Carbon 171 (2021) 857-868

Contents lists available at ScienceDirect

## Carbon

journal homepage: www.elsevier.com/locate/carbon

### Research Article

# Hunting the elusive shallow n-type donor – An *ab initio* study of Li and N co-doped diamond



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#### ARTICLE INFO

Article history: Received 29 June 2020 Received in revised form 29 August 2020 Accepted 15 September 2020 Available online 25 September 2020

Keywords: Semiconductors Diamond n-type doping Co-dopants Density functional theory

#### ABSTRACT

We report calculated energetics (at the GGA, and at the B3LYP, HSE06 hybrid density functional levels of theory) and electronic properties (B3LYP, HSE06) of Li and N co-doped diamond containing LiN<sub>x</sub> clusters (x = 1-4). Defect energies and structural properties at the GGA and hybrid (B3LYP, HSE06) levels of theory are very similar. Our results for isolated Li and N dopants agree with previous work in that they are not suitable candidates for shallow n-type donors. Here, we investigate the effects of addition of N atoms as a co-dopant with Li, which vary with the Li:N ratio and whether Li is interstitial or substitutional. Reaction energies, geometries, bonding and densities of states (DOS) are examined. Nitrogen stabilises substitutional rather than interstitial lithium. Atom-projected DOS show that the chief contributions to the defect states are not from orbitals on the dopant atoms themselves but from states associated with neighbouring carbon atoms. Only the tetrahedral 1:4 cluster, Li<sub>c</sub>N<sub>4</sub>, with substitutional Li is likely to behave as a shallow donor. We propose a non-molecular synthetic route for preparation of the Li<sub>c</sub>N<sub>4</sub> clusters are not.

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#### 1. Introduction

Due to their increasing availability and affordability from a number of commercial suppliers, chemical vapour deposition (CVD) diamond films are beginning to find a growing number of technological applications [1]. Doping diamond with boron allows p-type semiconducting material to be deposited, with films ranging in conductivity from highly insulating to near metallic [2]. This has enabled the fabrication of many simple p-type devices, such as sensors and electrochemical electrodes [3–5]. But for more complex devices, such as microprocessors, both p-type and n-type behaviour are required. Unfortunately, it has proven extremely difficult to find a dopant for diamond that imparts n-type

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semiconducting characteristics suitable for use in electronic devices [6].

Nitrogen is one possible dopant [7,8] and nitrogen-doped diamond has been successfully synthesised using various CVD techniques [9,10]. However, the experimental nitrogen energy level in diamond lies 1.7 eV below the conduction band minimum (CBM) [11]. This high excitation energy classifies N as a deep donor, and is therefore useless for most electronic devices [12]. Alternatively, phosphorus has been used to n-dope CVD diamond films successfully, but the films are not sufficiently conducting for many device applications [13]. Lithium has also been suggested as a possible shallow donor. Kajihara et al. concluded that interstitial Li atoms should act as shallow donors in diamond, with an excitation energy of only 0.1 eV required to move an electron from the donor level to the CBM [14]. The large energy associated with incorporating Li leads to theoretical predictions of very low solubility of Li in diamond [14]; this can be overcome to some extent using diffusion techniques in which Li concentrations as high as  $10^{19}$  cm<sup>-3</sup> have been reported [15]. In addition, under the standard high-







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temperature conditions used for CVD growth of diamond films, Li has a relatively high mobility [14,16,17], leading to non-uniform concentration profiles, inhomogeneous doping, and formation of Li clusters.

In addition to occupying interstitial sites, Li can also substitute for C or fill a vacancy. It then acts as a deep acceptor, rather than as a shallow donor [18]. Interstitial and substitutional Li can compensate each other, leading to electrical inactivity. This selfcompensation led previously to a decline of interest in such an apparently "unpromising" dopant [16,19,20]. The selfcompensation would, of course, not occur if only one mode of incorporation could be achieved experimentally, but, unfortunately, this has proven extremely difficult [21].

Recently, in pursuit of the elusive shallow donor for n-type diamond films, attention has again focused on possible incorporation of other elements less commonly used as dopants [22], and interest in Li doping has been rekindled, both as a single Li atom, or together with a second element (co-doping) such as boron [23]. Theoretical studies by Mainwood *et al.* and Goss *et al.* have examined the incorporation of Li [16–18,24–27]. In particular, the effects of clustering, which inhibits the ability of Li atoms to perform as shallow donors, have been investigated [16]. These studies also concluded that the lowest-energy interstitial position is a tetra-hedral site, and that incorporation of Li at a pre-existing vacancy site is preferred to interstitial incorporation by at least 1.7 eV [16,18,24,25].

Li can also play an interesting role at the diamond surface. Lithiation, *i.e.* addition of LiO groups to the diamond surface, has been shown both experimentally [28] and theoretically [29,30], to induce negative electron affinity (NEA), as does hydrogenation. Moreover, such lithiated surfaces remain stable at temperatures up to 1000 K. These NEA surfaces enhance the emission of electrons from the bulk diamond into the vacuum [28,30,31], which is a key requirement for field emission devices and thermionic energy converters. Other groups have used DFT to explore the effects of a variety of surface-terminating species, such as MgO, which has also recently been studied experimentally [32], F, Cl, Br, CoO, CuO, and TiO [29,33–37]. Even so, many applications of NEA diamond, particularly thermionic emission, still require n-type doping of the bulk, making identification of potential shallow-donor dopants particularly important [38–41].

Co-doping is a potential way forward. Possible combinations include B–N [42,43], B–P [44] and Li–N. Experimental incorporation of both Li and N in CVD diamond films has been reported [45,46]. High concentrations of Li ( $\sim 10^{19}$  cm<sup>-3</sup>) and N ( $\sim 10^{21}$  cm<sup>-3</sup>) were achieved simultaneously, but with no control of the mode of incorporation. No correlation was found between the Li/N content and the film conductivity, and the films remained electrically inactive.

In this paper, results of both density functional theory (using the generalised gradient approximation, GGA) and hybrid density functional theory (B3LYP and HSE06) calculations are reported to obtain a better understanding of the behaviour – both thermodynamic and electronic – of Li and N in diamond in order to guide new experimental work. While GGA, the simpler DFT implementation, gives good values for the defect thermodynamics that are in agreement with the hybrid methods, it includes a poorer representation of the exchange interaction, plus the band gap is severely underestimated, which is precisely the location of the defect states. We consider in detail the results of co-doping with Li and N and identify shallow donor atom clusters  $Li_CN_x$  (<0.2 eV) that could be formed in the diamond lattice. A  $\text{Li}_{\text{C}}\text{N}_4$  complex comprising a substitutional Li and four adjacent N atoms has been suggested by Moussa et al. as a shallow donor on the basis of calculated activation energies [47], but these authors did not carry out hybrid calculations, consider interstitial Li, examine the

thermodynamics and bonding in detail, present the density of states or consider other arrangements of the nitrogen atoms. We examine the electronic structure and energetics of this and other complexes in detail, and also propose a preparative route to the most promising complex, which should aid future experimental doping studies and realisation of n-type diamond.

#### 2. Theoretical/computational methods

Periodic DFT calculations on doped diamond used a 64-atom (cubic) supercell constructed from a  $2 \times 2 \times 2$  expansion of the cubic diamond unit cell. To assess any supercell size effects and check convergence with cell size, some calculations were repeated with 144-atom (tetragonal), 216-atom (cubic) and 512-atom (cubic) supercells. Details are in the Supplementary Information. For each functional tested, we first generated fully optimised (lattice parameter and basis atom positions) structures for bulk intrinsic diamond. We subsequently optimised supercells of Li-doped, Ndoped and Li-N co-doped diamond. For substitutional dopants, carbon atoms in the supercell were directly replaced by the dopant of interest (Li or N). We label a substitutional Li as Li<sub>C</sub>. Interstitial lithium, which we label with an additional subscript Li<sub>i</sub>, were placed at the T<sub>d</sub> position (tetrahedral site), consistent with previous work [16,24,25,48]. To examine co-doping, between 1 and 4 of the nearest-neighbour carbon atoms surrounding the Li atom were replaced by N. as illustrated in Fig. 1 for LicN<sub>4</sub>.

The energies of the doped supercells were minimised with respect to the positions of the atoms only. Optimisations with fixed unit-cell dimensions strictly give internal energy changes at constant volume. Nevertheless, such defect energies serve also as an excellent approximation for defect enthalpies at constant pressure [49]. Here, calculations relaxing all degrees of freedom gave almost identical results to those in which the supercell lattice parameters were fixed. No symmetry constraints were applied in any of the supercell optimisations. Where relevant, energies of the different possible spin states were calculated. We shall see that these different spin states generally have the same energy to within a few meV.

First, periodic (DFT) calculations were carried out using the plane-wave code CASTEP [50,51], with a plane-wave valence basis set of energy up to 800 eV, and ultra-soft core electron pseudo-potentials. The generalised gradient approximation (GGA) with the Perdew-Burke-Ernzerhof (PBE) exchange-correlation functional was used [52]. The Monkhorst-Pack grid for *k*-point sampling of the Brillouin zone was set at  $4 \times 4 \times 4$ ; convergence with respect to the number of *k*-points was checked. During optimisation, the geometries were deemed to be converged when the total forces on all the atoms were <0.005 eV/Å. The *OptaDOS* utility [53] was used to plot the densities of states (DOS).

Second, we used the *ab initio* periodic code CRYSTAL17, on grounds of computational cost, for hybrid density-functional-theory–Hartree-Fock calculations (B3LYP, HSE06) [54,55]. A number of doped-diamond supercell calculations using the CRYSTAL program have been reported previously [42,43,56–64]. This code uses localised Gaussian basis sets to evaluate energies and associated electronic properties [65–67]. The basis sets used were 6–21 (d-G) for C [68] and pob-TZVP for Li and N [69]. For the calculation of the Coulomb and exchange integrals, tolerance factors of 7, 7, 7, and 14 were used [67]. The convergence criterion for the energy was  $10^{-7}$  au and the *k*-point sampling  $4 \times 4 \times 4$  [70] and satisfactory convergence with respect to these parameters is demonstrated explicitly in the Supplementary Information.

All the defects and defect clusters considered in this paper are neutral (*i.e.* uncharged). The formation energies of the defects,  $\Delta E_{f}$ , are calculated [71] using:



**Fig. 1.** Two LiN<sub>4</sub> clusters studied: (a) substitutional Li (Li<sub>C</sub>N<sub>4</sub>) and (b) interstitial (Li<sub>i</sub>N<sub>4</sub>) with the interstitial atom at the T<sub>d</sub> site. C atoms are yellow, N atoms are blue and Li atoms are green. (A colour version of this figure can be viewed online.)

$$\Delta E_{\rm f} = E_{\rm tot}^{\rm Def} - E_{\rm tot}^{\rm In} - \sum n_i \mu_i \tag{1}$$

where  $E_{tot}^{Def}$  is the energy of the doped (defective) diamond supercell,  $E_{tot}^{ln}$  is the energy of the undoped (intrinsic) diamond supercell of the same size, n the number of atoms added to (+) or removed from (–) the supercell, and  $\mu_i$  the chemical potential of the atom *i* added or removed. The chemical potential for a Li atom was determined from a calculation of bulk Li metal, so that all our chemical reactions relate to incorporation of Li from Li metal. Experimentally, solid Li<sub>3</sub>N [46] or gaseous organo-lithium precursors [72] are often used as the source of Li, and it may well be that it is free Li atoms created in the aggressive hot CVD environment which enter the diamond; we comment later on the consequences for our calculated values of  $\Delta E_{\rm f}$ . For N, we have chosen as reference an isolated N atom rather than molecular N<sub>2</sub> because it is the more relevant species in the growth process in the CVD reactor [73,74]. The C chemical potential was taken from the calculations on pure diamond. All terms in Equation (1) were calculated at the same level of theory.

#### 3. Results and discussion

#### 3.1. Bulk diamond

For validation purposes, we start with the structure and properties of bulk (undoped) diamond. The calculated GGA, B3LYP and HSE06 optimised unit cell parameters are 3.568 Å, 3.594 Å and 3.571 Å, respectively, all in good agreement with experiment (3.567 Å) [75,76]. The corresponding C–C bond length is 1.55 Å (experiment 1.54 Å).

The GGA calculated band gap is 4.06 eV, in agreement with the value (~4.1 eV) calculated using similar methods by Lombardi *et al.* [24] This is significantly lower than the experimental value of 5.45 eV [76], a well-known discrepancy due to the limitations of DFT in its treatment of exchange and correlation, as mentioned in the introduction. Hybrid functionals that incorporate a proportion of exact exchange give more accurate electronic properties and, while accurate calculation of band gaps remains challenging [77,78], such methods offer many advantages over GGA [79]. The variation in band gap from one system to another should also be reliable with hybrid methods.

Fig. 2a shows the density of states (DOS) of pure diamond calculated with HSE06; the band gap calculated with HSE06 is 5.19 eV, which is in much better agreement with the experimental value (5.45 eV) than the GGA result. For comparison, the B3LYP

band gap is 5.72 eV. For reasons of space, we present only HSE06 DOS in this paper since the conclusions from the similar B3LYP DOS are the same.

#### 3.2. Nitrogen in diamond

The energy,  $\Delta E_{\rm f}$ , of substitution of a C atom in the 64-atom supercell by N, where the initial state of N is the atomic gas (Equation (2) in Table 1), is -0.8 eV per N incorporated (GGA), -1.1 eV (B3LYP) and -1.3 eV (HSE06), indicating that the reaction is energetically favourable. Note that if the reactant is the nitrogen molecule N<sub>2</sub> rather than the N atom, as would be appropriate for ambient, rather than CVD reactor, conditions, all energies involving nitrogen in Table 1 increase by ~4.9 eV, making direct incorporation of nitrogen into undefective diamond extremely unfavourable. We repeated the B3LYP calculation for a diamond supercell containing 144 atoms and the value remained unchanged to two significant figures, confirming convergence with supercell size; further tests of convergence with system size are given in the Supplementary Information.

There is considerable symmetry breaking in the optimised structures, regardless of method. The symmetry at the nitrogen site is reduced to  $C_{3v}$ . The three nearest-neighbour C–N bonds in the optimised HSE06 structure are 1.47 Å, which is 0.08 Å shorter than the C–C bonds in diamond, while the fourth is much longer, 2.03 Å, reflecting repulsion between the lone pair on nitrogen and the unpaired electron in the "dangling" bond. Changes in the surrounding C(nearest-neighbour) – C(next-nearest-neighbour) distances are small, approximately  $\pm 1.0\%$ . The lengths of C–C bonds situated further away from the dopant atom are found to remain unchanged, showing that the structure is strained only very locally within a few atoms of the dopant position. Our calculated geometry is in very good agreement with previous work [60].

If there is a pre-existing vacancy in the diamond structure that is subsequently occupied by N (Equation (3) in Table 1), the formation energy is reduced dramatically to -7.5 eV (GGA), -8.4 eV (B3LYP) or -9.3 eV (HSE06). These negative formation energies suggest that addition of N atoms into the diamond structure depends mainly on the availability of vacancy sites [80,81]. Such vacancies arise as a direct result of the CVD growth process, which typically takes place at 1100 K, and have been observed both experimentally [82,83], at concentrations up to 50 ppm [84], and in computer simulations of diamond growth [85–87].

The hybrid (HSE06) DOS in Fig. 2c for the nitrogen-doped supercell shows a deep donor level in the centre of the band gap consistent with experiment [6] and also a minority-spin acceptor



**Fig. 2.** Hybrid (HSE06) DOS of (a) pure diamond, (b) carbon vacancy and (c) N-doped diamond ( $S = \frac{1}{2}$ ) with substitutional nitrogen for the 64-atom supercell. (d) Hybrid (HSE06) DOS of Li<sub>i</sub>-doped diamond (interstitial, T<sub>d</sub> site) for the 65-atom supercell ( $S = \frac{1}{2}$ ). (e) Hybrid (HSE06) DOS of Li-doped diamond (substitutional) from the 64-atom supercell. In (c), (d), (e) and (f) both the total DOS and the projected DOS for the dopant atom are shown. In this and all subsequent figures, the energy of the highest occupied state is arbitrarily set to zero and indicated by the vertical dashed lines. The upper and lower frames show the DOS for spin-up and spin-down electrons, respectively. (A colour version of this figure can be viewed online.)

level, approximately 4.2 eV above the valence band maximum (VBM). Similar features have been noted by Ferrari *et al.* [60] in B3LYP calculations. In agreement with these authors, we too see, interestingly, that the majority of both the donor-band and the acceptor-band states arise not from states on *nitrogen* but on the nearest-neighbour *carbon* atom at 2.03 Å with the unpaired electron in the dangling bond. The Mulliken charges are -0.33e on N, +0.17e on the three C neighbours at 1.47 Å and +0.05e on the C at 2.03 Å and are negligible on all atoms further from the nitrogen. The bond populations between N and the three equivalent carbon neighbours are +0.27e; in contrast between N and the fourth C the bond population is -0.09e indicating an antibonding interaction. The spin density is largest on the carbon at 2.03 Å (+0.76e); on the N itself it is only +0.17e.

#### 3.3. Lithium in diamond

#### 3.3.1. Interstitial Li, Li<sub>i</sub>

Previous work has suggested that the  $T_d$  site is the most favourable for interstitial Li, and so we have used this site in this work [16,24,25,48]. Our solution is metallic, with Li<sub>i</sub>-C bond lengths

of 1.63 Å, 1.67 Å (2×) (HSE06), thus reducing the tetrahedral symmetry to  $C_{2v}$ . There is substantial variation in some of the C(nearest-neighbour) - C(next-nearest-neighbour) bond lengths which vary from 1.49 Å to 1.94 Å. Overall, this suggests that the diamond lattice is significantly strained by the introduction of interstitial lithium, more so than for nitrogen doping, and consistent with the higher formation energies.

Fig. 2d shows the HSE06 DOS. The dopant band, largely comprising states from one of the neighbouring carbons at 1.63 Å and one of the next-nearest carbons at 1.81 Å, merges with the conduction band, giving rise to the Fermi level lying within and close to the bottom of the conduction band. Experimentally realistic, lower Li concentrations, require a larger supercell than we are able to study; these could well be non-metallic and consistent with shallow-donor activity.

#### 3.3.2. Substitutional Li, Li<sub>C</sub>

When Li is substitutional, the tetrahedral symmetry at the site is lost and the symmetry becomes  $C_s$ ; the four nearest-neighbour carbons lie at distances from Li of 1.762 Å, 1.769 Å, 1.769 Å and 1.784 Å (HSE06); these are all~0.1 Å larger than those for Li<sub>i</sub>-C.

#### Table 1

Chemical equations and calculated GGA, B3LYP and HSE06 energies for incorporation of single dopants in diamond. Energies are given in eV per Li or N atom incorporated.  $V_C$  denotes a carbon vacancy while Li and Li<sub>i</sub> denote substitutional and interstitial Li, respectively. Reference states of nitrogen and lithium are the gaseous N atom, and solid Li, respectively.

	GGA	<b>B3LYP</b>	HSE06	
Nitrogen-doped diamond From intrinsic diamond:				
$C_{64}(s) + N(g) \rightarrow C_{63}N(s) + C(s)$	-0.8	-1.1	-1.3	(2)
$C_{144}(s) + N(g) \rightarrow C_{143}N(s) + C(s)$		-1.1		
From pre-existing vacancy				
$C_{63}V_C(s) + N(g) \rightarrow C_{63}N(s)$	-7.5	-8.4	-9.3	(3)
$C_{143}V_C(s) + N(g) \rightarrow C_{143}N(s)$		-8.4		
Li-doped diamond (interstitial Li, Li <sub>i</sub> )				
From intrinsic diamond:				
$C_{64}(s) + Li(s) \rightarrow C_{64}Li_i(s)$	8.4	9.1	9.5	(4)
$C_{144}(s) + Li(s) \rightarrow C_{144}Li_i(s)$		9.1		
Li-doped diamond (substitutional Li)				
From intrinsic diamond:				
$C_{64}(s) + Li(s) \rightarrow C_{63}Li(s) + C(s)$	8.3	8.6	9.1	(5)
$C_{144}(s) + Li(s) \rightarrow C_{143}Li(s) + C(s)$		8.4		
From pre-existing vacancy				
$C_{63}V_{C}(s) + Li(s) \rightarrow C_{63}Li(s)$	1.6	0.8	1.1	(6)
$C_{143}V_{C}(s) + Li(s) \rightarrow C_{143}Li(s)$		0.7		

Accompanying this increase is a decrease in the surrounding C(nearest-neighbour) - C(next-nearest-neighbour) distances by around 4.0%. The lengths of C–C bonds situated further away from the dopant atom are found to remain largely unchanged.

The HSE06 DOS for lithium-doped diamond with substitutional Li (Fig. 2e) is markedly different from that for interstitial Li (Fig. 2d). Removal of a carbon from the unit cell creates four dangling bonds, one on each of the neighbouring C atoms. Insertion of a Li atom at this site supplies one electron and formally removes one dangling bond. The new states which appear at the top of the valence band in the doped system arise from the four nearest-neighbour carbon atoms, with only a small contribution from Li. There are three unpaired electrons – the lowest energy spin state is S = 3/2 which is only  $\approx$  15 meV lower in energy than  $S = \frac{1}{2}$ . The unoccupied spindown peaks form an acceptor band 1-2 eV above the valence band; the states in this band all also come from the four nearestneighbour carbons, with only approximately 5% from the Li. Thus, this structure produces neither an acceptor nor a donor, in agreement with previous work [18,24,25]. We note that as with the substitutional nitrogen defect, while the new levels in the band gap arise from the presence of nitrogen or lithium, the levels are not, per se, N or Li levels.

The Mulliken charges are all very small: +0.08e on Li, -0.06e on the four nearest carbon neighbours, and very small but positive (+0.01e) on the next-nearest carbon neighbours. The spin densities vary rather more on the four nearest-neighbour carbons, decreasing with distance in the order +0.63e, +0.61e, +0.61e and +0.54e. On the Li itself, the spin density is +0.25e. Li–C bond populations are +0.144e, +0.152e, +0.152e and +0.174 increasing with increasing bond length.

#### 3.3.3. Thermodynamics of Li incorporation

GGA energies,  $\Delta E_f$ , for interstitial and substitutional incorporation of Li (Equation (4) and (5), respectively, in Table 1) taking solid Li as the initial Li state are 8.4 eV and 8.3 eV per Li, respectively; the corresponding HSE06 hybrid energies are slightly higher at 9.5 and 9.1 eV, while B3LYP gives intermediate values. Taking gaseous Li as the initial Li state, these energies are reduced by 1.6 eV.

The Li–C bonds in molecular compounds such as methyl lithium (2.31 Å) [88] and phenyl lithium (2.24–2.33 Å) are much longer than those found in Li-doped diamond (at both interstitial and

substitutional sites) [89], suggesting that the Li–C bonds in diamond are greatly compressed and the resulting strain contributes to high positive energies of Li incorporation in the absence of preexisting vacancies.

If the Li dopant occupies an existing vacancy site (Equation (6) in Table 1), the formation energy is significantly reduced to only 1.6 eV (GGA), 0.8 eV (B3LYP) and 1.1 eV (HSE06) taking solid Li as the initial Li state. With the reference state of Li taken as the gaseous atom, the energy of incorporation is again further reduced by 1.6 eV per Li atom, reducing this energy to near-zero or negative values. The formation energy is much lower for this process than incorporation of Li where there is no pre-existing vacancy, and so Li introduced during diamond growth will predominantly occupy such vacancy sites [18]. Unless such vacancy sites are present, or are generated during synthesis, only a small amount of Li is likely to be incorporated, and any introduced Li will occupy both interstitial and substitutional sites due to their similar formation energies.

#### 3.4. Li-N Co-doped diamond

The combination of N deep-donor properties and Li shallowdonor properties through introduction of a Li<sub>C</sub>N<sub>4</sub> complex in diamond has been predicted to promote shallow donor properties on the basis of calculation of donor and acceptor activation energies [47]. However, there has previously been no systematic study of LiN<sub>x</sub> defects for different *x*, no hybrid DFT level calculations, and no investigation of the energetics of dopant incorporation. Thus, we turn to diamond co-doped with Li (interstitial and substitutional) and N, Li<sub>i</sub>N<sub>x</sub> and Li<sub>C</sub>N<sub>x</sub> (x = 1-4), and discuss the synthetic implications. Table 2 lists the calculated formation energies for the formation of LiN<sub>x</sub> defects, with both substitutional and interstitial Li.

We first consider LiN. The energy of formation of Li<sub>i</sub>N co-doped diamond from intrinsic diamond, with Li at the T<sub>d</sub> interstitial site adjacent to the nitrogen (Table 2, Equation (7)), is high (5.0 eV for HSE06), and the energy for the incorporation of interstitial Li into diamond already N-doped even higher (Table 2, Equation (8)). However, the formation energy values are lower than for the Lidoped diamond, suggesting that the strain introduced by the long Li–C bonds is reduced by the presence of nitrogen. The energy of insertion of Li and N atoms into adjacent substitutional sites forming Li<sub>C</sub>N is  $\approx$  2 eV (Equation (9), Table 2), over 2 eV lower than for interstitial Li. The formation energy values for co-doping are again lower than those for Li-doped diamond. Thus, the presence of N favours the incorporation of Li into diamond, and strongly favours substitutional rather than interstitial Li, in contrast to Li incorporation in intrinsic diamond where there is little difference between the formation energies of substitutional and interstitial Li (Equation (4) and (5), Table 1). For N-doped diamond containing a vacancy adjacent to the nitrogen, the formation energy for Li<sub>C</sub>N co-doped diamond reduces even further to ~1 eV (Equation (11), Table 2). Thus, all the different methods indicate that, while still endothermic, Li<sub>C</sub>N co-doped diamond is formed more readily than Lidoped diamond.

Increasing the number of nitrogen atoms in the cluster leads to a successive overall lowering of formation energies and increased favouring of substitutional over interstitial Li (Table 2). Even for Li<sub>C</sub>N<sub>2</sub>, the formation energy from intrinsic diamond is exothermic (Equation (13), Table 2). Even though the formation energies of Li<sub>C</sub>N<sub>3</sub> and Li<sub>C</sub>N<sub>4</sub> are so much more negative than that of Li<sub>i</sub>N<sub>3</sub> and Li<sub>i</sub>N<sub>4</sub>, respectively, the formation energies of the latter are, nevertheless, negative and they could possibly be kinetically stable.

The Supplementary Information contains a detailed survey of the spin states, geometries and electronic structures of  $\text{Li}_i N_x$  (x = 1-3) and  $\text{Li}_C N_x$  (x = 1-3). None of these give rise to a shallow donor band. We now turn to consider in detail the suggested cluster of

#### Table 2

Chemical equations and calculated GGA, B3LYP and HSE06 energies for the formation of  $Li_iN_x$  and  $Li_CN_x$  defects in a 64-atom diamond supercell. Energies are in eV per Li atom incorporated. NV denotes nitrogen-doped diamond, with a vacancy  $V_C$  adjacent to the nitrogen. Reference states of nitrogen and lithium are the gaseous N atom, and solid Li, respectively.

	GGA	B3LYP	HSE06	
LiN				
Interstitial Li, from intrinsic diamond:				
$C_{64}(s) + Li(s) + N(g) \rightarrow C_{63}LiN(s) + C(s)$	4.5	4.9	5.0	(7)
Interstitial Li, from N-doped diamond:				
$C_{63}N(s) + Li(s) \rightarrow C_{63}LiN(s)$	5.3	5.9	6.2	(8)
Substitutional Li, from intrinsic diamond:				
$C_{64}(s) + Li(s) + N(g) \rightarrow C_{62}LiN(s) + 2C(s)$	1.8	2.0	3.1	(9)
Substitutional Li, from N-doped diamond:				
$C_{63}N(s) + Li(s) \rightarrow C_{62}LiN(s) + C(s)$	2.6	3.0	4.3	(10)
Substitutional Li, from NV in diamond:				
$C_{62}NV_{C}(s) + Li(s) \rightarrow C_{62}LiN(s)$	1.2	0.9	2.0	(11)
LiN <sub>2</sub>				
Interstitial Li, from intrinsic diamond:				
$C_{64}(s) + Li(s) + 2 N(g) \rightarrow C_{62}LiN_2(s) + 2C(s)$	2.4	2.7	2.6	(12)
Substitutional Li, from intrinsic diamond:				
$C_{64}(s) + Li(s) + 2 N(g) \rightarrow C_{61}LiN_2(s) + 3C(s)$	-4.6	-4.4	-4.5	(13)
LiN <sub>3</sub> defect				
Interstitial Li, from intrinsic diamond:				
$C_{64}(s) + Li(s) + 3 N(g) \rightarrow C_{61}LiN_3(s) + 3C(s)$	-1.0	0.2	-0.1	(14)
Substitutional Li, from intrinsic diamond:				
$C_{64}(s) + Li(s) + 3 N(g) \rightarrow C_{60}LiN_3(s) + 4C(s)$	-10.8	-10.7	-11.1	(15)
LiN <sub>4</sub> defect				
Interstitial Li, from intrinsic diamond:				
$C_{64}(s) + \text{Li}(s) + 4 \text{ N}(g) \rightarrow C_{60}\text{LiN}_4(s) + 4C(s)$	-2.6	-1.3	-2.6	(16)
Substitutional Li, from intrinsic diamond:				
$C_{64}(s) + \text{Li}(s) + 4 \text{ N}(g) \rightarrow C_{59}\text{LiN}_4(s) + 5\text{C}(s)$	-14.7	-13.6	-14.5	(17)

Moussa *et al.* [47], Li<sub>C</sub>N<sub>4</sub>, and briefly other clusters involving Li and four nitrogens.

#### 3.4.1. Li<sub>i</sub>N<sub>4</sub> and Li<sub>C</sub>N<sub>4</sub>

The relative stabilities of Li<sub>i</sub>N<sub>4</sub> and Li<sub>C</sub>N<sub>4</sub> are reflected in their geometries. Li<sub>i</sub>N<sub>4</sub> is extremely distorted and the corresponding DOS with its mid-gap defect states (see Supplemetary Information), shows that this cluster does not generate shallow donor states.

In contrast,  $Li_CN_4$  is tetrahedral and local  $C_{3v}$  symmetry is preserved at the nitrogens in  $Li_CN_4$ . Fig. 3 collects together the calculated bond lengths, Mulliken charges and bond populations. All Li–N bonds are 1.72 Å, an increase of 0.08 Å from that in  $Li_CN$ , while all N–C bonds are 1.49 Å, 0.02 Å larger than in the system doped with nitrogen alone. The next-nearest-neighbour carbon atoms from the nitrogen are all ~1.55 Å from the nearest-neighbour carbons, showing that the perturbation of the lattice is very local and decays rapidly.

Calculated Mulliken charges are: +0.35e on Li, -0.25e on each of the four nearest nitrogen neighbours, and positive (+0.14e) on the carbons bonded to N. Bond populations are +0.074e for Li–N, reflecting the ionic nature of the bond, +0.231e for N–C, close to +0.27e for that in the purely N-doped system and +0.377e for C(nearest-neighbour) - C(next-nearest-neighbour), this last value reflecting the shorter bond length compared to that in pure diamond.

The DOS of  $Li_CN_4$  is shown in Fig. 4. At the very high dopant concentrations in the range imposed by the supercell calculations, one might expect shallow donor levels to merge with the conduction band, with a diamagnetic solution lower in energy. This is indeed so for both the 64-atom and 144-atom supercells shown in this figure. It is interesting to note that these levels are not associated with orbitals on the Li or N atoms but from the carbons directly bonded to the  $Li_CN_4$  cluster. With larger supercells of 216 atoms and 512 atoms, these occupied states start to separate off from the conduction band. Corresponding band structures are given in the Supplementary Information.

We have also estimated a  $\text{Li}_{C}N_{4}$  donor activation energy of 0.37 eV using a 512-atom supercell, using the empirical marker method [27,90,91] with the known P-donor activation energy of 0.6 eV. Overall, our results suggest that at lower, more experimentally relevant concentrations,  $\text{Li}_{C}N_{4}$  is a shallow donor with an activation energy of 0.2–0.4 eV. This is consistent with the arguments of Moussa *et al.* [47]. Once again, the vast majority of the new states concerned do not involve states on Li or N; they originate from states on the *carbons* bonded to the nitrogen atoms.

We have examined two different geometrical arrangements of the Li<sub>C</sub>N<sub>4</sub> cluster in which one of the nitrogen atoms is further away from the central Li (see Fig. 5). In the initial structures, this nitrogen atom occupied a next-nearest neighbour site (which we label as Cluster 2) or a next-next-neighbour site (Cluster 3). The optimised geometries of the two clusters are shown in the Supplementary Information. There is substantial relaxation, particularly in Cluster 2, where an initial N-N distance of 1.55 Å increases to 1.9 Å. These are both considerably higher in energy than the tetrahedral cluster: 1.14 eV (Cluster 2) and 1.75 eV (Cluster 3), respectively – so under thermodynamic control the tetrahedral cluster will dominate. The DOS for Clusters 2 and 3 (also given in the Supplementary Information), are very different from that in Fig. 4b, with no shallow dopant levels. The DOS for Cluster 2 is broadly similar to that expected for a combination of LiN<sub>3</sub> and LiN (see Supplementary Information), and that for Cluster 3 to a combination of LiN<sub>3</sub> and N.

#### 3.4.2. Stability of Li<sub>C</sub>N<sub>x</sub> clusters

Table 3 includes energies for dissociation of  $Li_CN_x$  into separate  $Li_SN_{x-1}$  and N defects for the 64-atom supercell. All these energies are large and positive, consistent with the decrease in formation energies with increasing N content. For  $Li_CN_4$  (Equation (22), Table 3) we have also examined dissociation to separate Li and four tetrahedrally arranged N atoms (N<sub>4</sub>), a reaction which is also extremely endothermic.



**Fig. 3.** Bond lengths (in italics) and Mulliken net charges (in *e*) in  $Li_cN_4$ . The labels *a*, *b* and *c* denote symmetrically equivalent carbon atoms. (A colour version of this figure can be viewed online.)

Although a single Li<sub>C</sub>N<sub>4</sub> defect is thermodynamically stable with respect to isolated N and Li<sub>C</sub>N<sub>3</sub> defects by 2–3 eV (Equation (21), Table 3), combining reactions (13), (15) and (17) shows that the reaction between Li<sub>C</sub>N<sub>4</sub> and Li<sub>C</sub>N<sub>2</sub> to form two Li<sub>C</sub>N<sub>3</sub> clusters is exothermic by  $\approx 2$  eV. Thus, in principle, Li<sub>C</sub>N<sub>4</sub> clusters will be only kinetically stable but this is unlikely to be a problem in practice because of the large dissociation energy of Li<sub>C</sub>N<sub>4</sub>, Equation (21), and its consequent low mobility.

#### 3.4.3. Lithium-rich clusters

Lithium-rich clusters ( $Li_yN$ , y = 2-4) were also investigated, in which all dopants were positioned at substitutional sites, with a central N atom surrounded by y nearest-neighbour Li atoms. The formation energies for all three such  $Li_yN$  clusters from pure diamond are extremely high (>10 eV). Thus, a lithium-rich cluster configuration is unlikely to be achieved using known techniques.

#### 3.5. Implications for experimental design

Of the clusters we have examined, our calculations suggest, like those of Moussa *et al.* [47], that  $Li_CN_4$  clusters are the most promising as shallow donors. In this section we consider the thermodynamics of their formation in more detail.

The overall formation of  $Li_CN_4$ -doped diamond (with substitutional Li) from diamond containing a vacancy ( $C_{63}V_C$ ) is strongly exothermic:

$$C_{63}V_{C}(s) + \text{Li}(s) + 4 \text{ N}(g) \rightarrow C_{59}\text{LiN}_{4}(s) + 4C(s)$$

$$\Delta E_{(B3YLP)} = -21.3 \text{ eV}$$
(23)

Based on this, Fig. 6 shows a possible pathway for the formation of  $Li_CN_4$  defects via the  $N_4V$  defect. The vacancy formation energy in diamond – the energy to remove one carbon atom – is high,  $\approx 8 \text{ eV}$  for a single vacancy, in agreement with previous work [92,93]. Nevertheless, formation of such vacancies is inevitable during the diamond CVD growth process [82–84]. The energies of formation of  $N_2V$ ,  $N_3V$  and  $N_4V$  from a pre-existing vacancy (*i.e.* here  $C_{63}V$ ) and N atoms are all exothermic (Fig. 6).

The B3LYP energies in Fig. 6 show that clusters  $C_{63-x}N_xV_C$  for x = 2,3,4 but not x = 1 are thermodynamically stable with respect to elimination of the vacancy:

$$C_{63-x}N_xV_C(s) + (x - 1)C_{64}(s) + C(s) \to xC_{63}N(s)$$
(24)  

$$x = 1, \Delta E = -1.1eV$$
  

$$x = 2, \Delta E = 2.3 eV$$
  

$$x = 3, \Delta E = 7.3 eV$$
  

$$x = 4, \Delta E = 12.1 eV$$

Fig. 6 thus shows that under thermodynamic control,  $N_2V$ ,  $N_3V$  and  $N_4V$  centres will all be present, with a lower concentration of NV [94,95]. Consistent with our thermodynamic analysis is the common occurrence of  $N_3V$  and  $N_4V$  clusters in natural diamond and also in synthetic high-pressure high-temperature diamond, where they are known as N3 centres and B centres, respectively [96–99]. Thus, as well as CVD, high-pressure heat treatment of single-substitutional N-doped diamond may provide an alternative route to forming  $N_2V$ ,  $N_3V$  and  $N_4V$  clusters [100]. Indeed, a huge amount of work has been performed, using techniques such as irradiation of the diamond by high-power electrons or laser light, or implantation of N ions, to increase the concentration of vacancies in N-doped diamond in order to produce the  $N_xV$  centres that are proposed for many single-photon quantum applications [101].

The final step in the pathway is to trap a Li atom at the vacancy site in N<sub>4</sub>V under thermodynamic control. Starting from lithium *metal*, this is an endothermic process by 2-3 eV. However, starting from Li *atoms* reduces this energy by 1.5 eV, making the process achievable. Furthermore, if pre-existing substitutional and interstitial Li defects in diamond are mobile, the energy of the defect reactions:



**Fig. 4.** (a) and (b): Hybrid (HSE06) DOS for Li<sub>C</sub>N<sub>4</sub> co-doped diamond (substitutional) for (a) 64- and (b) 144-atom supercells, C<sub>59</sub>Li<sub>C</sub>N<sub>4</sub> and C<sub>143</sub>Li<sub>C</sub>N<sub>4</sub>, respectively. The total DOS and the projected DOS for the dopant atoms are shown. The highest occupied states are arbitrarily assigned a value of zero and are indicated by vertical dashed lines. Note these are diamagnetic conducting solutions. (A colour version of this figure can be viewed online.)



Fig. 5. Unrelaxed geometries of two further possible arrangements of Li<sub>C</sub>N<sub>4</sub> clusters (Cluster 2 and Cluster 3). Cluster 1 is shown in Fig. 1(a). C atoms are yellow, N atoms blue and Li green. (A colour version of this figure can be viewed online.)

#### Table 3

Chemical equations and corresponding GGA, B3LYP and HSE06 energies for the dissociation of  $Li_cN_x$  clusters in a 64-atom diamond supercell. Energies are in eV.

	GGA	<b>B3LYP</b>	HSE06	
Li <sub>c</sub> N				
$C_{62}LiN(s) + C_{64}(s) \rightarrow C_{63}Li(s) + C_{63}N(s)$	5.8	5.6	4.8	(18)
$Li_{C}N_{2}$			<u> </u>	(10)
$C_{61}LIN_2(s) + C_{64}(s) \rightarrow C_{62}LIN(s) + C_{63}N(s)$	5.6	5.4	6.3	(19)
LICIN3 Cool iN $_2(s) + Coa(s) Cool iN_2(s) + CooN(s)$	54	52	53	(20)
$Li_{CN_4}$	5.1	5.2	5.5	(20)
$C_{59}\text{LiN}_4(s) + C_{64}(s) \rightarrow C_{60}\text{LiN}_3(s) + C_{63}N(s)$	3.1	1.8	2.2	(21)
$C_{59}LiN_4(s) + C_{64}(s) \rightarrow C_{60}N_4(s) + C_{63}Li(s)$	16.1	17.6	18.1	(22)

$$C_{59}N_4V + C_{63}Li + C \rightarrow C_{64} + C_{59}LiN_4$$

$$\Delta E = -5.9 \text{ eV}$$
(25)

$$C_{59}N_4V + C_{64}Li_1 \rightarrow C_{64}(s) + C_{59}LiN_4$$

$$\Delta E = -6.2 \text{ eV}$$
(26)

are both highly exothermic.

The calculated energies indicate that temperature control is important to ensure that Li atoms are mobile, while isolated vacancies and the  $Li_CN_4$  clusters remain immobile, since, for example, the energy of the reaction

$$2C_{59}N_4Li + C_{63}V \rightarrow C_{57}LiVN_5 + C_{60}LiN_3 + C_{64}$$
  
$$\Delta E = -3.4 \text{ eV}$$
(27)

is exothermic by -3.4 eV. There is also the possibility of Li<sub>C</sub>N<sub>4</sub> clusters forming larger aggregates that are not shallow donors. While other possible routes involving molecular precursors containing LiN<sub>4</sub> groups have been considered by Moussa *et al.* [47], such precursors and these groups will be rapidly broken down under CVD conditions.

We thus have a potential window-of-opportunity to prepare



**Fig. 6.** B3LYP energies of possible experimental pathways for the synthesis of Li<sub>C</sub>N<sub>x</sub> co-doped diamond. C atoms are yellow, N atoms are blue and Li atoms are green. (A colour version of this figure can be viewed online.)

 $Li_CN_4$  if  $N_4V$  is present in sufficiently high concentration. Direct incorporation of Li into diamond has been demonstrated in our previous experimental reports [45,46] but this has yet to be performed using diamond already containing high concentrations of  $N_4V$  clusters.

#### 4. Conclusions

We have shown that thermodynamic properties of Li–N codoped diamond, calculated at the GGA level are in good agreement with results from hybrid B3LYP and HSE06, even though, as is well known, GGA is unreliable for electronic properties such as band gaps.

Addition of the co-dopant nitrogen to Li-doped diamond

reduces defect cluster-formation energies and stabilises the presence of substitutional rather than interstitial lithium, which we rationalise in terms of the strain present in the different defective structures. Key to this is the reduction in strain due to the absence in the co-doped material with substitutional Li of long nearestneighbour N–C distances of around 2.0 Å. These are present in Ndoped diamond due to the repulsion between the unpaired electron in a carbon dangling bond and the lone pair of electrons on nitrogen. This lone pair now participates in the Li–N bond which is much closer in length to the C–C bond length in undoped diamond. In all the examples we have studied, strain effects are highly localised within a few atomic spacings of the clusters.

It is interesting that the atom-projected densities of states of the Li-, N- and  $Li_CN_4$ -doped diamond show that the chief contributions

to the defect states are not from orbitals on the dopant atoms themselves but from states associated with the neighbouring carbon atoms. We conclude that only the 1:4 cluster,  $Li_CN_4$ , with substitutional Li, is likely to behave as a shallow donor, consistent with the suggestion of Moussa *et al.* [38]. A detailed thermodynamic analysis shows that under typical preparative conditions, different clusters  $Li_CN_x$  (x = 1-4) will be present. We propose a non-molecular synthetic route for the preparation of the  $Li_CN_4$ doped material via prior generation of the N<sub>4</sub>V centre at temperatures at which lithium atoms are mobile and isolated vacancies and  $Li_CN_4$  clusters are not.

The calculations suggest that the  $Li_CN_4$  cluster is, at least, a promising potential candidate for producing n-type diamond. We hope our results will prompt related experimental work. Our future computational work will concentrate on how to assist such experiments in the interpretation of spectroscopic data to distinguish between the different clusters present in order to optimise the concentrations of the  $Li_CN_4$  defect cluster.

#### **CRediT authorship contribution statement**

Sergio Conejeros: Investigation, Visualization, Writing. M. Zamir Othman: Investigation, Writing - original draft. Alex Croot: Investigation. Judy N. Hart: Investigation. Kane M. O'Donnell: Conceptualization, Writing - review & editing. Paul W. May: Conceptualization, Supervision, Writing - review & editing. Neil L. Allan: Conceptualization, Supervision, Writing - review & editing.

#### **Declaration of competing interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

#### Acknowledgements

The authors are grateful to the Public Service Department, Government of Malaysia, EPSRC grant EP/K030302/1 for financial support and Ben Truscott for his help with the software. S.C. gratefully acknowledges the Becas Chile program (CONICYT PAI/INDUSTRIA 74150058) for a postdoctoral grant at the University of Bristol, and FONDECYT Grant 11171063. This work was carried out using the computational facilities of the Advanced Computing Research Centre, University of Bristol http://www.bris.ac.uk/acrc. The raw data for these calculations can be found on the University of Bristol data repository with d.o.i. 10.5523/bris.2ypb34qlp93ox2spy3k3ql4trd.

#### Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.carbon.2020.09.065/.

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