Satellite measurements of the global mesospheric sodium layer

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Abstract

Optimal estimation theory is used to retrieve the absolute Na density profiles in the mesosphere/lower thermosphere from limb-scanning measurements of the Na radiance at 589 nm in the dayglow. Two years of observations (2003 and 2004), recorded by the OSIRIS spectrometer on the Odin satellite, have been analysed to yield the seasonal and latitudinal variation of the Na layer column abundance, peak height, and peak width. The layer shows little seasonal variation at low latitudes, but the winter/summer ratio increases from a factor of $\sim 3$ at mid-latitudes to $\sim 10$ in the polar regions. Comparison of the measurements made at about 06:00 and 18:00 LT shows little diurnal variation in the layer, apart from the equatorial region where, during the equinoxes, there is a two-fold increase in Na density below 94 km between morning and evening. This is most likely caused by the strong downward wind produced by the diurnal tide between $\sim 02:00$ and $10:00$ LT. The dramatic removal of Na below 85 km at latitudes above $50^\circ$ during summer is explained by the uptake of sodium species on the ice surfaces of polar mesospheric clouds, which were simultaneously observed by the Odin satellite.

1 Introduction

Meteoric ablation is the source of the layer of neutral sodium atoms that occurs globally in the upper mesosphere/lower thermosphere (MLT) between 80 and 105 km (Plane, 2003). The Na layer was discovered nearly 80 years ago: radiation at 589 nm was observed in the night sky spectrum (Slipher, 1929), and later identified as emission from neutral sodium atoms within the atmosphere (Bernard, 1939). In the 1950s, the layer was observed from resonance fluorescence at 589 nm (Na$(3^2P_J - 3^2S_{1/2})$) during twilight (Hunten, 1967). The variation of the fluorescence signal as the solar terminator passed through the layer enabled the layer profile to be determined. The absolute Na abundance was estimated by performing radiative transfer calculations on the

Measurements of the Na $D$-line emission in the dayglow were then performed using a Zeeman photometer, which utilized an oscillating magnetic field around a sodium vapour cell to filter out the Na $D$-line spectrum and hence discriminate against Rayleigh-scattered sunlight (Blamont and Donahue, 1961). The Na layer was also studied during the day by ground-based photometric measurements of atomic Na absorption (Chamberlain, 1956), and by rocket-borne photometric measurements of the dayglow (Donahue and Meier, 1967). However, because of the passive mode of photometric measurements, where solar radiation is the only excitation source, the observations were confined to the twilight or dayglow.

The invention of the tunable laser enabled enormous progress to be made with the development of the lidar (light detection and ranging) technique in the late 1960s (Bowman et al., 1969). Lidar provided many advantages over photometers. Firstly, by active excitation of the atmospheric Na, observations could be performed over a full diurnal cycle: initially, night-time measurements were performed (Gibson and Sandford, 1971), but modified receivers to exclude scattered sunlight then enabled daytime measurements to be made (Gibson and Sandford, 1972). These measurements quickly showed that the apparent enhancement of the Na layer during daytime, reported from photometric measurements of the dayglow (Blamont and Donahue, 1961), was not correct. Long-term observations of the Na layer provided more details of seasonal, latitudinal and diurnal variations (e.g. Megie and Blamont, 1977; Clemesha et al., 1979; Kirchhoff and Clemesha, 1983; Gardner et al., 1988; Tilgner and von Zahn, 1988; von Zahn et al., 1988; Plane et al., 1999; She et al., 2000; Clemesha et al., 2004; Gardner et al., 2005).

Of course, an important limitation with lidar measurements is that the instruments are usually ground-based, and thus provide only localized coverage (airborne lidars partially overcome this difficulty (Kane et al., 1991; Plane et al., 1998a), but only for short campaigns). Global coverage can only be provided by space-borne instruments.
In fact, the first satellite-borne observations of the sodium nightglow were made two decades ago (Newman, 1988), although the conclusions of that study were subsequently criticized (Clemesha et al., 1990). The Na nightglow is generated from a chemiluminescent cycle involving Na, O₃ and O (Chapman, 1939), and so the complication of using nightglow observations is that the Na atom density is derived indirectly (Xu et al., 2005). Recently, Fussen et al. reported global measurements of the mesospheric Na layer by the GOMOS stellar occultation instrument (i.e., the Na density was measured by atomic absorption) on the ENVISAT satellite (Fussen et al., 2004). Unfortunately, as we will discuss below, the retrieved Na column abundances, layer density profiles, and the seasonal and latitudinal variations are in poor agreement with the available data from ground-based lidars.

We have recently developed a new method (Gumbel et al., 2007) using optimal estimation theory (Rodgers, 2000) to retrieve the absolute mesospheric Na density profile from limb-scanning satellite observations of the Na D-lines in the dayglow. The method has been validated by comparing Na profiles, retrieved from the OSIRIS spectrometer on the Odin satellite, with profiles measured during overflights of the Na lidar at Ft. Collins, Colorado. The details of the methodology and error characterization are contained in our earlier paper (Gumbel et al., 2007). OSIRIS measures irradiance in the limb at wavelengths between 280 and 800 nm, with a resolution of ~1 nm (Llewellyn et al., 2004). Odin is in a Sun-synchronous orbit at a height of ~600 km, with an ascending node at ~18:00 h local time (LT) and descending node at ~06:00 LT (Murtagh et al., 2002).

In this paper, we have used this method to analyse two years of Odin data (2003 to 2004), in order to obtain the global seasonal distribution of the Na layer. During these two years there was a high frequency of mesospheric measurements in July/August, because during these northern hemisphere summer periods Odin was dedicated to observing polar mesospheric clouds (PMCs) by limb-scanning between 70 and 110 km.
Global Latitudinal and Seasonal Variations

Figure 1 illustrates the global coverage of the retrieved Na total abundance or column density (in units of Na atom cm$^{-2}$), where the retrieved Na density profiles have been integrated from 70 to 120 km. The data is zonally averaged to increase the data available in some latitude bins. In order to avoid artefactual distortions introduced by contouring the data, this map is plotted in a grid format, with each grid box corresponding to 10$^\circ$ in latitude. The number inside each grid box indicates the number of integrated Na density profiles that have been averaged to produce the resulting column density. The blank grid boxes show occasions when mesospheric dayglow measurements could not be made because the solar zenith angle was larger than 92$^\circ$. This occurred in the winter hemisphere at mid- to high latitudes. The blank grid boxes at very high latitudes (>$85^\circ$) occur because of these regions are not covered by the satellite orbits.

As many previous ground measurements have reported, the Na layer exhibits seasonal variations at all latitudes higher than 20$^\circ$: the total Na column density decreases to a minimum near the summer solstice and reaches a maximum in early winter. During mid-summer and the whole winter period, the Na abundance exhibits relatively small fluctuations compared with the much greater variability during the spring and autumn equinoxes. Figure 1 shows that the seasonal variation from summer minimum to winter maximum becomes larger at higher latitudes. Although the data from mid-winter months is absent (except at low latitudes), we can still estimate the seasonal ratio by employing the available data from late autumn/early winter. The data in Fig. 1 is in good accord with ground-based measurements at $\sim$40$^\circ$ N (Megie and Blamont, 1977; Plane et al., 1999; States and Gardner, 1999; She et al., 2000), at 51$^\circ$ N (Gibson and Sandford, 1971), at 23$^\circ$ S (Clemesha et al., 1979), at 43$^\circ$ S (Hunten et al., 1964), and at 90$^\circ$ S (Gardner et al., 2005). The maximum Na layer density usually occurs in about October in the northern hemisphere (NH), and in about April in the southern hemisphere (SH). At low latitudes, this ratio is $\sim$2, and rises to $\sim$3 at middle latitudes. In polar region, this ratio increases up to nearly 10. The minimum monthly averaged Na abundance in
Fig. 1 occurs in July at 80° N, and the maximum is during October at 80° N. In the equatorial region (±10°), by contrast, the seasonal variations are very small: the seasonal transitions between northern and southern hemisphere merge here, with a minimum in about March/April and a maximum in January/February. The seasonal ratio is typically no more than 1.5.

In order to understand the seasonal variation and latitudinal distribution of the Na abundance, we need to consider mesospheric sodium chemistry. In the mesosphere and lower thermosphere (MLT), Na+ ions and sodium bicarbonate (NaHCO₃) are the major reservoir species above and below the atomic Na layer, respectively (Plane et al., 1999; Plane, 2004). Due to the long residence time of ablated sodium in the region between 80 and 100 km (~7 days) (Plane, 2004), and the comparatively short lifetimes of the rate-determining chemical reactions that convert sodium between atomic Na and these reservoirs, the chemistry reaches a photochemical steady-state on the timescale of vertical transport by eddy or molecular diffusion (Plane et al., 1998b). Consequently, atmospheric chemistry plays a key role in determining the atomic Na distribution. Na is converted to NaHCO₃ via the following sequence of reactions (Plane, 2004):

\[ \text{Na} + \text{O}_3 \rightarrow \text{NaO} + \text{O}_2 \]  \hspace{1cm} (1)

\[ \text{NaO} + \text{H}_2\text{O} \rightarrow \text{NaOH} + \text{OH} \]  \hspace{1cm} (2)

\[ \text{NaOH} + \text{CO}_2(+\text{M}) \rightarrow \text{NaHCO}_3(\text{M}=\text{N}_2\text{orO}_2) \]  \hspace{1cm} (3)

All of these reactions are fast, with small temperature dependences. NaHCO₃ is converted back to Na by the reaction:

\[ \text{NaHCO}_3 + \text{H} \rightarrow \text{Na} + \text{H}_2\text{CO}_3 \]  \hspace{1cm} (4)

In contrast, this reaction has a large activation energy (Cox et al., 2001); that is, at higher temperatures it becomes much faster, and the steady-state balance shifts from NaHCO₃ to Na. A secondary effect is that O₃ forms from the recombination of O and O₂, and this reaction gets slower at higher temperatures: the result is that smaller O₃
concentrations slow down the conversion of Na to NaO, and the larger O concentrations reduce NaO back to Na, enhancing the effect of Reaction (4) becoming faster.

At high latitudes, several factors further reinforce the positive temperature dependence of the Na density below 96 km. The first of these is the removal of sodium species on the ice surfaces of PMCs, which form between 82 and 85 km during summer at high latitudes when the temperature decreases below ∼150 K (Gardner et al., 2005; Murray and Plane, 2005; She et al., 2006). These low temperatures are caused by upwelling air (via adiabatic expansion) which also transports H$_2$O from the lower mesosphere to above 80 km (Livesey et al., 2003), thereby increasing the rate at which Na is converted to NaHCO$_3$ (Reaction 2). Conversely, during winter there is strong downward transport by the meridional wind (Portnyagin et al., 2004; Gardner et al., 2005), which has two effects: the temperature increases because of adiabatic heating, and Na is transported downwards from the peak of the layer.

Figure 2 illustrates the latitudinal distribution of the correlation coefficient between the retrieved Na density and temperature (taken from the MSIS model, Picone et al., 2002), as a function of height. At middle and high latitudes, a strong positive correlation between temperature and Na density is seen below 96 km. This correlation is expected in view of the effects of temperature on the neutral sodium chemistry (see above), and has also been seen in mid-latitude lidar data (Plane et al., 1999).

At altitudes above 96 km, the correlation coefficient abruptly turns negative. This is because ion chemistry dominates on the topside of the Na layer. Na$^+$ ions are formed by charge transfer with ambient NO$^+$ and O$_2^+$ ions and, to a lesser extent, by photo-ionization (Plane, 2004). These processes are essentially temperature-independent. In contrast, the neutralization of Na$^+$ begins by forming a cluster ion with N$_2$:

Na$^+$ + N$_2$(+M) → Na$^+$.N$_2$, where M=N$_2$ or O$_2$  

This cluster ion can switch with more stable ligands such as CO$_2$ or H$_2$O, or be reduced back to Na$^+$ by atomic O, which slows the overall process (Cox and Plane, 1998). However, the cluster ions eventually undergo dissociative electron recombination to yield Na. Reaction (5) is an association reaction with a negative temperature dependence.
dependence (Cox and Plane, 1998): that is, at lower temperatures the cluster ions form more rapidly, and so the steady-state balance between $\text{Na}^+$ and Na shifts towards the neutral atom.

At low latitudes ($\pm 20^\circ$), the Na density does not exhibit a consistently strong correlation with temperature either below or above 96 km. In contrast, there is a strong diurnal variation in this region which appears to be driven by the diurnal tide (see below).

Figure 3 illustrates the seasonal variability of the Na layer: the top panel shows the monthly Na profiles, zonally averaged in four latitude bands. The corresponding monthly temperature profiles are shown in the bottom panel. Once again, the positive correlation between Na and temperature below 96 km, and the anti-correlation above this altitude, is clearly visible at latitudes higher than $20^\circ$ N and S. Figure 3a shows the seasonal variation of the Na density in the Arctic ($70^\circ$ N). Note the removal of Na on both the top and bottom of the layer during summer, which is much more pronounced than at lower latitudes. As discussed above, the removal on the bottom side is caused by the stability of the chemical reservoir NaHCO$_3$ at lower temperatures. However, this is greatly amplified by the uptake of sodium species on PMC ice particles (discussed in more detail below). Because of vertical mixing, Na is also removed on the topside of the layer. The seasonal temperature variation becomes smaller at mid- and low latitudes, leading to less removal of Na during summer.

Figure 3(b) shows the Na profile at $40^\circ$ N. This latitude was chosen for direct comparison with the Na layer seasonal dependence at $40^\circ$ N reported recently from stellar occultation measurements made by the GOMOS instrument on ENVISAT (Fig. 5 in Fussen et al., 2004). There are several major discrepancies with the results of the present study. First, the GOMOS retrievals show a second peak in the Na density during mid-summer, whereas the OSIRIS result is a mid-summer minimum, in accord with the lidar observations of the night-time Na layer at Ft. Collins (Colorado) (She et al., 2000) and Urbana (Illinois) (Plane et al., 1999; States and Gardner, 1999). Second, the maximum Na density in the GOMOS retrievals is $\sim 2500$ cm$^{-3}$ in March, compared with just over 5000 cm$^{-3}$ during October/November for OSIRIS and the mid-latitude lidar...
observations. Third, the width of the Na layer retrieved by GOMOS at 40° N is highly variable: it is narrowest in March (extending from 85–105 km), when the Na density at the layer peak is largest, and is much broader (75–110 km) in late summer when the peak density is a minimum. In contrast, Fig. 3b shows that the layer thickness retrieved by OSIRIS at 40° N is fairly constant, with the layer being somewhat broader in early winter when the peak density is a maximum (again in good accord with the lidar observations).

The reasons for these very marked discrepancies between the results from OSIRIS and GOMOS are not clear. Although the GOMOS results are for 2003, and the OSIRIS results combine data from 2003 and 2004, there are no significant differences in the OSIRIS Na layer data between these successive years. The OSIRIS measurements were made at 06:00 or 18:00 LT, whereas most of the GOMOS measurements were at around 09:00 or 23:00 LT. This might indicate that some diurnal effect could be responsible for the differences. Indeed, a limited number of GOMOS measurements were made over the whole range of LT, from which the diurnal behaviour of the Na layer could be examined (Fussen et al., 2004). The data showed very little change in the layer at 06:00, 12:00 and 18:00 LT, and then a large decrease, by ~40% between 70 and 90 km, at 24:00 LT. Strangely, such diurnal behaviour has not been observed by lidars located over a range of latitudes (Gibson and Sandford, 1972; Kirchhoff and Clemesha, 1983; States and Gardner, 1999; She et al., 2000). In conclusion, the GOMOS results are not in accord with either the OSIRIS measurements or ground-based lidars, but it is unclear why this is the case.

Figure 4 illustrates the distribution of the full width at half maximum (FWHM) and peak height of the Na layer, as a function of latitude and month. Due to the limited vertical resolution in the retrieved profiles, these parameters cannot be determined with the precision of a ground-based lidar. Nevertheless, useful results are obtained. Unsurprisingly, the FWHM is a minimum in polar summer, because of removal of Na on both the top and bottom of the layer (see above), whereas in winter the FWHM more than doubles at mid- to high latitudes. The peak height of the Na layer lies mostly
between 90 and 92 km. However, it is often higher (~94 km) in the equatorial region, as a result of the strong diurnal tide (see below).

3 Diurnal variation in the equatorial region

The Odin satellite takes measurements at about 06:00 LT (descending node) and 18:00 LT (ascending node), and the above analyses of seasonal properties of the Na layer combine the morning and evening observations. Indeed, at mid- and high latitudes there is no significant diurnal difference. In contrast, a marked difference between the morning and evening profiles is observed in the tropics, especially at the equinoxes. This is illustrated in Figs. 5a and b, which show the zonally averaged morning and evening profiles, respectively, and Fig. 5c, which shows the difference between the evening and morning, as a function of latitude. The Na layer in the tropical region (15° N–30° S) is strongly depleted in the morning. Most of the depletion occurs below 95 km, peaking at 88 km, and the morning layer column density is about 50% that in the evening.

Figure 6 shows the latitudinal dependence of the vertical tidal wind velocity at an altitude of 90.6 km (close to the peak of the Na layer), as a function of local time. This is calculated from the Global Scale Wave Model (GSWM) (Hagan and Forbes, 2002). Note that the largest wind velocities are in the tropical region (20° N–20° S). The strong downward wind, which actually starts around 100 km at midnight, appears at 91 km around 04:00 LT, reaches a maximum of −20 cm s\(^{-1}\) around 07:00 LT, and then dissipates after 10:00 LT. Between 14:00 LT and 18:00 LT, when the evening Odin measurements were made, the vertical wind reverses but its velocity is small, between −5 and +5 cm s\(^{-1}\).

This tidal behaviour in the tropics should affect the Na layer in several ways which explain the diurnal behaviour of the tropical layer (Fig. 5). First, the strong downward wind starting at midnight will cause a vertical displacement of 3–4 km by the time of the Odin measurement at 06:00 LT. Atomic Na, close to the layer peak at 91 km, will be
displaced into a region of higher pressure where the termolecular reaction

\[ \text{Na} + \text{O}_2 (+\text{M}) \rightarrow \text{NaO}_2 \]  

will form sodium superoxide, a reasonably stable reservoir species (Plane, 2004). Reaction (3), which is pressure-dependent, will also increase and produce the more stable reservoir NaHCO$_3$.

The strong downward tidal wind also affects the constituents O and H. For example, during September the atomic O concentration derived from the Wind Imaging Interferometer (WINDII) on the Upper Atmosphere Research Satellite exhibits a dramatic decrease (by over 50%) in the equatorial region between 02:00 LT and dawn, at altitudes up to about 95 km (Russell et al., 2005). Atomic O and H exert a major control on the sodium chemistry through Reaction (4) and the reactions:

\[ \text{NaO}_2 + \text{O} \rightarrow \text{NaO} + \text{O}_2 \]  

\[ \text{NaO} + \text{O} \rightarrow \text{Na} + \text{O}_2 \]

The resulting decrease in the rates of Reactions (4), (7) and (8) then reinforce the conversion of Na to NaO$_2$ and NaHCO$_3$, causing the observed depletion of Na atoms at 06:00 LT (Fig. 5).

4 Polar Depletion of the Na Layer

The global minimum of the Na column abundance occurs at high latitudes during summer, and can be an order of magnitude lower than the maximum (Fig. 1). At the same time, the FWHM width of the Na layer in the summertime polar region is also a minimum (Fig. 4a), about half the width of the layer during winter or at lower latitudes. In this section the reasons for this extreme seasonal behavior in the polar regions will be discussed in more detail.
Figure 7 displays the sodium density profiles at mid- and high latitudes between March and September, straddling the NH summer solstice. A white horizontal line has been drawn on each of the plots at 80 km to aid the eye in observing changes to the underside of the Na layer. Figure 8 compares the monthly variation of the mean Na layer profiles at latitudes of 40°N and 80°N, from spring equinox to the summer solstice. The undersides of the Na profiles during the solstice show a clear cutoff below 85 km.

Previous ground-based lidar measurements have observed the very low minimum Na abundance in the Arctic region during summer (e.g. Gardner et al., 1988; von Zahn et al., 1988; Gardner et al., 2005). Recent research has focused on the removal of metal atoms (and their compounds) on the surfaces of ice particles (Lubken and Hoffner, 2004; Plane et al., 2004), following a laboratory study which demonstrated that Fe, Na and K atoms are all removed very efficiently on low-temperature ice surfaces (Murray and Plane, 2005). Simultaneous measurements of mesospheric Na, PMCs and polar mesosphere summer echoes (PMSE) (She et al., 2006) also found that polar Na depletion is highly correlated with PMCs, whereas the smaller-sized ice particles detected by radar as PMSE exhibit a weaker anti-correlation with the Na density around 85 km.

The satellite retrievals enable the full process of polar Na layer depletion to be tracked. In April and May, the Na layer shifts gradually to a summertime profile, mainly due to the temperature-dependent Na chemistry discussed above, where a larger fraction of the sodium is in the form of NaHCO$_3$. In June, July and August the underside of the Na layer exhibits a sharp cut-off at about 85 km. In the June and July profiles, the lower edge of the layer gradually rises northwards from $\sim$50°N. The Na depletion on the underside disappears abruptly in September.

In a very recent paper (Petelina et al., 2006), the occurrence of PMC was derived from the same OSIRIS spectra that we have used to retrieve the Na density profiles. The results can therefore be used to explore the relationship between them. Although it is difficult to determine precisely the altitude of a PMC when viewed in the limb, the
PMC peak heights were found to be located mostly between 83–84 km (Petelina et al., 2006). PMCs were observed from 50° N to higher latitudes, and from the beginning of June until the end of August. The clouds appeared most frequently immediately after the summer solstice (Petelina et al., 2006). These PMC statistics are in complete accord with the polar Na profiles: the underside of the Na layer below 85 km disappears completely from June until the end of August. In fact, since the bottom of the Na layer in polar regions is very sensitive to the presence of PMCs, it may be useful as an indicator of the presence of PMCs which are not optically detectable.

5 Conclusions

This paper demonstrates the ability of our new algorithm (Gumbel et al., 2007), based on optimal estimation theory (Rodgers, 2000), to retrieve absolute Na density profiles from limb-scanning measurements of the Na D-line radiance at 589 nm in the dayglow. These global measurements provide a very useful complement to ground-based lidar observations. Particular highlights of the present data set are the first observations of the Na layer in the equatorial region. We find that the layer exhibits little seasonal variability, but has a very marked diurnal variation, particularly during the equinoxes. This is almost certainly driven by the diurnal tide. The measurements in the polar regions confirm the relationship between depletion of Na below 85 km and the presence of PMCs during the summer at latitudes above 50°.

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References


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**Fig. 1.** Grid plot of the monthly-averaged total Na column density (units: atom cm$^{-2}$), zonally averaged in 10° latitude bins. The data is compiled from OSIRIS limb-scanning radiance measurements during 2003 and 2004. The number inside each grid box is the number of integrated Na profiles that were averaged to yield the mean column density.
Fig. 2. Grid plot of the correlation coefficient between the MSISE-00 model temperature (Picone et al., 2002) and the retrieved Na density, versus latitude and altitude.
Fig. 3. Seasonal variation of the zonally-averaged Na density profile (top panel, units: atom cm$^{-3}$), and the MSISE-00 atmosphere temperature profiles (Picone et al., 2002) (bottom panel, units: K), from 75 km to 110 km at four latitude bands centred at: (a) 70° N, (b) 40° N, (c) the equator, and (d) 20° S.
Fig. 4. Top panel: seasonal variation of the full width at half maximum of the Na layer (in km), as a function of latitude and month. Bottom panel: seasonal variation of the peak height of the Na layer (in km, with ±1 km uncertainty), as a function of latitude and month.
Fig. 5. The zonally-averaged Na layer (units: atom cm$^{-3}$) in September, as a function of latitude: (a), averaged Na profile around 06:00 LT; (b), averaged Na profile around 18:00 LT; and (c), difference between the evening and morning profiles.
Fig. 6. Vertical tidal velocity at 90.6 km during September, as a function of local time (LT) and latitude. Data taken from the GSWM (Hagan and Forbes, 2002). The vertical white lines indicate the approximate local times of the Odin measurements.
Fig. 7. Contour plots of the Na density height profile (units: \( \text{atom cm}^{-3} \)) between 40° and 80° N, during (a) March, (b) April, (c) May, (d) June, (e) July, (f) August, and (g) September, 2004.
Fig. 8. Monthly Na profiles from March to July, 2004, at a latitude of (a) 40° N, and (b) 80° N.