On the Interpretation of the Optical Spectra of L-type Dwarfs

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ABSTRACT. We present synthetic optical spectra in the red and far-red (640–930 nm) of a sample of field L dwarfs suitably selected to cover this new spectral class, and the brown dwarf GL 229B. We have used the recent “dusty” atmospheres by Tsuji (2000) and by Allard (1999), and a synthesis code (Pavlenko et al. 1995) working under LTE conditions which considers the chemical equilibrium of more than 100 molecular species and the detailed opacities for the most relevant bands. Our computations show that the alkali elements Li, Na, K, Rb, and Cs govern the optical spectra of the objects in our sample, with Na and K contributing significantly to block the optical emergent radiation. Molecular absorption bands of oxides (TiO and VO) and hydrides (CrH, FeH and CaH) also dominate at these wavelengths in the early L-types showing a strength that progressively decreases for later types. We find that the densities of these molecules in the atmospheres of our objects are considerably smaller by larger factors than those predicted by chemical equilibrium considerations. This is consistent with Ti and V atoms being depleted into grains of dust.

In order to reproduce the overall shape of the optical spectra of our observations an additional opacity is required to be implemented in the computations. We have modelled it with a simple law of the form \( a_0 \left( \nu/\nu_0 \right)^N \), with \( N = 4 \), and found that this provides a sufficiently good fit to the data. This additional opacity could be due to molecular/dust absorption or to dust scattering. We remark that the equivalent widths and intensities of the alkali lines are highly affected by this opacity. In particular, the lithium resonance line at 670.8 nm, which is widely used as a substellarity discriminator, is more affected by the additional opacity than by the natural depletion of neutral lithium atoms into molecular species. Our theoretical spectra display a rather strong resonance feature even at very cool effective temperatures (\( \sim 1000 \) K); depending on the effective temperature and on the amount of dust in the atmospheres of very cool dwarfs, it might be possible to achieve the detection of lithium even at temperatures this cool. Changes in the physical conditions governing dust formation in L-type objects will cause variability of the alkali lines, particularly of the shorter wavelength lines.

Key words: Stars: atmospheres; fundamental parameters; low-mass, brown dwarfs; individual: BRI 0021–0214, Gl 229B, Kelu 1, DenisP J1228–1547, DenisP J0205–1159

1. Introduction

The optical spectra of the recently discovered very cool dwarfs present new challenges to theoretical interpretation. Their spectral characteristics are drastically different from those of the well known M-dwarfs and this has prompted the use of a new spectral classification, the so-called L-dwarfs (Martín et al. 1997a, 1999; Kirkpatrick et al. 1999a). In principle, the study of the optical spectral energy distribution may allow a better understanding of their physical properties, effective temperatures, gravities, atmospheric composition, etc. The main molecular absorbers at optical wavelengths in early- to mid-M dwarfs are significantly depleted at the lower temperatures present in L dwarfs because of the incorporation of their atoms into dust grains. This process should start in the latest M-dwarf atmospheres (Lunine et al. 1989; Tsuji, Ohnaka & Aoki 1996; Tsuji et al. 1996; Jones & Tsuji 1997; Allard et al. 1997), considerably reducing the strength of TiO, VO and other molecular bands, and producing significant changes in the overall properties of
the optical spectra. Naturally, the appearance of dust will modify the temperature structure of the atmosphere, significantly affecting the formation of the emerging spectrum (Allard et al. 1997).

In this paper we follow a semi-empirical approach to understand the relevance of different processes on the resulting spectral energy distributions in the optical for L dwarfs and Gl 229B. We have obtained far-red optical spectra of several of these cool dwarfs and compared them to synthetic spectra generated using the latest models by Tsuji (2000) and Allard (1999). These models have been successfully used in the interpretation of the near-infrared spectra of these objects (Tsuji, Ohsaka & Aoki 1993; Kirkpatrick et al. 1999). We have taken into account the depletion of some relevant molecules associated with the formation of dust and we have investigated the effects of dust scattering and/or absorption on the formation of the optical spectra. Remarkably strong alkali lines are present in the spectra and dominate its shape in the 600–900 nm region, providing major constraints to the theoretical modelling. We present the observations in section 2, models and synthetic spectra in section 3, and the role of alkalis, molecular bands and dust scattering and/or absorption is considered in section 4. In section 5 we discuss the implications of our study on the determination of effective temperatures and gravities as well as the formation of the lithium resonance line which is a key discriminator of substellar nature in these objects. Finally, conclusions are presented in Section 6.

### 2. Observations and data reduction

We have collected intermediate-resolution spectra in the 640–930 nm range for Kelu 1 (Ruz, Leggett & Allard 1997), Denis-P J1228–1547 and Denis-P J0205–1159 (Delfosse et al. 1997) using the 4.2 m William Herschel Telescope (WHT; Observatorio del Roque de los Muchachos, La Palma). The coolest L dwarf in our sample, Denis-P J0205–1159, has also been observed with the KeckII telescope (Mauna Kea Observatory, Hawaii). Table 1 summarizes the log of the observations. The instrumentation used was the ISIS double-arm spectrograph at the WHT (only the red arm) with the grating R158R and a TEK (1024×1024pix²) CCD detector, and the Low Resolution Imaging Spectrograph (LRIS, Oke et al. 1995) with the 600 groove mm⁻¹ grating and the TEK (1024×1024pix²) CCD detector at the KeckII telescope. The nominal dispersion and the wavelength coverage provided by each instrumental setup were similar and are listed in Table 1. Slit projections were typically 2–3 pix giving spectral resolutions of 6–8 Å. Spectra were reduced by a standard procedure using IRAF, which included debiasing, flat-fielding, optimal extraction, and wavelength calibration using the sky lines appearing in each individual spectrum (Osterbrock et al. 1999). Finally, the observed WHT spectra were corrected from instrumental response making use of the spectrophotometric standard stars BD +26°2606 and HD 19445 which have absolute flux data available in the IRAF environment. No flux standards were observed at the KeckII telescope, and thus the instrumental signature of the KeckII spectrum of Denis-P J0205–1159 was removed by matching it with the WHT spectrum. Our observations are displayed in Fig. 1 together with spectra of the other two objects in our sample: BR10021–0214 (M9.5, Irwin, McMahon, & Reid 1991), and Gl 229B (a brown dwarf with methane absorptions, Nakajima et al. 1993; Oppenheimer et al. 1993). Our sample covers a wide range of spectral types, from the transition objects between M and L types down to the latest types.

### 3. Model atmospheres, spectral synthesis code, chemical equilibrium and opacities

We carried out the computations using the LTE spectral synthesis program WITA5, which is a modified version of the program used by Pavlenko et al. (1995) for the study of the formation of lithium lines in cool dwarfs. The modifications were aimed to incorporate “dusty effects” which can affect the chemical equilibrium and radiative transfer processes in very cool atmospheres. We have used the set of Tsuji’s (2000) “dusty” (C-type) LTE model atmospheres. These models were computed for the case of segregation of dust–gas phases, i.e. for conditions $r_{dust} > r_{crit}$, where $r_{dust}$ is the size of dust particles and $r_{crit}$ is critical...
size corresponding to the gas–dust detailed equilibrium state. In a previous study (Pavlenko, Zapatero Osorio & Rebolo 2000) we had used Tsuji’s (2000) B-type models which are computed for the case of $r_{\text{dust}} = r_{\text{crit}}$. In this paper we have also used a grid of the NextGen “dusty” model atmospheres computed recently by Allard (1999). The temperature-pressure stratification of Allard’s models lie between those of the C-type and B-type models of Tsuji (2000) as it can be seen in Fig. 2.

Chemical equilibrium was computed for the mix of $\approx 100$ molecular species. Alkali-contained species formation processes were considered in detail because of the important role of the neutral alkali atoms in the formation of the spectra. Constants for chemical equilibrium computations were taken mainly from Tsuji (1973).

In the high pressure conditions of the atmospheres of L-dwarfs some molecules can be oversaturated (Tsuji et al. 1996a); in this case such molecules should undergo condensation. To take into account this effect, we reduced the abundances of those molecular species down to the equilibrium values (Pavlenko 1998). The constants for computations of saturation densities were taken from Gurwitz et al. (1982).

We used the set of continuum opacity sources listed in Table 2 where we also give the original sources for the opacity computation codes. That opacity grid allows us to obtain reasonable fits to a variety of stars (see e.g. Martín et al. 1997b; Israelian, García López & Rebolo 1998; Yakovleva & Pavlenko 1998). Opacities due to molecular band absorption were treated using the Just Overlapping Line Approximation (JOLA). Synthetic spectra of late M-dwarfs using both the continuum opacities listed in Table 2 and the JOLA treatment for molecular band absorptions have been already discussed in Pavlenko (1997).

The alkali line data were taken from the VALD database (Piskunov et al. 1995; see their Table 3). At the low temperatures of our objects we deal with saturated absorption lines of alkalis. Their profiles are pressure broadened. At every depth in the model atmosphere the profile of the absorption lines is described by a Voigt func-

**Fig. 1.** Far-red optical spectra for our sample. New observations are those of Kelu 1, Denis-P J1228–1547 and Denis-P J0205–1159 (KeckII spectrum). Data for BRI 0021-0214 and Gl 229B have been collected from Martín, Rebolo & Zapatero Osorio (1996) and from Schultz et al. (1998), respectively. Spectral types for the L-dwarfs are given following the classification of Martín et al. (1999). Spectra in the left panel have been shifted by 0.8 units for clarity. Identification of some atomic and molecular features is provided in the top.
Table 2. List of the opacity sources used in our computations

| Opacity source                                      | Subprogram | Author          |
|-----------------------------------------------------|------------|-----------------|
| Bound-free absorption of H                          | HOP        | Kurucz (1993)   |
| $\text{H}_2$ absorption                             | H2PLOP     | Kurucz (1993)   |
| $\text{H}^+$ ion absorption                         | HMINOP     | Kurucz (1993)   |
| Rayleigh scattering by H atoms                      | HRAYOP     | Kurucz (1993)   |
| He I bound-free absorption                          | HE1OP      | Kurucz (1993)   |
| He II bound-free absorption                         | HE2OP      | Kurucz (1993)   |
| He$^-$ absorption                                   | HEMIOP     | Kurucz (1993)   |
| Rayleigh scattering by He                           | HERAOOP    | Kurucz (1993)   |
| Bound-free absorption of Mg I, Al I, Si I, Fe I, OH and CH | COOLOP     | Kurucz (1993)   |
| Bound-free absorption of N I, O I, Mg II, Si II, Ca I | LUKEOP     | Kurucz (1993)   |
| Bound-free absorption of C II + N II + O II         | HOTOP      | Kurucz (1993)   |
| Electron Thompson scattering by $e^-$                | ELECOP     | Kurucz (1993)   |
| Rayleigh scattering by $\text{H}_2$                  | H2RAOP     | Kurucz (1993)   |
| Absorption due to Hydrogen lines                    | HLINOP     | Kurucz (1993)   |
| $\text{H}_2$ quasimolecule absorption               | QUASIH     | Pavlenko (1999a) |
| C$^-$ ion absorption                                | CMINUSOP   | Pavlenko (1999a) |
| Bound-free of C I                                   | C1OPAC     | Pavlenko (1999a) |

3.1. Molecular opacities

Detail molecular opacities have been computed for all the molecules listed in Table 2 of Pavlenko et al. (1995) as well as for CrH and CaH. To compute the opacity due to absorption of VO and TiO bands we followed the scheme presented in Pavlenko et al. (1993) and Pavlenko (1997). However, for the $B^4\Pi(v) - X^4\Sigma^-$ band system of VO we used a more complete matrix of Franc-Condon Factors computed by the FRANK program (Cymbal 1977) with account of rotational-vibrational interaction in the Morse-Pekeris approximation modified by Schumaker (1969, see Pavlenko 1999b for more details). Furthermore for this band system we used oscillator strength $f_e$ from Allard & Hauschildt (1995). For the $\epsilon$ band system of TiO the parameters of Schwenke (1993) were used.

The CrH molecular band opacity was also considered, the data required for the computations of the elec-

Table 3. List of atomic lines used in our computations

| $\lambda_{\text{vac}}$ (nm) | $gf$  | $E''$ (eV) |
|-----------------------------|-------|------------|
| Li I 610.5164               | 0.126 E+01 | 1.840      |
| Li I 610.5275               | 0.229 E+01 | 1.850      |
| Li I 610.5290               | 0.257 E+00 | 1.850      |
| Li I 670.9551               | 9.790 E-01 | 0.000      |
| Li I 670.9701               | 4.900 E-01 | 0.000      |
| Li I 812.8408               | 0.216 E+00 | 1.850      |
| Li I 812.8629               | 0.432 E+00 | 1.850      |
| Na I 589.1518               | 1.288 E+00 | 0.000      |
| Na I 589.7489               | 6.457 E-01 | 2.105      |
| Na I 818.5443               | 1.690 E+00 | 2.100      |
| K I 766.6961                | 1.340 E+00 | 0.000      |
| K I 770.1031                | 6.760 E-01 | 2.105      |
| Rb I 780.2348               | 1.370 E+00 | 0.000      |
| Rb I 794.9729               | 6.800 E-01 | 0.000      |
| Cs I 894.5900               | 7.800 E-01 | 0.000      |
| Cs I 852.3285               | 1.620 E+00 | 0.000      |
tronic transition $A^6\Sigma^{(+)}-X^6\Sigma^{(+)}$ being taken from Huber & Herzberg (1979). Franc-Condon factors were computed by the FRANK program (Cymbal 1977). For this band system we used $f_e = 0.001$, determined by Pavlenko (1999b). Molecular bands of the $A^6\Sigma^{(+)}-X^6\Sigma^{(+)}$ of CrH lie in the wide wavelength region 500–1200 nm, and a head of the strong (0,1) band lies near the core of the K i atomic line. A head of the (0,0) band of $A^6\Sigma^{(+)}-X^6\Sigma^{(+)}$ system of CrH at 860 nm is blended with the heads of (1,0), (3,2), (2,1) bands of the $B^4\Pi(v)-X^4\Sigma^-$ system of VO. For the spectral region 650–710 nm we took into account the absorption due to the $B^2\Sigma-X^2\Sigma$ band system of CaH for which we adopted $f_e = 0.05$. Frank-Condon factors were computed with the FRANK program. Although FeH is an important absorber at wavelengths around 870 nm and 990 nm for the coolest dwarfs, it is not yet included in our calculations due to the lack of appropriate laboratory data. In the wavelength region presented in this paper, the blue band of FeH is blended with CrH and VO.

4. Analysis and interpretation

4.1. Atomic features

Our spectral synthesis for model atmospheres in the temperature range 2200–1000 K confirms the relevant role of atomic features due to the Na i and K i resonance doublets in the spectral range 640–930 nm. The strength of these doublets increases dramatically when decreasing the effective temperature ($T_{\text{eff}}$). In Fig. 3 we display synthetic spectra for different $T_{\text{eff}}$ and gravity values. These computations do not include any molecular or grain opacity in order to show how alkali absorptions change with these parameters. In addition, we can see in the figure how the overall shape of the coolest L-dwarf spectra is governed by the resonance absorptions of Na i and K i. Even the less abundant alkali (i.e. Li, Rb, Cs) produce lines of remarkable strength. Note the increase in intensity of K i and Na i lines with decreasing $T_{\text{eff}}$ and with increasing atmospheric gravity. Although this different behaviour can, in principle, make it difficult to disentangle these parameters from the optical spectra, simple physical considerations give an upper limit to gravity of log $g = 5.5$ for brown dwarfs with lithium and therefore the large broadening of the K i lines seen in some of our objects cannot be attributed to higher aphysical values of gravity.

Our computations provide a qualitative explanation of the far-red spectral energy distributions presented in Fig. 1. The equivalent widths (EWs) of the K i and Na i resonance doublets may reach several thousand Angstroms, becoming the strongest lines so far seen in the spectra of astronomical objects. This is mainly caused by the high pressure broadening in the atmospheres of the coolest dwarfs, where damping constants of the absorption lines vary from 0.001 in the uppermost layer of the atmosphere to 2–4 in the deepest regions. The subordinate lines of Na i at 819.5 nm, clearly seen in all the L-dwarfs in our sample, become weaker as $T_{\text{eff}}$ decreases. These computations also show that the subordinate Li i line at 812.6 nm may be detected in early/mid L-dwarfs with equivalent widths not exceeding 1 Å, and that the triplet at 610.3 nm appears completely embedded by the wings of the K i and Na i resonance lines.
4.2. Molecular features

In the optical spectra of very cool dwarfs one expects the presence of bands of VO, TiO and indeed, our synthetic spectra show that these bands play an important role in the cases of BRI0021−0214 and Kelu1. In Fig. 4 (upper panel) we compare the observed spectrum of Kelu1 with a synthetic spectrum obtained using the Tsuji C-type model for $T_{\text{eff}} = 2000\, \text{K}$ and $\log g = 5$. The natural depletion of molecules resulting from the chemical equilibrium is not sufficient to get a reasonable fit to the data. The discrepancies with respect to the observed spectrum can be notably reduced (Fig. 4, lower panel) if we impose a depletion of CaH, CrH, TiO and VO molecules which should account for the condensation of Ca, Cr, Ti and V atoms into dust grains. Since we do not have an appropriate theoretical description of the processes of dust formation at present, we use the simple approach of modifying the chemical equilibrium for all molecules. We implemented this “extra” depletion simply by introducing a fac-
Fig. 6. Kelu 1’s observed spectrum (full line) compared to the best fits (dash-dotted line) obtained using Tsuji’s (2000) C-type models (upper panel, $T_{\text{eff}} = 2000$ K and $\log g = 5$), and Allard’s (1999) dusty models (lower panel, $T_{\text{eff}} = 1800$ K and $\log g = 5$). Both predicted spectra with a resolution of 10 Å have been computed considering the \(R\) depletion factors given in Fig. 4 and the AdO law described in the text with \(a_0 = 0.03\).

Fig. 7. DenisP J1228–1547 and DenisP J0205–1159’s observed data (full line) compared to the best fits (dash-dotted line) obtained using Tsuji’s (2000) C-type models (upper panel: $T_{\text{eff}} = 1600$ K and $\log g = 5$; lower panel: $T_{\text{eff}} = 1200$ K and $\log g = 5$). Both predicted spectra with a resolution of 10 Å have been computed considering the \(R\) depletion factors given in Fig. 5 and the AdO law described in the text with \(a_0 = 0.006\).

4.3. The need for additional opacity (AdO)

Our first attempts to model the spectra including atomic and molecular features showed in Figs. 4 and 5 were only modestly successful. Although we could reproduce reasonably well the red wing of the K I line, the theoretical fluxes in the blue part of our synthetic spectra (640–750 nm) are too large. In fact, we cannot fit the observations by taking into account only the opacity provided by the Na I and K I resonance doublets and the continuum opacity sources listed in Table 2.
for a range of $T_{\text{eff}}$, gravities, $a_0$ and depletion factors $R$ for TiO, VO, CaH and CrH. The previous figures display those syntheses which better reproduce the observed spectra. From our computations we infer that VO is less efficiently depleted than TiO at a given $T_{\text{eff}}$. The depletion of these oxides appears to increase very rapidly as we go from BRI 0021–0214 and Kelu 1 to the lower temperature objects DenisP J0205–1159 and DenisP J0205–1159. Note the good fit to the observed spectra at 860 nm provided by the (0,0) band of the $\Delta^{0}\Sigma^{(\pm)}\rightarrow\chi^{0}\Sigma^{(\pm)}$ system of CrH. From the study of this band we also find that the depletion factor of CrH increases from the warmer to the coolest objects in the sample. Finally, we also find that atomic lines become weaker and narrower when increasing the amount of AdO in the atmospheres.

We have also studied whether we can explain the optical spectrum of Gl 229B using the following Tsuji’s model: C-type, $T_{\text{eff}} = 1000$ K, and $\log g = 5.0$. and the AdO law of index $N = 4$ used above. In Fig. 8, we show several spectral synthesis reproducing the optical spectrum of this object, and showing the effect of different $a_o$ parameters which is related to the amount of dust in the atmosphere. In Gl 229B we need the highest value of $a_o$, which is interpreted as evidence for the most “dusty” atmosphere in our sample. We have not attempted to reproduce the CsI lines in Gl 229B. According to our hypothesis, the inclusion of AdO avoids the contribution of high pressure regions to the formation of these lines. The shorter wavelength CsI line at 894.3 nm is very affected by dust opacity.

5. Discussion

5.1. $T_{\text{eff}}$ for L-type dwarfs

Our computations provide a reasonable description of the far-red optical spectra of L-dwarfs and provide a physical basis for a progressively decreasing $T_{\text{eff}}$ for the proposed spectral classifications (Martín et al. 1996; Kirkpatrick et al. 1999d). Effective temperatures for a few field L-dwarfs and for Gl 229B have been derived from spectra at IR wavelengths (Allard et al. 1996; Marley et al. 1996; Matthews et al. 1994; Tsuji et al. 1993; Jones et al. 1996; Kirkpatrick et al. 1999b). Here we study to what extent we can use the broad energy spectral distribution in the optical to infer the $T_{\text{eff}}$ for the cool dwarfs in our sample. Our best estimates using Tsuji’s models (see Table 4) are in good agreement with those found from IR data for objects of similar spectral types and for Gl 229B. Using Allard’s models and the simple approach of AdO described in section 4.3, we are also able to reproduce the observed spectra, albeit we require lower $T_{\text{eff}}$’s by up to several hundred degrees for the coolest L-dwarfs. This is mainly due to the hotter stratification of Allard’s model atmospheres for the potassium and sodium lines forming layers (see Fig. 4). Opposite to IR-based temperature determinations, the estimation from optical spectra is very
Table 4. \( T_{\text{eff}} \) estimations for our sample adopting Tsuji’s (2000) C-type models and \( \log g = 5.0 \).

| Object            | Sp. Type | \( I - J \) | \( T_{\text{eff}} \) (±200 K) | Other measures | Source          |
|-------------------|----------|-------------|-------------------------------|----------------|-----------------|
| BRI 0021–0214     | M9.5     | 3.30        | 2200                          | 1980, 2300     | TMR93, LAH98    |
| Kelu 1            | L2 (L2)  | 3.50        | 2000                          | 1900, 1900     | B99, RLA97      |
| DenisP J1228–1547 | L4.5 (L5)| 3.81        | 1600                          | 1800           | B99             |
| DenisP J0205–1159 | L5 (L7)  | 3.82        | 1200                          | 1700, 1800     | B99, TK99       |

NOTES. Spectral types are given in Martín et al. (1999). Those spectral types in brackets come from Kirkpatrick et al. (1999b). \( (I - J) \) colors have been taken from Leggett, Allard & Hauschildt (1998). References: TMR93 = Tinney, Mould & Reid (1993); LAH98 = Leggett et al. (1998); B99 = Basri et al. (1999); RLA97 = Ruiz et al. (1997); TK99 = Tokunaga & Kobayashi (1999).

Sensitive to the input physics parameters, like dust formation, molecular equilibrium, sources of opacity and atmospheric models, which limits the accuracy of the estimates to \( \sim 200 \text{K} \). On the other hand, the optical spectra provide an opportunity to test the reliability of the physical description of the atmospheres.

Cs\( _1 \) lines in the optical spectra have been recently used to infer the \( T_{\text{eff}} \) of some L-dwarfs (Basri et al. 1999). In spite of the sensitivity of these lines to many input parameters in the models (i.e. to the amount of opacity, chemical equilibrium, etc.) we find in general a good agreement with their estimated temperatures for the earlier L-dwarfs (see Table 4). However, for the latest L-type objects in our sample, DenisP J1228–1547 and DenisP J0205–1159, we find temperatures up to 400 K lower. We may attribute this discrepancy to the effects that dust opacity has in the formation of optical lines of alkali. This produces a kind of “veiling” of the atomic lines, reducing their intensities. Basri et al. (1999) did not consider this effect, and therefore they required hotter models in order to explain the strength of the observed Cs lines. Another possible reason for the discrepancy is that the temperatures in Table 4 have been obtained for gravity \( \log g = 5 \); if we increased gravity by 0.5 dex we would have to increase the temperatures of our models by about 200 K. In this case the spectral synthesis does not reproduce the observations so well but they are still acceptable.

5.2. Alkali lines: the case of lithium

Recently, several new Gl 229B-like objects have been discovered by the SDSS survey (Strauss et al. 1999) and the 2MASS collaboration (Burgasser et al. 1999). Based on their near-IR spectra, these authors suggest that these new cool brown dwarfs may be warmer than Gl 229B. In order to estimate the optical properties of these objects we have computed synthetic spectra using a Tsuji’s C-type model of \( T_{\text{eff}} = 1200 \text{K} \) and \( \log g = 5.0 \), and we have adopted the basic prescriptions that were followed in section 4, i.e total depletion of VO and TiO and the AdO law with \( N = 4 \). In Fig. 9 we plot the resulting spectra considering different amounts of dust opacity. As expected, the overall shape of the spectrum is intermediate between that of DenisP J0205–1159 and Gl 229B. In the absence of dust absorption \( (\alpha_o = 0.0) \) alkali lines are clearly seen (including the lithium resonance doublet), and the spectrum is governed by the sodium and potassium lines. If we consider dust opacities comparable to those in Gl 229B, the alkali lines of Cs and Rb become weaker, but still detectable with intermediate resolution spectroscopy. Remarkably, the subordinate Na\( _i \) doublet at 819.5 nm is very sensitive to the incorporation of dust opacity due to the larger depths of its formation as compared with resonance lines. For very high dust opacities these lines may become undetectable.
Fig. 10. Densities of lithium and the most abundant species containing lithium are plotted for the C-type Tsuji’s (2000) model atmospheres with $T_{\text{eff}} = 2000$ K (upper panel) and 1200 K (lower panel) and $\log g = 5$. Computations have been performed under the assumption of complete chemical equilibrium for a solar elemental mixture.

The effects of additional dust opacity on the formation of Li i lines (resonance and subordinate ones) also deserve detailed consideration since they play a major role as a discriminator of substellar nature for brown dwarf candidates (see Rebolo, Martín, & Magazzù 1992; Magazzù, Martín, & Rebolo 1993). Most of the known brown dwarfs are actually recognized by the detection of the Li i resonance doublet in their spectra (Rebolo et al. 1998; Martín et al. 1997; Rebolo et al. 1998; Tinney 1998; Kirkpatrick et al. 1999). The chemical equilibrium of lithium contained molecules have been considered in all our syntheses (Fig. 10 depicts the density profiles for lithium species for two C-type models by Tsuji 2000). Our computations show that both, the resonance line at 670.8 nm, and the subordinate lines at 601.3 nm and 812.6 nm are very sensitive to the AdO that we need to incorporate in the spectral synthesis if we want to explain the observed broad spectral energy distribution. Among the subordinate lines the doublet at 812.6 nm is more easily detectable, but the predicted EWs, assuming fully preserved lithium, are rather small, ranging from $\text{EW} = 0.4$ Å to 0.04 Å for $T_{\text{eff}}$ values in the range 2000 K down to 1200 K. These EWs are considerably reduced by the inclusion of the AdO described in the previous section, which makes their detection rather difficult. In Table 3 we give the predicted EWs of the Li i resonance doublet at 670.8 nm for several of the coolest model atmospheres (2000–1000 K) considered in this work. First, we note that in the absence of any AdO (second column in the table), we would expect rather strong neutral Li resonance lines in the spectra of objects as cool as DenisP J0205–1159 and Gl 229B. The chemical equilibrium of Li-contained species still allow a sufficient number of Li atoms to produce a rather strong resonance feature; one reason for this is that Cl and O
Table 5. Equivalent widths (Å) of the Li resonance doublet at 670.8 nm computed for the C-type Tsuji’s (2000) model atmospheres, cosmic Li abundance (log \( N(\text{Li}) = 3.2 \)) and gravity \( g = 5.0 \).

| \( T_{\text{eff}} \) (K) | \( a_0 \) | EW (Å) |
|------------------------|----------|--------|
| 1000                   | 0.00     | 0.6    |
| 1200                   | 0.01     | 0.7    |
| 1400                   | 0.10     | 0.9    |
| 1600                   | 0.12     | 1.6    |
| 2000                   | 0.17     | 3.6    |

atoms should be also bounded into other molecules (e.g. NaCl, KCl, H\(_2\)O, etc.). Our computations indicate that objects like DenisP J0205–1159 and cooler objects with moderate dust opacities should show the Li resonance doublet if they had preserved this element from nuclear burning, and consequently the lithium test can still be applied. Furthermore, even in very dusty cool atmospheres like that of Gl229B for which we have inferred a high value of the opacity parameter of \( a_0 = 0.1 \) (fourth column in Table 5), the lithium resonance line could be detected with an EW of several hundred mÅ (high S/N data would be required).

Another effect that we shall consider is whether small changes in the AdO (which could be originated as a consequence of some “meteorological” phenomena occurring in these cool atmospheres) can lead to detectable variations in the EWs of the lithium lines. In particular, weak lithium lines do not necessarily imply a depletion of this element. The observed Li\(_i\) variability in Kelu 1 (with changes in EW by a factor 5), could be an indication of meteorological changes in the atmosphere of this rapidly rotating cool object. In Fig. 11 we present several spectral synthesis showing the sensitivity of the lithium line to the AdO in the atmosphere. The AdO parameters which give the best fit for the lithium line in Kelu 1 (EW = 6.5±1.0 Å) coincide with those also providing the best fit to the whole optical spectrum (see Fig. 10). Anyway, the obtained lithium abundance is consistent with complete preservation of this element.

6. Conclusions

In this paper we have attempted to model the far-red spectra (640–930 nm) of several L-dwarfs suitably selected to cover this new spectral class. We have used model atmospheres from Tsuji’s (2000) and Allard’s (1999), as well as an LTE spectral synthesis code (Pavlenko et al. 1997) which takes into account chemical equilibrium for more than 100 molecular species, and detailed opacities for the most relevant bands. We have arrived to the following conclusions:

1) Alkali lines play a major role governing the far-red spectra of L-dwarfs. At early types, this role is shared with TiO and VO bands, which dominate this spectral region in late M-dwarfs. As we move to later spectral types we need to incorporate progressively higher depletions of these oxides and of the hydrides CrH and CaH, consistently with the expectation that Ti and V atoms are depleted into grains; and we also require additional opacity to reproduce the overall shape of the spectra. This additional opacity could be either due to molecular/dust absorption or to dust scattering.

2) We have shown that a simple law for this additional opacity of the form \( a_0 (\nu/\nu_0)^N \), with \( N = 4 \), gives a sufficiently good fit to the observed spectra of L-dwarfs and Gl229B. For this late object we require the highest value of \( a_0 \) consistent with a very dusty atmosphere. The strength of alkali lines is highly affected by this opacity.

3) From the best fits to our spectra, we derive the most likely \( T_{\text{eff}} \) values for our sample of L-dwarfs. For the warmer objects, our values are consistent with those obtained by other authors, however we find lower \( T_{\text{eff}} \)’s by several hundred degrees for the coolest L-dwarfs. Because the optical spectra are very much affected by the input physics, a more reliable \( T_{\text{eff}} \) scale should be obtained by fitting the IR data of these cool objects.

4) After detailed consideration of chemical equilibrium, we find that the lithium resonance doublet at 670.8 nm can be detected in the whole spectral range (down to 1000K). In the coolest L-dwarfs the strength of the resonance line is more affected by the amount of additional opacity needed to explain the spectra than by the depletion of neutral lithium atoms into molecular species. In those atmospheres where the additional opacity required is low, the lithium test can provide a useful discrimination of substellar nature. Changes in the physical conditions governing dust formation in L-dwarfs, will cause variability of the lithium resonance doublet. Taking into account the need for additional opacity in Kelu 1, we find that the lithium abundance can be as high as log \( N(\text{Li}) = 3.0 \), i.e. consistent with complete preservation.

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