Journal of Materials Chemistry A



PAPER

View Article Online
View Journal | View Issue



Cite this: *J. Mater. Chem. A*, 2025, **13**, 26364

ZnGa₂Te₄ thin-film absorbers for photoelectrochemical CO₂ reduction†

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Photoelectrochemical (PEC) carbon dioxide reduction reaction (CO₂RR) has been considered as a promising route to convert and store solar energy into chemical fuels. It is crucial to find suitable photoelectrode materials that are photo-catalytically active and exhibit excellent photochemical stability. One of the promising contenders is ZnTe with the \sim 2.26 eV band gap and prolonged stability under CO₂RR PEC conditions. Herein, a new telluride based thin-film ZnGa₂Te₄ photocathode with lower band gap and stronger visible light absorption compared to ZnTe is synthesized and characterized using a combinatorial sputtering technique. A two-step annealing method with excess Te supply is implemented to synthesize nearly stoichiometric ZnGa₂Te₄ absorber material with a zincblende-derived tetragonal crystal structure confirmed by synchrotron X-ray and electron diffraction. Theoretical calculations show that $ZnGa_2Te_4$ has suitable direct bandgap (~ 1.86 eV) and high absorption coefficient ~10⁵ cm⁻¹, in agreement with experimentally prepared films. Transient absorption spectroscopy reveals the biexponential decay dynamics, with time constants, $\tau_1 \sim 0.04$, and $\tau_2 \sim 0.65$ µs in microsecond time scales and provides the optical transition pathways for this semiconductor thin film. PEC measurements show that the ZnGa₂Te₄ photocurrent densities are comparable to the widely investigated ZnTe photocathodes or even surpass it under simulated sunlight condition. ZnGa₂Te₄ samples demonstrate promising photoelectrochemical stability, maintaining consistent performance under illumination. The inclusion of diaryliodonium additive substantially increases its CO_2RR selectivity to $\sim 60\%$. These findings open a new avenue for the synthesis of telluride-based thin-film photocathodes for further exploration and will motivate future research to integrate this potential photocathode material into PEC devices.

Received 11th April 2025 Accepted 7th July 2025

DOI: 10.1039/d5ta02891d

rsc.li/materials-a

Introduction

Photoelectrochemical (PEC) carbon dioxide reduction reaction (PEC CO₂RR) technology is one of the emerging pathways to advanced energy solutions. PEC CO₂RR based-on semiconductor photoelectrodes can convert solar energy into chemical fuels. This process mimics natural photosynthesis and it provides a solution to fulfill our world's increasing energy demand. To achieve PEC CO₂ reduction, it is important to find a photoabsorber with specific optical properties including appropriate band edge positions, robustness in harsh aqueous environments, and high activity for CO₂RR. A range of different semiconductor materials such as Si, oxides (Cu₂O,

Fe₂O₃, BiVO₄, CeO₂), metal organic frameworks (MOFs) and nitrides, *i.e.* ZnTiN₂ have been studied as potential photoelectrodes for PEC CO₂RR.⁵⁻¹⁰

Recently chalcogenide-based semiconductor materials have received attention due to their interesting photovoltaic characteristics. Cu(In,Ga)(S,Se)₂,¹¹ CZTS¹² and CuGa₃Se₅ ¹³ chalcopyrite photocathodes have shown promising results for PEC applications. The zinc blende phase of ZnTe has been tested and exhibits photocatalytic activity toward both H₂ evolution (HER) and CO₂RR.¹⁴ A recent study showed that tailoring the surface composition of ZnTe photocathodes could result in an improved interfacial charge transfer, effectively modifying its CO₂RR activity and selectivity before adding any electrocatalysts.¹⁵ The optoelectronic properties of sputtered ZnTe could be improved by chloride treatments and the measured faradaic efficiency (FE) can be increased using organic additives.¹⁶ However, the ~2.26 eV band gap of ZnTe is wider than ideal for the operation as a photocathode, calling for new material searches.

There have been several computational screening studies for materials that may be suitable as cathodes for CO₂RR. A recent

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[†] Electronic supplementary information (ESI) available. See DOI: https://doi.org/10.1039/d5ta02891d

study identified 17 telluride photocathodes among 52 candidates with excellent aqueous stability under operational conditions.17 Another theoretical screening study emphasizes the promising CO₂RR activity of tellurium-containing semiconductors and suggests Te-based photo absorbers possess more thermodynamically favorable conditions to convert CO2 to chemical fuels.18 Based on the optical properties and considering excitonic effects, 4 telluride photocathode materials that are suitable for visible-light photocatalysis have been shortlisted, including GaTe, Cd(GaTe₂)₂, LiInTe₂, and Zn(GaTe₂)₂.¹⁹ Most of these tellurides have been synthesized using the Bridgman technique, CVT and also solid state reaction for powder and bulk characterization.20-22

Herein, we develop a synthesis route of tellurium based thinfilm photocathodes ZnGa₂Te₄ using combinatorial sputtering techniques. We demonstrate that adjusting the cation (Zn, Ga) to anion (Te) ratio, facilitate a phase transition from cubic $(F\bar{4}3m)$ to tetragonal $(I\bar{4})$ crystal symmetry during postdeposition annealing. Notably, excess of Te in as-deposited film is required to form stable tetragonal ZnGa₂Te₄ structure after annealing. DFT calculation show that ZnGa2Te4 is direct bandgap semiconductor (bandgap 1.86 eV) with high absorption coefficient (105 cm-1) that are consistent with optical measurements and suitable for effective sunlight absorption compared to ZnTe (2.26 eV band gap). Transient absorption (TA) spectroscopy at time scales from 100 fs through 10 μs shows the photoexcited carrier kinetics of the ZnGa₂Te₄ photocathode with a promising carrier lifetime for the photoelectrode applications. Photoelectrochemical (PEC) analyses are conducted, revealing that ZnGa₂Te₄ exhibits performance comparable to other chalcogenide-based semiconductors such as ZnTe benchmark photoelectrode. Finally, the ZnGa₂Te₄ films show a promising photo-current density with efficient fuel production and inclusion of diaryliodonium additive suppresses HER and significantly improves CO2RR to become the major reaction. Notably, ZnGa₂Te₄ maintained stable photocurrent and consistent product formation during PEC stability test.

Results and discussion

 $(ZnTe)_x(Ga_2Te_3)_{1-x}$ materials family

Among these predicted tellurides, several contain Zn, Ga and Te, together forming a $(ZnTe)_x(Ga_2Te_3)_{1-x}$ family of materials with related tetrahedrally bonded crystal motifs. ZnTe, a II-VI compound, is usually formed in a cubic, zinc blende structure (Fig. 1a). Here, divalent Zn and hexavalent Te form an octet. On the other hand, Ga₂Te₃, containing trivalent Ga, is also based on tetrahedral atomic coordination motifs, but one third of the cation sites, i.e. gallium in Ga₂Te₃ are empty, generating disordered vacancy sites on the zinc blende lattice (Fig. 1c).23 The II-III₂-VI₄ compounds which are the focus of this paper, *i.e.* ZnGa₂Te₄, can be derived from these binary ZnTe and Ga₂Te₃ compounds. For example, ZnGa2Te4 has been observed in a vacancy-ordered zincblende-derived structure (Fig. 1b). Based on this observation, we propose that there exists an octetconserving compositional tie-line, $(ZnTe)_x(Ga_2Te_3)_{1-x}$ $(0 \le x \le x \le x)$ 1), where the relationship, built upon the zinc blende lattice, shown in Fig. 1d is conserved. This sum of valence electrons from Zn ([Ar] 3d¹⁰4s²), Ga ([Ar] 3d¹⁰4s²4p¹), and Te ([Kr] 4d¹⁰5s²5p⁴) is conserved at 96 valence electrons, and can be demonstrated through the following equations:

$$12 \text{ ZnTe} = 12 \text{ Zn(II)} + 12 \text{ Te(VI)} = 96 \text{ valence electrons}$$

$$3 \operatorname{ZnGa_2Te_4} = 3 \operatorname{Zn(II)} + 6 \operatorname{Ga(III)} + 3 \operatorname{V_{Cation}} + 12 \operatorname{Te(VI)} = 96 \text{ valence electrons}$$

$$4 \text{ Ga}_2\text{Te}_3 = 8 \text{ Ga}(\text{III}) + 4 \text{ V}_{\text{Cation}} + 12 \text{ Te}(\text{VI}) = 96 \text{ valence electrons}$$

The II-III₂-VI₄ compounds can be characterized by tetrahedral cation-to-anion co-ordination, as an ordered vacancy compounds (Fig. 1b). Whenever all cation sites are tetrahedrally coordinated, tetragonal defective structures are formed.24 However, despite known structures, this suitable candidate material in this telluride family has never previously been experimentally investigated for CO2RR, and its further investigation is desirable.

Thin films of $(ZnTe)_x(Ga_2Te_3)_{1-x}$ (where, x = 0 to 1) were synthesized on EXG glass substrates by RF combinatorial cosputtering using ZnTe, Ga₂Te₃ and Te target precursors (see Experimental section). Deposition was carried out in an Ar atmosphere, with a substrate heater setpoint temperature of 200 °C. The whole combinatorial film was post-annealed at 500 °C by rapid thermal annealing (RTA) under atmospheric nitrogen pressure. Throughout this paper, we refer to these combinatorial films as $(ZnTe)_x(Ga_2Te_3)_{1-x}$, as they span the zincblende-derived compositional tie-line (Fig. 1d). However, note that Te concentration is a free variable which is at first in

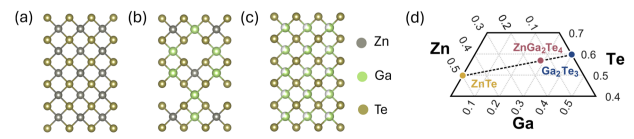


Fig. 1 Crystal structure (a) zinc blende derived ZnTe (b) tetragonal ZnGa₂Te₄ (c) defective Ga₂Te₃ (d) chemical formulas and ternary phase diagram showing the compositional tie-line between zincblende-derived Zn-Ga-Te materials $(ZnTe)_x(Ga_2Te_3)_{1-x}$ thin film growth.

excess and then after annealing pins close to the targeted composition on the tie-line. A cubic to tetragonal phase transition has been observed upon changing cation-anion composition.

Fig. S1† shows the results of X-ray diffraction (XRD) patterns of as-deposited (ZnTe) $_x$ (Ga $_2$ Te $_3$) $_{1-x}$ thin films with varying cation composition, measured by X-ray fluorescence (XRF). According to the XRD results, the as-deposited (ZnTe) $_x$ (Ga $_2$ -Te $_3$) $_{1-x}$ films (deposited at $T_{\rm dep} \sim 200$ °C) formed an amorphous/nanocrystalline Te phase with a wide range of elemental compositions (Zn/(Zn + Ga) = 0.20–0.95, Te $\sim 80\%$) (Fig. S1†). No Ga $_2$ Te $_3$ or other secondary phases were observed in this growth condition. However, it is important to note that, when no extra Te target source was used, the as-deposited films become amorphous and peeled off on further annealing (Fig. S2†).

To convert the amorphous precursor into the product with desired zincblende-derived structure, we post-annealed the asdeposited film. After post-annealing the films at 300 °C for 5 minutes, partial crystallization was observed, characterized by the emergence of zinc blende ZnTe-like phases, although the Te phase remained predominant (Fig. S3†). Notably, no distinct Ga-containing secondary phases were detected at this stage. However, when the annealing temperature was increased to 500 °C, a structural transformation was observed, resulting in zincblende-derived ZnGa₂Te₄ structure as the main phase. Formation of a well-crystallized, phase-pure ZnGa₂Te₄ at higher

annealing temperatures enhances structural ordering which is crucial for improved photoelectrochemical performance. Postannealing helped to transform amorphous/Te nanocrystal structure into phase-pure cubic and tetragonal (ZnTe)x(Ga2- Te_3 _{1-x} (Fig. 2a and b), depending on the anion and cation ratio. Within the composition range of Zn/(Zn + Ga) = 0.65-0.85 and \sim 40% Te composition, all reflections in the XRD pattern can be indexed to cubic $F\bar{4}3m$ symmetry. The tetragonal zincblendederived structure in (ZnTe)x(Ga2Te3)1-x is observed across a range of cation stoichiometries ranging from 0.35-0.65, with Te fraction \sim 50%. When the Zn/(Zn + Ga) ratio reaches \geq 0.8, an unidentified peak emerges at approximately $2\theta \approx 35^{\circ}$ (Fig. 2a – upper panel). It is interesting to see the tetragonal structure still emerges even though the annealed combinatorial films are Tedeficient. The fact that the Te concentration pre-and postannealing straddles the tie-line of interest suggests that further iterative process optimization can be used to fine-tune the film composition. Formation of a well-crystallized, phasepure ZnGa₂Te₄ at higher annealing temperature enhances structural ordering which is crucial for improved photoelectrochemical performance.

The deposition temperature ($T_{\rm dep}$) was varied from 200 °C to 300 °C (maintaining other deposition parameter same) to control the composition of volatile Zn and Te elements. The 200 °C growth temperature was optimal for achieving better crystallinity for the post-annealed ZnGa₂Te₄ thin film (Fig. S4†).

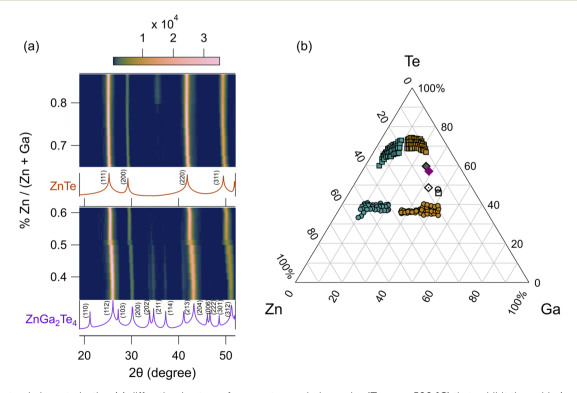


Fig. 2 Structural characterization (a) diffraction heatmap from post annealed samples ($T_{\rm growth} \sim 500$ °C) that exhibit the cubic (upper panel)—tetragonal (lower panel) crystal symmetry ($\lambda = 1.54$ Å) (b) ternary phase diagram showing the full composition of films (filled square and circle shape represents as-deposited sample and annealed samples, respectively) on EXG glass. Filled diamond shape (magenta) represent nominal stoichiometric ZnGa₂Te₄ composition. Other symbols represent the ZnGa₂Te₄ composition synthesized on FTO glass for photoelectrochemical measurement.

This improvement is attributed to better composition control during sputter deposition.

Due to the high vapor pressure of Te, loss of Te increases with increasing deposition temperature. At deposition temperature of $T_{\rm dep} \sim 300$ °C, significant Te loss was observed during film growth (Zn: Ga: Te $\sim 12.9: 28.3: 58.7$), compared to $T_{\rm dep}$ \sim 200 °C (Zn: Ga: Te \approx 6.3: 11.6: 81.9; Table S1†). This Te loss during deposition impacted the film crystallinity. At 300 °C, the crystallinity of ZnGa2Te4 film decreased, with amorphous/nano crystalline grains (Fig. S5c†). However, at $T_{\rm dep} \sim 200-250$ °C, the excess Te promoted diffusion and nucleation kinetics, enabling the formation of larger grains (~120 nm; Fig. S5a and b†) with significantly improved crystallinity for vacancy-ordered zincblende-derived structure.

To further demonstrate the photoelectrochemical measurement, a uniform, stoichiometric, and phase-pure tetragonal film was prepared on FTO coated soda lime glass. To grow a uniform film on FTO glass, the deposition temperature was further decreased to 180 °C, which helps to maintain near stoichiometric condition for the post annealed samples. (Fig. 2b - filled diamond shape (grey)). A few calibration experiments were required (Fig. 2b open shapes, Table S2†) before achieving this final composition.

Structural characterization

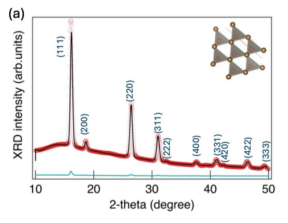
To determine the phase purity of cubic and tetragonal structure, post-annealed combinatorial films were characterized by synchrotron grazing incidence wide angle X-ray scattering (GIWAXS) and integrated to generate a powder diffraction pattern. The experimental data and a LeBail whole pattern refinement against the reported $F\bar{4}3m$ and $I\bar{4}$ structure is presented in Fig. 3a and b, respectively, alongside the two crystal structure models. The fit returns lattice parameters for cubic structure of a = 6.09639 Å and for tetragonal structure of a =5.94994 Å, c = 11.98607 Å.

To better understand the structural properties of ZnGa₂Te₄ thin films deposited on EXG glass substrate, (scanning) transmission electron microscopy ((S)TEM) analysis was carried out on a specimen prepared by focused ion beam (FIB) milling. Integrating across five selected area electron diffraction (SAED) patterns acquired from different locations along the TEM lamella (Fig. 4a) didn't produce an obvious ring pattern due to the relatively small sampling volume of SAED. However, 4D scanning transmission electron microscopy (4D-STEM) analysis was conducted to construct a virtual SAED pattern (Fig. 4b) by summing thousands of diffraction patterns across the film. This virtual SAED pattern shows more defined diffraction rings that align well with the ZnGa₂Te₄ structure, especially when comparing the azimuthally-integrated 1D experimental diffraction pattern with the simulated diffraction pattern from the ZnGa₂Te₄ structure (Fig. 4c). The (112), (224) and (204) planes of the tetragonal structure reflections can be assigned from 4D-STEM and are consistent with the XRD results. STEM energy-dispersive X-ray spectroscopy (EDS) was also performed to provide spatially resolved elemental information about the presence of Zn, Te and Ga elements within the film with large crystallized grains (Fig. 4d).

The composition of the film is as expected for ZnGa₂Te₄, though there is also an increased presence of oxygen at the film surface that could be attributed to oxidation occurring over time due to ambient exposure (Fig. S6†). Note that the tungsten (W) signal towards the film surface is from the protective W layer deposited during FIB preparation. There is also Ga present in the protective W layer, which comes from Ga implantation from the Ga-ion beam used for FIB preparation. We do not expect a significant amount of Ga implantation in the film or substrate because the final thinning of TEM lamella down to electron transparency was conducted using an Ar-ion beam. Quantitative EDS linescans (Fig. S6†) indicates an absence of Ga signal at film surface. Fig. 4e shows the top view scanning electron microscopy (SEM) image of corresponding ZnGa₂Te₄ absorber.

Electronic band structure calculations and optical property measurements

To gain atomistic insights into ZnGa₂Te₄ properties, we have performed first-principles calculations of ZnGa₂Te₄ electronic



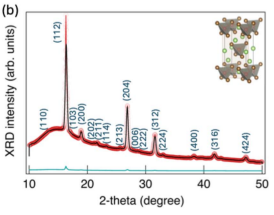


Fig. 3 Wide-angle X-ray analysis integrated wide-angle X-ray scattering pattern along with a LeBail fit to the (a) cubic Ga:ZnTe and (b) tetragonal $ZnGa_2Te_4$ structure (here, black, red and cyan color represent calculated, experimental and difference spectrum of the refinement spectra) (λ 0.97625 Å)

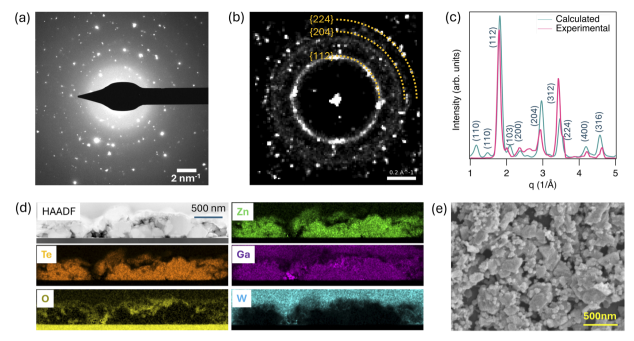


Fig. 4 Electron microstructure (a) TEM SAED pattern, (b) 4D-STEM virtual SAED (Bragg vector map) from the ZnGa₂Te₄ film, (c) experimental 1D diffraction pattern azimuthally integrated from 4D-STEM along with the simulated diffraction pattern, (d) STEM HAADF image and corresponding EDS elemental maps and (e) SEM top view of ZnGa₂Te₄ thin films.

structure using the screened hybrid functional of Heyd-Scuseria-Ernzerhof (HSE06). As shown in Fig. 5a, the calculated band structure and density of states confirm that ZnGa2Te4 is a nearly direct bandgap semiconductor, with a bandgap of 1.86 eV. A strong presence of Te (p) states in the occupied region indicates that Te (p) orbitals play a dominant role in shaping the valence band. Meanwhile, the overlap of Te (p) and Ga (s) states near the Fermi level in the unoccupied region suggests that both elements contribute to the conduction band. This interaction is crucial for charge carrier dynamics, as it facilitates electron transition into the conduction band, which could influence the electronic and optoelectronic properties of the material. For ZnTe, both Te (p) and Zn (s) states contribute significantly to the valence and conduction bands (Fig. S7†). Fig. 5b shows that the computed optical absorption coefficient (α) of ZnGa₂Te₄ exceeds 10⁵ cm⁻¹, with an absorption onset at \sim 1.8–1.9 eV, making it an efficient solar absorber.

Further structural, optical, and PEC characterization has been conducted to evaluate the suitability of ZnGa₂Te₄ as a photocathode, comparing it with well known ZnTe films. Raman spectroscopy has been performed on cubic ZnTe/FTO and tetragonal ZnGa₂Te₄/FTO film, to further understand the structural changes from cubic to tetragonal phase conversion. As shown in Fig. 6a, for ZnTe, the peaks observed at 205, 410, 619 and 824 cm⁻¹ could be assigned to first, second, third and fourth order longitudinal optical (LO) phonon scattering,

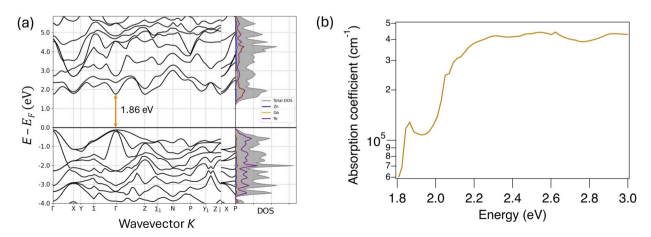


Fig. 5 DFT calculation (a) electronic band structure of ZnGa₂Te₄ with total and element-resolved density of states (b) theoretically calculated absorption coefficient (α) for ZnGa₂Te₄.

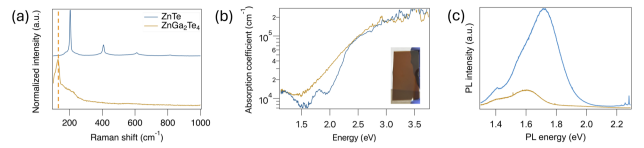


Fig. 6 Optical characterization (a) Raman scattering – dashed line indicates the A1 vibrational mode for tetragonal ZnGa₂Te₄ (b) absorption coefficients ($\alpha(E)$) spectra (inset is a photograph of uniform ZnGa₂Te₄ thin film on FTO substrate) (c) room-temperature PL spectra.

respectively, which correspond to the cubic ZnTe structure.²⁵ However, when Zn/(Zn + Ga) ratio reaches to \sim 0.35 with Te fraction \sim 47%, most of the LO phonon mode disappears. As ZnGa₂Te₄ structure resembles with tetragonal ZnGa₂Se₄ structure and in ZnGa₂Se₄, the most dominant A1 mode appears at 142.8 cm⁻¹ for ZnGa₂Se₄, while for ZnGa₂Te₄ tetragonal structure, A1 mode appears \sim 126.0 cm⁻¹.²⁶ The substitution of Se with Te should cause a shift of the peaks toward lower wavenumbers, primarily due to the incorporation of the heavier Te atom replacing Se atom.

The absorption spectra of ZnTe/FTO and ZnGa₂Te₄/FTO thin films were measured to investigate their optical properties. In Fig. 6b, $ZnGa_2Te_4$ shows absorption edges within \sim 1.7-1.8 eV lower that 2.26 for ZnTe. The absorption coefficient is 10⁵ cm⁻¹, which is consistent with calculated results (Fig. 5b) and comparable to that of ZnTe. A defect center luminescence appeared around 1.70 eV for cubic ZnTe (Fig. 6c). This center luminescence in ZnTe could be attributed to a V_{zn}- or some other donor defect complexes.27 Interestingly, 3 sharp peaks are identified in the high energy region ~2.26 eV in PL spectra of ZnTe, which could be assigned as resonance Raman mode, as we observed longitudinal phonon modes in Raman spectra (Fig. 6a). These enhanced peaks could be interpreted as the multi-longitudinal optical phonon bands of cubic $F\bar{4}3m$ structure enhanced by the near-bandedge PL band.28 However, no resonance peak was observed and PL peak emission was less intense for ZnGa₂Te₄ structure (Fig. 6c). The shift in the ZnGa₂Te₄ PL peak could be attributed to defect-induced emission. This less intense luminescence can be explained with the thicker ZnTe (\sim 350 nm) film compared to ZnGa₂Te₄ film (\sim 140 nm) (Table S2†).

Photogenerated carrier dynamics

To monitor the charge carrier dynamics of ZnGa₂Te₄/FTO film, TA spectroscopy has been performed across a range of time scales from picoseconds to microseconds. Fig. 7a and b shows TA spectra for different delay times for ZnGa₂Te₄ films on the picosecond and microsecond time scales. There is a fast initial evolution within 1 ps which may indicate ultrafast trapping but could also be thermalization of photoexcited carriers to the band edge.²⁹ Beyond 1 ps, ZnGa₂Te₄ has a consistent spectrum throughout the 5 ns window, and even up to 1 μs. The TA spectra from 1–500 ps exhibits one broad absorbance peak

centered at \sim 1.80 eV, corresponding to the direct band to band transitions in ZnGa₂Te₄.³⁰ This peak decays substantially on the picosecond time scale to reveal a photoinduced absorption with increased intensity at high energies. This evolution could be the resultant effect of trapped charges in various states with energies from deep state to band edge. Trap-mediated electron–hole recombination might be a significant loss pathway for ZnGa₂Te₄ based semiconductor which should be further optimized.

TA kinetics were extracted from both the pico- and micro second experiments and stitched together (Fig. 7c). Kinetics were extracted at a probe energy of 1.62 eV, 1.87 eV and 2.30 eV. At probe energy of 1.62 and 1.87 eV, they display similar kinetic decay trends with a complex decay kinetics, requiring triexponential function ($\tau_1 \sim 0.38 \pm 0.026$, $\tau_2 \sim 19.9 \pm 0.935$, and $\tau_3 \sim 566 \pm 32.87$ ps) to fit (Table S3†). These complex multiexponential components can be explained by trapmediated pathways with different trap depths. This τ_1 and τ_2 can be correlated with the rapid relaxation of photoexcited electrons from the band edge to exciton-mediated trap states, while τ_3 corresponds to the decay of dissociated free carriers occurring on a nanosecond timescale.31,32 The 0.38 ps decay in 1.62 eV is correlated with a growth at 1.87 eV. Although these features both appear as a positive ΔA , the growth at 1.87 eV may be related to loss of the ground state bleach which is superimposed on a photoinduced absorption, explaining the inverted shape of this feature. The signal at 2.30 eV does not decay in the picosecond window but matches dynamics at 1.87 eV after 1 ns (1000 ps). On the microsecond time scale, decay traces obtained at 2.30 eV are fit to two components ($\tau_1 \sim 0.04 \pm 0.006$, $\tau_2 \sim 0.65$ \pm 0.3 μ s) (Table S4†).

Specifically, carrier lifetimes for ZnTe thin-film absorbers, which have been heavily researched, are similar. For our ZnTe sample, the transient absorption kinetics (spectrum not shown), at a probe energy of 1.70 eV also required fitting with a tri-exponential function, on picosecond time scale, yielding lifetimes of $\tau_1 \approx 5.5 \pm 0.6$, $\tau_2 \approx 61 \pm 7.5$, and $\tau_3 \approx 854 \pm 97$ ps, which are comparable to those observed in our ZnGa₂Te₄ films. Xiong *et al.* demonstrated that ZnTe films grown on FTO substrates by thermal evaporation exhibit complex charge carrier relaxation dynamics. Transient absorption spectroscopy (TAS) measurements, with decay traces recorded at 800 nm, were fitted using a tri-exponential model, yielding lifetimes of $\tau_1 \approx 2.6 \pm 0.2$, $\tau_2 \approx 24.5 \pm 2.5$, and $\tau_3 \approx 363 \pm 32$ ps.³³ It is

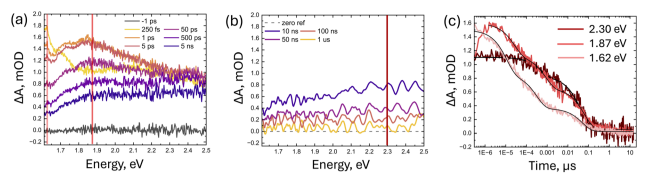


Fig. 7 Carrier dynamics transient absorption spectra of $ZnGa_2Te_4$ photoexcited at 2.48 eV (100 nJ per pulse) at pump-probe delays spanning 250 fs-5 ns (a) and 10 ns-1 μ s (b) TA kinetics measured at 2.30 eV (dark red), 1.87 eV (red), 1.62 eV (light red). Energies where kinetics were taken are indicated by the appropriately colored lines in panels (a) and (b). Kinetics were stitched together from the picosecond and microsecond experiments to show dynamics from 100 fs-10 μ s (c).

important to study and optimize these photoexcited carrier relaxation process of $ZnGa_2Te_4$, to enhance the migration of charge carriers for photocatalytic reactions.

Photoelectrochemical properties

Mott-Schottky analyses at four different frequencies were carried out in CO₂-bubbled 0.1 M KHCO₃ solution (pH 6.8) to experimentally characterize the energy band positions of ZnTe and ZnGa₂Te₄, and the resulting plot is shown in Fig. 8a. The negative slope in the Mott-Schottky plot indicates both ZnTe and ZnGa₂Te₄ are p-type semiconductors. Flat band potential $(E_{\rm Fb})$ is estimated from the x-axis intercepts of Mott-Schottky plot and is determined to be 1.07 V and 0.7 V vs. Ag/AgCl (or -5.71 and -5.34 eV) for ZnTe and ZnGa₂Te₄, respectively. For a p-type semiconductor, the flat band potential approximates the valence band edge potential (VB) under flat band conditions and these measured VBs agree well with the theoretically predicted ones.17 With the known band gap energy discussed earlier, the conduction band (CB) edges are calculated to be -3.45 and -3.48 eV for ZnTe and ZnGa₂Te₄, respectively (Fig. 8b). According to the reported potentials for CO₂R, ¹⁷ the CB position of ZnTe and ZnGa2Te4 are less negative than the electrochemical CO₂ conversion to CH₄ (-4.2 eV), CH₃OH (-4.04

eV), HCHO (-3.96 eV), CO (-3.91 eV), and HCOOH (-3.83 eV). While these band positions can cause simultaneous competing hydrogen evolution reaction (HER), the two photocathodes investigated in the present work are capable of reducing CO₂ under illuminations.

The chopped (dark/light) linear scanning voltammetry (LSV) scans of the ZnGa₂Te₄ and ZnTe thin-film photo cathodes are presented in Fig. 9a, b, S8a and b,† respectively. The HT-ANEC instrument with fiber-optic-coupled photodiodes for front side illumination is used to evaluate the performance of photocathodes investigated in this study.34 To achieve higher photocurrents for accurate and reliable quantifications of CO2RR product, we used a high energy LED at 455 nm (\sim 2.70 eV within the visible light region) to compare these two photocathodes. ZnTe and ZnGa₂Te₄ compounds displays similar photocurrent over the entire operating potential range and reach a photocurrent density close to -2 mA cm^{-2} at -1.0 V vs. reversible hydrogen electrode (RHE) (Fig. 9a and S8a†). As the bandgap of these two are determined to be \sim 1.86 and \sim 2.26 eV, respectively, and the energy band position characterized by Mott-Schottky above shows both are suitable for PEC CO₂R and HER, the 2.70 eV LED has photon energy higher than the bandgaps of both ZnGa₂Te₄ and ZnTe, which could contribute to their

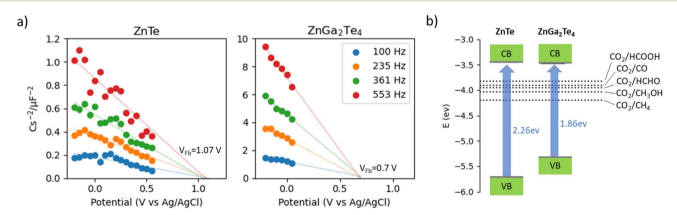


Fig. 8 Mott–Schottky analysis (a) energy band position characterization for ZnTe and ZnGa₂Te₄. Measurements are carried out in CO₂-bubbled 0.1 M KHCO₃ (pH = 6.8) and at multiple frequencies to validate the measured flat band potentials. (b) Band potentials and bandgap energy for ZnTe and ZnGa₂Te₄.

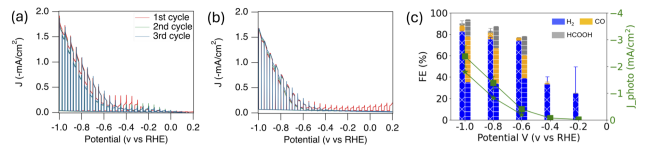


Fig. 9 Photoelectrochemical performance CV results under 455 nm LED illumination (a) without additive and (b) with additive, and (c) CAs at multiple potentials without additive (1st bars; truncated) and with additive (2nd bars; square) for ZnGa₂Te₄ photocathodes.

similar photocurrents. For ZnGa₂Te₄, however, only the near stoichiometric but slightly Te-rich sample shows both high photocurrents and negligible dark currents. Other offstoichiometric, Te-poor samples do not show such promising PEC results (Fig. S9 and Table S2†). It is noted that all samples, including both ZnTe and ZnGa2Te4 have a reductive current between -0.25 and -0.6 V vs. RHE, but present only in the 1st cycle of the cathodic sweep. These reductive currents could be caused by reduction of residues on the surface (e.g., ZnO_x, GaO_x, or TeO_r).

CV scans with a white LED (Doric dual LED, LEDC2-W35_SMA, Fig. S10†) providing broad spectrum illumination was also used to compare the performance of ZnGa2Te4 and ZnTe. In contrast to illumination under 455 nm (2.70 eV), ZnGa₂Te₄ which possess a narrower bandgap (ZnGa₂Te₄ vs. ZnTe: 1.86 vs. 2.26 eV) shows significant photocurrents over the entire CV scan range (Fig. S11†). The additional lower energy photons from the white LED compared to 455 nm LED (i.e. >455 nm) suggest ZnGa₂Te₄ absorbs visible light below the bandgap energy of ZnTe.

Chronoamperometry (CA) at multiple potentials following cyclic voltammetry (CV) was conducted to reveal products formed (Fig. 9c and S8c† - left side bars). The products accumulated during CA measurements were analyzed by GC and HPLC in the HT-ANEC system. Results show that both ZnTe and ZnGa₂Te₄ produce only H_2 at lower applied bias (less than -0.4 V vs. RHE). At higher applied bias (> $-0.6 \text{ V} \text{ } \nu \text{s}$. RHE), both photocathodes start to produce C1 products (including carbon monooxide, CO and formate, HCOOH) in addition to H2. ZnGa2Te4 shows FE of CO₂RR less than 10% while ZnTe sample shows >20%. Ga has been reported to be a HER catalyst. 35,36 In contrast, Zn is known as an efficient catalyst for reducing CO₂ into CO and HCOOH. More specifically, Zn-rich surfaces on ZnTe have been reported to not only facilitate the charge transfer but also acts as an electrocatalyst to enhance the selectivity for carbon products.³⁷ The substitution of Ga for Zn in ZnGa₂Te₄ could be responsible for the decrease in CO₂RR and increase in HER selectivity. Nevertheless, increasing applied bias increases CO2RR selectivity on both photocathodes.

The effects of organic additives

Motivated by our previous study, 10 mM additive diaryliodonium has been included in the electrolyte for ZnTe and

ZnGa₂Te₄ CO₂RR investigations. ^{13,16} The first cycle of CV scans shows reductive currents at very positive potentials (Fig. 9b and S8b†). This could result from polymerization/deposition of additives onto the electrode surface, since they don't occur in subsequent cycles. Due to illumination by 2.7 eV LED, which is higher than the bandgaps of both materials, ZnTe and ZnGa₂-Te₄ show similar photocurrents in the presence of additives, as observed above with no additives present. The additive, however, results in slightly decreased photocurrents compared to the no additive environment for both electrodes.

When the additive diaryliodonium is present in the electrolyte, ZnTe shows a moderate enhancement of CO₂RR selectivity at the potentials where CO2RR occurred along with suppression of HER (Fig. S8c† - right side bars). This change agrees with previously reported effects of additives on ZnTe PEC CO2RR performance.16 Interestingly, the addition of additive greatly increases FE of CO₂RR including CO and HCOOH on ZnGa₂Te₄ from \leq 10% to \sim 50% with suppression of HER (Fig. 9c - right side bars). This CO₂RR selectivity enhancement upon addition of additives makes its performance comparable to that of ZnTe under 455 nm (i.e. 2.70 eV) illumination conditions. The photocurrent values and corresponding product distributions for each CA experiment are summarized in Table S5.†

The organic additive has been shown to reduce and dimerize/oligomerize into a nonconductive layer on the electrode surface to enhance CO₂RR product selectivity.³⁸ It was reported that the diaryliodonium additive may help increase CO₂RR selectivity not just by suppressing HER but could further increase CO2RR catalytic activity. It boosts the partial current density of CO₂RR sufficiently to maintain total current density not only for dark electrolysis on Cu-based electrodes but also for PEC on photocathodes ZnTe. 13,39,40 Mechanisms proposed to account for the change in selectivity include slow diffusion of proton carriers to the electrode, lower H₂O and increased CO₂ concentration within the films, nanostructuring of the electrode, and interactions of CO2 reduction intermediates on the electrode with the film.41,42 For the diaryliodonium additive specifically, rotating-disk-electrode (RDE) studies from Watkins et al.39 indicate that the film acts as a barrier toward the diffusion of proton sources to the electrode surface, supporting that decreased proton availability near the electrode surface suppresses HER. The grafted film from diaryliodonium grows perpendicular to the electrode surface and creates a low-density

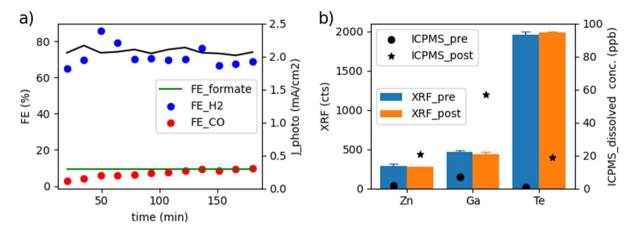


Fig. 10 (a) Stability test of $ZnGa_2Te_4$ without additives using a traditional H-cell at -0.8 V vs. RHE with periodic headspace measurements. Note that formate shown in the figure is the average over entire measurement. (b) Corrosion study by ICP-MS and XRF for pre-and post-PEC run samples.

film with channels, enabling facile substrate transport through the film and maintaining catalytic activity. The observed slightly lower photocurrents from our tests could be due to the attenuation of incident light intensity by the additives.

The reduced bandgap of $ZnGa_2Te_4$ (i.e. \sim 1.86 eV), which is in the range needed for PEC applications should facilitate visible light absorption from the solar spectrum. SEM image shows grain size $\sim 120~\text{nm}$ for ZnGa_2Te_4 but for ZnTe grain size less than 60 nm (Fig. S12a and b†). Larger grain size facilitates efficient charge transport, with less recombination, thereby enhancing photoelectrochemical (PEC) performance. Also, we determined roughness of the (ZnTe, ZnGa₂Te₄)/FTO film using AFM, which shows the root-mean-square (RMS) surface roughness of the ZnGa2Te4 and ZnTe film around ~11.2 nm and \sim 9.7 nm, respectively (Fig. S13a and b†). These properties contribute to efficient PEC activity of ZnGa2Te4 (Fig. 9 and S11†). With the assistance of organic additives to increase its CO₂RR selectivity, CO₂RR performance of ZaGa₂Te₄ becomes comparable to, and even surpasses, that of the widely investigated ZnTe.

Long-term photoelectrochemical stability

Inductively coupled plasma mass spectrometry (ICP-MS) was performed on post-reaction electrolytes to quantify corrosion from the electrode into solution. X-ray fluorescence (XRF) measurements on the electrode surface were performed preand post-PEC to confirm the changes in elemental molar loadings of the electrode. Fig. S14† shows ZnTe does not show noticeable corrosion under the conditions used for the PEC measurements. XRF confirms the pre- and post-run composition (Table S6†) barely changes, in agreement with previous reports. Table S6† barely changes, in agreement with previous reports. As each photocathode was first tested at -0.2 V, the relatively high concentrations of elements seen at -0.2 V is likely due to reduction of surface oxides. Nevertheless, increasing Ga corrosion was observed with increasing applied bias. Theoretical prediction suggests that this ZnGa₂Te₄ could

be stable up to -0.5 V vs. RHE. The HT-ANEC measurement is 15 min at each potential.

To further understand how this corrosion of Ga could influence the performance of ZnGa₂Te₄ in PEC CO₂R, we conducted a longer PEC measurement using a conventional H-cell setup without additive at -0.8 V vs. RHE. To produce photocurrent densities consistent with prior HT-ANEC measurements with the white LED, and to assure sufficient photocurrent for reliable product quantification, a mercury-xenon lamp calibrated for 3-sun illumination intensity was used (Fig. S10b†). Fig. 10a shows that the photocurrent of ZnGa2Te4 remained stable over the entire measurement. The product selectivity generally agrees with that from HT-ANEC measurements at this potential, with H₂ as the major product and with minor CO and formate product formation. It is noted that formic acid could only be analyzed after the longer duration measurement concluded and the value shown is the average over the entire test period. CO increases initially and levels out at \sim 10%.

ICP-MS, as well as XRF characterization, was carried out after the H-Cell PEC measurement (Fig. 10b). Similar to HT-ANEC measurements, corrosion occurs, especially for Ga, and this loss may be responsible for the slight increase in CO production at the beginning of the measurement. Interestingly, the dissolved metal concentrations including Zn, Ga, and Te from this 180 min measurement are very similar to those of the 15 min HT-ANEC measurements. This suggests the corrosion mostly occurs at the beginning of PEC and the dissolved metals concentration at the given electrochemical potential and pH reaches equilibrium,⁴³ consistent with the observed relatively stable photocurrents and product formation.

Conclusion

By introducing the Te rich synthesis route, we have successfully synthesized thin films of a telluride material, ZnGa₂Te₄ with tetragonal structure, as a photocathode for PEC CO₂RR. The structural, optical, and photoelectrochemical properties of the tetragonal ZnGa₂Te₄ films were systematically investigated. Its

suitable direct bandgap of 1.86 eV and relatively high carrier lifetime measured by transient absorption spectroscopy makes this thin film absorber a promising candidate for photocathode applications. With a stable current density of approximately -2mA cm⁻², this material is an interesting candidate for further study. There are many opportunities to improve this ZnGa₂Te₄ photocathode performance by varying the structure, morphology and defect properties, we demonstrate that its CO₂RR selectivity can be strongly enhanced by organic additives. Overall, this work demonstrates the successful synthesis of ZnGa₂Te₄ thin film as a potential photoelectrode for PEC CO₂R application.

Experimental section

Synthesis

The Zn-Ga-Te phase space was explored using combinatorial radio-frequency (RF) cosputtering methods. The chamber base pressure was 1×10^{-8} torr prior to depositions. During depositions, the pressure was maintained at 2.7 mTorr with 16 sccm Ar. Substrates were clamped to platens heated by a serpentine SiC resistance coil. Following a period of pre-sputtering, the combinatorial alloy films were deposited for 90 minutes at 200 ° C on EXG glass.

 $(ZnTe)_x(Ga_2Te_3)_{1-x}$ thin films were synthesized by radio frequency (RF) co-sputtering from alloyed 50.8 mm diameter ZnTe and Ga₂Te₃ targets, and one Te target, with ZnTe and Ga₂Te₃ power density of 0.74 and 2.22 W cm⁻², respectively, and Te 0.49 W cm⁻² in a Ar gas environment. All the films were annealed in RTA at 500-550 $^{\circ}\text{C}$ in N_2 environment for 5 minutes. However, to grow uniform film on EXG and FTO, substrate temperature was fixed 200 and 180 °C, respectively. Target power density was optimized for uniform ZnGa₂Te₄

ZnTe films were grown by RF sputtering of ZnTe target and deposited on FTO glass substrate. Substrate was deposited at 170 °C temperature for 70 min, and followed by annealing at \sim 435 °C.

Characterization

XRD/XRF. Combinatorial X-ray diffraction (XRD) measurements were conducted on a Bruker D8 diffractometer using Cu K_α radiation and a 2D detector. Patterns were integrated to generate an intensity vs. 2θ pattern. Synchrotron grazing incidence wide angle X-ray scattering (GIWAXS) measurements were performed at beamline 11-3 at the Stanford Synchrotron Radiation Lightsource, SLAC National Accelerator Laboratory. The data were collected with a Rayonix 225 area detector using a wavelength of $\lambda = 0.97625$ Å, a 3° incident angle, a 150 mm sample-to-detector distance, and a beam size of 50 µm vertical × 150 μm horizonal. The diffraction images were calibrated with a LaB₆ standard and integrated with the Nika SAS package. X-ray fluorescence (XRF) measurements were performed using a Rh anode at 50 keV. GIXRD data (Fig. S4†) were collected using the high-throughput, automated grazing incidence XRD beamline 2-1 ($\lambda = 0.729 \text{ Å}$).

SEM. Scanning electron microscopy (SEM) was operated by using a Hitachi S-4800 with accelerating voltage 3 kV and a working distance of 7 mm. Scanning electron microscopy (SEM) was operated by using a Hitachi S-4800 with accelerating voltage 3 kV and a working distance of 7 mm. (Scanning) transmission electron microscopy ((S)TEM) high-angle annular dark-field (HAADF) images and selected area electron diffraction (SAED) patterns were acquired with a Thermo Fisher Scientific Spectra 200 transmission electron microscope operating at an accelerating voltage of 200 keV. Specimens for TEM were prepared from deposited films via in situ focused ion beam lift-out methods44 using a Tescan Solaris Ga focused ion beamscanning electron microscope (FIB-SEM) workstation. Elemental mapping was performed in STEM mode using the Super-X energy-dispersive X-ray spectroscopy (EDS) system equipped with four windowless silicon drift detectors, allowing for high count rates and elemental sensitivity (down to 0.5-1 at%). The EDS data were quantified using a multi-polynomial parabolic background and absorption correction in Velox. 4D-STEM datasets were acquired with a 0.565 mrad convergence angle and approximate probe size of 1.5 nm. Dataset processing and analysis was done using the py4DSTEM python package45 and Cornell Spectrum Imager46 in Fiji47 to generate virtual selected area electron diffraction patterns from specific regions of interest and index these patterns to crystallographic phases.

AFM. Atomic force microscopy (AFM) images were obtained by using an AFM (Bruker) in the non-contact mode with a silicon AFM probe. The cantilever was driven under a resonant frequency of \sim 330 kHz and 0.4 N m⁻¹ spring constant.

Raman/PL. PL and Raman measurements were performed using the Renishaw inVia confocal system by using the 532 nm laser line and the $20 \times long$ working distance (0.5 mm) objective. The scattering light from the sample was directed by a grating with 1800 lines per mm for Raman or 600 lines per mm for PL prior to the CCD detector.

Transient absorption spectroscopy. Transient absorption (TA) measurements were collected in a standard pump-probe configuration using a Ti:Sapphire regenerative amplifier with an 800 nm fundamental and 90 fs pulse width at a repetition rate of 1 kHz. For ultrafast (fs-ns) experiments, the pump and probe paths are split at the outset and the pump (2.48 eV) is generated via sum frequency generation in an optical parametric amplifier (Light Conversion, TOPAS-C). The pump passes through a chopper spinning at 500 Hz to modulate the pump on/pump off cycle. The probe is focused into a sapphire crystal to generate white light via supercontinuum generation. The probe is delayed relative to the pump using a mechanical delay stage. Pump and probe are focused and spatially overlapped at the sample and changes in the absorption spectrum are monitored in a fiber-coupled CMOS spectrometer. Ultrafast data is collected using Helios software provided by ultrafast systems. For microsecond experiments, the same pump pulse is used, but the white light is created through continuum generation in a diode-laser pumped photonic crystal fiber and digitally delayed relative to the pump (EOS, Ultrafast Systems). Data analysis was conducted using Surface Xplorer to chirp or time zero correct the data and then further analyzed in Origin (OriginLabs).

Photoelectrochemical measurement (PEC). 0.1 M KHCO $_3$ electrolyte, with or without 10 mM diaryliodonium additive, 455 nm light-emitting diode (LED) illumination at 11 mW cm $^{-2}$ was used for this study. PEC chopped-light cyclic voltammetry (CV) was conducted at 10 mV s $^{-1}$ sweeps from +0.2 to -1.0 V ν s. RHE.

A 455 nm LED (Thorlabs M455F3) was selected as a major light source in this report because (1) it has higher energy than the band gap of both ZnGa₂Te₄ and ZnTe is in the range of visible light, not ultraviolet, and (2) could generate greater photocurrents for more accurate product quantifications by GC and HPLC. A LED with white light (intensity \sim 6.94 mW cm⁻²) spectrum was also used in some cases (Fig. S10†) to distinguish the PEC performance between the two materials in the broad visible light regions. We performed multipotential measurements (by CA) following CV (i.e. CA and CV was on two different sample spots) to investigate the CO₂ RR product distribution. We carried out CA measurements in the order -0.2, -0.4, -0.6, -0.8, and -1 V vs. RHE for 15 min (900 s) unless the film mechanically delaminates. The light was on for the first 882 s and then shut off for the rest 18 s. The ending 18 s under the dark provides the baseline for calculating J_photo and J_FE and can be used to evaluate if dark currents exist due to any film delaminations (the substrate is FTO and could show electrochemical activity at the potential range used in this report). The high-throughput analytical electrochemistry (HT-ANEC) instrument³⁴ we used to test CO₂RR performance requiring rapid flow to generate suitable and reproducible mass transport conditions. The rapid flow could cause the predeposited additive film to delaminate over time. For the tests with additive, the electrolyte contained additives at all times to ensure that the film remains on the electrode; hence, the thickness of diaryliodonium is not directly controlled. For this initial study, we focus on the discovery of the combined improvement in selectivity and activity. The impact of diaryliodonium thickness on stability and selectivity optimization will be the subject of future work. At the end of each (photo)electrolysis, we sampled gaseous and liquid products by the robotic sample handling system and was analyzed by gas chromatography (GC; Thermo Scientific TRACE 1300) and high-performance liquid chromatography (Thermo Scientific UltiMate 3000), respectively. Detailed product detection methods are can be found at our previous publication.34

For PEC CO_2 reduction measurements in an H-Cell, the cathode had an area of 1 cm 2 . CO_2 saturated 0.10 M potassium bicarbonate without additive was used as the electrolyte. A Pt foil anode was used behind a bipolar membrane (Fumasep® FBM single film, Fumatech) membrane. A leakless Ag/AgCl electrode was used as a reference electrode. A mercury–xenon lamp calibrated for 3-sun illumination intensity was used (Fig. S10b†). Carbon dioxide was provided to the electrochemical cell at a flow rate of 5 sccm using an Alicat flow controller. The gas exhaust stream of the electrochemical cell was passed through a liquid trap before flowing through the gas

sampling loop of an Agilent 7820a GCMS/TCD with an Alicat flow meter connected to its exhaust.

ICPMS. Inductively coupled plasma mass spectrometry (ICP-MS) by Thermo Fisher Scientific iCAP™ RQ instrument was used to determine the concentration of dissolved metals in electrolytes used for photo-electrochemistry.

Mott–Schottky analysis. BioLogic Potentiostat (SP-200) was used to conduct Mott–Schottky experiments. The electrolyte used was a $\rm CO_2$ -bubbled 0.1 M KHCO $_3$ solution with pH = 6.8. Mott–Schottky analysis was carried at multiple frequencies (2 kHz, 1 kHz and 0.5 kHz) at applied potential scan from 1 to -1 V νs . Ag/AgCl.

Density functional theory (DFT) calculations. To investigate the electronic and optical properties of ZnTe and ZnGa₂Te₄, spin-polarized Density Functional Theory (DFT) calculations were performed using the Vienna Ab initio Simulation Package (VASP)48 and accelerated through the circuit of Materials Project automation software: atomate2, Fireworks, Custodian, and Jobflow. 49-51 The crystal structure for both ZnTe and ZnGa2Te4 was extracted from the Materials Project database. 52 A double relaxation procedure was applied to obtain the ground state crystal structure of ZnTe and ZnGa2Te4. The projector augmented wave method was used to model the electronic wavefunctions and the Perdew-Burke-Ernzerhof (PBE)53 functional was used to approximate the exchange-correlation energy. The Brillouin zone integration was carried out using the tetrahedron method with Blöchl corrections. A plane-wave basis set was constructed with a kinetic energy cutoff of 680 eV. For geometry optimizations, a k-spacing of 0.22 was used for k-space sampling. Electronic energies were converged to within 1 \times 10⁻⁵ eV, and structural relaxations were performed using the conjugate gradient algorithm until all atomic forces were reduced below -0.02 eV Å^{-1} . Then, the PBE-optimized geometries were further optimized at the meta-GGA level using the r2SCAN functional using the atomate2 relaxation sets.50,54 To compute the full band structure of ZnTe and ZnGa2Te4, a static calculation was performed using the Heyd-Scuseria-Ernzerhof (HSE) hybrid functional, which offers the necessary accuracy in band gap calculations despite its increased computational expense.55,56 To determine the optical properties of ZnTe and ZnGa₂Te₄, we calculated the frequency-dependent dielectric functions with the independent-particle approximation with the same double-relaxed crystal structures used in the previous HSE calculation.57 The real and imaginary components of the dielectric functions were then used to compute the optical absorption spectra of the material.

Data availability

The data supporting this article have been included as part of the ESI.†

Author contribution

Conceptualization, S. Q., S. R. B., and A. Z.; methodology, S. Q., S. R. B., and A. Z.; investigation, S. Q., Y. C. L., M. G., J. M., W. Z., R. Y., M. A., C. P. M., S. R. B.; writing—original draft, S. Q.

writing—reviewing and editing, S. Q., Y. C. L., M. G., J. M., W. Z., S. R. B., A. Z. and all co-authors; funding acquisition A. Z., K. P., J. H.; supervision A. Z.

Conflicts of interest

The authors declare no conflict of interest.

Acknowledgements

This work was authored in part by the National Renewable Energy Laboratory for the U.S. Department of Energy (DOE) under contract no. DE-AC36-08GO28308. Funding provided by the Liquid Sunlight Alliance, which is supported by the U.S. Department of Energy, Office of Science, Office of Basic Energy Sciences, Fuels from Sunlight Hub under award number DE-SC0021266. Use of the Stanford Synchrotron Radiation Lightsource, SLAC National Accelerator Laboratory, is supported by the U.S. DOE, Office of Science, Office of Basic Energy Sciences under Contract No. DE-AC02-76SF00515. Thanks to Dr Nicholas Strange for support with GIWAXS measurements. The views expressed in the article do not necessarily represent the views of the DOE or the U.S. Government.

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