PHYSICS

Novel 2D CaCl crystals with metallicity, room-temperature ferromagnetism, heterojunction, piezoelectricity-like property and monovalent calcium ions

Lei Zhang,† Guosheng Shi,† Bingquan Peng,†, Pengfei Gao,†, Liang Chen,†, Ni Zhong,§ Liuhua Mu,§ Lijuan Zhang,§ Peng Zhang,§ Lu Gou,§ Yimin Zhao,§ Shanshan Liang,§ Jie Jiang,§ Zejun Zhang,§ Hongtao Ren,§ Xiaoling Lei,§, Ruobing Yi,§, Yinfeng Zhang,§ Xing Liu,§ Minghong Wu,§ Long Yan,* and Haiping Fang,*

ABSTRACT

Under ambient conditions, the only known valence state of calcium ions is +2, and the corresponding crystals with calcium ions are insulating and nonferromagnetic. Here, using cryo-electron microscopy, we report direct observation of two-dimensional (2D) CaCl crystals on reduced graphene oxide (rGO) membranes, in which the calcium ions are only monovalent (i.e. +1). Remarkably, metallic rather than insulating properties are displayed by those CaCl crystals. More interestingly, room-temperature ferromagnetism, graphene–CaCl heterojunction, coexistence of piezoelectricity-like property and metallicity, as well as the distinct hydrogen storage and release capability of the CaCl crystals in rGO membranes are experimentally demonstrated. We note that such CaCl crystals are obtained by simply incubating rGO membranes in salt solutions below the saturated concentration, under ambient conditions. Theoretical studies suggest that the formation of those abnormal crystals is attributed to the strong cation-π interactions of the Ca cations with the aromatic rings in the graphene surfaces. The findings highlight the realistic potential applications of such abnormal CaCl material with unusual electronic properties in designing novel transistors and magnetic devices, hydrogen storage, catalysts, high-performance conducting electrodes and sensors, with a size down to atomic scale.

Keywords: 2D CaCl crystals, monovalent calcium ions, room-temperature ferromagnetism, heterojunction, coexistence of piezoelectricity-like property and metallicity

INTRODUCTION

Calcium ions are present in rocks, bones, shells, biominerals, geological deposits, ocean sediments and many other important materials [1]. Calcium ions in solution also play major roles in the retention of carbon dioxide in natural waters, water hardness, signal transduction and tissue generation [2–4]. As one of the alkaline earth metals, the calcium atom has two valence electrons according to the octet rule. Up to now, the only known valence state of calcium ions under ambient conditions was +2, and the corresponding crystals with calcium ions are insulating and nonferromagnetic [2]. If the valence of the calcium ion could be modified from divalence, which breaks with conventional theory, it could generate revolutionary new properties for the crystals with calcium ions, inducing a wide range of applications and novel insights.

Since the discovery of C60 [5], carbon nanotubes [6] and graphene [7], carbon has continued to surprise us. Up to now, most studies have focused on the morphologies and properties of those...
carbon-based structures and their complexes with other components having normal stoichiometry. Very recently, Na–Cl and K–Cl systems with abnormal stoichiometries on graphite/graphene under ambient conditions were reported [8]. The crystals are two-dimensional (2D) as they are formed in unsaturated solution on the graphene surfaces. Unfortunately, the atomic structures of these 2D Na–Cl crystals have not been experimentally observed and the corresponding properties have not been explored, partly because of the relatively weak cation–π interaction of the Na cations with the aromatic rings in the graphene surfaces resulting in relatively weak structural stability of the 2D Na–Cl crystals. We note that for divalent cations, the cation–π interactions are much stronger than the interactions for monovalent cations [9]. Here, using cryo-electron microscopy (cryo-EM), we directly observed 2D CaCl crystals on rGO membranes in unsaturated solution under ambient conditions. More importantly, the calcium ions in the CaCl crystals are monovalent (i.e. +1), and consequently the crystals display unexpected metallicity, room-temperature ferromagnetism, and resultant graphene–CaCl heterojunction, coexistence of piezoelectricity-like property and metallicity, together with distinct hydrogen storage and release capability under ambient conditions.

RESULTS AND DISCUSSION

2D Ca–Cl crystals were obtained by soaking ultra-thin rGO membranes (thickness <10 nm) in CaCl₂ solution below the saturated concentration under ambient conditions. This is quite similar to preparation of 2D Na–Cl crystals with Na₂Cl and Na₃Cl crystals in the rGO membranes in NaCl solution below saturated concentration [8], but the rGO membranes here were ultra-thin. The ultra-thin rGO membranes were fabricated by directly forming ultra-thin GO membranes on suitable supporting substrates, which were treated using glow-discharge and desorption methods, and then the ultra-thin GO membranes were reduced to rGO membranes under vacuum and high-temperature conditions (details in Supplementary data, section PS1). To prevent surface contamination from atmospheric air, the freshly prepared membranes were kept in vacuum and inert gas conditions. The supporting substrates were carbon holey films with hole sizes of ∼1.2 μm and center-to-center distances between neighboring holes of ∼2.5 μm. As the ultra-thin rGO membranes are quite flexible, the holey films support as well as provide substrate-free areas with optimized shapes and sizes for rGO membranes to be imaged by TEM. The ultra-thin membranes were then transferred to CaCl₂ solution under protection of argon atmosphere conditions, and were incubated in 5.0 mol/L (M) CaCl₂ solution overnight under ambient conditions. Preparation of Ca–Cl@rGO membranes of such thinness, corresponding to only several layers of the graphene sheets, allows clear characterization of the 2D Ca–Cl crystal structures and minimization of the unfavorable effects of inhomogeneous frozen rates on the membranes during the freezing process.

The ultra-thin membranes coated with original salt solution were then flash-frozen at 20°C with 100% humidity and analyzed by cryo-EM [10,11] to study the in situ formation [12–14] of Ca–Cl crystals (Fig. 1a) in unsaturated solution. As the Ca–Cl crystals are 2D, and very sensitive to the high-energy electrons, the transmission electron microscopy (TEM) measurement conditions required to be optimized. We used 80-kV TEM high-tension together with low-dose mode, which has shown the best balance between the electron-beam damage increased at higher high-tensions and the resolution decreased at lower high-tensions. As a result, a stable single-crystal diffraction pattern was observed in the ultra-thin membranes (Figs 1b and S1). High-resolution cryo-EM images show that such crystals have a lattice spacing of 4.29 ± 0.14 Å (Fig. 1bi), corresponding to a graphene-like honeycomb lattice with a side length of 2.86 ± 0.09 Å. Electron diffraction and fast Fourier analyses [15] of the Ca–Cl lattice yielded a hexagonal lattice with first-order maximal points, that is (1−100) reflections, at (1 ± 0.03)/4.29 Å⁻¹. Further, double-orientated crystals with the same lattice structure of the single Ca–Cl crystal were also observed both on TEM diffraction patterns and high-resolution images (Fig. S2). In Figs 1bi and S1, we can see that there are about five sets of hexagonal diffraction spots at ∼1/2.13 Å⁻¹, too, which correspond to the (1−100) reflections of the graphene. Notably, the (2−200) reflections of the Ca–Cl crystals are coincident with these (1−100) reflections of graphene (Figs 1bii and S1).

Then, we measured the atomic ratio of Ca to Cl and the properties of this Ca–Cl crystal. The ultra-thin Ca–Cl@rGO membranes, which include only several layers of graphene sheets, are too thin to perform elemental analysis and it is too difficult to perform measurements of the electrical, magnetic and hydrogen storage properties. We fabricated thick freestanding rGO membranes (thickness >10 μm) [8,16,17] from an rGO suspension (see Supplementary data, section PS1). The membranes were incubated in 5.0 M CaCl₂ solutions overnight and then centrifuged to remove free solution and dried at 70°C for 4 hours. We called the
Figure 1. Two-dimensional (2D) Ca–Cl crystals from salt solution below the saturated concentration. (a) Schematic drawings of the sample preparation processes. (b) (i) Cryo-EM image of the Ca–Cl crystals in the ultra-thin reduced graphene oxide (rGO) membrane. Inset shows a zoomed-in area of the high-resolution image together with the CaCl crystal structure from the molecular model I, in which Ca and Cl atoms are shown as larger cyan spheres and smaller green spheres, respectively. (ii) Diffraction pattern of a typical crystal structure by cryo-EM in electron diffraction mode, showing a hexagonal lattice pattern with six first-order maxima points at $(1 \pm 0.03)/4.29 \text{Å}^{-1}$. (iii) Fourier transform of the entire bright-field image showing the same hexagonal lattice as in (ii). (c) Atomic ratio of Ca to Cl as a function of the etching time measured by XPS during etching by argon ion for a sample of the dried Ca–Cl@rGO membrane. (d) One stable structure from the molecular model I of CaCl crystal modules adsorbed on graphene sheet from theoretical computations. This structure exhibits P3m1 symmetry (space group: 156), with a lattice constant of 4.92 Å, a thickness of 6.65 Å from Cl to graphene, and a bond length of 3.01 Å from Cl to Ca. Ca, Cl, and C atoms are in cyan, green, and brown, respectively. (e) Electrical resistivity measured using the multimeter with two electrodes connecting with the up and down surfaces of the dried rGO and GO membranes, respectively. (f) Room-temperature ferromagnetism of the dried rGO membrane (red dots) and the dried Ca–Cl@rGO membrane (blue squares) on the magnitude of the applied strains induced at different bending angles were measured. (g) Heterojunction behavior of the dried Ca–Cl@rGO membrane. The current–voltage plot shows typical rectification behavior of the dried Ca–Cl@rGO membrane (red curve) compared with the dried rGO membrane (blue curve) under ambient conditions. (h) Piezoelectricity-like property of the dried Ca–Cl@rGO membrane under ambient conditions. Piezoelectricity-like resultant membranes the ‘dried Ca–Cl@rGO membranes.’ Double- and multiple-orientated Ca–Cl crystals were observed on conventional TEM images of these dried Ca–Cl@rGO membranes (Fig. S4). The hexagonal $(2-200)$ reflections of the Ca–Cl crystals are also consistent with the $(1-100)$ reflections of the graphene sheets, demonstrating that the structure of the Ca–Cl crystals in the dried Ca–Cl@rGO membranes is consistent with the structure of the Ca–Cl crystals observed above in the ultra-thin Ca–Cl@rGO membranes.

Atomic ratios of Ca to Cl in the dried Ca–Cl@rGO membranes were measured by conventional (non-cryogenic) TEM energy-dispersive X-ray spectroscopy (EDS) analysis and X-ray photoelectron spectroscopy (XPS) (see Supplementary data, section PS1). The dark-field high-angle annular dark field scanning TEM (HADDF-STEM) image shows typical distributions of the Ca–Cl crystals in the membranes (Fig. S7). Interestingly, the corresponding TEM EDS analysis (Fig. S7) shows that, in addition to the formation of regular CaCl$_2$ crystals, there are some stable structures with a Ca : Cl ratio of $\sim 1 : 1$, which we denoted as CaCl. To further investigate the stoichiometries of Ca to Cl in the membranes, we performed XPS elemental analysis by etching the membrane with argon ions. The depth profile shows that the Ca : Cl ratios vary from a value below $0.6 : 1$ at the membrane top surface to a stable $\sim 1.2 : 1$ in the inner sheets (Fig. 1c). The membrane surface may exhibit regular CaCl$_2$ because of evaporation of adsorbing salt solution, but on the inner sheets there are mainly CaCl crystals with a stable Ca : Cl ratio of $\sim 1 : 1$.

To discuss the possible structures of the crystals, we used density-functional theory (DFT) [18,19] combined with CALYPSO and USPEX methods [20,21] (see Supplementary data, sections PS1 and PS3) to obtain a stable structure with a Ca : Cl ratio of $1 : 1$, which could be denoted as CaCl crystal (Figs 1d and S11a, model I). Interestingly, as shown in Fig. 1b1, the atomic-resolution TEM images of the CaCl crystal consistent with this structure were experimentally obtained. In this structure, the Ca occupies a plane parallel to the graphene sheet and is located in the center of nonadjacent aromatic rings. Difference charge density shows that there is significant charge transfer between Ca and Cl (Fig. S15). The charge is almost all around the Cl, which shows strong ionic bond characteristics, and the charges of graphene and Ca are assembled to the space between Ca and graphene, demonstrating significant
Figure 2. Electronic structure of model I. (a) Projected band structure and projected DOS on each element in the primitive cell of CaCl alone. (b) Projected band structure and projected DOS on each element in the primitive cell of full model. The size of the points in (a) and (b) indicates the weight of each element’s contribution. Inset: the high symmetry q point path in the Brillouin Zone. $E_F$, Fermi energy.

cation-$\pi$ interaction between Ca and the aromatic rings in graphene. Numerical Bader charge analysis [22] shows that Ca transfers $\sim0.85$ (close to 1) electron to Cl and shares $\sim0.65$ electron with graphene (Table S3). However, the charge shared with graphene is not induced by normal covalent bond but the electron delocalization via cation-$\pi$ interaction, resulting in calcium acting as an ion of +1 valence state (see Supplementary data, section PS4). Chemical bonding analyses of Ca in the Ca–Cl@rGO membranes show that there is only one ionic bonding between the Ca cation and the Cl anion, that is the Ca cation is monovalent, and there is no chemical bonding between the Ca cation and the graphene as the cation-$\pi$ interaction has been named as the non-covalent interaction [23,24] (see Supplementary data, section PS4). Further, there are other possible structures of Ca–Cl, including another CaCl (model II and Fig. S11a) and regular CaCl$_2$, Ca$_3$Cl$_4$, Ca$_3$Cl$_2$ (Figs S11b and S11c), on the graphene surface or confined between graphene sheets.

These CaCl crystals with unusual electronic structure of +1 calcium ions exhibit revolutionary new properties for the crystals with calcium ions, such as metalliclicity, ferromagnetism, heterojunction, coexistence of piezoelectricity-like property and metalliclicity, and adsorption capacity with hydrogen storage. The projected electronic band structure and density of states (DOS) of model I shows that CaCl crystals have distinct metallicity in which Ca plays a dominant role (Fig. 2a). The complexes of graphene with CaCl crystals also have strong metallicity, and the Ca and Cl together contribute $\sim37.7\%$ of the total electronic states near the Fermi level, indicating that CaCl crystals are also metallic in this system (Fig. 2b). More remarkably, compared with pristine 2D graphene and 2D CaCl crystal, the complex shows greatly enhanced electrical conductivity along the horizontal direction, as well as the vertical direction in the case of membrane (Figs S16–18 and Table S5).

Abnormal valence state change of the CaCl crystals was analyzed and confirmed by soft X-ray absorption spectroscopy (near-edge X-ray absorption fine structure spectra, NEXAFS). In addition to the L$_{2,3}$-edge peaks for regular CaCl$_2$ at 347.82 and 351.12 eV, there are new Ca L$_{2,3}$-edge peaks at 348.02 and 351.22 eV (Figs 3, S21 and S22). These new L$_{2,3}$-edge peaks are clearly different from the calcium metal with Ca–Ca bonding and the regular CaCl$_2$ crystals [25+](Fig. 3). Importantly, the photo energy of the new L$_{3}$-edge peak of Ca from the dried Ca–Cl@rGO (~348.02 eV) membrane is larger than both that of pure calcium metal (~347.92 eV) and of regular CaCl$_2$ crystals (~347.82 eV) (see further discussion in Supplementary data, section PS4). To further verify this phenomenon of valence state change, we performed NEXAFS experiments on the dried Cu–Cl@rGO membrane, which was prepared by incubating the rGO membranes with CuCl$_2$ solutions. The L$_{2,3}$-edge peaks for Cu in the dried Cu–Cl@rGO membrane are consistent with the L$_{2,3}$-edge peaks in the regular CuCl crystal, demonstrating that the valence state of Cu has been changed from +2 to +1 in the dried Cu–Cl@rGO membrane (see further discussion in Supplementary data, section PS5).

Figure 3. Calcium near-edge X-ray absorption fine structure spectra (NEXAFS) of the regular calcium metal, the regular CaCl$_2$ crystal and the CaCl crystals in rGO membranes.
Metallic properties of CaCl crystals were confirmed experimentally by measuring the electrical resistivity (see Supplementary data, sections PS1 and PS6). Figure 1e shows that the resistivities of pure dried rGO membranes and Ca–Cl@rGO membranes are both less than ~4.0 × 10^{-2} Ω·cm, indicating their good conductivity. We note that our result on rGO membranes is consistent with metallic properties reported in the literature [25]. In contrast, for dried GO membranes with high resistivity, the average resistivity of dried Ca–Cl@rGO membranes was reduced by at least one order of magnitude (Fig. 1e). The metallicity of CaCl crystals was further demonstrated using conductive atomic force microscopy (AFM) on the graphene sheets (see Supplementary data, sections PS1 and PS6).

Room-temperature ferromagnetism of the CaCl crystals in rGO membrane was also experimentally observed. Figure 1f shows that the saturation magnetic moment (\(M_s\)) of the dried CaCl@rGO membrane is ~2.1 emu/g at 300 K, whereas \(M_s = \sim 0.4\) emu/g for the rGO membrane at the same temperature, showing a ~400% enhancement of magnetism by introducing only ~5% (wt, Fig. S8) calcium to the membrane. For a system with periodic model I configuration, Stoner instability analysis and polarized DFT calculations [26] do not suggest magnetism. Further DFT calculations revealed that the possible origin of such strong ferromagnetism is the edge or defect sites providing unexpected piezoelectricity-like property based on current experimental observations (see Supplementary data, section PS9).

A graphene–CaCl heterojunction was experimentally demonstrated. The current–voltage curve of the dried Ca–Cl@rGO junction membrane under positive and negative gate voltages is obviously asymmetric, indicating typical rectification behavior (Fig. 1g). Further, the +1 valence in these CaCl crystals provides distinct adsorption capacity with hydrogen storage, which not only can fill the insufficient adsorption of metallic calcium atoms, but also overcome the exorbitant absorption of +2 calcium ions (see Supplementary data, section PS10).

The metallic CaCl crystals in rGO membrane possess unexpected piezoelectricity-like property. DFT calculations show that, for 5% geometric elongation along the x direction, the charge distribution of the Ca, Cl, and C atoms in model I has significant relocation (Fig. S29). Our experiment, where a dried Ca–Cl@rGO membrane was connected by two Cu foil electrodes (Figs 1h and S33a), shows a maximal output voltage of ~4 mV when the membrane is bent with an angle of 90° (Fig. S33c) to generate large enough strains in the membrane, and the voltage decreases, reaches zero and may even overshoot to negative values when we remove the bending force. In contrast, we cannot detect a visible output voltage for the rGO membrane with the same bending angles. The voltage of the Ca–Cl@rGO membrane increases with increasing bending degree (Fig. 1h). Although our theoretical calculations strongly support the origin of such phenomenon being the piezoelectricity of the Ca–Cl@rGO membrane, we cannot exclude the possibility that the visible voltage outputs may also be partly induced by other effects during sample bending. Therefore, we refer to this phenomenon as piezoelectricity-like property based on current experimental observations.

**CONCLUSION**

Nature continuously produces surprises, and our results demonstrate another one. 2D Na–Cl crystals with unconventional stoichiometries on graphite or graphene [8] and at extreme conditions, such as high pressure [27] have been reported. Here we observed 2D CaCl crystals with abnormal stoichiometries directly on rGO membranes in unsaturated solution, and remarkably, demonstrate experimentally the +1 valence of Ca ions together with various unexpected properties and applications of the crystals, including metallicity, room-temperature ferromagnetism, and resultant graphene–CaCl heterojunction, coexistence of piezoelectricity-like property and metallicity, as well as distinct hydrogen storage and release capability. Such abnormal CaCl crystals are attributed to the strong cation–π interactions between Ca cations and aromatic rings in the graphene surfaces. As strong cation–π interactions also exist between other metal cations (such as Mg^{2+}, Fe^{2+}, Co^{2+}, Cu^{2+}, Cd^{2+}, Cr^{3+}, and Pb^{2+}) and graphitic surfaces [8,9,23,24,28–32], similar 2D crystals with abnormal valence state of the metal cations on graphitic surface and corresponding abnormal properties as well as novel applications are highly expected. In fact under ambient conditions, CuCl crystals with +1 copper ions can exist stably for several days (see Supplementary data, section PS5) and have room-temperature ferromagnetism (see Supplementary data, section PS9). This would help to overcome the difficulty of the short lifetime in application as a catalyst for the +1 copper ion [33]. We note that it is expected that every metal element has room-temperature ferromagnetism via formation of the corresponding abnormal 2D crystals with an unpaired valence electron in the metal ions. Further, considering the
wide distribution of metallic cations and carbon on Earth, such nanoscale ‘special’ compounds with previously unrecognized properties may be ubiquitous in nature.

METHODS
See Supplementary section PS1 for details.

SUPPLEMENTARY DATA
Supplementary data are available at NSR online.

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AUTHOR CONTRIBUTIONS
H.F. and S.Z. designed the project. L.Z., B.P., L.G., R.Y. and Y.Q. performed the cryo-EM and conventional TEM experiments; G.S., L.C., L.-J.Z. and Y.Z. performed the NEXAFS experiments and analyzed data with L.M.; P.G., L.Z., P.Z. and G.S. performed theoretical analysis of the structure and electrical properties; B.P., L.Y., L.C., N.Z. and C.D. carried out the electrical measurements; G.S., L.C., L.-J.Z. and Y.Z. performed the NEXAFS experiments with liquid-phase and cryogenic TEM. Nat Rev Mater 2018; 10: 776–9.

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