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




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Citation for the original published paper (version of record):

Rosati, R., Schmidt, R., Brem, S. et al (2021). Dark exciton anti-funneling in atomically thin semiconductors. Nature Communications, 12(1). <http://dx.doi.org/10.1038/s41467-021-27425-y>

N.B. When citing this work, cite the original published paper.

Dark exciton anti-funneling in atomically thin semiconductors

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Transport of charge carriers is at the heart of current nanoelectronics. In conventional materials, electronic transport can be controlled by applying electric fields. Atomically thin semiconductors, however, are governed by excitons, which are neutral electron-hole pairs and as such cannot be controlled by electrical fields. Recently, strain engineering has been introduced to manipulate exciton propagation. Strain-induced energy gradients give rise to exciton funneling up to a micrometer range. Here, we combine spatiotemporal photoluminescence measurements with microscopic theory to track the way of excitons in time, space and energy. We find that excitons surprisingly move away from high-strain regions. This anti-funneling behavior can be ascribed to dark excitons which possess an opposite strain-induced energy variation compared to bright excitons. Our findings open new possibilities to control transport in exciton-dominated materials. Overall, our work represents a major advance in understanding exciton transport that is crucial for technological applications of atomically thin materials.

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The mechanical exfoliation of atomically thin nanomaterials has initiated a new research field¹. Since then, in particular transition metal dichalcogenides (TMDs) have been intensively studied^{2–4}. The family of these truly two-dimensional materials exhibits a strong Coulomb interaction giving rise to the formation of excitons, which are Coulomb-bound electron-hole pairs. Having binding energies of a few hundreds of meV, they dominate the optoelectronic response of TMDs even at room temperature³. Recent studies have shown that besides the regular bright excitons accessible in optical spectra, dark excitonic states play a significant role for understanding the optical response and non-equilibrium dynamics in TMDs^{5–8}. Here, the Coulomb-bound electrons and holes either exhibit an opposite spin or are located at different valleys (K , K' , Λ) in momentum space. The required spin-flip or momentum-transfer cannot be provided by photons, making these states dark and difficult to access in optical spectra. Very recently, momentum-dark $K\Lambda$ excitons (with the hole localized at the K and the electron at the Λ valley) could be visualized in angle-resolved photoemission spectroscopy⁹.

Being atomically thin, flexible, and exhibiting strong absorption and ultrafast non-equilibrium dynamics, TMDs have been considered as promising candidates for next-generation optoelectronic applications including light-emitting, detecting, and harvesting devices¹⁰. Optical excitation, relaxation dynamics and transport of carriers represent the crucial many-particle mechanisms governing the efficiency of such devices. The optical response^{8,11} and non-equilibrium dynamics¹² of excitons in TMDs have been well studied. Exciton transport, however, has still remained elusive. In conventional materials, external electrical fields are used to control the transport of optically excited carriers¹³. However, deeply-bound excitons as neutral particles (consisting of negatively charged electrons and positively charged holes) are only weakly affected by in-plane electrical fields, e.g., via Stark shifts and exciton dissociation. Only in the case of spatially separated interlayer excitons in van der Waals

heterostructures, an electrical control of exciton transport has been demonstrated¹⁴ via the energetic tuning allowed by the dipole moment¹⁵. Thus, strain engineering has been introduced to manipulate the propagation of excitons in atomically thin semiconductors and could be exploited to e.g., boost the energy conversion efficiency in photovoltaics devices¹⁶. TMDs are remarkably sensitive to strain resulting in a large tunability of their energy landscape. In particular, strain was shown to significantly shift exciton resonances^{17,18}. As a result, strain-induced energy gradients allow us to control and manipulate exciton propagation. Excitons have been observed to funnel up to a micrometer range towards spatial regions with the strongest strain gradient, where the energy is minimal^{16,19–24}.

Tungsten-based TMD monolayers (WS_2 , WSe_2) are indirect semiconductors, where dark excitons are the energetically lowest states^{9,25}. Although these excitons are not directly accessible in optical spectra, they have been demonstrated to crucially influence optics, dynamics, and even diffusion in TMDs^{3,4,26}. However, their role on exciton funneling has still remained in the dark. In this work, we combine spatially and temporally resolved photoluminescence (PL) measurements with a microscopic many-particle theory to investigate exciton funneling in presence of spatially inhomogeneous strain gradients in the exemplary case of a WS_2 monolayer. Surprisingly, we find an inverse exciton funneling (*anti-funneling*) toward spatial regions with low strain (Fig. 1)—opposite to the conventional funneling behavior observed so far^{20–24}. Shedding light on the many-particle processes behind the funneling process, we explain this counter-intuitive behavior. In presence of strain, lattice distortions induce valley-dependent variations of energies, i.e., bright and momentum-dark excitons shift in different directions (Fig. 1c). In particular, the energy of momentum-dark $K\Lambda$ excitons increases with strain, and thus this type of excitons funnels opposite to the strain gradient toward spatial regions with minimal strain (Fig. 1e). Our joint experiment-theory study represents a major

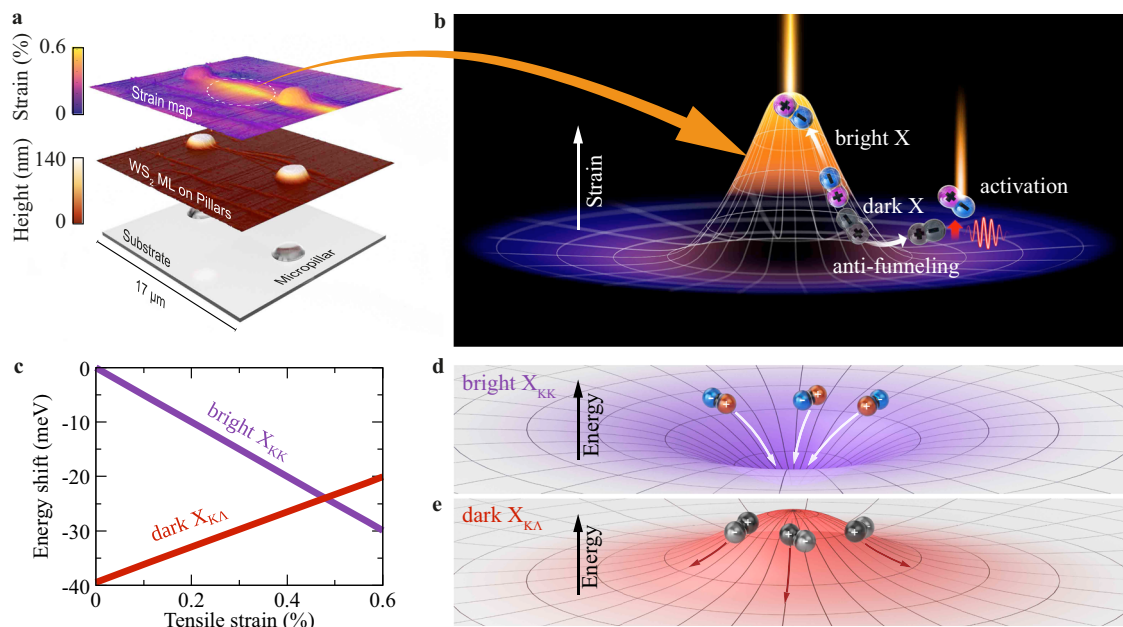


Fig. 1 Dark and bright exciton funneling in an atomically thin semiconductor. **a** Creation of an inhomogeneous strain profile in a WS_2 monolayer via transfer on an array of polymer micropillars. The lowest graph shows the patterned substrate, the middle one the sample topography with the WS_2 monolayer, and the top one the resulting strain profile in the experiment. **b** Illustration of funneling of bright excitons and anti-funneling of dark excitons in the strained material. Dark excitons can be activated by phonons and/or defects making them visible in optical spectra. **c** Strain-induced shift of dark $X_{K\Lambda}$ and bright X_{KK} exciton resonances. **d,e** Opposite shifts of $X_{K\Lambda}$ and X_{KK} give rise to reverse spatial energy gradients resulting in an unconventional anti-funneling of dark $X_{K\Lambda}$ excitons away from regions with maximum strain.

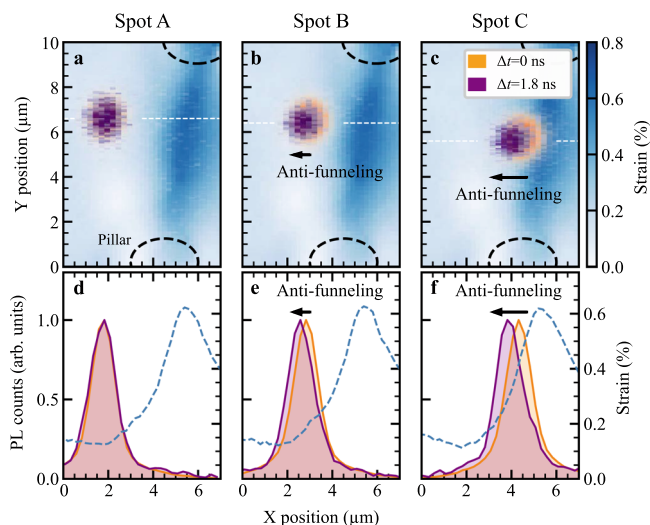


Fig. 2 Exciton anti-funneling. **a–c** Spatially resolved photoluminescence from an inhomogeneously strained WS₂ monolayer upon pulsed excitation at a delay time 0 ns (orange) and after 1.8 ns (purple). The blue colormap depicts the measured strain gradient induced by the micropillars. **d–f** Cuts along the *x* direction (horizontal white dashed line) in figures **a–c**. The dashed blue line shows the strain distribution along the *x* direction. We find that excitons do not propagate in absence of a strain gradient (spot A). Surprisingly, they funnel towards lower-strain regions (opposite to conventional funneling), when the excitation spot is subject to a considerably large strain gradient (spot B and C). The supplementary material includes PL movies further illustrating this surprising anti-funneling behavior.

advance for the microscopic understanding of exciton funneling in atomically thin semiconductors including the entire landscape of bright and dark excitons.

Results

Spatiotemporal exciton funneling. We investigate spatiotemporal exciton dynamics in a strongly spatially dependent strain profile. To create such a strain gradient, we use the concept of a nanopatterned surface, which has previously been exploited to create single-photon emitters in TMD monolayers^{27–29}. We transfer the WS₂ monolayer on an array of polymer micropillars (120 nm height and 2 μm width) on a borosilicate substrate (cf. Methods and Supplementary Information). Mechanical strain strongly modifies the electronic band structure and shifts exciton resonances (Fig. 1b). As a result, the applied strain can be mapped by measuring the energetically lowest exciton energy in spatially resolved optical absorption spectra (cf. SI for more details). Stamping the monolayer onto the micropillar array creates an inhomogeneous strain profile with the highest strain values directly between two pillars (Fig. 2a), allowing us to study exciton propagation in spatial regions of the highest strain gradients without interfering with the micropillar.

To investigate the time-resolved exciton propagation in the experiment, we spatially map the photoluminescence (PL) from the WS₂ monolayer with picosecond time resolution. We pump the monolayer with ultrashort laser pulses (≈200 fs) at 2.1 eV. An objective lens focuses the laser onto a fixed sample position. By scanning a second objective lens in transmission geometry, we image the PL distribution onto a streak camera. In that way, we achieve a temporal and a two-dimensional spatial resolution of the PL (cf. Methods for details on the experimental setup and measurement routine). From this data, we obtain ultrafast PL movies tracking exciton motion within the strain profile created

by micropillars (see the movies in the SI). Figure 2a–c shows cuts of the spatiotemporal dynamics highlighting three different excitation spots (A, B, C). Here, the orange color illustrates the exciton PL created by optical excitation (0 ns), while purple corresponds to the drifted exciton PL due to the funneling after 1.8 ns (cf. Supplementary Video S1 for the whole evolution). The blue colormap represents the strain field induced by the micropillars. For better visualization of the funneling process, we also show the PL and the strain profile along the *x* axis, cf. Fig. 2d–f. Interestingly, due to the stamping procedure a maximum strain of 0.6% is reached at the center between two micropillars, rather than directly at the position of the pillars. In these regions, the monolayer is lying flat on the substrate, which ensures a homogeneous dielectric environment. First, we optically pump at a location where the strain profile is flat (spot A). As a result, we find no observable exciton propagation (orange and purple line overlap in Fig. 2d). At locations B and C, we excite excitons in spatial regions with a significant strain gradient (dashed line in Fig. 2d–f). Here, we find a clear exciton drift, which is larger for a higher strain gradient (cf. Fig. 2e, f, and Supplementary Information). Unexpectedly, we observe exciton funneling towards low strain regions in contrast to conventional funneling^{20–24}.

Dark-exciton anti-funneling in WS₂. To be able to explain this intriguing and unexpected anti-funneling behavior, we complement our experiments with microscopic calculations of the spatiotemporal dynamics of excitons. Here, we exploit a generalized drift-diffusion equation for exciton densities $N_v(\mathbf{r}, t)$ yielding

$$\begin{aligned} \dot{N}_v(\mathbf{r}, t) = & \nabla(D(\mathbf{r})\nabla N_v(\mathbf{r}, t)) + \frac{1}{k_B T} \nabla(D(\mathbf{r})N_v(\mathbf{r}, t)\nabla E_v(\mathbf{r})) \\ & - \gamma_t N_v(\mathbf{r}, t) + \dot{N}_v^{\text{th}}(\mathbf{r}, t). \end{aligned} \quad (1)$$

Here, we explicitly take into account the entire exciton landscape, where the valley index v includes the energetically lowest 1s states of bright and momentum-dark excitons (such as KΛ, KK', ΓK). The total exciton density $N(\mathbf{r}, t) = \sum_v N_v(\mathbf{r}, t)$ is initially spatially localized with a Gaussian profile reflecting the confined optical excitation pulse in the experiment. The spatial dependence of all other quantities stems from the applied strain gradient $s(\mathbf{r})$, which in particular affects exciton energies $E_v(\mathbf{r})$. The latter are obtained by solving the Wannier equation⁷ starting from the unstrained single-particle dispersion relation⁵ and adding strain-induced shifts³⁰, cf. Methods and Supplementary Information. The spatial variation of energy (i) impacts the stationary diffusion coefficient $D(\mathbf{r})$ by altering the occupation of single exciton states and opening/closing exciton-phonon scattering channels²⁶ (see also Supplementary Information) and (ii) induces a drift force $-\nabla E_v(\mathbf{r})$ for excitons, see Methods. The first term in Eq. (1) corresponds to the standard Fick's equation for diffusion, i.e., it leads to a spatial broadening of the exciton occupation N_v due to the optically induced spatial gradient ($\nabla N_v(\mathbf{r}, t)$). This occurs already for homogeneous ($s(\mathbf{r}) = s_0$) or vanishing strain ($s = 0$). The appearing diffusion coefficient $D(\mathbf{r})$ is microscopically obtained via a full evolution of the Wigner function²⁶, cf. the Supplementary Information. In the second term in Eq. (1), the energy E_v plays the role of a scalar potential, whose spatial variation induces a drift force driving excitons toward spatial regions with a minimal E_v . This force is present only for spatially inhomogeneous strain and leads to exciton funneling (Fig. 1b). Since with larger strain E_v decreases for bright ($v = \text{KK}$) and increases for dark ($v = \text{K}\Lambda$) excitons (Fig. 1c), we expect an opposite funneling behavior (cf. Supplementary Information). As a result, momentum-dark KΛ excitons could provide an

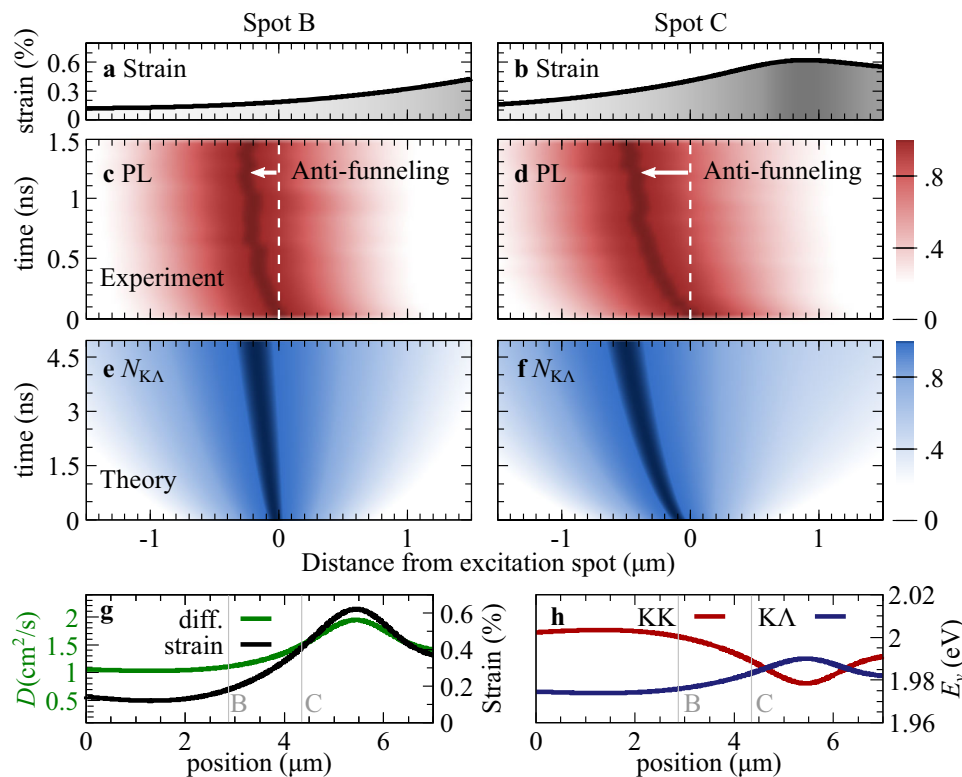


Fig. 3 Exciton anti-funneling: experiment-theory comparison. **a,b** Spatial strain profiles at the spots B and C (illustrated in Fig. 2b,c). **c,d** Experiment: Spatiotemporal evolution of the PL signal imaged with the streak camera after an optical excitation at the spots B and C. **e,f** Theory: Temporally and spatially resolved dynamics of momentum-dark $K\Lambda$ excitons obtained by solving the drift-diffusion dynamics [cf. Eq. (1)] assuming the same conditions as in the experiment. **g** Spatially dependent strain profile (black line, extracted from the experiment, cf. Fig. 2e) and theoretically predicted diffusion coefficient (green line) and **h** energy of bright KK and dark $K\Lambda$ excitons. The position of the investigated spots B and C are denoted by thin vertical lines.

explanation for the anti-funneling behavior observed in the experiment (Fig. 2).

Previous studies have shown that the interaction with high-dipole molecules³¹ or phonons⁸ can brighten dark exciton states and make them visible and even dominate the low-temperature PL of tungsten-based TMDs. Through funneling, dark $K\Lambda$ excitons can reach spatial regions, where the spectral separation of dark and bright states $E_{KK} - E_{K\Lambda}$ becomes small, significantly enhancing the phonon-driven intervalley scattering. As a direct consequence, a large amount of originally dark $K\Lambda$ excitons scatters into bright KK states and becomes visible in PL spectra (cf. Figs. 1b and SI). Additional defect-induced activation mechanisms could further boost the effect. More details on the theoretical approach (including the total decay rate γ_t and intervalley thermalization \dot{N}_v^{th} appearing in Eq. (1) as well as funneling-induced activation of dark excitons) can be found in the Supplementary Information.

To further illustrate the anti-funneling behavior, we show the spatiotemporal evolution of the experimentally measured PL signal in Fig. 3c, d. We focus on spatial cuts along the x axis (cf. the dashed white line in Fig. 2b, c) for optical excitation close to the maximum strain gradient (corresponding to spot B and C). For a better orientation with respect to the funneling direction, we also plot the spatially dependent strain profile at these spots (cf. Fig. 3a, b). Opposite to the expected behavior, we see a clear leftward propagation away from regions with maximal strain, i.e., an *anti-funneling* of excitons. The propagation is more pronounced in the spot C due to the presence of a higher strain gradient. A quantitative analysis shows that excitons propagate for a few hundreds of nm within the first nanoseconds. This is sensitive to the strain profile, i.e., at spots with a larger strain

gradient, the drift efficiency is higher (cf. the Supplementary Material). The anti-funneling length is in the same range as the conventional exciton propagation observed so far in literature^{20,21}.

The PL at room temperature is usually associated with bright excitons^{3,8}. However, these states cannot explain the observed anti-funneling behavior, since they propagate toward regions of high strain, where their energy is minimal (cf. Fig. 1c). In contrast, the energy of momentum-dark $K\Lambda$ excitons decreases with strain and is thus the driving force for the observed anti-funneling behavior. To illustrate this, we show in Fig. 3e, f the spatially and temporally resolved occupation $N_{K\Lambda}$ of momentum-dark $K\Lambda$ excitons in direct comparison to the PL measurements in Fig. 3c, d. The theoretical data is obtained by numerically evaluating the drift-diffusion Eq. (1) exactly at the conditions of the experiment, i.e., assuming the same spatially inhomogeneous strain profile shown in Fig. 2e, f. In both experiment and theory, we find a clear exciton motion to the left of the excitation spot, i.e., towards spatial regions with a smaller strain gradient (cf. the strain profile in Fig. 3a, b). This indicates that the surprising anti-funneling observed in the PL can indeed be directly ascribed to the propagation of dark excitons. This can only occur at excitation spots, where momentum-dark $K\Lambda$ excitons are energetically lower or close to the bright KK states (Fig. 3h) and thus exhibit a considerable occupation. We find a more pronounced funneling after optical excitation at location C closer to the maximum of the strain gradient, since here the spatial variation of strain is steeper, resulting in a stronger drift force $-\nabla E_{K\Lambda}(\mathbf{r})$. In addition, we observe both in experiment and theory that interestingly the funneling is initially faster and then it slows down approaching a stationary value for the propagation. This effect is well reproduced by the theoretically predicted evolution of dark-

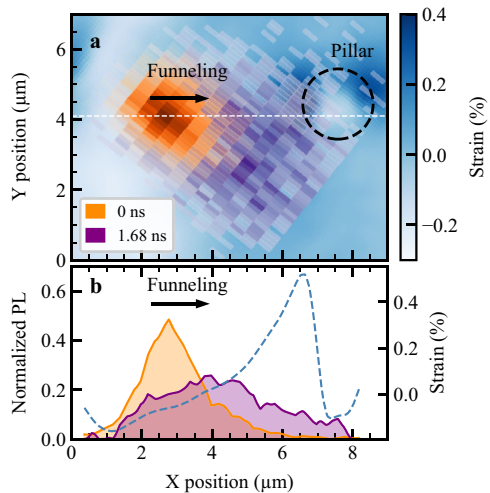


Fig. 4 Conventional exciton funneling in MoSe₂ monolayers. a

Spatiotemporal photoluminescence of MoSe₂ shown at $\Delta t = 0$ ns (orange area) and for a delay time of 1.68 ns (purple area) in presence of a strain profile (blue scale). The dashed black circle denotes the position of the micropillar. **b** Cuts along the white dashed line in **a**. The dashed blue line shows the spatial strain profile. We find a conventional funneling towards region of higher strain in MoSe₂ (see also Movie S2).

exciton occupation and can be understood as an interplay of two effects: First, the diffusion coefficient D is spatially-dependent and exhibits smaller values left from the excitation spots (cf. the green line in Fig. 3e). While funneling leftward, dark excitons move in regions with a smaller D , hence resulting in a decreasing funneling velocity. Second, dark excitons are subject to phonon-driven intervalley thermalization. A part of dark $K\Lambda$ excitons scatters into KK states, which funnel to the right and effectively slow down the overall exciton funneling.

Discussion

By combining spatiotemporal photoluminescence experiments with quantum-mechanic many-particle modeling we provide deep microscopic insights into the transport of excitons in atomically thin semiconductors. Creating a strain gradient via nanopatterned surfaces and investigating exciton propagation via spatiotemporal photoluminescence, we find an anti-funneling behavior in WS₂ monolayers. Complementing our experiments with microscopic theory, we attribute the anti-funneling to dark $K\Lambda$ excitons, which possess an opposite strain-induced energy variation compared to bright excitons and thus propagate in the opposite direction. To further prove that momentum-dark $K\Lambda$ excitons are the origin of the inverse funneling observed in our experiments, we have performed similar measurements on a molybdenum diselenide (MoSe₂) monolayer, cf. Fig. 4 and Supplementary Video S2. Contrary to the case of WS₂, we find a conventional exciton funneling, i.e., the PL signal moves from low- toward high-strain regions (cf. orange and purple area indicating initial and final distributions, respectively). This observation is in full agreement with our theoretical predictions, since in MoSe₂ dark $K\Lambda$ excitons are located above the bright states⁷. Therefore, dark states are weakly occupied and are expected to have only a minor impact on exciton funneling (cf. Supplementary Information). Remarkably, a quantitative analysis of the funneling behavior shows an efficient propagation of excitons exceeding 1 μm within 0.8 ns (cf. Supplementary Information), which is beyond the reported values of some hundreds of nanometers in previous studies performed on WS₂^{20,21}.

Our results demonstrate that strain engineering is an efficient tool to manipulate the propagation of neutral excitons. In particular, the interplay of dark and bright states offers new possibilities for tuning the optical response, dynamics, and transport, which are key processes for the realization of novel ultrathin and flexible optoelectronic devices.

Methods

Theoretical methods. The strain-dependent exciton energies entering Eq. (1) are obtained starting from the unstrained single-particle dispersion relation⁵, adding strain-induced shifts³⁰ and solving on microscopic footing the Wannier equation^{7,32}

$$\frac{\hbar^2 k^2}{2m_v} \Psi_v(\mathbf{k}) - \sum_{\mathbf{q}} W_{\mathbf{q}} \Psi_v(\mathbf{k} + \mathbf{q}) = E_v^b \Psi_v(\mathbf{k}), \quad (2)$$

where m_v is the reduced exciton mass in exciton valley v . Here, E_v^b corresponds to the exciton binding energy, which depends weakly on strain via m_v , cf. Table in Supplementary Information for the list of all appearing parameters. Furthermore, $\Psi_v(\mathbf{k})$ describes the excitonic wave function in momentum space, while $W_{\mathbf{q}}$ is the Coulomb interaction obtained via a modified form of the potential for charges in a thin film of thickness d surrounded by a dielectric environment³³. Taking into account anisotropic dielectric tensors and solving the Poisson equation with the boundary conditions described above yields $W_{\mathbf{q}} = V_{\mathbf{q}} \epsilon_{\text{scr}}(q)$ with the bare 2D-Fourier transformed Coulomb potential $V_{\mathbf{q}}$ and a non-local screening

$$\epsilon_{\text{scr}}(q) = \kappa_1 \tanh \left[\frac{1}{2} \left[\alpha_1 d q - \ln \left(\frac{\kappa_1 - \kappa_2}{\kappa_1 + \kappa_2} \right) \right] \right], \quad (3)$$

where $\kappa_i = \sqrt{\epsilon_i^{\parallel} \epsilon_i^{\perp}}$ and $\alpha_i = \sqrt{\epsilon_i^{\parallel} / \epsilon_i^{\perp}}$ account for the parallel and perpendicular component of the dielectric tensor ϵ_i of monolayer ($i = 1$)³⁴ and environment ($i = 2$)³⁵, see Table in Supplementary Information.

The exciton energy reads $E_v = E_v^0 - E_v^b$, where E_v^0 is the minimal energy separation between conduction and valence band in valley v . This quantity depends crucially on strain^{5,30}, see Table in Supplementary Information. Due to large spectral separations, we focus on the energetically lowest 1s states of the 6 most relevant exciton valleys ($KK, KK', K\Lambda, K\Lambda', \Gamma K, \Gamma K'$). These states exhibit a center-of-mass momentum \mathbf{Q} , such that the state in valley v and momentum \mathbf{Q} has an energy $E_{v,\mathbf{Q}} = E_v + \hbar^2 Q^2 / (2M_v)$ with M_v being the total exciton mass. These energies depend strongly on strain mostly via E_v , hence they are space-dependent in the case of a spatially-inhomogeneous strain profile, i.e., $E_{v,\mathbf{Q}} \equiv E_{v,\mathbf{Q}}(\mathbf{r})$. As explained in the main text, this induces (i) a spatially-dependent diffusion coefficient $D(\mathbf{r})$ that has been microscopically evaluated in Ref. 26 (cf. Supplementary Information for more details) and (ii) a drift force $-\nabla E_v$ with a generalization of the electrical mobility provided by $D/k_B T$ according to the Einstein relation^{36,37}. It is well known that trions^{38–41} can also be observed in the presence of space-dependent strain profiles²², however, we focus here on the intrinsic undoped regime. The impact of spin-dark states^{40,42} has also been neglected, since they are expected to only lead to quantitative changes due to their similar character as KK' excitons considered here.

In the presence of spatially-homogeneous strain, optically excited excitons reach an equilibrium characterized by a (local) thermalized energy distribution $N_v^0(\mathbf{r}, t)$ ⁴³ and a conventional diffusive regime⁴⁴. This occurs on a femtosecond timescale at room temperature. Since the timescale of exciton funneling is much longer, we have assumed that the stationary conditions are always fulfilled with $N_v^0(\mathbf{r}, t)$ corresponding to the Boltzmann distribution $N_v^0(\mathbf{r}, t) \propto \sum_{\mathbf{Q},v} \text{Exp}[-E_{\mathbf{Q},v}(\mathbf{r})/k_B T]$ with g_v as the valley-degeneracy factor ($g_v = 3$ for $v = K\Lambda, K\Lambda'$, otherwise $g_v = 1$). At every position, the relative amount $\bar{n}_v(\mathbf{r}) = N_v^0(\mathbf{r}, t) / \sum_v N_v^0(\mathbf{r}, t)$ of exciton occupation in valley v with respect to the total exciton population $\sum_v N_v^0(\mathbf{r}, t)$ is time-independent, while it depends on space via the excitonic energies. In particular, the ratio $\bar{n}_{K\Lambda} / \bar{n}_{KK}$ of $K\Lambda$ and KK excitons decreases for increasing strain due to the opposite strain-induced shifts of the exciton energy E_v , cf. Fig. 1c in the main text.

We describe the spatiotemporal evolution of exciton densities $N_v(\mathbf{r}, t)$ by exploiting the generalized drift-diffusion equation Eq. (1). As stated in the main manuscript, the first two terms provide diffusion and valley-intrinsic drift, respectively. The third term in Eq. (1) describes the decay of exciton occupation. Bright excitons localized in the light cone with a zero center-of-mass momentum decay via the radiative recombination rate $\gamma_{\text{rad}} = 4.5 \text{ meV}/\hbar$, which was previously calculated³⁰ and shows only small variations with strain. Once depleted via direct recombination, bright excitons are then refilled via scattering from states with a non-zero center-of-mass momentum. Since only a small amount of excitons resides in the light cone, the total excitonic density $N(\mathbf{r}, t)$ decays via an effective rate $\gamma_{\text{rad}} \bar{n}_0(\mathbf{r})$ with \bar{n}_0 being the fraction of the exciton occupation in the light cone at the position \mathbf{r} . This results in a slower decay through radiative recombination of the overall excitonic density, which takes place on a timescale of tens of nanoseconds for tungsten-based monolayers at room temperature⁴³. However, the PL signals decay typically much faster, suggesting that the leading decay mechanism in most TMDs is given by defects and impurities (except for hBN-encapsulated samples^{45–47}). Since this is an extrinsic process and varies for each sample, we assume a typical value of $\gamma_i = 1/\tau$, with $\tau = 0.35 \text{ ns}$ in accordance with recent strain experiments on WS₂¹⁸. The last term in Eq. (1) provides the intervalley

thermalization which can be written as $\dot{N}_v^{\text{th}}(\mathbf{r}, t) = \bar{\gamma}[N_v^0(\mathbf{r}, t) - N_v(\mathbf{r}, t)]$. The thermalization time $\bar{\tau} = \bar{\gamma}^{-1}$ depends on exciton valley and strain (and as such being in principle space-dependent), however, it occurs on a femtosecond timescale at room temperature^{43,44}. Compared to the much slower exciton diffusion and funneling taking place on a timescale of hundreds of ps to ns, we assume the thermalization to occur instantaneously at all spatial positions. Note that we have evaluated Eq. (1) along the x direction of strain variation exploiting the symmetry of the system.

Sample and experimental setup. To create inhomogeneous strain fields in an atomically thin semiconductor, we transfer a TMD monolayer onto a borosilicate substrate (170 μm thickness) with an array of micropillars on top^{28,29,48}. The arrays of polymer pillars (Novolak Allresist AR-N 7520.07, 10 μm distance between pillars) are fabricated using electron beam lithography. They have a height of 140 nm and circular or elliptical shapes with a diameter of $\approx 2 \mu\text{m}$. To characterize the created strain landscape, we record spectrally resolved white-light transmission images of the sample and extract the shift of the bright X_{KK} exciton for every sample position (cf. Supplementary Information). The strongest exciton shifts appear between two pillars (X -direction). The exact strain profile depends on the varying experimental conditions (angle, pressure, shear forces) during the transfer of the TMD monolayer onto the pillars and, hence, varies from sample to sample. From these exciton shifts, we can extract the strain in the monolayer using the uniaxial gauge factor for monolayer WS_2 ($-50 \text{ meV}/\%$)¹⁸.

To measure the temporally and spatially resolved exciton motion, we use a home-built optical transmission microscope. Femtosecond pulses from a tunable Er:fiber laser ($E = 2.1 \text{ eV}$, $\tau \approx 200 \text{ fs}$, $H = 0.9 \mu\text{J}/\text{cm}^2$) or from a Ti:Sapphire laser ($E = 1.71 \text{ eV}$, $\tau < 200 \text{ fs}$, $H = 3.8 \mu\text{J}/\text{cm}^2$) are focused onto the sample using an objective lens (Nikon TU Plan ELWD 50x, $NA = 0.6$) with a focus size $< 1 \mu\text{m}$. The photoluminescence is collected by a second objective lens (Nikon TU Plan 100x, $NA = 0.9$) and imaged onto the entrance slit of a streak camera (Hamamatsu C10910-01) with an attached sCMOS sensor. Residual pump light is removed using an appropriate long pass filter. The streak camera provides temporal and spatial resolution along the X dimension. To obtain a spatial resolution in the Y dimension, the collection objective lens is mounted on a piezo positioner and moved in the Y direction. This configuration allows us to spatially map the photoluminescence by scanning the objective lens in the vertical direction, projecting different sample positions onto the entrance slit of the streak camera, while exciting the monolayer at a fixed position. In that way, we obtain ultrafast videos of the exciton motion in monolayer TMDs.

Data availability

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

Code availability

The codes used to generate the data are available from the corresponding authors on reasonable request.

Received: 9 September 2021; Accepted: 16 November 2021;

Published online: 10 December 2021

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Acknowledgements

This work was supported by the Deutsche Forschungsgemeinschaft (DFG) through SFB 1083 (subproject B9) and Project No. 420760124 (BR 2888/8-1), the European Union's Horizon 2020 research and innovation program under grant agreement no. 881603 (Graphene Flagship) and the 2D Tech VINNOVA competence Center (Ref. 2019-00068).

Author contributions

R.R., S.B., R.P.C., and E.M. developed the theoretical model for exciton funneling. R.S., I.N., J.K., J.A.P., R.S., S.M.d.V., and R.B. devised and performed the spatiotemporal PL measurements. All authors contributed to the writing of the manuscript.

Funding

Open access funding provided by Chalmers University of Technology.

Competing interests

The authors declare no competing interests.

Additional information

Supplementary information The online version contains supplementary material available at <https://doi.org/10.1038/s41467-021-27425-y>.

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Peer review information *Nature Communications* thanks the anonymous reviewer(s) for their contribution to the peer review of this work.

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