

ARTICLE

https://doi.org/10.1038/s41467-019-09398-1

OPEN

Additive-free MXene inks and direct printing of micro-supercapacitors

Chuanfang (John) Zhang ^{1,2}, Lorcan McKeon^{1,3}, Matthias P. Kremer ^{1,2,4}, Sang-Hoon Park^{1,2}, Oskar Ronan^{1,2}, Andrés Seral-Ascaso^{1,2}, Sebastian Barwich^{1,3}, Cormac Ó Coileáin^{1,2}, Niall McEvoy^{1,2}, Hannah C. Nerl ^{1,3}, Babak Anasori⁵, Jonathan N. Coleman ^{1,3}, Yury Gogotsi ⁵ & Valeria Nicolosi^{1,2,4}

Direct printing of functional inks is critical for applications in diverse areas including electrochemical energy storage, smart electronics and healthcare. However, the available printable ink formulations are far from ideal. Either surfactants/additives are typically involved or the ink concentration is low, which add complexity to the manufacturing and compromises the printing resolution. Here, we demonstrate two types of two-dimensional titanium carbide $(Ti_3C_2T_x)$ MXene inks, aqueous and organic in the absence of any additive or binary-solvent systems, for extrusion printing and inkjet printing, respectively. We show examples of all-MXene-printed structures, such as micro-supercapacitors, conductive tracks and ohmic resistors on untreated plastic and paper substrates, with high printing resolution and spatial uniformity. The volumetric capacitance and energy density of the all-MXene-printed microsupercapacitors are orders of magnitude greater than existing inkjet/extrusion-printed active materials. The versatile direct-ink-printing technique highlights the promise of additive-free MXene inks for scalable fabrication of easy-to-integrate components of printable electronics.

¹ CRANN and AMBER Research Centers, Trinity College Dublin, Dublin 2, Ireland. ² School of Chemistry, Trinity College Dublin, Dublin 2, Ireland. ³ School of Physics, Trinity College Dublin, Dublin 2, Ireland. ⁴ I-FORM Advanced Manufacturing Research Centre, Trinity College Dublin, Dublin 2, Ireland. ⁵ A.J. Drexel Nanomaterials Institute and Department of Materials Science and Engineering, Drexel University, Philadelphia, PA 19104, USA. Correspondence and requests for materials should be addressed to C.Z. (email: zhangic@tcd.ie) or to Y.G. (email: gogots@drexel.edu) or to V.N. (email: nicolov@tcd.ie)

he recent boom in flexible, portable electronics and internet of things has greatly stimulated the design of advanced, miniaturized energy-storage devices¹⁻³. However, manufacturing of micro-supercapacitors (MSCs), in particular those that can achieve high energy density with a long lifetime or energy harvesting at a high rate, remains a significant challenge⁴. While elaborate patterning techniques, such as lithography, spray-masking and laser-scribing can partially resolve the problems⁵⁻⁷, the sophisticated processing and inefficient material utilization in these protocols limit the large-scale production of MSCs. In other words, incorporating nanomaterials with excellent charge-storage capability into low-cost manufacturing routes is in high demand.

Direct ink writing of functional materials offers a promising strategy for scalable production of smart electronics with a high degree of pattern and geometry flexibility⁸⁻¹². Compared with conventional manufacturing protocols, direct ink writing techniques, such as inkjet printing and extrusion printing, allow digital and additive patterning, customization, reduction in material waste, scalable and rapid production, and so on^{13,14}. An important advance for direct ink writing is the incorporation of functional inks with suitable fluidic properties, in particular surface tension and viscosity^{9,12,13}. Substantial progress has been made in the ink writing of various electronic/photonic devices^{10,12,15,16} based on graphene^{13–15,17,18}, molybdenum disulfide¹², black phosphorous¹⁶, etc^{19,20}. However, so far only limited success has been reported in achieving both fineresolution printing and high-charge-storage MSC performance⁸. In addition, in most printable inks, additives (such as surfactants or secondary solvents) are typically used to tune the concentration/rheological properties of the ink, as well as to improve the conductivity of the printed lines. The additional surfactant removal and thermal annealing steps complicate the device manufacturing process. In other words, the formulation of additive-free inks is of significance for a scalable, low-cost, yet efficient printing process.

MXenes are a family of two-dimensional (2D) carbides and nitrides of transition metals (M), where X stands for carbon or nitrogen²¹. The most extensively studied MXene, titanium carbide $(Ti_3C_2T_x)$ where T_x represents the terminated functional groups) possesses a high electronic conductivity up to ~10,000 S cm⁻¹ and a TiO₂-like surface²², resulting in ultrahigh volumetric capacitance (~1500 F cm⁻³) in $Ti_3C_2T_x$ hydrogel films and high areal capacitance (~61 mF cm⁻²) in $Ti_3C_2T_x$ MSCs²³⁻²⁵. While plenty of printed graphene MSCs have been reported with good areal capacitances, but fairly low volumetric capacitances^{15,18,26}, to date, there are just a few reports on inkjet printing of MXenes for sensors and electromagnetic interference shielding^{27,28}. The only reported MXene printing was achieved on a thermal HP printer²⁵, which limits the deposition of the MXene ink on a blank paper with a single pass, and is thus incompatible with most of micro- or nanofabrication procedures, which typically require multiple passes and/or deposits on curved surfaces. For scaling up production and industrial applications of flexible MXene-based devices, a controllable and scalable piezoelectric printing approach, which is compatible with the commercial manufacturing lines, is needed. To date, all-printed MXene MSCs with fine resolution using direct ink printing techniques have yet to be developed.

The main challenges of realizing precise MXene printing lies either in the MXene ink formulation or the solvent evaporation kinetics, or both. Typically, inkjet printing requires a high ink viscosity within a narrow range (1–20 mPa·s) and a suitable surface tension to ensure a stable jetting of singledroplets^{9,14,16,29,30}. In addition, surface tension has to be matched with the surface energy and texture of the substrate to allow good wetting¹⁶. To overcome the "coffee ring" issue^{9,14}, inks are usually mixed with surfactants or polymer stabilizers (e.g., ethyl cellulose)^{13,15,31}, and/or exchanged with low-boiling-point solvents (e.g., terpineol^{11,14,15} or ethanol¹⁴) to rapidly solidify the materials. Electronics printed in this way contain residual surfactants/polymers, which need to be removed either through high-temperature annealing^{11,15}, chemical treatment^{9,14} or intense pulsed light treatments¹³. These processes are not compatible with most substrates and are especially impractical for MXenes, which may oxidize upon heating in open air^{32,33}. In addition, MXene nanosheets typically suffer from a quick precipitation in low-boiling-point solvents, limiting the formation of concentrated MXene ink for efficient additive manufacturing.

Here, we report formulation and direct printing of additivefree, concentrated MXene inks, with high printing efficiency and spatial uniformity. Two types of $Ti_3C_2T_x$ MXene inks, aqueous and organic, in the absence of any additives, were designed for extrusion printing and inkjet printing, respectively. The all-MXene printed MSCs have exhibited excellent areal capacitance and volumetric capacitance. In addition, the protocols of MXene ink formulations as well as printing are general, i.e., ohmic resistors can be inkjet-printed, suggesting the great potential of this printing platform for scalable manufacturing of nextgeneration electronics and devices.

Results

Solvent selection criteria. To minimize the amount of defects on the MXene nanosheets, a less aggressive etching method, socalled minimally intensive layer delamination (MILD), is employed in this research (Supplementary Fig. 1)³⁴. The asobtained multi-layered (m-) Ti₃C₂T_x "cake", which swells after multiple washes (Supplementary Fig. 2), was subjected to vigorous manual shaking in water or bath sonication in organic solvent for delamination. Unlike other 2D materials, which typically require the addition of surfactants or polymer stabilizers^{12,13}, the negative electrostatic charge on the hydrophilic $Ti_3C_2T_x$ nanosheets leads to stable aqueous inks containing clean and predominantly single-layered flakes (Supplementary Figs. 3, 4). On the other hand, by selecting suitable organic solvents with a high polarity and a high dispersion interaction strength (dispersion Hansen solubility parameter)^{3,35}, stable, concentrated MXene organic inks can be formed. It is worth mentioning that solvents, such as methanol, with a low boiling point typically possess a low polarity index, and so poorly disperse MXene nanosheets and thus limit the dispersion stability. While a binarysolvent system with a low boiling point and low-medium polarity index would allow faster evaporation, the MXene ink concentration and, as a result, the printing resolution and efficiency, could be compromised compared with pure organic solvents with a high boiling point/polarity index. These two types of viscous inks are used for direct ink writing, namely, organic inks for inkjet printing and aqueous inks for extrusion printing, such as MSCs (Fig. 1).

Formulation of inkjet-printable MXene organic inks. We start by describing the formulation of inkjet-printable MXene organic inks. Four organic solvents, N-Methyl-2-pyrrolidone (NMP), dimethyl sulfoxide (DMSO), dimethylformamide (DMF) and ethanol that disperse MXene forming stable colloidal solutions³⁵ were used to delaminate the nanosheets and give MXene inks (Fig. 2a, Supplementary Figs. 5, 6), as detailed in Methods. Transmission electron microscopy (TEM) images and electron diffraction (ED) of the MXene nanosheets from the ink are shown in Fig. 2b and inset as well as Supplementary Fig. 6b–f. The MXene inks were found to be composed of monolayers to

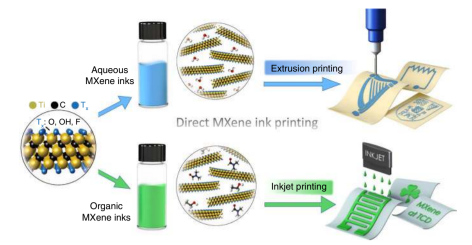


Fig. 1 Schematic illustration of direct MXene ink printing. The $Ti_3C_2T_x$ organic inks, i.e., $Ti_3C_2T_x$ -ethanol (molecules shown in the bottom panel) are used for inkjet printing of various patterns, such as MSCs, MXene letters, ohmic resistors, etc. The $Ti_3C_2T_x$ aqueous inks (with water molecules shown in the top panel) are designed for extrusion printing of MSCs and other patterns on flexible substrates. As for the MSCs, a gel electrolyte made of H_2SO_4 -PVA, was coated onto the as-printed patterns and dried naturally, forming all-MXene printed, solid-state MSCs

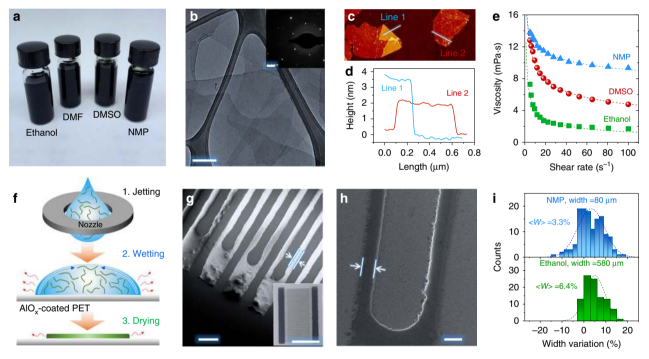


Fig. 2 Characterization of MXene organic inks. **a** Photos of various MXene organic inks. **b** TEM image of MXene nanosheets from NMP ink. Inset shows the selected area electron diffraction (SAED) pattern. Scale bar = 200 nm and = 2 1/nm in the inset. **c** AFM image and **d** the corresponding height profiles along the lines in (**c**). **e** Viscosity plotted as a function of shear rates for different MXene organic inks. The data were fitted according to the Ostwald-de Waele power law: $\eta = k\gamma^{n-1}$, where *k* and *n* are the consistency and shear-thinning index, respectively⁴⁶. **f** Scheme of fine resolution of inkjet printing of MXene organic inks. The curved green lines represent MXene nanosheets while the arrows indicate the inward (blue) and outward (red) flows of the droplet. Three critical steps, namely, stable jetting, good substrate wetting and droplet drying, control the spatial uniformity of the resultant printed patterns/lines. SEM image of the inkjet-printed MSC using, **g** the NMP ink (inset shows the whole device) and **h** the ethanol ink. Scale bar in (**g**) and (**h**) = 200 µm and = 1 cm in the inset of (**g**). The distances between the two arrows in (**g**) and (**h**) are 50 µm and 130 µm, respectively. **i** width variation of inkjet-printed MXene lines printed using NMP (top) and ethanol (bottom) inks

few-layered nanosheets. From the TEM histogram, the flakes in various organic inks possess a mean lateral dimension in the range of ~1–2.1 μm (Supplementary Fig. 7). The atomic force microscopy (AFM) analysis further confirms that the suspended nanosheets are predominantly single-layered (Fig. 2c, d), agreeing with previous reports^{24}.

By sealing all the organic inks in Ar-filled hermetic bottles and storing them in a refrigerator³², no changes in ink stability (that is, re-aggregation) have been observed over the course of 12 months except in the case of the DMSO ink (Supplementary Fig. 6g–j), which precipitated after 6 months. This necessitates future studies to reveal the possible reasons, as the colloidal stability of the ink is crucial for printing. For instance, measuring the extinction and absorption spectra of DMSO ink over time could provide the decay rate and other insights, however, this is beyond the scope of this work. Nevertheless,

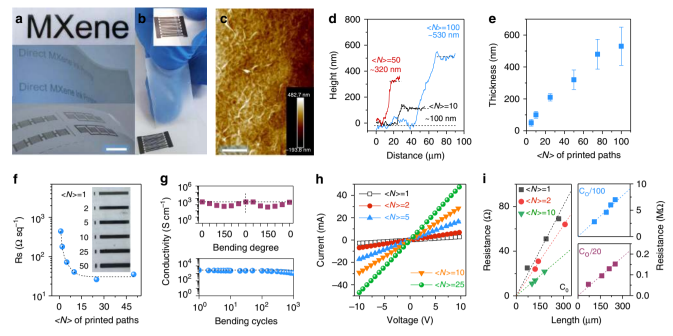


Fig. 3 Inkjet printing of MXene organic inks. **a** Optical images of inkjet-printed "MXene" word (top), "Direct MXene Ink Printing" word (middle) and MSCs (bottom) supported on AlO_x-coated PET. A total of 20 MSCs with different combinations and 80 letters were printed as a demonstration, indicating a high reproducibility of the inkjet printing. Scale bar = 1 cm. **b** Multiple peeling tests of an inkjet-printed MSC (inset) using scotch tape. No material is stuck to the tape after ten peels, indicating a strong adhesion between the printed lines and the AlO_x-coated PET substrate. **c** AFM image of the inkjet-printed line, showing a homogenous surface consisting of interconnected nanosheets. Scale bar = 10 µm. **d** The height profiles of inkjet-printed lines with different number of paths, *<N>*. **e** The line thickness plotted as a function of *<N>*. The lines in (**d**) and (**e**) were printed using NMP ink with a concentration of 12.5 mg mL⁻¹. **f** The sheet resistance, R_s , plotted as a function of ending degree (top) and number of bending cycles (bottom). One cycle is defined as bending the printed line to 150° then releasing to 0° (flat). **h** I-V curves of lines with different *<N>*. **i** Resistance of the printed lines with different *<N>* plotted as a function of length (left). An excellent linearity is observed in all the as-printed lines. The right panel shows resistance of lines with *<N>* = 1 using NMP ink, indicating a line resistivity of 620 kΩ µm⁻¹ and 26 MΩ µm⁻¹ achieved in the inks diluted by 20 times (-0.63 mg mL⁻¹, bottom right) and 100 times (-0.13 mg mL⁻¹, top right), respectively

the shelf-life of all these inks are over a timeframe that is viable for inkjet $printing^{16}$.

To reach a fine-resolution printing, MXene inks should be designed for stable jetting (that is, no secondary droplet formation after each electrical impulse)¹⁶. The inverse Ohnesorge number Z is commonly used as a figure of merit to predict if an ink will form stable drops: $Z = \sqrt{\gamma\rho D}/\eta^9$, where Z depends on surface tension (γ), density (ρ), viscosity (η) and nozzle diameter (D). The viscosity–shear rate plots indicate non-Newtonian characteristics and shear-thinning (pseudoplastic) behaviour in the organic inks (Fig. 2e and Supplementary Fig. 8)³⁶. Based on the inks' rheological properties (Supplementary Table 1), the Z~2.6 for ethanol ink is slightly higher than those of DMSO (Z~2.5) and NMP (Z~2.2) inks. The Z values of all-MXene organic inks are well within the optimal Z value range for stable jetting (1 < Z < 14)^{12,16}.

After jetting, proper substrate wetting and ink drying are crucial for uniform material deposition. Previous best practices suggested that the ink γ should be 7–10 mN m⁻¹ lower than the substrate surface energy¹⁶. While γ of ethanol (~22.1 mN m⁻¹) well matches that of common substrates like glass (~36 mN m⁻¹) and polyethylene terephthalate (PET, ~48 mN m⁻¹)¹⁶, the MXene-ethanol ink concentration is fairly low (0.7 mg mL⁻¹). DMF, NMP and DMSO on the other hand, possess high γ (~37.1, 40.8 and 43.5 mN m⁻¹, respectively), with MXene concentrations up to 12.5 mg mL⁻¹, but require additional treatment such as solvent transfer to match the substrate¹⁶. Here we choose an AlO_x-coated PET substrate (~66 mN m⁻¹)³⁷ to solve the substrate wetting issue for all organic inks, as shown in Fig. 2f.

The representative inkjet-printed lines using NMP and ethanol (Fig. 2g, h) inks showcase a high printing resolution without undesirable coffee ring effects; the nanosheets can be clearly seen on the smooth surface, forming a conductive film (Supplementary Fig. 9). By inkjet printing the NMP ink, a line (width, gap, spatial uniformity) of (~80 μ m, ~50 μ m, ~3.3%) was achieved, in contrast to (~580 μ m, ~130 μ m, 6.4%) in the ethanol-based inks (Fig. 2i and Supplementary Fig. 10). On the other hand, depositing NMP ink onto Kapton and glass substrates led to non-uniform lines (Supplementary Fig. 11), highlighting the importance of substrate selection in achieving high-resolution printing of MXene inks.

All-MXene inkjet-printed patterns. Figure 3a shows examples of inkjet-printed patterns, such as "MXene" word, printed using NMP-based MXene ink on an AlOx-coated PET substrate under ambient conditions. In particular, all-MXene MSCs can be produced in series and/or parallel (Fig. 3a). These MSCs are strongly adhered to the substrate, as confirmed by the clean scotch tape observed after peeling the MSC ten times (Fig. 3b, Supplementary Movie 1). These MSCs showcase an interconnected nanosheet network in the film, according to the atomic force microscopy (AFM, Fig. 3c). By adjusting the printing pass, both the thickness (obtained from AFM height profile, Fig. 3d) and sheet resistance $(R_{\rm s})$ can be effectively tuned. The line thickness and roughness increase linearly with the number of paths (<N>, Fig. 3e and Supplementary Fig. 12). For instance, a line thickness of $100 \pm$ 21.5 nm and 530 ± 120 nm is achieved when $\langle N \rangle = 10$ and 100, respectively. The R_s of the printed lines (2-cm-long) quickly

decreases from 445 Ω/sq ($\langle N \rangle = 1$) to 35 Ω/sq ($\langle N \rangle = 50$), as shown in Fig. 3f and the inset. In addition, the inkjet-printed lines become darker with increasing <N>, indicative of uniform printing²⁸. No oxide was observed on the as-printed lines while the film surface became rougher in the lines that were exposed to the ambient lab environment for 6 months (Supplementary Fig. 13). This is in good agreement with X-ray photoelectron spectroscopy (XPS) results (Supplementary Fig. 14a), as the deconvoluted Ti 2p core-level spectrum of the printed line produced using NMP ink $(\langle N \rangle = 2)$ is similar to that of fresh MXene³⁸, indicating that the pristine nanosheets are preserved after evaporation of the solvent. Although the C=O contribution is negligible in the deconvoluted C 1s core-level spectrum (Supplementary Fig. 14b), the N1s core-level spectrum can be deconvoluted into three peaks, which can be attributed to the trapped and adsorbed NMP molecules in the inkjet-printed MXene lines^{39–41}. This result suggests that some NMP molecules residue within the MXene nanosheets after the completion of printing.

The representative printed line ($\langle N \rangle = 5$) shows an electronic conductivity up to 2770 S cm⁻¹ initially and maintains 510 S cm⁻¹ after 6 months in ambient air, most probably due to the trapped water among the MXene layers. Moreover, printed MXene patterns are not damaged by substrate deformation, i.e., the conductivity of the as-printed lines can be fully recovered to the initial value upon releasing the bending force (Fig. 3g, top); after 1000 bending cycles, the conductivity decreases to 1093 S cm⁻¹

(Fig. 3g, bottom), probably due to the mismatch of the nanosheets (Supplementary Fig. 15).

In printed electronics, printing homogeneous lines is of great importance¹⁴. Here the reliability and reproducibility of inkjet printing technology allows us to print smooth, even and straight MXene lines on plastic substrates, and enables us to study their electrical properties. Figure 3h shows the current-voltage profiles of various lines with $\langle N \rangle$ ranging from 1 to 25, demonstrating ohmic characteristics. The resistance of these MXene lines is $\langle N \rangle$ and length dependent, showing an excellent linearity with negligible contact resistance (Fig. 3i, left). The line resistivity was determined to be 0.27, 0.21 and 0.12 $\Omega \mu m^{-1}$ for $\langle N \rangle = 1, 2$ and 10, respectively. Through diluting the initial ink concentration by a factor of 20 (~0.63 mg mL⁻¹) and 100 (~0.13 mg mL⁻¹), the line resistivity sharply increases to $620 \text{ k}\Omega \text{ }\mu\text{m}^{-1}$ and $26 \text{ }M\Omega$ μ m⁻¹, respectively at $\langle N \rangle = 1$ (Fig. 3i, right). This indicates that by adjusting ink concentration, line length and thickness, our direct MXene ink writing technique may offer a simple route to print resistors/conductive wires, with resistance values ranging from a few Ω to several M Ω , on flexible substrates, which are essential components of printed analogue circuits¹⁴.

Extrusion printing of all-MXene patterns. We further demonstrate the extrusion printing using MXene aqueous ink due to its suitable fluidic properties³⁶, including a viscous nature (Fig. 4a), a high MXene ink concentration (~36 mg mL⁻¹) and an apparent

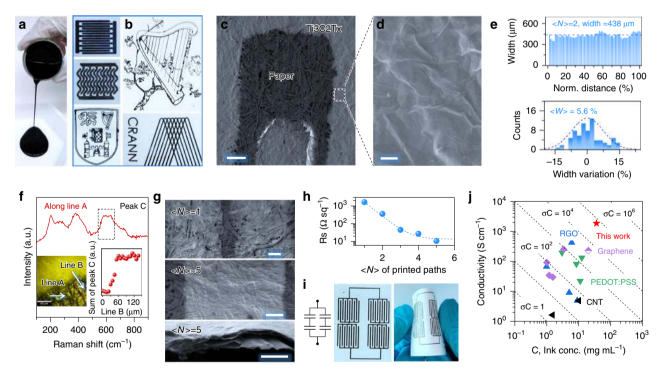


Fig. 4 Extrusion printing of MXene viscous aqueous inks. **a** Photo of MXene aqueous ink, showing its viscous nature. **b** Optical images of all-MXene printed patterns, including MSCs with various configurations, on paper. The yield is high using the extrusion printing method, producing > 70 MSCs ($\langle N \rangle = 1$) on paper based on 1 mL of the MXene ink. **c** Low-magnification SEM image of printed MXene MSC. Scale bar = 200 µm. **d** High-magnification SEM image of MXene MSC of the framed area in (**c**), showing stacked, interconnected MXene nanosheets forming a continuous film. Scale bar = 500 nm. **e** The width distribution and the resultant width spatial uniformity of the extrusion-printed MXene MSCs. **f** Raman spectrum of the extrusion-printed lines along line A, inset shows the sum intensity of peak C along line B. The MXene characteristic peaks are well retained, indicating no oxidation occurred during the extrusion printing. **g** SEM images of printed MSCs with $\langle N \rangle = 1$ (top) and $\langle N \rangle = 5$ (middle). Cross-sectional SEM image of MSC with $\langle N \rangle = 5$ is shown on the bottom, demonstrating a well-stacked, continuous nanostructure. Scale bar = 100 µm (top and middle) and = 1 µm (bottom). **h** The sheet resistance, R_s , plotted as a function of $\langle N \rangle$, showing a great flexibility (right panel). **j** Comparison of the conductivity of the as-printed lines plotted as a function of ink concentration, showing the advantage of our work in printing highly concentrated inks for highly conductive networks. The data in (**j**) come from ref. ¹³ and references within

viscosity of ~0.71 Pa·s (Supplementary Fig. 16). No sedimentation is observed over the course of 12 months in the aqueous ink (sealed in Ar-filled bottles and placed in a refrigerator). Figure 4b shows various fine-printed patterns. For instance, printing 2 paths gives MSCs with line gap ~120 μ m (Fig. 4c), width ~438 μ m and spatial uniformity within ~5.6% (Fig. 4d). The as-printed lines consist of interconnected nanosheets, forming a continuous metallic network (Fig. 4e). Well-resolved characteristic Raman peaks of MXene are detected when probing the line along different directions (Fig. 4f and insets), showing no signs of oxidation during extrusion printing.

While the thickness of the lines printed on paper cannot be precisely measured due to the rough substrate surface, we found, generally, that a higher *<N>* corresponds to a thicker continuous film (Fig. 4g and Supplementary Fig. 17), forming a wellpercolated nanosheet network that provides a much lower sheet resistance. Indeed, increasing <N> from 1 to 5 leads to an exponentially-decayed R_s from 2000 to $10 \Omega \text{ sq}^{-1}$ of the extrusion-printed lines (Fig. 4h). The metallic conductivity of the compacted printed lines eliminates the need for additional current collectors and conductive agents, while strong adhesion of stacked MXene sheets due to hydrogen bonds between the layers eliminates the need for a polymeric binder, enabling the scalable production of all-MXene-printed MSCs. For instance, highresolution, flexible MXene-based tandem MSCs can be extrusionprinted and connected in series and parallel, to meet the energy or power requirements (Fig. 4i). In addition to porous paper, the viscous MXene aqueous ink can be extrusion-printed on solid Al foil without any pre-treatment (Supplementary Fig. 18), showing a homogenous surface and a spatial uniformity to within 3.6% (Supplementary Fig. 19).

To achieve efficient printing, the ideal ink formulation should possess both, high concentration (C) and high electronic conductivity (σ). Thus, a figure of merit, FoM = σC (S cm⁻¹ · mg mL⁻¹), is commonly used to describe the electronic network properties of a printable ink¹³. A higher FoM value is preferable, as it requires less printed paths to obtain similar electrode conductivity. A combination of MXene bulk behaviour and ultrahigh nanosheet concentration results in a record-high FoM (66,996 S cm⁻¹ · mg mL⁻¹) in this work (Fig. 4j), much higher than those of other printable inks such as graphene (FoM = 6000 S cm⁻¹ · mg mL⁻¹)¹³. The ultrahigh FoM in the MXene inks enables both high-resolution printing and excellent chargestorage performance in the printed MSCs, as discussed below.

Charge-storage performance of printed MSCs. To demonstrate the possible use of the direct MXene ink printing technique for producing miniature energy-storage devices, the charge-storage performance of both inkjet- and extrusion-printed all-MXene MSCs was evaluated using a sulfuric acid (H₂SO₄)-poly(vinyl alcohol, PVA) gel electrolyte²³. The normalized cyclic voltammetry (CV) and galvanostatic charge-discharge (GCD) curves in Fig. 5a, b as well as the rate response (Supplementary Fig. 20) indicate pseudo-capacitive and high rate behaviour of a typical extrusion-printed MSC (line gap ~89 μ m, <N> = 3). By optimizing the <N> as well as the printed line gap, the electrochemical performance of extrusion-printed MSCs can be changed (Supplementary Fig. 21). In general, increasing the *<N>* results in an enhancement of areal capacitance while minimizing the line gap leads to a substantially reduced time constant, indicative of simple ion diffusion paths (Fig. 5c and Supplementary Fig. 22). For instance, the C/A improves from 3.5 to 43 mF cm^{-2} upon depositing the MXene ink from $\langle N \rangle = 1$ to 5 (Fig. 5c). Electrochemical impedance spectroscopy (EIS, Fig. 5d) indicates that pseudo-capacitive behaviour is gradually improved upon cycling,

agreeing with the CV and GCD results. It is worth noting that the voltage of all-MXene MSCs is limited to 0.5 V due to some parasitic reactions at low rates, highlighting the necessity of printing asymmetric MSCs to enlarge the voltage window.

Similarly, all inkiet-printed MSCs, with a variety of line $\langle N \rangle$ and solvents, demonstrate pseudo-capacitive responses and rate capability up to $1 V s^{-1}$ (Supplementary Figs. 23–25). In particular, the MSC printed using NMP ink exhibits the highest capacitance, reaching 1.3 and 12 mF cm⁻² when $\langle N \rangle = 2$ and 25, respectively (Fig. 5c and Supplementary Fig. 25). Importantly, our inkjet- and extrusion-printed all-MXene MSCs have outperformed most other printed MSCs^{10,15} in terms of areal capacitance (C/A) and volumetric capacitance (C/V), which are two practical metrics for evaluating the charge-storage performance (Fig. 5e)⁴². For instance, printed graphene MSCs typically display a volumetric capacitance below 100 F cm^{-3 4,5,10,15}, much lower than our inkjet-printed MXene MSC using the NMP ink $(562 \text{ F cm}^{-3}, \langle N \rangle = 25, \text{ Fig. 5f})$. The calculated energy density of the extrusion-printed MXene MSC ($\langle N \rangle = 5$) reaches as high as $0.32 \,\mu\text{W}\,\text{h}\,\text{cm}^{-2}$ at a power density of $11.4 \,\mu\text{W}\,\text{cm}^{-2}$, and $0.11 \,\mu\text{W}\,\text{h}\,\text{cm}^{-2}$ is still maintained at a high power density of $158 \,\mu\text{W}\,\text{cm}^{-2}$ (Fig. 5g). The achieved energy density is an order of magnitude higher than that of MSCs based on printed graphene⁴³, sprayed graphene/MXene⁴⁴, etc. We note that by optimization, such as minimizing the line gap, increasing the <N>, and/or introducing more pseudo-capacitive sites on the MXene nanosheets, the MSC performance could be further enhanced. There are also dozens of other than $Ti_3C_2T_x$ MXenes to choose from ref.²¹.

Both as-printed MSCs showcase excellent mechanical flexibility (Fig. 5h, i) and electrochemical cycling performance (Fig. 5j), with capacitance retention of ~97% and ~100% in the extrusion and inkjet-printed MSCs, respectively. The excellent mechanical properties of MXene films can be attributed to the high mechanical strength of $Ti_3C_2T_x$ flakes⁴⁵ and strong adhesion between the flakes. These high-performance, all-printed MXene MSCs can be arbitrarily connected in series or parallel, forming tandem devices which exhibit capacitive responses, to satisfy specific energy/power demands (Fig. 5k and Supplementary Figs. 26, 27).

Discussion

We demonstrate additive-free MXene inks and direct printing of high-performance, all-MXene micro-supercapacitors with a high resolution. The printed flexible MSCs demonstrate excellent electrochemical performance, including volumetric capacitance up to 562 F cm⁻³ and energy density as high as $0.32 \,\mu\text{W}$ h cm⁻², surpassing all other printed MSCs, to the best of our knowlege. The direct MXene ink printing technique is of fundamental importance to fields beyond energy storage and harvesting, including electronics, circuits, packaging and sensors, where cheaper and easy-to-integrate components are needed. Of equal importance is that the MXene ink formulation can be achieved by means of a scalable, facile and low-cost route. The additive- and binary-solvent-free, low-temperature printing technique suggests new possibilities for applications in smart electronics, sensors, electromagnetic shielding, antennas and other applications.

Methods

Preparation of Ti₃C₂T_x aqueous inks. Thirty-five millilitres of DI-water was added to the above mentioned m-Ti₃C₂T_x "cake", followed by vigorous shaking by hand/vortex machine for 20 min. This process delaminates the m-Ti₃C₂T_x into single- or few-layered nanosheets well dispersed in water. Then, the mixture was centrifuged at 3500 rpm for 30 min. The top 80% supernatant was collected, and further centrifuged at 5000 rpm for 1 h. After decanting the supernatant, which contains relatively small nanosheets and/or impurities, 10 mL of DI-water was

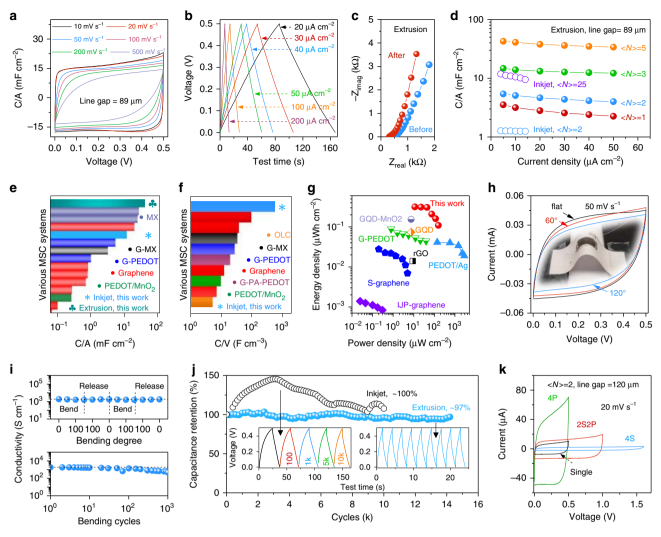


Fig. 5 Electrochemical response of inkjet- and extrusion-printed MXene MSCs. **a** Normalized cyclic voltammograms (CV) profiles and **b** galvanostatic charge-discharge (GCD) curves of a typical extrusion-printed MSC (line gap -89 μ m, <*N*> = 3). **c** Electrochemical impedance spectroscopy of extrusion-printed MSC before and after the CV tests at various scan rates. **d** Areal capacitance of inkjet- and extrusion-printed MSCs with different <*N*>. An areal capacitance of 1.3 and 12 mF cm⁻² is achieved with inkjet printing of <*N*> = 2 and 25, respectively. **e** Areal capacitance (C/A) and **f** volumetric capacitance (C/V) comparison of this work to other reported MSC systems, showing much higher C/V of our printed MXene MSCs than other reports. **g** Ragone plot comparison of this work (extrusion-printed MSC with <*N*> = 5) to other MSC systems. Detailed references and specific values in (**e**)-(**g**) can be found in the Supplementary Information (Supplementary Table 2-4). **h** CVs of extrusion-printed MSC supported on a paper substrate (inset) under different bending degrees. **i** Electronic conductivity of the extrusion-printed lines plotted as a function of bending degree (top) and bending cycles (bottom). **j** Long-term cycling of inkjet- and extrusion-printed MSCs with current densities of 14 and 200 μ A cm⁻², respectively. Insets are the typical GCD curves, showing capacitive behaviour during cycling, indicating that the excellent electrochemical performance is not due to parasitic reactions. **k** Typical CV curves of the as-printed tandem devices, such as printing four MSCs in series and in parallel, and two in series and in parallel. The as-formed tandem devices exhibit capacitive responses, showing the great flexibility of this approach to satisfy different energy/power demands

added to the sediment for redispersion by vigorous shaking, resulting in ${\rm Ti}_3{\rm C}_2{\rm T}_x$ aqueous inks.

Preparation of Ti₃C₂T_x organic inks. In this work, various organic inks were prepared using a solvent-transfer strategy. Typically, the as-prepared Ti₃C₂T_x aqueous dispersion was centrifuged at 10,000 rpm for 1 h. After decanting the supernatant, 20 mL of NMP was added to 0.1 g of Ti₃C₂T_x and the dispersion was sonicated for 30 min. A low speed (1500 rpm, 30 min) centrifugation was then employed to separate the well-dispersed flakes from the aggregated platelets. The supernatant, the sediment was re-dispersed in NMP. The ink concentration can be easily controlled by varying the volume of added NMP. DMSO, DMF and ethanolbased inks were prepared following a similar procedure.

Inkjet printing of micro-supercapacitors and resistors. The $Ti_3C_2T_x$ organic inks, without any additives, were inkjet-printed using a Dimatix DMP2800 Material printer on a variety of substrates, such as AlO_x-coated PET (NB-TP-

3GU100, Mitsubishi Paper Mills Ltd.), glass, Kapton, etc. The printer was fitted with a cartridge (DMC 11610), producing 10 pL droplets with spacing defined by rotating the print head to a pre-defined angle. The nozzle array consists of 16 identical nozzles 21 µm in diameter spaced 254 µm apart. The substrate was placed onto the vacuum plate of the printer. During printing, the substrate was heated up to 60 °C, while 70 °C was maintained at the ink ejection point in the print head. Patterns, resistors and micro-supercapacitors were printed at a droplet spacing of 25 µm on the coated PET substrate and 100 µm on the glass substrate. Micro-supercapacitor devices with a range of film thicknesses were produced by changing the print pass of the print head. Resistors were printed with different paths (ranging from 1 to 25 Ps) using the pristine Ti₃C₂T_x–NMP ink as well as NMP inks diluted by 20 and 100 times. Resistance was measured as a function of the channel length of the resistor.

Extrusion printing of micro-supercapacitors. The $Ti_3C_2T_x$ aqueous inks, without any surfactants, were extrusion-printed on paper substrates using a 3D printer (Voxel8 Inc., USA). The temperature of the substrate was set to 60 °C while the

ARTICLE

print head was held at room temperature. The ink was loaded inside the print head and squeezed through the 200 μ m-wide nozzle and deposited onto the substrate and then quickly solidified. Printing paths were designed by CAD drawings (SolidWorks 2016, Dassault Systèmes) and converted into G-code by a Voxel8's proprietary tool path generator to command the x-y-z motion of the printer head. Various patterns and micro-supercapacitors were printed with different print pass, line spacing, width, length, etc.

Materials characterization. The rheological properties of MXene aqueous inks as well as organic inks, were studied on the Anton Paar MCR 301 rheometer. Morphologies and microstructure of the as-printed lines and devices were studied by scanning electron microscopy and Raman spectroscopy. The surface chemistry of MXene was studied using XPS and the composition analysed by energy-dispersive X-ray spectroscopy (EDX). The electrical conductivity and flexibility of the as-printed lines were evaluated using a two-point probe technique. A detailed description of characterization can be found in Supplementary Methods.

Electrochemical characterization. Both inkjet- and extrusion-printed MSCs were coated with a layer of gel-like polymer electrolyte made of 3 M sulfuric acid (H₂SO₄)-Poly(vinyl alcohol, PVA) gel electrolyte, followed by natural drying. The electrochemical performance of the as-printed MXene MSCs was evaluated on a potentiostat (VMP3, BioLogic) in a voltage window of 0.5 V. CV, GCD, long-term cycling, as well as flexibility of the device were assessed. A detailed description of characterization can be found in Supplementary Methods.

Data availability

The datasets generated during and/or analysed during the current study are available from the corresponding author on reasonable request. We also provide a source data file to include all the source data except the images.

Received: 23 November 2018 Accepted: 8 March 2019 Published online: 17 April 2019

References

- Beidaghi, M. & Gogotsi, Y. Capacitive energy storage in micro-scale devices: recent advances in design and fabrication of micro-supercapacitors. *Energy Environ. Sci.* 7, 867–884 (2014).
- Tang, H. et al. In situ formed protective barrier enabled by sulfur@titanium carbide (MXene) ink for achieving high-capacity, long lifetime Li-S. *Batter. Adv. Sci.* 4, 1800502 (2018).
- Zhang, C. (John) & Nicolosi, V. Graphene and MXene-based transparent conductive electrodes and supercapacitors. *Energy Storage Mater.* 16, 102–125 (2019)..
- Pech, D. et al. Ultrahigh-power micrometre-sized supercapacitors based on onion-like carbon. Nat. Nanotechnol. 5, 651–654 (2010).
- El-Kady, M. F. & Kaner, R. B. Scalable fabrication of high-power graphene micro-supercapacitors for flexible and on-chip energy storage. *Nat. Commun.* 4, 1475 (2013).
- Zhang, C. (John) et al. Liquid exfoliation of interlayer spacing-tunable 2D vanadium oxide nanosheets: High capacity and rate handling Li-ion battery cathodes. *Nano Energy* 39, 151–161 (2017)..
- El-Kady, M. F., Strong, V., Dubin, S. & Kaner, R. B. Laser scribing of highperformance and flexible graphene-based electrochemical capacitors. *Science* 335, 1326–1330 (2012).
- Liu, Y. et al. Development of graphene oxide/polyaniline inks for high performance flexible microsupercapacitors via extrusion printing. *Adv. Funct. Mater.* 28, 1706592 (2018).
- 9. Hu, G. et al. Functional inks and printing of two-dimensional materials. Chem. Soc. Rev. 47, 3265–3300 (2018).
- Li, J. et al. Scalable fabrication and integration of graphene microsupercapacitors through full inkjet printing. ACS Nano 11, 8249–8256 (2017).
- Song, D. et al. High-resolution transfer printing of graphene lines for fully printed, flexible electronics. ACS Nano 11, 7431–7439 (2017).
- McManus, D. et al. Water-based and biocompatible 2D crystal inks for allinkjet-printed heterostructures. *Nat. Nanotechnol.* 12, 343–350 (2017).
- Secor, E. B., Ahn, B. Y., Gao, T. Z., Lewis, J. A. & Hersam, M. C. Rapid and versatile photonic annealing of graphene inks for flexible printed electronics. *Adv. Mater.* 27, 6683 (2015).
- 14. Li, J. et al. Efficient inkjet printing of graphene. Adv. Mater. 25, 3985 (2013).
- Li, L. et al. High-performance solid-state supercapacitors and microsupercapacitors derived from printable graphene inks. *Adv. Energy Mater.* 6, 1600909 (2016).

- Hu, G. et al. Black phosphorus ink formulation for inkjet printing of optoelectronics and photonics. *Nat. Commun.* 8, 278 (2017).
- Liu, Z. et al. Ultraflexible in-plane micro-supercapacitors by direct printing of solution-processable electrochemically exfoliated graphene. *Adv. Mater.* 28, 2217 (2016).
- Hyun, W. J. et al. Scalable, self-aligned printing of flexible graphene microsupercapacitors. Adv. Energy Mater. 7, 1700285 (2017).
- Zhou, K.-G., Mao, N.-N., Wang, H.-X., Peng, Y. & Zhang, H.-L. A mixedsolvent strategy for efficient exfoliation of inorganic graphene analogues. *Angew. Chem. Int. Ed.* **50**, 10839–10842 (2011).
- Kang, J. et al. Solution-based processing of optoelectronically active indium selenide. Adv. Mater. 30, 1802990 (2018).
- Anasori, B., Lukatskaya, M. R. & Gogotsi, Y. 2D metal carbides and nitrides (MXenes) for energy storage. *Nat. Rev. Mater.* 2, 16098 (2017).
- Zhang, C. J. et al. Transparent, flexible, and conductive 2D titanium carbide (MXene) films with high volumetric capacitance. *Adv. Mater.* 29, 1702678 (2017).
- 23. Lukatskaya, M. R. et al. Ultra-high-rate pseudocapacitive energy storage in two-dimensional transition metal carbides. *Nat. Energy* **2**, 17105 (2017).
- Zhang, C. J. et al. Stamping of flexible, coplanar micro-supercapacitors using MXene inks. Adv. Funct. Mater. 28, 1705506 (2018).
- 25. Quain, E. et al. Direct writing of additive-free MXene-in-water ink for electronics and energy storage. *Adv. Mater. Technol.* **4**, 1800256 (2019).
- Sun, G. et al. Layer-by-layer printing of laminated graphene-based interdigitated microelectrodes for flexible planar micro-supercapacitors. *Electrochem. Commun.* 51, 33–36 (2015).
- Zheng, J. et al. An inkjet printed Ti₃C₂-GO electrode for the electrochemical sensing of hydrogen peroxide. *J. Electrochem. Soc.* 165, B227–B231 (2018).
- Vural, M. et al. Inkjet printing of self-assembled 2D titanium carbide and protein electrodes for stimuli-responsive electromagnetic shielding. *Adv. Funct. Mater.* 28, 1801972 (2018).
- Aleeva, Y. & Pignataro, B. Recent advances in upscalable wet methods and ink formulations for printed electronics. J. Mater. Chem. C. 2, 6436–6453 (2014).
- Secor, E. B., Prabhumirashi, P. L., Puntambekar, K., Geier, M. L. & Hersam, M. C. Inkjet printing of high conductivity, flexible graphene patterns. *J. Phys. Chem. Lett.* 4, 1347–1351 (2013).
- Zhang, L. et al. Inkjet printing high-resolution, large-area graphene patterns by coffee-ring lithography. Adv. Mater. 24, 436–440 (2012).
- 32. Zhang, C. J. et al. Oxidation stability of colloidal two-dimensional titanium carbides (MXenes). *Chem. Mater.* **29**, 4848–4856 (2017).
- Lipatov, A. et al. Effect of synthesis on quality, electronic properties and environmental stability of individual monolayer Ti₃C₂ MXene flakes. *Adv. Electron. Mater.* 2, 1600255 (2016).
- 34. Alhabeb, M. et al. Guidelines for synthesis and processing of two-dimensional titanium carbide ($Ti_3C_2T_x$ MXene). *Chem. Mater.* **29**, 7633–7644 (2017).
- Maleski, K., Mochalin, V. N. & Gogotsi, Y. Dispersions of two-dimensional titanium carbide MXene in organic solvents. *Chem. Mater.* 29, 1632–1640 (2017).
- Akuzum, B. et al. Rheological characteristics of 2D titanium carbide (MXene) dispersions: a guide for processing MXenes. ACS Nano 12, 2685–2694 (2018).
- Struller, C. F., Kelly, P. J. & Copeland, N. J. Aluminum oxide barrier coatings on polymer films for food packaging applications. *Surf. Coat. Technol.* 241, 130–137 (2014).
- Halim, J. et al. X-ray photoelectron spectroscopy of select multi-layered transition metal carbides (MXenes). *Appl. Surf. Sci.* 362, 406–417 (2016).
- Charlier, J., Cousty, J., Xie, Z. X., Poulennec, C. V.-L. & Bureau, C. Adsorption of substituted pyrrolidone molecules on Au(111): an STM and XPS study. *Surf. Interface Anal.* **30**, 283–287 (2000).
- Peng, H. et al. Effect of transition metals on the structure and performance of the doped carbon catalysts derived from polyaniline and melamine for ORR application. ACS Catal. 4, 3797–3805 (2014).
- Sun, H. et al. Binder-free graphene as an advanced anode for lithium batteries. J. Mater. Chem. A 4, 6886–6895 (2016).
- Gogotsi, Y. & Simon, P. True performance metrics in electrochemical energy storage. *Science* 334, 917–918 (2011).
- Sollami Delekta, S., Smith, A. D., Li, J. & Östling, M. Inkjet printed highly transparent and flexible graphene micro-supercapacitors. *Nanoscale* 9, 6998–7005 (2017).
- 44. Li, H. et al. Flexible all-solid-state supercapacitors with high volumetric capacitances boosted by solution processable MXene and electrochemically exfoliated graphene. *Adv. Energy Mater.* **7**, 1601847 (2017).
- Lipatov, A. et al. Elastic properties of 2D Ti₃C₂T_x MXene monolayers and bilayers. *Sci. Adv.* 4, eaat0491 (2018).
- Zhang, C. et al. Highly porous carbon spheres for electrochemical capacitors and capacitive flowable suspension electrodes. *Carbon N. Y.* 77, 155–164 (2014).

Acknowledgements

We thank the SFI-funded AMBER Research Centre (SFI/12/RC/2278) and the Advanced Microscopy Laboratory for the provision of their facilities. V.N. thanks the European Research Council (StG 2DNanocaps and CoG 3D2D print) and Science Foundation Ireland (PIYRA) for funding. V.N. and M.P.K are supported by the Science Foundation Ireland (SFI) under grant number 16/RC/3872 and is co-funded under the European Regional Development Fund and by I-Form industry partners. We also acknowledge Prof. Matthias E. Möbius and Conor Patrick Cullen (Trinity College Dublin, Ireland) for the useful discussions on rheological tests and XPS. Y.G. and B.A. were supported by the U.S. Department of Energy (DOE), Office of Science, Office of Basic Energy Sciences, grant #DE-SC0018618.

Author contributions

C.F.Z., Y.G. and V.N. conceived the project. C.F.Z. synthesized materials and prepared the MXene ink, S.B. performed the rheology tests, L.M. performed the inkjet printing, M. P.K. conducted the extrusion printing, M.P.K. and B.A. created the 3D schemes and graphics, A.S., O.R. and H.C.N. performed the electron microscopy analysis, S.H.P. measured the conductivity. N.M. performed Raman analysis, C.C. conducted AFM studies, C.F.Z. performed electrochemical characterizations, C.F.Z., J.N.C., Y.G. and V.N. analysed electrochemical data. C.F.Z. wrote the manuscript with contributions from all co-authors, all authors discussed the results and commented on the manuscript.

Additional information

Supplementary Information accompanies this paper at https://doi.org/10.1038/s41467-019-09398-1.

Competing interests: The authors declare no competing interests.

Reprints and permission information is available online at http://npg.nature.com/ reprintsandpermissions/

Journal peer review information: *Nature Communications* thanks the anonymous reviewers for their contribution to the peer review of this work.

Publisher's note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/ licenses/by/4.0/.

© The Author(s) 2019