Metal halide based organic-inorganic perovskite semiconductors have attracted a great deal of attention for their optical and electrical properties [1–13]. Three-dimensional (3D) perovskites have recently been used to produce solar cells with efficiencies of up to 15% [1–4], while two-dimensional (2D) perovskites have recently been used to produce solar cells with 10% efficiency [5–7]. The structure of a typical 2D lead iodide (PbI2) perovskite is shown in Fig. 1(a), consisting of alternating layers of corner-sharing PbI6 octahedra and interdigitating RNH3 molecules (where R is an organic moiety). Excitons are formed and trapped in the inorganic layers, where the reduction in dimensionality (quantum confinement) and low refractive index organic layers (dielectric confinement) lead to binding energies in excess of 200 meV [5]. Therefore, such perovskite semiconductors exhibit strong excitons at room temperature, as seen in the sharp strong absorption and photoluminescence peaks [Fig. 1(b)]. These materials are easily processed from solution [6] and can thus be incorporated into a variety of nanostructures. In addition, due to their high binding energy and oscillator strength, these perovskites are ideal candidates for the production of new mixed light-matter states at room temperature as a result of strong coupling [7–12].

Recently there has been much interest in the mixed states of excitons and surface plasmon polaritons (SPPs), where the spectral properties of plasmonic gratings: “photonic” modes caused purely by interference from the periodicity of the structure, and “plasmonic” modes where SPPs also interact with the diffracted light. We present evidence for “image biexcitons” within organic-inorganic perovskite-coated silver gratings. These composite quasiparticles are formed by the interaction between an exciton and its image in the metal mirror below, with binding energy 100 meV at room temperature. By changing the polar and azimuthal angles of the incident light, we observe strong coupling between excitons and surface plasmon polaritons on the grating, with Rabi splittings of 150 and 125 meV for the exciton and biexciton, respectively. A detailed analysis of the field polarizations and dipole orientations shows how these Rabi couplings arise from the strongly modulated field volume.

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FIG. 1. (Color online) (a) Schematic of the 2D lead iodide perovskite structure. (b) Absorption and photoluminescence spectra of the CHPI thin film on a glass substrate at room temperature. (c) SEM image (top) and AFM profile (bottom) of the Ag grating. (d) AFM image (top) and profile (bottom) of the CHPI-coated grating. (e) Schematic cross section of the CHPI-coated Ag grating structure.

present rules out any influence from modified CHPI assembly in the grooves, which are in any case hundreds of times larger than the PbI layer spacing. In addition, the exciton diffusion length in 2D perovskites is of order 10 nm [24], therefore we do not expect any limiting effects due to the grating geometry. We note slight changes in the CHPI coverage alter the positions and intensities of dispersive grating modes [cf. Fig.3(a), with a thinner CHPI coating], however, the exciton modes remain essentially unchanged.

It is well known that the emitted energy of a dipole (exciton) is lowered when placed in front of a metallic surface due to interactions between the dipole and the reflected electromagnetic field [25–30]. Using the method of images, we can replace the metal and describe instead the coupling between an exciton in the CHPI ($\epsilon_1$) and its image exciton in the metal ($\epsilon_2$), modified by their respective dielectric environments. These spin-coated CHPI QWs are universally parallel to the substrate surface, as shown by x-ray diffraction [6,12], and Chance et al. [28] showed the redshift in the emitted energy of an exciton ($\Delta E_{\text{ex}}$) oriented parallel to the interface can be approximated by

$$\Delta E_{\text{ex}} \sim \frac{1}{l^3} \left( \frac{\epsilon_2 - \epsilon_1}{\epsilon_2 + \epsilon_1} \right) q \Gamma_0,$$

where $l$ is the distance between the exciton and a metal surface, $k_1$ is the wave number of light in CHPI, $q$ is the quantum yield of CHPI excitons (taken here to be 1), and $\Gamma_0$ is the inverse exciton radiative lifetime without the metal. Similar to the appearance of excitons in the spectra, we expect to observe such coupled “image biexcitons” as minima in the reflectivity, at a wavelength that differs from the uncoupled exciton according to Eq. (1). The strength of the coupling between the exciton and reflected electromagnetic field depends on the exciton dipole moment, which is controlled by the term $q \Gamma_0$. From this we can see the $l^{-3}$ dependence of the redshift, as shown in Fig. 2(d), where the experimentally observed $\Delta E_{\text{ex}} \sim 100$ meV corresponds to $l \sim 22$ nm, close to the experimentally determined CHPI thickness. Clearly $\Delta E_{\text{ex}}$ is also affected by the dielectric response of CHPI and Ag, and from Eq. (1) we see that $\Delta E_{\text{ex}}$ is maximized if $\epsilon_2 + \epsilon_1 \rightarrow 0$, i.e., when emission is resonant with an SPP on the metal-dielectric interface. The linewidth of the exciton is also affected by interactions with image charges in the metal, however, in our perovskite system this effect is not dominant due to tight planar confinement of excitons. We expect larger effects in systems that are less perfectly 2D, such as semiconductor heterostructures and J-aggregate systems, where surface charges play a much larger role.

The role of the SPP in this case is to outcouple the signal of the redshifted exciton. There are three main decay channels for dipole emission near a metal surface: direct emission to photons, emission to SPPs, and nonradiative processes such as the excitation of electron-hole pairs and lossy surface waves on the metal. Other nonradiative paths via defects or phonons are independent of $l$ and will be ignored in this analysis. Emission into SPPs provides an extra radiative decay channel as this signal can be extracted to the far field via the periodic nanostructure, and this mechanism has been used to improve the luminescence efficiency of light-emitting
devices (LEDs) [31,32]. The relative decay probability for each process is calculated as a function of $l$ [30] and shown in Fig. 2(d). Although these calculations are intended for SPPs propagating on planar metal surfaces, we can use them as approximations for our grating system, although we note such estimates are indeed expected to become less accurate with increasing structure depth. Up to a CHPI thickness of 25 nm, SPP-mediated emission is the most important radiative decay channel, with a maximum emission probability at 22 nm, matching the experimentally observed $\Delta E_{ex}$. Even for thicker CHPI films we expect the exciton modes to remain at the same positions, because SPP emission becomes weak at large $l$ where $\Delta E_{ex}$ is negligible.

In practice, we observe a range of bie exciton energies in our spectra (Figs. 2 and 3). There is a clear overall decrease in the reflectivity compared to Ag-only gratings (see the Supplemental Material [33]) for the wavelength range 490–550 nm, which coincides with the redshifted energies we would expect for the excitons in our 25 nm film according to Eq. (1). However, in all cases we still see the strongest signature from excitons that are 15–25 nm from the Ag interface, which have the largest SPP emission probability [Fig. 2(d)]. This corresponds to a wavelength of 490–510 nm, where we observe our second dip. Furthermore, the line shape of the bie exciton resonance does not resemble that of a single oscillator, which is due to the superposition of excitons from this range of distances.

In our MQW perovskite system, localized excitons in periodically spaced nearby QWs are optically coupled together to form many collective exciton-polariton states, each of which has an average distance $l$ from the Ag surface [34–37]. Therefore, in CHPI-coated Ag gratings, we observe both in-plane exciton polaritons, and of-plane interactions that lead to image bie excitons, which are outcoupled through SPP emission with a binding energy of 100 meV at room temperature [Fig. 2(e)]. For our grating system, the exciton and SPP modes become closer in energy with increasing $\phi$ (see below and Fig. 3), and, as a result, splitting between the exciton modes (indicated by the arrows in Fig. 3) increases to around 185 meV at $\phi = 90^\circ$. The azimuthal dependence of the exciton splitting reflects the tunable modification of the Coulomb interaction in this geometry, but, however, requires further theoretical development.

Besides the strong excitons, more dispersive grating modes can also be seen in the TM reflectivity scans of CHPI-coated Ag gratings (Fig. 3). Due to momentum and energy conservation, the dispersion of such grating modes (see the Supplemental Material) is given by

$$k_m^2 = k_i^2 \sin^2 \theta + G_m^2 \pm 2k_i G_m \sin \theta \cos \phi,$$

where $k_m$ is the wave vector of the measured grating mode, and $k_i$ is the wave vector of the incident light taking into account the refractive index of CHPI [33]. In our spectra we observe the $m = \pm 1$ plasmonic modes (gray dashed lines), and as these become resonant with the exciton and image exciton, the light-matter modes strongly couple and produce an anticrossing in the reflectivity of 0.25 eV. Extracting the mode positions from the $\phi = 90^\circ$ scan [Fig. 3(d)] allows them to be fit to a three oscillator model using the Hamiltonian

$$\hat{H} = \begin{pmatrix} E_{ex} & 0 & \Omega_{ex}/2 \\ 0 & E_{bx} & \Omega_{bx}/2 \\ \Omega_{ex}/2 & \Omega_{bx}/2 & E_{pl} \end{pmatrix},$$

where $E_{ex}$, $E_{bx}$, and $E_{pl}$ are the energies of the exciton-polaritons, image-bie excitons, and plasmonic grating modes, respectively, while $\Omega_{ex}$ and $\Omega_{bx}$ represent the interaction between the SPP and exciton/image bie exciton. From this we find Rabi splittings of $\Omega_{ex} = 150$ meV and $\Omega_{bx} = 125$ meV. These are greatly enhanced because of the large confinement of the plasmonic optical field in the thin PbI QW layers. The Rabi splitting is given by $\Omega \propto \sqrt{f_{osc} N_{QW} / V}$, where the oscillator strength ($f_{osc}$) of the CHPI is assumed to be similar for coupling to photons or plasmons, the number of QWs ($N_{QW}$) is proportional to the CHPI thickness, and the mode volume ($V$) is here proportional to the optical mode size. Comparing to Fabry-Pérot planar CHPI microcavities in strong coupling [12] which have a CHPI thickness of 72 nm, a cavity length of 407 nm, and a Rabi frequency of $\Omega_{FP} = 65$ meV, the simple scaling above predicts $\Omega_{SPP} \sim \Omega_{FP} \sqrt{(22/72)(407/22)} = 156$ meV, in excellent agreement with our measurements. Using SPPs to strongly couple to the excitons thus dramatically reduces the cavity length, thus enhancing the light-matter coupling. We note that in contrast to this scaling between cavity and plasmonic enhancements, comparable Rabi splittings are produced by J-aggregate layers on both arrays of Ag holes and inside microcavities [21], because field confinements are not similarly concentrated.

We calculate the full eigenstates of the system using finite element method simulations. These confirm the anticrossings observed, and provide the optical field profiles. In the case of strong coupling at $\phi = 90^\circ$, the time-averaged near field shows the strongest intensity inside the CHPI which coats the
mediate SPP interactions. The polariton states mix excitons coupled into the layered perovskite system, where the excitons strongly couple with the SPP mode. Far-field light is directly responsible for mixing the dipole orientations, enabling the that the coupling between the excitons and their images is exciton is only seen for the in-plane dipole. It thus appears strong coupling is seen for both dipole orientations, the bare dipoles are shown in Figs. 4(d) and 4(e), respectively. While Simulated the same as films spin coated onto a flat substrate [6,12]. QWs are polarized parallel to this interface, which is indeed to the metal-dielectric interface, while excitons in CHPI are laterally confined by the grating as well as being trapped inside the surface layers where it couples to the excitons. Although CHPI is known to align planar to any local surface, it is possible that coverage may be different on vertical and horizontal surfaces of the grating. However, the strongest SPP field intensity is found at the bottom grating surface. Here the SPP E-field direction is primarily perpendicular to the metal-dielectric interface, while excitons in CHPI QWs are polarized parallel to this interface, which is indeed the same as films spin coated onto a flat substrate [6,12]. Simulated $\phi = 90^\circ$ spectra for in- and out-of-plane exciton dipoles are shown in Figs. 4(d) and 4(e), respectively. While strong coupling is seen for both dipole orientations, the bare exciton is only seen for the in-plane dipole. It thus appears that the coupling between the excitons and their images is responsible for mixing the dipole orientations, enabling the strong coupling with the SPP mode. Far-field light is directly coupled into the layered perovskite system, where the excitons mediate SPP interactions. The polariton states mix excitons within the perovskite, which are delocalized across many PbI monolayers, with SPPs which are tightly confined to the CHPI layer above the Ag grating and laterally localized in the grating slits by the coupling of standing waves. Such light-matter polaritonic quasiparticles thus combine organic, inorganic, and plasmonic components in an unusual fashion.

Strong coupling has previously been observed between inorganic or organic excitons and Au nanoslit gratings at low temperature. The coupling constants in these systems are much smaller compared to CHPI at room temperature: 55 meV for 50 nm J-aggregate films at 77 K [16], and 8 meV for 10 nm GaAs QWs at 10 K [15]. More recently room-temperature strong coupling has been seen for J-aggregate film overcoated Ag hole arrays, with splittings exceeding 600 meV at room temperature due to their large oscillator strengths [21]. One of the key challenges is to produce long-lifetime devices from both CHPI and J-aggregate nanostructures, which, although both are stable over many months at room temperature, age over longer times, which thus limits current applications. Additionally, electrical pumping, which is required in many device applications, has now been achieved in perovskite LEDs, which benefit from the layered and well-ordered sheets of CHPI interacting to form well-defined exciton polaritons. Another key difference with traditional semiconductors is that for III-V semiconductors the QWs have to be spaced at least 20 nm from the metal surface to maintain their optical quality. In contrast, our 25 nm thick CHPI film is prepared directly on the metal, and still gives strongly radiative exciton modes because the organic sandwich protects the PbI QW layers. Theoretically, Fig. 2(d) shows that excitons remain radiative via SPP coupling for film thicknesses above 10 nm. Hence the perovskite system is well suited to manipulate light-matter interactions.

In conclusion, we report evidence of image bieexcitons in perovskite-coated silver gratings with binding energies of 100 meV at room temperature. Such quasiparticles arise from the interaction between excitons and their images in the metal, and are outcoupled from the grating structure via SPP emission. These out-of-plane bieexciton states mediate coupling between in-plane QW excitons and out-of-plane SPP grating modes. This enables the observation of strong coupling at room temperature with Rabi splittings of 150 and 125 meV for the exciton and image biexciton, respectively. Such a modification of exciton behavior is of great interest for other layered van der Waals semiconductors, such as derivatives of graphene and transition metal dichalcogenides, particularly for future optoelectronic devices that demand large field enhancements by coupling to SPPs.

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