The $\lambda6614$ diffuse interstellar absorption band: evidence for internal excitation of the carrier

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1 INTRODUCTION

Identification of the carriers of the enigmatic Diffuse Interstellar Bands (DIBs) is often referred to as the biggest current challenge in astronomical spectroscopy. The observational data comprise over 500 interstellar absorption features, predominantly at visible wavelengths, which are observed towards bright O and B stars seen though significant foreground material. Although this field of research was established nearly one hundred years ago, efforts to identify the carriers over the years have not met with success. However, in a significant development, based on the results of gas-phase laboratory experiments, (Campbell et al. 2013) conclude that they have positively identified two near-infrared diffuse interstellar absorption bands as due to transitions of $\text{C}_n^+$. The overall subject area has been reviewed by Herbig (1974) and Sarre (2006).

Some of the diffuse bands have been known to exhibit distinct shapes for at least forty years (Herbig 1974; Westerlund & Krelowski 1988a,b). Diffuse band substructure in the form of a triplet was first observed in the $\lambda6614$ band by Hibbins et al. (1994) towards $\mu$ Sgr, $\sigma$ Sco, $\beta^1$ Sco and $\zeta$ Oph. Using the Ultra-High-Resolution-Facility ($\text{RP} = 600,000$) on the Anglo-Australian Telescope, the diffuse bands $\lambda6614$ and $5797$ were found to have distinct components within the profiles, which provides support for a molecular rather than a condensed-phase origin (Sarre et al. 1993; Kerr et al. 1996; Kerr et al. 1998). Rotational contour modelling of the $\lambda6614$ triplet profile towards $\mu$ Sgr using R, Q and P-type rotational branches yielded planar oblate symmetric top molecules of polycyclic aromatic hydrocarbon (PAH) or cyclic form as potential carriers (Kerr et al. 1998). It was noted by Kerr et al. (1996) that an additional absorption found in spectra of $\lambda6614$ towards $\mu$ Sgr was not accounted for in ‘R, Q, P’ triplet interpretation or by the contour modelling, but that it might be explained in terms of ‘vibrational hot bands, isotopic modifications or splitting of the level structure by magnetic (e.g. spin-orbit) or Jahn-Teller interactions’. An additional feature labelled ‘c’ that appears on the redward side in spectra of $\lambda5797$ (Kerr et al. 1995) probably has a similar origin.

Triplet and doublet structure for $\lambda6614$ and $\lambda5797$, respectively, was reported by Ehrenfreund & Foing (1994) at $\text{RP} = 70,000$ as well as a doublet for $\lambda6379$; the separation between ‘R’ and ‘P’ peaks enabled estimates of rotational constants of the carriers to be obtained which are consistent with carriers being PAH molecules with more than 40 carbon atoms, 12-18 carbon chains or compounds of $\text{C}_{60}$ (Ehrenfreund & Foing 1996). This was further developed by Cami et al. (2004) who noted variation in the relative positions of the peaks within $\lambda6614$ according to sight-line, which could be interpreted in terms of the rotational excitation temperature of a molecular carrier. An alternative interpretation for the multiplet structure has been offered by Webster (1996) in which the substructure peaks are attributed to $^{13}\text{C}$ isotopic modifications of the carrier.

Newly discovered extended red tails for some of the diffuse bands recorded through an infrared-illuminated region towards Herschel 36 have also been interpreted in terms of molecular rotational contours (Dahlstrom et al. 2013; Oka et al. 2013). In their analysis the extended contours of $\lambda5780$, $5797$ and $6614$ are considered to be due to polar

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molecules with three to six heavy carbon-like atoms. In contrast, λλ5850, 6196 and 6379, which do not have pronounced tails, were suggested to be likely due to non-polar molecules or large polar molecules with a low relative change in rotational constant on electronic excitation. For both contour types only a single vibronic band was included in the calculations. While attributing the contour shapes to single rotational contours of relatively small molecules, the authors did not rule out other interpretations including the role of low-frequency vibrations.

In a recent study Bernstein et al. (2015) concluded that the overall profiles of λλ6614 detected in diffuse and translucent clouds result from accidentally overlapping diffuse bands centred at 6613.6 Å (the strongest component of the triplet) and at 6614.2 Å (a single highly asymmetric red degraded band), where these independent bands are due to different carriers - based on a suggestion made earlier by Galazutdinov et al. (2002). This interpretation differs markedly from the internal excitation ‘hot band’ origin as put forward by Kerr et al. (1999) which involves a single carrier and is further considered and modelled here.

Particularly in the light of high-quality HARPS data we have re-examined the hot-band suggestion, including line-of-sight profile comparison, co-addition of principal and hot band profiles, and molecular rotational contour calculations. We describe a model in which the absorption on the redward side of λλ6614 for e.g. HD 147889 – which is enhanced in comparison with HD 179406 – is attributed to ‘hot band’ contributions, as would be expected to arise due to a higher internal ‘temperature’ in a molecule with low-lying vibronic states. For simplicity we choose to use the term ‘hot band’ irrespective of the precise vibrational, electronic etc. origin of the excited level(s) which is unknown.

2 DIFFUSE BAND PROFILES AND MOLECULAR ABSORPTION FEATURES

In diffuse clouds the rotational temperature of optically detected polar interstellar molecules such as CN is generally close to that of the cosmic microwave background radiation temperature of 2.72548 ±0.00057 K (Fixsen 2009), though is generally found to be slightly higher, probably due to the effect of electron collisions (Black & van Dishoeck 1991, Ritchey et al. 2011, Krelowski et al. 2012, Leach 2013, Harrison et al. 2013). A notable exception is the line-of-sight towards Herschel 36 with rotational temperatures of 6.7 K and 14.6 K determined for CH and CH⁺, respectively. The highly unusual display of diffuse band shapes most probably arises from a higher-than-usual degree of diffuse band carrier internal excitation. Electronic absorption spectra of non-polar molecules in diffuse clouds including H₂, C₂ and C₃ carry information on the ground-state rotational level populations which are determined by the balance between collisional and radiative excitation processes; effective rotational ‘temperatures’ of up to c. 200 K have been found. However, the physical interpretation of the rotational level column densities is not straightforward as discussed by Cassu & Cecchi-Pestellini (2012), Hupe et al. (2012), Welty et al. (2012) and in citations of earlier work.

Observations of variation in diffuse band shapes include both blue and red shifts and band broadening where these have been shown to be unrelated to Doppler effects from thermal or turbulent motion or due to multiple line-of-sight clouds. Early examples include λλ5780 (Porceddu et al. 1992) for which unusually large widths were found towards three stars in Orion, and small red shifts for λλ6614, 5780, 5797, 6284 and 5705 towards HD 37022 and HD 37061 which are also in Orion (Krelowski & Greenberg 1999). Comparative blue shifts have also been reported for HD 37048 and stars in the Sco OB1 association (Galazutdinov et al. 2003, 2008, 2013). High-resolution HARPS spectra have revealed marked differences in diffuse band profiles, for example towards HD 179406, HD 163800, HD 147165 and HD 147889 (Galazutdinov et al. 2008). A good overall summary of studies of diffuse band profile variation is presented in section 4.2 of Dahlstrom et al. (2013).

3 OBSERVATIONAL DATA

The star HD 179406 (20 Aql) lies behind a ζ-type interstellar cloud (W(5797)/W(5780) = 0.48, Eᵣ₋ᵥ = 0.31), as does HD 147889 (W(5797)/W(5780) = 0.42, Eᵣ₋ᵥ = 0.30). Of particular importance, as inferred from the velocity profiles of the K₁ and CH absorption lines, there is a single dominant velocity component and hence very likely a single interstellar cloud in each case (Galazutdinov et al. 2008). However, this does preclude the presence of regions with differing velocity dispersions or conditions within the cloud as the spectrum recorded is integrated along the line-of-sight. The profile of λλ6614 is known to differ significantly with line-of-sight - see fig 3. of Galazutdinov et al. (2008), both in band width and in the relative strengths of the triplet components. The observational data considered here were obtained by Galazutdinov et al. (2008) using the High Accuracy Radial Velocity Planet Searcher (HARPS) échelle spectrometer, with sightlines selected where possible based on their having a single CH 4300 Å line with low velocity dispersion (Galazutdinov et al. 2008). The laboratory rest frequency for the carrier of λλ6614 is obviously not known and it is assumed that it shares the same radial velocity shift as for CH as diffuse band strengths are generally found to correlate well with the column density for this molecule, when observed.

4 PROFILES OF λλ6614 TOWARDS HD 179406 AND HD 147889

4.1 Profile comparison

The high resolving power (RP ∼ 115,000) of the HARPS instrument means that the substructure and shapes of the individual components can be seen in some detail as shown in the upper panel of fig 1. In this figure the two bands are constrained to have a common normalised integrated intensity (equivalent width) W. This reveals that the band

1 In the molecular interpretation a diffuse band comprises many unresolved rotational lines each with their own quantum mechanical line strength, and as discussed here, can also include hot bands – each with their own Franck-Condon factor. It follows that when comparing spectra arising from carriers with different levels of internal excitation the diffuse band equivalent widths are
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The basis for our interpretation is that the main triplet structure arises from R, Q and P rotational branches of the origin band of a molecular electronic transition (Sarre et al. 1995). Considering the overall profiles and starting from an onset at c. 6612.9 Å, there is a sharp increase in absorption which suggests the presence of a rotational (R) branch head. The second and always strongest feature (Q) has a less steep long-wavelength side towards HD 147889. For HD 179406 a third possible (P) shoulder near $\sim 6613.8$ Å is quite weak; this is a common characteristic of partially resolved red-degraded molecular rotational profiles. However, for HD 147889 this third feature is quite narrow and strong, particularly when comparing its intensity relative to the R and Q features towards HD 179406; the relative strengths of the first (R) and third ('P') peaks are in fact reversed between these two lines-of-sight. Finally, the redward extension of $\lambda 6614$ for HD 147889 is significantly stronger than for HD 179406 and a fourth narrow absorption feature at $6614.2$ Å is far more prominent towards HD 147889. Other examples of $\lambda 6614$ profiles are given in Galazutdinov et al. (2002).

The difference between the spectra towards HD 179406 and HD 147889 is less deep but is broader than that recorded towards HD 179406. Towards HD 147889 which results in an R-branch which extends to a slightly shorter wavelength than for HD 179406; the second difference is largely accounted for through addition of a second weaker hot band, the result being shown in the lowest panel (fig. 2 (d)). The hot band shifts reflect differences in the spacings of the low-frequency intervals in the ground and excited electronic states. Inclusion of further weak hot band contributions could achieve a yet better fit to the tail in a technical sense, but there is no certainty that only one vibrational mode contributes, and the co-addition procedure implicitly assumes that the rotational constants and rotational populations – and therefore rotational profiles, do not change either between vibronic level or from those for HD 179406. In Section 4.2 we show that there is likely a small contribution to the profile from a hot band even for HD 179406. Notwithstanding these limitations, we find the fact that simple co-addition of shifted copies of the HD 179406 profile accounts for most of the profile towards HD 147889 as in fig. 2 to be supportive of the proposed hot band model.

An excellent laboratory example of the behaviour of low-frequency hot bands in electronic spectra is seen in a jet-cold spectrum of the cyclic $C_3$ molecule, where evidence for additional vibrational bands both overlapping and to the redward side of the origin band was found by varying the sample temperature (Maier et al. 2009). Analogous astrophysical examples include radiation-pumped population transfer to excited vibrational and rotational levels of H$_2$, and observed non-thermal population distributions for high rotational levels of $C_2$ and $C_3$ which are found to vary with line-of-sight. However, not all reddened lines-of-sight with diffuse band features including $\lambda 6614$ have detectable $C_2$ or $C_3$.
5 CONTOUR MODELLING INCLUDING HOT BANDS

5.1 Background

Rotational contour modelling of diffuse band profiles was first conducted by Danks & Lambert (1974) and has become a popular way in which to seek to narrow the range of possible diffuse band carriers. It was first applied to λ6614 for μ Sgr (Kerr et al. 1996) and further applications to this diffuse band involve cumulenes (Schulz et al. 2000), contaminated H2 clusters (Bernstein et al. 2013), small linear molecules (Oka et al. 2013) and accidentally overlapping diffuse bands including a prolate top (Bernstein et al. 2013).

5.2 Contour modelling

While the co-addition approach of section 4.2 is a useful first step in exploring the impact of vibrational hot bands on diffuse band profiles, it is preferable to model the profiles of A6614 towards HD 179406 and HD 147889 with molecular rotational contours. It was already demonstrated that a planar oblate symmetric molecule could provide a good match to the profile towards μ Sgr (Kerr et al. 1996) and we make use of this here, but now including vibrational hot band structure. This was undertaken initially for HD 179406 as it appears to be the simplest and most likely the coolest spectrum. The model adopted is that reported by Kerr et al. (1996) for a vibronically allowed 1A2 → 1X1A transition of a planar oblate symmetric top about the size of the coronene molecule. The same molecular parameters – B" = 0.00947 cm\(^{-1}\), ΔB = -0.42% and Coriolis parameter ζ = -0.43 were employed and constrained to these values, while the rotational temperature and line width were allowed to vary in a PGOPHER fit (Western 2014). The fitting result for a single vibronic band for HD 179406 is shown in fig. 3(a) where the parameters deduced are: Gaussian line width FWHM = 0.43(1) cm\(^{-1}\), T (band origin, air) = 15.120.45(1) cm\(^{-1}\) and T\(_{rot}\) = 18.5(3) K. The FWHM line width of 0.43 (1) cm\(^{-1}\) corresponds to a velocity dispersion of 8.5 kms\(^{-1}\) which is slightly higher than the value of 5.65 kms\(^{-1}\) reported for CH by Kázmiernczak et al. (2009). Inclusion of a very weak hot band results in a slight improvement to the fit (fig 3(b)) and suggests that there is a small contribution from a hot band even for HD 179406.

The molecular constants employed in the fitting of the HD 179406 data were constrained to these values and then used in fitting the spectrum recorded towards HD 147889; a very small shift in the origin of +0.08 cm\(^{-1}\) (1.6 kms\(^{-1}\)) was needed but falls well within the FWHM velocity profile of the (reference) CH spectrum with FWHM of 5.65 kms\(^{-1}\) for HD 147889 (Kázmiernczak et al. 2009). Unsurprisingly, given the co-addition results for HD 179406 shown in fig. 2, hot band contributions are essential and the result of PGOPHER fitting is shown in fig. 4(a) (upper panel); the parameters floated were the rotational temperature, the line width and the weighting of two hot bands. The rotational temperature is found to be slightly higher than for HD 179406 with a value of 22.0(2) K. The longer wavelength region is generally well accounted for and while the overall resultant fit is not quite as good on the long-wavelength side as that attained by simple co-addition of copies of HD 179406 (fig. 2) described in Section 4.2 this probably reflects deficiencies in the molecular rotational contour model. These range from the choice of molecular geometry and electronic state symmetries to the non-inclusion of centrifugal distortion effects and use of a single rotational temperature across all J and K values. However, the major change in the relative strengths of the three main ‘triplet R, Q and P’ absorption peaks between HD 179406 and HD 147889 is well accounted for. The relative weights of the three vibrational bands (origin, first and second hot band) for HD 147889 were found to be 0.75 : 0.20 : 0.05 which is in accord with an exponential Boltzmann fall-off law for thermally populated equally spaced non-degenerate vibrational levels. The line width determined for HD 147889 is FWHM = 0.43(1) cm\(^{-1}\).

While an exponential fall-off in the relative weight of the hot bands is an obvious starting point, the situation is complicated by the fact that (i) the level populations are not necessarily thermal and are probably determined by a blend of collisional and radiative excitation processes, (ii) the excited levels may in fact be electronic (e.g. Jahn-
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Figure 3. Upper panel (a) – $\lambda$6614 towards HD 179406 compared with result of PGOPHER modelling with no hot band included. Lower panel (b) - as for (a) but with one hot band included with a redward shift of -0.66 cm$^{-1}$ and relative weight of 0.07.

Figure 4. Upper panel (a) – $\lambda$6614 towards HD 147889 compared with result of PGOPHER modelling with two hot bands included (redward shifts of -0.66 cm$^{-1}$ and -1.54 cm$^{-1}$) and a Boltzmann fall-off in hot band weightings of 0.75:0.20:0.05. Lower panel (b) – result of fitting with floated weights for three hot bands with shifts of -0.66 cm$^{-1}$, -1.54 cm$^{-1}$ and -2.64 cm$^{-1}$ with weights of 0.66:0.14:0.10:0.10 (see text).

A non-thermal population distribution among vibrational (and rotational) levels might be expected for a non-polar planar oblate top molecule given data from rotationally resolved electronic absorption spectra of non-polar molecules such as C$_2$ and C$_3$ along the same lines-of-sight. Rotational populations for C$_2$ ($X^1\Sigma^+_g^+$) are available for both HD 179406 and HD 147889 [Każmierczak et al. 2010] and are presented in fig 5. In neither case is the plot linear which would hold for thermal population of rotational levels, and high rotational levels in particular are significantly populated for HD 147889. Comparisons of diffuse band profile characteristics with these populations for C$_2$ has to be treated with some caution as some lines-of-sight such as the $\sigma$ line-of-sight HD 147165 ($\sigma$ Sco) have readily detectable $\lambda$6614 absorption but C$_2$ falls below the detection limit. Nevertheless, to the extent that the $\lambda$6614 carrier and the C$_2$ molecule share some common spatial regions and conditions, it would appear that the 'hotter' profile of $\lambda$6614 towards HD 147889, as inferred in this paper, correlates with population of the higher J levels in C$_2$. Given that the higher-J tail for C$_2$ is normally attributed to radiative pumping, this suggests that the population of excited vibronic levels of the ground state of the $\lambda$6614 carrier towards HD 147889 originates in the same way.

Figure 5. Rotational level populations of the C$_2$ molecule towards HD 179406 (triangles, red) and HD 147889 (squares, blue) [Każmierczak et al. 2010] illustrating non-linear non-thermal distributions with an extended high-J tail for HD 147889.
6 SUMMARY AND CONCLUSIONS

We have conducted an investigation into the origin of the λ6614 fine structure, its overall band shape and its change between the lines-of-sight, HD 179406 and HD 147889. The interpretation offered is founded in terms of the population distribution among the rotational, vibrational and potentially low-lying electronic states of a medium-sized (Nc ~ 20) planar PAH-type molecule. The λ6614 carrier in the line-of-sight towards HD 147889 is found to have significantly higher internal excitation which accounts for its greater width and additional structure. We consider that this may be a general feature of diffuse band profiles, but one that clearly depends on the presence, or otherwise, of suitable low-lying vibrational (or vibronic) states. Moreover, the extent to which hot bands are evident in diffuse band spectra will depend on whether these transitions are significantly wavelength-shifted or simply overlap the origin band. However, in cases where no low-lying (populated) vibronic levels exist, slightly altered absorption profiles may still occur due to changes in rotational contours with temperature.

This interpretation differs from others recently put forward which consider the λ6614 profile in terms of a single rotational contour (Oka et al. 2013) or invoke accidentally overlapping diffuse bands (Bernstein et al. 2013). In the study of Herschel 36 by Oka et al. (2013) an extraordinary variety of diffuse band profiles were found which were grouped broadly in terms of the polarity of the band carrier, those with pronounced red tails including λ6614, λ5797 and λ5780 being attributed to polar molecules, with high rotational levels being populated due to irradiation by the nearby IR source Her 36 SE. In the interpretation presented in this paper the extended red tails would arise principally from vibrational hot bands that are red-shifted with respect to the origin band, and bands without extended red tails would be examples where the carrier does not possess low-lying vibronic states and/or any hot bands present are not significantly wavelength-shifted from the origin band.

One aspect arising from these considerations is that the measured equivalent widths of a diffuse band such as λ6614 along different lines-of-sight do not necessarily give the exact column densities of the carrier. This arises because transfer of population between levels which have different intrinsic transition strengths (e.g. Franck-Condon factors for excitation from the lowest compared with excited vibrational levels) can result in weaker (or stronger) absorption for the same carrier column density. In this context it is of interest to investigate whether this might play a role in the close, but not perfect, correlation of λ6614 and λ6196 (Moutou et al. 1992, Galazutdinov et al. 2002, McCall et al. 2010), the results of which will be reported elsewhere.

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