Phonon modes of MgB$_2$: super-lattice structures and spectral response

Jose A. Alarco,*ab Alison Chou,ab Peter C. Talbotab and Ian D. R. Mackinnona

Micrometre-sized MgB$_2$ crystals of varying quality, synthesized at low temperature and autogenous pressure, are compared using a combination of Raman and infra-red (IR) spectroscopy. These data, which include new peak positions in both spectroscopies for high quality MgB$_2$, are interpreted using DFT calculations on phonon behaviour for symmetry-related structures. Raman and IR activity additional to that predicted by point group analyses of the $P6/mmm$ symmetry are detected. These additional peaks, as well as the overall shapes of calculated phonon dispersion (PD) models are explained by assuming a double super-lattice, consistent with a lower symmetry structure for MgB$_2$. A $2 \times$ super-lattice in the $c$-direction allows a simple correlation of the pair breaking energy and the superconducting gap by activation of corresponding acoustic frequencies. A consistent physical interpretation of these spectra is obtained when the position of a phonon anomaly defines a super-lattice modulation in the $a$–$b$ plane.

1. Introduction

Phonon properties of MgB$_2$ materials have been intensively investigated by optical techniques, such as Raman$_{1-15}$ and infra-red (IR)$_{16,17}$ spectroscopies, and are also complemented by experiments using inelastic X-ray scattering (IXS)$_{18-20}$ and inelastic neutron scattering (INS)$_{21-23}$ Derived phonon information extends to the electron–phonon interaction and superconducting energy gaps.$^{10,15,19,24}$ While these studies have described many properties of MgB$_2$, the spread of experimental results for phonon frequencies is wide and not always consistent. This inconsistency is likely due to variable quality of the investigated materials and to different data acquisition conditions. Anharmonicity,$^{1,2,18,22,23,25,26}$ phonon anomalies,$^{7,19,27}$ presence of impurities,$^{9,11,12,17}$ off stoichiometry from Mg deficiency,$^{19}$ defects,$^{19}$ or unreacted boron,$^{27}$ structural disorder or distortion$^{5,11,14-16}$ and multi-phonon contributions$^{16,20,25}$ are among the cited manifestations or reasons for divergent results on MgB$_2$ property–structure relations.

We have recently synthesized MgB$_2$ under conditions that result in a reproducible, dense and coarse-grained microstructure.$^{28}$ This coarse-grained MgB$_2$ has excellent electrical, magnetic and transport properties. By varying conditions of this synthesis process, we have also produced inferior quality MgB$_2$ which shows a lower superconducting transition temperature ($T_c$ = 38.0 K) and lower density, finer-grained material.$^{28}$ This latter type of material is similar in form and property to that for which several spectroscopic observations on MgB$_2$ are reported.$^{5,11,15}$ This work examines differences in optical properties of two different forms of MgB$_2$, in order to evaluate potential causes for disparity of results in the current literature.$^{1,2,18-20,25-27}$

Density functional theory (DFT) is used to estimate the phonon behaviour of MgB$_2$ based on well-determined crystallographic data using neutron and X-ray diffraction.$^{29-31}$ Point group analysis of the $P6/mmm$ symmetry for MgB$_2$ predicts one Raman and two IR active peaks (see Table 4). However, other studies$^{1,2,8,15,32,33}$ as well as this one, suggest that additional Raman or IR peaks are observed. $Ab\textit{ initio}$ DFT has been used to calculate the phonon density of states (PDOS) and the phonon dispersion (PD) of MgB$_2$. $^{21,32,34-36}$ However, this literature also appears to show somewhat divergent results between computational models and experimental data.$^{1,32,34-36}$ In this work, experimental data are considered in light of modelling to estimate phonon frequencies, density of states and dispersion relations in MgB$_2$ with a particular focus on high quality MgB$_2$ crystals.

2. Methods

2.1 Experimental

Samples for this work are obtained by reaction of Mg metal and NaBH$_4$ precursors using a 50 ml Parr reactor as described by Mackinnon et al.$^{28}$ All materials are handled in a controlled atmosphere glove box containing Argon (99.99%). Starting materials are pre-heated in an evacuated chamber adjacent to the glove box in order to reduce to the dehydrated state and
then transferred to an inert atmosphere. Water and oxygen content in the glove box is normally less than 1 ppm. After MgB₂ is produced via this low temperature, autogenously pressured method, the reaction chamber is cooled to room temperature and opened in the argon-filled glove box via slow pressure equilibration using a gas release valve. In general, material removed from the reaction chamber is placed immediately on substrates or contained within controlled atmosphere environments for subsequent characterisation.

For these Raman experiments, the samples, while still in the glove box, are placed inside a gas-tight Teflon sample chamber fitted with an optical (glass) window for Raman measurements. For IR measurements, the samples are handled in a similar manner to reduce exposure to air at all times up to the initiation of data collection. We have evaluated samples degraded in air over long times (e.g. days) to confirm Raman and IR characteristics of these materials. Data reported in this work are for samples with minimal degradation. Two types of MgB₂ samples are produced by the reactions described by Mackinnon et al. and a summary of essential characteristics is given in Table 1.

Table 1 Characteristics of MgB₂ samples used in this study

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Sample A</th>
<th>Sample B</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maximum temp. of reaction (°C)</td>
<td>500</td>
<td>500</td>
</tr>
<tr>
<td>Heating profile hold temp. (°C)</td>
<td>At 50 and 250</td>
<td>No hold</td>
</tr>
<tr>
<td>Superconducting temp., Tc (°C)</td>
<td>38.5</td>
<td>38.0</td>
</tr>
<tr>
<td>Particle size (μm)</td>
<td>40–80</td>
<td>3–10</td>
</tr>
<tr>
<td>Particle morphology</td>
<td>Euhedral hexagon or bipiramidal</td>
<td>Thin platelet discs</td>
</tr>
<tr>
<td>Dominant color (incident light)</td>
<td>Gold</td>
<td>Black/grey</td>
</tr>
<tr>
<td>Lustre</td>
<td>Bright</td>
<td>Dull</td>
</tr>
</tbody>
</table>

Raman spectra are collected with an inVia Renishaw Raman microscope using two different excitation wavelengths: (i) a near-IR diode laser operating at 785 nm and (ii) a He–Ne laser operating at 532 nm. The laser power at the sample is 20 mW for the 785 nm laser and 10 mW for the 532 nm laser, focussed into a spot size of ~1 μm through a ×50 objective lens. Each Raman spectrum consists of 4 accumulations with a 20 s exposure per scan. Spectrum noise and background are corrected using the smoothing and baseline correction functions in the GRAMS software.

Attenuated total reflectance (ATR) FTIR measurements are recorded with a Nicolet i550 ATR-FTIR spectrophotometer internally purged with dry air. All measurements are performed at room temperature with the sample (exposed to air) in direct contact with a diamond ATR crystal. Data are collected over an accumulation of 8000 scans at a speed of 0.02 min per scan, energy step 0.428 cm⁻¹ and resolution of 4 cm⁻¹ over the range of 350–1800 cm⁻¹.

A Leica multi-focus, stereo optical microscope and a Zeiss Sigma variable pressure field emission SEM with Oxford instruments silicon drift detector (SDD) are used for microscopy observations and energy dispersive spectroscopy (EDS) elemental analysis. Samples are prepared for SEM/EDS by placing a thin layer of powder onto aluminium stubs with double-sided carbon tape. In general, samples are not coated with a conductive coating to avoid analytical interference(s).

2.2 Modelling calculations

The CASTEP module of Materials Studio 7.0 provides the functionality to calculate vibrational and associated properties such as IR and Raman spectra for a wide range of materials. Both linear response (also known as density functional perturbation theory, DFPT) and finite displacement (FD) methods are used in this work. Gradient-corrected exchange–correlation functionals are supported in linear response calculations, although the results may be less accurate than those obtained with local density approximation (LDA) and the computational cost is noticeably higher.

Most calculations are completed using Materials Studio 7.0 via the Microsoft Windows mode of a 12 core Mac Pro Xeon 64 bit workstation. This workstation operates with an Intel chip and mimics up to 24 cores. When memory requirements exceed the capacity of this computer, calculations are undertaken via the high performance computing (HPC) facility at QUT using multiples of 16 cores. The optimum choice for calculations on the HPC facility is 64 cores.

Specific functionals and the acronyms used to describe functionals in these CASTEP calculations (e.g. LDA–CA–PZ or GGA–PBE) for a different, albeit cubic, structure are described by Alarco et al. The DFPT method is implemented in CASTEP for norm conserving potentials (NCPs). On the other hand, FD methods, although in principle available for both NCPs and ultrasoft potentials (UPs) are more effective with UPs. Some FD calculations using NCPs are possible with coarse k-grids but use of a finer k-grid resulted in negative frequency values and difficulty with convergence.

For single cell calculations on MgB₂, the lattice parameters a = 0.3085 nm and c = 0.3523 nm have been used as input. For super-lattice calculations, appropriate multiples of these parameters have been used. Fig. 1 shows a schematic of the reciprocal space lattice for P6/mmm symmetry of MgB₂ where the green and grey spheres represent Mg and B atoms, respectively. Super-lattice constructions of the MgB₂ cell used in this work are multiples along the c-axis direction that retain similar reciprocal space directions.

3. Experimental results

In order to validate spectral data for both Raman and IR, a range of experiments are undertaken on each sample type to
evaluate key parameters that influence interpretation. These experiments are aimed at evaluating the quality of spectra and dependence on, for example, power rating and illumination, wavelength, incident intensity, particle morphology and exposure to air. This a priori approach to Raman and IR spectra for MgB$_2$ is not apparent in the current literature albeit other phases have been evaluated.$^{41-43}$

SEM images of Samples A and B are shown in Fig. 2. The difference in morphology between each sample is demonstrable albeit each sample shows a similar X-ray diffraction pattern for MgB$_2$.$^{28}$

### 3.1 Raman acquisition parameters

Fig. 3a shows a comparison of the Raman spectra for Sample A acquired with varying laser power and exposed to air during the acquisition of spectra. The presence of peaks in the Raman spectrum is apparent when the percentage of laser power is greater than 10%. Peak definition is optimal for Sample A at 100% laser power or $B$ 20 mW. Unlike some materials that, under high power conditions are often overheated or destroyed,$^{44-47}$ both samples appear unaffected in this manner.

Fig. 3b shows a comparison of Sample A and B under the same illuminating conditions. These spectra show that larger size grains enhance the observation of well-defined Raman peaks as noted in earlier work on different compounds by a number of authors.$^{41-43,47}$

Fig. 4 shows an example of Raman spectra from Sample A acquired at 532 nm with freshly prepared material and minimum exposure to air. Notice the peak at $\approx 230$ cm$^{-1}$, which is close to peaks observed and calculated in other Raman studies of MgB$_2$. Additional Raman peaks (labelled with ‘*’*) are also observed in Fig. 4b, presumably due to different orientations of these larger size grains. Minor amplitude variations due to instrument noise are also observed in Fig. 4b (arrowed).

Fig. 5 shows Raman spectra for Sample A collected on different days, at two laser excitations. The spectra show a close match of peak positions at low wavenumbers. Up to nineteen Raman active peaks as compiled in Table 2 can be identified in the spectra collected on this high quality sample of MgB$_2$.

Table 2 gives a summary of the peak positions detected in the Raman spectra from Sample A as shown in Fig. 3–5. Inspection of data in Table 2 shows that some frequencies – within experimental error – are grouped together or are common to both illumination wavelengths. Minor differences in explicit values may be attributed to slightly different temperature effects. However, for experiments summarized in Table 2, low energy peaks consistently occur and occur repeatedly with comparable intensity and peak width. This occurrence implies that these low energy peaks are active to similar extent and are not governed by $P6/mmm$ symmetry.
3.2 Infra-red acquisition parameters

Samples for IR characterization are briefly exposed to air due to the configuration of the sample holder and diamond window. In this method, a reflection configuration provides a path that enhances the IR signal from the interior of a sample which extends the time period for acquisition before sample degradation dominates the spectrum. This technique, which increases the absorption path, reduces the influence of surface effects on IR spectra compared with Raman spectroscopy. Fig. 6 shows IR spectra from a freshly exposed piece of Sample A and from Sample B.

Table 3 lists the approximate peak positions of IR spectra for samples A and B. For sample A, peak parameters are determined by Gaussian peak-fitting routines with GRAMS software.\(^37\) For sample B, peak positions are listed since Gaussian fits do not produce reasonable spectral shapes. This spectrum is also difficult to fit using combinations of Lorentzian and Gaussian peak shapes.

### Table 2: Characteristics of Raman Spectra for Sample A

<table>
<thead>
<tr>
<th>Peak</th>
<th>From Fig. 3(^a)</th>
<th>From Fig. 4</th>
<th>From Fig. 5</th>
<th>From Fig. 5 785 nm</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>230</td>
<td>246</td>
<td>247</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>299</td>
<td>300(^b)</td>
<td>297</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>382</td>
<td>393</td>
<td>398</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>406</td>
<td>406</td>
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<td></td>
</tr>
<tr>
<td>5</td>
<td>464</td>
<td>465</td>
<td>454</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>530–552</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>603</td>
<td>590</td>
<td>594</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>767</td>
<td>767</td>
<td>749</td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>803</td>
<td>800</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>908</td>
<td>906</td>
<td></td>
<td></td>
</tr>
<tr>
<td>11</td>
<td>940–973</td>
<td></td>
<td>958</td>
<td></td>
</tr>
<tr>
<td>12</td>
<td>1062–1073</td>
<td>1047</td>
<td></td>
<td></td>
</tr>
<tr>
<td>13</td>
<td>1385</td>
<td>1378</td>
<td>1345</td>
<td></td>
</tr>
<tr>
<td>14</td>
<td>2433</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

\(^a\) Laser power at 50% produces slightly higher peak shift positions compared with 100% laser power. \(^b\) This position is uncertain due to overlapping intensity at a shoulder peak.

3.3 Modelling results

The presence of additional peaks that may not be described by P6/\textit{mmm} symmetry in experimental data from both Raman and IR methods, suggests computational models may be used to provide higher accuracy and/or precision for interpretation. Thus, modelling in this work evaluates the reciprocal space \textit{k}-grid and relevant grid intervals, the space group and lower symmetry conditions for MgB\textsubscript{2}. Results from these modelling approaches are described below.

#### 3.3.1 \textit{k}-grid interval

The fineness of the \textit{k}-grid is known to affect the accuracy and convergence of phonon calculations as demonstrated in a number of publications including an
The region between 400 cm\(^{-1}\) and 650 cm\(^{-1}\) around the \(G\) (or \(I\)) point is a noteworthy feature (circled in Fig. 7) that varies considerably with change in \(k\)-grid interval. For Raman active frequencies, the region around the \(G\) direction shifts from a single degenerate “dip” at \(\sim 450\) cm\(^{-1}\) at the \(G\) centre point to a degenerate “Mexican hat” configuration (with two adjacent dips) located between the \(G\)–\(K\) and \(G\)–\(M\) directions with decreasing grid interval. In addition, the doublet minima, or approximately degenerate bands, shift(s) to higher frequency at \(\sim 550\) cm\(^{-1}\).

Table 4 shows the values for Raman active frequencies as the interval of the grid is decreased. A substantial change from \(425\) cm\(^{-1}\) to the \(550–650\) cm\(^{-1}\) range occurs with decreasing \(k\)-grid interval. These frequencies span the range of values reported in the literature for MgB\(_2\) under different experimental conditions as noted below.

Similar calculations have been attempted using the FD method, but convergence occurs only with use of UPs. For example, with a coarse \(k\)-grid at \(0.07\) Å\(^{-1}\), anomalous behaviour of the lower phonon bands occurs along the reciprocal space \(AH\)–\(HK\) and \(ML\)–\(LM\) directions. This anomalous behaviour becomes extreme (i.e. with negative values), when NCPs are used with either LDA–CA–PZ or GGA–PBE functionals. However, calculations using UPs eliminate these lower band anomalies in most cases. A strong peak at a very high frequency (\(\sim 890–910\) cm\(^{-1}\)) for the Raman active mode at the \(I\) (or \(G\)) point is maintained at fine grid scale.

Additional calculations with the FD method show that the lowest acoustic band tends to deviate from other acoustic bands when using the GGA–PBE functional and is less pronounced for the LDA–CA–PZ functional. This relative difference occurs for fine \(k\)-grids of \(0.02\) Å\(^{-1}\) and is shown in Fig. 8 as PD plots. However, the peaked Raman active mode reverses shape to form a “valley” near the \(G\) point. This form of the Raman active mode is similar to that determined from DFPT methods with corresponding calculated frequency values closer to experimental values (compare Fig. 7 and 8). Optimum results for the FD method are obtained with this \(k\)-grid interval and a super-cell cut-off radius of 5 Å.

Calculations using a larger cut-off radius and/or a finer \(k\)-grid to improve accuracy led to unviable, extremely long calculation times. These calculations are terminated once the pre-set time limit between three and five days is achieved without iteration progress.

### 3.3.2 Lower symmetry models

Table 5 shows a comparison of active and inactive modes for the space group \(P6_{3}mm\) with other related lower symmetry groups that contain symmetry elements of the parent space group. While the actual frequencies remain near constant, for lower symmetry the Raman and IR active peaks are significantly different. For example, the 413.3 to 416.9 mode is Raman active for \(P6\) and \(P6_{3}mm\) symmetries but not for all other symmetries listed in Table 5. Note that in some cases, the mutual exclusion rule for IR and Raman active modes in centrosymmetric groups is not upheld.39–51
3.3.3 Super-lattice structures. The use of superstructure modes to address the vibrational properties of MgB$_2$ is discussed by Tschauner et al.\textsuperscript{52} For example, a doubling of the $c$ axis parameter for MgB$_2$ and in space group $P\overline{6}_3mc$ allows for two Raman active modes, while space group $P\overline{3}c1$ allows for three modes, $P\overline{6}_3c2$ allows for six and $P\overline{6}_3cm$ allows for eight Raman active modes.\textsuperscript{52} In space group $P\overline{3}c1$, nine modes are both Raman and IR active.\textsuperscript{52} In this work, examples of lower symmetry which cannot be represented in a single unit cell have also been explored. Thus, to achieve the distribution of atoms, we have used super-lattice structures that reproduce the spatial arrangement of Mg and B atoms in the basic MgB$_2$ structure.\textsuperscript{29–31} For the double super-lattice construction in the $c$-direction to represent the primitive cell, a lower symmetry and separate insertion of two B atoms is required. This construct is not necessary for the higher $P\overline{6}/mmm$ symmetry, where inserting one B atom replicates all the other B atom positions. Thus, in this format where B atoms are linked by different symmetry elements compared to the basic $P\overline{6}/mmm$ symmetry of MgB$_2$,\textsuperscript{29–31} movement of adjacent B atoms is less constrained in order to conserve symmetry. A similar situation can be envisaged for Mg atoms under an appropriate choice of reduced symmetry as shown for $P\overline{3}c1$.

![Fig. 7](image1.png)

**Table 4** MgB$_2$ Raman active frequencies for $P\overline{6}/mmm$ symmetry

<table>
<thead>
<tr>
<th>Minimum k-vector [Å$^{-1}$]</th>
<th>Raman active mode [cm$^{-1}$]</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.04</td>
<td>425.89</td>
</tr>
<tr>
<td>0.03</td>
<td>644.61</td>
</tr>
<tr>
<td>0.02</td>
<td>582.29</td>
</tr>
<tr>
<td>0.015</td>
<td>551.04</td>
</tr>
</tbody>
</table>

![Fig. 8](image2.png)

In this work, examples of lower symmetry which cannot be represented in a single unit cell have also been explored. Thus, to achieve the distribution of atoms, we have used super-lattice structures that reproduce the spatial arrangement of Mg and B atoms in the basic MgB$_2$ structure.\textsuperscript{29–31} For the double super-lattice construction in the $c$-direction to represent the primitive cell, a lower symmetry and separate insertion of two B atoms is required. This construct is not necessary for the higher $P\overline{6}/mmm$ symmetry, where inserting one B atom replicates all the other B atom positions. Thus, in this format where B atoms are linked by different symmetry elements compared to the basic $P\overline{6}/mmm$ symmetry of MgB$_2$,\textsuperscript{29–31} movement of adjacent B atoms is less constrained in order to conserve symmetry. A similar situation can be envisaged for Mg atoms under an appropriate choice of reduced symmetry as shown for $P\overline{3}c1$.\textsuperscript{52}
Table 5  Active and inactive modes for P6/mmm and related lower symmetries

<table>
<thead>
<tr>
<th>Group</th>
<th>P6</th>
<th>P6</th>
<th>P6/m</th>
<th>P622</th>
<th>P6mm</th>
<th>P6m2</th>
<th>P6/m/mmm</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>168(^b)</td>
<td>174(^b)</td>
<td>175(^b)</td>
<td>177(^b)</td>
<td>183(^b)</td>
<td>187(^b)</td>
<td>191(^b)</td>
</tr>
<tr>
<td>Frequency (cm(^{-1}))</td>
<td>IR</td>
<td>R</td>
<td>R</td>
<td>IR</td>
<td>IR</td>
<td>IR</td>
<td>IR</td>
</tr>
<tr>
<td>342.8–344.3 (dd)</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
</tr>
<tr>
<td>413.3–416.9</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>N</td>
<td>Y</td>
<td>N</td>
<td>Y</td>
</tr>
<tr>
<td>425.9–644.7 (dd)</td>
<td>N</td>
<td>Y</td>
<td>Y</td>
<td>N</td>
<td>Y</td>
<td>N</td>
<td>Y</td>
</tr>
<tr>
<td>711.6–716.9</td>
<td>N</td>
<td>N</td>
<td>N</td>
<td>N</td>
<td>N</td>
<td>N</td>
<td>N</td>
</tr>
</tbody>
</table>

\(^a\) These values display strong dependence on the dimension of the k-grid. Other frequencies display smaller variations (<6 cm\(^{-1}\)) that are largely attributable to minor differences in lattice parameters calculated during geometry optimization. (dd) indicates a doubly degenerate mode. \(^b\) Value is the space group number.

and can be constructed for the other forms of lower symmetry described in this work.

Fig. 9 shows the PD as a function of decreasing interval of k-grid for the lower group symmetry P3c1. The broad features of the PD for P6/m/mmm symmetry as shown in Fig. 7 and 8 are largely maintained albeit a higher band population along similar reciprocal directions appears. The “Mexican hat” form of the near degenerate bands in the region 500–600 cm\(^{-1}\) still remains. At the smaller k-grid interval (Fig. 9b), an additional band at ~650 cm\(^{-1}\) occurs but is not bifurcated nor a doublet. Fig. 9b also shows that the optical and acoustic bands are in closer proximity compared with Fig. 9a at higher k-grid interval. As expected, along the super-lattice \(\Gamma A\) line direction, substantial creation of new branches and/or breaking of degeneracy can be observed as a result of the folding of the Brillouin zone (in similar fashion to illustrations in Fig. 17 (ref. 42)).

Table 6 shows the frequencies and respective IR and Raman active modes for the lower symmetry super-lattices determined by these PD calculations. The majority of frequencies that correspond to symmetry P6/m/mmm are reproduced. Remarkably, additional frequencies between 230 cm\(^{-1}\) and 240 cm\(^{-1}\) occur. These frequencies are similar to values determined by the experimental measurements noted above. Additional higher frequencies also occur and appear similar to experimentally observed frequencies.

![Phonon dispersion (PD) for group P3c1 calculated with increasing fineness of k-grid.](image)

**Fig. 9**  Phonon dispersion (PD) for group P3c1 calculated with increasing fineness of k-grid: (a) \(k = 0.04\) A\(^{-1}\), (b) \(k = 0.02\) A\(^{-1}\).

4. Discussion

Experimental Raman results from this work show that MgB\(_2\) spectra are sensitive to acquisition and sample condition such as the grain size of the material as well as the energy and power of laser excitation (see Fig. 3). This sensitivity is consistent with prior reports on grain size dependence of Raman peak shapes\(^{41–43,47}\) and partly explains the disparity of Raman results in the literature, as many characterized materials are fine grained. Few large crystals have been analyzed, and of those, primarily along high symmetry orientations.\(^{2,5,9,10,20}\) In other cases, the presence of impurities may influence the quality of observed spectra. For example, in the work by Rafailov \et al.\(^{11}\) MgB\(_2\) formed in a quartz ampoule at 900 °C from Mg powder and B powder shows a grain size of about 45 nm in diameter.\(^{11}\) However, the presence of MgO impurity is indicative of a potential interaction with the quartz tube. Thus, Raman data from this work\(^{11}\) may be difficult to interpret due to impure composition and very fine particle size.

To place this study into perspective, we summarize below prior publications on Raman and IR spectroscopy of MgB\(_2\), in conjunction with prior modelling of MgB\(_2\) phonon behaviour.
A consistent interpretation of these data through model and experiment is possible with an evaluation of symmetry conditions for the MgB$_2$ structure.

### 4.1 Prior spectroscopy

#### 4.1.1 Raman properties.

The general consensus is that MgB$_2$ belongs to space group $P6/mmm$ ($D_{6h}$), as determined from refinements of X-ray and neutron diffraction data.\textsuperscript{24-31} For this space group, factor-group analysis predicts four modes at the $\Gamma$ point: $E_{1u}, A_{2u}, E_{2g}$ and $B_{1g}$, where only the $E_{2g}$ mode is Raman-active, the $E_{1u}$ and $A_{2u}$ modes are IR active, and the $B_{1g}$ mode is silent.\textsuperscript{1,7} According to Quilty,\textsuperscript{10} MgB$_2$ belongs to the $D_{6h}$ point group, with Raman-active symmetry species $A_{1g}, E_{1g}$ and $E_{2g}$, which appear to include Raman activity of the $A_{1g}$ and $E_{1g}$ modes at the $\Gamma$ point.\textsuperscript{1} The $E_{2g}$ Raman-active phonon mode is doubly degenerate at $\Gamma$ and involves in-plane, anti-phase stretching and hexagon-distorting displacements of the B atoms.\textsuperscript{7,22,25,53} Kunc et al.\textsuperscript{1} have shown the decomposition of the coordinate representation and the phonon displacement patterns at both $\Gamma$ and $A$. The branch $E_{2g}(\Gamma)-E_{2u}(A)$ in the phonon dispersion exhibits the strongest electron-phonon coupling.\textsuperscript{1,18,25,34,54} The $B_{1g}(\Gamma)$ is the $z$-displacement analogue of $E_{2g}(\Gamma)$.\textsuperscript{1}

While most Raman experiments detect the frequency $\sim 75$ meV ($= 604.9$ cm$^{-1}$), which corresponds to the $E_{2g}$ mode, measurements also identify peaks with magnitude and temperature dependence that appear closely related to the pair breaking energy or superconducting gap.\textsuperscript{1,6,9,10,24,55} For example, upon cooling of an MgB$_2$ sample below 50 K, Kunc et al.\textsuperscript{1} have observed a build-up of additional scattering intensity in the range 50–300 cm$^{-1}$ (Fig. 8(a) of ref. 1). This additional intensity can be modeled by two Gaussian peaks located at 128 and 226 cm$^{-1}$, respectively, in the 2 K spectrum.\textsuperscript{1} These peaks are observed only for parallel polarizations of the incident and scattered light. With increasing temperature, these peaks vanish at a temperature between 74 K and 100 K; values that are well above $T_c$.\textsuperscript{1}

Additional peaks are also observed in other Raman studies.\textsuperscript{5,9,11} These additional peaks are not attributable to the $E_{2g}$ mode nor to the pair breaking energy or superconducting gap.\textsuperscript{1,6,9,10,24,55} These peaks are of weak magnitude and do not appear to show strong electron–phonon coupling. In addition, the conditions for detection of these weak peaks are limited, easy to confuse and readily attributed to an artifact. For example, Hlinka et al.\textsuperscript{5} have observed humps in Raman spectra of large crystals from thick films at about 300 cm$^{-1}$, 400 cm$^{-1}$, 600 cm$^{-1}$, 750 cm$^{-1}$ and 830 cm$^{-1}$. These results correspond approximately to the reported peaks for PDOS computed in this study and documented above. Rafailov et al.\textsuperscript{11} investigated the Raman spectra of MgB$_2$ and systematically compared the data with potential impurities. In their work,\textsuperscript{11} peaks at 250 cm$^{-1}$, 600 cm$^{-1}$ and 750 cm$^{-1}$ are assigned to MgB$_2$. However, in the same study, peaks at 255 cm$^{-1}$, 517 cm$^{-1}$, 1370 cm$^{-1}$ and 1590 cm$^{-1}$, and another set at 380 cm$^{-1}$, 560 cm$^{-1}$ and 960 cm$^{-1}$ are measured as the light and dark areas (which we interpret to represent Mg and MgO), respectively, of an unreacted Mg stripe.\textsuperscript{11}

#### 4.1.2 IR properties.

IR spectroscopy studies on MgB$_2$ are not as abundant as Raman spectroscopy experiments. Sundar et al.\textsuperscript{16} measured the IR spectra of MgB$_2$ in the range of 125 cm$^{-1}$ to 700 cm$^{-1}$ as a function of temperature to 5 K. They found that the absorption spectrum is characterised by a broad band centered at 485 cm$^{-1}$, with shoulders at 333 cm$^{-1}$ and 387 cm$^{-1}$. Additional further sharp features are observed at 542 cm$^{-1}$, 592 cm$^{-1}$ and 633 cm$^{-1}$, which appear to match modes of $\beta$-rhombohedral boron. These features are attributed to the activation of boron-like modes in MgB$_2$ by disorder. Other high frequency modes manifested as broad bands centered at 1040 cm$^{-1}$, 1442 cm$^{-1}$ and 1635 cm$^{-1}$ appear to be a multiple of the absorption band at 485 cm$^{-1}$ and hence, may be due to a combination of modes.\textsuperscript{16} In a subsequent publication,\textsuperscript{17} the small features at 333 cm$^{-1}$, 387 cm$^{-1}$, 542 cm$^{-1}$, 592 cm$^{-1}$ and 634 cm$^{-1}$ are associated with small amounts of unreacted B, as can be inferred from comparison with the spectrum for crystalline B.

### 4.2 Prior computational modelling of phonons

*Ab initio* DFT has previously been used to calculate the PDOS and the PD of MgB$_2$.\textsuperscript{21,32,34,36} Kong et al.\textsuperscript{34} use the linear-response, full-potential, Linear-Muffin-Tin-Orbital (LMTO) DFT method, developed earlier by Savrasov.\textsuperscript{56,57} Shukla et al.\textsuperscript{32} use DFT GGA with NCPs and linear response to compute harmonic frequencies and to determine a theoretical PD that matches their experimental data from IXS. Bohnen et al.\textsuperscript{35} calculate the PD of MgB$_2$ to compare with that of isostructural AlB$_2$, using a mixed-basis pseudopotential method. This mixed pseudopotential model is a linear response approach similar to that of Kong et al.\textsuperscript{34} Osborn et al.\textsuperscript{21} have used a simple Born–von Karman (BvK) model for the PDOS to represent the main peaks in their INS experiments. Other works, which limit the calculation to phonon frequency shifts, often use the frozen phonon approach.\textsuperscript{1,26,54} This approach is only valid when coupling between different modes is negligible and anharmonic interaction does not change significantly over the Brillouin zone.\textsuperscript{26}

According to Parlinski,\textsuperscript{36} far from the $\Gamma$-point, the acoustic PD relations reflect the vibrations of the Mg atoms in MgB$_2$. The optic branches describe vibrations of the graphite-like boron network. Moreover, the branches of symmetry $A_{2u}$ and $B_{1g}$ at the $\Gamma$-point correspond to the out-of-plane boron vibrations, while four remaining curves with symmetry $E_{1u}$ and $E_{2g}$ at the $\Gamma$-point determine the in-plane boron motion. The out-of-plane and in-plane polarizations are approximately along the same branches all over the Brillouin zone.\textsuperscript{36}

For reference, Fig. 10 shows the directions of vibration for boron atoms in the MgB$_2$ structure defined by the $E_{2g}$ mode. A dynamic description of vibration modes for MgB$_2$ is given by Yildirim.\textsuperscript{33}

### 4.3 Spectral data – symmetry implications

Our experimental results from both Raman and IR spectroscopy as described above indicate that a higher number of
peaks than are predicted by group theory analysis for $P6/mmm$ symmetry occur for Sample A. Conventional analysis for the $P6/mmm$ space group predicts one Raman active mode and two IR active modes. In comparison, our data suggest at least seven modes are active in both Raman and IR spectra obtained from Sample A. In addition, some peak frequencies appear to be both Raman and IR active. Several reasons for this spectral response are examined below.

A random orientation of large grained, polycrystalline, materials and larger crystal faces of Sample A likely contribute to ready detection of additional peaks compared with Sample B albeit additional peaks are identified. The presence of extra peaks in Raman spectra of MgB$_2$ is not exclusive to this work alone (see Section 4.1.1). In addition, several peaks appear to be both Raman and IR active. This outcome indicates not only that the symmetry is reduced with respect to $P6/mmm$, but also that the reduced symmetry may be non-centrosymmetric. For centrosymmetric materials, the rule of mutual exclusion for Raman and IR spectra requires a peak to be either Raman or IR active. $^{49-51}$ However, this need not apply for non-centrosymmetric materials.

Earlier calculations of MgB$_2$ vibrational behaviour $^{26,54}$ demonstrate use of the frozen phonon approach. However, as mentioned by Kunc et al.,$^1$ the frozen-in displacements required to perform these calculations lower the rotational symmetry of the system from $D_{6h}$ (or $P6/mmm$) to $D_{3h}$, among other possibilities. Our calculations with reduced symmetry soften the constraints on Raman and IR activity without significant change to the frequency values as shown in Table 5.

Quantitative determination of specific factors such as anharmonicity,$^{1,2,18,22,23,25,26}$ phonon anomalies,$^{7,19,27}$ presence of impurities,$^{9,11,12,17}$ and multi-phonon contributions$^{16,20,25}$ on the perturbation of first order vibrational features of MgB$_2$ is outside the scope of this article. However, it is worth noting that the Raman spectra from these experiments show fluorescence in the 1000–4000 cm$^{-1}$ range (see Fig. 3a). This fluorescence is likely to be related to the generation of second order vibrational features.$^{58}$

Constructing a super-lattice to model structures with reduced symmetry appears to match earlier determinations of MgB$_2$ phonon dispersions.$^{21,32,34-36}$ However, the match between model and experiment is reasonable only if we consider that the multiplicity or near degeneracy of certain bands is partly hidden by experimental errors, or limited resolution, of the data. For instance, the $E_{1u}$ energy at the $\Gamma$ point in Fig. 3 of the article by Shukla et al.$^{32}$ appears to be the same energy as the top region of the highest energy acoustic mode at the $M$ point. This study (Fig. 6 and 8) shows that a super-lattice with reduced symmetry reproduces the experimental data of Shukla et al.$^{32}$ This alternative interpretation is informative, given that phonon dispersions in the work reported by Shukla et al.$^{32}$ are constructed to fit experimental data from IXS,$^{2,18-20,32}$ and INS,$^{21-23}$ respectively.

Symmetry conditions may also influence the accuracy of computational models for MgB$_2$ spectra and the nature of Raman or IR activity. For example, experimental results and models more closely align when the 6-fold rotation axis is reduced to a 3-fold rotation axis, resulting in a doubling of the unit cell in the $z$ direction. This difference in symmetry description – which invokes a super-lattice in the $z$ direction – predicts a higher number of Raman and IR active modes than with conventional six-fold symmetry. This outcome suggests that the Mg atoms have also lost 6-fold symmetry. The manifestation of this lower symmetry is an alternating Mg atom position towards and away from the plane containing boron atoms. Furthermore, the reduced symmetry super-lattice model predicts an additional frequency at about 235 cm$^{-1}$. This frequency is observed in a number of experiments including this work as shown in Table 2.

### 4.4 Comparison of DFPT and FD modelling methods

Within DFT, the calculation of lattice dynamics is, in essence, the change in electronic charge density induced by the presence of a phonon with wave vector $q$. $^{56}$ Within the framework of the linear-response (or DFPT) method, calculation of the phonon spectra and the electron–phonon interaction is reduced to finding the first-order variations in the one-electron wave functions, the charge density and the effective potential induced by the presence of a phonon with a given wave vector $q$. $^{57}$ In addition, the linear response method provides an analytical means to compute the second derivative of the total energy with respect to a given perturbation.

On the other hand, the FD method, as exemplified by the extension of the LMTO method described by Savrasov, $^{56}$ is an analytical version of a finite-difference approach within a super-lattice. When applied to the same problem, the results of both approaches, DFPT and FD, must be the same except for errors that may be introduced by taking finite differences.

As noted, $^{39}$ phonon spectra in metallic compounds are very sensitive to details of the Fermi surface. To obtain reasonable spectra using the linear response method for metals, the $k$-point sampling must be dense and significantly higher density than the Fine mesh setting in CASTEP.$^{39}$ Based on experience, we recommend setting the separation parameter on the $k$-points tab of the CASTEP electronic options dialog to less than 0.02 Å$^{-1}$. A clear sign of insufficient quality of $k$-point sampling is the presence of imaginary acoustic modes near the $\Gamma$-point.
In these calculations, the pseudo potentials mimic how valence electrons experience the screened core potential. In practice, agreement is expected between the integrated real and pseudo charge densities outside a chosen core radius for each valence state. The norm conservation criteria essentially ensures that the total charge of each pseudo wave function equals the charge of the all-electron wave function potential. For both DFPT and FD methods, changes in the $E_{2g}$ region of the PD are accomplished by changes in the $\Gamma A$, $\Gamma M$ and $\Gamma K$ directions. However, these changes are more pronounced in different phonon branches.

The PD calculated from the DFPT model, when using a sufficiently dense grid ($k < 0.03 \text{A}^{-1}$) displays the typical “Mexican-hat” topology for the mode around $E_{2g}$ as shown in Fig. 6. According to Kunc et al., this mode corresponds to vibrations in the boron plane. Such topology has been extensively discussed for systems displaying the Jahn–Teller (JT), or pseudo-Jahn–Teller effect and may be a signature of this effect in the MgB$_2$ system. Moreover, calculations with LDA and GGA functionals result in identical PDs in DFPT, while LDA and GGA clearly differ in the FD approach. Apparently, the charge redistribution in the DFPT method, required with the charge renormalization for NCPs, is accounted for primarily by the “Mexican-hat” changes in the PDs. In comparison, charge re-distribution is not imposed to the same extent in the FD method and results in larger modifications of the PD in the acoustic band region.

4.5 Phonon anomalies

Phonon anomalies, primarily in acoustic branches, are associated with improvement in the superconducting transition temperature ($T_c$) of structurally related materials. In the case of MgB$_2$, Wood et al. argue for the existence of an acoustic phonon anomaly in the $\Gamma A$ direction on the basis of DFT calculations. Their results resemble the crossing of the second and third branches that are observed in PD calculations shown in Fig. 7 and 9.

Baron et al. identify the anomaly of the $E_{2g}$ mode as a Kohn anomaly, which originates from partial screening of lattice vibrations by the conduction electrons. This screening changes rapidly on certain surfaces in the space of phonon $q$-vectors and therefore, on these surfaces the frequencies vary abruptly with $q$. Baron et al. also report the presence of an anomalous optical mode in MgB$_2$, similar in energy to that of the $E_{2g}$ mode, but with a different line width and symmetry. This mode is longitudinal along $\Gamma A$ and not predicted by the theory used at the time. Remarkably, calculations of the PD shown in Fig. 9 (and listed in Table 6) for the double cell symmetry predict a flat branch at $\sim 650 \text{ cm}^{-1}$, which is similar to values associated with the $E_{2g}$ mode.

4.6 Reduced symmetry dynamics

The inference from these calculations that adjacent Mg atoms may move out of phase for a 3-fold reduced symmetry indicates that Mg atoms may also respond to in-plane boron atom vibrations. Therefore, the $E_{2g}$ mode would take place simultaneously with, or include a strong coupling to, adjacent Mg atomic movements. For example, coupling with $A_{1g}$ modes is a reasonable expectation and may be evident in high resolution neutron diffraction investigations on Mg$_{1-x}$Al$_x$B$_2$ by Campi et al. In Fig. 3 from this investigation, it is shown that for no doping ($x = 0$), the radial distribution corresponding to the fourth periodicity in the $c$-direction is not four times the $c$-lattice parameter, in contrast to the first three periodicities (1, 2 and 3) which when divided by the respective multiples, all converge onto the same $c$-lattice parameter.

This work highlights the dynamic nature of structural symmetry when phonons are included. Phonons are equivalent to dynamical perturbations of the lattice oscillating at the frequency $\omega$, as pointed out by d’Astuto et al. As such, the phonon self-energy varies with time. To properly account for this, it is important to go beyond the adiabatic approximation to make use of time dependent perturbation theory. According to Calandra et al., X-ray and Raman measurements are both explicable if dynamic effects beyond the adiabatic Born–Oppenheimer approximation as well as electron self-energy effects are included in the determination of phonon self-energy.

The use of reduced symmetry approximations as described in this study improves the description of phonon behaviour within the limitations of the adiabatic approximation. The potential JT-like PD that displays a “Mexican-hat” topology may be related to oscillatory electronic bands that are predicted to swing above and below the Fermi level as the lattice vibrates, with a corresponding charge population re-distribution when the band crosses the Fermi level.

4.7 Phonon behaviour and superconductivity

The phenomenon of emergence and disappearance of Raman peaks as a function of temperature across the superconducting transition temperature, $T_c$, is indicative of a strong link to superconductivity. The above discussion suggests that a peak at $\sim 116 \text{ cm}^{-1}$ in Raman spectra may be attributed to a vibration mode in a lower symmetry structure. The energy of this peak at $\sim 116 \text{ cm}^{-1}$ is approximately equal to the superconducting gap $2D = 4k_BT_c$ ($\sim 111 \text{ cm}^{-1}$) for $T_c \sim 40 \text{ K}$, where $k_B$ is Boltzmann’s constant.

In this work, experimental data and computational models show that $\sim 230 \text{ cm}^{-1}$ is an $A$-zone boundary frequency for the original $P6/mmm$ unit cell (see Fig. 6 and 7). This frequency becomes part of the $F$-centre frequencies of the $2\times$ super-lattice in the $c$-direction, following a procedure similar to that discussed by Kunc et al. and shown in Fig. 9. Inspection of the $A$-zone boundary for the $2\times$ super-lattice also shows that a frequency $\sim 116 \text{ cm}^{-1}$ has been created at the new $A$-zone boundary (see Fig. 9). Therefore, an extension of the super-lattice symmetry to a $4\times$ cell in the same direction as the $2\times$ super-lattice results in a frequency value at the $F$-centre zone of $116 \text{ cm}^{-1}$. This analysis establishes a correlation between key phonon frequencies and a super-lattice with a commensurate modulation in the $c$-direction. Such a modulation...
is likely to be of a dynamic nature and involve electron transfers to bonds and spin polarizations that define the super-lattice.

A fourfold modulation in the $c$-direction (i.e. a $4 \times$ super-lattice) corresponds to approximately $4 \times 3.5 \, \text{Å} = 14 \, \text{Å}$. This value is approximately half the coherence length for MgB$_2$ in the $c$-direction.$^{72,73}$ If we consider that the phonon anomaly minimum also corresponds to a super-lattice modulation in the $a$-$b$ plane, we obtain approximately $7 \times 3.08 \, \text{Å} = 21.56 \, \text{Å}$, for the period of modulation. A factor of 7 times arises because the bottom of the anomaly is located at $\sim 2$ calculation points from the $\Gamma$-centre zone out of a total of 14 points that describe the direction to the $K$-zone boundary (see Fig. 6). A modulation in the $a$-$b$ plane of $\sim 22 \, \text{Å}$ is approximately half the coherence length of about $44 \, \text{Å}$ for MgB$_2$ in this direction.$^{72,73}$

At the new zone boundary created by a super-lattice modulation in the $a$-$b$ plane, extension of a line along the k-vector that corresponds to the minimum of the phonon anomaly determines the frequencies of the acoustic bands. This linear extension intersects the acoustic bands at frequencies $\sim 50 \, \text{cm}^{-1}$, $\sim 75 \, \text{cm}^{-1}$ and $\sim 120 \, \text{cm}^{-1}$, respectively (see Fig. 9). These three frequencies have been detected in Raman investigations of MgB$_2$ with end values discussed in terms of pair breaking mechanisms and two superconducting gaps.$^{15,55}$

### 4.8 Energy gaps and phonon relaxation

Tables 5 and 6 show that the following approximate balance of energies applies

$$A_{1g} (= B_{1g}) = E_{2g} + 116 \, \text{cm}^{-1}$$

where $B_{1g}$ and $A_{1g}$ are the highest frequencies in Tables 5 and 6, ($B_{1g}$ and $A_{1g} \approx 710$–715 cm$^{-1}$) and $E_{2g}$ is $\sim 600 \, \text{cm}^{-1}$, respectively. $B_{1g}$ and $A_{1g}$ may convert with each other via accompanying charge movements that transforms group symmetry from $P6/mmm$ to the lower symmetry double super-lattice. This equivalence provides not only a balance of energy, but an approximate mechanism where movements confined to the $a$-$b$ plane may convert into movement in the $z$-direction leading to the creation of a switchable (on–off) dynamic super-lattice with corresponding boundary energy gaps.

An atom with three degrees of freedom has $k_B T/2$ thermal energy per degree of freedom.$^{74,75}$ Since MgB$_2$ has three atoms per unit cell, the thermal energy per unit cell of MgB$_2$ is $4.5k_B T$. At $T_c$, this thermal energy provides energy just above the superconducting gap $2\Delta = 4k_B T_c$, and thus, suggests that excitation of phonon modes that are separated by a gap energy is likely to be a significant determinant of superconductivity.

Fig. 11 shows a plot of the calculated frequencies for the $2 \times 2 \times 2$ super-lattice using spectral values from Table 6, assuming integer multiples of the energy equivalent to the vibration mode of the lower symmetry structure noted in 4.7 above (i.e. $\sim 116 \, \text{cm}^{-1}$). These $ab\ initio$ calculated frequencies define a linear trend based on the major acoustic mode for this super-lattice symmetry with other nearly parallel, slightly offset values. This trend suggests that the frequencies of the MgB$_2$ structure at the $\Gamma$-point have some similarity to those of an 'harmonic oscillator', where the allowed frequencies are expressed as integer multiples of a basic vibration.$^{76–79}$

Coherent phonon relaxation has been investigated in semiconductors$^{50,80}$ and is the basis for coherent phonon spectroscopy.$^{81}$ Coherent phonon relaxation is described in terms of phonon decay time and dephasing.$^{50,80,81}$ Decay times are typically longer at lower temperatures and dephasing may induce anharmonicity. Primary mechanisms in coherent phonon relaxation are up-conversion of modes by thermal phonons as well as down-conversion by decay of optical phonons into acoustic phonons.$^{50,80,81}$ In this latter case, acoustic phonons carry half the energy of the predecessor optical phonon but with opposing $k$-vectors, thus conserving energy and momentum while avoiding incoherent scattering.

The linear integer proportionality of phonon frequencies at the $\Gamma$-point for MgB$_2$ noted above suggests that conversion of modes – where energy and momentum as well as phase coherence are conserved – is inferred from our combined experimental data and DFT calculations. For example, if the temperature cannot excite up-conversion from a particular energy level to the next higher energy mode, all remaining relaxation options are decay modes with phase coherent mechanisms.

This mechanism is exemplified by the available conversion processes from the important $E_{2g}$ mode ($\sim 581 \, \text{cm}^{-1}$). For example, when temperature remains above $T_c$, excitation into the $712 \, \text{cm}^{-1}$ mode is enabled by thermal phonons. Down-conversion from this higher level leads to scattering and resistive losses because decay paths with exclusively coherent options cannot be established. Once the temperature cannot excite up-conversion to the $712 \, \text{cm}^{-1}$ mode, the only remaining options for relaxation are decay modes within major phase coherent mechanisms.

Decay from the $E_{2g}$ mode at $\sim 581 \, \text{cm}^{-1}$ via two acoustic modes at $\sim 116 \, \text{cm}^{-1}$ (with opposing momentum) lands at an energy level of $351 \, \text{cm}^{-1}$ which is observed in Raman spectra of MgB$_2$ (as shown in Table 2), and is also calculated for the $2 \times 2 \times 2$ super-lattice (as shown in Table 6). The energy at $351 \, \text{cm}^{-1}$ belongs to the set of multiple integer values for MgB$_2$. Such a mechanism is not accessible for decay from the $712 \, \text{cm}^{-1}$
mode, since two acoustic modes at \( \sim 116 \text{ cm}^{-1} \) do not land within available integer values. It is worth noting that the 351 \text{ cm}^{-1} mode energy is approximately equivalent to half the B_{1g} and A_{1g} energy levels (at \( \sim 712 \text{ cm}^{-1} \)), which may also be important for energy conservation.

5. Conclusions

Combined Raman, IR and DFT investigations of high quality micrometer-sized MgB_2 allows a consistent explanation for experimentally-determined optical properties. Peaks in excess of those determined by group analysis of the \( \overline{P6/mmm} \) symmetry have been detected using both Raman and IR spectroscopies. Some peaks display both Raman and IR activity, which indicates that the symmetry is reduced from \( \overline{P6/mmm} \) to a non-centrosymmetric type.

PDOS calculations are insensitive to subtle changes in the PD produced by both commensurate and incommensurate super-lattice modulations. Similarly, frozen phonon approaches, which are limited to accurate determination of vibration frequencies at the \( \Gamma \) point, also mask crucial information at super-lattice positions. Computationally, the use of UPs appears to mask fine-scale charge re-distributions affecting the PD. Such charge re-distribution is associated with fluctuations of the nearly flat bands running parallel to the Fermi energy. 69–71

DFT calculations of PDs and vibrational frequencies associated with a double cell in the c-direction explain the observed experimental data not only in this study but from several others which use different analytical techniques. PDs calculated with a \( 2 \times \) super-lattice produce a better overall shape and match to PDs determined from IXS and INS experimental data. Use of super-lattice symmetries for MgB_2 is a better-modelled approximation to dynamic modulations introduced by phonons.

Energy conservation through conversion of phonon energies by coherent relaxation may be manifest in the set of multiple bands running parallel to the Fermi energy. 69–71 DFT calculations of PDs and vibrational frequencies associated with a double cell in the c-direction explain the observed experimental data not only in this study but from several others which use different analytical techniques. PDs calculated with a \( 2 \times \) super-lattice produce a better overall shape and match to PDs determined from IXS and INS experimental data. Use of super-lattice symmetries for MgB_2 is a better-modelled approximation to dynamic modulations introduced by phonons.

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