Mechanical and liquid phase exfoliation of cylindrite: a natural van der Waals superlattice with intrinsic magnetic interactions

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Abstract

We report the isolation of thin flakes of cylindrite, a naturally occurring van der Waals superlattice, by means of mechanical and liquid phase exfoliation. We find that this material is a heavily doped p-type semiconductor with a narrow gap (<0.85 eV) with intrinsic magnetic interactions that are preserved even in the exfoliated nanosheets. Due to its environmental stability and high electrical conductivity, cylindrite can be an interesting alternative to the existing 2D magnetic materials.

The study of van der Waals (vdW) heterostructures and superlattices is undoubtedly among the most active fields within the 2D materials research [1–6]. The capability of fabricating materials with tailored electrical and optical properties by assembling dissimilar 2D materials is extremely appealing [7–13]. Up to now, one of the most widespread approaches to fabricate these vdW heterostructures consists in the stacking of individual layers one-by-one using deterministic placement methods [10, 14–21]. Although these methods constitute a powerful route to fabricate stacks of virtually any combination of 2D materials, they suffer from some severe drawbacks such as the presence of interlayer contaminant adsorbates [16, 22, 23] and the difficulty of precisely controlling the orientation angle between the stacked layers [24–26]. The exfoliation of naturally occurring vdW superlattices formed by alternating layers of two different 2D materials originated by a phase segregation process during their formation has emerged as an alternative method to fabricate thin layers of vdW heterostructures avoiding the above mentioned issues. The sulfosalt mineral franckeite has been recently reported as the first example of exfoliated naturally occurring vdW superlattices [27–32]. Interestingly, the sulfosalt family has other examples of natural vdW superlattices that could be exfoliated as well.

Here we present the first study on mechanically and liquid phase exfoliated (LPE) cylindrite flakes, another mineral member of the sulfosalt family. The isolated flakes are first characterized structurally by TEM and Raman spectroscopy. We then fabricated field effect devices and photodetectors to study their electrical and optical properties. Interestingly, although cylindrite is a superlattice formed by the stacking of two large band gap semiconductors, we find that cylindrite flakes are narrow band gap semiconductors (<0.85 eV). This illustrates very well how a superlattice differs from the trivial sum of the properties of the individual constituents. Finally, we also provide a magnetic characterization of bulk and exfoliated cylinders which display magnetic correlations below 20 K. Cylindrite and other intrinsically magnetic natural heterostructures could therefore be an interesting alternative to expand the exiguous family of the 2D magnetic materials [33–35]. Cylindrite presents the additional advantage of being stable in ambient conditions and conducting and therefore of potential interest for magneto-transport [36, 37], in contrast with other insulating magnetic 2D materials.

Cylindrite belongs to the sulfosalt mineral family and it has an approximate formula Pb₃Sn₄FeSb₂S₁₄.
Within this family, it is part of a subclass of minerals called misfit compounds that are characterized by a peculiar crystal structure composed of stacks of alternating SnS₂-like octahedral (O) and PbS-like pseudo-tetragonal (T) layers [38–40]. This feature is originated by a phase segregation of the two crystalline phases during the rock formation. Interestingly, the growth conditions to reproduce the structure of cylindrite can be replicated in the laboratory to synthesize artificial crystals with cylindrite structure but with user-tailored composition [41, 42]. Figure 1 shows a simplified representation of the crystal structure of cylindrite where for the sake of simplicity substitutional Fe, Sb and Sn atoms are not displayed.

Figure 2(a) shows an optical image of cylindrite mineral displaying its characteristic appearance with clusters of small diameter (<1 mm) and long (~3–5 mm) cylinders. The individual cylinders can be easily extracted from the rock (the inset of figure 2(a) shows one of these cylinders held at the tip of lab tweezers). Since the cylinders are formed by concentric layers of material with vDW mediated interlayer interactions (see figure 2(b)), they can be easily exfoliated with adhesive tape (mechanical exfoliation) or by LPE (figure 2(c)). Figure 2(c) shows the different steps to carry out the LPE of cylindrite [32, 43–45]: (1) several cylindrite cylinders were ground carefully in an agate mortar until a black powder was obtained, (2) the powder was dispersed in iPrOH at a 1 mg·ml⁻¹ concentration, (3) this dispersion was sonicated for 1 h in an ultrasonic bath kept at 20 °C, (4) the as-prepared suspension was centrifuged (990 g, 30 min, 20 °C, Beckman Coulter Allegra X-15R, FX6100 rotor, radius 9.8 cm) in order to eliminate thicker (and thus heavier) flakes. After this process, the supernatant was collected carefully to obtain a pale orange-colored exfoliated cylindrite suspension (see figure 2(d)). From the Gel-Film, the cylindrite flakes can be transferred to an arbitrary substrate by an all dry deterministic transfer method [15]. Figure 2(d) also shows an optical image of the same flake after being transferred onto a SiO₂/Si substrate. The topography of the cylindrite flakes have been studied with atomic force microscopy (AFM) to determine their thickness (figure 2(e)).

We characterized the structure of the mechanically and LPE cylindrite flakes by transmission electron microscopy (TEM). Figure 3(a) shows a low magnification TEM image of a mechanically exfoliated cylindrite flake transferred onto a holey silicon nitride TEM grid. A striped pattern of darker and lighter areas can be seen in the low magnification image. This pattern arises from the rippled structure of the cylindrite structure that is due to a deformation of the lattice that occurs to force the commensuration of the two incommensurate lattices. HRTEM images, as shown in figure 3(b), show the characteristic stacking of the pseudo-tetragonal (T) and octahedral (O) layers of the cylindrite [14]. The fringes are approximately spaced 3.56 nm along the c direction. Figure 3(c) shows the indexed fast Fourier transform (FFT) of the image in figure 3(b) in which the reflections corresponding to the T and O layers are marked with yellow and red circles respectively. Figures 3(d) and (e) shows the corresponding low-magnification TEM and HRTEM images of a LPE cylindrite flake showing the same structural features as the mechanically exfoliated flakes. In addition, the energy-dispersive x-ray spectroscopy (EDS) shown in figure 3(f) unveils the presence of Fe in the nanosheets, expected in cylindrite also after exfoliation.

In order to characterize the electrical and optical properties of cylindrite we have fabricated simple field effect devices and photodetectors. Figure 4(a) shows the current versus voltage (IV) characteristic of a cylindrite device (see inset). The corresponding source-drain current versus gate voltage trace is displayed in the bottom inset in figure 4(a), showing a marked p-type doping without reaching the OFF state of the device (the conductance decreases monotonically upon increasing the gate voltage without vanishing).

Interestingly, cylindrite shows a sizeable photoresponse even for the illumination wavelength of 1550 nm, indicating that its bandgap is <0.85 eV. Figure 4(b) shows the photocurrent (I_{ph}) generated upon pulsed illumination with a wavelength of 1550 nm at different power densities ranging from 0.10 to 0.51 W cm⁻². The relationship between the photocurrent and the light intensity is shown in figure 4(c). It can be fitted well with a power law: I_{ph} \sim P^{\Theta}, where exponent \Theta determines the photo response to light intensity. By fitting the curve, \Theta = 0.82 is obtained, which indicates the presence of charge carrier traps in the flake. From this plot one can extract the power dependence of the responsivity R = I_{ph}/(P_{d} + A), where I_{ph} is the photocurrent, P_{d} is the power density of the illumination and A is the active area of the device. The
Figure 2. (a) Cylindrite samples: bulk mineral and an individual cylinder extracted from the rock (inset). (b) SEM image showing the base of a single cylindrite cylinder. The inset shows an amplified region around the revolution axis of the cylinder. The layered structure and the kinks source of the cylindrical shape are clearly observed. (c) Cylindrite samples. Left: bulk mineral; middle: powder material obtained after grinding several cylinders in an agate mortar; right: suspension of exfoliated material prepared by sonication of a 1 mg·ml⁻¹ powder dispersion in iPrOH. (inset in c) Optical images of macroscopic cylindrite flakes peeled off from the cylindrite cylinder with Scotch tape. (d) Optical images of a cylindrite flake on a Gel-Film stamp before its transfer (left) and after transferring it onto a SiO₂/Si substrate (right). (e) AFM topographic image of the area highlighted with a dashed square in (d). The corresponding thicknesses of different cylindrite flakes.

Figure 3. (a) Low magnification TEM image of a mechanically exfoliated cylindrite flake. (b) HRTEM image of an ultrathin mechanically exfoliated cylindrite layer. (c) FFT of b, consistent with the stacking of the pseudo-tetragonal (T: red) and octahedral (O: yellow) layers. (d) Low magnification TEM image of a LPE cylindrite flake. (e) HRTEM image with atomic resolution of a LPE cylindrite flake. (f) EDS spectrum taken on a single ultrathin cylindrite nanosheet. The arrows highlight the presence of Fe in the flake.
maximum responsivity we obtain at 1550 nm is 0.83 mA W\(^{-1}\), which is comparable with other reported narrow bandgap 2D photodetector materials such as black phosphorus [46–48].

In order to study the magnetic properties of cylindrite, we first measured the magnetism of bulky cylinders mechanically extracted from the mineral rock and we compared them to those of dispersed thin nanosheets stemming from LPE. Note that the very same cylinders isolated for the measurement in bulk are used to produce the LPE dispersion. This is important to ensure strictly the same chemical composition in both samples, as a natural mineral is the source to prepare them. The concentration of our suspension, 1 mg ml\(^{-1}\), allows a ‘macroscopic’ sample of exfoliated thin flakes of cylindrite dispersed in the selected solvent. In spite of its potential interest to explore low-dimensional magnetism in layered and nanoscale materials, LPE has been scarcely used to explore the magnetism of other bidimensional materials different form functionalized graphene. Indeed, to the best of our knowledge, only WS\(_2\) dried precipitates have been explored after exfoliation in the liquid phase [49].

The magnetism of the bulk cylindrite and LPE nanosheets has been studied using a SQUID magnetometer from Quantum Design equipped with a 5 Tesla coil. Figures 5(a) and (b) show the temperature dependence of the magnetization measured in a set of three cylindrite bulk cylinders with similar aspect ratio and packed together with their long axes aligned parallel. In a zero-field cooled (ZFC) measurement, the sample is first cooled down in the absence of a magnetic field. The magnetization is thereafter measured while ramping the temperature up under an applied magnetic field \(H = 1000\) Oe parallel (figure 5 (a)) or perpendicular (figure 5(b)) to the cylinders’ axial direction.

The magnetization monotonically increases while decreasing temperature down to \(T_k = 10\) K. Below \(T_k = 10\) K a sharp increment of \(M\) is triggered that peaks at \(T_{g,\parallel} = 5.3\) K when the field is parallel to the cylinders’ axis. A similar peak at \(T_{g,\perp} = 5.1\) K but with a less pronounced kink is observed for the perpendicular orientation of \(H\). This anomaly in the magnetization can be attributed to a slowing down of the spin dynamics in magnetically disordered systems, the hallmark of spin glass-like behavior. It has been reported for cylindrite at similar temperatures [50]. The magnetic frustration could be originated by the random distribution of Fe in the magnetically diluted cylindrite lattice. The spin glass-like interactions manifest also in the magnetization measured as a function of the magnetic field, shown in figures 5(d) and (e). A small, elongated sigmoidal-like hysteresis, typical of spin glass-like systems, opens with a coercive field \(H_c,\parallel \approx 1.58\) kOe for \(H\) parallel to the cylinders’ long axis. This value reaches up to \(H_c,\perp \approx 5.05\) kOe for the perpendicular configuration (figure 5(e)). The two orders of magnitude difference in ZFC magnetization and the different \(H_c\) between parallel and perpendicular configurations is originated by the magnetic anisotropy experienced by the Fe ions in the lattice [51].

After the bulk measurements are completed, the same three cylinders are used to obtain cylindrite nanosheets dispersed in iPrOH by LPE as detailed before. A droplet of the dispersion containing roughly hundreds of nanoflakes is mounted in a sealed plastic diamagnetic capsule and thereafter frozen for SQUID measurements. Figure 5(c) shows the resulting ZFC magnetization curve measured as a function of temperature. The background contribution to the magnetic signal of solvent and capsule is corrected by subtracting a control measurement on a capsule containing only iPrOH. Again, a small peak is observed at around \(T_k = 4.3\) K, close to bulk \(T_c\), followed by a sharper increment of \(M\) down to the lowest measured
temperatures. The small peak seems related to the spin-glass-like transition observed in bulk while the latter increment in $M$ may be associated to an increase of isolated Fe ions resulting from the LPE process behaving as a paramagnet, although an underestimated contribution to the background cannot be ruled out. In addition, the $M$ versus $H$ measurement (figure 5(f)) shows a small hysteresis with reduced $H_c \approx 76$ Oe compared to bulk signaling somehow a ‘magnetic softening’ of the exfoliated flakes. This points to a decrease of harder magnetic interactions that, in turn, could be explained by the lack of interlayer interactions in the 2D flakes, as compared to the tridimensional material. The persistence of a peak in the ZFC magnetization and the small hysteresis indicate therefore that the magnetic interactions are preserved in the exfoliated nanosheets. The reduced $H_c$ and the slightly decrease of $T_g$ could be also ascribed to the decrease in the effective size of the sample during the LPE synthesis, not only in the thickness but also in the lateral dimension. This behavior has been reported in other magnetic materials when their size is reduced to the nano/micro-scale [52–54] or their magnetic lattice is repeatedly diluted [55] and therefore the number of interacting spins is dramatically reduced.

In summary we demonstrated the isolation of thin flakes of cylindrite by mechanical and liquid-phase exfoliation of cylindrite bulk crystal and we provided a first structural, magnetic, electrical and optical characterization. In particular, we showed that cylindrite nanosheets are highly doped p-type semiconductors with a narrow bandgap. These features, in combination with its inherent environmental stability, make cylindrite an alternative material to black phosphorus. We also found that cylindrite presents magnetic interactions that are preserved even in the exfoliated nanosheets. In contrast with other reported magnetic 2D materials, cylindrite has the advantage of being stable in ambient conditions and it is electrically conducting. Moreover, we demonstrated that liquid-phase exfoliation of magnetic layered materials is a powerful tool to explore the magnetism of layered heterostructures at low-dimensionalities and at the nano-scale.

Materials and methods

Materials
The cylindrite crystal source used in this work is a natural mineral from San José mine (Oruro, Bolivia).

Scanning electron microscopy (SEM)
The SEM images are recorded by a secondary electrons detector mounted in a Carl Zeiss AURIGA Scanning Electron Microscope. The acceleration voltage is 2 kV and the working distance is 8 mm.

Transmission electron microscopy (TEM)
High resolution transition electron microscopy (HRTEM) observations were carried out in an aberration-corrected JEOL JEM-GRAND ARM300F operated at 120 kV and equipped with a cold field emission gun and a Gatan OneView camera.

Atomic force microscopy (AFM)
AFM images were carried out in a Nanotec Cervantes AFM operated in tapping mode with cantilevers with a spring constant of 40 N m$^{-1}$ and 300 kHz of resonance frequency.

Optical microscopy images
Optical microscopy images have been acquired with an AM Scope BA310 MET-T upright metallurgical
microscope equipped with an AM Scope mu1803 camera with 18 megapixels.

Transport measurements

Transport measurements have been carried out in a homebuilt probe station equipped with a source measuring unit (2450 Keithley) and a probe with a multimode optical fiber illuminator. A 1550 nm fiber coupled laser (KL5150 Thorlabs) was used to measure the photoresponse of cylindrite in the short-wavelength infrared regime.

Magnetic measurements

The magnetization in the samples is measured in a Quantum Design SQUID magnetometer mounted in a low temperature cryostat. The bulk cylinders are stacked together with kapton tape in a diamagnetic capsule.

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References

[1] Novoselov K S, Mishchenko A, Carvalho A and Castro Neto A H 2016 2D materials and van der Waals heterostructures Science 353 aac9439
[2] Liu Y, Weiss N O, Duan X, Cheng H -C, Huang Y and Duan X 2016 Van der Waals heterostructures and devices Nat. Rev. Mater. 1 16042
[3] Geim A K and Grigorieva I V 2013 Van der Waals heterostructures Nature 499 149–26
[4] Jariwala D, Marks T J and Hersam M C 2017 Mixed-dimensional van der Waals heterostructures Nat. Mater. 16 170–81
[5] Ponomarenko L A, Gorbatchev R V, Yu G L, Elias D C, Jalil R, Patel A A, Mishchenko A, Mayorov A S, Woods C R and Wallbank J R 2013 Cloning of Dirac fermions in graphene superlattices Nature 497 598–602
[6] Dean C R, Wang L, Maher P, Forsythe C, Ghahari F, Gao Y, Katoh J, Ishigami, M, Moon P and Koshino M 2013 Hofstadter’s butterfly and the fractal quantum Hall effect in moire superlattices Nature 497 598–602
[7] Yan R et al 2015 Esaki diodes in van der Waals heterojunctions with broken-gap energy band alignment Nano Lett. 15 5791–8
[8] Furchi M M, Pospischil A, Libisch F, Burgdörfer J and Mueller T 2014 Photovoltaic effect in an electrically tunable van der Waals heterojunction Nano Lett. 14 4785–91
[9] Lopez-Sanchez O, Alarcon Llado E, Koman V, Fontcuberta i Morral A, Radenovic A and Kis A 2014 Light generation and harvesting in a van der Waals heterostructure ACS Nano 8 3042–8
[10] Frisenda R, Navarro-Moratalla E, Gant P, De Lara D P, Jarillo-Herrero P, Gorbatchev R V and Castellanos-Gomez A 2018 Recent progress in the assembly of nanodevices and van der Waals heterostructures by deterministic placement of 2D materials Chem. Soc. Rev. 47 53–68
[11] Cui X et al 2015 Multi-terminal transport measurements of MoS 2 using a van der Waals heterostructure device platform Nat. Technol. 10 534–40
[12] Massignette M, Schmidt P, Via I, Schäder K G, Reserbat-Plantey A, Watanabe K, Taniguchi T, Hofstätter’s butterfly and the fractal quantum Hall effect in graphene superlattices Nature 497 598–602
[13] Withers F et al 2015 Light-emitting diodes by band-structure engineering in van der Waals heterostructures Nat. Mater. 14 301–6
[14] Frisenda R, Molina–Mendoza A J, Mueller T, Castellanos-Gomez A and van der Zant H S J 2018 Atomically thin p-n heterojunctions of bilayer graphene and hexagonal boron nitride ACS Nano 12 42–6
[15] Castellanos-Gomez A, Buscema M, Molenaar R, Singh V, Jansen L, van der Zant H S J and Steele G A 2014 Deterministic transfer of two-dimensional materials by all-dry viscoelastic stamping 2D Mater. 1 011002
[16] Uwanno T, Hattori Y, Taniguchi T, Watanabe K and Nagashio K 2015 Fully dry PMMA transfer of graphene on h-BN using a heating/cooling system 2D Mater. 6 01002
[17] Zomer P J, Dash S P, Tombrors N and Van Wees B J 2011 A transfer technique for high mobility graphene devices on commercially available hexagonal boron nitride Appl. Phys. Lett. 99 232104
[18] Zomer P J, Guimaraes M H D, Brant J C, Tombrors N and Van Wees B J 2014 Fast pick up technique for high quality heterostructures of bilayer graphene and hexagonal boron nitride Appl. Phys. Lett. 105 13101
[19] Pizzocchero F, Gammelgaard L, Jenson B S, Caridad J M, Wang L, Hone J, Beggild P and Booth T J 2016 The hot transport properties of single-layer graphene Nat. Commun. 7 11371
[20] Frisenda R, Molina-Mendoza A J, Mueller T, Castellanos-Gomez A and van der Zant H S J 2018 Atomically thin p-n heterojunctions of bilayer graphene and hexagonal boron nitride ACS Nano 12 42–6
[21] Castellanos-Gomez A, Buscema M, Molenaar R, Singh V, Jansen L, van der Zant H S J and Steele G A 2014 Deterministic transfer of two-dimensional materials by all-dry viscoelastic stamping 2D Mater. 1 011002
[22] Uwanno T, Hattori Y, Taniguchi T, Watanabe K and Nagashio K 2015 Fully dry PMMA transfer of graphene on h-BN using a heating/cooling system 2D Mater. 6 01002
[23] Zomer P J, Dash S P, Tombrors N and Van Wees B J 2011 A transfer technique for high mobility graphene devices on commercially available hexagonal boron nitride Appl. Phys. Lett. 99 232104
[24] Zomer P J, Guimaraes M H D, Brant J C, Tombrors N and Van Wees B J 2014 Fast pick up technique for high quality heterostructures of bilayer graphene and hexagonal boron nitride Appl. Phys. Lett. 105 13101
[25] Pizzocchero F, Gammelgaard L, Jenson B S, Caridad J M, Wang L, Hone J, Beggild P and Booth T J 2016 The hot transport properties of single-layer graphene Nat. Commun. 7 11371

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pick-up technique for batch assembly of van der Waals heterostructures Nat. Commun. 7 11894
[20] Friese N and Castellanos-Gomez A 2018 Robotic assembly of artificial nanomaterials Nat. Nanotechnol. 13 641–2
[21] Masubuchi S, Morimoto M, Morikawa S, Onodera M, Asakawa Y, Watanabe K, Taniguchi T and Machida T 2018 Autonomous robotic searching and assembly of two-dimensional crystals to build van der Waals superlattices Nat. Commun. 9 1413
[22] Kretinin AV et al 2014 Electronic properties of graphene encapsulated with different two-dimensional atomic crystals Nano Lett. 14 3270–6
[23] Haigh SJ, Gholinia A, Jalil R, Romani S, Britnell L, Elias DC, Novoselov KS, Pontomarenko LA, Geim AK and Gorbachev R 2012 Cross-sectional imaging of individual layers and buried interfaces of graphene-based heterostructures and superlattices Nat. Mater. 11 764–7
[24] Huang S, Ling X, Liang L, Kong J, Terrones H, Meunier V and Dresselhaus MS 2014 Probing the interlayer coupling of twisted bilayer MoS2 using photoluminescence spectroscopy Nano Lett. 14 5580–8
[25] Liu K, Zhang L, Cao T, Jin C, Qiu D, Zhou Q, Zettl A, Yang P, Liu K, Zhang L, Cao T, Jin C, Qiu D, Zhou Q, Zettl A, Yang P, et al 2014 Evolution of interlayer coupling in twisted molybdenum disulfide bilayers Nano Lett. 14 5496–66
[26] Nayak PK, Horbachtenko Y, Ahs N, Kim G, Lee J-U, Ma KY, Jang A-R, Lim H, Kim D and Ryu S 2017 Probing evolution of twist-angle-dependent interlayer excitons in MoSe2/WSe2 van der Waals heterostructures ACS Nano 11 4041–50
[27] Molina-Mendoza AJ et al 2017 Franscheke as a naturally occurring van der Waals heterostructure heterostructures Nat. Commun. 8 14409
[28] Vildcýk M et al 2017 Exfoliation of natural van der Waals heterostructures to a single unit cell thickness Nat. Commun. 8 14410
[29] Prando G 2017 Van der Waals heterostructures: the natural way Nano. Technol. 12 191
[30] Ganti P, Ghasemi F, Maeso D, Munuera C, López-Eliava E, Frisenda R, De Lara D P, Rubio-Bollinger G, García-Hernandez M and Castellanos-Gomez A 2017 Optical contrast and refractive index of natural van der Waals heterostructure nanosheets of franscheke Beilstein J. Nanotechnol. 8 2357–62
[31] Ray K, Yore AE, Mou T, Jha S, Smithe K KH, Wang B, Pop E and Newaz AK M 2017 Photoreponse of natural van der Waals Heterostructures ACS Nano 11 6024–30
[32] Burzuri E, Vera-Hidalgo M, Giovannelli E, Villalba P, Castellanos-Gomez A and Pérez EM 2018 Simultaneous assembly of van der Waals heterostructures into multiple nanodevices Nanoscale 10 7966–70
[33] Gong C et al 2017 Discovery of intrinsic ferromagnetism in two-dimensional van der Waals crystals Nature 546 263–69
[34] Huang B et al 2017 Layer-dependent ferromagnetism in a van der Waals crystal down to the monolayer limit Nature 546 270–3
[35] Bonilla M, Kolekar S, Ma Y, Diaz HC, Kalappattil V, Das R, Eggers T, Gutierrez H R, Phan M-H and Batzill M 2018 Strong room-temperature ferromagnetism in VSe2 monolayers on van der Waals substrates Nat. Nanotechnol. 13 289
[36] Arai M, Moriya R, Yabuki N, Masubuchi S, Ueno K and Machida T 2013 Construction of van der Waals magnetic tunnel junction using ferromagnetic layered dichalcogenide Appl. Phys. Lett. 107 103107
[37] Hardy W J, Chen C-W, Marcinkova A, Ji H, Sinova J, Natek D and Morosan E 2015 Very large magnetoresistance in Fe0.28TaS2 single crystals Phys. Rev. B 91 54426
[38] Williams T B and Hyde BG 1988 Electronic microscopy of cylinder and franscheke Phys. Chem. Miner. 15 521–44
[39] Makovicky E 1976 Crystallography of cylindrite, Part I. Crystal lattices of cylindrite and incaite Neues Jahrb. f. Mineral. Abhandlungen 126 304–26
[40] Wang W S and Buseck PR 1992 Cylindrite: the relation between its cylindrical shape and modulated structure Am. Mineral. 77 758–64
[41] Meinecke C, Vogt J, Kaden R, Bente K and Buta T 2007 Investigation of synthetic cylindrite microstructures Nucl. Instruments Methods Phys. Res. B 260 317–20
[42] Kaden R, Wagner G, Sturm C, Schmidt–Grund R, von Wencensl H, Prager A, Bente K and Grundmann M 2010 Synthesis and physical properties of cylindrite micro tubes and lamellae Phys. Status Solidi 214 1335–50
[43] Coleman JN et al 2011 Two-dimensional nanosheets produced by liquid exfoliation of layered materials Science 331 568–71
[44] Nicolosi V, Chhowalla M, Kanatzidis MG, Strano MS and Coleman JN 2013 Liquid exfoliation of layered materials Science 340 1226419
[45] Backes C, Higgins TM, Kelly A, Boland C, Harvey A, Hanlon D and Coleman JN 2016 Guidelines for exfoliation, characterization and processing of layered materials produced by liquid exfoliation Chem. Mater. 28 243–55
[46] Buscema M, Groenendijk DJ, Blanter S L, Steele GA, van der Zant HS J and Castellanos–Gomez A 2014 Fast and broadband photoreponse of few-layer black phosphorus field-effect transistors Nano Lett. 14 3347–52
[47] Yuan H et al 2015 Polarization–sensitive broadband photodetector using a black phosphorus vertical p–n junction Nano. Technol. 10 707–15
[48] Engel M, Steiner M and Avouris P 2014 Black phosphorus photodetector for multispectral, high-resolution imaging Nano Lett. 14 6411–7
[49] Mao X, Xu Y, Xue Q, Wang W and Gao D 2013 Ferromagnetism in exfoliated tungsten disulfide nanosheets Nanoscale Res. Lett. 8 430
[50] Salyer PA and Ter Haar LW 1997 Magnetic properties of the mineral, cylindrite (FePb3Sn4Sb2S14) J. Appl. Phys. 81 5163–5
[51] Salyer P A and ter Haar L W 2000 Single–crystal magnetic studies of cylindrite (FePb3Sn4Sb2S14) J. Appl. Phys. 87 6025–7
[52] Sun L, Seanson PC and Chien CL 2000 Finite–size effects in nickel nanowire arrays Phys. Rev. B 61 R6463
[53] Schneider CM, Bessler P, Schuster P, Kirscher J, De Miguel JM and Miranda R 1990 Curie temperature of ultrathin films of fcc–cobalt epitaxially grown on atomically flat Cu (1 0 0) surfaces Phys. Rev. Lett. 64 1059
[54] Farle M, Baberschke K, Stetter U, Aspelmeyer A and Gerhardt F 1993 Thickness–dependent Curie temperature of Gd (0 0 1)/W (1 1 0) and its dependence on the growth conditions Phys. Rev. B 47 11571
[55] Reich D H, Rosenbaum TF, Aepli G and Guggenheim HJ 1986 Ferromagnetism, glassiness, and metamagnetism in a dilute dipolar-coupled magnet Phys. Rev. B 34 4956