

Optical absorption and luminescence in diamond

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Abstract

The optical properties of diamond are important in theoretical solid-state physics, geology, electronics and diamond synthesis.

This review outlines some relevant aspects of theoretical physics—molecular orbital and other calculations of the properties of point defects, symmetry and group theory, zero-phonon lines and the configuration coordinate diagram.

The experimentally observed spectra can conveniently be divided into infrared (< 1 eV), which are observed in absorption and are mainly vibrational, and 'visible' (1–5.5 eV), which are vibronic and are observed in both absorption and luminescence. 'Visible' spectra include those present naturally, those induced by irradiation, and those induced by irradiation plus annealing.

This review covers all the important point defects observed optically in diamond, and includes some results which are not generally available. The relevance of the experimental results to the current controversy about how to calculate the properties of point defects is brought out.

This review was received in April 1978.

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1. Introduction

1.1. Why study optical absorption and luminescence in diamond?

Diamonds ought to be transparent over most of the spectrum, but many are not. They may absorb and luminesce naturally, or after particle bombardment, and we want to know why. Secondly, diamond is a testing ground for theoretical calculations on point defects, which are responsible for this absorption and luminescence. Thirdly, the mystery of the vanishing interstitial in silicon (Watkins 1965) may be solved by reference to diamond. Fourthly, the study of the different spectra manifested by natural and synthetic diamonds may tell us more about how the former were created, and how to grow the latter more efficiently.

So there are good reasons for studying diamond; but there are difficulties too. Synthetic diamonds are available (§1.7) but nitrogen and boron are the only impurities which can be introduced into them with any degree of control.

1.2. Previous work

There are a number of reviews on the physical properties of diamond, including Berman (1965), Burls (1967) and Field (1979). Reviews particularly relevant to this article are those by Clark (1965), Clark and Mitchell (1971a, b, 1976) and Clark *et al* (1978) on radiation damage, by Davies (1972a, b) on nitrogen in diamond, and by Kemmey and Wedepohl (1965), Vavilov (1975) and Vavilov and Konorova (1976) on semiconducting diamond. Davies (1977a) covers similar ground to the present review.

1.3. The classification of diamond

No two diamonds are identical, but many are sufficiently similar for a *qualitative* classification system to be useful, although some crystals do not fit comfortably into any one category. The system was proposed by Robertson *et al* (1934) and elaborated by various workers. It is outlined in table 1. Although the system was based on measurable macroscopic properties of diamonds, it has become clear that nitrogen and boron impurities are the microscopic bases of the classification.

This is the generally accepted classification scheme. It is worth emphasising that types Ib and IIb are definitive categories, with a particular qualification for membership. Types Ia and IIa are residual classes (although they contain the vast majority of natural diamonds). Type Ia in particular is a mixed bag. As table 1 shows, such diamonds contain aggregates on (100) planes (the 'platelets') which are directly observable in the electron microscope (Evans and Phaal 1962a, b) and which give rise to the anomalous x-ray diffraction 'spikes' (Hoerni and Wooster 1955). They also exhibit defect-induced optical absorption of at least five types: the A, B1 and B2 bands in the infrared, and the N3 system and the secondary absorption edge in the uv (Davies 1972a, b). These are all discussed more fully in §3, where it is shown that not all these properties correlate. Consequently, extensions to the classification scheme have been suggested by Sobolev *et al* (1969a), Davies (1972a, b) and Klyuev *et al* (1972). None of these modifications are generally accepted and they are not consistent with each other.

Table 1. The classification of diamonds. Crystals of mixed types may occur (see figure 1).

Type	Defining feature	Optical absorption			Typical electrical resistance
		Visible	Infrared	ESR	
Ia	Aggregated nitrogen, x-ray diffraction spikes, 'platelets'	Secondary absorption edge at about 300 nm, N3 (415 nm)	A bands B1 and B2 peaks	—	$> 10^{15} \Omega \text{ cm}$
Ib	Paramagnetic singly substitutional nitrogen	Strong featureless absorption increasing in intensity at higher photon energy	Peaks at 0.14 and 0.167 eV	Paramagnetic nitrogen	$> 10^{15} \Omega \text{ cm}$
IIa	—	Fundamental absorption edge, 5.5 eV, often with weak featureless 'tail'	Lattice bands only	Single isotropic line	$> 10^{13} \Omega \text{ cm}$
IIb	p-type semiconductor	Fundamental absorption edge, 5.5 eV, absorption at red end of spectrum	Acceptor bands, free carrier absorption	None	150 $\Omega \text{ cm}$

The best compromise therefore seems to be to adopt the subclassifications IaA, IaB1, IaB2 and IaB3, the characteristics of which are detailed in table 2. Of course, for many purposes the simple Ia category will suffice.

Although they are not given a special category, 'smoky-brown' diamonds appar-

Table 2. Proposed subclassification of type Ia diamonds and the impurity-induced A, B1 and B2 infrared absorption bands. Different authors quote slightly different values of wavenumber. It is not known whether the 780 cm^{-1} feature is B1 or B2; according to Charette (1961a) it is neither. Sobolev *et al* (1969a) refer to N3-containing diamonds as type IaB3, but G Davies (personal communication) has pointed out that there is no evidence for a subclass of diamonds containing predominantly N3 centres.

Type	Optical absorption		
	IR	UV	Other
IaA	480, 1100, 1215, 1282 cm^{-1}	N5, N6, N7, N8 and secondary absorption edge at $\lambda < 330 \text{ nm}$	
IaB1	1010, 1100, 1175, 1332 cm^{-1}	(N9 ?)	
IaB2	330, 1365, 1430 cm^{-1} , 1.25 eV	Bands at 260–280 nm	x-ray spikes, platelets
IaB3	—	N2, N3, N4	

ently form a distinct subclass, with specific EPR, optical and other properties (Shcherbakova *et al* 1975).

Finally, we note that many diamonds are inhomogeneous. That is, parts of the diamond may be type II, other parts type I (Berman 1965, Takagi and Lang 1964, Evans 1976, Hanley *et al* 1977); some parts may be 'platelet-rich' (type IaB2), others 'platelet-free' (type IaA or IaB1) and so on (figure 1 (plate)). This inhomogeneity has bedevilled many attempts to correlate particular features of different diamonds; experimenters must be wary of it.

1.4. Impurities in diamond

Many impurities have been detected in diamond, though one should emphasise that gem-quality crystals are surprisingly pure considering their origin. Apart from nitrogen and boron no element has been found in a concentration greater than 100 ppm (the carbon atom is so small that few atoms can fit onto a normal lattice site without a great deal of strain, and solubilities are therefore low). The fact that so many elements have been detected is a tribute to the sensitivity of the methods used.

The early work used spectrographic analysis or mass spectrometry and residual gas analysis. More recently, the powerful technique of activation analysis has extended earlier results. Fesq *et al* (1973) have conveniently tabulated the elements observed by themselves and by previous workers, so it will be sufficient here to discuss only the impurities of particular interest. Nitrogen is dealt with in §§1.3, 3 and 4. Hydrogen is discussed by Sellschop *et al* (1977).

Boron is the element responsible for p-type semiconductivity in diamond (Chrenko 1973, Lightowers and Collins 1976). Aluminium was originally thought to be the acceptor but was almost certainly present as metallic inclusions in synthetic diamond (Chrenko 1973) and as submicroscopic aluminosilicate droplets in natural crystals (Sellschop 1975, Fesq *et al* 1975) rather than as dispersed single atoms. We cannot say that aluminium cannot be an acceptor in diamond (although recent calculations by Astier and Bourgoin (1976) make it unlikely), but boron plays the dominant role.

Oxygen in diamond has been detected by vacuum fusion and by activation analysis (Kaiser and Bond 1959, Sellschop *et al* 1974, Sellschop 1975). Some of the oxygen is present as a gas in visible inclusions (Melton and Giardini 1974). Some is in chemical combination as submicroscopic aluminosilicate inclusions, though there is a persistent excess over and above what can be contained in the aluminosilicate (Sellschop 1975). This excess presumably could be present substitutionally or interstitially in the diamond lattice (oxygen is a well-known impurity in silicon).

More recently, in a study of 'inclusion-free' diamonds, Melton and Giardini (1976) have claimed that oxygen, and not nitrogen, is the dominant impurity in diamond. This seems unlikely given the internal consistency of Kaiser and Bond's results, and the corroboration by Lightowers and Dean using a different technique, but further work is certainly desirable to resolve the discrepancy (Walker 1976b).

1.5. Nomenclature of optical systems

It is usual to label only the zero-phonon line (§2.1) of an optical system. Clark *et al* (1956a, b) introduced letters and numbers to label all the optical absorption lines they observed (table 3). However, many other zero-phonon lines have since been observed, particularly in luminescence, and the nomenclature adopted, though often descriptive,

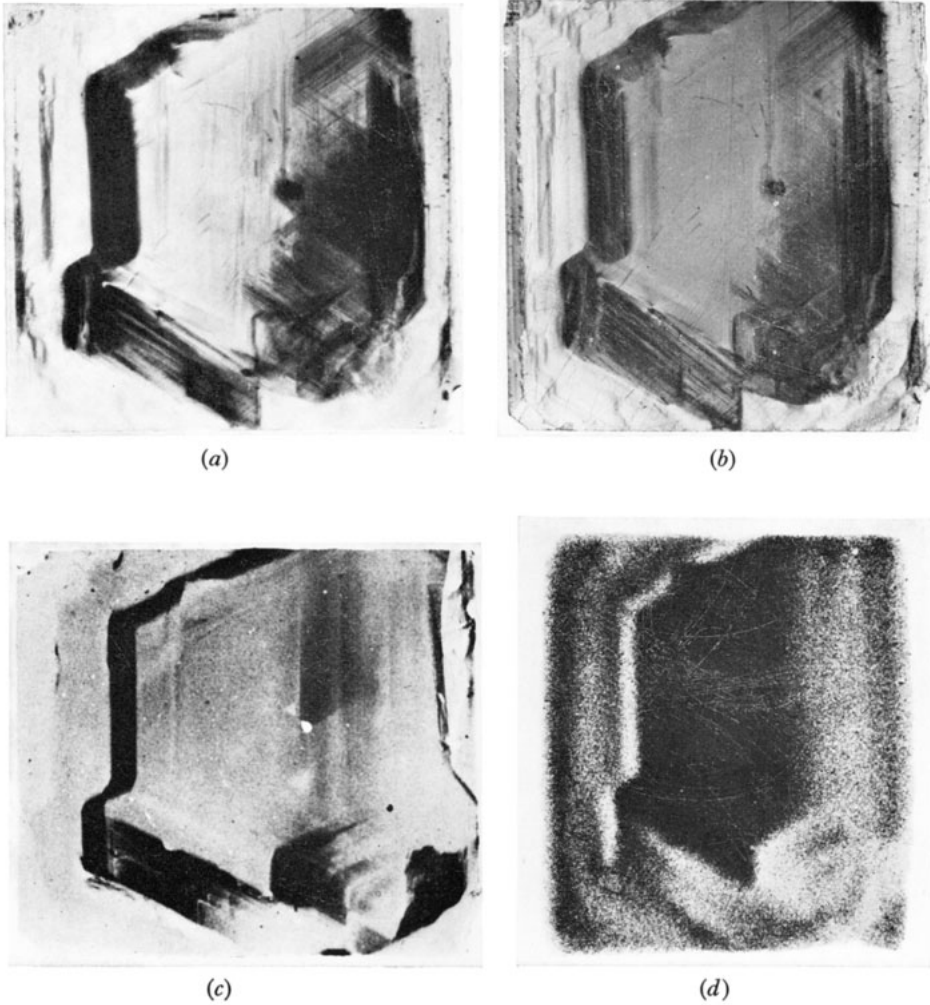


Figure 1. An inhomogeneous diamond, as studied by various techniques. (a) Bragg topograph, MoK α radiation, (b) Bragg topograph, CuK α radiation, (c) ultraviolet transmission topograph, (d) 'spike' topograph. These topographs are discussed in detail by Takagi and Lang (1964), from whose paper they are taken (see also Hanley *et al* 1977).

Table 3. The nomenclature for zero-phonon lines introduced by Clark *et al* (1965a, b). R10 is more frequently called ND1.

Letter	Significance	Range
N	Natural diamond	N1 (1.50 eV) to N9 (5.26 eV)
GR	General radiation (induced in all diamonds by irradiation)	GR1 (1.67 eV) to GR8 (3.00 eV)
R	Radiation (masked in type Ia diamond by the secondary absorption edge)	R9 (3.04 eV) to R11 (3.99 eV)
TR	Type II radiation (not observed in type I)	TR12 (2.64 eV) to TR17 (2.83 eV)
H	Heat treatment (preceded by irradiation)	H1 (0.18 eV) to H18 (3.56 eV)

is not consistent with Clark *et al*. Some are labelled by letters, others by their wavelength or photon energy.

1.6. Paramagnetic resonance

Electron paramagnetic resonance (EPR) or electron spin resonance (ESR)—the terms are used interchangeably because it is usually spin paramagnetism one is concerned with in solids—is a powerful technique for the identification and study of point defects in solids. It has been used with great success in silicon (Watkins 1965) and diamond (Loubser and van Wyk 1978).

1.7. Synthetic diamond

The successful synthesis of diamond was reported by Bundy *et al* (1955). Progress since that time has been reviewed by Bundy *et al* (1973). The production process is now commonplace and for some industrial applications the synthetic material is better than the natural. Synthetics are usually of type Ib, which is surprising because Ib natural crystals are very rare. Semiconducting crystals can be made by doping with boron (the natural variety again being rare), confirming that boron forms the acceptor centre in natural semiconducting diamonds. Weakly semiconducting crystals can be made by doping with aluminium; it is usually assumed that this metal is not electrically active but 'getters' any nitrogen present, thus allowing any boron to remain uncompensated.

Because of the relatively poor quality and small size of synthetic diamonds they are not generally convenient for optical or ESR experiments though a certain amount of work has been reported (du Preez 1965, Bourgoïn *et al* 1972, Wight *et al* 1971, Bourgoïn and Massarani 1976, Massarani and Bourgoïn 1976), and, apart from nitrogen and boron, it is not possible at present to introduce impurities in a controlled manner, which is a considerable handicap. This is one of the reasons why irradiation is so important, as we will see.

1.8. Ion implantation and irradiation

Because of the difficulties of doping diamonds with impurities by diffusion or during growth, some workers have tried ion implantation. In silicon the technique

has been used successfully but in diamond progress has been slower, the major problem being that annealing after implantation to remove radiation damage may result in conversion to graphite; the successes claimed in early work were exaggerated. The induced conductivity was almost certainly due to radiation damage or graphite rather than genuine impurity doping (Clark and Mitchell 1971a, b, Bourgoïn and Walker 1976). However, Vavilov *et al* (1974) apparently succeeded in implanting boron, and lithium appears promising (Vavilov *et al* 1972). But there are still difficulties. Potential dopants have atomic radii larger than carbon, so solubilities will be low.

Irradiation is the only technique which has met with significant success in modifying the optical properties of diamond. Electrons are most convenient because their direction, energy and flux can be easily controlled, and because they produce individual point defects, resulting in sharp lines in the crystal. Neutrons give broad lines, presumably because many defects are produced close together, leading to distorted surroundings and strains. Annealing after irradiation allows defects to become mobile and to migrate through the crystal, becoming trapped at impurities to form new complexes and hence new spectra. This implies that the initial irradiation should be at as low a temperature as possible, to inhibit migration during irradiation, which might lead to confusion about which are the primary defects (i.e. vacancies and interstitials). Most studies have used room-temperature irradiations, but one defect at least is mobile below that (Lomer and Wild 1971, Massarani and Bourgoïn 1976, Lomer and Welbourn 1976).

We will see in §4.3 that some spectra have been ascribed to the vacancy trapped at specific nitrogen complexes, but only one radiation-induced defect has been linked to boron—the 2BD centre (§4.2). This is presumably because natural type IIb diamonds are rare, and furthermore contain only a low concentration (typically 10 ppm) of boron acceptors. Although synthetic diamonds are not generally suitable for optical experiments (§1.7), it seems worthwhile to study irradiated synthetic type IIb to see if they display characteristic spectra.

2. Theory

2.1. The configuration coordinate diagram, zero-phonon lines and the Huang–Rhys factor

The configuration coordinate diagram (figure 2(a)) is useful in explaining optical spectra of defects in solids. A simplified treatment is given here. For further details see Sturge (1967), Curie (1975) and Stoneham (1975).

At $T=0$ K only the lowest ‘vibronic’ (vibrational–electronic) state ($m=0$) is populated. The normalised transition probability to the n th level of the excited state is:

$$W_{n0} = S^n e^{-S}/n!$$

which leads to an absorption spectrum consisting of peaks at $E = E_0 + n\hbar\omega$ (figure 2(b)). In diamond these peaks are usually resolved, even at room temperature; in some cases (e.g. F centres in alkali halides), the individual transitions are broadened and only their envelope is seen. The most probable (Franck–Condon) transition is the vertical one, A \rightarrow B in figure 2(a), with $n=S$. The defect can then relax to C by emission of S phonons, decay to D by emission of a photon, and relax back to A by emission of another S phonons. S —the Huang–Rhys factor—is a measure of the strength of the electron–phonon coupling.

A very important transition occurs between the levels $m=0$ and $n=0$. This is the zero-phonon line (Z in figure 2(b))—a purely electronic transition. Since no phonons are involved, it is very sharp. In principle, its width is determined by the lifetime of the excited state, though in practice random strains in the crystal broaden it to values of about 1 meV. Nonetheless it is considerably sharper than the phonon-assisted transitions, and the way it splits under various perturbations gives valuable information about the symmetry of the defect (§2.5). At high temperatures or large S (strong electron-phonon coupling) the zero-phonon line is weak compared to the

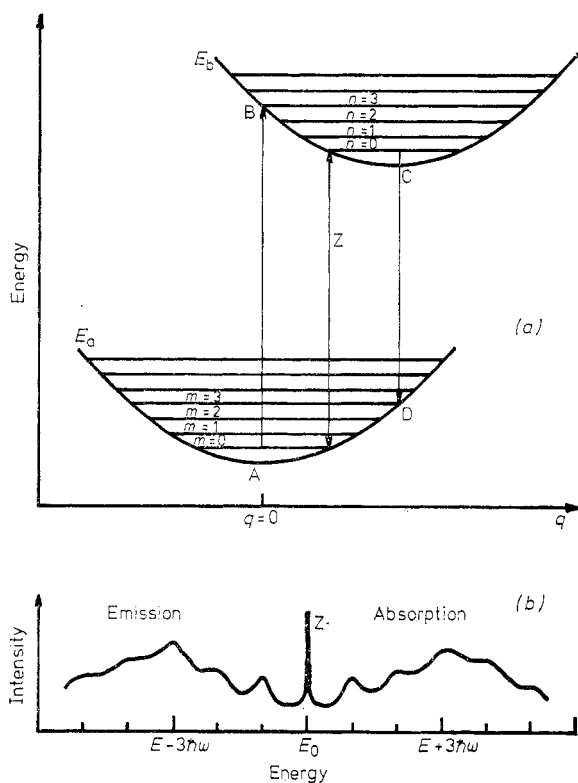


Figure 2. (a) A configuration coordinate diagram showing the ground (E_a) and excited (E_b) electronic states with their vibrational sublevels. Z is the zero-phonon line. (b) The corresponding absorption and emission bands. Note that their maxima correspond to the transitions $A \rightarrow B$ and $C \rightarrow D$. See text.

phonon sidebands. For $S > 6$ it is usually thought to be unobservable, though G Davies (personal communication) has pointed out that the uv zero-phonon lines of the IaA nitrogen defect (§3.2.1.1) are plainly visible though S is greater than 10.

In absorption the phonon-assisted transitions occur at higher energy than the zero-phonon line, whereas in emission they are at lower energy (the Stokes shift) (figure 2(b)). On the simple model, there should be mirror symmetry about the zero-phonon line. If there is not, non-linear coupling or a Jahn-Teller effect must be present. This is the case for some defects in diamond (§4).

The Huang-Rhys factor is an important characteristic of an optical system. It indicates the fractional intensity of the zero-phonon line, the magnitude of the Stokes

shift and the width of the phonon sideband (all measured at $T=0$ K). Thus:

$$I_{zpl} = I_{tot} \exp(-S)$$

$$E_{abs} - E_{emiss} = 2S\hbar\omega$$

$$\Delta^2 = 5 \cdot 6(\hbar\omega)^2 S.$$

In some cases (for example, the GR1 system—see §4.2), the values of S calculated by these three methods are not in agreement, indicating that the simple model does not apply—there is therefore electronic degeneracy, a Jahn–Teller effect (§2.3) or a change in force constants.

2.2. Symmetry and group theory

It is impossible to deal here with group theory and crystal symmetry, but some brief explanations of the notation to be used will be given. For further details, see Tinkham (1964) and Heine (1964).

The diamond lattice is face-centred-cubic, with two atoms per unit cell, at 0,0,0 and $a/4, a/4, a/4$. (Alternatively, we may say that the structure contains two interpenetrating face-centred-cubic lattices with origins at 0,0,0 and $a/4, a/4, a/4$.) The space group is O_h^7 . The point group of the *lattice* is O_h ; the point group of an individual lattice atom site is T_d . The distinction is important because some point defects may have symmetries which are not subgroups of T_d , though they must be subgroups of O_h (Henry and Lonsdale 1965 p340). For example, the A centre, which is probably a nitrogen pair, has D_{3d} symmetry (§3.2), as does the so-called ‘hexagonal’ interstitial site (Weigel *et al* 1973).

The defining symmetry elements of T_d and its subgroups are given in table 4. The non-cubic subgroups of O_h are D_{4h} , D_4 , C_{4v} , D_{2d} , C_{4h} , C_4 , S_4 (tetragonal); D_{3d} , D_3 , C_{3v} , C_{3i} , C_3 (trigonal); D_{2h} , D_2 , C_{2v} (rhombic); C_{2h} , C_2 , C_s (monoclinic); C_i , C_1 (triclinic).

In a solid the normal spectroscopic labels of atomic energy levels cannot be used because the crystal electrostatic field quenches orbital angular momentum. The maximum orbital degeneracy is three in T_d , two in D_{2d} and C_{3v} , and one in the other subgroups of T_d . The spectroscopic labels used in solids are group-theoretical ones—the characters of the irreducible representations of the point groups. There are various systems of notation. In the Mulliken system T denotes triply degenerate orbital levels, E doubly degenerate levels, and A and B non-degenerate levels. Subscripts can be added to distinguish different labels with the same degeneracy, and (where appropriate) to denote odd or even parity. In T_d there are five labels: A_1 , A_2 , E, T_1 and T_2 . The allowed transitions are A_1 - T_2 , A_2 - T_1 , E- T_1 , E- T_2 , T_1 - T_1 , T_2 - T_2 and T_1 - T_2 . For lower symmetries the labels and allowed transitions are given by Tinkham (1964), Heine (1964) and Kaplyanskii (1964a,b). Superscripts ($2S+1$) are used in the conventional way to denote spin degeneracy, e.g. 3T_1 or 1E . Another system uses $\Gamma_1, \Gamma_2, \dots, \Gamma_5$ instead of A, E and T, plus extra labels $\Gamma_6, \Gamma_7, \Gamma_8$ for the double group 2T_d .

2.3. The Jahn–Teller effect

Several defects in diamond manifest a Jahn–Teller effect. The theorem states that any symmetric molecule or point defect with degenerate electronic energy levels

Table 4. The table uses the Schoenflies notation (Tinkham 1964). C_n and S_n denote proper and improper n -fold notations about a specific axis; σ_h and σ_v are reflections in the 'horizontal' and 'vertical' axes ('vertical' is the axis of highest symmetry); σ_d is a σ_v which bisects two two-fold axes. The point groups are labelled according to the symmetry elements they contain, T denoting tetrahedron (more than one axis with $n > 2$), D dihedral (more than 1 axis with $n = 2$). π , σ and V denote linear (parallel to the axis of the centre), planar (perpendicular to the axis of the centre), and volume oscillators. See Kaplyanskii (1964a, b) for further details.

Symmetry system	Point groups	Defining symmetry elements	Electric dipole oscillators
Tetrahedral	T_d	$C_4 \langle 100 \rangle$ $C_3 \langle 111 \rangle$	V (isotropic)
Tetragonal	D_{2d} S_4	$C_4 \langle 100 \rangle$ $S_4 \langle 100 \rangle$	$\left\{ \begin{array}{l} \pi \langle 100 \rangle \\ \sigma \langle 100 \rangle \end{array} \right.$
Trigonal	C_{3v} C_3	$C_3 \langle 111 \rangle$	$\left\{ \begin{array}{l} \pi \langle 111 \rangle \\ \sigma \langle 111 \rangle \end{array} \right.$ V
Rhombic I	C_{2v} D_2	$C_2 \langle 100 \rangle$ C_2 or $\sigma_v \langle 011 \rangle$ C_2 or $\sigma_v \langle 0\bar{1}1 \rangle$	$\pi \langle 100 \rangle$ $\pi \langle 011 \rangle$ $\pi \langle 0\bar{1}1 \rangle$
Rhombic II	C_{2v} D_2	$C_2 \langle 100 \rangle$ C_2 or $\sigma_v \langle 010 \rangle$ C_2 or $\sigma_v \langle 001 \rangle$	$\pi \langle 100 \rangle$ $\pi \langle 010 \rangle$ $\pi \langle 001 \rangle$
Monoclinic I	C_2 σ_h	C_2 or $\sigma \langle 110 \rangle$	$\pi \langle 110 \rangle$ $\sigma \parallel \langle 110 \rangle$
Monoclinic II	C_2 σ_h	C_2 or $\sigma \langle 100 \rangle$	$\pi \langle 001 \rangle$ $\sigma \parallel \langle 001 \rangle$
Triclinic	C_1	E	V

will distort in order to lift the degeneracy and lower its energy. The only exceptions are linear molecules and Kramers degeneracy. The operation of the theorem is indicated schematically in figure 3, which shows a doubly degenerate state of energy E_0 which splits as a result of a distortion Q . The splitting for a small distortion is linear in Q , so one level has a lower energy E_- than the undistorted configuration. Elastic

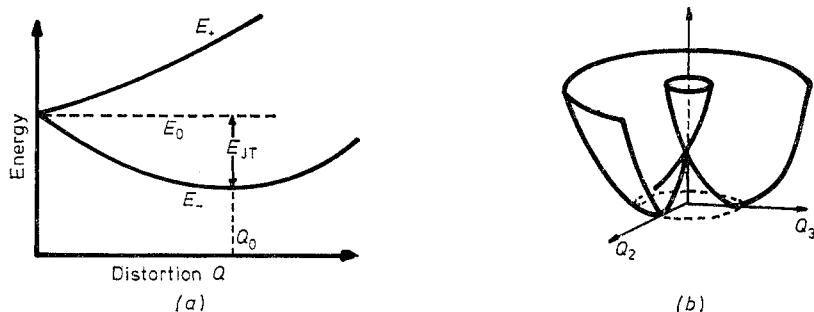


Figure 3. (a) Schematic representation of the lowering in energy E_{JT} resulting from the splitting of a degenerate electronic level due to a Jahn-Teller distortion. (b) Revolution of (a) about the vertical axis—the 'Mexican Hat'.

restoring forces which are quadratic in Q limit the distortion to a finite value Q_0 , and the consequent energy reduction is the Jahn–Teller energy E_{JT} . See Sturge (1967), Ham (1972), Englman (1971) and Stoneham (1975) for recent review articles.

There are four main ways in which the Jahn–Teller effect may manifest itself in point defects. Firstly, the symmetry observed may be lower than expected. This is the static Jahn–Teller effect. The substitutional nitrogen donor in diamond is an example (§3.2). It might be expected to be tetrahedral, but the extra electron occupies an antibonding orbital localised on the nitrogen and one single carbon atom, resulting in a distortion to trigonal symmetry. There are four equivalent distortions, corresponding to the four $\langle 111 \rangle$ directions. Secondly, if the temperature is high enough or the energy barrier low enough, reorientation between equivalent distortions may occur. This is the dynamic Jahn–Teller effect. The original symmetry has effectively been restored, the electronic degeneracy being replaced by an equivalent vibronic degeneracy. It happens to the nitrogen donor in diamond at about 800 K (Shul'man *et al* 1967, Loubser and Van Ryneveld 1967), and is thought to be important at the GR1 centre (§4.2).

The third manifestation of the Jahn–Teller effect was not recognised as such until much later, from the work of Ham (1972), who showed that a dynamic Jahn–Teller effect can cause a ‘quenching’ of orbital angular momentum, spin–orbit splitting and uniaxial stress splitting (the Ham effect). Fourthly, it may affect vibronic band shapes (§2.1).

2.4. Calculations of the properties of defects

The spacing and symmetry of energy levels in atoms and molecules can now be calculated with considerable accuracy. But the analogous problem in semiconductors—the energy levels of point defects—has been solved only in certain special cases. ‘Shallow’ defects (i.e. with energy levels within 0.1 eV of band edges) can be treated successfully using the effective mass approximation (Stoneham 1975).

‘Deep’ levels are another matter, and diamond, though a favourite ‘model crystal’, is difficult in that the band gap is wide (5.5 eV), so that all defects are ‘deep’. Nonetheless, a great deal of work has been carried out, and if there is doubt as to which methods of calculation are most appropriate, and disagreements about symmetries of ground states and stabilities and configurations of defects, the calculations have nonetheless provided valuable insights. They have also been handicapped by lack of identification, or in some cases misidentification, of defects. It is not appropriate to discuss the calculations here; for more details, consult Stoneham (1975) and the references therein. However, it does seem that the ‘defect molecule’ calculations of Coulson and his co-workers successfully predict the energy levels of vacancies in diamond (§§4.1 and 4.2).

Some attempts have been made to calculate the one-phonon infrared spectra associated with defects (Dawber and Elliott 1963, Szigeti 1963), particularly the ‘nitrogen platelets’. Since it now appears that the platelets do not contain the bulk of the nitrogen these calculations could not be expected to be successful. However, some of the defects have now been correctly identified (§3), and further calculations are desirable.

One notable result is that the peaks of the defect-activated one-phonon absorption are not expected to coincide exactly with the phonon frequencies of the perfect lattice. If they did, then all defects would produce the same absorption spectrum,

which would differ only in intensity from one crystal to another (Szigeti 1963); this is obviously not the case (§§1.3 and 3). In fact, for a given defect, the absorption coefficient is given by:

$$\alpha(\omega) = \frac{2\pi^2 D}{3nc} A^2(\omega) S(\omega)$$

where D is the concentration of impurities, n is the refractive index, $A(\omega)$ is a dipole moment due to the defect and $S(\omega)$ is the density of unperturbed lattice modes.

2.5. Symmetry lowering perturbations

The symmetry of a defect is an important clue to its identity. For example, in the diamond lattice any intrinsic defect having tetrahedral symmetry can only be a vacancy or a tetrahedral carbon interstitial. This has been important in confirming the vacancy as the defect responsible for the GR absorption system (see §4.2.1).

As in atomic spectra, one can in principle study the Stark (electric field) and Zeeman (magnetic field) splitting of zero-phonon lines (Runciman 1969, Kaplyanskii *et al* 1971). In practice, neither of these techniques has been very successful, largely because zero-phonon lines are relatively broad (typically > 1 meV), but valuable information can be obtained from magnetic circular dichroism (§4.2.1).

The N3 zero-phonon line is the only one known to display a resolvable Stark splitting (Kaplyanskii *et al* 1970). Other defects have been studied using a modulation technique to resolve splittings within the linewidth (Kaplyanskii *et al* 1971), but at least some of the results are questionable (see GR1 and H3 in §4).

Another symmetry determining technique, polarisation of luminescence, has been used on the H3, H4 and N3 centres. However, it cannot be applied to non-luminescent centres, and some symmetries cannot be distinguished.

The most powerful symmetry determining technique in practice in solids is uniaxial stress, in which the crystal is mechanically compressed along selected symmetry axes. Uniaxial stress has been used to study many centres in diamond. Unlike the Stark and Zeeman effects, resolvable splittings of zero-phonon lines can be obtained with easily achievable stress (Kaplyanskii 1964a, b, 1967, Runciman 1965, Hughes and Runciman 1967).

3. Infrared spectra (mainly vibrational)

We have already seen (§1.3) that the different types of diamond can be distinguished by their infrared spectra. In this context 'infrared' means wavelengths greater than $2 \mu\text{m}$, energies less than 0.6 eV or 5000 cm^{-1} . (Other spectra, at about $1 \mu\text{m}$ in the IR, are more conveniently dealt with in §4 because they are vibronic, like the visible spectra; most of the spectra considered in §3 are vibrational in origin.)

Thus, all diamonds show specimen-independent absorption in the $2\text{--}6 \mu\text{m}$ region (Sutherland *et al* 1954). This is intrinsic lattice absorption and is due to two- and three-phonon processes, since one-phonon transitions are forbidden by the symmetry of the perfect diamond lattice (Lax and Burstein 1955). Type IIa diamonds show only this infrared absorption. The other types contain impurities which destroy the local symmetry of the lattice and allow one-phonon processes to occur in the $7\text{--}13 \mu\text{m}$ region (Ia and Ib), or which cause electronic transitions around 0.3 eV (IIb). The

absorption due to both one- and two-phonon processes reflects the phonon density-of-states function, modified in the former case by a frequency-dependent factor characteristic of the defect (§2.4).

3.1. The lattice bands

The 2–6 μm bands ($1500\text{--}5000\text{ cm}^{-1}$) (figure 4) were recognised as due to the perfect diamond lattice by Sutherland *et al* (1954), Collins and Fan (1954) and Lax and Burstein (1955). They increase in intensity with temperature, as expected for lattice bands, whereas the defect-activated one-phonon bands in type I diamond do not change. The 3660 and 3120 cm^{-1} lines are due to three-phonon processes, the others being two-phonon.

In 1970, Solin and Ramdas, using laser-excited Raman spectroscopy plus neutron scattering data, assigned all the features of the infrared and Raman spectra to specific phonons.

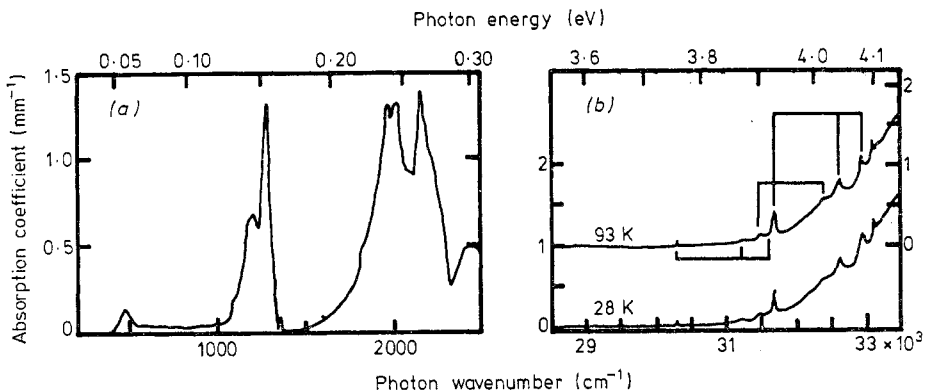


Figure 4. Absorption spectra of a type Ia diamond. (a) Infrared—below 0.2 eV are the impurity-induced group A absorption bands; above 0.2 eV the lattice bands shown by all diamonds. (b) The group A ultraviolet absorption lines and the ‘secondary absorption edge’. The true absorption edge is at 5.5 eV (after Davies 1976).

3.2. Defect-induced absorption

3.2.1. The A, B1 and B2 bands—type Ia diamond. These bands (figures 4 and 5) were discovered by Robertson *et al* (1934). In 1954 Sutherland *et al* noted that there were two uncorrelated series, which they labelled A and B, plus two independent lines at 1520 and 1540 cm^{-1} . The B bands were subdivided into the independent B1 and B2 groups by Sobolev *et al* (1968, 1969a, b, c) (table 2), a discovery foreshadowed by Clark (1965) and Charette (1961a), who observed that the 1370 cm^{-1} peak did not correlate with some of the other B bands. Diamonds with only A bands are common, and consequently the A spectrum is well known. Davies (1971, 1972a) has mathematically analysed superimposed A and B bands, and his band B spectra have been experimentally confirmed by Klyuev *et al* (1970) (figure 5). So we now know the spectra of each of these three series; we will discuss the A, B1 and B2 defects separately.

3.2.1.1. The A defect. Kaiser and Bond (1959), using mass spectrometry and chemical analysis, showed that the absorption coefficient of the most prominent A

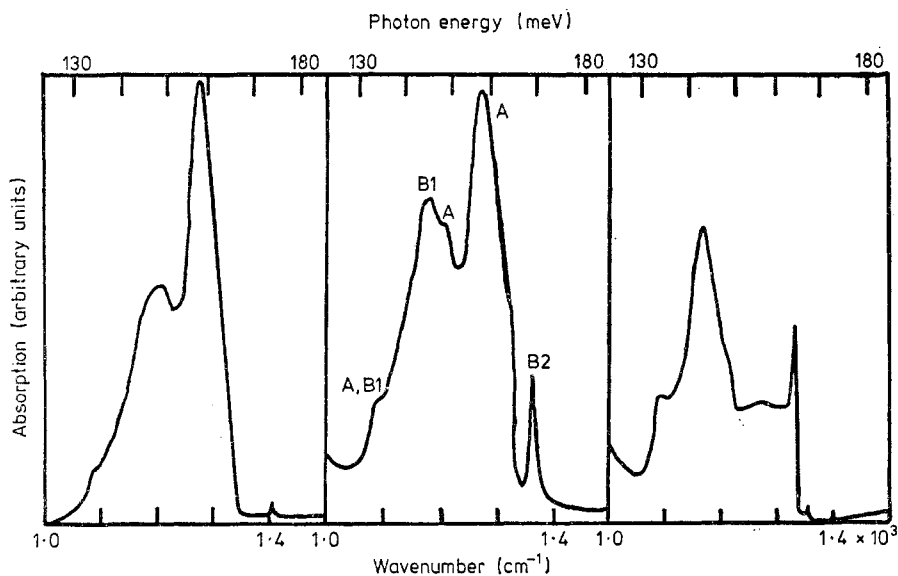


Figure 5. Impurity-induced infrared absorption spectra of three type Ia diamonds: left—the group A lines (type IaA); middle—mixed group A, B1 and B2 lines; right—primarily group B1 lines (type IaB) (after Davies 1971).

peak at $7.8 \mu\text{m}$ (1282 cm^{-1}) correlated with nitrogen content of the diamond; this result was confirmed by Lightowers and Dean (1964) using activation analysis. Both sets of workers reported a correlation between absorption coefficients at 1282 cm^{-1} and 306.5 nm (4.046 eV), with $\alpha(306.5) = 0.5 \alpha(1282)$. However, according to Summersgill (1972), the true correlation is between the absorption coefficient of the A feature at 1282 cm^{-1} (i.e. after subtraction of any absorption due to the B1 feature) and the secondary absorption edge coefficient at 306.5 nm , with $\alpha(306.5) = 0.65 \alpha_A(1282)$. Kaiser and Bond further showed that the 1282 cm^{-1} peak correlated with the increase in lattice constant of the diamond (see also Lisoivan and Sobolev 1974), but that the density did not vary. This implies that the nitrogen occupies a substitutional position—the greater mass of nitrogen relative to carbon is compensated by the lattice expansion.

It was assumed that this nitrogen gathered in large aggregates ('platelets') which were observable in the electron microscope and caused anomalous features ('spikes') in x-ray Laue photographs. However, in 1968, Sobolev *et al* showed that the 1282 cm^{-1} peak correlated neither with the x-ray spike intensity nor with the platelet concentration measured by electron microscopy, but there was a good correlation of spike intensity with the 1370 cm^{-1} peak, now labelled B2 (§3.2.1.3).

What then was the configuration of the nitrogen responsible for the 1282 cm^{-1} peak? Sobolev *et al* assumed it was '... an association of the type of N_2 '. The hypothesis received partial confirmation from Shcherbakova *et al* (1969), who observed an EPR signal from a pair of nitrogen atoms.

Further confirmation of the nitrogen pairs was provided by Davies (1976), who studied some ultraviolet absorption lines known to be related to the A centre. Although they are not in the infrared it is convenient to discuss them here.

Robertson *et al* (1934, 1936) noted that the diamonds with one-phonon infrared absorption also showed broadband uv absorption above 3.7 eV (figure 4). Sutherland

et al (1954) correlated this 'secondary absorption edge' with the A infrared bands. (The true absorption edge is at 5.5 eV.) Clark *et al* (1956a) discovered that superimposed on the secondary edge were some sharp lines which they labelled N5, N6, N7 and N8 (table 5) that correlated with N5 and with the slope of the secondary edge.

Table 5. The group A absorption lines in the ultraviolet. The defect is trigonal, with an A ground state. Note that the 159 meV (1282 cm^{-1}) phonon is the most prominent in the infrared spectrum also (see table 2 and figures 4 and 5) (after Clark *et al* 1956a, Davies 1976, Nazaré and Davies 1977).

Name	N5	—	—	—	N6	—	N7	—	N8
Energy (eV)	3.757	3.871	3.901	3.916	3.928	4.014	4.042	4.084	4.19
Assignment	Zero- phonon A—A	N5 + 113 meV	Zero- phonon A—E	N5 + 159 meV	Zero- phonon A—E	3.901 + 113 meV	N6 + 113 meV	N6 + 159 meV	?

Davies (1976) and Nazaré and Davies (1977) have discovered and correlated several additional lines not recorded by Clark *et al* (table 5). N5, N6 and the 3.901 eV peak are zero-phonon lines at a trigonal defect. The others are one-phonon sidebands, the phonon energies being 113 and 159 meV. The 'secondary absorption edge' is probably due to internal transitions at the A centre, rather than to transitions into the conduction band.

Davies shows that these results, together with previous data, severely limit the number of possible models for the A centre. It is a pair of substitutional nitrogen atoms in nearest-neighbour positions (figure 6(a))—a nitrogen 'molecule'.

The most prominent A infrared band at 159 meV (1282 cm^{-1}) is probably due to the LO phonon at the L point (the $\langle 111 \rangle$ zone boundary), shifted from its perfect lattice value of 1252 cm^{-1} by the mechanism explained in §2.4. Thus the molecular nature of the A centre now seems to be established.

The centre displays the phenomenon of sharp absorption lines superimposed on a continuum like the GR centre. It may therefore be that it displays similarly anomalous photoconductivity behaviour (§4.2.1). High-resolution photoconductivity studies of the uv lines are desirable.

What we do not know is why most natural diamonds are type Ia and contain nitrogen pairs, whereas synthetics are usually Ib, containing single nitrogen atoms; nor what role, if any, the A centre plays in forming the 'platelets'. It may be the initial synthesis conditions (Klyuev *et al* 1973), or the subsequent annealing and pressure treatments (Chrenko *et al* 1977). Further work is necessary.

3.2.1.2. The group B1 bands. The B1 spectrum is shown in figure 5 and the wavelengths and energies of the peaks are given in table 2. There are two questions to consider: the microscopic nature of the B1 centre, and possible correlations with other defects. We will consider the latter first.

Lightowers and Dean (1964) and Davies and Summersgill (1973) showed that there was no correlation between this centre and N3, but there apparently was a good correlation with N9, a result also stated or assumed (but not explicitly demonstrated) by many Russian workers (Sobolev and Lisoivan 1972, Sobolev *et al* 1969a, Klyuev *et al* 1972). However, Davies and Summersgill (1973), who did demonstrate the B1/N9 correlation, argued that it was fortuitous, because a typical diamond might

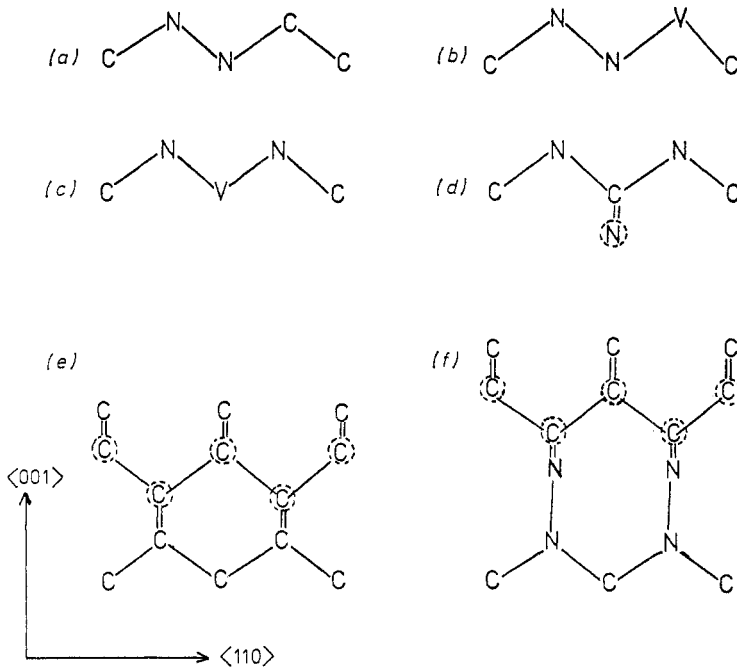


Figure 6. The diagram shows a {110} plane. (a) The nitrogen pair—thought to be the A aggregate. (b) A vacancy trapped by a nitrogen pair. The defect apparently rearranges to (c) an N-V-N configuration—thought to be the H3 defect (after Davies 1976). (d) The Mitchell (1964) model of the N3 defect—three nitrogen atoms. (e) The normal diamond lattice. (f) The Lang (1964) model for the platelets. The dotted circles in (d), (e) and (f) indicate atoms lying out of the plane of the paper.

contain an estimated 10^{22} N9 centres m^{-3} , and more than 10^{24} B1 centres. The N9 zero-phonon line, being electronic, is detectable at much lower concentrations than the vibrational infrared bands. This assumes that N9 is an allowed electric dipole transition, which seems likely. We must therefore conclude that the common origin of the N9 and B1 bands is not proven.

Davies (1972a) has deduced from the activation analysis results of Lightowers and Dean (1964) that the B1 centre contains nitrogen. Thus:

$$3.7 \times 10^{18} K_{1175} > C > 1.4 \times 10^{18} K_{1175}$$

where C is the nitrogen concentration in atoms cm^{-3} and K_{1175} is the absorption coefficient at 1175 cm^{-1} . Experimentally, Sobolev and Lisoivan (1972) find that:

$$C = 7.6 \times 10^{18} K_{1175} = 1.2 \times 10^{18} K_{236.5}$$

where $K_{236.5}$ is the absorption coefficient due to the N9 defect. The agreement is reasonable, since Davies' result is an estimate. However, the situation is unsatisfactory for several reasons. Firstly, Davies' estimate is based on only one diamond. Secondly, Davies has already rejected the correlation between the B1 and N9 defects which is implicit in the latter equation. Thirdly, Sobolev and Lisoivan give virtually no experimental details. Fourthly, the N9 zero-phonon line width is specimen-dependent, so the integrated absorption should be used, not the peak height.

Sobolev and Lisoivan (1972) propose their ' $N_S(111)$ ' model of the B1 centre—a dislocation loop stabilised by nitrogen atoms, the dislocation loops being due to the condensation of vacancies left behind after nitrogen atoms have precipitated into the platelets (Evans and Phaal 1962a). The model is consistent with results reported by Berman and Martinez (1976) and Evans *et al* (1976), but Klyuev *et al* (1977) were unable to confirm it, and Evans and Phaal (1962a, b) have observed dislocation loops in type IIa as well as Ia crystals, whereas B1 centres occur only in Ia. The $N_S(111)$ model has also been proposed to account for N9 and band A (§4.1).

What we do know about the B1 defects is that, like B2 centres (platelets) but unlike A centres, they do not affect the lattice constant and they must be very small (Sobolev and Lisoivan 1972). They must have a well-ordered molecule-like structure because they have a well-defined infrared spectrum and give rise to a well-defined vibronic system (H4) after irradiation and annealing (Davies 1972b, Davies and Summersgill 1973). If nitrogen is involved in the B1 centres, the atoms must be present in pairs because no EPR has been detected. They do not contain aluminium (Sobolev *et al* 1969a, b, c).

In conclusion, the B1 defect has not yet been identified. It is probably a small nitrogen aggregate, possibly associated with dislocation loops, and possibly correlating with the N9 centre. Further work is necessary.

3.2.1.3. The group B2 bands—the platelets. The platelets in type Ia diamonds are interesting because of the length of time they have remained a mystery and because of the wealth of experimental techniques brought to bear on them, though we still do not know their atomic arrangement and how they are formed.

In 1940 Raman and Nilakantan observed anomalous 'spikes' around some of the Laue spots in their x-ray diffraction pictures and it was soon realised that the defects responsible had to be ordered along the $\{100\}$ planes (Hoerni and Wooster 1955, Lonsdale and Smith 1942). Frank (1956) proposed segregated $\{100\}$ planes of impurity atoms, probably silicon, analogous to Guinier–Preston zones in age-hardened alloys, though subsequently Caticha-Ellis and Cochran (1958) found that the silicon concentration in their diamonds was too low to account for the x-ray spikes.

Kaiser and Bond (1959) demonstrated that in type I diamonds the intensity of the group A infrared peak correlated with the concentration of nitrogen impurity. This led Elliott (1960) to speculate that the platelets were composed of nitrogen. Unfortunately, the atomic models he proposed were based on Frank's (1956) calculations of atomic displacements, which were erroneous (Frank 1964). Another model based on nitrogen, and complying with the requirements of valency and of the unchanged density of platelet-containing diamond, was advanced by Lang (1964) (figure 6). Since the platelets had been observed directly by electron microscopy (Evans and Phaal 1962a), the problem appeared to have been solved. The platelets seemed to be nitrogen.

However, there were some discrepancies. Grenville-Wells (1952) and Sutherland *et al* (1954) showed that the ultraviolet and infrared absorption—known from Kaiser and Bond (1959) to be correlated with nitrogen content (§3.2.1.1)—did not correlate with the presence or absence of x-ray spikes. The breakthrough came when Sobolev *et al* (1968) demonstrated that the x-ray spike intensity correlated not with the 'nitrogen' infrared peak at 1282 cm^{-1} (A), nor with the 1175 cm^{-1} (B1) peak, but with the B2 line at 1370 cm^{-1} . 'The currently popular view that the impurity nitrogen is mostly located in the platelets . . . is erroneous'. Their results were confirmed by

Evans and Rainey (1975a, b), and Davies (1970a) put an upper limit of 10% (and no lower limit) on the nitrogen in platelets as opposed to smaller aggregations.

Confirmation of the Russian work came from a totally different technique—thermal conductivity. Turk and Klemens (1974), analysing the thermal conductivity variation with temperature of one diamond, deduced that 97% of the nitrogen it contained was in small 'point-like' aggregates, and only 3% in the planar platelets. Their results were confirmed by Berman *et al* (1975) and Berman and Martinez (1976).

We thus have a puzzle. Platelets are found only in type Ia (i.e. nitrogen-containing) diamonds; nitrogen is the only impurity known to be present in sufficient concentrations to account for the platelets; yet the platelets contain little if any nitrogen. Since the 1370 cm^{-1} platelet peak has never been observed without the A and/or B features it appears that nitrogen is a necessary but not sufficient condition for platelet formation. Evans (1973) therefore proposed that the platelets are primarily carbon; they are formed during growth by interstitial nitrogen atoms pairing with substitutional nitrogens to give a substitutional nitrogen pair and an interstitial carbon atom. The carbon interstitials then aggregate into platelets.

Some support for this hypothesis came from Berman *et al* (1975) and Berman and Martinez (1976). Their platelet concentration as measured by thermal conductivity was in rather better agreement with that deduced from the 1370 cm^{-1} peak if it was assumed that the platelets were stacking faults rather than nitrogen aggregates. Additional support came from the electron microscopy studies of Woods (1976), who studied 'giant' platelets up to 5 μm long (the usual size is 0.1 μm or less). He concluded that they were interstitial layers of carbon. However, his conclusions have been questioned by Lang (1977). Other workers (Bursill *et al* 1977), using phase contrast electron microscopy, have found that the Lang (1964) nitrogen platelet model (figure 6) fits their results best. Hence the question is still open. The answer may come from high-temperature high-pressure annealing studies (Nikitin *et al* 1969, Klyuev *et al* 1972, Evans *et al* 1976, Chrenko *et al* 1977).

The only optical property of the platelets that we have been concerned with so far is the absorption peak at 1370 cm^{-1} . However, it has recently become apparent that platelets give rise to visible and infrared luminescence. In 1971 Wight *et al* found a correlation between a broad featureless cathodoluminescent band peaking at 1.25 eV and the integrated intensity of the 1370 cm^{-1} peak. Hanley *et al* (1977) and Kiflawi and Lang (1977) have observed this luminescence from individual 'giant' platelets (figure 7 (plate)) and have shown that it is highly polarised (as one would expect from a planar aggregate). It is not clear whether this luminescence is intrinsic to the platelets or is due to decoration by other defects, though the former seems more likely since the luminescence is apparently not observed independently of the platelets.

Visible cathodoluminescence also occurs due to decoration of the platelets by H3 defects (Hanley *et al* 1977). Whether the H3 defects, which are thought to be a vacancy trapped by an A nitrogen pair, have migrated to the platelets or whether they are an intrinsic part of the structure, is not clear. It may be significant that both the Lang (1964) model of the platelets and the Davies (1976) model of the H3 centre involve a pair of nitrogen atoms (Walker 1977a).

3.2.2. The single nitrogen substitutional centre—type Ib diamond. This defect was first observed by Smith *et al* (1959) using paramagnetic resonance. Nitrogen could be unambiguously identified because ^{14}N , the most common isotope (99.6% abundant), is virtually the only nuclide with unit nuclear spin, and because nitrogen is a

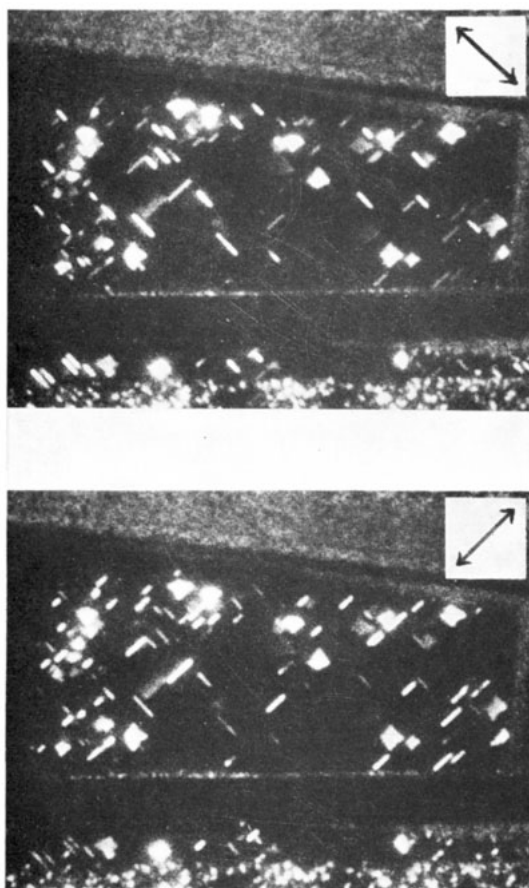


Figure 7. Polarised infrared (1.25 eV) cathodoluminescence from 'giant' {001} platelets. The arrows, length 100 μm , are parallel to E vector. The rectangles are platelets in the plane of the diagram; the lines are 'edge-on' platelets. Smaller platelets can be seen along the lower margins of the photographs (after Kiflawi and Lang 1977).

common impurity in diamond. The nitrogen atom substitutes for a carbon lattice atom, and the paramagnetic electron is located in an antibonding orbital between the nitrogen and one of the nearest-neighbour carbon atoms. The defect thus has $\langle 111 \rangle$ (trigonal) symmetry.

About 0.1% of all clear natural diamonds contain a homogeneous distribution of paramagnetic nitrogen and show characteristic optical absorption. This was discovered by Dyer *et al* (1965), who called them type Ib to distinguish them from the diamonds containing nitrogen aggregates (§§1.3 and 3.2.1). Type Ib crystals are yellow or green, depending on their nitrogen concentration. Dyer *et al* showed that they display one-phonon absorption in the infrared (figure 8) and a continuum starting at about 2 eV (600 nm) and increasing in intensity towards the ultraviolet. There are sharp lines on the continuum at 4.07, 4.15, 4.24, 4.58 and 4.64 eV, which are resolved only at 80 K, and a broad (0.5 eV) peak at 4.4 eV (280 nm) which could be their phonon band, as well as other broad peaks below 4 eV. The integrated intensity of the EPR line correlates with the integrated intensity of the infrared band and with the slope of the absorption at 2.6 eV.

Although natural type Ib diamonds are rare, synthetics usually fall into this category unless nitrogen is deliberately excluded from the reaction or other dopants added (§1.7). The spectra of Dyer *et al* have been confirmed by work on synthetic crystals (Charette 1961b, Sobolev *et al* 1969c, Chrenko *et al* 1971, Klyuev *et al* 1975a). There is only order-of-magnitude agreement on the relative strengths of the EPR and optical spectra. An absorption coefficient of 1 cm^{-1} at 1130 cm^{-1} is produced by $0.8 \times 10^{24} \text{ atoms m}^{-3}$ of nitrogen (Sobolev *et al* 1969c, Klyuev *et al* 1975a) or by $4 \times 10^{24} \text{ atoms m}^{-3}$ (Chrenko *et al* 1971). The latter used larger crystals, which implies greater reliability, but independent results, preferably using natural crystals, are desirable.

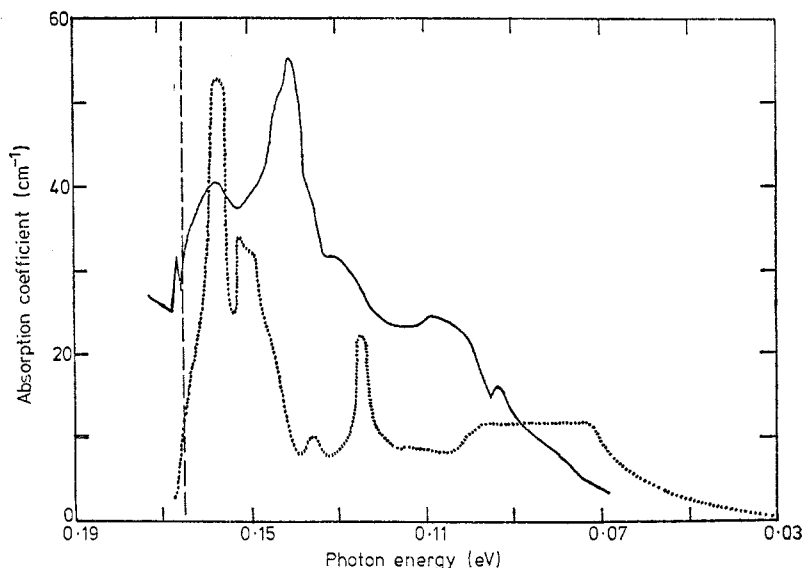


Figure 8. Full curve: infrared absorption spectrum of a type Ib diamond; dotted curve: calculated density of states (after Angress *et al* 1968). The peaks at 93 and 105 meV are as yet unconfirmed by other workers; Davies (1977a) has suggested that the 167 meV peak is an independent feature.

To investigate further the properties of nitrogen in diamond, some synthetics have been doped with ^{15}N ($I = \frac{1}{2}$). The earliest attempt (Cannon 1971) had limited success in that some changes were produced in the EPR spectrum. Later workers (Klyuev *et al* 1975b, Samoilovich *et al* 1975) measured the hyperfine parameters of ^{15}N , which were in excellent agreement with the calculated values. Furthermore, there was a change in the infrared spectrum. The 1135 cm^{-1} peak had shifted to 1120 cm^{-1} . No changes were observed in the other IR peaks. This suggests that the 1120 cm^{-1} peak is an 'in-band resonance', due to a vibration characteristic of the defect but falling within the range of diamond lattice vibrations because nitrogen has a higher mass than carbon (as opposed to a 'local mode' due to a lighter atom which would lie above the maximum lattice frequency—see Dawber and Elliott (1963)). The other peaks, since they do not change, are presumably due to perfect lattice phonons. Their energies will not coincide exactly with those of the perfect lattice (§2.4).

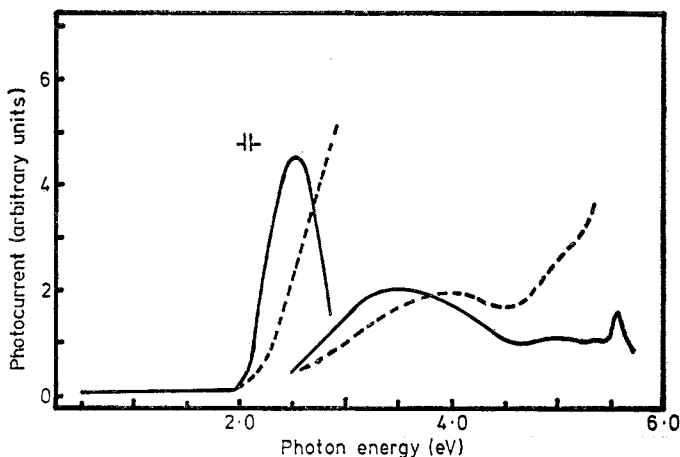


Figure 9. The room-temperature photoconduction spectrum of a synthetic type Ib diamond. Full curve: uncorrected; broken curve: corrected for intensity of exciting sources (after Farrer 1974). Two light sources were used.

Nitrogen, being a group V element, should be a donor in diamond, and experiments on type Ib crystals confirm this. Optical and thermal measurements indicate that the donor level is 2 eV below the conduction band (figure 9) and the thermoelectric effect indicates n-type conductivity (Farrer 1974, Vermeulen and Farrer 1975). (In fact, the thermal activation energy is 1.7 eV, whereas the optical threshold is at 2.2 eV. The difference is a consequence of the Jahn-Teller effect and is close to the value of 0.7 eV deduced for the Jahn-Teller energy from EPR measurements (Shul'man *et al* 1967, Loubser and Van Ryneveld 1967).) It seems that the surmise by Dyer *et al* that the UV absorption is due to transitions into the conduction band is correct. Earlier work had placed the donor level at 4.05 eV below the conduction band (Dean 1965, Denham *et al* 1967). This level needs further investigation, but it is not the single nitrogen donor.

Thus, the nitrogen substitutional centre in diamond appears to be well-characterised, both from the EPR and optical points of view. Further work is desirable, particularly on the UV peaks; they may be another example of anomalously sharp lines on a photoconduction continuum, like the GR2-8 lines.

3.2.3. *The boron acceptor centre—type IIB diamond.* There have been several recent reviews of the electrical and optical properties of type IIB diamonds which are p-type semiconductors (Vavilov 1975, Vavilov and Konorova 1976, Bourgoïn and Walker 1976, Davies 1977a, Collins and Lightowlers 1978). The acceptor impurity was originally thought to be aluminium, but it is now known to be boron (Chrenko 1973, Lightowlers and Collins 1976).

Electrical measurements (Wedepohl 1957a) show that type IIB diamonds are normal p-type semiconductors. The ground state of the acceptor centre is about 0.37 eV above the valence band, and the Hall mobility is 0.1–0.15 m² V⁻¹ s⁻¹ at room temperature. Unrealistic values of the hole effective mass are obtained unless it is assumed that some of the acceptor centres are compensated for by donors. Typical concentrations in natural diamonds are 10²²–10²³ acceptors m⁻³ and 10²¹–10²² donors m⁻³. At larger concentrations (10²⁴ m⁻³) and low temperatures (< 200 K), where holes cannot be excited into the valence band, conduction can occur by the hopping of carriers from occupied to unoccupied boron sites (Williams *et al* 1970, Bourgoïn and Walker 1976). In such a case the activation energy for conduction will be much lower—typically 0.2 eV.

Wedepohl (1957a, b) established that the acceptor centre displayed absorption peaks at 307, 346, 363, 514, 537 and 675 meV which correlated in intensity with each other, with a continuum absorption starting at 370 meV and extending to 2 eV, and with the number of acceptors uncompensated by donors ($N_A - N_D$). The three lower-energy peaks were recognised as bound state transitions, and the continuum as ionisation of the centre leading to photoconductivity (process 1 of figure 12).

The 307 meV peak is kept broad by a lifetime effect due to the proximity of a nearby level at 266 meV (Smith and Taylor 1962). Transitions to this level from the ground state are forbidden, but are observed in highly-doped synthetic crystals (Collins *et al* 1965) and in photoconductivity (Collins *et al* 1969). Below 300 K the 346 and 363 meV peaks show fine structure (figure 10), some of which disappears at 4 K; this was explained by Crowther *et al* (1967) using effective mass theory (figure 11). Their calculations were confirmed by their own uniaxial stress work and by the Zeeman and Stark effect measurements of Bagguley *et al* (1966) and Anastasakis (1969).

The broader absorption peaks at 514, 537 and 675 meV which are above the ionisation energy (370 meV), plus others at 462 and 625 meV, are phonon replicas of the lower-energy peaks, the phonon energy being approximately 160 meV (Smith and Taylor 1962). Hence photons with energies greater than 370 meV can be absorbed without creating free carriers (process 2 of figure 12). Since this is competitive with direct ionisation (process 1 of figure 12), the photoconductivity spectrum ought to show troughs at these absorption peaks, which is indeed the case (Hardy *et al* 1962, Hardy 1962a, Collins and Lightowlers 1968, Lightowlers *et al* 1968).

The photoconductivity spectrum also shows dips which do not correspond to absorption peaks. They are due to the capture of free holes into excited states of the acceptor centre with emission of phonons (process 3 of figure 12) (Hardy *et al* 1962). Later work (Collins and Lightowlers 1968, Lightowlers *et al* 1968, Collins *et al* 1969), taking measurements up to 2.6 eV, showed that up to 14 phonons could be emitted by a hole before it was captured by the 304 and 348 meV excited states. Furthermore, holes could be captured by excited states to which transitions from the ground state are forbidden. This confirmed the existence of the level at 266 meV, which was mentioned earlier, plus others at 240 and 289 meV.

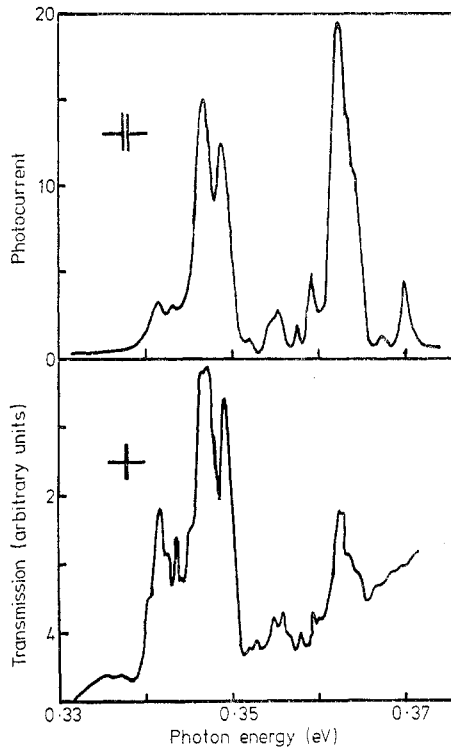


Figure 10. Comparison of photoconductivity at 120 K (upper curve) with absorption at 80 K (lower curve) of two natural IIb diamonds. Note that the photoconductivity peaks nearest to the ionisation energy (0.37 eV) are enhanced relative to the absorption peaks, due to photothermal ionisation (process 4 of figure 12) (after Collins and Lightowlers 1968).

There is yet another photon/phonon process observed in photoconductivity at the acceptor centre—photothermal ionisation—in which, unlike the other two processes, absorption *peaks* coincide with photoconductivity *peaks* (figure 10). It was observed and explained by Collins and Lightowlers (1968) and Lightowlers *et al* (1968) and is shown as process 4 of figure 12. Holes are raised to the bound excited states, and the lifetime is sufficient for them to then be thermally excited into the conduction band. In photoconductivity the peaks at 364 meV are enhanced relative to the 348 meV peaks because less thermal energy is necessary to ionise them. The activation energies deduced from the photothermal process (24.5 meV for the 348 meV peaks and 8.5 meV for the 364 meV peaks) tie in well with the thermal ionisation energy from the ground state of 373 meV (Lightowlers and Collins 1966). The photoconductivity peaks are usually better resolved than the absorption peaks because unperturbed excited states, having longer lifetimes, are most likely subsequently to undergo thermal ionisation.

At 5 K there is not enough energy for the photothermal ionisation mechanism to operate. Nonetheless, weak photoconductivity peaks are observed, and they are dramatically enhanced if the crystal is irradiated with 'bias light' of energy greater than 370 meV (i.e. within the ionisation continuum). The photocurrent is due to tunnelling of holes between excited levels of adjacent acceptors. The bias light, by ionising centres, provides vacant acceptor sites to which holes can tunnel from occupied sites (Collins and Lightowlers 1968, Lightowlers *et al* 1968).

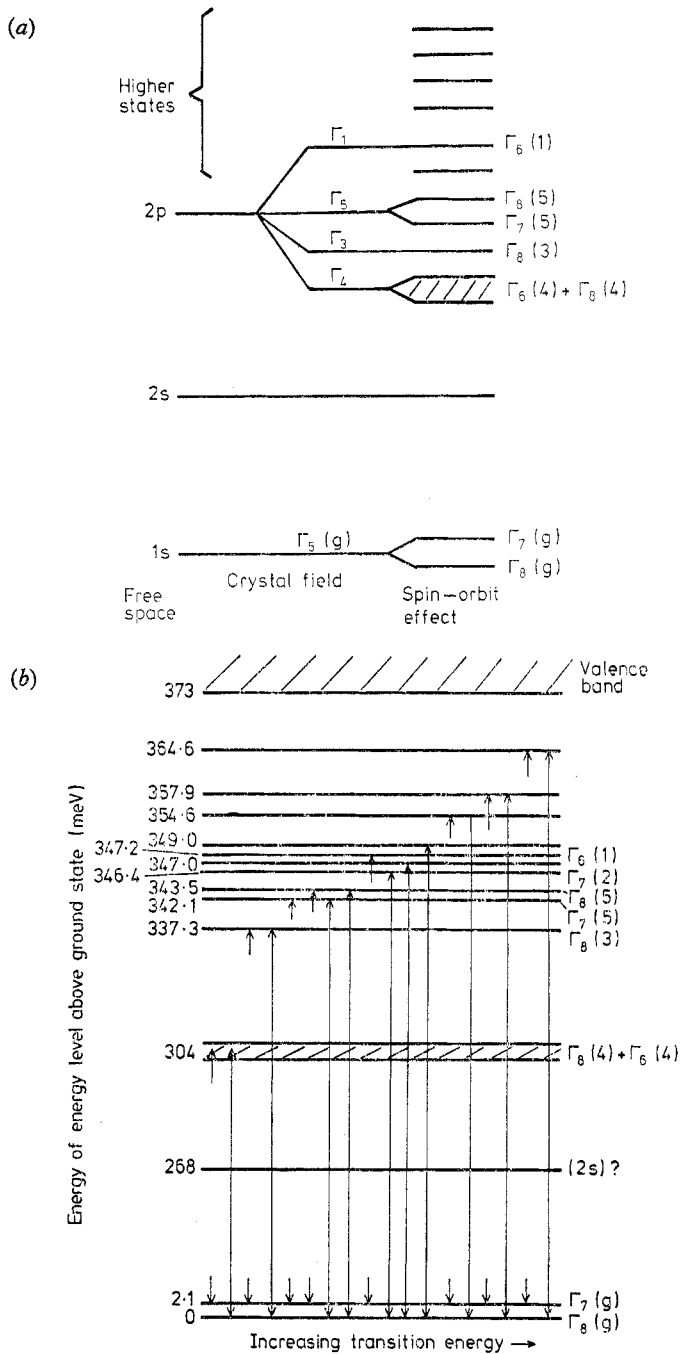


Figure 11. (a) A schematic diagram showing the effect of the diamond crystal field and the spin-orbit interaction on the boron atom. The bracketed characters indicate the parentage of the levels (see Crowther *et al* 1967, Anastassakis 1969). (b) The energy levels of the boron acceptor in diamond. All transitions are allowed excepted $\Gamma_6 \rightarrow \Gamma_6$ and $\Gamma_7 \rightarrow \Gamma_7$, and virtually all are observed (arrows). The unlabelled levels arise from more highly excited states than 2p. The diagram is based on table I of Crowther *et al* (1967). The levels at 347, 349 and 364.6 are thought to be four-, two- and four-fold degenerate, respectively (Anastassakis 1969, Bagguley *et al* 1966). Transitions to the level at 268 meV (and to others at 240 and 289 meV) are forbidden from the ground state (Collins and Lightowlers 1968, Collins *et al* 1969). The split ground state explains why some of the acceptor transitions disappear at low temperature (these transitions are shown as incomplete lines).

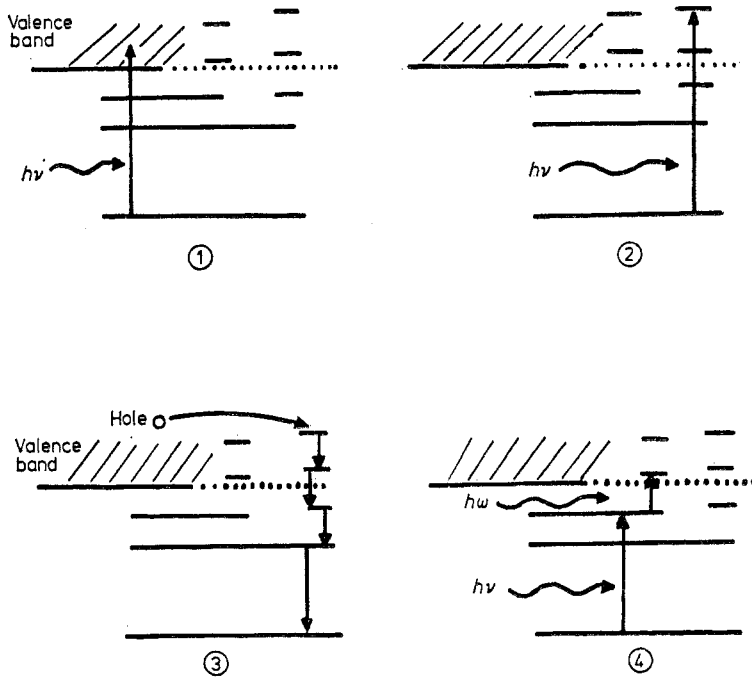


Figure 12. Photoprocesses at the acceptor centre: $h\nu$ and $h\omega$ indicate photon and phonon energies, respectively. (1) absorption leading to photoconductivity—photon absorbed, free hole created. (2) Absorption without photoconductivity—photon excites higher vibronic levels. (3) Hole capture into higher vibronic levels causing photoconductivity minima. (4) Photothermal ionisation—absorption of photon plus phonon leads to photoconductivity.

Another unusual feature of the acceptor centre is a weak absorption band at 160 meV first recorded by Charette (1961a) which Smith and Taylor (1962) studied in detail. It sharpens considerably on going from 293 K to 85 K, and displays structure reminiscent of the A and B infrared bands due to nitrogen. Smith and Taylor proposed that it had the same origin; a normally forbidden one-phonon process activated by the impurity. However, it differs significantly in two respects from the A and B bands. Firstly, the A and B bands change very little with temperature. Secondly, the acceptor centre is very efficient in inducing single-phonon absorption. Only 3×10^{22} acceptors m^{-3} are needed to give a 1 cm^{-1} absorption coefficient, compared with 7×10^{25} nitrogen atoms or radiation defects. This is a consequence of coupling between the electronic and nuclear motions (Hardy 1962b, Davies 1977a).

Free carrier absorption in semiconducting diamond is seen in the infrared at wavelengths above $10 \mu\text{m}$ ($< 0.1 \text{ eV}$), particularly if the diamonds are heated to populate the valence band with holes (Clark *et al* 1960).

The identity of the donor which compensates some of the acceptor centres is currently unknown. The question is discussed by Collins and Lightowers (1978), Davies (1977a) and Collins (1978).

Wedepohl (1957b) discovered that irradiation of semiconducting diamonds decreased their conductivity, hole concentration and hole mobility (see also Clark *et al* 1961). The infrared peaks were correspondingly reduced in intensity. His

conclusion was that the irradiation introduced extra donors (see also Dyer and Ferdinando (1966) and §4.2.1). Wedepohl discounted the possibility that the acceptors were displaced from their lattice sites, because both the irradiation dose and the acceptor concentration were small; this would rule out a direct collision process.

However, we now know that in silicon the aluminium acceptors can be ejected from the substitutional sites by migrating silicon interstitials (Watkins 1965). In diamond, most authors follow Wedepohl and assume, explicitly or otherwise, that the number of acceptors stays constant during irradiation. In the light of the evidence from silicon it would be advisable to re-examine this assumption. It could account for difficulties in interpretation experienced by Wedepohl and others (see also §4.2.7 for a defect which might involve a carbon or boron interstitial).

Thus we have seen that the acceptor centre in diamond displays many interesting physical effects. Some of these are qualitatively different from acceptors in silicon or germanium, because the depth of the acceptor level in diamond is significantly greater than the phonon energy. Scope for further work lies not so much with the acceptor centre itself (though calculations might be made on the higher excited states, extending the work of Crowther *et al*) as with irradiation of boron-doped synthetic crystals in order to identify impurity-related defects. Other areas of interest are the possible production of devices if n-type diamond can be synthesised, and the elucidation of whether elements other than boron can form acceptor centres.

3.2.4. Irradiation-induced absorption. Neutron or electron irradiation induces single-phonon infrared absorption similar in character to the A and B bands. Peaks are observed at 148, 139 and 125 meV, with weaker maxima at 165 and 159 meV, and a line above the Raman energy at 190 meV (compare table 2 and figures 4 and 5). The absorption coefficient per defect is much the same for radiation damage as it is for nitrogen. Annealing at 1000 K considerably reduces the absorption, as it does the visible spectra induced by irradiation (Smith and Hardy 1960, Smith 1961, Smith *et al* 1962, Evans and Wild 1967).

4. Visible and near-visible spectra

This section deals with electronic and vibronic spectra from 0.8 eV (1500 nm) to the fundamental absorption edge at 5.5 eV (220 nm). Energies and wavelengths quoted in this section will usually refer to zero-phonon lines. It is convenient to subdivide spectra into those present naturally, those created by irradiation, and those induced by subsequent annealing. Lack of space has precluded the discussion of several defects (see Davies 1977a).

4.1. Unirradiated diamond

4.1.1. The N3 centre (415 nm, 2.99 eV). The N3 absorption and photoluminescence is predominant in many natural diamonds, which has ensured that this centre has been much studied. It has previously been discussed by Clark (1965), Davies (1972a, 1977a) and Davies and Summersgill (1973).

The N3 spectra are shown in figure 13 and are an excellent example of mirror-image vibronic spectra. Dyer and Matthews (1957) found the centre in all the type Ia and IIa diamonds they studied, though not in type IIb. N3 correlates in intensity

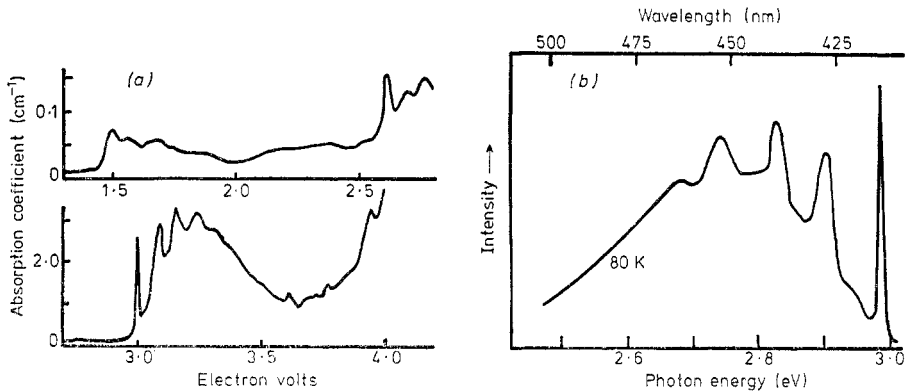


Figure 13. (a) N3 absorption spectrum showing also N1, N2 and N4 (after Clark *et al* 1956a). (b) N3 photoluminescence spectrum (after Dyer and Matthews 1957). Note the almost perfect mirror symmetry of the N3 absorption and luminescence spectra.

with N2 (477.6 nm, 2.596 eV) and N4 (344.2 nm, 3.603 eV), but not with N5 to N8. N1 (825 nm, 1.50 eV) is weak and has not been correlated with the other lines (Clark *et al* 1956a). Suggestions that N3 correlates with N9 (Raal 1959) and the group B infrared lines (Sutherland *et al* 1954) have been refuted by Lightowers and Dean (1964), Dean and Male (1964b), Sobolev *et al* (1970) and Davies and Summersgill (1973). A luminescence line at 2.001 eV may be a transition at the N3 centre (Dyer and Matthews 1957, Clark 1965).

The N3 absorption and luminescence bands have been studied in detail by Sobolev *et al* (1969b), Nedzvetskii and Dymke (1970) and Nedzvetskii and Gaisin (1974a), who have assigned all the phonon peaks using neutron scattering data and have shown that departures from mirror symmetry are small. Coupling is predominantly to symmetry-maintaining distortions along the trigonal axis; there is no evidence of Jahn-Teller effects (Davies 1974b).

Polarised luminescence and uniaxial stress agree in assigning N3 to a transition from an A ground state to an E excited state at a trigonal defect (Clark *et al* 1962, Clark 1965, Runciman 1965, Crowther and Dean 1967, Clark and Norris 1970). There is a small splitting of 0.59 meV in the excited state, presumably due to the spin-orbit effect (Davies and Summersgill 1973). (This is not to be confused with the variable splitting observed in luminescence by some authors and explained as self-absorption by Dyer and Matthews (1957).) The Stark effect rules out the trigonal point groups D_{3d} and D_3 , and hence N3 must have C_{3v} symmetry. It is the only defect in diamond to display a resolvable Stark effect in a static electric field (Kaplyanskii *et al* 1970, 1971).

The excitation spectrum of N3 contains a peak corresponding to N4, and depolarisation of the N3 luminescence is observed when N4 is excited. This is explained if N4 is an A to A transition ('a π $\langle 111 \rangle$ oscillator') at the same centre, and if the N4 excited state decays, radiatively or otherwise, by a transition to the N3 excited state (Clark and Norris 1970). Hence there should be no luminescence counterpart of the N4 absorption, as Dean (1965) found. This implies that N4 is a forbidden transition. Consistent with this, Welbourn and Davies (1977) have noted that both N2 and N4 are broad (typically 215 cm^{-1} compared to 30 cm^{-1} for N3). They suggest that these lines are vibronic in origin, the zero-phonon lines being forbidden.

Thus the N3 energy level diagram appears to be as shown in figure 14. It suggests

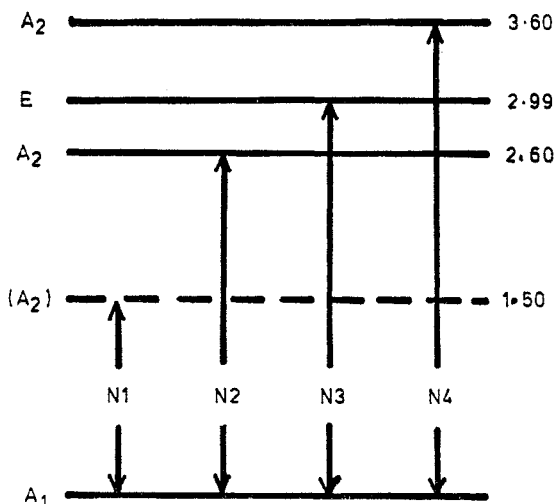


Figure 14. The N3 energy level diagram. The ground state is assumed to be A_1 (if it is A_2 then subscripts 1 and 2 are interchanged). The only forbidden transition in C_{3v} is A_1-A_2 . The N1 level has not been confirmed as belonging to the centre. It is assumed that N2 and N4 are forbidden transitions.

that transitions between the various excited states should be observable in luminescence, at energies between 0.4 and 2.1 eV. Many unidentified lines have been observed in this region, including the one at 2.001 eV already mentioned, though none have yet been correlated with N3. The diagram may also explain why N3 is not usually observed in cathodoluminescence; the most-favoured decay path for the excited state may not include $E \rightarrow A_1$. Dean's (1965) model for the N3 centre now seems unlikely (Collins 1974). An alternative model, proposed by Mitchell (1964), consists of three nitrogen atoms in next-nearest-neighbour positions, bonded to a common carbon atom (figure 6). This has the requisite C_{3v} symmetry and is consistent with N3 being found in nitrogen-rich diamonds. Other evidence relating this model to N3 comes from paramagnetic resonance (Welbourn and Davies 1977, Loubser and Wright 1973a, Loubser 1977), though further work is necessary to clarify certain discrepancies.

Until recently the N3 system was seen only in natural diamonds; now Chrenko *et al* (1977) have demonstrated that it can be induced in synthetic Ib diamonds by suitable heat and pressure treatment.

4.1.2. The N9 centre (236 nm, 5.26 eV). N9 was christened by Raal (1959), who also refers to earlier work. He studied the centre in absorption in very thin samples—it is often obscured by the 'secondary absorption edge' due to the A nitrogen aggregate. N9 luminescence can be found in most Ia and IIa diamonds, though not in types Ib and IIb (Dean and Male 1964a,b, Lightowers and Dean 1964, Wight and Dean 1967, Crowther and Dean 1967).

Raal proposed that N9 correlated with N3; this has been disproved by various authors (see §4.1). As we have already seen in §3.2.1.2, some authors claim a correlation between N9 and the B1 infrared bands, though the evidence is conflicting and the relationship doubtful.

In absorption the N9 lines are often broad; in luminescence and luminescence

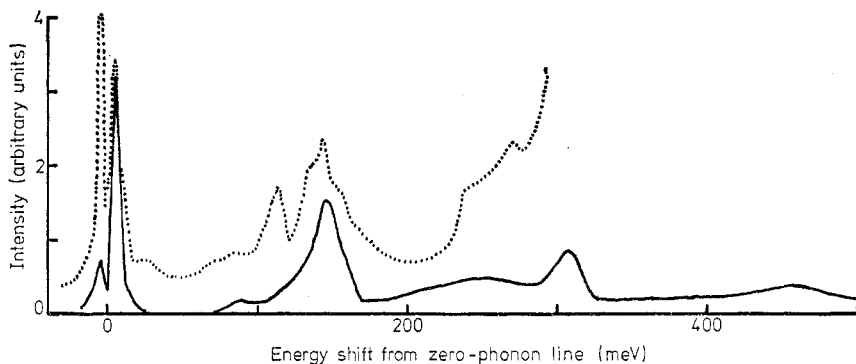


Figure 15. N9 luminescence at 100 K (full curve) and 80 K luminescence excitation (dots) spectra, plotted in terms of the energy shift from the zero-phonon doublet. The shift is negative for the lower spectrum (after Wight and Dean 1967, Dean and Male 1964a). The increase in luminescence excitation intensity at 300 meV corresponds to the fundamental absorption edge.

excitation they are sharper and more easily studied (figure 15). They broaden relatively little at higher temperatures; the zero-phonon line is a doublet (5.254 and 5.263 eV), and the splitting is still resolvable at room temperature (Nahum and Halperin 1962). In luminescence the higher energy component of the doublet is very weak, indicating that the splitting occurs in the excited state. The thermal activation energy is 10 meV, consistent with the optical splitting (Wight and Dean 1967). The phonon sidebands in luminescence show less structure than in excitation, partially because one zero-phonon line is almost frozen out (figure 15). Some of the extra excitation peaks may be zero-phonon lines (Wight and Dean 1967); on the other hand, there may be a genuine departure from mirror symmetry. This could indicate a Jahn–Teller effect, though according to Davies (1974b) coupling is primarily to symmetric modes.

Structure beyond the fundamental absorption edge at 5.5 eV is thought to be associated with N9. It is not detectable directly in absorption, but can be seen in luminescence excitation and in photoconductivity (Crowther and Dean 1967, Wight and Dean 1967, Denham *et al* 1967).

There is a weak band in excitation at 5.17 eV (240 nm), which Sobolev and Lisoivan (1972) claim correlates with the 5.26 eV peaks, though Crowther and Dean (1967), denying this, label it N10. Another weak line at 5.28 eV is visible in the excitation spectrum, though not in luminescence, and is probably due to a forbidden transition to a higher excited state. It is enhanced by uniaxial stress, consistent with this interpretation (Wight and Dean 1967, Crowther and Dean 1967). But Crowther and Dean were unable to determine the symmetry of N9 because of the closeness of the zero-phonon doublet and the inducing of the forbidden transitions. This is unfortunate, since the symmetry of the centre is an excellent guide to a model. The uniaxial stress results should be repeated in luminescence (*not* excitation) at liquid helium temperature; one member of the doublet would then be frozen out, as would transitions from the forbidden level, making the spectra easier to interpret; the 80 K spectra of Crowther and Dean would then be simpler to unravel.

There are two models for the N9 centre. The first, if one accepts the correlation with the B1 centre, is a dislocation loop on (111) planes which somehow involves nitrogen—N_S (111) (Sobolev and Dubov 1975). (It is discussed in §3.2.1.2). The

second is a nearest-neighbour donor-acceptor pair in its neutral charge state (Dean 1965) (§4.1.3).

Since N9 is found principally in type Ia diamonds it probably involves a nitrogen aggregate, but further work is clearly necessary.

4.1.3. Band A luminescence. A broadband luminescence between 1 and 3.7 eV can be excited in virtually all diamonds by energetic particles (α , electrons) or photons (UV or x-ray) (figure 17). It may display some structure due to superimposed luminescence or absorption of other systems such as N3, but is itself basically featureless. The bandwidth is typically 0.8 eV. The peak occurs at different energies for different types of diamond—2.2 eV (type Ib synthetics), 2.8 eV (natural IIb), 3.1 eV (Ia and IIa)—and shifts to higher energy at high excitation density. Some crystals may show two peaks. The luminescence is prominent below room temperature, but is totally quenched at 400 K, the activation energy for quenching being 0.36 eV. It is also quenched by radiation damage. Type IIa material may show weaker luminescence than type Ia (Dean and Male 1964b, Dean 1965, Crossfield 1974, Kiflawi and Lang 1974, Sobolev and Dubov 1975).

According to Dean (1965), band A luminescence is caused by recombination of electrons and holes at donor-acceptor pairs. This model predicts that the photon energy is given by:

$$E = E_g - (E_A + E_D) + \frac{e^2}{\epsilon r} - \frac{ea^5}{\epsilon r^6}$$

where E_g is the indirect energy gap (5.5 eV), E_A and E_D are the depths of the acceptor and donor states from the valence and conduction bands, ϵ is the dielectric constant and r is the separation of the donor and acceptor. The final (polarisation) term, in which a is an adjustable parameter, is negligible for large r . A consequence of this model is that the larger the separation r between donor and acceptor, the lower the photon energy and the longer the decay time. Experimentally this does seem to be the case with band A luminescence (Dean 1965, Crossfield 1974, Sobolev and Dubov 1975). It seems that the 2.8 eV peak is due to nearest-neighbour pairs, the 2.2 eV peak to all the other, more distant, pairs. The supposition is that in natural crystals the donors and acceptors have drifted together, whereas in synthetics a more nearly statistical distribution obtains. Using an acceptor energy of 0.37 eV (§3.2.3) and a donor energy of 4 eV (Denham *et al* 1967) in the equation gives a threshold for luminescence of 1.1 eV, as observed (Dean 1965, Crossfield 1974).

The model for the band A defect suggested by Sobolev and Dubov (1975) seems unable to account for the properties displayed. Dean's model accounts reasonably well for band A luminescence, though the acceptor is boron, but the 4 eV donor seems unlikely to be the single nitrogen atom, since that lies 2 eV deep (§3.2.2). Further work is clearly necessary on this question and on the relationship, if any, between band A and N9.

4.1.4. The S1 centre (503.4 and 510.7 nm, 2.463 and 2.429 eV). This centre occurs in the photoluminescence spectrum of some natural diamonds. At 85 K there are two zero-phonon lines at 2.463 and 2.429 eV; at 4 K the former freezes out and the latter is stronger (Sobolev *et al* 1969b, Nedzvetskii and Gaisin 1973a).

S1 is particularly interesting because the main zero-phonon line at 2.463 eV almost coincides in energy with the H3 and 3H zero-phonon lines (§§4.2 and 4.3).

But the three centres are quite distinct; their phonon replicas in luminescence are quite different, and only S1 shows the 2.429 eV line. Furthermore, both S1 zero-phonon lines show a linear Stark effect (Kaplyanskii *et al* 1971), whereas for H3 it is quadratic.

H3 and S1 can coexist in the same crystal (Nedzvetskii and Gaisin 1973a). These authors also claim that S1 contains three other zero-phonon lines, at 2.393, 2.363 and 2.344 eV, but this has not been confirmed by other workers. Lines at similar energies are produced by irradiation (Burton 1972, Davies 1977a), so these three lines may be independent features.

According to Klyuev *et al* (1973), S1 tends to occur in crystals containing the single nitrogen centre, i.e. type Ib crystals. An EPR centre of C_{2h} symmetry was discovered by Klingsporn *et al* (1970) in strongly luminescent type Ib diamonds. Shcherbakova *et al* (1972) found that all diamonds containing this EPR centre showed S1 luminescence, suggesting that they were the same defect. But the former shows a linear Stark effect, implying no inversion symmetry, whereas the EPR centre has a C_{2h} point group, which contains the inversion operation.

Clearly, further work is necessary on the S1 centre, e.g. symmetry determination by polarised luminescence and uniaxial stress, measurement of thermal activation energy and determination of its absorption spectrum. The two zero-phonon lines should be correlated explicitly with each other and with the EPR centre, and their bleaching behaviour examined. Although 3H, H3 and S1 are distinct centres, the similarity in energy of their zero-phonon lines may indicate a connection between them; this should be investigated.

4.1.5. Cathodoluminescence from dislocations. A blue luminescence, commencing at 2.4 eV and extending to at least 3.1 eV, has been observed from dislocations in natural and synthetic diamonds. It is more than 90% polarised, the E vector being parallel to the line of the dislocation, and is best observed in regions of the crystal where the blue band A luminescence is weak. Whether it is intrinsic to the dislocations or due to decoration by impurities is unclear. Dislocations can certainly be decorated by H3 defects, and hence emit the characteristic greenish luminescence. In synthetic crystals some dislocations emit in the yellow and red, though the luminescence bands have not been identified (Kiflawi and Lang 1974, 1976, Woods and Lang 1975, Hanley *et al* 1977).

4.1.6. The 1.40 eV centre (884 nm). This defect is seen in cathodoluminescence, but not in photoluminescence or absorption, in natural and synthetic type Ib crystals (figure 16). The zero-phonon line at 1.40 eV, which contains one-fifth of the intensity of the system, is a doublet of 2.9 meV separation. There is no change in relative intensity of the two components between 20 and 80 K and hence the splitting occurs in the ground state. In diamond the zero-phonon line energy usually decreases as T increases (Davies 1972b), but in this case it increases. The defect's identity is unknown, but it probably involves the nitrogen donor (Dean 1965, Wight *et al* 1971).

4.1.7. Bands B and C. Both these bands appear in cathodoluminescence (figure 17) but have not been recorded in absorption or photoluminescence. Band B occurs in natural type II crystals, though it may be obscured in type I crystals by competing luminescence (Wight *et al* 1971).

Band C (which does not correlate with band B) is seen in all except type Ib. It is unusual in that the apparent zero-phonon line couples to a single very sharp

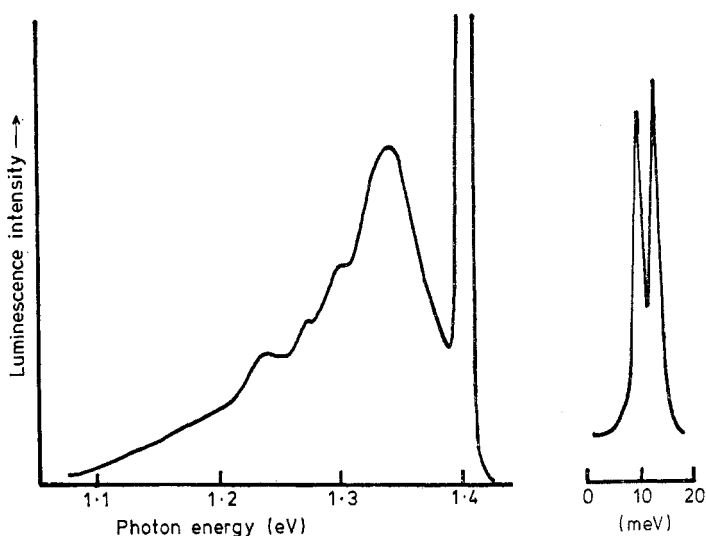


Figure 16. The 1.40 eV centre cathodoluminescence at 80 K. The 1.40 eV zero-phonon line is a doublet, as shown on the right (after Wight 1968).

53 meV phonon, at least eleven replicas being resolved (Wight *et al* 1971). This phonon must be a resonant mode characteristic of the band C defect, implying either a defect of large mass or locally softened force constants.

4.1.8. The amber centre (0.77 eV). This is common in, but not exclusive to, type Ib crystals. There is a zero-phonon line at 0.77 eV with phonon replicas at 0.83 and 0.89 eV, and continuous absorption up to at least 3.5 eV with structureless broad bands at 2.2 and 3.3 eV (figure 18). In the infrared there is also a peak at 0.52 eV and a characteristic single isotropic EPR line. The defect may involve nitrogen (du Preez 1965).

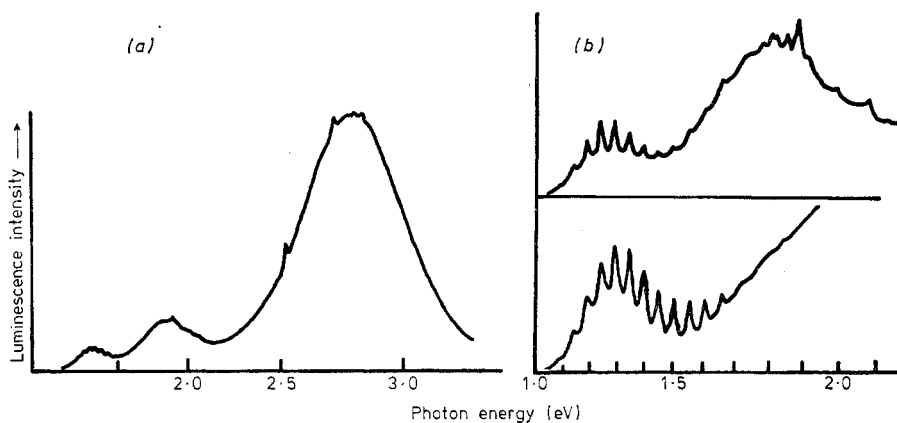


Figure 17. (a) The cathodoluminescence bands A, B and C. (b) Upper: same specimen as (a), but scale expanded. Lower: band C in a different crystal (after Wight 1968).

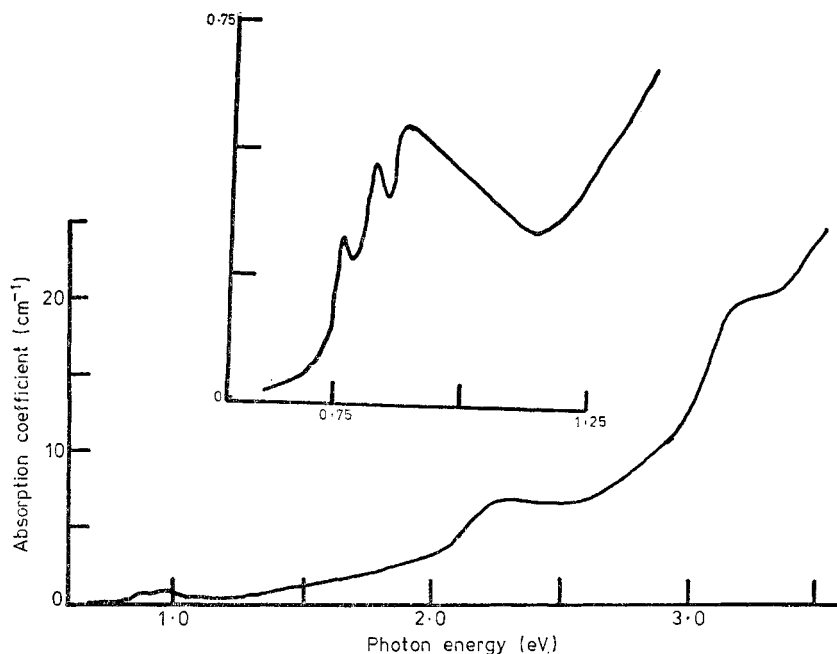


Figure 18. 80 K absorption spectra of the amber centre (after du Preez 1965). The scales are expanded in the inset.

4.2. Irradiated diamond

The wealth of spectra observed after room-temperature irradiation without annealing strongly suggests a mobile defect. The corollary is that any radiation defects produced without a high-temperature anneal must be either intrinsic or must involve a trapped interstitial, since the vacancy (the GR1 centre) is immobile below 900 K.

4.2.1. The GR1 centre (741 nm, 1.673 eV). There are a pair of zero-phonon lines at 1.665 and 1.673 eV (744.5 and 741.0 nm), a broad vibronic band peaking at 2 eV ('the GR1 band') and further zero-phonon lines between 2.8 and 3.1 eV (GR2, GR3, . . . , GR8) (figures 19 and 20) superimposed on a broad band ('the uv band') peaking at 4 eV. It is now generally accepted that the GR1 defect is the neutral vacancy. It is interesting and important for the anomalous effects it shows, and for its relevance to theoretical solid-state physics. Its properties have previously been reviewed by Clark and Mitchell (1971a, b, 1976), Clark *et al* (1979) and Davies (1977a).

Irradiation of any diamond produces a blue-green coloration due to the absorption bands in the red and violet (figure 19). The strength of the GR1 band is linearly related to the irradiation dose (Clark *et al* 1956a, 1961). This indicates an intrinsic lattice defect—no impurity is involved.

Irradiation of type IIb diamonds leads to the disappearance of the acceptor peaks in the infrared, and the early work indicated that the GR1 absorption band did not appear until all the acceptors had been compensated. Hence it is usually assumed that the GR1 centre is a donor and compensates the acceptors (Wedepohl 1957b, Clark *et al* 1961, Dyer and Ferdinando 1966; see also §3.2.3). (This incidentally

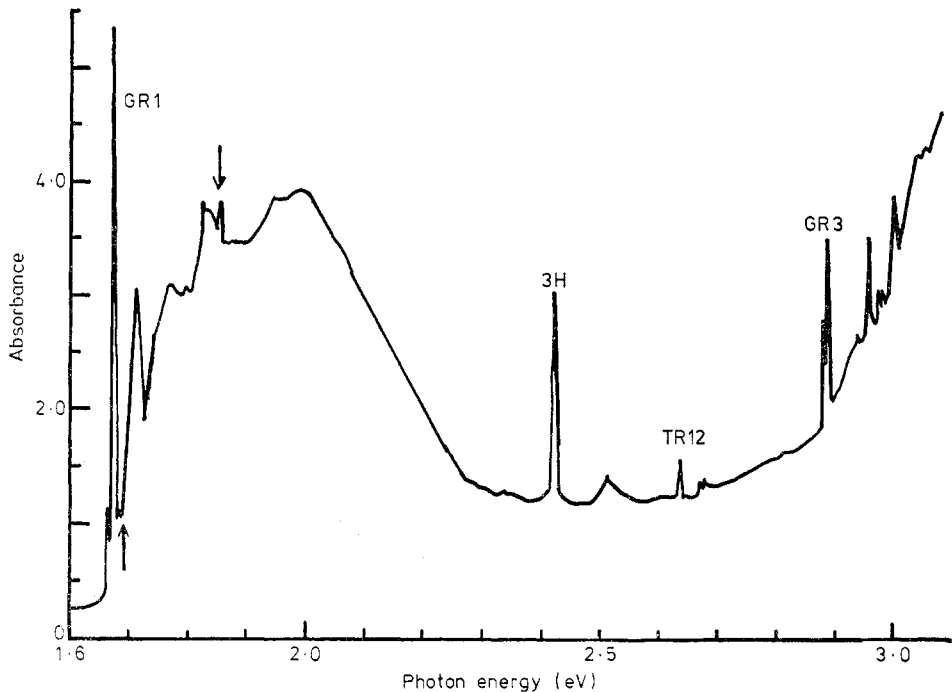


Figure 19. 80 K absorption spectrum of an electron-irradiated type IIb diamond. The two arrowed peaks belong to the tetragonal defect. The 3H and TR12 spectra are indicated; otherwise, all the structure is due to the GR defect. Before irradiation there was negligible absorption in this region.

implies that positively charged and hence paramagnetic vacancies are present in such crystals, though they have apparently not yet been observed.) However, Collins (1977) has detected the GR1 zero-phonon line in diamonds in which not all the acceptors have been compensated, and the GR1 centre annealed out below 1100 K, whereas the acceptor spectrum re-appeared only above 1200 K in the same crystal. Furthermore there is evidence that the GR1 centre ionises by emitting a hole (Vermeulen *et al* 1974), i.e. acceptor-like behaviour—this is discussed in more detail below. Hence the picture is somewhat ambiguous at present; it may be that the defect can be either a donor or an acceptor, depending on the Fermi level, like the vacancy in silicon (Watkins 1965).

Clark *et al* (1961) deduced for diamond a displacement energy (i.e. the energy needed to displace a lattice atom from its site) of 80 eV. They used both the production of the GR1 band at room temperature in type IIa diamond and the carrier removal rate at 120 K in natural semiconducting diamond (though the samples were warmed to room temperature after irradiation, and conductivity measurements were made between 200 and 300 K). Bourgoïn and Massarani (1976), using synthetic boron-doped crystals, irradiating at 15 K and measuring the conductivity at 12 K, deduced a displacement energy of 35 eV, which they attributed to the formation of vacancy-interstitial pairs. There is not necessarily a conflict if we assume that the result of Clark *et al* applies to isolated vacancies, particularly as we know that there are annealing stages below room temperature (Lomer and Wild 1971, Massarani and Bourgoïn 1976) and that temperature affects the damage rate (Collins 1977), but further work is desirable to explain the discrepancy.

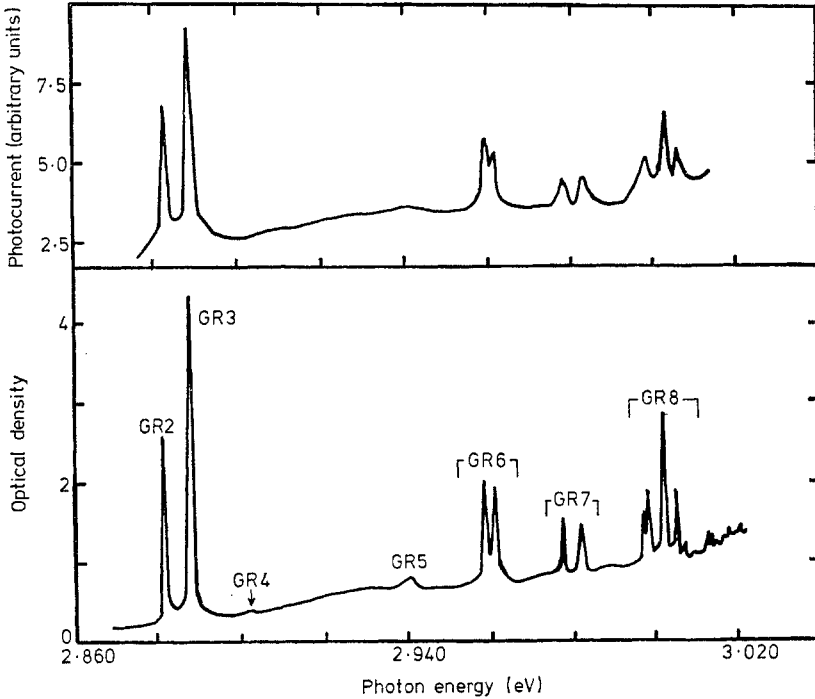


Figure 20. Photoconductivity (upper) and absorption spectrum (lower) at 40 K of the GR2-GR8 lines. Instrumental resolution is lower in the photoconductivity spectrum (after Vermeulen *et al* 1974). Compare with figure 19.

The GR1 absorption is strong; the oscillator strength is about 0.1. The luminescence is weak, but is an approximate mirror image of the absorption, and is unpolarised—an important result. The luminescence excitation spectrum shows peaks corresponding to GR3 and GR6, with a broad band at higher energies—the first evidence of a relation between these transitions and GR1 (Clark and Norris 1971).

It has been suggested (see Clark and Mitchell 1971) that the broad peak in the GR1 absorption spectrum at 1.71 eV is a zero-phonon line, but this now seems unlikely. A corresponding peak appears in luminescence at 1.63 eV indicating that it is a phonon replica of the main zero-phonon line at 1.673 eV. Such a low phonon energy may be due to local softening of the crystal round the defect (Ritter 1970, Larkins and Stoneham 1971). Accurate measurements (Wight *et al* 1971, Nedzvetskii and Gaisin 1973b) find that this first phonon energy is 41 meV in absorption but 36 meV in luminescence. Such a discrepancy also occurs with the H3 and H4 defects, consistent with them being due to the vacancy trapped at an impurity.

The 1.673 eV zero-phonon line has a weak satellite at 1.665 eV which is visible at 80 K but freezes out at 4 K (Lannoo and Stoneham 1968, Clark and Walker 1972, 1973). Both lines have been studied under uniaxial stress, and the energy level diagram derived from this and from thermal studies is shown in figure 21 (Clark and Walker 1973, Davies and Penchina 1974, Davies 1974a, 1975a). The defect has the full T_d symmetry of the diamond point group.

A tetrahedral defect would have unpolarised luminescence (table 4), as Clark and Norris (1971) found for the GR1 luminescence band as a whole, and B J Parsons

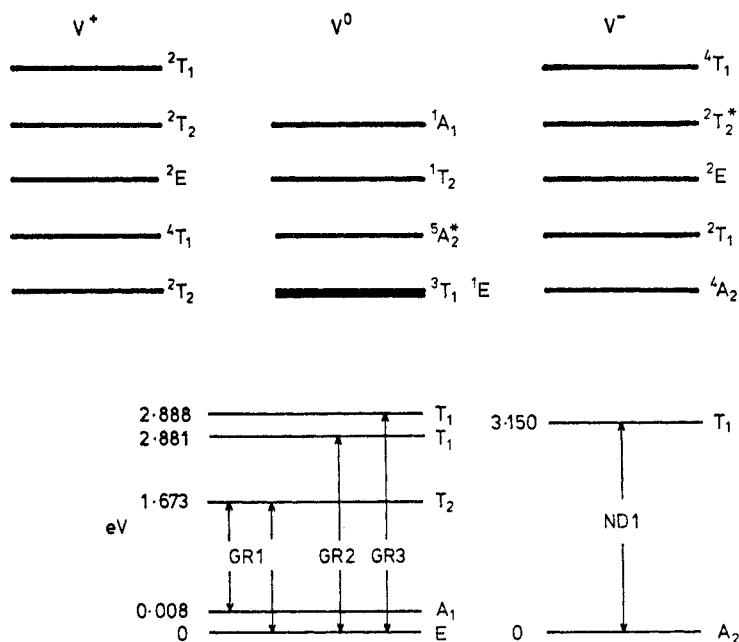


Figure 21. Upper: the energy levels of the positive, neutral and negative vacancies in diamond according to the 'defect molecule' calculations of Coulson and Larkins (1971). For some values of the parameters used in the calculations the asterisked levels may be the ground state. Lower: the experimentally determined energy levels for the GR and ND1 defects.

(1972 personal communication) confirmed for the zero-phonon lines at 1.665 and 1.673 eV. The 1.673 eV line shows a linear Stark shift (Kaplyanskii *et al* 1971), consistent with tetrahedral symmetry. The GR1 ground state is known to be diamagnetic (Clark *et al* 1964). Magnetic circular dichroism measurements (Douglas and Runciman 1977a) indicate a degenerate excited state and confirm the diamagnetic ground state, consistent with the T and E assignments already discussed.

Thus, all the experiments agree in assigning the GR1 energy levels as a 1E ground state, with a 1A_1 just above and a 1T_1 at 1.673 eV (figure 21). The experiments cannot distinguish between A_1 and A_2 , and T_1 and T_2 , but since these results are very similar to the ones predicted for V^0 by the molecular orbital calculations (Coulson and Larkins 1971, Stoneham 1975) A_1 and T_2 are usually chosen ($A_1 - T_1$ is forbidden). Hence these recent experiments confirm the long-standing belief that the GR1 defect is the neutral vacancy. Furthermore they indicate that the 'defect molecule' approach is a reasonable way of calculating the energy levels of point defects in semiconductors—a very important result.

Since the GR1 ground state is degenerate, a Jahn-Teller effect is expected; the fact that the Huang-Rhys factors derived from the zero-phonon line intensity, the Stokes shift and the bandwidth (§2.1) are not equal (3.7, 5.7 and 9.2, respectively) supports that expectation (Stoneham 1975). It appears that the E and A states (figure 21) are vibronic states derived from an E electronic state coupled to lattice vibrations of e symmetry. The T₂ state also couples to e modes. The Jahn-Teller energy is 0.63 eV and the phonon energy is 130 meV. Another level is predicted 20 meV above the ground state, but it has not been observed (Lowther 1975, 1976,

Stoneham 1977a). An earlier suggestion (Clark and Walker 1973), that the A state derived from the 3T_1 state (figure 21) as a result of spin-orbit splitting, now seems unlikely (Davies 1974a, Lowther 1975, 1976, Stoneham 1977a). The low g value in the T state (Douglas and Runciman 1977a) is a result of Jahn-Teller quenching (Lowther and Stoneham 1978).

The GR1 absorption and luminescence now appear to be understood. This is not yet the case for the GR2-GR8 transitions (figures 19, 20 and 21). These lines, although recorded by Clark *et al* (1956a) and other authors, received little attention until recently. We now know that they are all transitions at the GR1 defect (because they correlate in intensity with the GR1 zero-phonon line, show similar photoconductivity and thermal behaviour, and the most prominent of them appear in the GR1 excitation spectrum). There are more than seven lines; GR6 and GR7 are doublets and GR8 a quintet (figure 20). GR2 and GR3 have been studied under uniaxial stress and confirmed as E-T transitions at the GR1 defect; the upper state is T_1 if the GR1 upper state is T_2 (and vice versa), which explains why they, unlike the 1.673 eV line, do not have a satellite—it is forbidden (Clark and Norris 1971, Clark and Walker 1972, Walker *et al* 1974, Davies and Foy 1978). Douglas and Runciman (1977b) have suggested that GR5 and GR8 may not be transitions at the GR1 centre, but the differences they found between a Ia and a IIa crystal are probably due simply to large linewidths in the former.

Farrer and Vermeulen (1972) studied the photoconduction behaviour of all the GR lines and found photoconductivity troughs corresponding to the GR1 absorption peaks, as expected—photons absorbed by the localised levels of the GR1 centre are not available for photoconductivity (cf process 2 of figure 12). However, there were sharp photoconductivity peaks corresponding to the GR2-GR8 absorption peaks (figure 20)—a totally unexpected and currently not completely explained phenomenon. (The ND1 centre shows similar behaviour.) When photoconduction peaks are observed they are usually broad (1 eV or more), corresponding to quasi-localised levels within a conduction or valence band. Photothermal ionisation (process 4 of figure 12) cannot be the explanation because the GR2 excited state must be at least 120 meV from the conduction band (i.e. 120 meV below GR8), but photoconductivity is observed at 40 K (where $kT=3.4$ meV); and furthermore the relative intensities of the lines are the same in photoconductivity and absorption whatever the temperature, which would not be the case in photothermal ionisation (compare figures 10 and 20).

Another puzzling feature is that the GR centre has been regarded as a donor, as we have seen, since irradiation leads to compensation of the acceptor centres. Yet the evidence is that the GR2-8 photocurrent is due to holes (Vermeulen *et al* 1974, Vermeulen 1976). It seems unlikely that the GR2-8 excited levels can be within the conduction or valence bands, and autoionisation is probably involved. Three mechanisms have been proposed. Stoneham (1977b) has suggested that GR2 is a two-electron transition; it de-excites by autoionisation in which an electron is lifted out of the valence band. He further suggests that GR3-GR8 correspond to vibronic structure on the GR2 transition. This seems unlikely since they are so sharp, but detailed calculations using this model are desirable.

Dunn (1976), using the energy levels calculated for V^0 , V^+ and V^- by Coulson and Larkins (1971) (figure 21) but taking 3T_1 as the ground state of V^0 , has proposed that the GR lines are bound-exciton transitions from the ground state of the neutral vacancy to the ground state (GR1) or excited states (GR2-8) of a V^- which has trapped a hole. These discrete transitions overlap with continua due to emission

of holes from V^+ and V^0 , and hence autoionisation can occur. However, the GR1 ground state is not paramagnetic, as demanded by this model, and it is not clear whether the calculations will work with a 1E ground state for V^0 .

Lowther (1977) has proposed a similar model to Dunn, involving transitions between a neutral vacancy and a hole combined with a negative vacancy. He points out that GR2–GR8 are very sharp, which suggests that the hole is located in one of the basal orbitals of the nearest-neighbour carbon atoms surrounding the vacancy (i.e. the orbitals connecting these carbon atoms to the next-nearest neighbours). These orbitals lie very close to or just within the valence band, permitting photoconductivity while allowing the absorption lines to remain sharp. The hole de-excites into the 4A_2 ground state (figure 21) of V^- , and luminescence from GR2–8 should be significantly different to that from GR1. Although the model is not without its difficulties, it is the most detailed and the most promising one to date. However, the luminescence prediction is difficult to test. Although Clark and Norris (1971) have shown that GR1 luminescence can be excited by photon energies corresponding to GR2–8, the author was unable to detect any GR2–8 luminescence (nor any luminescence corresponding to transitions between the GR1 and GR2–8 excited states, despite the fact that T_1 – T_2 transitions are allowed—see figure 21).

Dyer and du Preez (1965) proposed that the uv band was due to transitions from the GR1 centre into the conduction band. However, this would lead to problems in explaining the sharpness of GR2–8, as we have seen. A more likely explanation is that part, if not all, of it is the vibronic sideband of the GR2–8 zero-phonon lines (Davies 1975c).

As Davies (1977a, b) has emphasised, there is much we still do not know about the GR centre. Further work is desirable, particularly on its possible connection with ND1 (see §4.2.2).

4.2.2. The ND1 centre (394 nm, 3.15 eV). The defect can be created in all types of diamond, though in IIa and IIb it is normally weak. In Ib it is created during an 80 K irradiation; in Ia, anneals of 620 K are necessary after irradiation and its strength qualitatively correlates with the nitrogen concentration. Coupling is predominantly to a 'breathing-mode' phonon of 80 meV (Clark *et al* 1956a, Dyer and du Preez 1965, 1967, Davies and Lightowlers 1970, Davies 1974b).

Dyer and du Preez (1965, 1967) and Davies (1977a) have investigated in detail the photochromic and thermochromic behaviour of ND1. Heating at 770 K enhances the intensity of ND1 and of a single isotropic EPR line which is induced by irradiation, whilst GR1 and the uv band are diminished. Ultraviolet light of energy less than 3.15 eV has the same effect, but if the energy is greater than 3.15 eV (i.e. within the ND1 absorption system) it is ND1 that is bleached and GR1 enhanced (figure 22). Annealing above 800 K destroys ND1.

The ND1 defect has tetrahedral symmetry, the ground state being A and the excited state T (Runciman 1965, Davies and Lightowlers 1970, Kaplyanskii *et al* 1971, Douglas and Runciman 1977b). The centre is not luminescent (Davies and Lightowlers 1970, Davies 1972b), contrary to earlier reports. This is presumably because, like GR2–8, there are photoconductivity *peaks* corresponding to the absorption peaks; these are present even at 8 K, which rules out photothermal ionisation, and they are still unexplained (Farrer and Vermeulen 1972).

A tetrahedral defect must be either a single vacancy, a single interstitial or a substitutional impurity. Davies and Lightowlers (1970) proposed a nitrogen interstitial,

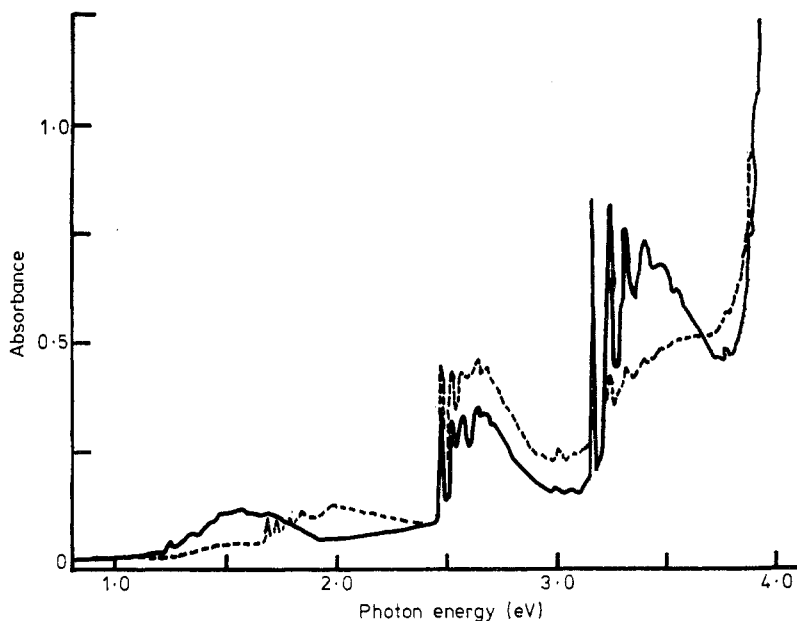


Figure 22. 80 K absorption spectrum of an irradiated type Ia diamond. Full curve: after heating in the dark; broken curve: after subsequent exposure to unfiltered mercury light. The GR1 (1.673 eV) and ND1 (3.15 eV) bands vary in a complementary fashion, as do H2 (1.25 eV) and H3 (2.46 eV) (after du Preez 1965).

since the centre is created in strength only in nitrogen-containing diamonds. But Sobolev and Eliseev (1976) and Davies (1977b) have recently suggested that ND1 is a negative vacancy. Davies' arguments are as follows. Firstly, GR1 is very weak in type I compared to type II diamonds, the ratio GR1/ND1 being 18, 1.3 and $\ll 0.1$ in a IIb, Ia and Ib irradiated under similar conditions. Since GR1 is a very stable defect this suggests that vacancies are being created in a different charge state. Secondly, the molecular orbital calculations (see figure 21) find that V^- has a $4A_2$ ground state with a $4T_1$ state several eV above, similar to ND1. Thirdly, the photochromic effects are more easily explained as a single defect changing its charge state, and the carrier emitted when the GR defect ionises does appear to be a hole (Vermeulen *et al* 1974), consistent with a change from V^0 to V^- . The carrier released when ND1 ionises has not been determined explicitly, but Vermeulen's (1976) work is consistent with electrons. Fourthly, the H3 centre is thought to contain a vacancy, and its growth on annealing may match the decrease in either GR1 or ND1. Fifthly, a $4A_2$ ground state, with spin 3/2, should be EPR-active; Dyer and du Preez (1965) have indeed found an EPR signal correlating with the ND1 absorption, though it has not been analysed in detail. Finally, Lowther and Stoneham (1978) have shown that the ND1 uniaxial stress parameters are consistent with a negative vacancy. Hence, Davies' hypothesis seems to fit the facts, and if correct it is a very important corroboration of the Coulson molecular orbital calculations (Coulson and Larkins 1971). Further tests of the hypothesis, by determining the sign of the ND1 charge carriers, and particularly by EPR studies, are desirable.

4.2.3. The 3H centre (503.5 nm, 2.462 eV). This centre is produced in all types of diamond by irradiation, even at 80 K, though more readily at room temperature.

Its intensity is enhanced by short anneals at 600 K, but longer times or higher temperatures cause it to disappear, as does UV irradiation. This is a charge transfer effect rather than a genuine anneal, because the centre can be restored by x-irradiation, and is stable to at least 900 K. In type Ia its strength is proportional to the nitrogen content, and it is particularly strong in type Ib crystals. Whether this is a Fermi level effect or implies that nitrogen is involved in the centre is not clear. Phonon coupling is weak ($S \sim 0.7$), the major phonon replica being at 67 meV (figure 23). Uniaxial stress indicates a $\pi \langle 110 \rangle$ oscillator at a rhombic I centre (table 4) (Palmer 1961, Davies 1974a, Walker 1974, 1976a, 1977b, Nedzvetskii and Gaisin 1975).

The centre was labelled 3H because its zero-phonon line is at virtually the same energy as H3. (In earlier work, 3H was probably assumed to be H3.) Both centres have the same rhombic I symmetry (Walker 1974, Davies *et al* 1976). However,

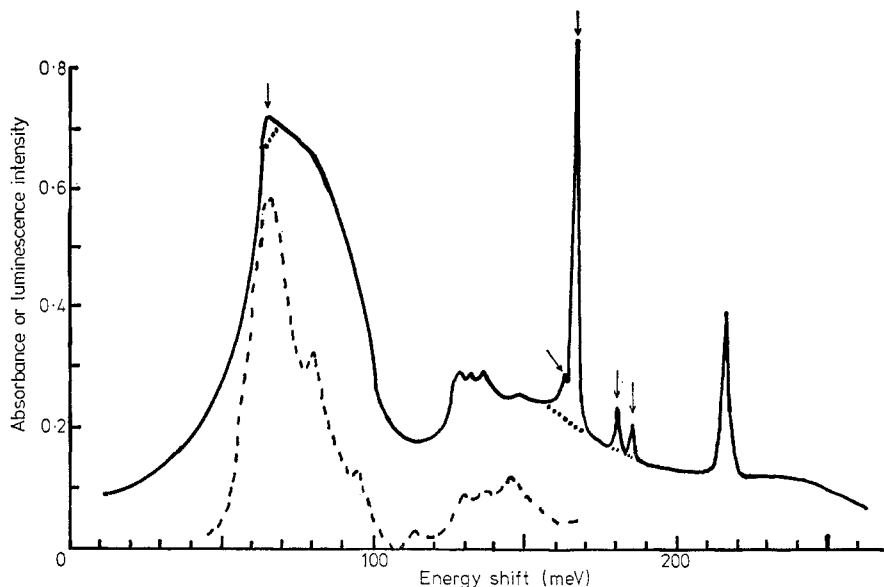


Figure 23. A comparison of the 3H absorption (broken curve) and photoluminescence (full curve) at 80 K. The abscissa shows shift in energy from the zero-phonon line at 2.462 eV and is negative for the luminescence spectrum. The arrowed peaks may be due to the laser used to excite the luminescence (after Walker 1977b).

there is no evidence of any relationship between them; H3 can be produced only in type Ia diamonds, whereas 3H can occur in considerable strength in all types, and their luminescence spectra are quite different (figures 23 and 28) (Walker 1976a, 1977b). For similar reasons 3H is presumably not the same defect as S1, which also has a 503.5 nm zero-phonon line (§4.1.4).

Hanley *et al* (1977) have observed 3H in cathodoluminescence in the surface layers of natural diamonds, presumably a result of natural irradiation. They report that on one occasion, using an intense electron beam, the 3H luminescence faded, to be replaced by the 575.5 nm system (§4.3). Since this latter can also be produced in any diamond by irradiation, a connection between them seems possible.

There is only one, speculative, model for 3H, an oxygen-vacancy pair (Walker 1976a). Because 3H can be bleached, it must be paramagnetic in at least one charge state, and this should be investigated. Further studies of its luminescence spectrum

and Stark shift are desirable, to confirm beyond doubt that it is not the same centre as S1.

4.2.4. The S2 centre. This centre was observed in the 80 K photoluminescence spectra of a few natural crystals. It has five zero-phonon lines A–E at 2.371, 2.537, 2.597, 2.626 and 2.637 eV, the three strongest of which (A, B and C) display phonon replicas. Going from 80 K to 140 K causes B to weaken dramatically, while C and D increase in intensity and A and E stay approximately constant (Il'in *et al* 1971). These changes are presumably related to population and depopulation of electronic levels.

The S2 system has also been recorded in absorption (J Walker, unpublished work) in Ia diamonds, though not in IIa or IIb, after room-temperature electron irradiation. The strength was specimen-dependent. Only A and B were strong enough to measure, though weak peaks were present at energies corresponding to C, D and E. There were phonon replicas of A at 32 and 72 meV (in good agreement with Il'in *et al*), each of intensity one-quarter that of A, giving a Huang–Rhys factor of 0.4. A was clearly observable at 80 K but decreased in intensity at lower temperature, indicating that it is a transition from an excited state about 10 meV above the ground state. B was 2.5 times as strong as A, but less sharp (16 meV compared to 4 meV). Its intensity did not vary with temperature, and at 17 K it looked like an unresolved triplet or quartet. Further irradiation caused S2 to decrease in intensity, as did annealing at 600 K; whether it could be restored by illumination was not investigated.

Other authors (Clark *et al* 1956a, Dean and Male 1964b, Wight 1968, Burton 1972, Davies 1974a, Davies and Hamer 1976) have recorded zero-phonon lines very close in energy to all of the S2 transitions, particularly in irradiated type Ib crystals. Further work is clearly necessary, firstly to confirm that the five lines A–E genuinely belong to the same centre, secondly to ascertain their relationship to the other work and, thirdly, to investigate the suggestion of Klyuev *et al* (1972, 1973) that S2 is related to the N9 and B1 centres.

4.2.5. The TR defect. This defect was first studied by Clark *et al* (1956a, b), who observed it in absorption and luminescence in irradiated type IIa crystals (figure 19). TR12, TR12a and TR17 (2.638, 2.645 and 2.828 eV) are zero-phonon lines; the others, including TR13–TR16 (figure 24), though relatively well resolved, are probably phonon replicas of TR12. The Huang–Rhys factor is 0.8. There are also several infrared peaks between 1.52 and 1.54 eV which appear to correlate qualitatively with TR12. Short anneals at 800 K enhance the system; longer times or higher temperatures destroy it. The luminescence spectrum is not a mirror image of the absorption, which suggests a Jahn–Teller effect. Uniaxial stress identifies TR12 as a σ oscillator at a monoclinic I defect (table 4) (Walker 1974, 1977b).

The TR absorption is strengthened by short 800 K anneals but longer times or higher temperatures destroy it. It occurs in type IIa and IIb crystals, but not in type Ia (Walker 1977b). Nor was it detected in absorption or cathodoluminescence in irradiated synthetic type Ib diamonds (Burton 1972), though it was plainly visible in both absorption (Walker 1977b) and cathodoluminescence (Collins 1977) after lower doses in type IIb. G Davies (1976, personal communication) has observed a weak TR12 line in a type Ia crystal, but this may have been due to specimen inhomogeneity.

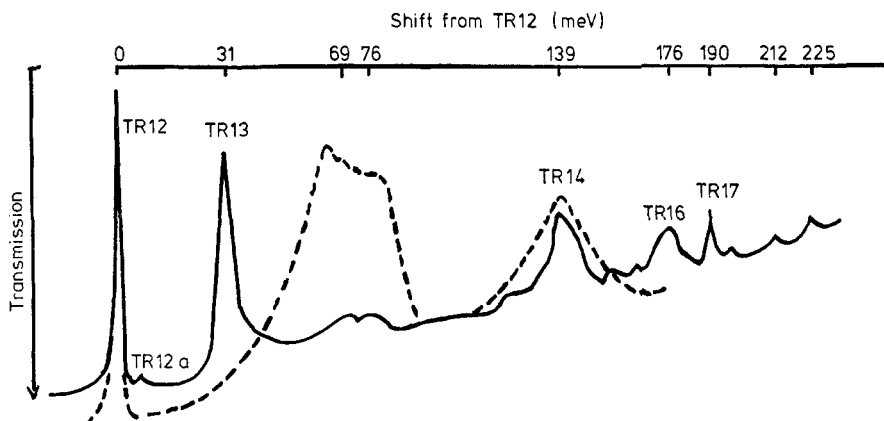


Figure 24. A comparison of the TR absorption (full curve) and photoluminescence (broken curve). The abscissa shows shift in energy of the various peaks from the TR12 zero-phonon line at 2.638 eV and is negative for the luminescence spectrum. Note the absence of 'mirror symmetry' (after Walker 1977b).

The defect is thought to be an interstitial complex, possibly involving an impurity (Walker 1977b). The failure to produce it in type I crystals was ascribed to the presence of alternative traps for the carbon interstitials, but in view of Davies' (1977b) results for GR1 and ND1 it may be that the TR defect is simply in a different charge state in such diamonds.

4.2.6. The 5RL defect (270.6 nm, 4.581 eV). This unusual system (figure 25) was discovered in cathodoluminescence by Wight (1968). The main zero-phonon line C at 4.581 eV, together with its phonon structure down to 4.4 eV and two other zero-phonon lines H and I at 4.407 eV and 4.389 eV, are reproduced very sharply at intervals of 236 meV. The defect has been observed after electron or neutron irradiation in 'intermediates' and types IIa and IIb (natural and synthetic), but not in types Ia or Ib, possibly because of competing absorption. Wight suggests that the

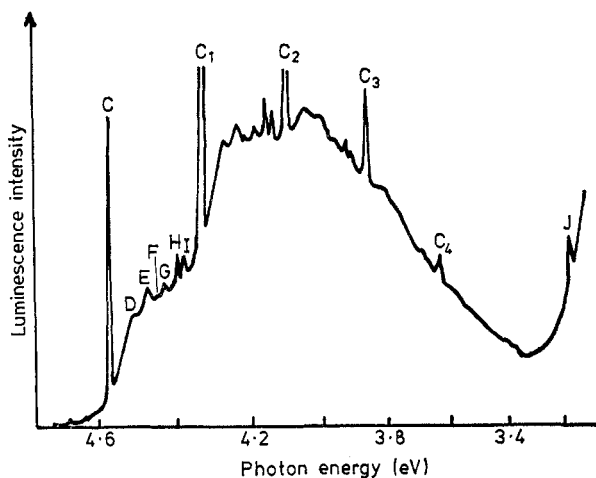


Figure 25. 100 K cathodoluminescence spectrum of the 5RL system in a type IIb diamond (after Wight 1968).

defect is intrinsic, though perhaps an aggregate. Collins (1977, 1978) has observed that the centre anneals below 1200 K, and has recorded the zero-phonon line in absorption.

The phonon energy of 236 meV is well outside the Raman energy of 165 meV. It must be a local mode phonon, though there is no record of a peak at 236 meV in the infrared after irradiation (Smith 1961). Further work on the 5RL centre is highly desirable.

4.2.7. The 2BD defect. This defect was discovered by Wight (1968) in cathodoluminescence (figure 26) in type IIb diamonds containing the 5RL system, and hence is thought to be due to irradiation ('2BD' stands for 'type IIb damage'). There appear to be two zero-phonon lines, F and G. F consists of four components, the strongest being at 4.777 eV, with weaker peaks at 4.781, 4.803 and 4.830 eV; they couple to a sharp 210 meV phonon. G(4.698 eV) is a similar multiplet, the phonon

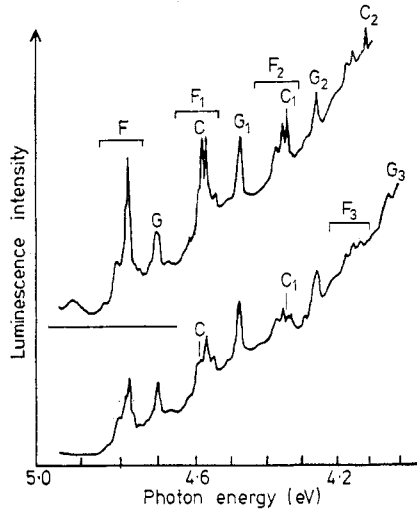


Figure 26. The 2BD cathodoluminescence spectrum in two diamonds, at 50 K (upper) and 100 K (lower). Peaks C, C₁ and C₂ are part of the 5RL system (see figure 25) (after Wight 1968).

energy in this case being 220 meV. (Both these are above the Raman energy of 165 meV.) Wight tentatively suggested that the defect was an acceptor–vacancy pair. No other author has studied this system.

The acceptor impurity, boron, has all its atomic spectral transitions in the ultra-violet. Hence F and G may derive from these transitions, in the free atom, particularly from the intense doublet at 4.96 eV. The local mode phonons suggest a defect with a mass lower than carbon, or increased force constants. A boron–vacancy pair seems unlikely, since vacancy-related defects usually have low phonon energies (e.g. GR1, H3, the 1.944 eV centre). The centre could be a boron interstitial or a boron substitutional atom which has trapped a carbon interstitial.

Further work is clearly necessary on this interesting and neglected centre. In particular, it is not certain that F and G are due to the same defect. It may also have a bearing on whether the GR defect is a donor, and why the acceptor infrared absorption disappears in irradiated semiconducting diamonds (§§3.2.3 and 4.2.1).

4.2.8. *The tetragonal defect (736 and 667 nm, 1.685 and 1.859 eV).* These two zero-phonon lines are observed in absorption in all types of diamond, but not apparently in cathodoluminescence (figure 19). They were incorrectly assumed to be part of the GR1 system by some authors, but they anneal out at about 700 K whereas GR1 does not and they do not appear in the GR1 excitation spectrum. The 1.685 eV line freezes out below 80 K, indicating that it is a transition from an energy level about 6 meV above the ground state (Wight *et al* 1971, Clark and Walker 1972, 1973, Davies 1974a, 1977b, Walker 1977c).

A radiation-damage defect seen in EPR (the R2 system (Faulkner and Lomer 1962, Lomer and Wild 1973)) also anneals at about 700 K (Clark *et al* 1964) and Walker (1977c) has shown that it correlates in intensity with both the 1.685 and 1.859 eV lines. Furthermore R2 has tetragonal symmetry, and uniaxial stress has indicated this symmetry for the optical lines (figure 27). Hence the optical and EPR centres must be

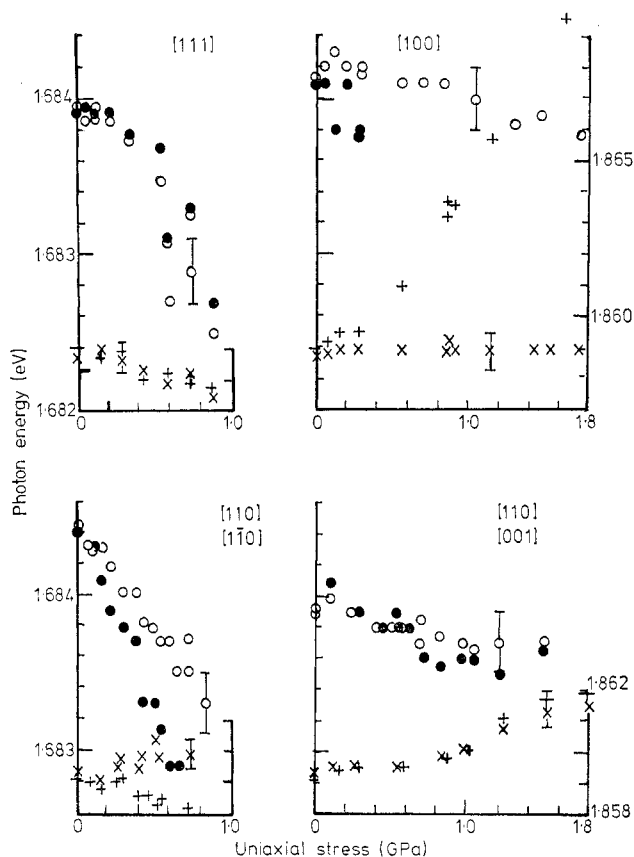


Figure 27. The 1.685 (circles) and 1.859 eV (crosses) absorption lines under $\langle 100 \rangle$, $\langle 111 \rangle$ and $\langle 110 \rangle$ stresses. Open circles and \times indicate parallel polarisation, and filled circles and $+$ indicate perpendicular polarisation. Note the similarity in the splitting patterns, and in particular that parallel and perpendicular components are not resolved under $\langle 111 \rangle$ stress and under $[110]$ stress, $[001]$ view—a result characteristic of an A to A transition in tetragonal symmetry. The 1.685 eV perpendicular component under $\langle 100 \rangle$ stress (filled circles) is too weak to observe above 0.4 GPa. Note that the left-hand scale applies to circles, the right-hand scale to crosses (Walker 1975).

one and the same. The oscillator strengths for the optical lines can be estimated as 10^{-2} , indicating allowed electric dipole transitions.

The identity of the defect has not been established; it is probably an interstitial complex involving an impurity (Walker 1977c). Further work is desirable, particularly in EPR.

4.3. Irradiated and annealed diamond

4.3.1. The H lines. These absorption lines were investigated by Clark *et al* (1956b) in irradiated and annealed type Ia diamonds. The more intense ones (H2, H3 and H4) have been much studied and are due to three independent but related centres. H13 may be a transition at the H3 defect (Clark and Norris 1970). Nothing is known of the other H lines.

4.3.1.1. The H2 centre (991.8 nm, 1.25 eV). This centre is produced by heating irradiated diamonds containing more than 10^{25} nitrogen atoms m^{-3} . The anneal temperature (900 K) seems to be critical; lower or higher temperatures do not result in H2.

Heating pre-annealed diamonds at temperatures of 500–600 K in the dark enhances H2 and reduces H3; illumination with mercury light (2.84 or 3.40 eV) reverses the process. Figure 22 shows this in a diamond containing GR1 and ND1 also, but it happens when only H2 and H3 are present. H2 cannot be produced in strength in type Ib or type II diamonds, and there is apparently no EPR associated with it (du Preez 1965, Dyer and du Preez 1967).

Wight (1968) has observed in cathodoluminescence a zero-phonon line at 1.264 eV with a 60 meV phonon replica in an irradiated and annealed diamond. It is possible that this is the emission counterpart of the H2 absorption system. No high-resolution H2 absorption spectrum has been published.

By analogy with Davies' (1977b) suggestion for ND1 and GR1, it may be that H3 and H2 are different charge states of the same defect, since they also show complementary bleaching behaviour. If so, at least one charge state will be paramagnetic, and it should be possible to detect hyperfine interactions due to the two nitrogen atoms contained in the H3 centre. The model suggested for H2 by Dyer and du Preez, a vacancy and an interstitial trapped at a 'nitrogen' platelet, now seems unlikely.

4.3.1.2. The H3 centre (503.2 nm, 2.464 eV). The H3 centre is one of the most studied in diamond (figure 28). Its properties have been reviewed recently by Davies (1977c). It can be observed in absorption, photoluminescence and cathodoluminescence after irradiation followed by annealing at 800 K or above, and is stable to at least 1500 K. Although strongest in type Ia crystals, it has been observed in luminescence in types Ib and IIa, though not in type IIb. The centre has generally been assumed to be a vacancy trapped at a nitrogen aggregate, since it appears when the GR1 centre anneals out, and this is still the current view (Clark *et al* 1956b, Dyer and Matthews 1957, Dean *et al* 1960, Wight 1968, Davies 1976).

The symmetry of the centre is either rhombic I or monoclinic I (table 4), most likely the former. The first (polarised luminescence) experiments indicated a $\pi \langle 110 \rangle$ oscillator at a $\langle 110 \rangle$ centre, provided that low excitation energy (2.84 eV) was used (Elliott *et al* 1958, Clark *et al* 1962, Clark and Norris 1970). The most recent uniaxial stress results (Davies *et al* 1976) are in excellent agreement with rhombic I

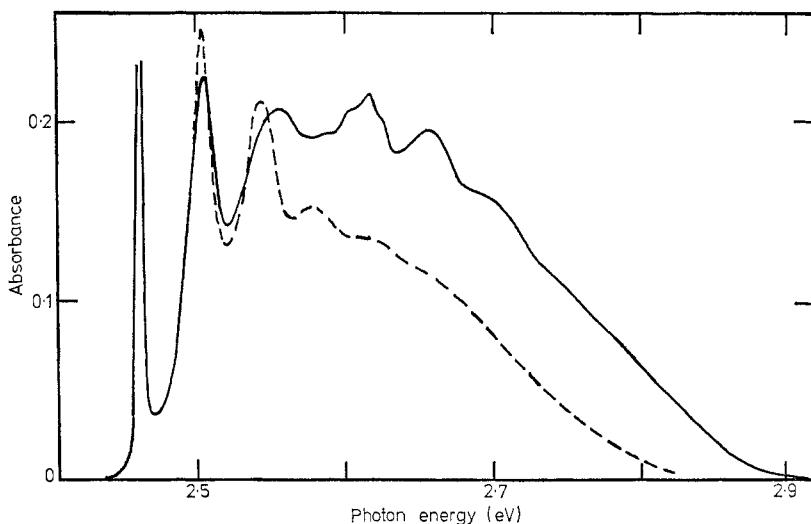


Figure 28. 80 K absorption spectrum of H3 (full curve) compared with mirror image of luminescence (broken curve) (after Davies *et al* 1976).

symmetry. However, there is still a problem in that the Stark effect (Kaplyanskii *et al* 1971) indicates inversion symmetry, whereas C_{2v} , the only rhombic subgroup of T_d , is inversionless.

The uniaxial stress work of Davies *et al* revealed a forbidden transition at the H3 centre. It appears under $\langle 110 \rangle$ and $\langle 111 \rangle$ stress, and is a transition from the A_1 ground state to an A_1 excited state at 2.479 eV, 16 meV above the B_1 state at 2.463 eV. This transition is responsible for the departure from mirror symmetry of the absorption and luminescence bands (figure 28). In luminescence at 80 K the 2.479 eV level is not populated; the emission spectrum is simpler and can be fitted by totally symmetric phonon coupling (Davies 1974b, Davies *et al* 1976). The phonon energies in absorption and emission are slightly different (figure 28), as they are at the GR1 centre (Nedzvetskii and Gaisin 1974b, Davies *et al* 1976). The current model for H3 is therefore a vacancy (i.e. the GR defect) trapped by a nitrogen pair (figure 6); to have rhombic I symmetry the vacancy must exchange with one of the nitrogens, giving an N-V-N arrangement (Davies 1976). The Wedlake model for H3 (see Clark and Mitchell 1971), an N-V pair, was based on Runciman's (1965) incorrect uniaxial stress results, and is now untenable.

Many authors have observed weak H3 luminescence from nominally unirradiated diamond. Recent topographical studies (Hanley *et al* 1977) demonstrates that it is emitted preferentially from slip traces, dislocations and platelets. The luminescence appears to be intrinsic to these features rather than to H3 defects decorating them. Furthermore the H3 centres have been produced without any radiation damage (H3 and H4 can occur in the radiation-damaged rinds of some diamonds, but only at depths of less than 30 μm , corresponding to the range of α -particles). These facts carry implications for the structure of these defects. For example, both the Lang platelet model and the Davies H3 model involve pairs of nitrogen atoms (figure 6) (Walker 1977a). EPR centres observed particularly in plastically deformed diamonds have been shown to contain two nitrogen atoms (Loubser and Wright 1973b, Shcherbakova *et al* 1975, Welbourn and Woods 1977); a possible connection with H3 should be investigated.

We have seen that H13 appears in the excitation spectrum of H3 (Clark and Norris 1970) and is therefore probably a transition at the same centre. H2 and H3 show complementary bleaching behaviour (§4.3.1.1 and figure 22). Hence they may be different charge states of the same defect. H4 is an independent centre, but apparently very similar to H3, as are other centres with zero-phonon lines at 2.417 and 2.536 eV (Davies 1977c, de Sa and Davies 1977). The other H lines may or may not be related to H2, H3 and H4; correlation studies and excitation spectra are necessary. A luminescence line at 2.3 eV apparently accompanies H3 (Dean *et al* 1960, Wight 1968, Sobolev *et al* 1969b, Collins 1974) and could be a transition between excited states of the centre. Because H3 can be bleached, it must be paramagnetic in at least one charge state, though du Preez (1965) was unable to detect any EPR associated with either H3 or H2.

Two other zero-phonon lines, due to the 3H and S1 centres, occur at 503 nm, and the 3H centre also has rhombic I symmetry. As we have already seen, the three centres are distinct but may be related; further work is necessary.

4.3.1.3. The H4 (2.498 eV), 2.417 and 2.536 eV bands. It is convenient to discuss these three centres together because of their similarity to each other and to H3 (figure 29). All of them are produced by irradiation and 900 K annealing of type IaB diamonds. H4 is stable to at least 1200 K. Unlike H3, it has not been observed in synthetic crystals, nor in unirradiated diamond (Clark *et al* 1956b, Dyer and Matthews 1957, Nedzvetskii and Dymke 1970, Sobolev *et al* 1969b, de Sa and Davies 1977, Hanley *et al* 1977).

Davies (1972b) showed that H4 was produced when the B nitrogen aggregate

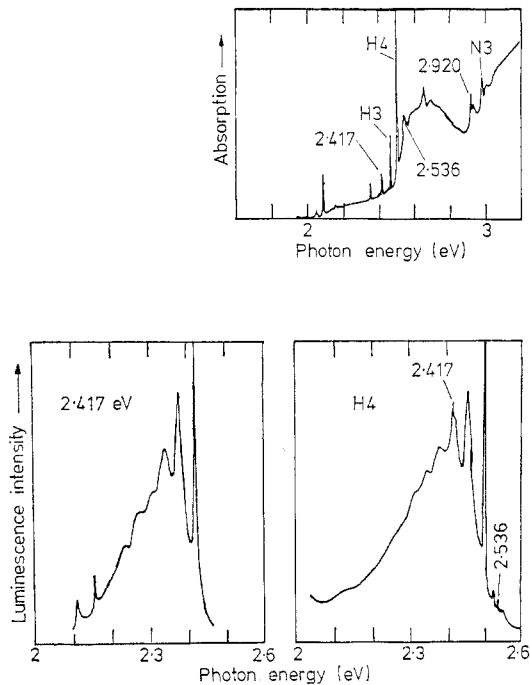


Figure 29. Absorption (upper) and photoluminescence (lower) spectra of the H4, 2.417 and 2.536 eV defects. Various other independent zero-phonon lines can be seen (after de Sa and Davies 1977).

trapped a radiation damage product (now thought to be the vacancy). Runciman (1965) was unable to identify its symmetry, but polarised luminescence (Clark and Norris 1971) indicated either rhombic I or monoclinic I.

de Sa and Davies (1977) have shown that H4 and the 2.417 eV and 2.536 eV centres all have monoclinic I symmetry (with stress coupling coefficients very similar to those of the H3 centre). Although they do not correlate quantitatively, and hence are distinct centres, they typically occur with relative intensities 100: 10: 2. H4 was investigated in most detail and was shown to have a pair of nearly degenerate excited states, to the upper of which transitions are forbidden. This results in a departure from mirror symmetry; a peak at 154 meV in absorption does not appear in the luminescence spectrum (figure 29), exactly like H3 (figure 28). The 2.417 eV centre also shows this asymmetry.

The first phonon peak in absorption is at a higher energy shift than its counterpart in luminescence (de Sa and Davies 1977), a characteristic shared by H3 and the GR1 centre (Nedzvetskii and Gaisin 1974b). This, and the evidence already recounted, strongly suggests that all these centres are aggregates of nitrogen atoms which have trapped a vacancy (i.e. the GR defect) (de Sa and Davies 1977).

Further work could involve the Stark effect, to pin down the point group of the centres (suggested to be C_{1h} rather than C_2 or C_{2h} by de Sa and Davies) and investigation of possible correlations with other nitrogen-dependent centres (e.g. N3, N9 and the EPR centres described by Shcherbakova *et al* (1975)).

4.3.2. The 575 nm system (2.15 eV). This system (figure 30) was discovered by Ralph (1960) using cathodoluminescence. It is produced in all diamonds by irradiation followed by annealing, usually at 900–1100 K, though lower temperatures may suffice, and anneals out at 1400 K. The luminescence can also be photo-excited. In some crystals the cathodoluminescence is present before laboratory irradiation (Ralph 1960, Wight 1968, Davies 1970b, Collins 1977, Hanley *et al* 1977).

Alternatively, continued exposure to the cathodoluminescence electron beam after irradiation will induce and enhance the luminescence, even at 80 K. This

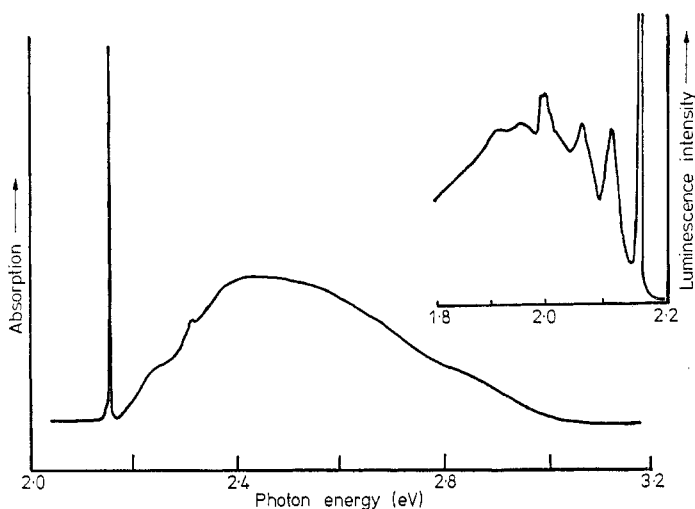


Figure 30. Absorption and cathodoluminescence spectra of the 575 nm system (after Wight 1968, Davies 1977d).

suggests an athermal creation mechanism (Wight 1968, Davies 1970b, 1975b). On one occasion the 3H cathodoluminescence was observed to disappear, the 575 nm system replacing it (Hanley *et al* 1977). This suggests that the latter is created when 3H anneals, or that they are due to different charge states of the same defect; consistent with these possibilities both 3H and the 575 nm system can be created in all diamonds.

Davies (1972b) has observed a weak 575 nm zero-phonon line in absorption in types Ia and Ib crystals. Hence the transition cannot occur between two excited states. The symmetry of the centre is trigonal, the transition being between an A excited state and an E ground state. A low-lying A level is strongly coupled to the E state by stress, implying a Jahn–Teller effect; this is confirmed by the absorption band, which is not a mirror image of the luminescence (Davies 1977d).

No model is currently proposed for the centre. Dean's (1965) suggestion that it was the GR1 defect may now be discounted. Its symmetry, ubiquity and anneal temperature suggest a divacancy.

4.3.3. The 1.944 eV system (637.7 nm). This centre (figure 31) was discovered by du Preez (1965). It is strongest in irradiated and annealed (900 K) type Ib diamonds, both natural and synthetic, though it also occurs in type Ia. The spectrum can be seen in absorption (crystals appear mauve in colour) and in photoluminescence (red), but not apparently in cathodoluminescence. Polarised luminescence indicated trigonal or (less likely) monoclinic I symmetry. Coupling is primarily to totally symmetric

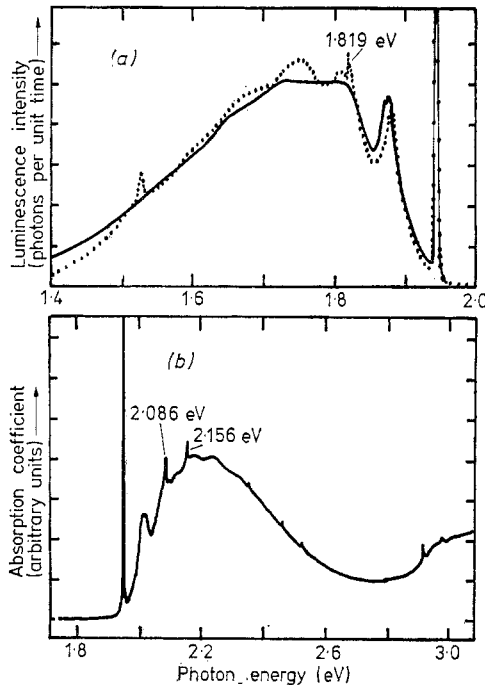


Figure 31. The 1.944 eV centre: (a) measured 80 K luminescence spectrum (dots) compared with luminescence spectrum calculated from the absorption spectrum (full curve). (b) Measured 80 K absorption spectrum. Some independent zero-phonon lines are also present (after Davies and Hamer 1976).

phonons; there is no evidence of Jahn–Teller effects (du Preez 1965, Clark and Norris 1971, Burton 1972, Collins 1974, Davies 1974b, 1977a).

du Preez noted that the 1.944 eV centre appeared as GR1 and ND1 (thought to be vacancies) annealed out. His model, a vacancy trapped at a single nitrogen atom, has been confirmed by Davies and Hamer (1976), whose uniaxial stress experiments show that the 1.944 eV line is due to an A (ground) to E transition at a trigonal defect. Although the E state is degenerate, uniaxial stress confirms that there is no Jahn–Teller effect. de Sa (1977) has shown that a zero-phonon line at 4.328 eV is due to an A–E transition at a trigonal centre, and correlates with the 1.944 eV line. It therefore must be a transition at the same centre.

Loubser and van Wyk (1977) have observed an EPR signal which appears to come from an excited triplet state of the 1.944 eV centre; their work confirms the N–V model.

Klingsporn *et al* (1970) identified a monoclinic (C_{2h}) EPR centre containing nitrogen in type Ib crystals. These crystals fluoresced red and yellow under UV excitation (P E Klingsporn, personal communication). Shcherbakova *et al* (1972) have already associated the EPR centre with the yellow-fluorescing S1 system. It is possible that the 1.944 eV system, since it involves nitrogen, is a different charge state of the S1 defect and gave rise to the red fluorescence in the crystals used by Klingsporn *et al*.

7. Summary and conclusions

I have included in this review the most important defects seen in optical absorption and luminescence in diamond. I have indicated where specific experiments are needed to clear up ignorance or disagreement, and where there are possible correspondences between optical and EPR centres. The last few years have seen a great deal of progress in identifying defects, but there is still much work to be done. Combined optical and EPR experiments are particularly promising. Doped synthetic crystals will permit additional advances, though the area of ‘diamond electronics’ is not in the foreseeable future.

The relevance of the work on diamond to the controversy about how to calculate the energy levels of ‘deep’ defects in semiconductors has already been mentioned. It is to be hoped that recent experimental verifications of ‘defect molecule’ predictions will stimulate further calculations.

Acknowledgments

I would like to thank the Royal Society for the award of a European Programme Fellowship with the Groupe de Physique des Solides de l’Ecole Normale Supérieure, Université Paris VII, during the tenure of which most of this review was written, and Jacques Bourgoïn for making my stay so pleasant and profitable. I am indebted to Professor H H Hopkins, FRS, for making facilities available at the J J Thomson Physical Laboratory, University of Reading, where the review was completed.

I am grateful to the colleagues who have given me help and advice, or have commented on the manuscript: in particular, C D Clark, A T Collins, G Davies, J E Lowther, E W J Mitchell, R C Newman and A M Stoneham.

The authors and publishers who have generously given me permission to reproduce their diagrams here are acknowledged in the captions. I am grateful to Mme Girard for preparing the typescript and to my wife for preparing the diagrams.

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