Elemental and Mineralogical Composition of the Western Andean Snow (18°S–41°S)

Juan A. Alfonso1,2, Raul R. Cordero1, Penny M. Rowe1,3, Steven Neshyba4, Gino Casassa5,6, Jorge Carrasco6, Shelley MacDonell7, Fabrice Lambert8,9, Jaime Pizarro1, Francisco Fernandoy10, Sarah Feron1,11, Alessandro Damiani1,12, Pedro Llanillo1, Edgardo Sepulveda1, Jose Jorquera1, Belkis Garcia2, Juan M. Carrera2, Pedro Oyola13 & Choong-Min Kang14

The snowpack is an important source of water for many Andean communities. Because of its importance, elemental and mineralogical composition analysis of the Andean snow is a worthwhile effort. In this study, we conducted a chemical composition analysis (major and trace elements, mineralogy, and chemical enrichment) of surface snow sampled at 21 sites across a transect of about 2,500 km in the Chilean Andes (18–41°S). Our results enabled us to identify five depositional environments: (i) sites 1–3 (in the Atacama Desert, 18–26°S) with relatively high concentrations of metals, high abundance of quartz and low presence of arsenates, (ii) sites 4–8 (in northern Chile, 29–32°S) with relatively high abundance of quartz and low presence of metals and arsenates, (iii) sites 9–12 (in central Chile, 33–35°S) with anthropogenic enrichment of metals, relatively high values of quartz and low abundance of arsenates, (iv) sites 13–14 (also in central Chile, 35–37°S) with relatively high values of quartz and low presence of metals and arsenates, and (v) sites 15–21 (in southern Chile, 37–41°S) with relatively high abundance of arsenates and low presence of metals and quartz. We found significant anthropogenic enrichment at sites close to Santiago (a major city of 6 million inhabitants) and in the Atacama Desert (that hosts several major copper mines).

Elemental and mineralogical composition of snow allows detecting deposition of pollutants and particulate matter. Several studies have conducted chemical composition analysis of snow samples in Europe1–5, North America6,7 and Asia8–11. However, efforts that have targeted the Andes12–14 have been mostly conducted in snow on glacier surfaces at few points. Further analysis are needed, especially at snow-covered areas close to Santiago, Chile, a city of 6 million inhabitants (~33°S), with high levels of urban pollution and aerosols15,16.

Multiple climate regions can be identified along the Andes17–19. In our study area, the climate is modulated by the permanent influence of the South Pacific Anticyclonic (SPA) circulation that causes drier conditions in the Atacama Desert (18–32°S)20. The prevalent westerly circulation and the seasonal displacement of the SPA cause drier conditions in summer and wet conditions in winter in northern Chile (29–32°S), as well as in central Chile (33–37°S). Mountain snow precipitation associated with passing frontal systems (and cut-off lows) occurs mainly during the wet season (mid-April to September)20. Westerly circulation prevails in southern Chile (37–41°S) where frontal weather systems travel eastward generating snow precipitation as they cross the Andes21.

1Universidad de Santiago, Av. B. O’Higgins 3363, Santiago, Chile. 2Instituto Venezolano de Investigaciones Científicas (IVIC), Apartado 20632, Caracas, 20632, Venezuela. 3NorthWest Research Associates, Redmond, USA. 4Department of Chemistry, University of Puget Sound, Tacoma, USA. 5Unidad de Glaciología y Nieves, Ministerio de Obras Públicas, Santiago, Chile. 6Centro GAIA Antártica, Universidad de Magallanes, Punta Arenas, Chile. 7Centro de Estudios Avanzados en Zonas Áridas (CEAZA), La Serena, Chile. 8Department of Physical Geography, Pontificia Universidad Católica de Chile, Santiago, Chile. 9Center for Climate and Resilience Research, Universidad de Chile, Santiago, Chile. 10Universidad Nacional Andrés Bello, Viña del Mar, Chile. 11School of Earth, Energy and Environmental Sciences, Stanford University, Stanford, USA. 12Center for Environmental Remote Sensing, Chiba University, Chiba, Japan. 13Centro Mario Molina, Antonio Belle 292, Santiago, Chile. 14Harvard School of Public Health (HSPH), Boston, Massachusetts, USA. Correspondence and requests for materials should be addressed to R.R.C. (email: raul.cordero@usach.cl)
The snowpack is an important source of water for many Andean communities in central Chile (33–37°S), where also agriculture and power generation depend on the Andean streams22. The snowpack in central and southern Chile (33–41°S) has been affected by mid-tropospheric warming23 as well as by a persistent drought in 2010–201524 that has resulted in decreasing river flows25. In addition to anomalies in precipitation, the Andean snowpack (18–41°S) has been impacted by particulate matter transported by near-surface winds flowing from the valleys west of the Andes towards the mountains26,27. The prevailing circulation affecting the high Andean mountains is mainly westerly airflow, ranging from northwesterly to southwestery direction28. The subsidence inversion (strengthened by the SPA and the local topography) confines the particles within the boundary layer in the Atacama Desert and central Chile. However, the same factors lead to near-surface local circulations (i.e. mountain-valley breeze), which enable particle transport/dispersion from sources (such as major mines) that operate over the Andes27.

Emissions from transport, industrial pollution and residential heating in Santiago (~33°S) have been found to affect the Andean snowpack14,28. However, the extent of anthropogenic enrichment in the Andean snowpack remains to be thoroughly tested. In the case of the Atacama Desert and central Chile, potential sources affecting the Andean snow include the mining industry29; Chile is the top producer of copper and molybdenum; it accounts for approximately one third of the global copper production while ranks third in molybdenum production30. In the case of southern Chile, impurities in the snow may result from wood stoves (the preferred heating system for the population in that area)29 as well as from volcanic ash deposition30. Indeed, dozen of volcanoes are active in the Chilean Andes31,32. Because of the importance of the Andean snow for Chile, elemental and mineralogical composition analysis of the snowpack is a worthwhile effort.

In this paper, we report on the chemical composition analysis (major and trace elements, mineralogy, and chemical enrichment) of surface snow sampled during consecutive Austral winters (2015 and 2016) across a transect of about 2,500 km in the Chilean Andes (18–41°S). By combining the elemental and mineralogical composition of the snow samples, we were able to identify the five depositional environments shown in Fig. 1: (i) sites 1–3 (in the Atacama Desert) with relatively high presence of metals and quartz and low abundance of arsenates, (ii) sites 4–8 (in northern Chile,) with relatively high abundance of quartz and low presence of metals and arsenates, (iii) sites 9–12 (in central Chile) with anthropogenic enrichment of metals, relatively high values of quartz and low abundance of arsenates, (iv) sites 13–14 (also in central Chile) with relatively high values of quartz and low presence of metals and arsenates, and (v) sites 15–21 (in southern Chile) with relatively high abundance of arsenates and low presence of metals and quartz. As explained below, modest anthropogenic influence found in northern Chile and southern Chile suggests that element concentrations measured at sites in these zones may be useful as baselines for future studies.

**Material and Methods**

**Sampling.** Sampling was conducted during consecutive Austral winters (2015 and 2016) at 21 sites across a transect in the Chilean Andes, from Putre to Volcán Osorno; see Table 1 and Fig. 1 for further details. Sampling was carried out at the end of the accumulation season. Sampling sites were selected to represent broad regions at locations with no known local aerosol sources. At each site, two samples from the surface layer (up to 15–20 cm depth) and separated by 1–2 m distance were collected. Samples ranged from 1 to 2 kg each.

Although some additional samples were collected from the subsurface layer, we focused on the surface layer since prior efforts have shown consolidation of pollutants in the surface layer with snowmelt31. Indeed, prior efforts at the same sampling sites allowed us to verify that subsurface layers tend to be cleaner than the surface26. Meteorological data suggest a lack of snowfall for at least five days before the sampling at most sampling sites. At sites 3 and 4, cold fronts brought snowfall to the area immediately prior to sampling. Although in these cases the surface layer was cleaner than the subsurface layer, element concentrations were found to be comparable (of the same order).

Researches conducting sampling in the field took care to prevent sample contamination. As also explained by Rowe et al.26, a stainless steel spatula was used and samples were placed into plastic bags, in turn packed in Whirlpack bags. Styrofoam coolers were used for transportation to the laboratory. In the laboratory deionised water was used for washing tools and containers before filtering, while latex gloves and lab coats were worn during filtering. Snow samples were transferred to a glass beaker and snowmelt was thereafter vacuum-filtered, leaving the insoluble material on the filter. Stainless-steel funnels and 0.4-µm Nucleopore filters were used. Filters were afterwards placed in sterile petri dishes.

**Chemical composition.** Elemental composition of the insoluble material in the snow samples was carried out using Energy dispersive X-ray fluorescence (EDXRF) spectrometry. EDXRF is a technique widely used for elemental analysis of particles on filter media34,35. Elemental analysis was conducted at the Harvard School of Public Health (HSPH) using an Epsilon 5 EDXRF spectrometer (PANalytical, The Netherlands). The procedure at HSPH for analysing elements on filters is described by Kang et al.35; it involves using 49 MicroMatter XRF calibration standard polycarbonate films (Micromatter Co., Vancouver, Canada) spanning from Na (atomic number 11) to Pb (atomic number 82). When intercomparing measurements of standard films, the HSPH EDXRF spectrometer has exhibited good performance: “2% errors in precision (suggesting a good reproducibility) and 4% errors in accuracy (showing a proper calibration)”35. In the case of the lower concentrations associated with field samples, the HSPH EDXRF spectrometer has exhibited a reproducibility within 15% for most of the elements35. For this study, concentrations of Al, Mg, Si, P, Ca, Ti, S, K, V, Cr, Mn, Ni, Cu, Fe, Co, Zn, Rb, Sr, Zr, Mo, Ba and Pb were determined to be significant.

Mineralogy was assessed using X-ray powder diffraction (XRD)36. A Siemens D-5005 x-ray diffractometer fitted with parallel beam geometry and Cu Kα radiation, was applied. XRD patterns were collected every 0.02°,
from 5° to 90°. Minerals were identified at the Instituto Venezolano de Investigaciones Científicas (IVIC) from their characteristic peaks.

**Statistical analysis.** Cluster analysis and principal components analysis (PCA) were used to assess element assemblages. Before applying the multivariate statistical analysis, concentrations rendered by EDXRF were log-transformed and standardized (by subtracting the mean and dividing the outcome by the standard deviation)\(^37\). An orthogonal method (the varimax method) was applied for factor rotation\(^37\). We evaluated the following cluster analysis methods: Ward, complete linkage, single linkage, weighted pair-group average, unweighted pair-group average and unweighted pair-group centroid. STATISTICA\(^38\) was used to conduct the calculations.

In order to detect relative contributions of anthropogenic sources, we applied Crustal enrichment factors (EF\(_{\text{UCC}}\))\(^3\),\(^36\)–\(^4\). EF\(_{\text{UCC}}\) was taken as the ratio between the concentration of an element and the concentration an element originated from rocks and/or dust. As pointed out by Cantonati et al.\(^4\) and Veyssye et al.\(^4\), enrichment factors within the range 0.1–10 allow discarding the significant presence of elements other than those from rocks and dust, while EF\(_{\text{UCC}}\) values higher than 10 do suggest contributions from different sources (natural and/or anthropogenic); EF\(_{\text{UCC}}\) values within the range 10–500 (moderate enrichment) indicate input sources additional to crustal material, while EF\(_{\text{UCC}}\) values higher than 500 (high enrichment) suggest pronounced anthropogenic contribution\(^4\). In this study, EF\(_{\text{UCC}}\) values were calculated for each sampling site. In addition, a second technique for estimating metal enrichments was applied. It implied normalizing the metal concentrations against an element of detrital origin\(^4\). Following prior efforts\(^4\)–\(^6\), we normalized the element concentrations against Al.

**Back-trajectory computation.** Air-parcel backward trajectories were applied to sampling sites where we found anthropogenic enrichment. We used the Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) model\(^47\)–\(^49\) fed with the Global Data Assimilation System (GDAS) archive (available in a resolution of three hours and a 1° latitude-longitude)\(^50\). At each site we computed 36-h backward trajectories by using

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**Figure 1.** (a) Sampling Sites. Numbers identify the 21 sampling sites indicated in Table 1. Color code indicates the five different depositional environments identified in this study. (b) Sampling sites 9–11 nearby Santiago (33°27′S; –70°41′W); dotted line indicates the metropolitan area of this major city. (c) Elevation along latitude 33°27′S. Plots generated by using PYTHON’s Matplotlib Library (https://matplotlib.org)\(^62\).
15–21 (in southern Chile, 37–41°S). Dendograms from the cluster analysis for other linkage methods (the unweighted pair-group average method) consistently suggest the same two groups shown in Fig. 3.

(ii) group 2 consists of sites 4–8 (in northern Chile, 29–32°S), sites 13–14 (in central Chile, 35–37°S), and sites 15–21 (in southern Chile, 37–41°S). Dendograms from the cluster analysis for other linkage methods (the unweighted pair-group average method, Two groups are identified: (i) group 1 consists of sites 1–3 (in the northern Atacama Desert, 18–26°S), sites 9–11 (relatively close to Santiago, ~33°S) and site 12 (close to Curicó, ~35°S).

Statistical analysis. Figure 3 shows the dendrogram obtained from the cluster analysis (of elemental concentrations) using the unweighted pair-group average method. Two groups are identified: (i) group 1 consists of sites 1–3 (in the northern Atacama Desert, 18–26°S), sites 9–11 (relatively close to Santiago, ~33°S) and site 12 (close to Curicó, ~35°S); (ii) group 2 consists of sites 4–8 (in northern Chile, 29–32°S), sites 13–14 (in central Chile, 35–37°S), and sites 15–21 (in southern Chile, 37–41°S). Dendograms from the cluster analysis for other linkage methods (the single linkage method, complete linkage method, unweighted pair-group centroid method, Ward’s method, and weighted pair-group average method) consistently suggest the same two groups shown in Fig. 3.

Table 1. Sampling sites.

| Sites     | Location      | Latitude | Longitude | Elevation (m) | Sampling date |
|-----------|---------------|----------|-----------|---------------|--------------|
| 1 Putre   | S 18°06'08.5" | W 69°31'14.3" | 5318 | 2015/07/05    |
| 2 Cerro Toco | S 22°57'09.4" | W 67°46'38.1" | 5370 | 2015/07/10    |
| 3 La Ola  | S 26°27'17.1" | W 69°03'07.2" | 3578 | 2015/07/14    |
| 4 Valle del Elqui | S 29°20'42.6" | W 70°04'17.9" | 2341 | 2015/07/17    |
| 5 La Ramada | S 30°57'38.1" | W 70°33'21.2" | 1757 | 2015/07/19    |
| 6 Valle Choapa | S 32°03'49.2" | W 70°35'42.4" | 1919 | 2015/07/20    |
| 7 Porilllo | S 32°50'1.4"  | W 70°79'9.2"  | 2800 | 2015/07/21    |
| 8 Los Hornos | S 32°53'32.2" | W 70°12'64.1" | 2322 | 2015/07/22    |
| 9 Valle de Maipo | S 33°39'45.8" | W 70°15'35.0" | 2302 | 2015/07/26    |
| 10 Valle Nevado | S 33°21'58.3" | W 70°15'16.3" | 2635 | 2015/07/24    |
| 11 La Parva | S 33°20'06"   | W 70°18'00.2"  | 2610 | 2016/07/25    |
| 12 Curicó  | S 35°08'10.8" | W 70°28'45.2" | 1860 | 2016/08/19    |
| 13 Maule   | S 35°59'19.7" | W 70°33'45.1" | 1860 | 2016/08/20    |
| 14 Chillán | S 36°54'26.4" | W 71°23'59.5" | 1963 | 2016/07/23    |
| 15 Antuco  | S 37°22'04.6" | W 71°22'57.3" | 1494 | 2016/08/23    |
| 16 Volcán Collaqui | S 37°52'35.1" | W 71°28'26.9" | 1142 | 2016/08/24    |
| 17 Corralco | S 38°24'21.1" | W 71°33'34.3" | 1601 | 2016/08/27    |
| 18 Volcán Llaima | S 38°31'37.8" | W 71°47'44.5" | 1830 | 2016/08/28    |
| 19 Volcán Villarrica | S 39°23'46.9" | W 71°57'53.0" | 1450 | 2016/08/29    |
| 20 Antillanca | S 40°47'12.5" | W 72°11'31.5" | 1349 | 2016/07/19    |
| 21 Volcán Osorno | S 41°07'12.0" | W 72°31'47.2" | 1326 | 2016/08/31    |

meteorological data over a 92-day period prior to the sampling date (see Table 1). Over that period, trajectories were computed every 3 hours, making a total of 736 trajectories per site. These trajectories were divided into clusters following a cluster analysis based on the total spatial variance (TSV)25.

Results

Chemical composition. Figure 2 shows the boxplots of the element concentrations (ng of the element per g of snow) in the snow samples collected in the five depositional environments indicated in Fig. 1. As shown in Fig. 2, for all sets, either Si or Fe exhibited the highest concentration followed in general by K or Ca, Al, Ti and Mg. Figure 2 also allows realizing the dispersion of element concentrations found in Andean snow (18–41°S). It can be observed that the concentrations of Co, Cu, Rb, Ba and V ranged within three orders of magnitude. Concentrations of Mg, P, K, Ni, Zn, Sr, Zr, Ti, Mn, Mo, Pb and Fe ranged within two orders of magnitude, while the other elements ranged only within one order of magnitude.

Table 2 shows the best estimates of the element concentrations (ng of element per g of snow) in the snow samples at each site while the associated uncertainty are shown in Table 3. We found relatively high concentrations of K, Ti, V, Mn, Fe, Co, Zn, Rb, Sr, Ni, Cu, Zr, Mo, Ba and Pb at site 11 (La Parva, nearby Santiago), while at site 2 (Cerro Toco, in the Atacama Desert) relatively high concentrations of Al and Si were detected.

We found relatively high concentrations of Mg and Ca at site 3 (La Ola, in northern Chile), whereas at site 10 (Valle Nevado, also near Santiago), 14 (Chillán, in central Chile) and 19 (Volcán Villarrica, in southern Chile), relatively high concentrations of Cr, S and P were measured, respectively.

The lowest concentrations of Si, K, Ti, Fe, Co, Ni, Cu, Zn, Sr, Zr and Ba were found at site 20 (Antillanca, in southern Chile), while at site 19 (Volcán Villarrica, also in southern Chile) we detected the lowest concentrations of V, Rb and Pb.

The lowest concentrations of Al and Mo were measured at site 15 (Antuco, central Chile), while at sites 2 (Cerro Toco, in the Atacama Desert), 6 (Valle Choapa, in northern Chile), 8 (Los Hornos, in northern Chile), 13 (Maule, in central Chile), 16 (Volcán Collaqui, in southern Chile) and 21 (Volcán Osorno, also in southern Chile), we found the lowest concentrations of P, Ca, Cr, Mg, Mn and S, respectively. See Table 2 for further details.

Statistical analysis. Cluster analysis and PCA were carried out on the dataset in Table 2 (after normalization and standardization). Table S1 shows the correlation matrix for the element concentrations.

Figure 3 shows the dendrogram obtained from the cluster analysis (of elemental concentrations) using the unweighted pair-group average method. Two groups are identified: (i) group 1 consists of sites 1–3 (in the northern Atacama Desert, 18–26°S), sites 9–11 (relatively close to Santiago, ~33°S) and site 12 (close to Curicó, ~35°S); (ii) group 2 consists of sites 4–8 (in northern Chile, 29–32°S), sites 13–14 (in central Chile, 35–37°S), and sites 15–21 (in southern Chile, 37–41°S). The two groups follow the one observed for the geographical distribution of the sites (Fig. 1).
PCA results are shown in Table 4, while Fig. 4 shows scores and loading. Two principal components accounted for 81% of the total variance. Component 1 (eigenvalue equal to 16.1) presented significant negative loads of Mg, Al, Si, S, K, Ti, V, Cr, Mn, Fe, Co, Ni, Cu, Zn, Rb, Sr, Zr, Mo, Ba and Pb. Component 2 (eigenvalue equal to 1.7) has a high positive load of P. The PCA results (see Fig. 4b) also distinguished the same two groups suggested by the cluster analysis.

Mineralogy. Table 5 shows the bulk mineralogy of the snow samples, as the percentage of each mineral in the insoluble material filtered from snow. The main mineralogical species were: silicates (labradiorite, microcline, albite), hydrated silicates (muscovite, laumontite, K-Fe-Mg-Al hydroxysilicate, biotite), quartz, hydrated sulphates, oxides, hydrated oxides, as well as hydrated arsenates at sites 15–21 (in southern Chile).

Enrichment factors. Table 6 shows the enrichment factors (EFUCC) of elements found in the snow. Mg, Si, P, S, K, Ca, Ti, Sr and Zr showed no significant enrichment (EFUCC < 10) in the snow samples. In the case of V, Cr, Mn, Fe, Co, Ni, Cu, Zn, Rb, Sr, Zr, Mo, Ba and Pb, EFUCC distributions ranged from no enrichment at some sites to moderate enrichment (10 < EFUCC < 500) at others. Cu and Mo presented in general relatively high enrichments, from moderate (10 < EFUCC < 500) to intense (EFUCC > 500). Our results show a high enrichment of Cu at site 2 (Cerro Toco, in the Atacama Desert) and both Cu and Mo at sites 9–11 (close to Santiago).

Figures S1–S5 show the correlation between Al concentration and the concentration of Ti, V, Cr, Mn, Fe, Co, Ni, Cu, Zn, Rb, Sr, Zr, Mo, Ba and Pb. The dashed lines in Figs S1–S5 represent 95% confidence interval. Values found well above these lines represent metals enrichment at a particular site. Again, our results indicate enrichment of Ni, Mn, Ti, V, Co, Cu, Zn, Zr, Ba and Pb at site 10 (Valle Nevado) and at site 11 (La Parva); of Fe, Rb and Mo at sites 9–11 (Valle de Maipo, Valle Nevado, and La Parva); and of Sr and Cr at site 10 (Valle Nevado). These results are consistent with significant enrichments previously reported in the case of snow sampled at Cerro Colorado (in the very same area of our sampling sites 10 and 11). The correlations in Figs S1–S5 also suggest enrichments of Cr and Ni at site 1 (Putre in the Atacama Desert), and of Mn at site 12 (close to Curicó).
Table 3. Best estimates of the element concentrations (ng of the element per g of snow) in the snow samples at each site.

| Sites | u(Mg) (ng/g) | u(Al) (ng/g) | u(Si) (ng/g) | u(P) (ng/g) | u(S) (ng/g) | u(K) (ng/g) | u(Ca) (ng/g) | u(Ti) (ng/g) | u(V) (ng/g) | u(Cr) (ng/g) | u(Mn) (ng/g) | u(Fe) (ng/g) | u(Co) (ng/g) | u(Ni) (ng/g) | u(Cu) (ng/g) | u(Zn) (ng/g) | u(Pb) (ng/g) |
|-------|--------------|--------------|--------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| 1     | 0.290        | 0.777        | 1.425        | 0.016      | 0.025      | 0.014      | 0.101      | 0.099      | 0.041      | 0.062      | 0.012      | 0.030      | 0.422      | 0.009      | 0.010      | 0.033      | 0.014      | 0.025      | 0.037      | 0.040      |
| 2     | 0.267        | 1.021        | 1.696        | 0.010      | 0.035      | 0.010      | 0.024      | 0.149      | 0.057      | 0.005      | 0.009      | 0.048      | 0.007      | 0.008      | 0.007      | 0.076      | 0.019      | 0.013      | 0.026      | 0.029      | 0.035      | 0.258      |
| 3     | 0.248        | 0.505        | 1.138        | 0.007      | 0.008      | 0.006      | 0.073      | 0.186      | 0.035      | 0.003      | 0.004      | 0.021      | 0.005      | 0.007      | 0.005      | 0.018      | 0.010      | 0.007      | 0.013      | 0.015      | 0.015      | 0.019      |
| 4     | 0.133        | 0.232        | 0.311        | 0.005      | 0.007      | 0.004      | 0.025      | 0.029      | 0.011      | 0.002      | 0.004      | 0.012      | 0.162      | 0.003      | 0.003      | 0.011      | 0.006      | 0.005      | 0.008      | 0.012      | 0.014      | 0.097      |
| 5     | 0.115        | 0.161        | 0.195        | 0.007      | 0.006      | 0.005      | 0.016      | 0.030      | 0.009      | 0.002      | 0.005      | 0.123      | 0.004      | 0.003      | 0.013      | 0.007      | 0.005      | 0.008      | 0.014      | 0.016      | 0.108      |
| 6     | 0.079        | 0.388        | 0.826        | 0.009      | 0.010      | 0.004      | 0.010      | 0.023      | 0.006      | 0.031      | 0.037      | 1.420      | 0.010      | 0.006      | 0.011      | 0.004      | 0.003      | 0.006      | 0.008      | 0.010      | 0.067      |
| 7     | 0.063        | 0.462        | 1.405        | 0.007      | 0.014      | 0.003      | 0.081      | 0.197      | 0.077      | 0.003      | 0.003      | 0.012      | 0.047      | 0.003      | 0.004      | 0.008      | 0.014      | 0.008      | 0.013      | 0.013      | 0.018      |

Table 3. Uncertainty associated with the element concentrations shown in Table 2.

Discussion

Possible Sources. As pointed out above, cluster analysis and PCA of the element concentrations in the snow samples suggested two main groups: i) group 1 consists of sites 1–3 (18–26°C) and sites 9–12 (33–35°C); ii) group 2 consists of sites 4–8 (29–32°C), sites 13–14 (35–37°C), and sites 15–21 (37–41°C). Group 1 (sites 1–3 and 9–12) exhibited significantly higher concentrations of Mg, Si, S, K, Ti, V, Cr, Mn, Fe, Co, Ni, Cu, Zn, Rb, Sr, Zr, Mo, Ba and Pb than group 2 (sites 4–8 and 13–21). No significant difference was found for P. Except for Cu and Mo
(see details below), the metal concentrations found in snow samples of group 2 are comparable to metal concentrations (attributed to deposition of background atmospheric aerosols) previously reported for snow samples collected in remote areas in Europe (Eastern Alps, Central Pyrenees, and French Alps), and North America (Alaska).

The two principal components (that explained 81% of the total variance in the PCA) suggest the origin of the elements in each group. Component 1 is characterized by high negative loads of Mg, Al, Si, S, K, Ti, V, Cr, Mn, Fe, Co, Ni, Cu, Zn, Rb, Sr, Zr, Mo, Ba and Pb, while Component 2 exhibits a high positive load of P. Component 1 includes elements that are abundant in the crust (Al, Ti, Mn and Fe), which pointed at terrigenous dust deposition. Moreover, Component 2 includes P, often used in herbicides and pesticides. The position of sites 13 (Maule)
and 17 (Corralco) in Fig. 4b highlights the loading of P in snow samples at these sites. However, the low crustal enrichment factor for P (see Table 6) does not support significant effects of the agricultural activity on the snowpack in the area of Maule and Corralco.

In all of the samples, Cu and Mo exhibited the highest enrichment factors (EFUCC), from moderate to intense. Element concentrations of both Cu and Mo show a very good correlation (Pearson coefficient = 0.98, see Table S1), which suggests that they have a common origin. This result is consistent with the fact that concentrations of hypogene Cu (0.40–0.86%) and Mo (0.01–0.025%) have been reported for the Andes in 18 major deposits (in Chile, Argentina and southern Peru)52. Chile is currently the top producer of copper (accounting for about one third of the global copper production) and ranked third in molybdenum production53.

Figure 5 shows the 36-h back trajectory clusters for sites where we found significant anthropogenic enrichment. Back trajectories were computed every 3 hours using meteorological data over a 92-day period prior to the
sampling date making a total of 736 trajectories for each site. Back trajectories from sites 9–12 were divided into 3 clusters, while trajectories from sites 1–3 were divided into 4 clusters.

Figure 5a,b show the back trajectory clusters for sites 9–11, which are situated less than 20 km east of heavily populated urban areas of Santiago (clusters for Valle Nevado and La Parva are shown in a single plot since these sites are very close each other). Cluster 2 was the largest in Fig. 5a,b. This cluster suggests a heavy influence from Santiago, as a majority of the trajectories (49% in Fig. 5a and 60% in Fig. 5b) originated in or passed over Santiago, and are characterized by being short, with low wind speeds. The second most frequent cluster in Fig. 5a (Fig. 5b) was number three, which contained 27% (21%) of all trajectories. These trajectories typically indicated a source

Table 6. Crustal enrichment factor (EFUCC).

| Sites | Mg | Si | P | S | K | Ca | Ti | V | Cr | Mn | Fe | Co | Ni | Cu | Zn | Rb | Sr | Zr | Mo | Ba | Pb |
|-------|----|----|---|---|---|----|----|---|----|----|----|----|----|----|----|----|----|----|----|----|----|
| 1     | 0.2| 1  | 1 | 3 | 1 | 3  | 3  | 14| 2  | 3  | 15 | 9  | 31 | 6  | 2  | 3  | 1  | 103 | 2 | 5  |
| 2     | 0.2| 1  | 0.1| 4 | 2 | 1  | 3  | 7  | 8  | 4  | 5  | 29 | 6  | 627 | 18 | 3  | 4  | 1  | 399 | 4 | 29 |
| 3     | 0.5| 1  | 1  | 1  | 1  | 2  | 4  | 7  | 8  | 7  | 6  | 37 | 6  | 157 | 22 | 4  | 4  | 2  | 56  | 2 | 14 |
| 4     | 1  | 1  | 0.4| 2 | 1  | 1  | 2  | 3  | 20 | 5  | 4  | 20 | 13 | 31 | 8  | 2  | 1  | 1  | 92  | 1 | 14 |
| 5     | 0.5| 1  | 2  | 2  | 1  | 1  | 3  | 4  | 41 | 5  | 5  | 39 | 22 | 78 | 15 | 1  | 2  | 1  | 80  | 1 | 18 |
| 6     | 0.3| 1  | 1  | 1  | 7  | 1  | 1  | 3  | 5  | 26 | 3  | 4  | 28 | 15 | 493 | 17 | 2  | 2  | 1  | 421 | 1 | 11 |
| 7     | 0.4| 1  | 3  | 1  | 3  | 4  | 3  | 39 | 8  | 5  | 34 | 24 | 186 | 45 | 1 | 2  | 2  | 207 | 4 | 23 |
| 8     | 0.5| 1  | 1  | 2  | 1  | 1  | 3  | 5  | 39 | 7  | 6  | 29 | 21 | 201 | 14 | 0.4 | 2  | 2  | 253 | 0.3 | 22 |
| 9     | 0.3| 1  | 0.5| 6  | 3  | 0  | 4  | 11 | 10 | 7  | 9  | 61 | 6  | 734 | 19 | 6  | 2  | 2  | 596 | 5 | 23 |
| 10    | 0.3| 1  | 0.3| 7  | 2  | 1  | 7  | 25 | 36 | 16 | 21 | 168 | 15 | 2055 | 61 | 13 | 8  | 8  | 1141 | 21 | 74 |
| 11    | 0.3| 1  | 1  | 3  | 2  | 1  | 5  | 15 | 9  | 9  | 12 | 89 | 9  | 1287 | 28 | 7  | 4  | 3  | 636 | 9 | 40 |
| 12    | 0.3| 1  | 1  | 1  | 1  | 4  | 6  | 9  | 9  | 5  | 35 | 4  | 135 | 12 | 2  | 2  | 2  | 186 | 2 | 26 |
| 13    | 0.1| 0.2| 6  | 12 | 1  | 3  | 3  | 4  | 27 | 8  | 4  | 23 | 13 | 57 | 38 | 1 | 2  | 2  | 166 | 3 | 12 |
| 14    | 0.1| 0.5| 1  | 8  | 1  | 1  | 4  | 2  | 13 | 2  | 3  | 13 | 10 | 21 | 8  | 0.4 | 1  | 2  | 66  | 1 | 6 |
| 15    | 0.2| 1  | 0.8| 6  | 1  | 3  | 5  | 4  | 72 | 6  | 4  | 22 | 42 | 161 | 49 | 1 | 2  | 3  | 132 | 0.2 | 24 |
| 16    | 0.2| 0.2| 2  | 4  | 0.4| 1  | 3  | 9  | 28 | 1  | 2  | 5  | 16 | 190 | 20 | 0.1 | 1  | 0.2 | 95  | 7 | 26 |
| 17    | 0.1| 0.4| 2  | 0.4| 0.3| 1  | 5  | 15 | 6  | 2  | 51 | 7  | 39 | 23 | 0.3 | 4  | 2  | 50  | 2 | 21 |
| 18    | 0.4| 1  | 5  | 6  | 1  | 4  | 3  | 5  | 36 | 5  | 4  | 13 | 40 | 79 | 76 | 0.2 | 2  | 3  | 249 | 1 | 56 |
| 19    | 0.1| 0.2| 6  | 4  | 0.3| 2  | 2  | 0.3| 29 | 2  | 1  | 12 | 18 | 30 | 21 | 0.1 | 1  | 0.5 | 180 | 1 | 6 |
| 20    | 0.1| 0.1| 1  | 2  | 0.2| 2  | 1  | 0.5| 24 | 1  | 1  | 11 | 3  | 20 | 6  | 0.2 | 0.4 | 0.4 | 112 | 0.2 | 9 |
| 21    | 0.3| 1  | 1  | 1  | 0.5| 3  | 5  | 7  | 24 | 6  | 5  | 31 | 11 | 23 | 7  | 1  | 3  | 2  | 56  | 10 | 5 |

Figure 5. 36-h back trajectory clusters. (a) Valle Nevado/La Parva; (b) Valle del Maipo; (c) Cerro Toco; (d) Putre; (e) La Ola; (f) Curicó. Plots generated by the using MeteoInfo software tools for meteorological data visualization.63
area in northern Chile but passing close to Los Bronces (a major copper mine). Sites 9 (Valle del Maipo), 10 (Valle Nevada) and 11 (La Parva) are located about 30 km south in the case of site 9 (and about 15 km south in the case of sites 10 and 11) of Los Bronces. Cu and Mo exhibited high enrichment factors (EF_{UCC} > 500) at sites 9–11.

Cu also exhibited a high enrichment factor (EF_{UCC} > 500) at site 2 (Cerro Toco). This site is located in the Atacama Desert, a region whose main economic activity is the copper mining industry. Figure 5c shows that Cluster 2 contained 39% of the trajectories for Cerro Toco. This cluster represented air coming out of the northwest passing close to several major copper mines (Radomiro Tomic, Ministro Hales and Chuquicamata, the latter being one of the largest copper mines in the world). Cerro Toco is located within 150 km of these major copper mines. Cluster 1 also contained 39% of all trajectories for Cerro Toco. These trajectories typically indicated a source area west of Cerro Toco and pass over important astronomical projects (powered by diesel generators) located only few km west of the sampling site.

Figure 5d,e shows the back trajectory clusters for site 1 (Putre) and site 3 (La Ola). Cluster 3 in Fig. 5d, containing 60% of all trajectories, was the largest. The routes for this cluster were typically short and indicated a slow wind speed. The majority of these trajectories originated in the area close to Tacna (a Peruvian city of about 300,000 inhabitants) and to a major gold mine (Pucamarcas) in the Peruvian side of the border. Site 1 is located about 70 km east of Tacna and about 40 km southeast of the Pucamarcas mine. In the case of site 3 (La Ola), Fig. 5e shows that Cluster 1 contained 55% of the trajectories. This cluster represented air coming out of the north. Some trajectories in this cluster originated from nearby a major copper mine (La Escondida), located about 220 km north of the sampling site. The second most frequent cluster in the case of La Ola was number two, which contained 21% of all trajectories. These trajectories typically indicated a source area about 60 km northwest of the sampling site, close to another major copper mine (El Salvador).

Figure 5f shows the clusters for site 12 (Curicó). Cluster 2 was the largest of the 3, containing 58% of the back trajectories. These trajectories were typically characterized by slow air movement, with the trajectories extending about 60 km from the west to the receptor site. Cluster 1 was the second most common path. This cluster contained 41% of the total trajectories. These trajectories were faster and pass over the most populated and industrialized areas in Chile. Still, due to the moderate EF_{UCC} values, an anthropogenic influence can be considered modest at site 12.

Volcanic activity. There are more than one hundred volcanoes in the Chilean Andes that have been active during the Holocene. Several of these volcanoes are currently active. Areas surrounding active volcanoes are subjected to volcanic ash deposition. In active geothermal systems in the southern Andes of Chile, As is present in borehole fluids. In thermal waters in the southern volcanic zone (SVZ), As is transported from geothermal dissolution of aluminosilicate glass dissolute, which is favored by extreme acid conditions; volatile acids (such as and Ca (Na) zeolites, minor quartz was also locally found. The relatively low quartz concentration at sites 15–21 suggests a reduction of amorphous silica by carbon monoxide during the frequent explosive eruptions in the zone.

Summary and Conclusions

We have carried out a chemical composition analysis (major and trace elements, mineralogy, and chemical enrichment) of surface snow sampled across a transect of about 2,500 km in the Chilean Andes (18–41°S).

Cluster analysis and principal component analysis (PCA) applied to the element concentration data, rendered consistent figures suggesting two main groups: (i) group 1 consists of sites 1–3 (in the Atacama Desert, 18–26°S), sites 9–11 (close to Santiago, 33°S) and site 12 (close to Curicó, 35°S); (ii) group 2 consists of sites 4–8 (in northern Chile, 29–32°S), sites 13–14 (in central Chile, 35–37°S), and sites 15–21 (in southern Chile, 37–41°S).

Group 1 exhibited significantly higher concentrations of several metals (Al, Si, Ti, Cr, Fe, Co, Ni, Cu, Zn, and Pb) than group 2. Indeed, with the exception of Cu and Mo, metal concentrations found in snow samples of group 2 are comparable to metal concentrations (attributed to the deposition of background atmospheric aerosols) previously reported for snow samples collected in remote areas elsewhere.

Cu and Mo exhibited the highest enrichment factors (moderate-intense) in all samples. The very good correlation between concentrations of both Cu and Mo is consistent with the fact that major metal deposits in the Andes contain both elements.

Our results confirm significant anthropogenic enrichment (Cu and Mo) at sites near Santiago (sites 9–11). These sites are located about 30 km south in the case of site 9 (and about 15 km south in the case of sites 10 and 11) of Los Bronces, a major copper mine. Anthropogenic enrichment (Cu) was also found at Cerro Toco in the Atacama Desert (site 2), located within 150 km of several major copper mines (Radomiro Tomic, Ministro Hales and Chuquicamata).

Moreover, our results confirm the influence of Santiago (a major city of 6 million inhabitants) on the snow-pack nearby. We found significant enrichment of Ni, Mn, Ti, V, Co, Cu, Zn, Zr, Ba and Pb at site 10 (Valle Nevada) and at site 11 (La Parva); of Fe, Rh and Mo at sites 9–11 (Valle de Maipo, Valle Nevada, and La Parva); and of Sr and Cr at site 10 (Valle Nevada). These results were expected since these sites are located less than 20 km east of heavily populated urban areas of Santiago.
We have also detected the fingerprint of volcanic ash deposition on snow samples taken in southern Chile (37°–41°S), an area with a dozen of active volcanoes. Our results at sites 15–21 exhibited a relatively low quartz concentration and high concentrations of arsenates, which is consistent with prior studies that showed the absence or minor amounts of quartz in ashes.

Our elemental and mineralogical analysis allowed us to identify five depositional environments: (i) sites 1–3 (in the Atacama Desert, 18°–26°S) with relatively high concentrations of metals, high abundance of quartz and low presence of arsenates, (ii) sites 4–8 (in northern Chile, 29°–32°S) with relatively high abundance of quartz and low presence of metals and arsenates, (iii) sites 9–12 (in central Chile, 33°–35°S) with anthropogenic enrichment of metals, relatively high abundance of quartz and low presence of arsenates, (iv) sites 13–14 (also in central Chile, 35°–37°S) with relatively high values of quartz and low abundance of metals and arsenates, and (v) sites 15–21 (in southern Chile, 37°–41°S) with relatively high presence of arsenates and low abundance of metals and quartz.

Our results have important implications for many Andean communities, especially in central Chile (33°–37°S), where we found significant anthropogenic enrichment. In this region the snowpack is important for urban water supply, power generation, and agriculture. Moreover, our findings are consistent with prior efforts that have shown that presence of higher concentrations of light-absorbing impurities in the Andean snow in the Atacama Desert and at locations nearby Santiago.

Data Availability
The datasets generated and analyzed during the current study are available from the corresponding author on reasonable request. For the trajectory analysis, we used the Global Data Assimilation System (GDAS1) Archive Information provided through the Real-time Environmental Applications and Display System (READY) (http://ready.arl.noaa.gov/gdas1.php).

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Author Contributions
Conceived and designed the experiments: R.R.C., P.M.R., S.N., G.C., J.C., E.L., J.P., A.D. and S.F. Performed the experiments: F.E., J.J., S.M., B.G., J.M.C., J.E., P.L. and E.S. Analyzed the data: J.A.A., R.R.C., P.O., C.M.K. and B.G. Wrote the paper: J.A.A., R.R.C., S.M., A.D., S.N. and S.F.

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