain the desired materials. Unlike its R–P homolog, the introduction of diamine cations in the interlayer results in the elimination of the van der Waals gap, preventing the intervention of air and water molecules, thus improving the phase stability of the material<sup>29,30</sup> and largely extending its working reliability in photovoltaic applications. For example, Sajjad Ahmad *et al.* developed a series of D–J phase 2D perovskites (PDA)(MA)<sub>*n*–1</sub>PbI<sub>3*n*+1</sub> (*n* = 1–4), which exhibited ultrahigh device stability. The devices without encapsulation retain over 95% efficiency on exposure to harsh environments.<sup>31</sup> Although D–J type 2DHPs have made tremendous progress in photoelectric devices, the current research focus on D–J 2DHPs with dielectric switching features is limited. Therefore, exploration of D–J 2DHPs with dielectric switching features and high photoresponsivity is an urgent step in the development of environmentally stable multifunctional materials.

Here, we have synthesized a D–J hybrid perovskite (BDA)MA<sub>2</sub>Pb<sub>3</sub>Br<sub>10</sub> (**1**, BDA<sup>2+</sup> is 1,4-butanediammonium and MA<sup>+</sup> is methylammonium), which exhibits reversible dielectric switching at 322 K. Furthermore, **1** exhibits excellent photoresponsivity, including a large on/off ratio of  $6.19 \times 10^3$ , a high detectivity of  $2.9 \times 10^{11}$  Jones, and a remarkable responsivity of 11.79 mA W<sup>–1</sup>. Moreover, **1** exhibits excellent fatigue resistance, as demonstrated by over 1200 photoresponse cycles and outstanding dielectric switching stability (maintained for two months). This work paves the way for the design of highly stable 2D D–J perovskites with dielectric anomalies and photo-

electric detection performance, enriching the family of multifunctional dielectric switches.

## Results and discussion

The single crystals of **1** which are up to  $20 \times 6 \times 1$  mm<sup>3</sup> in size are grown by the temperature-cooled solution method (Fig. S1†), and the phase purity of the sample is confirmed by the powder X-ray diffraction pattern (Fig. S2†). The quality of the crystal face is qualified using a scanning electron microscope (SEM), as shown in Fig. S3.† The image demonstrates the flat and smooth surface, indicating the high quality of the single crystal, which is beneficial for high-performance photo-detection. In order to understand the structure–function relationship, single-crystal X-ray diffraction was conducted to determine the structural details of (BDA)MA<sub>2</sub>Pb<sub>3</sub>Br<sub>10</sub>. As is demonstrated in Fig. 1a, **1** adopts a 2D trilayer perovskite architecture, in which the trilayer inorganic frameworks consist of corner-sharing PbBr<sub>6</sub> octahedra, and the BDA<sup>2+</sup> cations are arranged in a staggered manner within the interlayer space, forming an organic–inorganic alternate quantum well structure. Such a structure is beneficial for the dissociation of photogenerated electron–hole pairs and results in a longer lifetime of free carriers, leading to an improvement in photoconductivity. For 2D R–P perovskite, the organic cations are loosely connected by weak van der Waals interactions, inhibiting long-term operation and limiting their practical application. In contrast, the diamine moieties in D–J phase perovskites are connected to the inorganic sheets by hydrogen bonds, eliminating the van der Waals gap interaction and improving the optoelectronic detection performance.<sup>32,33</sup>



**Fig. 1** (a) A schematic diagram of the quantum-well-like structure of **1** at LTP. (b) The inorganic framework, with the MA<sup>+</sup> cation incorporated inside the cavity at LTP. (c) Single crystal structures of **1** at HTP. (d) The inorganic framework, with the disordered MA<sup>+</sup> cation inside the cavity at HTP; the two folds of cations have been distinguished in different colours.

We further explore variable temperature structural analysis at the low-temperature phase (LTP) and high-temperature phase (HTP) to investigate the phase transition mechanism of **1**. At the LTP, **1** crystallizes in the *Pnma* space group (Table S1†). As is described in Fig. S4 and Tables S2 and S3,† the  $\text{PbBr}_6$  octahedra show a distorted geometric configuration with an average Pb–Br–Pb bond angle of  $132.78^\circ$  and Pb–Br bond lengths of 2.84–3.15 Å. The coupling of the ordered organic cations ( $\text{MA}^+$  and  $\text{BDA}^{2+}$ ) and distorting Pb–Br sheets can be considered as a driving force to trigger the phase transition as the temperature increases (Fig. 1b). At the HTP, **1** crystallizes in the *Cmcm* space group (Table S1†). Both the organic cations  $\text{MA}^+$  and  $\text{BDA}^{2+}$  become disordered and are highly symmetrical relative to the mirror plane (Fig. 1c and d). In addition, the inorganic sheets adopt a highly symmetric configuration, which is confirmed by the average Pb–Br–Pb bond angle, calculated to be  $179.976^\circ$  (Fig. S4 and Table S4†). It can be concluded that the order–disorder transition of the organic cations and inorganic framework is the key factor in the temperature-induced reversible phase transition, which further generates the dielectric anomaly.

In order to further investigate the reversible phase transition process of compound **1**, differential scanning calorimetry (DSC) measurement was performed and is shown in Fig. 2a, and a pair of reversible thermal peaks at 329 K and 322 K can be observed during the heating and cooling process. Meanwhile, the temperature-dependent dielectric constant test is recorded in Fig. 2b. A step-like dielectric anomaly is observed around 322 K during the heating and cooling process, consistent with the DSC results, indicating the occurrence of a reversible phase transition. Moreover, the dielectric constants at different frequencies (at 10 kHz, 100 kHz, 300 kHz, 500 kHz, and 1000 kHz) show similar dielectric anomalous behaviour during the heating processes (Fig. S5a†) and the dielectric anomaly of the *a*-axis is more significant than that of the *b*- and *c*-axes in the 1000 kHz axial directions (Fig. S5b†), evidencing the anisotropies of **1**. The fundamental optical and semiconductor characteristics are verified by the UV–visible absorption spectrum, showing a sharp optical absorption cut-off at  $\sim 550$  nm. The band gap is estimated to be 2.50 eV by the *Tauc* equation. This value is comparable with those of other two-dimensional lead–bromine perovskites,

such as  $(\text{BA})_2(\text{DMA})\text{Pb}_2\text{Br}_7$  ( $\sim 2.65$  eV) and  $\text{BA}_2\text{EA}_2\text{Pb}_3\text{Br}_{10}$  ( $\sim 2.55$  eV),<sup>34,35</sup> and is suitable for photodetection in the visible region. Furthermore, the photoluminescence (PL) of **1** is recorded with a narrow emission peak at 510 nm (Fig. S6†), indicating that the electron–phonon coupling effect of **1** is weakened, which is favorable for photoelectric detection. In addition, we performed first-principles DFT calculations to investigate the band gap of **1**, indicating that **1** is a direct bandgap semiconductor with the valence band maximum (VBM) and conduction band minimum (CBM) located at the same *k*-point G, and the band gap was calculated to be 2.540 eV, in good agreement with the experimental results. The partial density of states (PDOS) reveals the contribution of different atomic orbitals to the band gap. As shown in Fig. 3c, the VBM mainly stems from the Pb 6s and Br 4p states, while the CBM is mainly determined by the Pb 6p state. Therefore, the semiconductor properties of **1** are chiefly governed by the inorganic layers. In addition, Fig. 3d describes the temperature-dependent conductivities along the three axes (*a*-, *b*-, and *c*-). During the heating process, the curve shows an upward trend, which also indicates the semiconductor properties of **1**. In particular, the conductivity rises sharply as the temperature approaches 322 K, which agrees well with its phase transition temperature, further validating that the distortion degree of the inorganic framework in HTP is smaller than that in LTP, and the conductivity is improved. These results above reveal the excellent semiconductor properties of **1**, which provide potential for application in photodetection.

To investigate the photodetection capability of **1**, a single crystal-based bi-lateral device is shown in Fig. 4a. The dark current is measured as  $4.2 \times 10^{-10}$  A under a  $V_{\text{bias}}$  of 10 V. When the light intensity is  $35.88 \text{ mW cm}^{-2}$ , the photocurrent increases to a saturated  $I_{\text{photo}}$  of  $2.6 \times 10^{-6}$  A (Fig. 4b), and the on/off ratio is about  $6.19 \times 10^3$ . This value is at the same level as those of the reported two-dimensional multilayer perovskite photodetectors, such as  $(\text{i-PA})_2\text{CsAgBiBr}_7$  and  $(\text{ALA})_2(\text{EA})_2\text{Pb}_3\text{Br}_{10}$ .<sup>36,37</sup> Fig. 4c shows the light intensity depen-

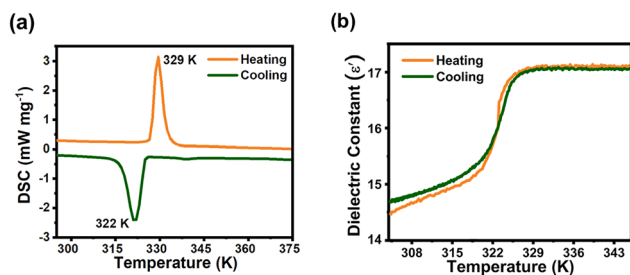


Fig. 2 (a) The DSC curve of **1** and (b) the dielectric constant curve of **1** at 1000 kHz.

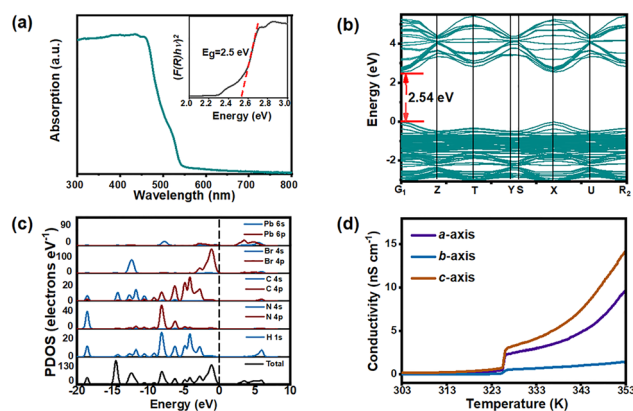
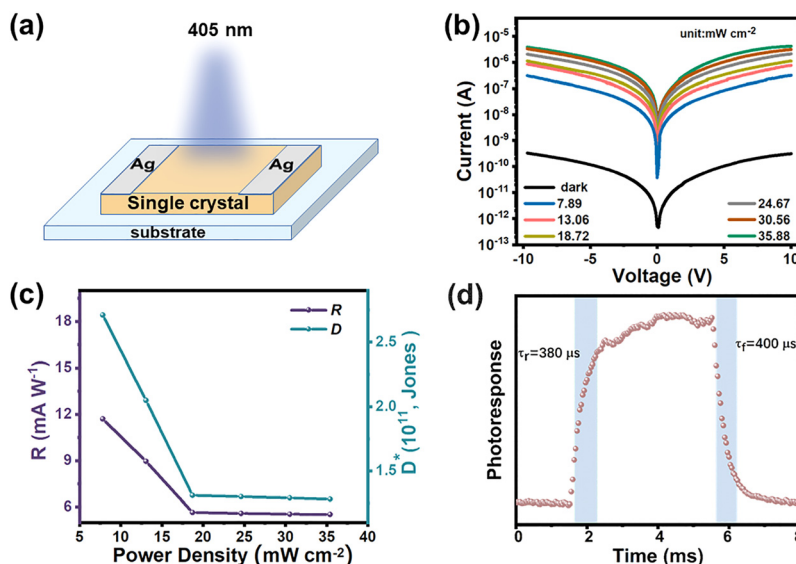


Fig. 3 (a) The UV-vis absorption spectrum of **1** (inset: the experimental bandgap). (b) The calculated bandgap diagram of **1**. (c) PDOS of **1**. (d) Variable temperature conductivity test along different axes of single crystals.

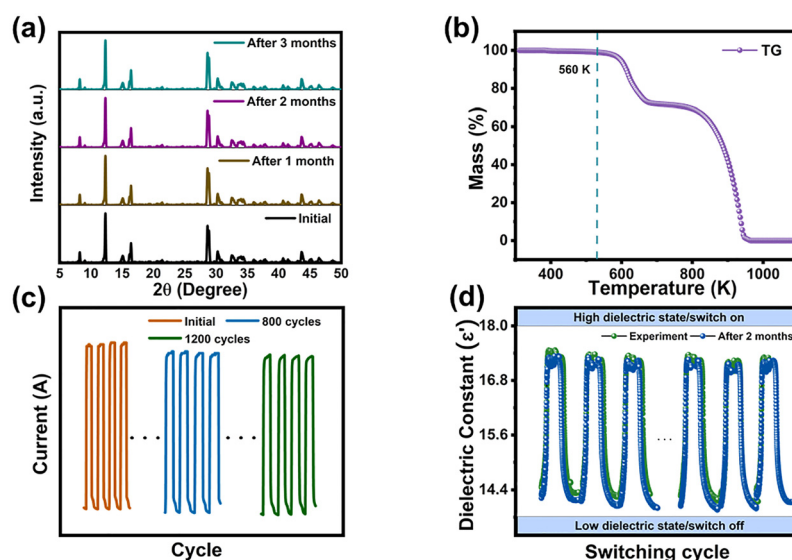


**Fig. 4** (a) The photodetection device schematic diagram of **1**. (b) The  $I$ – $V$  curves of **1** measured at 405 nm with different incidence powers. (c)  $R$  and  $D^*$  calculated from the photocurrent measured at 10 V bias. (d) The response time of **1**.

dent responsivity ( $R$ ) and detectivity ( $D^*$ ) of a single crystal device at 405 nm. And the maximum  $R$  and  $D^*$  are  $11.79 \text{ mA W}^{-1}$  and  $2.9 \times 10^{11} \text{ Jones}$ , respectively, comparable with those of other D–J-phase hybrid perovskites, such as (3AMPY)(EA)  $\text{Pb}_2\text{Br}_7$  and (HIS)(DMA)  $\text{Pb}_2\text{Br}_7$ .<sup>38,39</sup> The response time is another important parameter of photodetectors. As shown in Fig. 4d, the rise ( $\tau_r$ ) and decay ( $\tau_d$ ) photoresponse times are 380 and  $\sim 400 \mu\text{s}$ , respectively, during the switching “on–off” cycle, indicating that **1** is promising for future fast photoresponse devices.

To further explore the environmental stability and fatigue resistance, we evaluated the environmental stability of **1**, as

evidenced by the PXRD patterns of **1**, which are consistent with the initial results (maintained for three months). In addition, we analyzed the thermal stability of **1** by thermogravimetric analysis (TGA), as shown in Fig. 5b. The trace of **1** starts to decompose slowly at 560 K, which is much higher than its phase transition temperature. It is noteworthy that the temperature range from the phase transition temperature to the decomposition temperature is 345 K, indicating that **1** has high thermal stability as a dielectric switch. Such high environmental stability will be suitable for various application scenarios. To further investigate the high operating stability of the device, the durability of its dielectric switching and photo-



**Fig. 5** (a) The phase stability of **1** by the PXRD patterns. (b) TG curve of **1**. (c) The photoresponse capability of **1** after 800 and 1200  $I$ – $T$  cycles at 405 nm. (d) Periodic of repeatable switching measurement after several on/off cycles at 1 MHz.

responsivity were recorded. After 800 and 1200 photoresponse cycles at 405 nm (Fig. 5c), the saturated  $I_{\text{photo}}$  remains at a relatively stable level, indicating the favourable anti-fatigue properties of **1**. Dielectric cycling stability is another important parameter of stability. As revealed in Fig. 5d, the dielectric constant value remains constant after several dielectric cycles. Importantly, the device of **1** also exhibits excellent dielectric switches after exposure to air for 2 months. These features indicate that **1** has excellent environmental stability and excellent working stability, which will further enrich its application scenarios in multifunctional responsive materials.

## Conclusions

In summary, we have constructed a two-dimensional (2D) Dion–Jacobson (D–J) phase perovskite (BDA)MA<sub>2</sub>Pb<sub>3</sub>Br<sub>10</sub>, which exhibits obvious dielectric anomalies along different axes at 322 K. Additionally, **1** exhibits excellent photoelectric detection performance, with a large switching ratio of  $6.19 \times 10^3$ , a remarkable responsivity of  $11.79 \text{ mA W}^{-1}$ , and a high detectivity of  $2.9 \times 10^{11}$  Jones. In particular, based on the intrinsic structural stability of the D–J phase perovskites, single crystal devices of **1** reveal excellent thermal, environmental, and operating stability. The devices of **1** possess excellent photo-responsivity antifatigue merits (over 1200 photo-response cycles) and remarkable switching stability (maintained for two months). Our work presents a comprehensive methodology for ultra-stable optoelectronic applications and dielectric switches, which will inspire further intriguing research on multifunctional dielectric switches.

## Conflicts of interest

The authors declare no conflict of interest.

## Acknowledgements

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