Scalable and Transfer-Free Fabrication of MoS₂/SiO₂ Hybrid Nanophotonic Cavity Arrays with Quality Factors Exceeding 4000

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We report the fully-scalable fabrication of a large array of hybrid molybdenum disulfide (MoS₂) - silicon dioxide (SiO₂) one-dimensional, free-standing photonic-crystal cavities capable of enhancement of the MoS₂ photoluminescence at the narrow cavity resonance. We demonstrate continuous tunability of the cavity resonance wavelength across the entire emission band of MoS₂ simply by variation of the photonic crystal periodicity. Device fabrication started by substrate-scale growth of MoS₂ using chemical vapor deposition (CVD) on non-birefringent thermal oxide on a silicon wafer; it was followed by lithographic fabrication of a photonic crystal nanocavity array on the same substrate at more than 50% yield of functional devices. Our cavities exhibit three dominant modes with measured linewidths less than 0.2 nm, corresponding to quality factors exceeding 4000. All experimental findings are found to be in excellent agreement with finite difference time domain (FDTD) simulations. CVD MoS₂ provides scalable access to a direct band gap, inorganic, stable and efficient emitter material for on-chip photonics without the need for epitaxy and is at CMOS compatible processing parameters even for back-end-of-line integration; our findings suggest feasibility of cavity based line-narrowing in MoS₂-based on-chip devices as it is required for instance for frequency-multiplexed operation in on-chip optical communication and sensing.

Monolayer transition metal dichalcogenides (TMDs) have recently attracted great interest in the field of photonics because of their distinctive optical and spin properties¹-³. In contrast to bulk TMD materials, which are indirect bandgap semiconductors, monolayer TMDs are optically active due to their direct bandgap ranging between 1 and 2 eV⁴-⁵. In contrast to top-down fabrication methods like exfoliation⁶-⁷, which yields single TMD flakes of high purity, but does not permit scalability or systematic control of flake thickness and size, chemical vapor deposition (CVD) generates TMD islands and continuous films⁷-¹⁰ of uniform layer thickness and excellent optical and electrical properties. Remarkably, wafer-scale growth has been demonstrated¹¹. Even continuous spatial tuning of the optical bandgap within composite layers of different TMDs has been realized¹²,¹³.

The integration of optically active materials into photonic circuits, for example as light emitters or non-linear elements, requires efficient coupling of TMD excitons to optical fields. This can be dramatically enhanced by nanophotonic resonators. For instance, increased light extraction, signatures of Purcell enhanced emission, and even lasing were observed in systems of TMD monolayers transferred onto two-dimensional GaP-based photonic crystal cavities (PCCs)¹⁴-¹⁶ or whispering gallery microdisk resonators¹⁷. While these studies¹⁴-¹⁶ utilized exfoliation of a TMD layer and transfer onto the completed photonic crystal structure, large-scale applications will require direct growth of homogeneous TMD films onto the substrate as an essential processing step. Thermally grown SiO₂ is an excellent candidate substrate for hybrid TMD-PCC devices. It is the most established substrate for CVD growth of TMDs¹⁸ and an essential component of current state-of-the-art silicon technology. Moreover,
amorphous SiO$_2$ is non-birefringent and exhibits a large optical bandgap of ~9 eV. These properties allow SiO$_2$ to stand out over alternative materials, in particular the commonly-used indirect semiconductor GaP.

**Results**

Here we demonstrate a sequence of CVD growth of a monolayer MoS$_2$ film followed by fabrication of dozens of free-standing one-dimensional MoS$_2$-SiO$_2$ PCCs on a single substrate without any transfer steps in a fully-scalable approach. Across a broad library of different PCCs fabricated in a single step on the same substrate, we achieve wide linear tunability of the photonic mode spectrum throughout the entire emission band of MoS$_2$, simply by changing the periodicity of the photonic lattice. Both the tuning and the experimentally demonstrated quality ($Q$) factors, $Q > 4000$, are confirmed at high fidelity by finite difference time domain (FDTD) simulations.

Due to the relatively low refractive index of SiO$_2$ (~1.46), a ladder-type PCC design is highly desirable~\(^{19}\). The design used in this study is shown schematically in Fig. 1a. This geometry allows for wavelength-scale confinement of the optical modes along the beam direction ($x$) and tight confinement in the perpendicular directions due to total internal reflection. Specifically, our design consists of a 200 nm thick, free-standing, MoS$_2$-SiO$_2$ nanobeam with etched rectangular holes at a base periodicity $p$. The beam width is $w = 4.75 \ p$ and the hole sizes are $h_x = 0.5 \ p$ in $x$-direction and $h_y = 0.7 \ w$ in $y$-direction. As shown in the bottom part of Fig. 1b, the period of the first 7 holes in $\pm x$-direction, counted from the ladder center, is modulated parabolically from 0.9 $p$ to 1.0 $p$. The upper panels of Fig. 1b show the $E_y$ field component of the three dominant TE-polarized localized modes, separately calculated by FDTD simulations. As expected~\(^{20}\), the field distribution of the fundamental mode M1 shows...
Figure 2. Spatial confinement. (a) M1 mode of a cavity with period \( p = 310 \) nm. The integrated intensity in the spectral region of the cavity mode (left yellow-shaded region) is subtracted from the background (integrated counts of the right blue-shaded region). (b) The resulting intensity distribution of the mode M1 shows strong spatial confinement at the center of the PCC (the green outline represents the ladder rungs and the RIE-etched regions defining the outer border of the ladder).

Our fabrication process starts with CVD growth\(^1\) of MoS\(_2\) directly onto a 200 nm thermal oxide (SiO\(_2\)) layer of a Si substrate. After growth, we define the PCC pattern by electron beam lithography. The whole fabrication process is described in detail in the methods section. A scanning electron micrograph of a typical free-standing PCC with a period of \( p = 275 \) nm is shown in the inset in Fig. 1c. The total size of a single device, i.e. the area of the undercut region, is approximately \( 40 \mu m \times 40 \mu m \). In this work, we report on 32 devices fabricated within a rectangular area of \( 2 \) mm \( \times \) 3.5 mm. Closer packing of these relatively large individual devices would in principle allow for integration of more than 5000 devices on a typical 1 cm \( \times \) 1 cm substrate. We note that in our devices, the active MoS\(_2\) remains only within the rungs of the ladder, precisely at the positions of the antinodes of the cavity field. This unique feature ensures exclusive enhancement of the light-matter coupling. This is in strong contrast to hybrid devices fabricated by exfoliation and transfer of a 2D material on a prefabricated nanophotonic device\(^14\)–\(^16\).

Figure 1c shows the room temperature (\( T = 300 \) K) micro-photoluminescence (μPL) spectrum (see Methods section) of a typical PCC of period \( p = 275 \) nm. The broad background represents the PL emission of unstructured MoS\(_2\). In addition, the spectrum shows three sharp lines stemming from the expected localized photonic modes, M1, M2 and M3. Such cavity modes were detected from more than 50% of all studied PCCs. Lorentzian fits to these peaks (red, green and blue solid lines) reveal \( Q \)-factors of \( Q_{\text{M1}} = 1000 (\Delta \lambda = 0.62 \text{ nm}) \), \( Q_{\text{M2}} = 1700 (\Delta \lambda = 0.38 \text{ nm}) \) and \( Q_{\text{M3}} = 1570 (\Delta \lambda = 0.42 \text{ nm}) \), respectively, for this particular device. To confirm the TE-like character of these modes, we perform polarization dependent spectroscopy by placing a half-wave plate and a polarizer into the detection path of the μPL setup. Figure 1d shows the normalized and background corrected intensity of a M2 mode (symbols) as a function of the polarization angle \( \alpha \) that is fitted with a sin\(^2\) function with a period fixed to \( 180 \) degrees (green solid line). The polarization direction transverse to the beam (i.e., in \( y \)-direction) confirms the TE-like mode character. To further quantify the polarized nature of the cavity modes, we evaluate the degree of polarization, \( \text{DoP} = (I_{\text{max}} - I_{\text{min}})/(I_{\text{max}} + I_{\text{min}}) \), with \( I_{\text{min}} \) (\( I_{\text{max}} \)) being the minimum (maximum) intensity. From our data and fit, we obtain a value of \( \text{DoP} = 1.000 \pm 0.013 \) that is close to ideal.

Scanning μPL spectroscopy of a full structure is used to confirm the spatial localization and confinement of a PCC mode. For the data shown in Fig. 2a, a total area of \( 35 \mu m \times 35 \mu m \) was scanned at \( 1 \mu m \) steps. At each position, a full PL spectrum of the M1 mode was recorded. Figure 2a shows the spectrum recorded at the center of the nanocavity at position \( (x/y) = (19 \mu m/19 \mu m) \). We quantify the enhancement of the M1 emission by integration of the PL emission in the wavelength range of the cavity resonance (marked by the yellow box in Fig. 2a) and subtraction of the integrated intensity in a wavelength interval of identical width in close spectral proximity of the mode (marked by the blue box in Fig. 2a) for background correction. Performing this procedure at every point of the μPL scan, we are able to build a spatial emission enhancement map, which we normalize to its absolute maximum. Figure 2b shows this emission enhancement map. Superposition of the outline of the etched PCC structure reveals strong emission enhancement at the resonance wavelength of M1 precisely in the center of the structure. This finding directly validates the tight spatial localization of the PCC mode.

The PCC resonance wavelengths can be tuned by variation of the crystal period. Figure 3a shows the emission spectra recorded from a series of PCCs with periods tuned from \( p = 270 \) nm to 325 nm in steps of \( \Delta p = 5 \) nm. The PCC modes exhibit a continuous shift to longer wavelengths as \( p \) increases. For short periods \( p \leq 285 \) nm, modes M1, M2 and M3, marked by red, green and blue arrows, respectively, overlap with the emission band of MoS\(_2\).
Starting at $p = 290$ nm (green spectrum), an additional mode $M^*$ (black arrow) appears at shorter wavelengths, which also exhibits TE-like character in polarization-dependent measurements. As $p$ increases up to 325 nm, the entire mode spectrum of the PCC continuously shifts across the full MoS$_2$ emission band.

We analyzed the spectra obtained on each of the PCC structures in detail and extracted both the resonance wavelengths and the $Q$-factors of the PCC modes. For reference, Fig. 4a shows a typical PL spectrum of unstructured MoS$_2$. A composite image of a subset of the array of PCC cavities fabricated on a single chip is shown in Fig. 4b. Figure 4c and d aggregate the experimentally measured mode wavelengths and $Q$-factors as a function of the PCC periodicity, respectively. Experimental results are plotted as symbols with the symbol color matched to the mode assignment of Fig. 3. The error bars indicate the standard deviation of the mean value obtained by fabrication of more than one PCC at a given periodicity.

Figure 4c reveals clear linear tuning behavior of the resonance wavelengths for all observed cavity modes. Such linear tuning is precisely predicted by our FDTD simulations, which are detailed in the Methods section. The resonance wavelengths of $M1$ are calculated for SiO$_2$ thicknesses of $d = 200$ nm (the actual SiO$_2$ thickness) and $d = 300$ nm (to account for residual resist left on top of the PCC). The gray area in Fig. 4c shows the area where $M1$ is expected according to the simulations. The measured spectral shift follows faithfully these calculations.

Figure 4d shows the $Q$-factor analysis. The measured $Q$-factors are plotted as symbols using the same color-code as in Figs 3 and 4a. Most remarkably, we derive high $Q > 1000$ for almost all PCC modes resolved. However, our experimentally derived values are systematically lower than those observed by Gong and coworkers, presumably due to the thinner membranes used in our samples and increased optical absorption loss of MoS$_2$. Our measured $Q$-factors are in excellent agreement with those expected from FDTD modeling (gray shaded area).

Discussion

We designed and fabricated MoS$_2$-SiO$_2$ PCCs with $Q$-factors >4000. Simply by varying the periodicity of the photonic lattice, we are able to tune the PCC mode spectrum across the entire PL emission peak of MoS$_2$. The modes exhibit strong TE polarization and are tightly confined in the cavity region. We show that geometrical tuning of the cavity resonances follows simulations closely and that the measured $Q$ factors are in excellent agreement with our predictions for the ideal structure indicating the high quality throughout dozens of devices fabricated on the same chip – thus, highlighting the efficiency of our fabrication process. Our design can be natively transferred to other TMDs and TMD-based van der Waals heterostructures. In addition, their high $Q$-factor and strong field enhancement are ideally suited to enhance single-photon emission of quantum dot-like emission centers recently observed in the TMD WSe$_2$. The non-birefringent nature of SiO$_2$ is furthermore ideal for future applications in valleytronics. These cavity-enhanced light-matter interactions enable low power gating of the spin and valley degrees of freedom using a TMD coupled to an advanced non-polarizing cavity. Finally, the large localized electric fields in the cavity have the potential to offer unique access to non-linear optical effects such as second harmonic generation. In contrast to a recent report by Fryett and coworkers who used silicon cavities and an exfoliated TMD, the large optical bandgap of SiO$_2$ eliminates absorption losses of the visible/near-infrared photoluminescence of many TMDs.
Conclusions

We have demonstrated integration of a large array of fundamental nanophotonic devices onto an as-grown continuous film of MoS$_2$ using the most established and technologically-relevant growth substrate, SiO$_2$ on a Si wafer. Avoiding non-scalable transfer of material and demonstrating that MoS$_2$ single-layer film from a scalable technique meets homogeneity and quality requirements to realize cavity quality factors in excess of 1000, represents a substantial milestone towards technological advancements of MoS$_2$-based photonic devices. Our fully scalable approach is ideally suited for complex large-scale nanophotonic circuits built from a toolbox of functional design elements. In particular, simply by adding electrical contacts, the TMD layer can be transformed to an on-chip photodetector or a LED.

Figure 4. Tuning analysis and comparison to FDTD simulations. (a) A typical MoS$_2$ PL spectrum. (b) Optical microscope image of an array of 21 PCC devices fabricated on a continuous MoS$_2$ film. (c) Geometric tuning analysis – The PCC periods within the set of cavities of Fig. 3 are plotted as a function of the mode wavelength they support. M1, M2, M3 show excellent agreement with the linear tuning predicted by FDTD simulations for a SiO$_2$ thickness between $d = 200 \text{ nm}$ and $300 \text{ nm}$ (grey shaded area). (d) Experimentally observed Q-factor of the three dominant modes as a function of the wavelength the PCC supported. These values agree nicely with FDTD simulation results for M1 (grey shaded area).
Methods

CVD growth of MoS2. Monolayer MoS2 is grown onto the SiO2/Si substrate via CVD utilizing elemental sulfur and MoO3 powder as precursors.18 Alumina boats containing the precursors are placed at different positions in a quartz process tube. A molybdenum mesh resting on the edge of the MoO3-containing boat is used to support the substrate. The furnace with the process tube is gradually heated to 650–700°C. A continuous flow of N2 gas is utilized to transfer sulfur vapor from the boat closer to the furnace edge to the growth substrate near the furnace center. The substrate is held at the indicated temperature for ~10 min for monolayer MoS2 formation and then cooled-down gradually.

Lithographic Device Fabrication. The PCC pattern is written by electron beam lithography into 380 nm thick layer of spin-coated ZEP-520A resist. After development, this pattern is transferred into the MoS2-SiO2 layer by inductively coupled plasma reactive ion etching (ICP-RIE) using a CHF3:O2 chemistry (40sccm:1sccm). Subsequently, and without removing the remaining resist, we release the freestanding PCC nanobeam by etching the underlying Si via an isotropic 2 Torr XeF2 vapor phase process. The resist is not removed prior to optical characterization.

Optical characterization. All samples were characterized at room temperature using conventional µPL spectroscopy. The attenuated (50 µW) beam of a λ = 514 nm diode laser was focused to a ~1 µm diffraction limited spot by a 40×, NA = 0.6 microscope objective to locally excite the sample. Emission from the sample is collected by the same objective and spectrally analyzed by a 0.5 m grating monochromator equipped with a liquid N2 cooled Si CCD detector.

Finite Difference Time Domain Simulations. FDTD simulations were performed using the commercially available software package Lumerical FDTD. We used perfectly-matched-layer absorbing boundary conditions of the simulation region. The PCC was excited by electromagnetic pulses of a wide spectral range around the expected cavity resonance that are emitted by three randomly tilted dipole sources that are placed in the center of the cavity. The excitation maximum wavelength was slightly detuned from the expected resonance wavelength to avoid overlapping of excitation and emission signal. Time monitors around the cavity center record the time dependent electromagnetic field from 300 fs to 3000 fs after excitation and calculate the emission spectrum by Fourier transformation.

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Author Contributions
H.J.K. designed research. S.H. fabricated and characterized photonic crystal cavities. S.H. and H.M.M. developed fabrication process and FDTD modeling. A.E.N., D.M.-T. and S.N.A. performed CVD growth and initial material characterization. S.H., H.J.K. and L.B. wrote the manuscript. H.J.K. and L.B. supervised study.

Additional Information
Competing Interests: The authors declare that they have no competing interests.

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