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Electric-field-based control of molecular magnetism in TMPc/Sc₂CO₂ van der Waals systems

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Controlling the magnetic states of molecular structures through electric fields is a major challenge in the area of electron spin manipulation, but it can be achieved upon utilizing the sensitivity of the electron distribution at a heterostructure interface. Here, we found that the magnetic states of a transition-metal phthalocyanine (TMPc) can be effectively controlled through switching the polarization state of monolayer Sc₂CO₂ to achieve a transition between the nonmagnetic and magnetic states of some TMPc molecules. Moreover, the magnetic moments of some structures can be regulated through applying an external vertical electric field. These novel characteristics are caused by the effects of an intrinsic or external electric field on the d-orbital electron transfer and orbital splitting of TMPc molecules. In addition, the p–d hybrid orbital in AgPc will also be affected by the external vertical electric field, ultimately changing the magnetic moment. Therefore, we are very interested in these different magnetic effects, and the fascinating phenomena seen in this study could guide the development of ferroelectric control for use in high-performance electronic memory writing, high-density magnetic memory reading, sensors, and other spintronic devices.

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Introduction

The magnetoelectric effect refers to the coupling of ferroelectricity (FE) and ferromagnetism (FM), which has great potential for use in multifunctional devices.^{1–3} It is an effective method for controlling the electron spin states of nanostructures using an electric field.^{4,5} In general, there are two degenerate polarization states (P↑ and P↓) in two-dimensional ferroelectric materials, and the out-of-plane electric field depends on the polarization states; a polarized electric field can be used to effectively control the properties of magnetic materials. However, natural multiferroic materials are really rare and the magnetoelectric coupling is usually weak; on the other hand, traditional ferroelectric and ferromagnetic materials have different d-orbital electronic requirements.^{6,7} Although there are materials that violate the d₀ rule,⁸ in many cases, it is still pretty challenging to find intense coupling between FE and FM.

In recent years, many two-dimensional (2D) FE materials have been widely studied, such as SnTe,⁹ In₂Se₃,^{10,11} Sc₂CO₂,¹² AgBiP₂Se₆,¹³ CuInP₂S₆,^{14,15} monolayer WTe₂,¹⁶ monolayer group IV

chalcogenides,¹⁷ 2D honeycomb binary compounds,¹⁸ vertically polarized 2D ferroelectric BN, AlN, ZnO, MoS₂, GaSe,¹⁹ transition-metal dihalides,²⁰ and CrSnTe₃.²¹ In addition, theory predicts the existence of a variety of two-dimensional multiferrous materials with coexisting FE and FM, such as C₆N₈H organic networks,²² CrN,²³ Hf₂VC₂F₂,²⁴ and monolayer CrBr₃.²⁵

van der Waals heterostructures provide multiple possibilities for the development of artificial multiferroic materials, usually accompanied by novel physical phenomena and complex coupling principles. For example, the magnetic coupling of CrI₃ bilayers can be switched based on the Sc₂CO₂ polarization state,²⁶ *via* tuning the ferroelectric polarization state in In₂Se₃/FeI₂, the FeI₂ monolayer transforms from ferromagnetic to antiferromagnetic,²⁷ and the polarization state of CuInP₂S₆ monolayers can be switched using an electric field.²⁸ These reports indicate that the use of vdW heterostructures provides a method for achieving artificial multiferroicity.²⁹

In this paper, a two-dimensional artificial multiferroic heterostructure is proposed, which is formed *via* vdW interactions between a transition-metal phthalocyanine molecule (TMPc) and monolayer Sc₂CO₂. Monolayer Sc₂CO₂ has out-of-plane electric polarization of 1.60 μC cm⁻³, and it has been studied in previous research;³⁰ the position of the carbon atom mainly determines the polarization state, and the energy barrier between up (P↑) and down (P↓) polarization is 0.52 eV per unit.³⁰ Besides, phthalocyanine has been widely used in

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chemical catalysis^{31,32} and has been widely studied in spintronics.^{33,34}

We consider the case where the central atom of TMPc is a transition metal of the fourth or fifth period. *Via* first-principles calculations, we found that the magnetic state of TMPc can be switched through tuning the polarization state of the Sc_2CO_2 monolayer, the heterostructure can be switched from a magnetic state to a nonmagnetic state, and an external vertical electric field can control the magnetic moments of some heterostructures. Through further analysis, we know that the coupling variations between TMPc and Sc_2CO_2 are caused by a significant change in the structure arrangement and charge transfer between layers, mainly acting on the d and p orbitals. Moreover, an external electric field can further control charge transfer between layers so that the magnetic moment can be controlled. We also reveal the different origins of the magnetic variations of RuPc ($P\uparrow$) and AgPc ($P\uparrow$). Therefore, this research provides a novel method for designing artificial vdW multifunctional nanoelectronics and devices with strong magneto-electric coupling.

Calculation methods

All the results are obtained based on VASP (the Vienna Ab initio Simulation Package).³⁵ We use the projector-augmented wave (PAW) method³⁶ to indicate the ion potential. For dealing with exchange–correlation effects between electrons, we chose the generalized gradient approximation (GGA) method established based on the Perdew–Burke–Ernzerhof (PBE) functional;³⁷ the Hubbard U correction describes the partial filling of d orbitals, where $U_{\text{eff}} = U - J = 3 \text{ eV}$ ($U = 3, J = 0 \text{ eV}$),³⁸ and different U values do not affect the conclusions. The cutoff energy is 400 eV, and we use 25 Å vacuum space to minimize interactions between adjacent layers. In the relaxation structure, the convergence criterion is that the forces on all atoms are less than at least 0.02 eV \AA^{-1} , and that energy variations are less than at least 10^{-5} eV . Grimme's zero damping DFT-D3 method³⁹ is used to describe the vdW effects of interlayers. We formed a 5×5 Sc_2CO_2 supercell to reduce the interactions of structures; the lattice parameter of Sc_2CO_2 is 3.37 Å, which is similar to previous studies,²⁶ and dipole correction is considered. The differential charge distribution was drawn using Vesta software,⁴⁰ and Bader⁴¹ is used for charge transfer analysis.

Results and discussion

Firstly, we relax the structure to find the lowest-energy adsorption point of a TMPc molecule on Sc_2CO_2 , as shown in Fig. 1(c). The TMPc and Sc_2CO_2 relative monolayer positions can be determined *via* referring to the oxygen atom in the nearest neighbor layer. The relative positions can be divided into top and bottom arrangements. Through rotation and translation, we find that system energy in the top position is lowest. The following research is based on this situation.

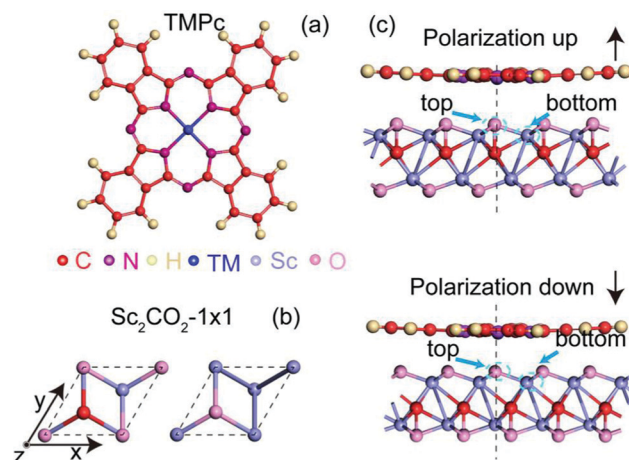


Fig. 1 (a and b) Top views of the metal phthalocyanine and a 1×1 cell of Sc_2CO_2 . (c) Side views of the TMPc/ Sc_2CO_2 system with the $P\uparrow$ and $P\downarrow$ configurations of the 5×5 Sc_2CO_2 supercell monolayer; the arrows point in the out-of-plane polarization direction.

As shown in Fig. 2(a and b), the transition metals of the fourth period show no evident magnetic moment variations on Sc_2CO_2 ; this may be due to the electronegativities of the fourth-period transition metals generally being weaker than those of the fifth-period transition metals. The energies of electrons occupying 4d and 5s orbitals are higher than those occupying 3d and 4s orbitals, and they can more easily participate in charge transfer processes. In terms of the D_{4h} symmetric ligand field structure of phthalocyanine, the magnetic variations can be explained based on the “4+1” splitting phenomenon.^{42,43} TMPc with a central magnetic atom has four suborbitals of similar energy in its d orbital, while one has much higher energy. For example, the bonding of the central atom and ligands in VPc ($3d^3 4s^2$) causes three electrons to be in close orbitals with lower energy, so the total magnetic moment is $3 \mu_B$, and with a further increase in d -orbital-filling electrons, such as in the case of MnPc (or with Fe, Co, Tc, Ru, and Rh), the electrons adopt opposite spin states, resulting in diminishing magnetic moments.

For fifth-period transition-metal atoms, especially Nb, Ru, and Rh, Sc_2CO_2 ($P\uparrow$) has a significant effect on the magnetic moment of the structure. In particular, the magnetic moment of the RuPc ($P\uparrow$) structure is $0 \mu_B$, which means that the introduction of Sc_2CO_2 transforms the magnetic state of RhPc molecules. We mainly analyze the magnetic state transition of RhPc ($P\uparrow$) and the magnetic moment variations of RuPc ($P\uparrow$) in the following discussion. In order to explore changes in the interlayer spacing of the heterostructures after Sc_2CO_2 reversal, we plot the fifth-period interlayer spacing ratios in more detail, as shown in Fig. 2(c); systems that are significantly affected by the polarization of the ferroelectric layer show obvious changes in the interlayer spacing ratio, further affecting the electron density distribution and magnetic moment. We can divide the fifth-period systems into three types, the magnetic moment of a type-a material increases as the electrons occupy adjacent quadruple orbitals, while the magnetic moment of a type-b



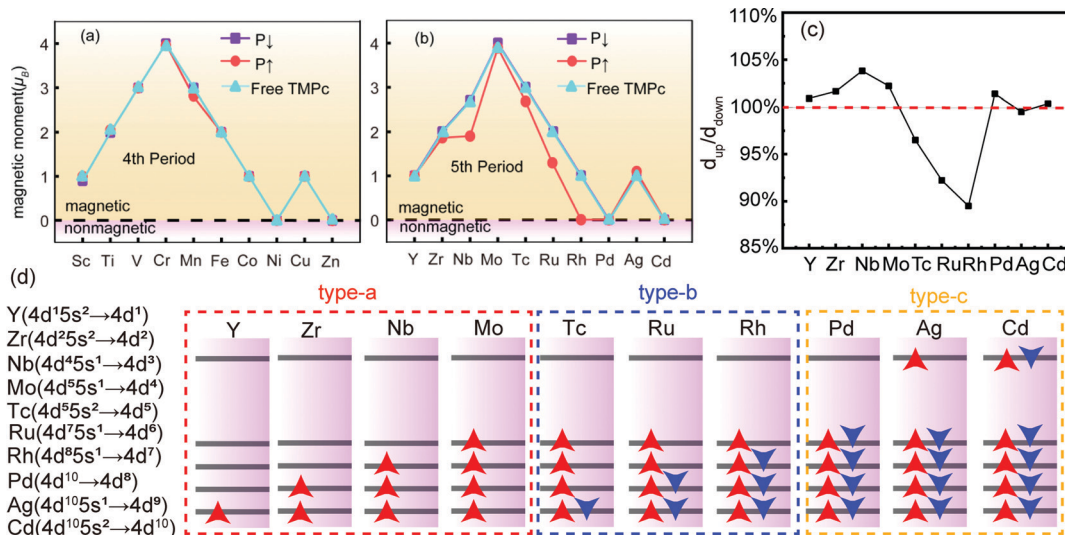


Fig. 2 (a and b) The effects of the substrate polarization direction on the magnetic moment when the central atom of the structure is a transition metal from the fourth or fifth period. (c) TMPc (\uparrow) and TMPc (\downarrow) layer distance ratios. (d) TMPc materials (fifth-period TMs) classified according to the change in magnetic moment.

material decreases as the electrons occupy the opposite spin states of the quadruple orbitals. When the quadruple orbitals are fully occupied, the magnetic moment is $0 \mu_B$, and magnetic moment changes in type-c materials are only related to the electron occupancy of the highest energy suborbitals.

To further assess the magnetic contribution from each d-orbital sublayer in the RhPc system, we provide the following

discussion. Fig. 3 illustrates the arrangements of the d-orbital sublayers. Based on similarities due to the “4+1” splitting phenomenon, the d_{xz} and d_{yz} orbitals of a pure RhPc molecule are degenerate. The two different spin states of the d_{z^2} orbital are evidently split, mainly contributing to the magnetism. The energy level of the $d_{x^2-y^2}$ orbital is significantly higher than those of d_{xy} , d_{xz} , d_{yz} , and d_{z^2} orbitals. When the RhPc molecule

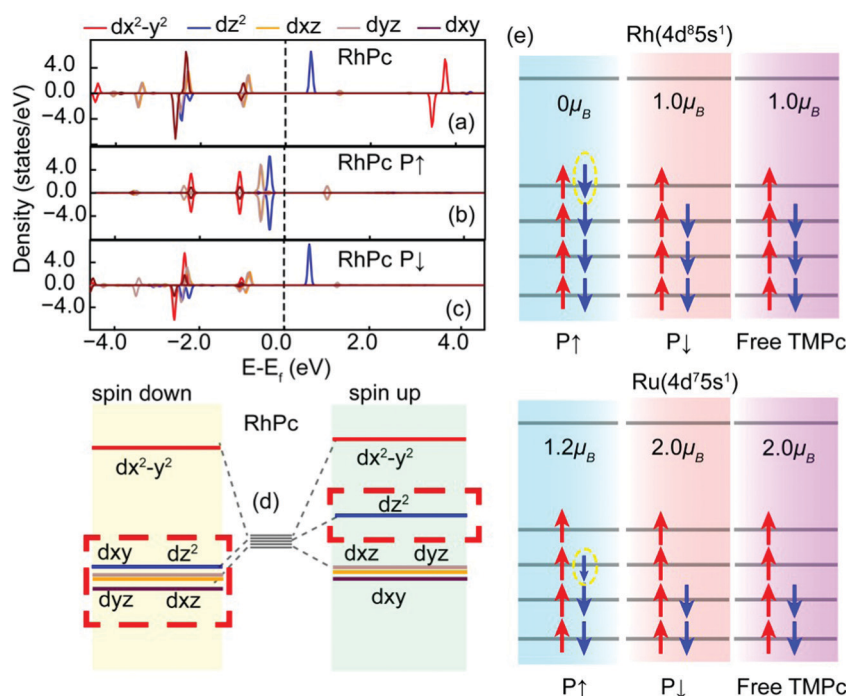


Fig. 3 (a) Density of states (DOS) and (d) energy levels of the d-orbital components of the TMPc central atom (Rh). (b and c) DOS of the d-orbital component of the TM atom (Rh) in two polarized structures. (e) The d-orbital electron arrangements of Rh and Ru in RhPc/Sc₂CO₂ and RuPc/Sc₂CO₂ ($P\uparrow$ and $P\downarrow$) and in RhPc and RuPc molecules without Sc₂CO₂.



is adsorbed on the Sc_2CO_2 monolayer, due to electrons migrating to the $d_{x^2-y^2}$ orbital, the $d_{x^2-y^2}$ orbital moves to a lower energy level and becomes degenerate with the d_{xy} orbital.

As shown in Fig. 3(e), firstly, the outer electron distribution of Rh is $4d^8 5s^1$; after bonding, the d orbital loses two electrons, then RhPc ($P\uparrow$) obtains one electron from the interaction between the ligand and Sc_2CO_2 , and the total magnetic moment is $0 \mu_B$. On the other hand, Rh does not obtain electrons in the $P\downarrow$ structure and there is still electron mismatch in the 4d orbital; the total magnetic moment of the structure is $1 \mu_B$. Secondly, the outer electron arrangement of Ru is $4d^7 5s^1$, and because the electronegativity of Ru is weaker than that of Rh, the number of electrons obtained in the $P\uparrow$ structure is slightly less than one, so the total magnetic moment is more than $1 \mu_B$; the $P\downarrow$ structure has two electrons which are unpaired, and the total magnetic moment of the structure is $2 \mu_B$.

Further analysis of charge transfer in RhPc reveals the physical mechanism. This is illustrated *via* analyzing the density of states, charge transfer, and molecular structures of the two polarization situations. Fig. 4(a and b) shows the density distributions of electron states in RhPc ($P\uparrow$) and RhPc ($P\downarrow$). It can be seen that Sc_2CO_2 ($P\uparrow$) transfers 0.798 electrons of charge to RhPc, which is much more than the 0.016 electrons of charge transferred from RhPc to Sc_2CO_2 ($P\downarrow$); the Fermi level is lower for Sc_2CO_2 ($P\uparrow$), showing a metallic state, which is different from the semiconductor state of Sc_2CO_2 ($P\downarrow$). This transition differs from that of $\text{MPz}/\text{In}_2\text{Se}_3$ ²⁹ charge transfer, but it is essentially the transfer and redistribution of charge under the effects of a polarized electric field. Fig. 4(c) shows the internal transfer trend of electrons, where C_S refers to the carbon in Sc_2CO_2 and C_T refers to the carbon in TMPc, and $C_{\text{Total}} = C_S +$

C_T . The number of electrons transferred has been subtracted from the intrinsic charges of the elements of TMPc and Sc_2CO_2 ; Rh gains more electrons in RhPc ($P\uparrow$). According to Fig. 4(d), the interlayer distance in RhPc ($P\uparrow$) is 2.95 Å, which is shorter than the distance of 3.12 Å between the layers in RhPc ($P\downarrow$); this means that interactions between TMPc and Sc_2CO_2 are more intense in RhPc ($P\uparrow$). The difference in interaction strengths leads to differences in charge transfer and orbital splitting, finally leading to the different magnetic states of TMPc in the two polarization situations.

Based on the response of TMPc to the Sc_2CO_2 polarized electric field, it can be predicted that the external electric field has a significant effect on the charge transfer and distribution, then affecting the total magnetic moment of the system. For the $P\uparrow$ situation, the interactions between TMPc and Sc_2CO_2 are more intense. Therefore, the electric field controls charge transfer more significantly.

Fig. 5(a) further explains the origin of the two different magnetic moments. At a position of 15.80 Å (above the Pc structure), the covalent bond between the Ag atom and the neighboring N atom is stronger. This is different from the almost complete lack of charge interaction between the Ru atom and the N atom. A more complex bond relationship can be seen at 14.75 Å. The main reason for the increase of the RuPc ($P\uparrow$) magnetic moment is that the charge of the Ru d orbital decreases, increasing the net magnetic moment, while the change of the AgPc ($P\uparrow$) magnetic moment is more sophisticated.

As seen in Fig. 5(b), the magnetic moments of AgPc ($P\uparrow$) and RuPc ($P\uparrow$) vary with the electric field, and it can be judged that the origins of the magnetic moment variations of the two structures are quite different according to the differential charge density variations. Fig. 5(c) illustrates charge transfer under an electric field

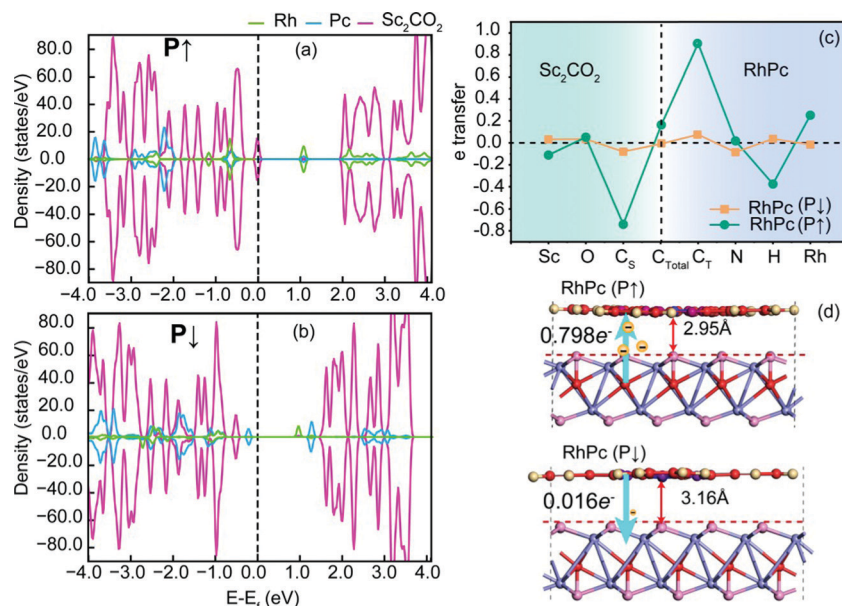


Fig. 4 The effects of substrate polarization direction ((a) up and (b) down) on the DOS of the RhPc/ Sc_2CO_2 structure. (c) The difference in charge transfer number between the RhPc/ Sc_2CO_2 heterostructure and RhPc and Sc_2CO_2 pure molecules. (d) The interlayer distance between the two polarization directions of the heterostructures.



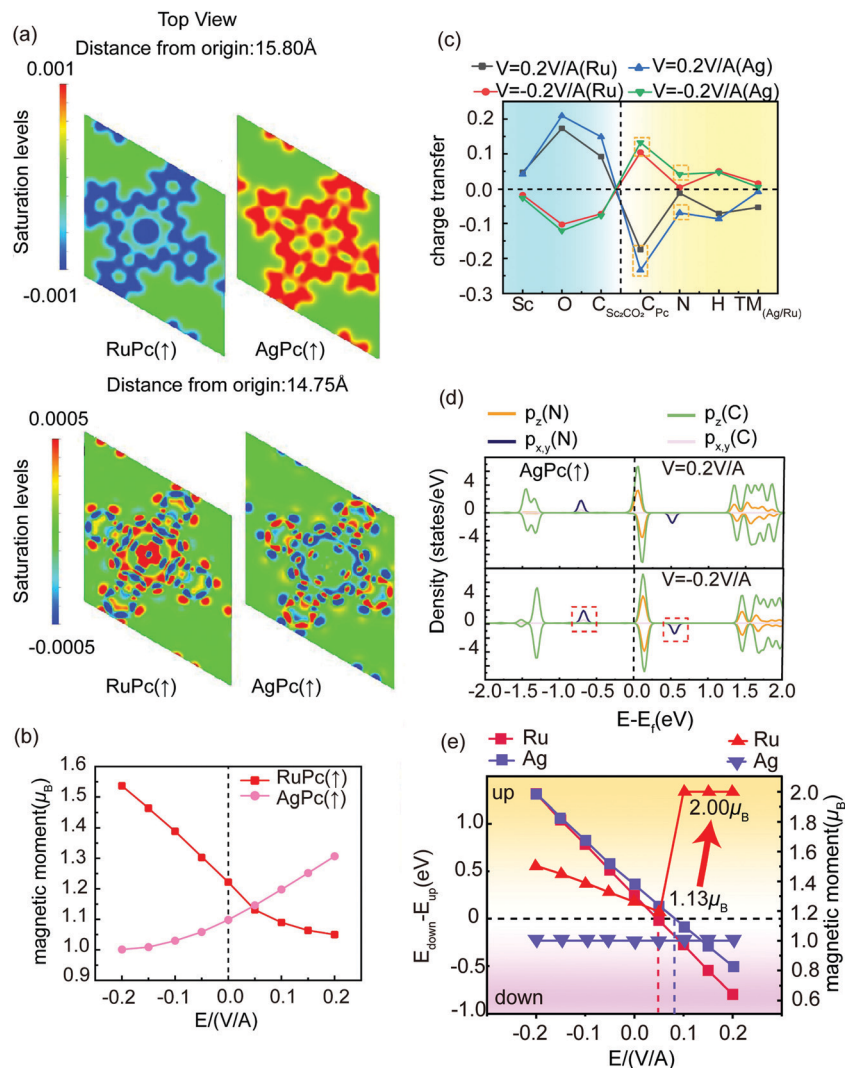


Fig. 5 (a) Differential charge distributions after switching the direction of the electric field when the magnetic moments of RuPc ($P\uparrow$) and AgPc ($P\uparrow$) increase (15.80 Å and 14.75 Å above Pc, respectively). (b) Magnetic moment variations of RuPc ($P\uparrow$) and AgPc ($P\uparrow$) in response to different electric fields. (c) Charge transfer between RuPc ($P\uparrow$) and AgPc ($P\uparrow$) in response to an electric field. (d) DOS of C and N in a AgPc molecule in different electric field directions. (e) The magnetic moment of RuPc is shown to significantly change due to the inversion of Sc_2CO_2 in response to a 0.05 V Å^{-1} electric field.

(the charge without the electric field has been subtracted). In opposite electric field directions, Ru loses different numbers of electrons, which dominates the variation of the magnetic moment, while Ag loses few electrons in the different electric fields, and the main contribution to the magnetic moment from the structure is from N and C_{Pc} , which can be explained based on the magnetic nature of carbonitride;^{44,45} as seen in Fig. 5(d), the p_x and p_y orbitals are degenerate, contributing part of the magnetic moment due to the asymmetry of electrons occupying spin-up and spin-down positions. AgPc ($P\uparrow$) exhibits a semiconductor state in the reverse electric field direction and a half-metallic state in the forward electric field direction. The increase in magnetic moment mainly comes from the splitting of the p_z orbitals of C and N near the Fermi level, which is different from the physical mechanism involved in the case of RuPc/ Sc_2CO_2 ($P\uparrow$).

In order to further explore the influence of the intensity of the interlayer exchange effect on magnetic variations, we

calculated the magnetic moment of the RuPc structure while changing the interlayer spacing of the heterostructure, as shown in Fig. 6; since there is no obvious difference in magnetic moments between the RuPc/ Sc_2CO_2 ($P\downarrow$) structure and RuPc molecule, we mainly study the RuPc/ Sc_2CO_2 ($P\uparrow$) structure, where $E_{\Delta} = E_{d=2.2\text{Å}} - E_d$, $d = 2.7 \text{ Å}$, marked in Fig. 6, is the optimized result for RuPc/ Sc_2CO_2 ($P\uparrow$) in this paper. When the interlayer spacing is too large, the result gradually approaches the behavior of two independent molecules in a vacuum. When the interlayer spacing is too small, the repulsion between electrons is strong, resulting in a drastic change in the structural distribution and the destruction of magnetism.

Conclusions

In summary, we propose and study two-dimensional van der Waals heterostructures composed of a transition metal



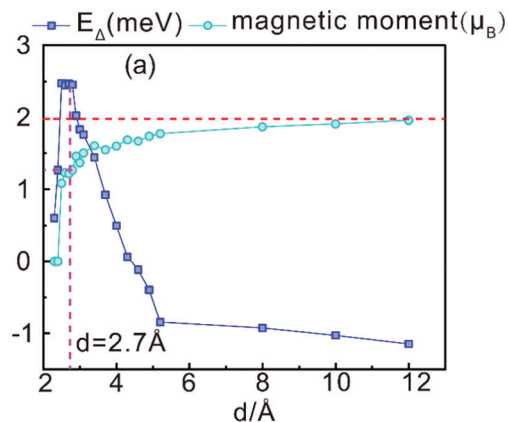


Fig. 6 The effects of the distance d between heterostructure layers on the magnetic moment and energy of the system.

phthalocyanine (TMPc) molecule and the ferroelectric monolayer Sc_2CO_2 via first-principles calculations. The magnetic state of RhPc and the magnetic moments of NbPc and RuPc can be controlled through switching the polarization of Sc_2CO_2 using an electric field. The physical origin of this phenomenon is that ferroelectric interfacial polarization leads to charge transfer and redistribution, which is similar to the “4+1” splitting phenomenon. The variations of the magnetic moments of AgPc ($P\uparrow$) and RuPc ($P\uparrow$) under an external electric field are predicted, and the different physical mechanisms are revealed. For the AgPc ($P\uparrow$) structure, carbon and nitrogen contribute to the magnetism and can be further regulated by the electric field. Therefore, our research provides a theoretical method for developing high-density nonvolatile magnetic memory and it may further accelerate research in related fields.

Conflicts of interest

There are no conflicts to declare.

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