Excitons, trions and Rydberg states in monolayer MoS\textsubscript{2} revealed by low-temperature photocurrent spectroscopy

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Exciton physics in two-dimensional semiconductors are typically studied by photoluminescence spectroscopy. However, this technique does not allow for direct observation of non-radiating excitonic transitions. Here, we use low-temperature photocurrent spectroscopy as an alternative technique to investigate excitonic transitions in a high-quality monolayer MoS\textsubscript{2} phototransistor. The resulting spectra presents excitonic peaks with linewidths as low as 8 meV. We identify spectral features corresponding to the ground states of neutral excitons (X\textsubscript{A}\textsuperscript{1s} and X\textsubscript{B}\textsuperscript{1s}) and charged trions (T\textsuperscript{A} and T\textsuperscript{B}) as well as up to eight additional spectral lines at energies above the X\textsubscript{B}\textsuperscript{1s} transition, which we attribute to the Rydberg series of excited states of X\textsuperscript{A} and X\textsuperscript{B}. The intensities of the spectral features can be tuned by the gate and drain-source voltages. Using an effective-mass theory for excitons in two-dimensional systems we are able to accurately fit the measured spectral lines and unambiguously associate them with their corresponding Rydberg states.
Two-dimensional transition metal dichalcogenides (2D-TMDs) are an excellent playground for studying and exploiting exciton physics. This family of materials presents unusually large exciton binding energies and lifetimes, even at room temperature and, in consequence, their optical and optoelectronic properties are largely dominated by excitonic transitions.

Until recent years, research of exciton physics in 2D-TMDs has mainly relied on photoluminescence (PL) spectroscopy measurements. This technique and its variants (time-resolved PL2, PL excitation3, etc.) allowed to obtain detailed information on the properties of light-emitting exciton transitions, including exciton binding energy4,5, lifetime6,7, spin and valley polarization8,9, etc. However, PL spectroscopy relies on spontaneous radiative decay of excitonic states. Thus, if a competing non-radiative exciton relaxation mechanism is present, it can cause a dumping, or even complete disappearance of the corresponding PL peaks, even if they are allowed by optical selection rules10. In monolayer (1L) MoS2, for example, standard photoluminescence spectroscopy reveals the two exciton states XA and XB, originating from the two spin-split band-edge optical transitions at the K point of the reciprocal lattice, but does not allow to access the higher-energy Rydberg states of these two excitons. Further, since this technique is mainly relied on photoluminescence (PL) spectroscopy16 have increasingly gained popularity for investigating excitonic states not accessible by PL.

Photocurrent spectroscopy (PCS)17,18 provides a simple, powerful, and yet largely underused complementary approach for studying excitonic transitions in 2D-TMDs. For this technique, the sample at study is exposed to monochromatic light and the light-induced change in conductivity (photoconductivity) is registered as a function of the illumination wavelength. Since the detection mechanism does not require for excitons to decay radiatively via spontaneous emission, PCS allows detecting exciton transitions regardless of the presence of dominant non-radiative relaxation mechanisms. Further, since this technique is sensitive to the electric charge, it is especially suited to study the transport properties and ionization mechanisms of photo-generated excitons, which are crucial for the development of exciton-based optoelectronics. Up to date, however, only few attempts have been made to use PCS for exciton characterization in 2D materials18,19, and measurements at cryogenic temperature and with high-quality samples are still missing in literature.

Here we investigate the excitonic properties of a monolayer MoS2 phototransistor by PCS at cryogenic temperature (T = 5 K). Our measurements allow us to fully resolve and identify the 1L-MoS2 neutral exciton states XA and XB as well as their associated charged trion states, TA and TB. Owing to the excellent quality of the fabricated device, in combination with the use of cryogenic temperature, we observe remarkably sharp exciton transitions, with bandwidths as low as 8 meV (full width at half maximum; FWHM), roughly one order of magnitude lower than in earlier PCS measurements18,19, and comparable to the bandwidths observed in low-temperature PL experiments for high-quality 1L-MoS2.20 By applying a gate voltage to increase the charge carrier density in the semiconductor channel we are able to tune the relative intensity of the exciton and trion transitions. This is observed not only for the XA and TA transitions—already reported in PL measurements—but also for the XB and TB features, not shown before. Moreover, we find that a drain-source voltage can also be used to modulate the relative intensity of exciton and trion spectral features in a similar way (shown in Supplementary Note 3). Besides the four mentioned peaks, the measured PC spectra also show eight additional features at energies above the XB transition, which we attribute to the Rydberg series of excited states of XA and XB3,15,21,22. Using a 2D effective-mass Hamiltonian with a non-hydrogenic Keldysh potential we are able to accurately reproduce the observed spectral features and unambiguously determine their origin.

Results

Device fabrication and electrical characterization. Fig. 1a schematically depicts the geometry of the studied 1L-MoS2 phototransistor. A full description of the device fabrication process and the identification of monolayer flakes is provided in the methods section and in Supplementary Notes 1 and 2. The 2D channel is fully encapsulated between top and bottom multilayer hexagonal boron nitride (h-BN) flakes in order to better preserve its intrinsic properties23. The Ti/Au electrodes follow an edge-contact geometry, as shown at the top-right inset of Fig. 1a and described in Supplementary Note 1.

We start our measurements by characterizing the electrical response of the device. Unless otherwise stated, all measurements reported below were performed in vacuum and at T = 5 K. Fig. 1b shows a transfer curve of the monolayer MoS2 phototransistor, obtained using the Si substrate as back gate. The curve shows a clear n-type behaviour, and the semiconductor channel conductivity increases as the back-gate voltage becomes larger than the threshold voltage (Vth = −9 V).

Figure 1c shows a two-terminal I–V curves of the device at different gate voltages. The curves present a back-to-back diode-like behaviour due to the presence of Schottky barriers at the contacts24,25. The different saturation currents for positive and negative voltages are caused by an asymmetry in the Schottky barrier heights. At low temperature (T = 5 K), thermionic transport can be neglected as a mechanism for conduction and thus, the source-drain current ID is mainly generated by tunnelling through the Schottky barriers.

Next, in order to characterize the 1L-MoS2 photoresponse, we expose the whole device to a homogeneous monochromatic light source. Fig. 1d shows the time-dependent source-drain current ID measured while the optical excitation is turned on and off at a frequency ν = 31.81 Hz, while applying a constant Vsd = 10 V and Vg = Vth = 2 V. The illumination energy is fixed at hν = 1.92 eV, in resonance with the XA exciton (as shown below). When the light is turned on, ID increases from its value in the dark, ID, by an amount IPD due to photoconductivity.

In 2D-TMD phototransistors, photoconductivity can emerge from two main mechanisms18,26,30, photoconductive effect, where light-induced formation of electron–hole pairs leads to an increased charge carrier density and electrical conductivity; and photovoltaic effect, where the light-induced filling or depletion of localized states causes a shift of the Fermi energy. When the characteristic relaxation times for these localized states are very long, photovoltaic effects are observed as photodoping, rather than as photoconductivity, and the Fermi energy shift remains for a long time, or even permanently, after the optical excitation is removed31.

As we further discuss below and Supplementary Note 7, we believe that in our case the observed photoconductivity is mainly dominated by photovoltaic effects26. It is worth remarking that, due to the presence of h-BN layers in contact with the 1L-MoS2 channel, the device is also affected by photodoping31, which can result in large and persistent shifts of Vth after exposure to light. We find that this effect can be prevented to a large extent by carrying out the measurements at a low illumination power density, below 2.5 mW cm−2. For such densities the light-induced shift of Vth during the spectrum acquisition process remains lower than 0.1 V. To further avoid inconsistencies in the measurements due to shifts of Vth, all the spectral data is presented as a function of Vg = Vth.
Photocurrent spectra. Figure 2 shows a typical low-temperature PC spectrum of the 1L-MoS\(_2\) phototransistor, acquired at \(T = 5\) K, \(V_{sd} = 10\) V and \(V_g - V_{th} = -20\) V (a room-temperature spectrum is shown in Supplementary Note 8 for comparison). To improve the signal-to-noise ratio, the measurement is performed while switching the illumination on and off at a fixed frequency of 31.81 Hz and the PC is registered using a lock-in amplifier. The PC spectrum is obtained by repeating this measurement while scanning the illumination wavelength in steps of 0.1 nm. A detailed description of the PCS setup can be found in Supplementary Note 4.

The resulting spectrum presents two main peaks at 1.918 eV and 2.060 eV, that we associate to the A and B excitonic ground states (X\(_A\)\(^{1s}\) and X\(_B\)\(^{1s}\)) of 1L-MoS\(_2\), with an energy splitting of 142 meV. We find that the linewidth of the observed peaks is at its lowest for gate voltages \(V_g\) well below \(V_{th}\). In these conditions the X\(_A\) peak bandwidth is found to be as low as 8 meV (FWHM), roughly one order of magnitude lower than in earlier reports, and comparable with the typical A-exciton bandwidths for low-temperature PL spectroscopy in h-BN encapsulated 1L-MoS\(_2\). This observed narrow bandwidth confirms the high quality of the 2D semiconductor channel.

It is worth remarking that, differently from optical spectroscopy, PC spectral features can only emerge if a certain optical transition results in a change in photoconductivity. Thus, charge-neutral excitons can only be observed by this technique in presence of an efficient exciton dissociation mechanism. In few-layer TMD phototransistors, exciton dissociation processes are typically largely enhanced in the vicinity of metal-semiconductor contacts, due to the large electric fields present near the interfaces. Thus, we expect that the observed PC is mainly produced in these regions.

For the spectral range between 1.85 eV and 2.15 eV the experimental PC spectrum can be very accurately fit by a multi-peak Lorentzian plus an exponential background, which accounts for the direct interband absorption edge. We find that the spectral profile is best reproduced by a quintuple Lorentzian function, with two main peaks centered at the energies of the X\(_A\)\(^{1s}\) and X\(_B\)\(^{1s}\) transitions discussed above plus three smaller peaks at 1.892 eV, 2.035 eV and 2.090 eV. We attribute the first
two of these smaller features to the A and B trion states, $T^A$ and $T^B$, expected to occur at energies 20–30 meV below $X_1$ and $X_1^*$ 35. Photogenerated trions can be either positively or negatively charged depending on the nature of the constituent charge carriers, however, in our sample the MoS$_2$ channel is strongly n-doped and, thus, we expect that the observed $T^A$ and $T^B$ features mainly account for negatively charged trions. While here we discuss our results in terms of excitons and trions, it is worth noting that recent works have proposed an alternative description for X and T transitions as originated from interactions between excitons and the Fermi sea of excess carriers (either electrons or holes) 36,37. Under this picture, the spectral features typically assigned to neutral excitons and charged trions correspond to the repulsive and attractive exciton-polaron branches respectively.

The feature at 2.090 eV is tentatively assigned to the first excited Rydberg state of the A exciton, $X_{2s}^A$, recently observed in 1L-MoS$_2$ by low-temperature micro-reflection and transmission spectroscopy measurements15,21. In fact, we also observe additional features lying at energies above 2.1 eV, indicated by arrows in Fig. 2, which we associate to the Rydberg series of excited states of $X^A$ and $X^B$, as further discussed below.

Although small, the three features assigned to $T^A$, $T^B$ and $X_{2s}^A$ are consistently reproduced in multiple spectra acquired at different bias voltages (see Supplementary Note 3). Furthermore, as we show below, these features become much more prominent when $V_g$ is increased to bring the MoS$_2$ Fermi energy above the edge of the conduction band.

**Gate modulation of photoconductivity and spectral features.** Fig. 3a shows the gate dependence of $I_{PC}$ for illumination with $h\nu = 1.92$ eV, on resonance with the $X^A$ transition. We find that the measured photocurrent becomes maximal for gate voltages close to the threshold voltage of the 1L-MoS$_2$ channel, i.e. when the Fermi energy approaches the edge of the conduction band, and largely decreases for sub-threshold voltages. This result is somewhat counterintuitive since for $V_g < V_{th}$ the electronic states at the conduction band should be completely depleted and thus, the probability of interband exciton absorption should be maximal. Indeed, for absorption spectroscopy experiments, the signal maximizes for Fermi energies below the edge of the conduction band, and decreases for larger gate voltages due to Pauli blockade18. For PC spectroscopy, however, the situation is more complex, as optical transitions will only be observed if they lead to a change in the device conductivity, either by a photoconductive effect or by a photovoltaic effect.

The observed decrease of PC for gate voltages below $V_{th}$ suggests that the main mechanism for photoreponse in our 1L-MoS$_2$ device is a photovoltaic effect similar to the one described by Furchi et al.26. For this effect, upon optical excitation, photoexcited carriers decay into localized states within the 1L-MoS$_2$ bandgap, resulting in a shift of the Fermi energy. Differently from the h-BN induced photodoping effects mentioned above31, the relaxation time for charge carriers in these impurities is very short, and thus the effect manifests as an increase of conductivity while the device is exposed to light. A characteristic signature of the photovoltaic effect is that the resulting photocurrent is proportional to the transconductance $G = dI_{PC}/dV_g$ of the semiconductor channel. As shown in Fig. 3a, the transconductance of the 1L-MoS$_2$ device, measured in the dark at $V_{sd} = 10$ V, markedly resembles the gate voltage dependence of $I_{PC}$, in consistence with the proposed mechanism for photoconductivity.

Next we address the effect of the gate voltage on the PC spectral features. For simplicity we restrict our discussion here to the five spectral lines discussed above (higher-energy spectral features not shown). Colormap of the measured photocurrent as a function of the excitation energy and the applied gate voltage. For each individual gate voltage the photocurrent data has been normalized to the value measured at $h\nu = 1.92$ eV, on resonance with the $X^A$ transition. c Individual photocurrent spectra acquired at different values of gate voltage $V_g$, with $V_g - V_{th}$ ranging from −30 to 24.2 V (grey lines). The dashed lines are fits of the measured spectra to a multi-peak Lorentzian plus an exponential background. For clarity, the spectra have been shifted vertically in steps of 0.6 nA.

Fig. 3 Gate dependence of the observed photocurrent. a Transconductance measured in dark (blue line, right axis) and gate-dependent photocurrent measured at source-drain voltage $V_{sd} = 10$ V for illumination on resonance with the $X^A$ exciton transition (orange dots, left axis). Error bars are smaller than the data points. b Colormap of the measured photocurrent as a function of the excitation energy and the applied gate voltage. For each individual gate voltage the photocurrent data has been normalized to the value measured at $h\nu = 1.92$ eV, on resonance with the $X^A$ transition. c Individual photocurrent spectra acquired at different values of gate voltage $V_g$ with $V_g - V_{th}$ ranging from −30 to 24.2 V (grey lines). The dashed lines are fits of the measured spectra to a multi-peak Lorentzian plus an exponential background. For clarity, the spectra have been shifted vertically in steps of 0.6 nA.
features will be discussed in the next section). Fig. 3b shows a color map of the photocurrent as a function of the excitation energy $h\nu$ and the gate voltage $V_g - V_{th}$. For each value of $V_g - V_{th}$ the photocurrent data has been normalized to the value measured at $h\nu = 1.92$ eV, on resonance with the $X_A$ transition. The five excitonic features described above are also apparent here, and their relative intensities are largely modulated by the gate voltage, as more clearly observed in the individual spectra shown in Fig. 3c and discussed below. A similar modulation of spectral features is also observed when tuning the drain-source voltage, as described in Supplementary Note 3.

When $V_g$ is increased, the trion transition $T_A$ becomes progressively larger, even becoming more prominent than $X_A$ for $V_g - V_{th} = 24.2$ V. A similar gate voltage modulation of the $X_A$ and $T_A$ spectral features in 1L-TMDs has been reported in literature for photoluminescence39,40, electroluminescence16, and absorption spectroscopy38,39,40,41 measurements. Typically, the $T_A$ spectral feature becomes more prominent when the Fermi energy is set above the conduction band edge, since in this situation excess electrons in the semiconductor channel can efficiently bind with photoexcited electron-hole pairs38. Similar to $T_A$, we find that the $T_B$ transition also becomes more prominent as the electron density in the conduction band is increased by the gate voltage.

As shown in earlier literature for monolayer TMDs, applying a gate voltage $V_g > V_{th}$ also results in an increased energy splitting between the $X_A$ and $T_A$ absorption peaks38,41. While this gate modulation should be visible in the PC spectra, here we only observe it very weakly, as we do not reach sufficiently large doping levels to fully resolve this effect (see Supplementary Notes 9 and 10).

Finally, as $V_g$ increases, we also observe a strengthening of the peak associated to the $X_{5s}^A$ excited state, which even becomes larger than $X_{5s}^B$ for a certain gate voltage range. As further discussed in next section and in Supplementary Note 6, we observe a similar gate modulation for excited Rydberg states of $X_A$ and $X_B$ lying at higher energies.

**Rydberg series.** We now turn to the study of the different spectral features observed for energies above $X_{5s}^A$. Fig. 4a shows the photocurrent spectrum of the 1L-MoS$_2$ device at $T = 5$ K, $V_{sd} = 10$ V and $V_g - V_{th} = 17.2$ V. In addition to the peaks associated to $X_{1s}^A$, $X_{1s}^B$, $T_A$ and $T_B$, we observe eight additional peaks, which we tentatively assign to the Rydberg series of excited states of $X_A$ and $X_B$, as labelled in the figure and enlisted in Table 1. Although some of these peaks are relatively weak compared to the $X_{1s}^A$ and $X_{1s}^B$ features, they consistently appear in spectra acquired for different gate and drain-source voltages (see Supplementary Note 6).

It is worth noting that, while $s$-type transitions are predicted to be dipole-allowed, and therefore visible in the linear photoconductivity spectra42, $p$ and $d$ excited states do not directly couple to light. Thus, in the following we restrict our discussion to $s$ states only.

Figure 4b shows the spectral position of the observed peaks and their tentatively associated quantum number $n$. In order to confirm unambiguously the spectral assignments of the peaks we fit their spectral positions using an effective-mass theory for excitons in 2D-TMDs43.

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**Table 1 Position of exciton transitions in the monolayer MoS$_2$ PC spectrum.**

| $T_A$ | $X_{1s}^A$ | $T_B$ | $X_{1s}^B$ | $X_{2s}^A$ | $X_{2s}^B$ | $X_{3s}^A$ | $X_{3s}^B$ | $X_{4s}^A$ | $X_{4s}^B$ | $X_{5s}^A$ | $X_{5s}^B$ |
|-------|-----------|-------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|
| $h\nu$ (eV) | 1.892 | 1.919 | 2.041 | 2.067 | 2.089 | 2.127 | 2.161 | 2.172 | 2.231 | 2.298 | 2.329 | 2.349 |

The energy position of the exciton transitions is extracted from the photocurrent spectrum in Fig. 4 by multi-Lorentzian fitting. For spectra acquired at different gate voltages we observe small fluctuations in the peak positions of, at most, 14 meV.
We solve numerically the effective-mass Schrödinger equation for the radially symmetric exciton states $n_s$, with energies $E_n$:

$$H\psi_n(r) = (E_n - E_g)\psi_n(r).$$

Here $E_g$ is the quasiparticle bandgap and $H$ is the Hamiltonian for the relative coordinate $r = r_e - r_h$, namely

$$H = -(\hbar^2/2\mu)\frac{\partial^2}{\partial r^2} + V(r).$$

For the A and B exciton effective masses we take $\mu_A = 0.27 m_0^{43}$ and $\mu_B = 0.28 m_0^{43}$, respectively, being $m_0$ the free electron mass. In thin semiconductor layers, the electron-hole interaction $V(r)$ cannot be simply modelled as a Coulomb potential because it is largely affected by nonlocal screening from the embedding medium. Instead, the screened electron-hole interaction is accurately described by the Keldysh potential

$$V(r) = -\left(\pi e^2/2r_0\right)[H_0(r/r_0) - Y_0(r/r_0)].$$

Here $H_0$ and $Y_0$ are zero-order Struve and Bessel functions and $\kappa$ is the dielectric constant of the embedding medium (h-BN in our case). The parameter $r_0$ is related to the screening length due to the 2D polarizability of the 1L-MoS$_2$. For the discussion below, it is worth mentioning that the Keldysh potential approaches the Coulomb potential $V(r) = -\epsilon^2/kr$ at large distance ($r \gg r_0/\kappa$) but diverges logarithmically at short distance ($r \ll r_0/\kappa$).

Thus, we are left with three fitting parameters for each excitonic Rydberg series, namely $E_n$, $\kappa$, and $r_0$. A first estimation of the quasiparticle bandgap $E_g$ can be deduced from the spectral position of the highly excited states ($n > 2$). For these states, the radius of the exciton lies in the region where $V(r) = -\epsilon^2/kr$ and the corresponding energy levels can be nicely fitted by 2D hydrogenic Rydberg series

$$E_n = E_g - \frac{\mu e^4}{(2\hbar^2\kappa^2)} \left(\frac{n}{n-1/2}\right)^2.$$

From the fitting we get $E_A^g = 2.203$ eV and $E_B^g = 2.387$ eV. We then use these values as an initial guess and perform a more accurate fit using the Keldysh potential and numerically solving the corresponding Schrödinger equation (discussed in Supplementary Note 5).

The fit reproduces with great accuracy the energies of the experimentally observed spectral features as shown in Fig. 4b, and allows us to estimate the opto-electronic parameters of 1L-MoS$_2$. The results of the fit are summarized in Table 2. For the X$^A$ Rydberg series we obtain a quasiparticle bandgap of 2.18 eV. Very recently Goryca et al. used absorption spectroscopy at high magnetic fields to estimate the quasiparticle bandgap of h-BN encapsulated MoS$_2$. There, they obtained $E_A^g = 2.16$ eV, in good agreement with the value reported here$^{43}$. For the dielectric screening parameters we get $r_0^B = 3.05$ nm and $\kappa = 4.4$, also similar to earlier results in h-BN encapsulated samples$^{43,44}$. Finally, we can estimate the binding energy of X$^A$ as $E_b^A = E_A^g - E_{1s}^A$. We get $E_b^A = 261$ meV, again in reasonable agreement with absorption and micro-reflectance spectroscopy measurements in h-BN encapsulated 1L-MoS$_2$$^{15,21}$.

For the X$^B$ Rydberg series information is scarce in literature and there are no earlier results for h-BN encapsulated samples. Hill et al. reported the observation of X$^B$ excited states in the room-temperature PL emission spectrum of 1L-MoS$_2$. There, they estimated a quasiparticle bandgap of $E_B^g = 2.47$ eV and an exciton binding energy of $\sim 440$ meV. From our measurements on h-BN encapsulated 1L-MoS$_2$ we get $E_B^g = 2.36$ eV and an exciton binding energy $E_b^B = 290$ meV, about 100 meV lower.

Finally, we use the obtained values of $E_A^g$ and $E_B^g$ to give an estimation of the MoS$_2$ valence band splitting $\Delta E_{VB}$. Assuming a conduction band splitting of $\Delta E_{CB} \approx 15$ meV$^{35,46}$ we get $\Delta E_{VB} = E_B^g - E_A^g - \Delta E_{CB} = 165$ meV, slightly above the value recently measured by ARPES for epitaxial 1L-MoS$_2$ on gold ($\Delta E_{VB} = 145 \pm 4$ meV)$^{47}$. Note that using the value for $E_b^B$ given in ref. 3 would yield an even larger $\Delta E_{VB} \approx 275$ meV.

**Discussion**

As we showed, low-temperature PCS allows us to observe very sharp excitonic spectral features, with linewidths as low as 8 meV (FWHM). While similar bandwidths for exciton features can also be achieved by PL or optical spectroscopy techniques, this typically requires the use of a microscope objective to concentrate the beam on a small area of the sample (in the order of 1 $\mu$m$^2$) to prevent peak broadening due to sample inhomogeneities. In our case however, the area of the sample that contributes to the observed spectrum is delimited by the spacing between the drain and source contacts (see Supplementary Note 4). This allows exposing the whole sample to light without losing spectral resolution, largely simplifying the experimental setup, as well as the procedure for optical alignment.

Using PCS we were able to address spectral features typically difficult to observe in PL due to their relatively low PL emission intensity. Owing to this fact, we could observe the electric field modulation of not only the X$^A$ and T$^A$ transitions, already described in literature for PL, but also of the X$^B$ and T$^B$ transitions, which to our knowledge was not reported in literature so far.

Finally, we were also able to clearly observe the excited Rydberg states of X$^A$ and X$^B$, up to $n = 5$. Similar spectral features have been also observed in previous experimental works by PL spectroscopy, PL emission and micro reflectance$^{3,15,48}$, as well as predicted in theoretical studies$^{35,42,43,49}$. However, earlier experiments only revealed excited states corresponding to the Rydberg series of either X$^A$ or X$^B$, but never both combined, making difficult to unequivocally label the observed spectral features. The PC spectra presented here, on the other hand, cannot be explained by considering only one series of Rydberg states, but require accounting for excited states of both X$^A$ and X$^B$. This strongly constrains the possible spectral assignments. The peak fittings also allow us to extract opto-electronic material

| Energy ($\text{meV}$) | Energy ($\text{eV}$) | $r_0^A$ (nm) | $r_0^B$ (nm) | $\kappa$ |
|----------------------|----------------------|---------------|---------------|---------|
| $E_A^g$              | 2.203                |                |               |         |
| $E_B^g$              | 2.387                |                |               |         |
| $E_1s^A$             | 2.16                 |                |               |         |
| $E_1s^B$             | 2.47                 |                |               |         |
| $\Delta E_{CB}$      | 15                   |                |               |         |
| $\Delta E_{VB}$      | 165                  |                |               |         |

Exponentially determined values of exciton binding energies ($E_b^A$ and $E_b^B$), quasiparticle bandgaps ($E_A^g$ and $E_B^g$) and dielectric screening parameters ($r_0^A$, $r_0^B$ and $\kappa$), for A and B excitons in monolayer MoS$_2$. For our work, typical error bars for exciton binding energies and quasiparticle bandgaps are $\pm 5$ meV. Typical error bars for $r_0^A$ and $r_0^B$ are $\pm 0.1$ nm. For $\kappa$ we get error bars of $\pm 0.1$. 

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**Table 2 Fundamental opto-electronic material parameters of monolayer MoS$_2$.**
parameters for encapsulated 1L-MoS2 that agree well with theoretical and experimental literature, further supporting the proposed peak assignments. Thanks to the simultaneous measurement of XA and XB, we could also estimate the valence potential for the study of optoelectronics and exciton physics in two-dimensional materials. In particular, the diversity of excitonic features identified here suggests that low-temperature PC spectroscopy could be favourable to other spectral techniques for observing excited excitonic states and exciton complexes with relatively small binding energies, since the larger exciton ionization rate for these states results in a strengthening of their associated spectral features. We thus expect that, in the near future, this characterization technique will progressively become more popular among the 2D optoelectronics community.

Methods

Device fabrication and contact geometry. Supplementary Figure 1 summarizes the main steps for the device fabrication. The process starts with the stacking of the heterostructures of single layer (1L) MoS2 completely encapsulated in hexagonal boron nitride (h-BN), using a dry-transfer method similar to the polypropylene carbonate (PPC) method for van der Waals heterostructures. We first exfoliate MoS2 and h-BN flakes by the standard Scotch-tape method and transfer the flakes onto SiO2 substrates. Then, we inspect the substrates through optical microscopy (Supplementary Fig. 1a), identify the 1L-MoS2 flakes by their optical contrast and confirm their thickness by micro-Raman spectroscopy as further detailed in section S2. We also use optical microscopy to identify and select two h-BN flakes with a thickness of 15–20 nm for the top layer h-BN and 25–30 nm for the bottom layer one.

Next, we start the stacking process by transferring the top h-BN layer onto the MoS2 flake (Supplementary Fig. 1b). Subsequently, we clean the substrate containing the top h-BN/MoS2 heterostructure with anisole, acetone and isopropanol (IPA) for few minutes. Both flakes are then picked up together using a PPP film, as described in ref. 5, and transferred onto the bottom layer h-BN, previously cleaned with isopropanol and IPA and annealed at 380 °C for 15 min in an Argon atmosphere. Once the entire heterostructure is assembled (Supplementary Fig. 1c) we perform a final cleaning step using anisole, acetone and IPA, followed by a second annealing in argon, with the same parameters described above. This final annealing step is crucial to remove contaminant and residual PPC, as well as eventual blisters from the heterostructure.

Once the h-BN/MoS2/h-BN heterostructure is assembled, the next step is the geometrical definition of the device by electron beam lithography (EBL) with a Raith Elphy Plus EBL system. We use a homemade PMMA (4% in chlorobenzene) as resist, spin coated at 4000 rpm for 1 minute and baked at 160 °C for 10 minutes. The use of chlorobenzene instead of commercial PMMA in anisole permits an easier and very homogeneous resist coating without the need of previous treatment, as well as a thicker coating, useful for the etching mask and the final lift-off process. After the EBL exposure (electron dose: 250 μC/cm² at 15 kV), we develop the resist with a mixture of 1 part MBK to 3 parts of isopropanol, which represents a good compromise between very good contrast and uniformity through sensitivity5,6. The resulting structure is shown in Supplementary Fig. 1d.

Next, (Supplementary Fig. 1e) we etch away the EBL-exposed areas by dry plasma etching with an ICP-RIE plasma process (100W, 1 mTorr and P=50 sccm, R=75 W, power pressure 6 mTorr and T=10 °C)5,6. The etching rate is fixed to 2 nm/s and controlled by a DC bias. As further discussed below, the sides of the etched structure have a pyramidal profile, fundamental for the consequent achievement of the edge contacts (Supplementary Fig. 1e). After this etching process, we clean the sample in acetone and IPA and carry a new annealing process, similar to the previous one, to remove residual contaminants of the PMMA resist. Further discussion on the etching process can be found in Supplementary Note 1.

After defining the stack geometry, a second EBL process (electron dose: 270 μC/cm² at 15 kV) is used to define the contact geometry (Supplementary Fig. 1f). For this step, special care was taken while designing the electrodes to avoid contact with eventual multilayer MoS2 flakes. Finally, we deposit titanium and gold (5/45 nm) by e-beam evaporation. The thickness of 15 nm allows for an easy sample alignment with micrometric resolution. In order to improve the signal-to-noise ratio of the optoelectronic measurements, the excitation signal is modulated by an optical chopper and the electrical response of the device is registered using a lock-in amplifier with the same modulation frequency. Further discussion on the spectral resolution of the measurement system can be found in Supplementary Note 4.

Data availability

The data that support the findings of this study are available from the corresponding author upon request.

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