Chemical and isotopic snow variability in East Antarctica along the 2001/02 ITASE traverse

Silvia BECAGLI,1 Marco PROPOSITO,2,4 Silvia BENASSAI,1 Onelio FLORA,3 Laura GENONI,3 Roberto GRAGNANI,4 Ombretta LARGIUNI,1 Simone Luca PILI,4,5 Mirko SEVERI,1 Barbara STENNI,3 Rita TRAVERSI,1 Roberto UDISTI,1 Massimo FREZZOTTI4

1Chemistry Department – Analytical Chemistry, Scientific Pole, University of Florence, Via della Lastruccia 3, I-50019 Sesto Fiorenti (Florence) Italy E-mail: silvia.becagli@unifi.it
2Museo Nazionale dell’Antartide, Strada del Laterino 8, I-53100 Siena, Italy
3Department of Geological, Environmental and Marine Sciences, University of Trieste, Via E. Weiss 2, I-34127, Trieste, Italy
4ENEA, Centro Ricerche, Casaccia, PO Box 2400, I-00100 Rome, Italy
5Department of Earth Science, University of Cagliari, Via Trentino 51, 1-09127 Cagliari, Italy

ABSTRACT. As part of the International Trans-Antarctic Scientific Expedition (ITASE) project, a traverse was carried out from November 2001 to January 2002 through Terre Adélie, George V Land, Oates Land and northern Victoria Land, for a total length of about 1875 km. The research goal is to determine the latitudinal and longitudinal variability of physical, chemical and isotopic parameters along three transects: one west–east transect (WE), following the 2150 m contour line (about 400 km inland of the Adélie, George V and Oates coasts), and two north–south transects (inland Terre Adélie and Oates Coast–Talos Dome–Victoria Land). The intersection between the WE and Oates Coast–Victoria Land transects is in the Talos Dome area. Along the traverse, eight 2 m deep snow pits were dug and sampled with a 2.5 cm depth resolution. For spatial variability, 1 m deep integrated samples were collected every 5 km (363 sampling sites). In the snow-pit stratigraphy, pronounced annual cycles, with summer maxima, were observed for nssSO42−, MSA, NO3− and H2O2. The seasonality of these chemical trace species was used in combination with stable-isotope stratigraphy to derive reliable and temporally representative snow-accumulation rates. The study of chemical, isotopic and accumulation-rate variability allowed the identification of a distribution pattern which is controlled not only by altitude and distance from the sea, but also by the complex circulation of air masses in the study area. In particular, although the Talos Dome area is almost equidistant from the Southern Ocean and the Ross Sea, local atmospheric circulation is such that the area is strongly affected only by the Ross Sea. Moreover, we observed a decrease in concentration of aerosol components in the central portion of the WE transect and in the southern portion of the Talos Dome transect; this decrease was linked to the higher stability of atmospheric pressure due to the channelling of katabatic winds.

INTRODUCTION

The aim of the International Trans-Antarctic Scientific Expedition (ITASE) programme is the study of the spatial variability and evolution over time (last 200 years) of parameters related to the atmospheric system, and recorded in the snowfall and upper layers of the Antarctic ice sheet (Mayewski and Goodwin, 1999). These data can be used to assess recent environmental changes in the drainage area extending from Dome Concordia (Dome C) to the Ross Sea and the Southern Ocean.

A detailed database is essential for determining the spatial significance of trends in the chemical and isotopic composition of surface snow. This information allows a better understanding of the effects of atmospheric conditions, air mass circulation and the strength and direction of dominant winds on the chemical and isotopic composition of snow. Moreover, such knowledge is essential for interpreting temporal changes in the vertical chemical profiles of deep ice cores drilled in this area. Due to the scarcity of climatological and glaciological data for some interior regions of the Antarctic continent, high-spatial-resolution records from inland sites are extremely important for understanding the chemical–physical processes that determine the concentration of chemical species as a function of altitude, distance from the sea, accumulation rates and other glaciological and climatological conditions.

The concentrations of the main ionic components can provide information about relative contributions of different atmospheric aerosol sources. The study of changes in chemical composition as a function of altitude and distance from the sea allows a better understanding of atmospheric aerosol transport and depositional processes.

Given that accumulation rates can modify the original composition of chemical markers in snow, by dilution in case of dominant dry deposition: Legrand and Mayewski, 1997) and post-depositional effects (particularly for volatile and unstable species like HCl, HNO3 and H2O2; McConnell and others, 1998; Röthlisberger and others, 2000; Traversi and others, 2000), it is necessary to understand if and where these potential palaeomarkers can be used to reconstruct the composition of past atmospheres.
For this purpose, we report the chemical and stable-isotope spatial variability in surface firn collected from Terre Adélie, George V Land, Oates Land and northern Victoria Land by sampling 1 m firn cores and snow pits at high spatial resolution along two north–south transects and one east–west transect. At selected sites, a detailed stratigraphy of the first 2 m of snow deposition was reconstructed with the aim of identifying seasonal patterns and the effects of post-depositional processes on some chemical markers.

SAMPLING AND ANALYSIS

In the framework of the ITASE project, France and Italy carried out a traverse along one west–east (D66–GV5) and two north–south (D59–D85 and GV7–Talos Dome–M4) transects from November 2001 to January 2002 (Fig. 1).

The west–east transect, named WE in the discussion, follows the 2150 m contour line, about 400 km from the Adélie, George V and Oates coasts; the first north–south transect (D59–D85, named NS-1) follows the central part of the previous French traverse from Dumont d’Urville to Dome C; the second north–south transect (GV7–Talos Dome–M4, named NS-2) follows the ice divide extending from the Oates Coast to Talos Dome, and then runs along the 2000 m contour line in the direction of Taylor Dome.

To limit the impact of wind-driven processes, the eight snow-pit sites (D66, GV1, GV2, GV3, GV4, GV5, GV7, TDN) were chosen in windless areas on the basis of satellite images and a digital terrain model at 1 km resolution provided by Rémy and others (1999). Field observations revealed that summer melt layers were not present at depth in any of the snow pits.

At site 31Dpt, shown in the map (Fig. 1), snow-pit samples were collected during the 1998/99 ITASE traverse (Frezzotti and Flora, 2002). The related chemical data (Becagli and others, 2003) are included in the discussion together with those of the sites along transect NS-2. The nine snow pits were hand-dug and sampled at 2.5 cm resolution up to 1.5–2 m depth by workers wearing non-particulating suits, polyethylene gloves and particle masks. Pit walls were cleaned by removing a 5–10 cm snow layer with a stainless-steel scraper just before sampling. Pre-cleaned polypropylene vials were inserted in the vertical snow walls; after removal, each sampling vial was labelled and sealed in a double polyethylene bag. Samples were stored below –15°C until melting immediately prior to chemical analysis. The snow density was determined along the pit wall by collecting horizontal core sections with a known volume and weighing them immediately after retrieval.

Surface snow was sampled every 5 km of the traverse (363 samples) down to 1 m by hand auger, then sealed in polyethylene bags and kept frozen. These 1 m cores were cleaned by removing the more external parts with a
stainless-steel scraper; each core was then melted (furnishing one sample) and analyzed. Dionex series instruments were used to analyze major cations (Na+, NH4+, K+, Mg2+, Ca2+) and anions (Cl-, NO3-, SO42-, F-, MSA) in all samples from snow pits and 1 m integrated firm cores. A detailed description of analytical methods is reported by Gagnon and others (1998) and Udisti and others (2004). H2O2 concentrations in snow-pit samples were determined using the peroxidase-based fluorescence method described in Largiuni and others (2002). Stable-isotope (δ18O) and tritium analyses were carried out following the procedures reported in Proposito and others (2002) and Stenni and others (2002), respectively. Stable-isotope composition was measured on 2.5 cm samples collected from six snow pits at the following sites: D66, GV1, GV2, GV3, GV5 and GV7. δ18O measurements were carried out on all surface snow samples (1 m cores), while tritium activity was measured every four samples during the time-consuming analytical method.

RESULTS AND DISCUSSION
Seasonal pattern and accumulation rates
In order to assess accumulation rate variability in the study area, stratigraphic dating of all snow pits was performed through the analysis of chemical and isotopic parameters with marked seasonal patterns (nssSO42−, MSA, H2O2, NO3− and δ18O). Dating was accomplished through an multi-parametric approach involving the calculation of a normalized profile for the seasonal behaviour of each chemical parameter. A detailed description of the multiparametric dating procedure is reported in Udisti (1996). The main advantage of this procedure is that it allows comparison of profiles presenting different absolute values and compensation of the time shift in the summer peaking of the different seasonal markers. For instance, year-round aerosol measurements in Antarctic coastal sites show that nitrate maxima occur mainly in spring due to the stratospheric–tropospheric exchanges (Legrand and others, 1999), while nssSO42− and MSA peak in full summer (Minikin and others, 1998).

Finally, in order to determine the accumulation rate for each year, the normalized sum profile from each snow pit was compared with the related δ18O profile, and the annual boundary (dashed line in Fig. 2) was put on each maximum identified in the normalized sum profile which is very often coincident with the δ18O corresponding maximum. All selected chemical markers for stratigraphic dating show a clear seasonal pattern in the region sampled, with summer maxima and winter minima; a few ambiguities were found in some sites for markers which are affected by post-depositional processes. For instance, H2O2 in most snow pits shows a weakening of the seasonal pattern due to decomposition reactions occurring at the snow surface and/or to post-depositional diffusion in snow layers (Wolff, 1996; McConnell and others, 1998). In particular, H2O2 content at site GV3 shows a gradually decreasing trend with depth, and there is an almost imperceptible seasonal pattern. In some pits (GV2, GV7 and TDN), nitrate also shows a decreasing trend with depth due to HNO3 post-depositional re-emission (Wolff, 1995; Wagnon and others, 1999; Röthlisberger and others, 2000); nevertheless, the seasonal pattern is still detected over the depth range analyzed. No evidence of post-depositional loss or smoothing of seasonal patterns was observed for nssSO42−, known as a conservative species in snow layers. The other biogenic marker (MSA) shows no evidence of movement towards the winter layers, despite some shift with respect to the chosen annual boundary. Such a post-depositional effect was found by Pasture and Mulvaney (2000) and Curran and others (2002) in shallow firm cores sampled at sites with higher accumulation rate located in the Antarctic Peninsula and at Law Dome. As for the isotopic profile, a clear seasonal signal can be observed in snow pits D66, GV1, GV5 and GV7, where δ18O and chemical records are in very good agreement. By contrast, at sites GV2 and GV3 post-depositional processes allowed the seasonal δ18O signal to be smoothed with respect to the other chemical markers. At GV2 and GV3, the H2O2 profile is heavily perturbed; this is probably not linked to the accumulation rates at these sites, which are not significantly lower than those at other sites (Table 1), but could be related to the cooler, windier climatic conditions in the central portion of the WE transect, as observed by Manzella and others (2004).

On the basis of the chosen annual boundaries (reported in Fig. 2) and density profile, accumulation rates as mm.w.e. were calculated for each year and are reported in Table 1 together with the averaged value for the sampled time interval. The sites are arranged according to their position in the two transects. The mean accumulation rates agree with those obtained from stake measurements and β-counting, taking into account site variability and the different time periods covered by the different methods (Becagli and others, 2004).

Along the WE transect, mean accumulation rates appear to be quite homogeneous, with minima in the central portion of the transect (GV1–GV3), located farthest from the Southern Ocean. Along the NS-1 transect instead, the mean accumulation rate decreases from the site closest to the Southern Ocean (GV7) to Talos Dome (TDN), and remains constant moving towards the Ross Sea (31°D). A detailed discussion on accumulation-rate variability as a function of site temperature is reported in Manzella and others (2004).

Stable-isotope variability
δ18O
Theδ18O values obtained from 361 surface snow samples (1 m cores) collected every 5 km along two NS transects and one WE transect are presented in Figure 3 as a function of altitude. The relationships between stable isotopes and temperature are discussed in a companion paper by Magand and others (2004). The expected negative correlation between altitude and δ18O is evident along the two NS transects; however, in the case of transect NS-2, this correlation only applies to the sampling sites from GV7 to TDN. After the ice divide, and toward M4, the data do not fall on a straight line, suggesting a differing climatic situation and/or a different origin for the moisture sources supplying precipitation to the area. This is confirmed by the lower temperatures observed between TD and M4 (Magand and others, 2004). Elevation gradients of −1.5‰ (100 m)−1 and −1.3‰ (100 m)−1 were calculated for the NS-1 and NS-2 transects up to Talos Dome, respectively; they are slightly higher than the −1‰ (100 m)−1 value observed along the 1998/99 Terra Nova Bay (TNB)–Dome C (DC) traverse

Becagli and others: Chemical and isotopic snow variability in East Antarctica
Fig. 2. Concentration vs depth profile of the chemical parameters used for dating, and the calculated normalized sum (see text for details) compared with the δ¹⁸O profile.

Becagli and others: Chemical and isotopic snow variability in East Antarctica
This is related to the decrease in temperature with altitude, creating a super-adiabatic lapse rate along the NS transects (Magand and others, 2004), in contrast with the near-dry adiabatic lapse rate of the TNB–DC traverse (Frezzotti and Flora, 2002). The surface snow samples of the WE transect do not show any correlation with altitude because they were collected along a limited range of altitudes; furthermore, the samples collected at 2100–2250 m (GV1–GV5) exhibit quite a large range of δ18O values as a consequence of the colder, windier conditions characterizing the central sector of this transect (Magand and others, 2004).

One must bear in mind the following limitations: (i) the snow deposited in a 1 m integrated sample may represent different years, depending on the accumulation rate; (ii) the seasonal distribution of accumulation and its interannual variability may affect mean δ18O values (Schlosser, 1999); (iii) wind-scouring phenomena may also introduce a bias in δ18O values.

Tritium
Tritium data from surface snow samples (1 m cores) collected along the two NS transects are displayed in Table 1. Annual accumulation rate (mm w.e. a–1) from snow pits

<table>
<thead>
<tr>
<th>Site</th>
<th>2001</th>
<th>2000</th>
<th>1999</th>
<th>1998</th>
<th>Average</th>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>D66</td>
<td>212.4</td>
<td>179.0</td>
<td>–</td>
<td>–</td>
<td>195.7</td>
</tr>
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<td>GV1</td>
<td>167.9</td>
<td>111.4</td>
<td>146.7</td>
<td>143.3</td>
<td>142.3</td>
</tr>
<tr>
<td>GV2</td>
<td>127.7</td>
<td>194.9</td>
<td>125.0</td>
<td>–</td>
<td>149.2</td>
</tr>
<tr>
<td>GV3</td>
<td>125.9</td>
<td>212.5</td>
<td>157.1</td>
<td>116.8</td>
<td>153.1</td>
</tr>
<tr>
<td>GV4</td>
<td>146.6</td>
<td>197.0</td>
<td>202.2</td>
<td>–</td>
<td>181.9</td>
</tr>
<tr>
<td>GV5</td>
<td>141.1</td>
<td>145.6</td>
<td>179.9</td>
<td>–</td>
<td>155.5</td>
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<td>North-south transect</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>GV7</td>
<td>241.7</td>
<td>279.3</td>
<td>–</td>
<td>–</td>
<td>260.5</td>
</tr>
<tr>
<td>GV5</td>
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<td>145.6</td>
<td>179.9</td>
<td>–</td>
<td>155.5</td>
</tr>
<tr>
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<td>145.6</td>
<td>129.8</td>
<td>147.2</td>
<td>104.2</td>
</tr>
<tr>
<td>31Dpt</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>112°</td>
</tr>
</tbody>
</table>

*Values from Stenni and others (2003).*

Figure 4 as a function of distance from the coast. Data from the 1998/99 TNB–DC traverse (Proposito and others, 2002) are also reported for comparison. In both datasets a positive trend in tritium activity is observed with increased distance inland. The positive correlation between tritium activity and distance from the coast/elevation has been observed in Antarctica by other authors (Merlivat and others, 1977; Jouzel and others, 1979, 1982) and may be related to a more efficient exchange between the stratosphere and troposphere over the more interior part of the continent. Jouzel and others (1982) suggested that tritium formed at high stratospheric altitudes is brought down within the circumpolar vortex, which is maximum in winter; a rapid exchange of air between the lower polar stratosphere and troposphere may occur during Antarctic winters due to absence of temperature inversion in the South Polar tropopause. Nevertheless, compared to data from the 2001/02 ITASE traverse, results from the TNB–DC 1998/99 traverse for the same distance from the coast show higher tritium concentrations. A probable explanation is that the 1998/99 traverse starts several degrees of latitude south of the 2001/02 traverse, being effectively closer to the interior of Antarctica. This is supported by the more ‘maritime’ condition of the new ITASE traverse, as suggested by the higher accumulation rates. The tritium content of atmospheric water vapour is indeed diluted by fresh oceanic moisture, which is characterized by low tritium activity.

Figure 5 plots all tritium data against latitude and includes those from the WE transect and two points reported by Jouzel and others (1982) for a coastal site (A3) in Terre Adélie and data from Dome C for the 1940–56 period. These authors showed a strong increase in tritium activity with latitude over Antarctica, with a maximum of 32 TU (tritium units) for the South Pole (outside the represented range).

Chemical variability
The concentrations of Na+, Cl−, nssSO42−, MSA and NO3− in surface snow are shown in Figures 6–8 for transects NS-1, NS-2 and WE, respectively. Ion concentrations are reported as a function of altitude S for the NS transects, and as a function of longitude E for the WE transect. Whereas elevation and distance from the sea remain fairly constant along the WE transect, they vary substantially along NS-1. NS-2 shows a particular feature: moving south, as the distance from the Southern Ocean increases, proximity to the Ross Sea firstly increases and then remains constant; in the transect, elevation increases up to the ice divide (Talos
Dome) and then decreases toward the last site (M4; Fig. 1). Elevation and distance from the Southern Ocean are reported on the top of Figures 6–8. In the same figures, circles are related to 1 m firm-core integrated samples, and dots represent mean values calculated from snow-pit samples; the vertical bars indicate the associated standard deviations. The data from the French traverse performed in 1982/83 (Legrand and Delmas, 1985) are shown as triangles in Figure 6.

Sea-spray components
Along transect NS-1 (Fig. 6) the primary sea-spray components (Na+ and Cl–) decrease sharply in the first 50 km from the coast, reaching nearly constant values around 17 and 38 $\mu$g L$^{-1}$, respectively. The French (Legrand and Delmas, 1985) and our profiles show similar data except for a few values at the start of the transect.

Wind-borne sea-salt particles are coarse and their concentrations rapidly decrease with increasing altitude and distance from the sea. The progressive decrease in Cl– concentrations vs altitude and distance from the coast suggests that these are critical parameters for penetration of sea salt toward inland regions (Minikin and others, 1994; Mulvaney and Wolff, 1994; Kreutz and Mayewski, 1999; Udisti and others, 1999). The same authors observed that the effect of elevation is more critical than distance from the sea.

Along transect NS-2 (Fig. 7), Na+ and Cl– quickly decrease between 70 and 130 km from the coast, reaching relatively stable values from 71°30’ to 73°30’ S, where the highest and most inland sites are located. In the part of the transect facing the Ross Sea, concentrations increase as elevation decreases, and proximity to the sea remains quite constant, suggesting a strong influence of the Ross Sea on the last part of the transect due to atmospheric circulation over this area. A good correspondence is observed between 1 m firm cores and snow-pit mean values, with large standard deviations due to intense marine events in the stations which are more affected by the Ross Sea. The snow pits sampled at these stations (31Dpt and TDN) are characterized by good Cl–Na and Mg–Na linear correlations, as demonstrated by good Cl–Na and Mg–Na linear correlations, as demonstrated by the value of the linear regression coefficient ($R$) reported in Figure 9, with a slope very close to the theoretical sea-water ratio (see dashed line in the plot). By contrast, although the two stations closer to the Southern Ocean (GV7 and GV5) show good Cl–Na and Mg–Na linear correlations, they are characterized by lower Cl–Na and Mg–Na ratios, reflecting different residence times for Cl– and Mg2+ with respect to Na+. The sites located in the area between the Southern Ocean coastline and the ice divide appear to be affected by marine input; however, air masses reaching this area do not come directly from the Southern Ocean, but probably follow a longer pathway along which some species undergo fractionation.

In transect WE (Fig. 8), Cl– and Na+ profiles exhibit a rather flat pattern, and mean values from snow pits fit very well with 1 m firm-core values; the high levels recorded around 135° E are at sites close to the Southern Ocean and at lower altitudes. Figure 10 illustrates the Cl–Na and Mg–Na linear correlation with respect to longitude E. All sites show good correlations with the slope of the regression line,
which is always lower than sea-water ratio, demonstrating that fractionation of Cl\(^-\) and Mg\(^{2+}\) occurs in the entire study area, and that all the stations behave similarly with respect to primary marine aerosol. This common pattern is due to the relatively constant high altitude and similar distance from the coastline.

**Sulphur compounds**

Principal component analysis (PCA) applied to snow-pit samples revealed that sea-spray components at all sites contribute to the first principal component. Sulphate, partly derived from the sea-spray input, contributes to this factor only at 31Dpt, the site most significantly affected by the Ross Sea. In the other snow pits, total sulphate always contributes to the second factor together with nssSO\(_4^{2-}\); this means that at all sites except 31Dpt, nssSO\(_4^{2-}\) is the most important contribution to the total sulphate budget. The average contributions of nssSO\(_4^{2-}\) to the total SO\(_4^{2-}\) budget are 70%, 76% and 78% in transects NS-1, NS-2 and WE, respectively. Therefore, only the nssSO\(_4^{2-}\) profile is shown in Figures 6–8.

The nssSO\(_4^{2-}\) profile for transect NS-1 shows quite constant background values, consistent with measurements performed by Legrand and Delmas (1985), reported as triangles in Figure 6. This pattern is typical of secondary aerosol components, mainly related to fine particles, at sites located at relatively high altitudes and distant from the sea,
where biogenic nssSO$_4^{2-}$ reaches quite constant background values (e.g. Minikin and others, 1994; Udisti and others, 1998; Becagli and others, 2003; Traversi and others, 2004). Transect NS-2 (Fig. 7), although starting nearer the coast and at lower altitudes with respect to transect NS-1, shows a constant level of nssSO$_4^{2-}$ at the first station, where a large part of the coarsest sea-salt particles was previously deposited. As distance/altitude from the coast increases, the gas-derived secondary aerosol (nssSO$_4^{2-}$) becomes the prevailing species because the diameter of particles shifts toward the finest size fraction which is mainly affected by long-range transport.

A gradually increasing trend is observed for nssSO$_4^{2-}$, as latitude S and distance from the Southern Ocean increase up to the ice divide; constant values were measured from the divide to 31Dpt, after which concentrations sharply increased over the last part of the transect. This behaviour could reflect changes in accumulation rates and in the strength of marine sources. The increase in concentrations between GV7 and TDN could be explained supposing a dominant dry deposition in this area, characterized by a generally low accumulation rate, leading to constant deposition fluxes as accumulation rates decrease. The southern part of the transect is about 250 km from the Ross Sea coast. Despite the relatively long distance from the sea, this part of the transect is strongly influenced by marine air masses (as shown by the Na$^+$ and Cl$^-$ pattern) due to the local atmospheric circulation (Cullather and others, 1996), thus causing an increasing concentration trend for nssSO$_4^{2-}$.

The nssSO$_4^{2-}$ concentration pattern for transect WE (Fig. 8) could also be driven by accumulation rates if a prevailing dry deposition is supposed; the highest concentrations are found in the central portion of the transect where accumulation is lowest (see Table 1), suggesting a constant flux in this area.

MSA concentrations show high variability along the three transects. In particular, they tend to increase slightly southward in the two NS transects. Due to its biogenic marine origin, the MSA distribution should decrease with distance inland and increasing altitude; however, the decrease is slower than that of sea salt because MSA is distributed in the finest size fraction of aerosol particles, thereby allowing MSA to reach the Antarctic Plateau (Legrand and others, 1992; Udisti and others, 1998; Kreutz and Mayewski, 1999).

The increasing pattern observed in transect NS-1 could be explained by a change in accumulation rates, which decrease moving inland, leading to a decreased dilution factor for the aerosol components deposited on the snow. Instead, in transect NS-2, the pattern is thought to be similar to that of nssSO$_4^{2-}$, with a gradual increase in concentrations toward the ice divide, again consistent with variations in accumulation rates, and a further increase until 31Dpt as the Ross Sea is approached. In the very last part of the transect, an anomalous diminishing trend is observed which must be confirmed by further data. The MSA pattern observed in profile WE is similar to that of nssSO$_4^{2-}$, with accumulation rates controlling MSA concentrations in this area.

NO$_3^-$

NO$_3^-$ concentrations show different patterns in transects NS-1, NS-2 and NS-3 (Figs 6–8). NO$_3^-$ concentrations in transect NS-1 are fairly constant and in good agreement with the values found in the shallow snow samples collected during the 1982–83 Terre Adélie traverse (Legrand and Delmas, 1985). The NO$_3^-$ profile for transect NS-2 shows a progressive increase up to 500 km from the coast. There are many spikes in transect WE, especially in the 140–145° E range, which are not reflected in the standard deviation of the corresponding pit samples.

Due to the decreasing trend in NO$_3^-$ concentrations with depth observed in some snow pits (see Fig. 2), snow-pit mean values are likely to reach lower values with respect to
1 m surficial firn cores. However, mean NO$_3^-$ concentrations are quite constant along the WE transect both for 1 m firm cores and for snow pits. As already described for nssSO$_4^{2-}$, the NO$_3^-$ concentration pattern results from the transport of fine aerosol particles and possible dilution by mean accumulation rates at the sampling site. However, the interpretation of the NO$_3^-$ profile is complicated by post-depositional processes (HNO$_3$ re-emission and photochemical decomposition; Wolff, 1995; Wagnon and others, 1999; Jones and others, 2000; Röthlisberger and others, 2000) occurring on surficial snow layers in low-accumulation sites. These processes may explain why nitrates in transect WE, unlike other secondary species (especially nssSO$_4^{2-}$), do not describe a parabola. Indeed, the sites located in the central portion of this transect are characterized by the lowest accumulation rates (see Table 1).

CONCLUSIONS

The chemical and isotopic dataset obtained from the traverse through Terre Adélie, George V Land, Oates Land and northern Victoria Land completed during the 2001/02 Antarctic campaign allowed the first detailed analysis of the spatial distribution of chemical tracers and of transport processes affecting this area. The seasonal patterns of nssSO$_4^{2-}$, MSA, H$_2$O$_2$, NO$_3^-$ and $\delta^{18}$O were quite evident in the stratigraphic profiles from snow pits, allowing reliable estimates of age and accumulation rates at all sampled sites. Considering natural site variability, the accumulation rates obtained with different measurements (stake farms and δ-out counting, as reported by Magand and others (2004), were consistent with our chemical and isotopic data.

The relationship between stable isotopes and altitude was examined for the two NS transects. Elevation gradients of $-1.5\%$(100 m)$^{-1}$ and $-1.3\%$(100 m)$^{-1}$ were calculated for the NS-1 transect and the northern part of the NS-2 transect, respectively. Stable-isotope data suggest different climatic conditions and/or different origins for the moisture sources supplying precipitation to the sampling sites along the transect from Talos Dome to M4. Surface snow samples from the WE transect show a large range of $\delta^{18}$O values that reflect the colder, windier climatic conditions characterizing the central sector of this transect (Magand and others, 2004).

Tritium concentrations in surface snow samples show a positive trend moving inland, in agreement with data obtained from the previous TNB–DC traverse carried out in the 1998/99 Antarctic campaign (Proposito and others, 2002). However, when considering the same distance from the coast, the 1998/99 traverse data show higher tritium activity compared to data from the 2001/02 traverse. This difference may be ascribed to the difference in latitude, probably linked to the more ‘maritime’ conditions in the area covered by the new ITASE traverse.

As for spatial variability along the three sampled transects, the NS transects highlighted the varying importance of marine sources as distance from the sea increases. In particular, transect NS-1 shows the typical decreasing trend in marine primary compounds with increasing altitude and distance from the sea. Concentrations and spatial patterns are consistent with those obtained along a previous traverse (Legrand and Delmas, 1985).

Transect NS-2 is characterized by two areas of influence: the Southern Ocean sector and the Ross Sea. The latter more strongly affects this area due to the atmospheric transport of marine air masses to the Talos Dome area. These atmospheric conditions are consistent with the modelled flux of moisture into the Ross Sea–East Antarctic region (Cullather and others, 1996).

Much of transect WE is swept by prevailing katabatic winds (Mancini and Frezzotti, 2003) which probably affect annual accumulation rates, with the lowest rates found at GV3. Moreover, high wind speeds result in more wind-blowed snow and facilitate snowpack ventilation. Wind-blowed snow contributes to scavenging conservative and volatile species, whereas concomitant sublimation may concentrate or release some chemical species. Wind pumping produces an air flux into the snow, favouring chemical exchanges between the atmosphere and snow (Pomeroy and Jones, 1996; Waddington and others, 1996). Surface and wind-driven sublimation, controlled by surface slope along the prevailing wind direction, has a huge impact on surface mass balance (Frezzotti and others, 2002).

Primary and secondary aerosol components along transect WE show no particular distribution patterns, with marine aerosol fractionation and low background values confirming the hypothesis of scarce oceanic input. This could be due to the persistence of higher pressures in this area, which block storms and marine air masses (Magand and others, 2004), and/or to the atmospheric circulation around the low-pressure centre that is located seaward of the Wilkes Land coast (Cullather and others, 1996).

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