Unravelling the interplay of geometrical, magnetic and electronic properties of metal-doped graphene nanomeshes

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Abstract

Graphene nanomeshes (GNMs), formed by creating a superlattice of pores in graphene, possess rich physical and chemical properties. Many of these properties are determined by the pore geometry. In this work, we use first principles calculations to study the magnetic and electronic properties of metal-doped nitrogen-passivated GNMs. We find that the magnetic behaviour is dependent on the pore shape (trigonal versus hexagonal) as dictated by the number of covalent bonds formed between the 3*d* metal and the passivating N atoms. We also find that Cr and V doped trigonal-pore GNMs, and Ti doped GNMs are the most favourable for spintronic applications. The calculated magnetic properties of Fe-doped GNMs compare well with recent experimental observations. The studied systems are useful as spin filters and chemical sensors.

Keywords: graphene, nanomesh, doping, magnetic and electronic properties, density functional theory

1. Introduction

Two-dimensional (2D) graphene can be considered the basis of different graphetic materials: fullerene, nanotubes and graphite. Due to promising applications, graphene has been the subject of intensive research recently, experimentally as well as theoretically [1–4]. Pristine graphene has a zero band gap which restricts its electro-optical applications. By reducing its dimensionality, quantum confinement results in a finite energy gap, as in quantum dots and nanoribbons [4]. In such structures the band gap depends on the geometry, size, and edge passivation, hence the band gap can be tuned and thus the systems can be used in electro-optical technology [5, 6].

Another graphene-based structure with a finite electronic band gap is a graphene nanomesh (GNM) [7, 8], a 2D system formed by creating a lattice of pores in a graphene sheet. GNMs have been fabricated with pore sizes between 5 and 20 nm [9]. Stable doping of GNMs can be achieved by controlled passivation of the pore edges followed by the chelation of donor or acceptor atoms [10]. GNM edges have also been used for gas separation [11], catalysis [12], sensing [13], and hydrogen storage [14, 15]. Crown ethers have been suggested as a new route towards chemically functionalized graphene [16]. For example, GNM nanostructures have the potential to become next-generation spintronic devices [17–19].

The recent theoretical studies of graphene nanomeshes [7, 10, 11, 14–17, 20–22] and, in particular, the advances in experimental preparation of such systems [9, 12, 13, 16, 18, 19] clearly suggest the need for further detailed investigations, in order to elucidate the structures with highest application potential. In this context, we study the magnetic and electronic structure of a dense regular array of pores in graphene, each of the pores being passivated by nitrogen, and doped with a 3d transition metal atom. While pore passivation by H and O also has been studied [10, 15, 16, 18, 19], our focus here is on N (see also [19]), which is known to form a stable C–N–metal bond, for example, in metal phthalocyanine [23]. Two pore shapes of high symmetry are addressed, namely a trigonal and a hexagonal one, as these are expected to be more stable, and easier to realize.

In section 2, the computational method is presented, then the thermodynamic stability is addressed on the basis of the binding energy, as well as the magnetic moments of different structures (section 3). Our results for the electronic structures are discussed in section 4. Finally, a summary is given in section 5.

2. Methodology and optimized N-GNM structures

Spin polarised density functional theory calculations are performed using the projector augmented-wave pseudopotentials in the Vienna ab initio Simulations Package [24, 25]. For the exchange-correlation energy density functional, the generalized gradient approximation [26, 27] in the scheme of Perdew-Burke-Ernzerhof [28] is utilized to obtain the optimized structures. Projector augmented-wave pseudopotentials for the valence electrons are employed. The wave functions are expanded in plane waves up to a cutoff energy of 600 eV. A Monkhorst–Pack k-point mesh [29] of $12 \times 12 \times 1$ is used for geometry optimisation, for which we employ the standard conjugate-gradient method, until the largest force on the atoms becomes smaller than $0.01\,\mathrm{eV}\ \text{Å}^{-1}$, and the tolerance of total energy reaches 10^{-6} eV. As initial position for the optimisation process, the dopant atom is placed above the centre of the respective pore, at a height of 4 Å. The nanomeshes are modeled as layered three-dimensional systems, with a vacuum of 15 Å in z-direction between the individual layers, in order to avoid any layer-layer interaction. The supercell dimensions in the x-y-plane, chosen to be 6×6 in this work, are kept fixed for all calculations.

In our study, we consider two GNM configurations. A trigonal (*t*) pore is formed by removing 6 C atoms, then passivating the pore edge by 3N atoms; we denote this configuration by *t*-N–GNM. A hexagonal (*h*) pore configuration is formed by removing 12 C atoms, passivating the pore edge with 6N atoms, denoted as *h*-N–GNM. Then 3*d* transition metals are placed in the pore, see figure 1, and their effect on the N–GNM magnetic and electronic properties is studied.

Since we are considering rather light metals in this work, we expect spin-orbit coupling to be of minor importance, as its effect increases with the fourth power of the atomic number. However, one should keep in mind that the spin-orbit

interaction can, in principle, have a distinct effect, especially for the states around the Fermi level, in certain 'critical' cases, e.g. for the half metal system MnBi [30]. In other cases, like NiMnSb, the modification is found to be very small, below 1% [30]. Some discussion of relativistic effects can also be found in a recent review paper [22].

3. Stability and magnetic moment

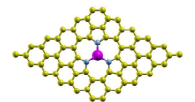
In the optimized structure, *t*- and *h*-N–GNMs have zero magnetic moments. Table 1 shows the bond lengths between metal and N atoms. The bond length decreases as the ionic size decreases from Sc to Cr in both pore configurations, while from Fe to Zn it is almost constant. A slight increase of the bond length is found for Mn (see below). The optimized structure is planar for metal-doped N–GNM in the case of the *h*-pore. However, for the *t*-pore the metal atom is located out of plane of the N–GNM sheet (see figure 1, side view) except for Zn. The vertical distance between the metal and the sheet plane decreases as the ionic size decreases, from 1.1 Å for Sc to 0.5 Å for Cu.

The binding energy (E_b) , table 1, is an indicator of the stability of the respective structure; it can be calculated using the following equation:

$$E_{\rm b} = E_{\rm Metal-N-GNM} - E_{\rm N-GNM} - E_{\rm Metal}, \tag{1}$$

where $E_{\text{Metal-N-GNM}}$, $E_{\text{N-GNM}}$, and E_{Metal} are the energy of the metal-doped N-GNM, the energy of the N-GNM, and the isolated metal in the same volume, respectively. The binding energy assesses the strength of the interaction between N-GNM and the transition metal atom. Roughly speaking, the binding strength ($\sim |E_b|$) increases when considering the series from Zn to Sc, i.e. from right to left in table 1 (with the exception of Co in the t-pore). This is consistent with the fact that along this series the number of empty outershell electron states on the metal increases, allowing for increasing hybridisation with N, and thus stronger binding. It is not surprising that the metals in the t-pore are more strongly bound than the corresponding ones in the h-pore, since there are three bonds with N in the former and only two bonds with N in the latter case. One should note, however, that stronger binding corresponds to longer bond lengths: the latter is consistent with the increase of the covalent radius from Zn ... Sc [31].

The magnetic moment, see table 1, along the sequence from Sc to Cr, with open *d* shells on the metal, is related to the number of remaining electrons after forming the metal—N bonds. Two and three electrons are shared with the N atoms in the *h*- and the *t*-pore, respectively, as shown in figure 1. Therefore the magnetic moment of the metal-doped *h*-N–GNM is larger than that of the *t*-N–GNM doped with the same metal by one. For Mn which is in the high-spin state only two electrons engage with the N atoms in both cases, *t*- and *h*-pore. Note that Mn in the high-spin state is known to have a larger covalent radius than in the low-spin state [31], in agreement with the higher Mn–N bond length in comparison with its 'neighbour' elements.



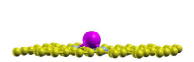




Figure 1. Optimal configuration for metal-doped N–GNM: (from left to right) trigonal (*t*) pore (top view), *t*-pore (side view), and hexagonal (*h*) pore (top view); golden, grey, and pink spheres represent carbon, nitrogen, and metal atoms, respectively.

Table 1. Bond lengths (Å), binding energies (E_b (eV)), and magnetic moments (μ_B), respectively, of metal-doped t- and h-N-GNM.

Metal		Sc	Ti	V	Cr	Mn	Fe	Co	Ni	Cu	Zn
Configuration		$3d^{1}4s^{2}$	$3d^24s^2$	$3d^34s^2$	$3d^54s^1$	$3d^54s^2$	$3d^64s^2$	$3d^{7}4s^{2}$	$3d^84s^2$	$3d^{10}4s^1$	$3d^{10}4s^2$
Bond length	t	1.98	1.90	1.88	1.85	1.92	1.82	1.81	1.82	1.82	1.80
	h	2.22	2.10	2.10	2.10	2.14	2.10	1.98	1.96	2.00	2.00
E_{b}	t	-10.0	-9.4	-8.5	-6.6	-6.6	-6.4	-7.3	-6.5	-5.3	-4.7
	h	-7.4	-6.8	-5.9	-4.3	-4.3	-4.5	-4.5	-4.4	-3.3	-1.4
Magnetic moment	t	0.0	1.0	2.0	2.6	4.8	0.0	3.5	2.0	0.4	0.0
	h	0.7	2.0	3.0	3.4	4.4	3.8	1.6	1.4	0.0	0.2

For Fe in the *t*-pore apparently all un-paired *d* electrons bond with the N atoms, thus the magnetic moment becomes zero. This agrees with the recent experimental observation [19] that Fe assumes a low-spin state if surrounded by a large number of N atoms. The spin state of Fe changes, however, when Fe is embedded in the *h*-pore where only two electrons are shared with the N atom. Thus Fe in the *h*-pore appears to be similar to Fe in a double vacancy [19]. We note that the magnetic moments in the sequence Sc ... Fe are larger for the *h*-pore that for the *t*-pore, which is consistent with general crystal field theory: a large hole should support higher spin states as compared to a small one [21].

For Co and Ni, the three and two un-paired electrons, respectively, bond with the N atoms in the *t*-pore. For these metals, which are near closed shell, the magnetic moment for the *h*-pore is *smaller* than the magnetic moment of the corresponding doped *t*-pore, which indicates a strong hybridisation of the outer shell electrons (*s* and *d* orbitals) with N. Accordingly, the bond lengths for Co and Ni are about 7% smaller than the bond length Fe–N. Regarding closed atomic shells, Cu and Zn are non-magnetic in *t*- and *h*-pores to a good approximation.

In comparison with previous work, we note that for the cases Sc, Fe, and Zn the magnetic moments of metal-doped *t*-N–GNMs are very similar to the same metal substituted into pristine graphene [21], and to graphene with a single vacancy [22]. For metal-doped *h*-N–GNMs our results are very similar for V, Fe, Co, Cu, and Zn embedded in graphene with a double vacancy [21]. For the latter case, the metal–N bonds have the same effect on the magnetic moment as metal–C bonds. Apparently, for the control of the magnetic state it is important how many C atoms are removed, how they are removed (which relates to the shape of the pores), and which passivating element is used.

4. Electronic structure

The density of states (DOS) of pristine graphene, which has been known for a long time [32], features a characteristic linear behaviour near the Fermi energy $(E_{\rm F})$, $\sim |E - E_{\rm F}|$. This behaviour is characteristically modified when passivated pores are created. Figure 2 shows the DOS for the *t*-N–GNM and *h*-N–GNM structures. For the *t*-pore a broad band of states is created around the Fermi energy (-0.2 to 0.7 eV), and the contribution of N states appears at -0.9, -0.7 eV, and around the Fermi energy (-0.1 to 0.2 eV). As compared with pristine graphene, t-N-GNM system is metallic, figure 2(a). For N–GNM with h-pore, the DOS shows semiconducting behaviour with a band gap of 0.8 eV (see figure 2(b)). The N states have a significant participations in the energy range -1.3to $-0.6\,\mathrm{eV}$, and a small contribution in the conduction band (1.2 to 1.5 eV). Clearly the shape of the pore, i.e. the number of removed C atoms and the passivation with N, crucially influences the density of states of the nanomesh.

The surprising (at least at first sight) observation is the metallic behaviour of t-N-GNM. However, the t-pore structure involves three closely interacting N atoms in the pore; this structure p-dopes the graphene by acquiring three electrons from it, as is confirmed by integrating the DOS from the (pristine) valence band edge to the Fermi energy. Hence the Fermi energy is moved compared to the pristine case, rendering the system metallic. From another point of view, we note that semiconducting behaviour is observed for O passivation, see figure 2(c) in [10]: starting from there and replacing O by N, the latter having three electrons less than the former, we arrive at the same conclusion. Of course, these results depend, as is well known [33, 34], on the concrete size of the pore and the supercell: for example, increasing the latter will generally decrease the effect of the pore, i.e. reduce the doping level, thereby shifting the Fermi energy towards the (pristine) valence band edge.

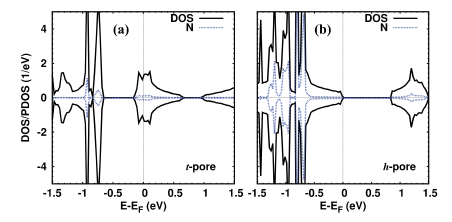


Figure 2. Density of states (DOS) and partial density of states (PDOS) of N atoms, for t-N–GNM (a) and h-N–GNM (b).

When adding Sc as a dopant to the *t*-pore, as compared with *t*-N–GNM (figure 2(a)), the system becomes a semiconductor with a band gap of $0.5 \, \text{eV}$, and the Fermi energy shifts towards higher energy, see figure 3(a). The Sc states appear only in the conduction band above $0.8 \, \text{eV}$, and the N states do not have a significant contribution in the DOS. The symmetry of spin up and spin down states results in a zero magnetic moment. With respect to Sc in the *h*-pore (figure 3(b)), the hybridisation between Sc and N is stronger than for Sc in the *t*-pore, and lies in a broad range (above $-0.3 \, \text{eV}$). This hybridisation creates covalent bonds between Sc and the N atoms. As compared with the DOS of *h*-N–GNM (figure 2(b)), the Fermi energy moves to the conduction band, and the structure becomes metallic. The asymmetry between spin up and spin down states leads to a magnetic moment of $0.7 \, \mu_{\text{B}}$.

The density of states for a t-pore doped with Ti is shown in figure 3(c), demonstrating the covalent bonds between Ti and N orbitals, in particular, at the Fermi energy. The Fermi energy is shifted further to higher energy (compared with the corresponding pore with Sc). A strong hybridisation between Ti and C states appears in the whole energy range of the conduction band. The spin up states at the Fermi energy originate from Ti states. Since the Ti atom has an additional d electron compared with Sc, the magnetic moment of this system is 1 $\mu_{\rm B}$, as we see also in the strong asymmetry of the two spin components. This system may be useful for spintronic devices due to the metallic spin-up states of Ti at the Fermi energy. In the case of the hexagonal pore with a Ti atom (figure 3(d)), the Fermi energy is shifted towards higher energy as compared with h-N-GNM, and covalent bonds due to the hybridisation between Ti and N states are formed. The 3d Ti states for spin up are located around the Fermi energy in the range -0.5 to 0.7 eV. The asymmetry between spin up and spin down components is higher than for Ti in the t-pore, so the magnetic moment becomes 2 $\mu_{\rm B}$. Due to the asymmetry of two spin components, this system can be used as spintronic device.

Figure 3(e) shows the density of states for V inside the *t*-N–GNM. A strong overlap between V and C states is seen in the valence and the conduction bands. In particular, there is a distinct overlap at the Fermi energy with a broad peak, showing that the metal and N atoms have a significant contribution in

one spin component. As compared with t-N-GNM (figure 2(a)), the contribution of V states leads to a splitting of the two spin components. For the h-pore containing V, as compared with h-N-GNM, the Fermi energy is shifted towards higher energy as shown in figure 3(f). Due to the contributions of V states in one spin direction (in the energy range -0.6 to $0.8\,\mathrm{eV}$), there is a clear spin splitting in the DOS in this range. There is no distinct contribution of V states at the Fermi energy, compared with V in t-N-GNM, and the N states appear in the energy -0.2 to $0.8\,\mathrm{eV}$.

Now filling the *t*-pore with a Cr atom, figure 3(g), the DOS is similar to the V doped *t*-N–GNM at the Fermi energy. The contribution of Cr states at the Fermi energy creates a clear spin splitting. Also, the Cr and N states contributions are apparent for one spin direction. The participation of Cr states is pronounced in the other spin component of the conduction band. Adding Cr in the *h*-pore, figure 3(h), a distinct peak contribution of Cr and N states appears at the Fermi energy for one spin. Due to the asymmetry of the two spin orientations, this system can be used as spin filter device.

The density of states of Mn in the t-N-GNM is illustrated in figure 4(a). The Mn states are located in the valence band, and a small splitting in spin components around the Fermi energy is observable. The bonding between N and Mn orbitals appears at $-1.5\,\text{eV}$, and in the energy ranges -1.1 to $-0.8\,\text{eV}$, and -0.4 to $-0.1\,\text{eV}$. The Mn dopant does not change the metallic states of t-N-GNM. For the h-pore doped with Mn, the Mn states contribute to the valence and the conduction bands (spin up only). A small contribution from Mn near the Fermi energy can be noticed. Some hybridisation between Mn and N states is seen in the whole energy range, see figure 4(b). The structure is metallic, in contrast to undoped N-GNM with h-pore.

The effect of Fe on the DOS of t-N-GNM system is addressed in figure 4(c). The Fe states span the energy range from -1.5 to 0.2 eV, and have a significant contribution near 0.5 eV and 1 eV for spin down. The N states are located in a small energy range from -0.4 to -0.1 eV. As in most cases for the t-pore, this system is metallic. For the t-pore with Fe, the metal states have a contribution in one spin direction only in the energy range -0.5 to 1.4 eV, with notable contributions from N states. In the same energy range for the other spin

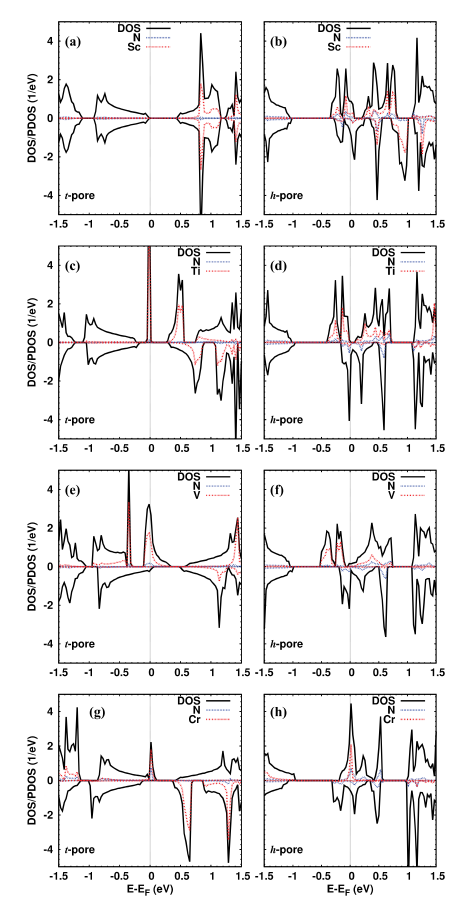


Figure 3. Density of states (DOS) and partial density of states (PDOS), for Sc (a), (b), Ti (c), (d), V (e), (f), and Cr (g), (h) doped N–GNM; *t*-pore (left) and *h*-pore (right).

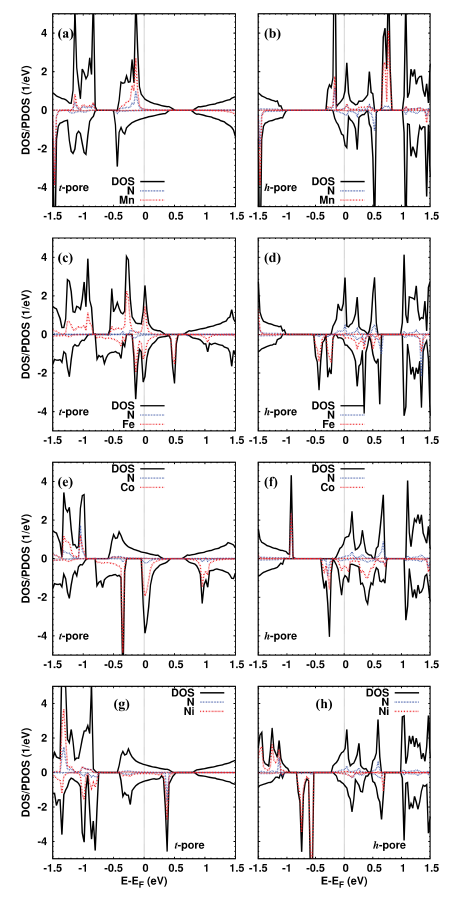


Figure 4. Density of states (DOS) and partial density of states (PDOS), for Mn (a), (b), Fe (c), (d), Co (e), (f), and Ni (g), (h) doped N–GNM; *t*-pore (left) and *h*-pore (right).

Table 2. Spin polarisation ratio in %.

Metal	Sc	Ti	V	Cr	Mn	Fe	Co	Ni	Cu	Zn
t-N-GNM	0	100	91	100	23	0	77	61	7	0
h-N - GNM	19	100	87	66	25	72	6	27	0	77

component, the N states contribute significantly. The splitting in the two spin components around the Fermi energy from -0.5 to $0.7 \, \text{eV}$ is apparent, see figure 4(d). Therefore the magnetic moment in this case is larger than that of Fe in the *t*-pore. Hence an Fe doped *h*-N–GNM is a promising system for spin filter applications.

When a Co atom is placed in a t-pore, see figure 4(e), the electronic structure is very similar to that of V, figure 3(e). However, there are stronger contributions of Co states around $-1.2\,\mathrm{eV}$ which enhance the magnetic moment to $3.5\,\mu_\mathrm{B}$, as compared with the V system. The contribution of Co states inside h-N-GNM in one spin direction is higher than in the other, see figure 4(f). The contribution of N states in the energy range -0.5 to $0.7\,\mathrm{eV}$ is similar to that of N in the h-pore with Fe (figure 4(d)). Metallic states appear at the Fermi energy, with a splitting between the two spin orientations due to the contribution of Co states.

Turning to closed 3d shell atoms (Ni, Cu, and Zn), we show the DOS for Ni doping only, see figures 4(g) and (h). In the tpore the Ni states contribute to the valence band, and at 0.4 eV, figure 4(g). The shift of the Fermi energy is smaller than in previous structures. The main contributions of the Ni atom in the h-pore, figure 4(h), are found in the valence band, and only a little in the conduction band. For Cu in the *t*-pore, the Cu states are found below 0.1 eV, and there is approximate symmetry in the two spin components, therefore the magnetic moment is small, 0.4 $\mu_{\rm B}$. The spin symmetry also appears for Cu inside the *h*-pore, where the Cu states are located in the valence band. For Zn in the t-pore, there is only a shift of the Fermi energy towards higher energy, as compared with undoped t-N-GNM, without any contributions from Zn states, so the system remains non-magnetic. For the h-pore, the Zn states contribute to the DOS at 0.3 eV, but approximate spin symmetry is observed; hence again the magnetic moment is small, $0.2 \mu_{\rm B}$.

For spintronic and spin filter devices, the spin resolved density of states at the Fermi energy is decisive. Hence we finally determine the spin polarisation ratio (P_{DOS}) at the Fermi level, $E_{\rm F}$:

$$P_{\text{DOS}} = \frac{D_{\uparrow}(E_{\text{F}}) - D_{\downarrow}(E_{\text{F}})}{D_{\uparrow}(E_{\text{F}}) + D_{\downarrow}(E_{\text{F}})},\tag{2}$$

where D_{\uparrow} and D_{\downarrow} denote the density of states of the spin up and spin down states, respectively [35, 36]⁶. The results are given in table 2. We conclude that the systems with Ti in both pores, and Cr in the *t*-pore can be used as spintronic devices. Furthermore, we can use V in both pores, Fe and Zn in the *h*-pore, and Co in the *t*-pore for spin filter applications. On the

other hand, Sc and Cu doped N-GNMs have only a low spin filter efficiency.

5. Summary

Using spin-polarised density functional theory, we find that the state of semi-metallic graphene is altered to metal and semiconductor in a graphene nanomesh passivated by N (N-GNM) with trigonal and hexagonal pores, respectively. The covalent bonds between N and the 3d metal are the basis for understanding the magnetic and electronic properties of metal-doped N-GNM. A splitting in the density of states of the two spin components is created due to the contributions of the 3d metals, except for Sc and Zn in the trigonal pore. We find doped N-GNMs to be magnetic for most 3d metals inside the trigonal pore (i.e. except for Sc, Fe, and Zn). In the case of the hexagonal pore, all doped N-GNMs are magnetic except for Cu. A 100% spin polarisation at the Fermi level appears when Ti is used as a dopant in both pores (trigonal and hexagonal), as well as for Cr in the trigonal pore. These structures will produce a pure spin current, which can be used in spintronics and nanoelectronics devices. In addition, we find V inside both pores, Co in the trigonal pore, and Zn in the hexagonal pore, to be useful for spin filter and chemical sensor applications.

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⁶ The 'spin polarisation ratio' is, of course, an often used concept, and it can refer to different quantities like the particle number, the density of states, the resistance, and others. Here, it is the 'DOS' version which is appropriate in view of transport applications. See, e.g. [35, 36].

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