**The 1991 explosive Hudson volcanic eruption as a geochronological marker for the Northern Antarctic Peninsula**

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**Abstract:** It is estimated that the explosive Hudson volcano eruption in Southern Chile injected approximately 2.7 km$^3$ of basalt and trachyandesite tephra into the troposphere between August 8-15, 1991. The Hudson signal has been detected in Antarctica at the eastern sector and in South Pole snow. In this work, we track the Hudson volcanic plume using a dispersion model, remote sensing, and a re-analysis of a high-resolution ice core analysis from the Detroit Plateau in the Antarctic Peninsula and sedimentary records from shallow lakes from King George Island (KGI). The Hudson eruption imprint in these records is confirmed by using a weekly resolved aerosol concentration database from KGI demonstrating that the regional impact of Hudson eruption predominates over the Mount Pinatubo/Philippines volcanic signal, dated from June 1991, in terms of particulate matter depositions. The aerosol elemental composition of Ca, Fe, Ti, Si, Al, Zn, and Pb increases from 2 to 3 orders of magnitude in background level during the days following the eruption of the Hudson volcano.

**Key words:** Aerosols, Antarctic Peninsula, Hudson volcano, Volcanism.

**INTRODUCTION**

The early years of the 1990 decade in Antarctica were characterized by important volcanic ash deposition over the continental ice sheets. Past volcanic activity can be recorded in ice cores as visible volcanic ash (tephra) layers and/or sulfate-rich layers, which have the potential to indicate large-scale atmospheric load (Cole-Dai & Mosley-Thompson 1999). Tephra layer analysis in ice cores can be useful for establishing ice core chronology, reconstructing past atmospheric circulation patterns, and investigating long-term links between volcanic aerosols and climatic change (Zhou et al. 2006).

The second-largest volcanic eruption of the 20th century occurred at Mount Pinatubo in the Philippines on June 12-15, 1991. The first detection of Pinatubo aerosols in Antarctica comes from McMurdo and South Pole Station reports in late 1991 (Cacciani et al. 1993, Deshler et al. 1994). Stratospheric horizontal winds played a significant controlling factor in the long-range aerosol dispersion of Pinatubo eruption products (Cole-Dai et al. 1999). However, no less relevant was the Hudson volcano (southern Chile) contribution during August 8-9, August 12-15, and September 9-15, 1991, due to the explosive character of its eruptions and its closest proximity to Antarctica.
The active stratovolcano Hudson is the southernmost volcano in the Chilean Andes. Contrary to Pinatubo ash dispersion, Hudson volcano injected more pyroclastic material into the troposphere with subsequent and rapid deposition (Inbar et al. 1995, Kilian et al. 1993, Legrand & Wagenbach 1999). It is estimated that during the 1991 explosive eruptions of Hudson volcano, approximately 2.7 km³ of basalt and trachyandesite tephra were ejected to a maximum height of 12 km above sea level (ASL) (Kratzmann et al. 2010). Most of the volcanic material had drifted southwards, reaching the Southern Ocean and the Antarctic continent. In contrast, Pinatubo sulfur-rich clouds reached 25-35 km containing around 1.8 million tons of SO₂ (Krueger et al. 1995), whose impact was spread out all around the globe. In a South Pole snowpit, measurement of sulfate concentration in snow layers dated 1991-1992 distinguished between the Hudson and the Pinatubo deposit (Cole-Dai et al. 1997a, 1999, Palmer et al. 2001). The Hudson peak was observed at that site during December 1991, while Pinatubo peaked in December 1992 and all along 1993.

By the time 1991 Hudson eruptions started, Pinatubo ash had already spread nearly worldwide, with a fine fraction starting to disperse poleward by late August/early September (Trepte et al. 1993). Examination of Pinatubo material dispersion, using the Total Ozone Mapping Spectrometer (TOMS) satellite sensor covering the period between August 13 and September 30, suggests that its impact around the Antarctic Peninsula was not as pronounced as in around the tropics and subtropics (Figure 1). The region encompassing Southern Patagonia, south of latitude 45°S, and

![Figure 1. Image in the upper part was acquired by the Stratospheric Aerosol and Gas Experiment II (SAGE II) flying aboard NASA’s Earth Radiation Budget Satellite (ERBS). It corresponds to the dispersion of Pinatubo (white square) ash around the globe (1020 nm Optical Depth). It shows composite data between August 13 to September 30 of 1991 (https://earthobservatory.nasa.gov/images/1510/global-effects-of-mount-pinatubo). In the window below, the sector where Hudson (green dot) ash had the most significant impact on August 12 of the same year (inferred from the aerosol index product TOMS/NASA sensor).](image-url)
the Southern Ocean – approximately between the longitudes 30°W and 120°W – was also less impacted. Although this period partially covers the Hudson eruption, no impact of the Hudson eruption was observed in the NASA SAGE II measurement of aerosol optical depth since it refers to the stratospheric dispersion of the Pinatubo signal.

A Dispersion model applied to the Hudson eruption using the Lagrangian ash tracking model PUFF proposed by Kratzmann et al. (2010) reproduced the aerial distribution extent and temporal evolution of the 1991 volcanic plume with prevailing ash and aerosol loads within the troposphere. Their model projected the presence of Hudson volcanic materials over the South Atlantic and the Southern Ocean. Another important and more obvious factor is the proximity between the southern Andean region and Antarctica compared to Mount Pinatubo/Philippines and the action of the westerly winds that blow Patagonian mineral particles (dust and volcanic ash) towards the Southern Ocean (Erickson et al. 2003, Gassó et al. 2010, Gassó & Stein 2007).

To better elucidate the impact of the 1991 volcanic events, we reanalyzed a weekly resolved aerosol elemental concentration database from King George Island/Admiralty Bay, obtained in the context of the Brazilian Antarctic Program; an elemental concentration database developed from a high-resolution ice core retrieved in the Northern Antarctic Peninsula/Detroit Plateau, obtained in the context of the US-Brazil-Chile project CASA (Climate of Antarctica and South America); and three shallow sediment cores in King George Island/Admiralty Bay that include the year 1991. We combined these databases with the modeling of volcanic ash dispersion to track the influence of Hudson volcanic eruptions.

**MATERIALS AND METHODS**

**Study site databases**

This work was conducted in two sites on the Northern Antarctica Peninsula: (1) King George Island/Admiralty Bay in the South Shetland Islands, and (2) Detroit Plateau in the Antarctic Peninsula, Figure 2. The Antarctic Peninsula is a relatively narrow mountain ridge that extends over 520,000 km² from the base of West Antarctica (Marie Byrd Land) at latitude 74°S towards its Northern limit in the Drake Passage (Figure 2a). The Bellingshausen Sea bounds the peninsula on the west side while the Weddell Sea lies east. At the tip of the Antarctic Peninsula is located the South Shetland Islands in the northern portion of the Bellingshausen Sea (Figure 2a). The climatic regime of the South Shetland Islands is defined as maritime subpolar (Simonov 1977). It is highly influenced by the migration of subpolar cyclones that bring to this region warm air, high nebulosity, and humidity from the South Pacific, resulting in precipitation as rain and snow (Bintanja 1995). King George Island is located between the Drake Passage and the Bransfield Strait between coordinates 61°50’S – 62°15’S and 57°30’W – 59°00’W. It comprises an area of 1,140 km² with 30 km width, a maximum extension of 79 km at the SW-NE direction, and a maximum altitude around 700 m a.s.l. (Figure 2a). The Antarctic Peninsula is the northernmost part of the Antarctic continent and is 80% ice-covered with an ice sheet of approximately 500 m thick (Bindschadler 2006). It has a sharp elevation gradient, with most glaciers flowing into the Weddell Sea at its Eastern Sector, which has been experiencing significant breakup since 2002 (Turner et al. 2009). These mountains form a significant barrier to the persistent westerly winds that transport moisture and warmer air masses to its location. In view of this physiography, the
regional climatic regime differs significantly at each side of the Antarctic Peninsula. During the austral summer, snow and ice cover more than 90% of its total surface area. Additionally, air mass advections from South America influence the South Shetland Islands since cyclonic migration through this region increases the meridional wind component southwards. This mechanism is responsible for the atmospheric transport of terrigenous particles and biotic material from South America (Hebel et al. 2018, Pereira et al. 2004).

Detroit Plateau (64°10’S; 60°0’W) is a major interior plateau of Graham Land, located in the northern sector of the Antarctic Peninsula, with a mean elevation of ~1900 m a.s.l.. Local mean annual accumulation is on average 2.5 m w.e. for the last 27 years (Potocki et al. 2016), allowing sufficient resolution to study events on a seasonal basis. Air mass advections from South America influence Detroit Plateau to some extent, albeit less so than for the South Shetland Islands.
Satellite products and dispersion model

The dispersion of the volcanic ash plumes was simulated daily by using the Hybrid Single-Particle Lagrangian Integrated Trajectory – HYSPLIT/NOAA (Stein et al. 2015). HYSPLIT is a dispersion model that uses a hybrid Lagrangian model approach and Eulerian methodology. In the particle model case for volcanic ash dispersion, a fixed number of particles are advected about the model domain by the mean wind field and spread using a turbulent component. The model’s default configuration assumes a 3-dimensional particle distribution. The model can be driven by several meteorological databases, and in this case, we used the NCEP/NCAR Reanalysis (Kalnay et al. 1996). HYSPLIT calculates the dispersion and deposition of volcanic material extending from the volcano summit to a user-specified maximum height. The model considers the terrain geomorphology and runs for spherical particles of density $2.5 \times 10^6$ g m$^{-3}$ with diameters ranging from 0.3 to 30 μm. Simulations were carried out for the August and September eruptions, with a maximum height as specified by reports of the height of the ash column. For the first set of simulations, each emission day was followed independently for consecutive 48 hours. A second simulation presents a cumulative view over ten days.

In addition to the dispersion model, we use the Aerosol Index (AI), from the TOMS sensor, onboard the Nimbus-7 (11/1978 to 4/1993) satellite. The AI is calculated based on the difference between the observed and the simulated backscattered ultraviolet (UV) radiation for an atmosphere without aerosols (Herman et al. 1997). The simulation also considers the surface UV reflectivity by using the climatology established over clear atmosphere observations. This method enables the detection of absorbing aerosols (smoke, dust, ash) over water, land, and snow/ice (Hsu et al. 1999a). The index is unitless, and its magnitude depends not only on aerosol optical properties and concentration but also on its altitude and therefore is used mostly in a qualitative way (e.g., Ginoux & Torres 2003, Hsu et al. 1999b), but relative use is also possible when considering short periods or short spatial scales. Here we use the TOMS AI data Version 8. All positive AI values indicate the presence of absorbing aerosols, but smaller values can represent the effects of uncertainty on the assumptions used in the simulation for an aerosol-free atmosphere, and we use only values greater than 0.7. The original data can be found on the Total Ozone Mapping Spectrometer website https://ozoneaq.gsfc.nasa.gov/data/toms/.

Aerosol sampling and elemental composition analysis

Aerosols were collected at the Laboratory of Atmospheric Science located 2 km away from the Brazilian Antarctic Station Comandante Ferraz (62°05'07"S 58°23'29"W). Samples were collected using a double filtration system of Nuclepore polycarbonate filters of 0.4 mm (fine aerosol mode) and 8.0 mm (course aerosol mode) porosity. Data presented here refers to the total elemental composition (fine+course fractions). Air filtration was carried out 24 hours per day and integrated for one week in 1991. A 47 mm-diameter aerosol filter holder was placed in an inlet outside the laboratory at approximately 10 m a.s.l. The pumping system (a Single-head GAST pump system) operated with rates between 10 to 16 L min$^{-1}$. The inlet has a microparticle cutoff size of 10 mm. Filters containing aerosols were analyzed using the PIXE (Particle-Induced X-Ray Emission) technique at the Institute of Physics at the São Paulo University (USP), Brazil. This technique is based on detecting characteristic X-rays emitted by elements in the sample target, as they are bombarded by high energy fluxes
of proton particles of 2.4 MeV generated by a compact isochrone cyclotron (CGR-MeV). The detection limit for PIXE reaches $10^{-7}$ g g$^{-1}$ and the range of element detectability varies from Na to U. Characteristic X-rays are measured by a Kevex Si(Li) detector.

**Ice core sampling and elemental composition analysis**

An ice core was retrieved from Detroit Plateau (64°05’07” S; 059°38’42” W; 1,937 m a.s.l.) in the Antarctic Peninsula in December 2007, within the US-Brazil-Chile joint research program CASA (Climate of the Antarctic and Southern America; https://www.polartropical.org/casa/). The Detroit Plateau ice cap temperature during coring activities was approximate -14°C. The ice core was transported frozen from Antarctica to CCI - Climate Change Institute/Maine University, United States of America.

An electromechanical system provided by the University of Maine drilled to 120 m depth through snow, firn, and ice. The ice core was 9 cm in diameter and was cut in situ into segments varying from 85 to 95 cm. Density values were obtained at the campsite. The upper 98 m of this core was processed using ultra-clean handling procedures. Ice core sampling was conducted inside a cold room (-20°C) at CCI, where the ice core sections were cut into 10 cm long samples, and the surface layer (~2 cm) was removed by an acid-washed Teflon knife. Ice samples for trace-element analysis were collected directly into acid-cleaned (10% trace metal grade HNO$_3$) polypropylene Nalgene jars. Before analysis, each sample was acidified to 1% with Optima double-distilled HNO$_3$ under a class-100 HEPA clean bench and left to digest for 60 days (Potocki et al. 2016). Then, at the Climate Change Institute (University of Maine), the samples were analyzed with an Inductively Coupled Plasma Mass Spectrometer (ICP-MS) for elemental composition. The dating of the ice core was based on H$_2$O$_2$ measurements that yielded a clear interannual sinusoidal variability (Potocki et al. 2016).

**Shallow lake sediment coring, chronology, and elemental analysis**

Periglacial shallow lakes selected for the present work are located at Admiralty Bay/King George Island. The choice of these lakes is related to the observed increase in susceptibility of the region to regional variability associated with dramatic ice mass losses (Sancho et al. 2017, Skvarca et al. 1998). These lakes receive direct atmospheric depositions and water melting from winter/spring fresh snow. Admiralty Bay is characterized by a high density of glacial drainage basins (nearly 32 according to Dani et al. 2012 and Bremer et al. 2004). Two of the three shallow lakes used in this work are located near the bay’s mouth (1. near the terminus of Ecology glacier: coordinate 62°10’39” S, 058°27’61” W with 17 cm long; and 2. at Agatha Point: coord. 62°10’8” S, 058°26’8” W with 16 cm long), the other sampling site is located in the inner part of the Bay at “Ipanema”: coord. 62°05’12” S, 058°25’1” W with 26 cm long). All sediment cores were sampled manually to avoid disturbances. We used acrylic tubes with an inner diameter of 7 cm and a wall thickness of 3 mm. During the field campaign, sediment layers varying from 0.5 to 1 cm were subsampled from the cores for radiometric dating and chemical analysis.

Sediment dating was performed using an extended range CANBERRA gamma spectrometry system with a 25 % relative efficiency co-axial GeHp detector. The gamma detection system is encased in 10 to 15 cm of low background lead shielding and internally coated with 1 mm copper sheets. The instrumentation is currently installed in the Laboratory of Radioecology and Global Changes (LARAMG) of the Rio de Janeiro
State University/Brazil. The gamma system provides a peak resolution of 1.8 keV at 1.33 MeV and 0.850 keV at 122 keV. The calibration standard to construct the detector efficiency curve, as a function of the gamma energy, was a liquid cocktail of radionuclides (\(^{133}\)Ba, \(^{57}\)Co, \(^{139}\)Ce, \(^{85}\)Sr, \(^{137}\)Cs, \(^{54}\)Mn, \(^{88}\)Y, and \(^{65}\)Zn) in 0.5 M of HCl. The standard was manufactured by the AEA Technology with a total activity concentration of 132 kBq and a volume of 100.76 g. Each sediment layer was dried for 24 hours at 50°C. After that, fine sediment samples (around 20 g) were placed in cylindrical plastic pots sealed and submitted to the counting of gamma-ray photons for 86400 seconds each. Radionuclides of dating interest were \(^{210}\)Pb, \(^{226}\)Ra, and \(^{137}\)Cs (Appleby & Oldfield 1992). To interpret \(^{210}\)Pb data in the sediment cores, the mathematical model was used known as Constant Rate of Supply (CRS). It assumes variations in sedimentation rates of a sediment core that cannot be explained by bioturbation phenomena or a physical mixture of sediments (Appleby & Oldfield 1992). This model is based on two premises: a) \(^{210}\)Pb Excess is supplied to the surface of the sediment at a constant rate; b) The radioactive \(^{210}\)Pb decay is constant due to its small post-depositional mobility. The \(^{210}\)Pb present in the rocks comes from \(^{238}\)U radioactive disintegration chain. At the time \(^{226}\)Ra disintegrates and forms \(^{222}\)Rn, a noble gas, part of it emanates from the rocks into the atmosphere, where it undergoes further disintegration up to \(^{210}\)Pb. As this radionuclide is solid, it precipitates in the surface by gravity or "washout" and joins with \(^{210}\)Pb formed in the crystalline structure of the sediment grains of the original rock. This deposited \(^{210}\)Pb is called excess or without support to distinguish it from the \(^{210}\)Pb derivative \textit{in situ} of the \(^{226}\)Ra decay (supported).

For chemical analysis, aliquots from each stratum were taken and dried at 40°C for 24 hours. Metal extraction was made by leaching the samples with 4 mL of HNO₃ at 0.5 M, and 1 mL of HCl at 0.5 M in 3 runs at 150°C. The extraction product was obtained in a sequential step by adding 1 mL of 0.5 M HCl and 25 mL of Milli-Q water to complete a final volume. All reagents used were Suprapur Merk®. Blanks underwent the same procedure and were measured with the samples to assess possible contamination. The quantification of metals occurred employing an Optical Emission Spectrometry with Inductively Coupled Plasma (ICP-OES Ultima II, from Jobin-Yvon). This multi-elementary technique uses an inductive plasma source (from the ionization of argon gas) to atomize and ionize the products of the chemical extraction. The ICP-OES measures the electromagnetic radiation emitted by the atoms or ions excited by the plasma.

In this work, the quantitative method was used. To increase sensitivity, an ultrasonic nebulizer was used to reduce the size of the particles that form the aerosol to be taken to the plasma flame. The applied power was 1000 Watts.

**RESULTS AND DISCUSSIONS**

Volcanic material depositions characterized the first years of the 1990 decade in Antarctica as a result of the Pinatubo and Hudson volcanic eruptions (Cole-Dai et al. 1999, Kilian et al. 1993, Legrand & Wagenbach 1999).

In our simulations of the dispersion of Hudson volcanic material originated at the eruptive episode on August 12-15, 1991, Figure 3, the plumes migrated in a general SE direction, thus directly impacting the South Shetland Islands (especially on August 13-14) and the adjacent Southern Ocean.

The Hudson 1991 eruption reached a Volcanoes Explosivity Index (VEI) of 5 (Global Volcanism Program, Smithsonian Institution,
An injection of a bulk tephra volume of more than 1 km³ into the troposphere and stratosphere. The eruptive event ejected about 2.7 km³ (dense-rock equivalent) of basalt and trachyandesite magma as tephra fell (Scasso & Carey, 2005). Hudson volcanic aerosols were transported poleward, mainly in the troposphere and lower stratosphere beneath the polar vortex directed by the westerlies (Legrand & Wagenbach 1999).

A combined analysis of the Aerosol Index and the transport and dispersion simulation for the volcanic eruption in the subsequent days of August 12-15, 1991 (Figure 4) illustrates the general dynamics of the plume around the Antarctic continent. It can be observed that a high load of volcanic material was deposited closer to the northern tip of the Antarctic Peninsula. In the Weddell Sea sector, an existing cyclonic system sped up the surface winds in clockwise movement impacting the Antarctic Peninsula from August 19 to August 23, 1991. In approximately one month, the plume from Hudson circled Antarctica, and due to prevailing winds (westerlies), most of the ash was deposited in the Southern Ocean. Similar effects of dilution and deposition hampering remote sensing observations from source to sampling site were described for dust emissions from Patagonia (Gassó et al. 2010). Chemical analysis of the Hudson volcano eruptive products revealed a composition of 58-63% of silica (SiO₂), medium-to high-K₂O content, relative enrichment of TiO₂, Na₂O, and enrichment in LIL (Large Ion Lithophile; i.e., K, Rb, Cs, Sr, Ba, and Pb), HFSE (High Field Strength Elements; i.e., Ti, Zr, Hf, Nb, and Ta),
and REE (Rare Earth Elements) (Inbar et al. 1995, Scasso & Carey 2005). Chemical analysis of the aerosol samples collected at the King George Island (Figure 5a), weekly resolution, reveals high enrichment in Ca, Fe, Ti, Si, and Al during the volcanic activity of Hudson, more notably after the 12-15 August event. Although with lower resolution (seasonal), concentrations of Fe, Ti, Si, and Ba are found to be highly elevated in the August 1991 snow layer in the Detroit Plateau ice core (Figure 5b). Concentrations of Pb and Zn in the sediment for this period are much higher than the baseline value (Figure 6).

The increase from 2 to 3 orders of magnitude over background for the days following the eruption in Southern Chile is noted in both aerosols and the ice core from the Antarctic Peninsula, suggesting a major contribution of the Hudson eruption. The combination of continuous aerosol monitoring before, during, and after the volcanic episode together with the dispersion model and Aerosol Index data constitute a definitive demonstration that the anomalous increases observed in the atmosphere, ice core, and in sediments of the pro-glacial environment of the Northern Antarctic Peninsula are related.
to the 1991 Hudson eruptive episode. Due to the recognized global-scale impact of Mount Pinatubo volcanism on the climate system, many researchers have attributed the changes in this part of the Southern Ocean, especially at the Northern Antarctic Peninsula, in 1991 exclusively to the Pinatubo eruption. Verona et al. (2019) have suggested that Pinatubo induced cooling in the Southern Ocean and warming up to 0.8°C off the Antarctic Peninsula. This happens for two reasons: (a) models in use do not assimilate most of the moderate-intensity volcanisms (Gao et al. 2008); (b) the real impact extension of Hudson volcanic event is not fully described for higher latitudes. Our results may shed some light on the significant impact of Hudson particulate matter precipitation at the Northern Antarctic Peninsula, despite our work not wanting to establish any direct relationship with the climate issue.

Another interesting aspect observed in this work is the interpretation of the abrupt increases in metal composition, like Al, Fe, Ti, and Ca, in the ice core. Such events are usually attributed to dust storm advection, but our observations show that explosive eruptions can substantially elevate concentrations of the elements.

Volcanic signal detection is observed in other locations around Antarctica, suggesting a widespread impact on the continent. Ice cores retrieved close to the Amundsen-Scott South Pole Station have significant deposition of $\text{SO}_4^{2-}$ related to the Hudson eruption event in September 1991 (Cole-Dai et al. 1997b). Similarly, the DT263 ice core, retrieved from an East Antarctic location, exhibited $\text{SO}_4^{2-}$ concentrations dated to 1991 and was identified as a Hudson
volcano contribution (Zhou et al. 2006). Although the Hudson volcanic eruption produced less SO$_2$ when compared to Pinatubo’s production (~8%), the volcanic aerosol mass from the former extended southward immediately and reached the Antarctica continent more effectively (Zhou et al. 2006). Also, the Hudson plume was lower in altitude than Pinatubo’s; this resulted in the deposition of Hudson ash almost immediately following the eruption, while the Pinatubo materials reached the atmosphere over Antarctica later in the year and deposited following that of the Hudson materials. The stratospheric SO$_2$ clouds dispersion resulting from the Hudson eruption was tracked by TOMS for many days and identified in East Antarctica by 20$^{th}$ August 1991 (Carn et al. 2003, Constantine et al. 2000).

Figure 7 summarizes the identified Hudson eruptions impacts around the whole Antarctic continent, depicting the widespread distribution of the volcanic aerosols due to the westerlies and a comparable result concerning the Aerosol Index data and transport/dispersion of volcanic material simulations presented here (Figure 4).

**CONCLUSIONS**

We tracked the Hudson volcanic plume using the HYSPLIT particle mode and the Aerosol Index data for the eruption episode between August and mid-September 1991. Remote sensing data indicates a rapid dispersion of the eruption plume from the Hudson volcano to the Antarctic Peninsula, with a more expressive impact from the second event, between 12-15 August 1991. Our simultaneous weekly resolved aerosol sampling from King George Island captured that event based on an anomalous increase in elemental Ca, Fe, Ti, Si, and Al, relative to the background in the days following the eruption, and compatible with the Hudson tephra chemical composition. The impact was also recorded in a simultaneous increase in Fe, Ti, Si, and Ba concentrations in an ice core from Detroit Plateau on the Antarctic Peninsula and in high Pb and Zn concentrations in shallow sediment cores retrieved on King George Island at the same time. Although our work has surveyed only the elemental compositions (not volcanic glass shards or cryptotephra), the above pieces
of evidence strongly suggest an association with the volcanic event.

Although the 1991 Pinatubo eruption had a greater global impact, its most distant location concerning Antarctica and the characteristic of its explosion led to the muted deposition of the eruption materials in the Antarctic ice sheet. Therefore, the 1991 Hudson eruptive event is of greater significance as a geochronological signal for the Northern Antarctic Peninsula. This work highlights Southern Andean volcanism as a key player in the West Antarctic and the Antarctic Peninsula climate, with immediate and direct impacts over the continent. Abrupt increases in metal composition in the ice cores, usually attributed to dust storm advection, can be observed following important eruptive events.

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