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GEOSCIENCES

Aerobiology in High Latitudes: Evidence of Bacteria Acting as Tracer of Warm Air Mass Advection reaching Northern Antarctic Peninsula

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Abstract: Despite the extent use of geochemical tracers to track warm air mass origin reaching the Antarctic continent, we present here evidences that microorganisms being transported by the atmosphere and deposited in fresh snow layers of Antarctic ice sheets do act as tracers of air mass advection from the Southern Patagonia region to Northern Antarctic Peninsula. We combined atmospheric circulation data with microorganism content in snow/firn samples collected in two sites of the Antarctic Peninsula (King George Island/Wanda glacier and Detroit Plateau) by using flow cytometer quantification. In addition, we cultivated, isolated and submitted samples to molecular sequencing to precise species classification. Viable gram-positive bacteria were found and recovered in different snow/firn layers samples, among dead and living cells, their number concentration was compared to northern wind component, stable isotopes of oxygen, d180, and the concentration of crustal elements (Fe, Ti and Ca). Use of satellite images combined with air mass back-trajectory analysis obtained from the NOAA/ HYSPLIT model corroborated the results.

Key words: extremophiles, atmospheric transport, Antarctica, atmospheric tracers, aerobiology.

INTRODUCTION

The aerial transport/dispersal of biological material is believed to be an important component of the input to remote locations such as the Antarctic region, and plays a crucial role in shaping patterns of biodiversity in such places (Fierer 2008, Pearce et al. 2010, 2016). Intimately linked with ecological and evolutionary processes in the colonized environments, the rates of airborne input, survival after transport process, and viability after arrival, is essential for understanding the origin and dynamics of several processes. The capacity of cells to live in extreme environments and remain viable for thousands and millions of years (Priscu et al. 1998, Karl et al. 1999, Christner et al. 2003, Bidle et al. 2007) has attracted the attention of the scientific community to microorganisms found in deep and old ice of the Earth's polar regions. Although the interest in the microbial communities of Antarctica has typically been turned toward ecologic, phylogenetic, and metabolic studies, microorganisms entrapped in snow/firn/ice layers in ice sheets could be looked from a different point of view, such as their origin and moment of deposition.

Considering the atmospheric transport events of short duration (time scale of days) between South America and the Antarctic Peninsula, the use of gaseous geochemical air mass tracers of crustal sources, for instance the ²²²Rn, have been successfully used, since the 80's decade (Pereira, 1999). Furthermore, combined ²²²Rn and Si, anthropogenic elements (Bi, Cd, Cr, Cu, Ni, V, and Zn) and biomass burning aerosols (black carbon) were also employed as tracers (Evangelista & Pereira 2002, Mishra et al. 2004, Pereira et al. 2006). Like the geochemical atmospheric tracers, the presence of South-American plants pollen was also observed in the seas of the Antarctic region since the early decades of the 20th century as well as a continuous arrival of pollen, by atmospheric transport, was firstly reported (Fritsch 1912 cited in Marshall 1996a).

The concept of monitoring biological material as atmospheric tracers is broader than the use of pollen species, since the Antarctic ice sheets contain a great diversity of microorganisms as viable and non-viable cells (Abyzov 1993), and potentially they may indicate exogenous origin. It has earlier shown that these microorganisms are found in several different Antarctic microhabitats and many of these species are not able to duplicate themselves in a normal life cycle (Warwick 2000). This finding suggests that these species may arrive in Antarctica coming from other regions of the globe, and may not derive from local communities (Warwick 2000). Additionally, the Antarctic habitats support many microorganism species of cosmopolitan nature (Warwick 1988). Therefore, the Antarctic continent, besides its geographic and climatic isolation, seams to continuously exchange biological matter with other regions of the planet (Warwick 1988, 2000). Microorganisms are transported to continental scales mostly attached to dust or organic fragments (Prospero 2005), but till now their atmospheric monitoring in Antarctica is still scarce and reduced to few summer campaigns. Basille et al. (1997) presented the first studies

using radiogenic isotope ratios of ⁸⁷Sr/⁸⁶Sr and ¹⁴³Nd/¹⁴⁴Nd in insoluble mineral particles in Antarctic ice cores to demonstrate that most dust reaching Central Antarctica derive from the Patagonian semi-desert region. For the Antarctic Peninsula, Dalia et al. (2004) and Pereira et al. (2006) employed simultaneous trace elements in aerosols, ground meteorological databases, acquisition of satellite images and atmospheric transport models to characterize the influence of the South American continent over the Northern Antarctic Peninsula in terms of dust emissions.

Although, the atmospheric dispersion of microorganisms reaching Antarctica have been reported before (Marshall 1996, 1997), their use as atmospheric tracers, based on quantitative and gualitative diversity in dated ice layers is still scarce. In this work, we have combined the microbiological analysis of bacteria entrapped in snow/firn/ice layers with concomitant tracers of lower latitude continental locations in order to investigate if microorganisms are transported to Antarctica jointly with warmer air masses. For King George Island we employed as tracer the geochemistry of dust material (Ca, Ti and Fe) and for Northern Antarctic Peninsula / Plateau Detroit the stable isotopes of oxygen that exhibit different signatures according to the latitudes.

MATERIALS AND METHODS

Study sites

Snow and firn samples were collected at 2 sites in Antarctica (Figure 1): (1) in the accumulation zone of an outlet glacier (Wanda Glacier) at Krakow Icefield / King George Island (61°50'-62°15'S; 57°30'-59°00'W) on November 2004, 350 m a.s.l., under the support of the Brazilian Antarctic Program; and at (2) Detroit Plateau located at **Graham Land** in Northern Antarctic Peninsula at approximately 2,000 m a.s.l. (64°05'S; 59°36'W) during CASA (Climate of the Antarctica and



Figure 1. (a) Sampling sites at Antarctic Peninsula for microbiological identification and (b-d) sampling of aerosols and snow/firn layers for microbiology at Detroit Plateau (Northern Antarctic Peninsula) and Wanda Glacier (King George Island).

Southern America) project expedition between November 7th, 2007 and December 14th, 2007 and and ITASE (International Trans Antarctic Scientific Expedition) research initiatives, a joint international scientific collaboration among United States, Chile and Brazil. Both snow pits at the two locations were manually setup by the scientific teams.

Samplings at King George Island / South Shetland Islands

The South Shetland Islands/Antarctic Peninsula are located approximately 550 km away from the Southern South America tip and is influenced by climatic regimes of several origins (polar, oceanic and continental, mainly derived from South America), Evangelista (1999). The wind structure is predominantly influenced by the high frequency of cyclonic systems that advect humid and warm air into that region, promoting strong winds and large volume of precipitation (Bintanja 1995). Evangelista & Pereira (2002) demonstrated that the cyclonic trajectories and their intrinsic energy are the key factors that contribute most to the apportionment of the atmospheric particulates from South America to that site. Along the year mean air temperature ranges from +2°C to -10°C. At King George Island, summer temperature remains above 0°C, typically from December to the end of March, which is responsible for intense melting processes well evidenced by the ionic and isotopic (d¹⁸O) signals in ice cores retrieved from its main domes, Simões et. al (2004). In the present work we sampled in November 2014 before air temperatures reach melting values in King George Island. Therefore, the sampled snow pack corresponded to the period of early autumn when accumulation of snow started in the region to late spring of 2004 and can be considered preserved. Attending the need of retrieving microorganisms from snow/firn

samples, three shallow pits (I, II and III) of 140 cm, 200 cm and 80 cm depth were dug on King George Island. Samples were collected on aseptic procedures, Figure 1, storage in autoclaved glass flasks and kept frozen until laboratory procedures in Brazil. Samples were collected at least 1 km away from outcrop rocks, the depth of snow pits were limited by the thick ice layer referred to the summer of the previous year. Samples of Pit I and II were collected in every 20 cm layer while pit III was sampled in each 10 cm. Pits II and III were only used to setup the microbiological method protocols while Pit I consisted of the main sample group which data comprise this work. Mean snow/firn volume for each layer was 150 mL and a total number of 25 samples obtained from the 3 pits.

Accumulation rate of the snow pack was based on the air temperature database obtained at the AWS (Automatic Weather Station) located at the Brazilian Antarctic Station Comandante Ferraz (~7 km from Wanda glacier). Both are located in the Admiralty Bay in King George Island. The positive to negative temperature turn over, when snow may deposit and start to accumulate, occurred approximately around March 15th which corresponded to the bottom chronology of the snow pack (140 cm from the top). Top of the pits corresponded to the sampling period. An estimate of the months along the snow profile was achieved considering a uniform deposition model. Figure 2 depicts inter-annual data of snow precipitation, wind velocity and air temperature for the study site, during 2004, available at http://www.cptec. inpe.br/antartica/. Instrumental precipitation inferred at the AWS presented an approximate uniform pattern as demonstrated by ANOVA (F=0.31; F_{critical}=4.07; P=0.82), calculated at 0.05 significant level for data in the shaded box at Figure 2 (bottom). Small decrease observed during July-August for precipitation may be

explained by the increased wind velocity at that period, which is mostly responsible for a drift of surface snow.

Detroit Plateau / Northern Antarctic Peninsula

Detroit Plateau is located at **Graham Land** in Northern Antarctic Peninsula, with heights above 1,500 m. Contrarily to King George Island, very few is known on the climatic and snow accumulation at the plateau. Sampling at Detroit Plateau adopted the same microbiological protocols as those established to Wanda glacier/King George Island. A 2m-pit was dug and snow layers' chronology was inferred by the d¹⁸O measurements in snow replicas. *In situ* meteorological data obtained at Detroit Plateau evidenced that summer temperatures do not reach the melting point.

Cultivation, Isolation and Sequencing of Microorganisms

All cultures were incubated in dark condition, without shaking at 37°C and 16°C for 120 days. Samples were melted at 4°C during a 6 hours period. Aliquots of 20 mL of each sample was taken for culture assays. Two different liquid media were used: Brain Heart Infusion (BHI) and Tioglicolate (TGI). The media were distributed in test tubes (5 mL per tube) and 2x concentrated in order to get the normal concentration after the sample inoculation. To the TGI tubes we added 2.5 mL of mineral oil before sterilization process. A volume of 5 mL of each sample was inoculated in two BHI tubes and two TGI tubes (for different incubation temperatures). After inoculation, BHI cultures were aired in hard shaker procedure for 2 minutes, while in TGI tubes the samples were gently added directly in the media with a pipette thought the oil layer.

Isolate samples were submitted to standard protocol for DNA extraction and purification with the GFX Genomic Blood DNA Purification





(Amersham Biosciences). The V3 hypervariable region of the 16S rRNA were amplified and sequenced using the primers 27F (5'AGA GTT TGA TCM TGG CTC AG 3'), (Lane, 1991) and L4101r (GCG TGT GTA CAA GAC CC) (Nudel et al. 1996). The PCR profile consisted of 5 min at 95°C. 30 cycles of 1 sec at 95°, 1 min at 60°C, and 2 min at 72°C, with a final extension step for 10 min at 72°C. Sanger sequencing reactions were performed at the MegaBACE 1000 (Amersham Biosciences) platform with the use of the DYEnamic ET Dye Terminator Kit (with the Thermo Sequenase™ II DNA Polimerase) , with 25 cycles of 10 sec at 94°C, 5 sec at 50°C and 4 min at 60°C, and the use of a internal primer 338F following Muyzer et al. 1993, 1995). All chromatograms were visually inspected for accuracy and to minimize missing data using the software Geneious v4.82. After the initial screening at the NCBI database using the BLASTn search algorithm, a Neighbor-Joining analysis was conducted with 1.000 bootstrap replicates. The chosen nucleotide substitution model for the dataset was the TN93 based on the Akaike criterion.

Denaturing Gradient Gel Electrophoresis (DGGE)

The PCR products were analyzed by denaturing gradient gel electrophoresis (DGGE) using the standard BioRad protocol with the DCode[™] Universal Mutation Detection System. The amplified fragments for the V3 region of the 16S rRNA were analyzed in a vertical polyacrylamide gel (8%) in a 1XTAE buffer, and a sustained 65 °C during the entire electrophoresis.

Flow Cytometry of Snow Samples

Aliquots of 100 mL from each sample were concentrated by freeze drying and re-suspended in 5 mL of PBS 1x pH 7.2. The nucleic acidspecific fluorochrome SYTO 13 green fluorescent nucleic acid stain in 5 mM solution in DMSO (Molecular Probes, Eugene, OR) was used in all experiments. The samples were stained with 20 μ M of SYTO 13, vortex to mix then incubate for 1-30 min in the dark at room temperature. The absorption wavelength of SYTO 13 was 488 nm and its emission was 509 nm (Guindulain et al. 1997) All the experiments used a single green fluorescent detector (FL1). Five thousand events were acquired in which sample. The samples were analyzed in an EPICS ALTRA flow cytometer (Beckman Coulter Inc., Hialeah, FL) and analyzed in Expo 32 software (Beckman Coulter).

Elemental Composition Analysis

Aliguots of 400 mL from each 20 cm of Pit I were concentrated by freeze drying process and resuspended in 10 mL of milliQ[™] water followed by filtration in 0.1 m Nuclepore™ filter. The filtered material was submitted to a Particle Induced X-Ray Emission (PIXE) analysis, owing the concentration of terrigenous elements (Ca, Ti and Fe), previously employed to track continental air masses at the studied site. In this technique, the filter is exposed to a 2 MeV proton flux bean generated by a Van De Graaff linear particle accelerator, installed at the Institute of Physics of Pontifícia Universidade Católica (PUC-RIO). Details of PIXE technique can be found at De Pinho et al. 1979. Metal analysis of Ca, Ti and Fe refer to King George Island samples. For Detroit Plateau aerosol samples were collected at daily resolution and analyzed in nuclepore filters (47 mm diameter and 0.4 m porosity), using an air sampling flux rate of 12 Lpm. Aerosol composition was based on S, Na, Fe and Al elements.

Isotopic analysis

Isotopes in water precipitation have proven to be a tracer in studies of the interaction between air masses and synoptic climatology, especially in terms of origin of the water vapor. The spatial distribution of global d¹⁸O, as provided by the Global Network of Isotopes in Precipitation (GNIP)/IAEA (accessible at http://isohis. *iaea.org*), show an isotopic signature of d¹⁸O ranging from -15 ‰ to -3.0 ‰ for the tropical to temperate sites of the South Hemisphere and values corresponding to -15 ‰ to -30 ‰ for polar air masses. Therefore d¹⁸O in snow may tag the presence of warmer or colder incursions of air masses at the Maritime Antarctica where our work was developed.

The device used for d¹⁸O analysis was an automatic interface of the DeltaPlus Advantage gas source mass spectrometer of the Thermo Finningan™ brand. A fraction of each liquid sample, referring to a 10 cm snow layer was processed for the analysis. The international standards used were: V-SMOW, IAEA-SLAP2 and IAEA-GISP2. d¹⁸O analysis refer to Detroit Plateau samples only.

Atmospheric transport model

In order to track the air mass past migrations towards Antarctica we have used the HYSPLIT (Hybrid Single Particle Lagrangian Integrated Trajectory) model. The HYSPLIT model is an atmospheric trajectory modeling tool widely used to help understand atmospheric transport, dispersion and deposition of mineral dust and pollutants. HYSPLIT is a platform developed by ARL (Air Resources Laboratory), belonging to NOAA (National Oceanic and Atmospheric Administration) and available online (http:// ready.arl.noaa.gov/HYSPLIT.php). The method of calculating the model is a hybrid between the Lagrange approach (using a mobile reference for advection and diffusion calculations as the trajectories move from their initial location) and the Eulerian methodology (which uses a fixed three-dimensional grid as a frame reference points for calculating pollutant concentrations in the air). The model output depicts trajectories of the center of the air masses backwards in time and allow inferring the air mass transit before reaching the study site.

RESULTS AND DISCUSSION

Identification of microorganisms for King George Island

From 20 inoculated snow samples from King George Island, bacterial growth was observed in 10 (3 only on BHI incubated at 37 °C, 1 only on BHI at 16 °C, 2 only on TGI at 16 °C, 1 on BHI and TGI 16 °C, 2 on BHI 16 °C and 37 °C; 1 on BHI and TGI 16 °C and 37 °C. All cultivated microorganisms were identified as gram-positive bacteria, and more than 80% of those were spore-forming *Bacillus*. They were isolated and submitted to **molecular sequencing** techniques which results are presented in Figure 3 and Table I. The resultant Neighbor-joining tree (Figure 3) show the isolates 1, 10, 4, and 3 recovered nested within the *Bacillus cereus* group, isolates 3 and 12 within the *Bacillus subtilis* group, the isolate 7 recovered nested within the *Paenibacillus* group, and the isolate 9, recovered in a group together with *Terribacillus goriensis* and *Virgibacillus picturae*.

Among the sequenced isolates, two of them were identified to the species level (*Bacillus*) and the remaining to the genus level (*Bacillus* and *Paenibacillus*). Among the *Bacillus* isolates, *Bacillus cereus* group if formed by about 11 closely related species with contributions for production of numerous enzymes (Chang et al. 2007), metabolites (Kevany et al. 2009),



0.06 0.05 0.04 0.03 0.02 0.01 0.00

removal of various heavy metals (Chen et al. 2016), persistent organic pollutants (Kazunga & Aitken 2000), and as probiotics for humans and animals (Gisbert et al. 2013). The *B. cereus* bacteria group occupy several habitats ranging from terrestrial and aquatic environments, and also including plants and animals, and presents strong survivability of spores allowing them to better withstand hostile conditions and to better disperse (Guinebretiere et al. 2008, Jensen et al. 2003).

The second isolated and identified *Bacillus* species, *B. subtilis* can be isolated from many environments such as terrestrial and aquatic. The species is ubiquitous and broadly adapted to grow in diverse settings within the biosphere. In response to nutrient deprivation, B. subtilis can form highly resistant dormant endospores (Sonenshein et al. 2002, Ricca et al. 2004). These spores could easily disperse by wind (Merrill et al. 2006, Jaenicke 2005) and migrate long distances. B. subtilis is also referred to as a soil dweller, likely presenting a saprophytic life history (Vilain et al. 2006). B. subtilis can also grow in close association with plant roots, and also within the gastrointestinal tract of several animals (Tam et al. 2006, Leser et al. 2008, Hong et al. 2005, Inatsu et al. 2006). In marine environments, although growth might occur, the abundance seems to be related with its observed association with the gastrointestinal tract of marine organisms (Newaj-Fyzul et al. 2007) and other biotic surfaces (Ivanova et al. 1999). Current results indicate that the apparent ubiquity of *B. subtilis* seems to be not only a result of spore persistence in these environments since *B. subtilis* seems to actively grow in diverse environments including ranging from soils, water, gastrointestinal tracts, plant roots.

The third identified isolate is related with *Paenibacillus*. The genus *Paenibacillus* (Ash et al. 1993) was created to receive the former 'group

3' species of the genus *Bacillus*. It includes about 35 species of facultative anaerobes and endospore-forming, neutrophilic, periflagellated heterotrophic, low G+C gram-positive bacilli. *Paenibacillus* species were recorded from different environments such as soils, roots, rhizosphere of various crop plants including wheat, sorghum, sugarcane, maize, barley (Guemouri-Athmani et al. 2000, von der Weid et al. 2000), trees such as pines (Holl and Chanway, 1992, Shishido et al. 1996), and also marine sediments (Ravi et al. 2007).

Yet those bacteria, such as B. subtilis or B. cereus, can be found in many different substrata and conditions, they do not have specific mechanisms to allow them to maintain a normal metabolism and cell division with respect to the relative low temperatures as those of the polar environments. The presence of those non-natural cold environmental flora microorganisms, and the long time in incubation (up to 60 days in some cases), contrasting with 24 hours of growth, after the re-inoculation, suggests its exogenous origin. After a long period deposited on ice, the cells and the spores remained viable but far from its optimum metabolic conditions, explaining their long time to recover in incubation.

Microorganism Counting, DGGE, and Elemental analysis for King George Island

Counting rate obtained by the flow cytometry showed a variable cell distribution with a pronounced peak at the 60-80 cm layer and two secondary peaks, at 20-40 cm and 120-140 cm, Figure 4. The major peak being four-fold higher than the remaining data. The concentration of unmarked particles and DNA marked cells had a similar variation for the analyzed layers. Of special interest was the data provided by the PIXE analysis which showed an impressive increase of insoluble inorganic elements (Fe, Ti

Isolate ID	Size (BP)	GenBank	Homology (%)
1	1310	Bacillus cereus	1304/1304 (100%)
2	503	Bacillus sp.	473/485 (97.5%)
3	1284	Bacillus subtilis	1278/1278 (100%)
4	874	Bacillus cereus	872/874 (99%)
7	1282	Paenibacillus sp.	1282/1285 (99.7%)
9	939	Bacillus sp.	934/939 (99.4%)
10	1289	Bacillus cereus	1289/1289 (100%)
12	1281	Bacillus subtilis	1281/1281 (100%)

Table I. Genetic identification of the isolated bacilli. The table presents the name of the isolate, the size in base pairs of the analyzed sequence, the GenBank classification for the sequence and its homology within the previous published strains.

and Ca) between 60-100 cm snow layers, and therefore enclosing the peak of microorganisms. This nearly concomitant peak strongly suggests that mineral particles and microorganisms were transported in the same synoptic event to the Antarctic region. Such association between alfresco bacterial communities and resuspended soil dust was previously reported and detailed in previous studied (Savoie et al. 1989, Li et al. 1996, Prospero 1996, Prospero & Lamb 2003, Griffin et al. 2001, 2003). Lighthart & Stetzenbach (1994) reported that bacteria and fungi can be transported by wind for long distances and remaining viable over normal environmental conditionings. Lighthart & Shaffer (1997) pointed out to the importance of aerosol droplet/particle in the maintenance of cell viability along an atmospheric transport.

As an attempt to investigate the atmospheric event associated with the peaks of microorganisms and crustal elements, we combined a set of information: *in situ* meteorological data (wind direction), satellite image galleries around latitude 60°S and the air masses back-trajectories from the HYSPLIT model. A detailed study of Evangelista (1999) concerning individual atmospheric transport events between South America and the Antarctic Peninsula stressed the high association between the meridional crustal material transport towards Antarctica and the northern wind component. On basis of this previous result, we have overlaid microorganism counting, the north wind component and crustal elemental concentrations assuming a uniform snow deposition chronology. The result shows a consistent association among the microbiological, meteorological and geochemical parameters.

According to the snow pit chronology, the peak of microorganisms occurs at middle winter, which confirms the hypothesis of atmospheric transport and deposition, during that season contamination due to weathering of ice-free areas by surface winds, presence of polar animals, and human occupation get their minimum values. Additionally, our DGGE results also suggest high diversity on layers from 100-140 cm. The strata correspond to March-April period, at the end of the Antarctic summer of 2004, when the Antarctic ice cap is not entirely formed and the resuspension of the local microbiota is expected due to the strong and constant surface winds.

Within this period, we have selected 215 satellite images from GOES satellite in ftp://



Figure 4. Microorganism counting from a shallow snow pit in Wanda Glacier (a) and relative north wind component at King George Island (according to snow deposition chronology); crustal elements Ca, Ti, Fe concentrations at same samples (b) (BDL: bellow detection limit); DGGE polyacrylamide gel (8%) obtained from the pit I at Wanda Glacier. The lanes correspond to sections: (2) 20-40 cm – 11 bands; (3) 60-80 cm –9 bands; (4) 80-100 cm – not detected; (5) 100-120 cm – 15 bands; (6) 120-140 cm – 10 bands; C+ (positive control). Each observed band is marked by a black arrow.

ftpantartica.cptec.inpe.br/pub/antartica/ GOES SubA/ at the Infra-Red and visible bands and we have run the HYSPLIT back-trajectories model for each day. From the image gallery and the atmospheric transport model, we have recognized events during July 2014 in which air mass trajectories previously have passed through the South American continent before reaching King George Island. One may observe that this period is coincident with the increased north wind component observed in the Meteorological Station at King George Island, Figure 4. In that circumstance, the air mass could be enriched by dust and microorganisms during their continental migration. From this method we have selected three potential events attending the above characteristics. Figure 5

depicts simultaneous GOES images and the HYSPLIT back-trajectories model output for three advective episodes linking South America and Antarctica Peninsula.

From Figures 4 and 5, it is clear the correspondence among the parameters used to track the atmospheric transport. The three events of southwards air mass advection originated in the meridional Patagonia semi-desert sector, may explain consistently the elevated concentrations of crustal elements found in the snow samples. Dust plume spread over the Southern Ocean has been detailed described from satellite measurements of MODIS and OMI/ NASA indicating the large influence of Patagonia to the delivery of dust to Sub-polar sites (Gasso & Stein 2007) and its relevance to the ocean



Figure 5. Satellite images and the HYSPLIT back-trajectories for 3 southwards air mass advection episodes between South America and Antarctic Peninsula. Episode (A1,B1) occurred in 8th July, 2004; (A2,B2) in 19th July, 2004; (A3,B3) in 22nd July, 2004. Small red dot is the location of King George Island. The superimposed arrows were defined on basis of corresponding synoptic charts analysis for the same days.

primary productivity through Fe deposition (Erickson et al. 2003). Although limited to one sequence of episodes of transport, this work provides an initial basis for further investigation in deeper ice core of that region, which could bring additional knowledge on microorganisms apportioning polar regions.

Microorganism counting and Isotope analysis for Detroit Plateau

During the Detroit Plateau mission, we collected aerosols *in situ* in order to characterize their behavior at high altitudes and its relation with the atmospheric transport. We sampled during 6 different days integrated for 24 hours. Elemental composition was based on Na and S as representative of marine influence and Fe and Al as representative of crustal mineral material. The results present in Figure 6 show that concentration of crustal material is closely related to the nature of the air mass. The episodes of 3, 4 and 5th December 2007 presented significantly higher concentrations of Al and Fe coincident to air masses which migration history is associated to the Southern Patagonia, evidencing a similar mechanism as observed to King George Island.

With respect the Detroit plateau 2m-snow pit. microorganism counts and corresponding oxygen isotopic data covariate well and corroborated the concept of atmospheric transport. Stable isotopes of oxygen, d¹⁸O, in precipitation is influenced by a combination of factors, such as precipitation amount, air temperature, moisture source, altitude and vapor transportation. The northern Antarctic Peninsula is particularly complex in the meteorological and synoptical point of view, since it is a place where exist multiple influences of different air masses (Polar. Southern Pacific and Southern American). Data of d¹⁸O in the snow pit of Detroit plateau varied from -14‰ to -27‰. Microorganism counts varied in a very close pattern as d¹⁸O, Figure 7. Increases of microorganism counts occurred in layers where d¹⁸O values were representative of lower latitudes in the Southern South America



Figure 6. Elemental composition for aerosols at Detroit Plateau during the 2007 campaign and associated air mass backward trajectories. "Terrigenous influence" refers to the migration of the air mass over continental land of South America.

while values bellow -20 ‰ reflect the polar influence in the region. Interesting to observe is that when the polar influence is present, microorganism counts is reduced by a factor between 2.5 and 3.0 probably indicating that the (sub)polar environment can be also a source of microorganism and constitute a background level at that region.

CONCLUSIONS

In this work we explored the use of molecular identification and counting in snow/firn layers of Antarctic ice sheets, by flow cytometry, that is a simple and technological facility well available in several laboratories around the world. We developed simultaneous measurements of microorganisms and other geochemical tracers successfully used in the literature as the elemental composition, representing tracers of crustal material based on Fe, Ti, Al and Ca, and



Figure 7. (left) Microorganism counts and d¹⁸O. Colored parts of d¹⁸O curve indicates possible warm air advection from lower latitudes; (right) average d¹⁸O in precipitation from AIEA/GNIP for January (summer) and July (winter) periods.

stable isotopes of oxygen. We found satisfactory associations between microorganisms and the geochemical tracers that were confirmed by atmospheric dispersion models and satellite images. We conclude that same events that transport dust to Antarctica are source of microorganisms that are probably associated to dust resuspension in the continents around Antarctica, especially South America. Therefore, microorganism counting in ice sheet layers can act as tracer of the continental terrigenous influence, their desertification evolution and wind dynamics between the continents.

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