Abstract
Understanding the mechanisms that govern ash aggregation is of critical importance in volcanology. Aggregation reduces the residence time of ash ($\leq$2 mm) in the atmosphere strongly, by enhancing the sedimentation of finer-grained material generated during explosive eruptions. To date, experimental studies have focused on the expectation that water provides the strongest bonds between particles to form spherical to oblate aggregates (typically less than or equal to a few mm, occasionally up to several cm) preserved in pyroclastic deposits. Under water-rich conditions, individual accreted particles rarely exceed 1 mm. In pyroclastic density current deposits produced during the 1982 eruption of El Chichón, Mexico (which emitted 7.5 Tg of SO$_2$ in the atmosphere), aggregate particles one to several millimeters, strongly cemented by a S-rich film, are common. They exhibit similarities with aggregates found in sulfur cones at Poás volcano, Costa Rica. We propose that sulfur is the binder between the silicate grains. Such a binding capacity implies a relatively fluid behavior of sulfur such as might be expected in the low viscosity temperature regime just above its melting point. If so, then the explosive ejection of sulfur during eruptions, combined with its ability to act as a cement for particles $>2$ mm, would imply that size fractions of lapilli can be efficiently removed from eruptive clouds a few kilometers from the vent. Such an aggregation mechanism would have important implications for pyroclast dispersal models and hazard assessment.

Introduction
In explosive eruptions, ash ($\leq$2 mm) typically represents a large fraction of the total erupted mass. It is dispersed in the atmosphere by vertical plumes and/or pyroclastic density currents (PDCs). Airborne measurements within relatively dilute outflow regions of eruptive clouds, where large particles have settled out, indicate that fine ash (1 $\mu$m to 1 mm) could yield number concentrations of up to hundreds per cubic centimeter (Hobbs et al. 1982, 1991).

Aggregation reduces the transportability of ash and thereby the distances at which a given concentration may be expected in volcanic clouds. This aggregation-induced reduction in atmospheric residence time results from the enhanced settling velocity of aggregates compared with single particles (Sparks et al. 1997).

For this reason, research has been undertaken: to understand the causes responsible for the formation of different types of aggregates, to reproduce experimentally their internal structure, and to model their transport and depositional behavior (see below).

Among the aggregates described in pyroclastic deposits worldwide, two particular occurrences, i.e., those described in PDC deposits emplaced during the 1982 eruption of El Chichón (Mexico), and the 2005 eruption of Iłamatepec (El Salvador), despite their otherwise similar character (in dimensions, morphology, and internal structure) to aggregates described elsewhere, differ from the others in exhibiting an orange to red, micrometer-thick film, cementing individual particles (Scolamacchia et al. 2005, 2010). Semiquantitative EDS analyses, as transects across the film, and compositional mapping of the aggregates indicate that the binding film consists of an amorphous phase rich in S and Fe and minor P, with variable amounts of K, Ca, Na, and Mg, depending on the particle to which they adhere. This particular compositional range has been attributed to the precipitation of an
amorphous phase from acid solutions within the volcanic clouds derived from the contemporaneous ejection of magmatic and hydrothermal fluids and groundwater (Scolamacchia et al. 2005).

Here, we propose an alternative explanation for the formation of these aggregates and for the origin of the binding substance between individual grains, based on textural characteristics and further analyses.

Background

The presence of aggregates has, in the past, been inferred from the observation of a polymodal particle size distribution of fallout deposits and is the most viable mechanism to explain the occurrence of secondary thickness maxima in such deposits (i.e., Sarna-Wojcicki et al. 1981; Carey and Sigurdsson 1982; Brazier et al. 1982, 1983).

In this respect, Durant et al. (2009) have proposed that the secondary thickening in fallout deposits of the 1980 eruption of Mount St. Helens, observed at 325 km ENE from the vent (Ritzville maxima of Sarna-Wojcicki et al. 1981), was produced primarily by the aggregation of particles with modal diameters of 19±3 μm, transported into the upper troposphere, where they initiated ice nucleation. The increase in mass loading of the eruptive cloud, due to ice formation, would have caused its subsidence until reaching an altitude of 6.9 km, where ice melting would have caused the fallout of the fine ash fraction within a few hours.

Spherical to oblate pellets (100 μm to several tens of mm in maximum diameter and, in some cases, few cm, Fig. 1), with or without an internal structure and/or a solid core, are commonly visible in fallout, and PDC deposits produced by a variety of eruption types (e.g. Reimer 1983; Sheridan and Wohletz 1983; Wilson 2001; Ritchie et al. 2002).

Observations during explosive eruptions have reported the fallout of 1 to 10-mm diameter aggregates, from plumes that are a few to tens of kilometer high, indicating that fine-ash particles frequently fall as larger aggregates (e.g., Brazier et al. 1982; Tomita et al. 1985; Gilbert and Lane 1994; Bonadonna et al. 2002).

Analysis of high-speed video recordings of ash falling from volcanic clouds during the 2010 eruption of Eyjafjallajökull volcano (Iceland) has yielded the interpretation that aggregation caused a tenfold increase in mass sedimentation at distances of 7 km from the vent (Taddeucci et al. 2011). Aggregates dominated over single particles in size fractions between 500 μm and 2 mm, and <250 μm, implying that without aggregation, only individual particles between 1.5 mm and 355 μm in size would have been able to settle from the cloud.

Several studies have attempted to determine the optimal conditions necessary for the scavenging of particles into aggregates, by experiments (Gilbert and Lane 1994; Schumacher and Schmincke 1995; James et al. 2002; Telling and Dufek 2012; Van Eaton et al. 2012a; Telling et al. 2013) and by numerical models (Veitch and Woods 2001; Textor et al. 2006a, b; Durant et al. 2009; Costa et al. 2010; Folch et al. 2010; Van Eaton et al. 2012b) in order to infer the areas in both volcanic plumes and pyroclastic density currents (PDCs), where aggregation could be favored.

Binding forces

Electrostatic forces between charged, recently fragmented, or colliding particles and capillary forces due to the action of liquid water have both been proposed to play a role in aggregation processes.

Lane et al. (1993) and Gilbert and Lane (1994) suggested that the growth of aggregates is controlled by collisions among liquid-coated particles, due to the ability of liquid bridges to dissipate the kinetic energy of particle collisions, promoting particle capture. Aggregation efficiency should thereby increase as particle diameter decreases, varying from nearly 100 to 5 % for particles 1 to 100 μm in size, respectively (Gilbert and Lane 1994). Liquid water could therefore explain the formation of sub-millimeter- to centimeter-sized aggregates found in the stratigraphic records (e.g., Schumacher and Schmincke 1991, 1995; Bonadonna et al. 2002; Brown et al. 2010, 2012), by providing stronger bonds between particles with respect to short-term, but long-range, electrostatic forces between charged, recently fragmented, ash, which by contrast would only form fragile, ephemeral, aggregates <1 mm in maximum size (James et al. 2002).

More recently, experiments indicated that a critical residence time (>50 min) in an atmosphere at relative humidity >71 % would allow the formation of a film of water around...
particles 90 to 150 μm in size, determining an increase in the aggregation efficiency by more than 60 % (Telling et al. 2013).

Various experiments agree that an optimum amount of liquid water, around 15 wt% (30 vol%), is required to form well-developed spherical aggregates from a polymodal ash population (Schumacher and Schmincke 1995; Van Eaton et al. 2012a). At such contents, aggregates would already be strong enough to survive impacts with the ground falling 1 m (Van Eaton et al. 2012a), without the presence of secondary minerals (e.g., gypsum and sodium chloride) precipitated from acid solutions (Gilbert and Lane 1994). At ~15 wt% water, the particles scavenged into aggregates are mostly <250 μm in size. Even for liquid contents >15 wt%, particles >200 μm are rarely incorporated into aggregates, and below 20–25 wt%, all particles >125–250 μm are also preferentially excluded (Van Eaton et al. 2012a). Aggregation would be even more strongly size-selective for water contents <5 wt%, at which only particles <63 μm are preferentially scavenged from the parent mixture (Van Eaton et al. 2012a). Such aggregates are fragile, and grain-dispersive forces can easily overcome the binding forces due to liquid water (Schumacher and Schmincke 1995).

Experimental results also indicate that particles ≥1 mm are preferentially excluded from aggregation, except for liquid water contents >20–25 wt% (45–50 vol%). At such contents, particles up to 2 mm are scavenged by pockets of oversaturated ash (Schumacher and Schmincke 1995; Van Eaton et al. 2012a, b), which nevertheless rapidly lose their strength, merging into slurries (liquid pellets, or AP3, of Gilbert and Lane 1994). At ~15 wt% water, the particles scavenged into aggregates are mostly <250 μm in size. Even for liquid contents >15 wt%, particles >200 μm are rarely incorporated into aggregates, and below 20–25 wt%, all particles >25 wt%, are also preferentially excluded (Van Eaton et al. 2012a). Aggregation would be even more strongly size-selective for water contents <5 wt%, at which only particles <63 μm are preferentially scavenged from the parent mixture (Van Eaton et al. 2012a). Such aggregates are fragile, and grain-dispersive forces can easily overcome the binding forces due to liquid water (Schumacher and Schmincke 1995).

When eruptive plumes rise in the atmosphere, water can undergo phase transformations. The formation of ice inside plumes can occur within the first 2–3 km of ascent, depending on the season and latitude (Sparks et al. 1997). Higher up, the formation of ice can be significant and dominate in comparison to liquid water, triggering aggregation processes (Textor et al. 2006b).

Heterogeneous ice nucleation triggered by volcanic ash occurs at lower supersaturation and higher temperatures, with respect to homogeneous ice nucleation in the presence of pure water (−35 and −40 °C). According to Durant et al. (2008), particles 1 μm to 1 mm in size are able to nucleate ice at temperatures between −10 and −20 °C; differences in the bulk silica content (i.e. basaltic vs. rhyolitic compositions) have only a weak effect on the onset of freezing.

Nevertheless, experiments using hailstones (Van Eaton et al. 2012a) indicate that solid ice preferentially scavenges particles <63 μm (mostly <34 μm), in the absence of a liquid phase. Particles are initially accreted by electrostatic forces generated by particle collisions; more permanent bonds develop only tens of seconds or minutes afterwards, due to ice sintering, at subfreezing temperatures. The timescale of ice sintering nevertheless limits the ice-binding ability because particle collisions, lasting only milliseconds in the absence of electrostatic or capillary forces, are insufficient to keep ice-coated particles in contact long enough for sintering (Van Eaton et al. 2012a).

Aggregate formation should therefore depend on the fraction of water or ice available at the particle surfaces. Hygroscopic compounds could accelerate sintering processes, promoting the development of thin liquid layers on ice surface, even at subfreezing temperatures (Van Eaton et al. 2012a). In fact, experiments by Textor et al. (2006a) indicated that acid solutions (i.e., mixtures of water and soluble gases like SO2, H2S, HCl, and HBr), or soluble salts, coating particles, are able to lower the freezing point of ice, increasing the mass of liquid water, accelerating aggregation processes, due to an increase in liquid water availability (Textor et al. 2006b). Nevertheless, according to Latham et al. (2011), the uptake of liquid water on particles <125 μm due to hygroscopic compounds would be unlikely below 100 % relative humidity (RH) and would not increase the aggregation efficiency even above 80 % RH.

Numerical models and field observations

The entrainment of ambient air, rich in water vapor, either at the fringes of eruptive plumes or during the horizontal movement of pyroclastic density currents (PDCs), can provide the amount of water necessary for aggregation, considering that the amount of water as exsolved volatiles during explosive volcanic eruptions is negligible in this process (Sparks et al. 1997).

Numerical models (Veitch and Woods 2001; Textor et al. 2006a, b; Durant et al. 2008; Folch et al. 2010; Costa et al. 2010) focused on moderate to large intensity explosive eruptions of (≤107 and 108 kg/s M.E.R.), which generate tens of kilometer-high eruptive columns that reach the stratosphere. Aggregation would be related to the concentration of particles within eruptive clouds, the fraction of liquid water or ice inside them, and the intensity of eruptions, which determines the height at which the process can occur (Veitch and Woods 2001; Textor et al. 2006b; Costa et al. 2010).

Even considering the accretion capacity of ice inside eruptive clouds (e.g., Textor et al. 2006a, b; Durant et al. 2008; 2009; Costa et al. 2010), all numerical models agree that liquid water provides the stronger binding forces between particles since they are based on experimental results, which use water exclusively as a binding agent. Therefore, they concentrate on size fractions ≤250 μm due to the higher probability of these particles’ sizes to be incorporated as stable aggregates (e.g., Veitch and Woods 2001; Textor et al. 2006a, b; Durant et al. 2008; Costa et al. 2010; Folch et al. 2010).
Based on the distribution of different types of aggregates within different layers of ignimbrite couplets in Tenerife, Brown et al. (2010) proposed that the internal structure of aggregates reflects the amount of condensing water with height inside thermally stratified PDCs. Structureless pellets (AP1 of Brown et al. 2012) made of particles with median grain sizes <100 μm, abundant in capping, laterally continuous, and elasto-supported fallout layers, form within the co-ignimbrite ash plumes. As they settle in the lower, more turbulent, and progressively hotter portions of the density-stratified current, they accrete layers of finer-grained particles (10 to 15 μm), forming pellets with a concentric structure (accretionary lapilli, Brown et al. 2010; or AP2 of Brown et al. 2012). These concentric pellets are preserved in the lower, matrix-supported, ignimbrite layers, together with fragments of them, indicating that fragmentation occurred via collisions during horizontal transport (Fig. 1). The accretion of finer-grained rims, requiring only a minimum of condensation, would therefore attest the higher temperatures and lower water contents existing in the lower portions of the currents. Similar vertical variations in the distribution of aggregates have been described by Schumacher and Schmincke (1991), in pyroclastic flow deposits at Laacher See (Germany), by Branney and Brown (2011) in meteorite impact ejecta deposits (Stac Fada Member) from an unknown source in Scotland and by Van Eaton and Wilson (2013) in the eruptive units of the 25.4 ka Oruanui eruption of Taupo (New Zealand). Based on the distribution of different types of aggregates, it would be possible to discriminate deposits emplaced by different portions of a density current (pyroclastic or impactoclastic) (i.e., upper, current-derived buoyant plume or unsteady lower body of the current). Nevertheless, according to Schumacher and Schmincke (1991), such a distinction is difficult to make for PDC deposits produced by hydromagmatic activity (i.e., base surges), because “mixed populations” of aggregates occur in the same depositional layer. Beyond supporting the model proposed by Brown et al. (2010), Van Eaton and Wilson (2013) point out that although linked to the passage of PDCs, the growth of extremely fine-grained (<10 μm) outer rims does not necessarily form within the laterally moving currents, being more likely dependent on the supply of fine particles rather than heat or water contents (Van Eaton et al. 2012a).

The size distribution of individual particles inside aggregates mimics to some degree the size distribution of the parent deposit and the sorting related to the specific emplacement mechanism (i.e., fallout or PDC). Analyzing the distribution of aggregates within different deposit types at Laacher See, Schumacher and Schmincke (1995) observed that aggregates within co-ignimbrite fall deposits showed a unimodal distribution of particles with maximum grain sizes between 250 and 350 μm, while aggregates within PDC deposits from hydromagmatic activity (i.e., pyroclastic surge) had coarser modes at >500 μm, in rare cases up to 2 mm. The occurrence of aggregates with a nucleus >500 μm found at distal sites from the vent was attributed to the increase in water vapor condensation during the horizontal spreading of PDCs, causing the greater amounts of water available to accrete coarser particles (Schumacher and Schmincke 1995).

**Aggregates at El Chichón (Mexico) and Ilamatepec (El Salvador) volcanoes**

Several types of aggregates (irregular clusters, spherical or elongated pellets, with or without a concentric laminar or a solid nucleus, following Brown et al. 2012), consisting of crystals and pumice clasts and cemented by a film rich in S and Fe, were found in deposits emplaced by PDCs of the 1982 eruption of El Chichón (Mexico) (Scolamacchia et al. 2005) (Fig. 2a, b) and the 2005 eruption of Ilamatepec (El Salvador) (Scolamacchia et al. 2010).

Such aggregates are only sometimes visible at the outcrop scale at both volcanoes, due to their covering of fine ash. Nevertheless, after being cleaned with an ultrasonic device (see Scolamacchia et al. 2005 for analytical methods used), such aggregates represent a significant fraction (30 up to 90 %) of the components of deposits produced during several eruptive phases by hydromagmatic activity of El Chichón. Individual particles in all aggregates have median grain sizes between 250 and 63 μm, reflecting the median granulometric distribution of the host deposits, but 2–3 wt% of particles of 2–6 mm in size not only functions as nuclei for the accretion of other particles, but also occurs in rims of pellets with a concentric structure (e.g., Figs. 8A, B of Scolamacchia et al. 2005). These finer-grained rims are composed of particles <63 μm (88 to 95 wt%), most of which (74–78 wt%) are finer than 16 μm (as observed in other aggregates worldwide). They also contain abundant particles with sizes of 125 μm (14 to 21 wt%), 250 μm (9–11 wt%), and 500 μm (4–6 wt%). In irregular clusters, particles between 500 μm and 3 mm represent between 40 and 60 wt% of the individual components. All particles are strongly bound to each other with a reddish-orange film (Fig. 2a, b), conferring an overall poor sorting of the aggregate (Fig. 3a, b). Strongly cemented irregular clusters, able to survive vigorous sieving by hand, represent up to 90 wt% of fractions >4 mm in some deposits. The distribution of all types of cemented aggregates shows greater abundance in proximal-medial areas (4–6 km from the crater) even if they are still present in smaller amounts (≤10 wt%) up to 9 km (Scolamacchia et al. 2005). Beyond these distances, aggregates consist only of structureless pellets (AP1 of Brown et al. 2012), which are free of the S-rich film, and consist mostly of particles finer than 16 μm (50 to 85 wt%) with maximum sizes of individual components of 100–300 μm (similar to...
aggregates described in other fallout and co-PDC deposits worldwide, e.g., Schumacher and Schmincke 1991; Bonadonna et al. 2002; Brown et al. 2010; Van Eaton and Wilson 2013). These pellets, which are also found associated with different types of cemented aggregates at lesser distances from the vent (i.e., Table 1 of Scolamacchia et al. 2005), have been inferred to have settled from phoenix clouds associated with PDC movement up to distances of 10.5 km (the maximum extension of diluted PDC deposits at El Chichón).

Cemented aggregates have also been found in PDC deposits of the 2005 eruption of Ilamatepec (mostly coated particles or pellets with a concentric structure, PC2 and AP2 of Brown et al. 2012), in which case, they represent more than 50 wt%, sometimes up to 90 wt%, of size fractions between 8 and 2 mm, with a distribution limited to distances up to 5 km from the eruptive vent (Scolamacchia et al. 2010). Scolamacchia et al. (2005) interpreted the nature of the S-Fe-rich film binding the particles to the precipitation of an amorphous phase from either S-rich solutions or condensing hygroscopic fluids, derived from the contemporaneous ejection of magmatic, hydrothermal fluids and phreatic water during several phases of the eruption.

Microanalytical investigation

An increasing number of field and experimental studies have demonstrated that the maximum sizes of individual particles accreted in aggregates are much smaller than those found in aggregates found at El Chichón and Ilamatepec (see Background section). Moreover, the amount of water uptake on ash surfaces due to hygroscopic soluble salts or aerosols does not increase except for the case of high RH conditions, which has been tested only on fine ash (<125 μm) (Latham et al. 2011). Given the above gaps in our understanding of these processes, here we reinvestigate the process of accretion by analyzing

Fig. 2 a View of an irregular cluster found in the 1982 PDC deposits at El Chichón volcano. Note the red cementing film among particles, consisting of crystals of different types and rounded pumice. Scale at the bottom in millimeter. b Detail of the film cementing particles in irregular clusters at El Chichón. Note the thickness of the film and the maximum sizes of particles, which commonly exceed 1 mm. Scale to the left in millimeter

Fig. 3 a Backscattered SEM image showing the contact between single particles in ash aggregates found at El Chichón. Note the poor sorting, coarser fragments, and abundant void spaces among particles, constituting the aggregate. A micrometric-thick film (white), exhibiting bubble-like shapes, holds the particles together. Compare with Fig. 4. b Backscattered SEM image showing a detail of the cementing film between different particles, mostly crystals of plagioclase, in ash aggregates found at El Chichón. The high contrast of the film, few microns thick, is due to its high Fe content, as well as S
transversal cross sections of aggregates. We used a CAMECA SX100 electron microprobe (LMU Munich) with an accelerating potential of 15 kV and a beam current of 5 nA.

In spherical and elongated pellets with a concentric structure, the cementing film was less than 1 μm thick and distributed within the fine-grained material (see Fig. 9E, F of Scolamacchia et al. 2005), preventing quantitative measurement by WDS.

In irregular clusters and in spherical pellets with a solid core, the film thickness was up to 5–6 μm. In this case, even when using a focused beam (100–200 nm), the beam thickness was 6–8 μm; therefore, a contamination from neighboring particles is expected. The contact between the film and the accreted particles (Fig. 4a, b) indicates that some of them (i.e., pumice) were still molten and mingled with the cementing material (Fig. 4b). Reaction rims/fronts occur between the film and the crystals (Fig. 4b), and an incipient vesiculation of the film between accreted glass and crystals is visible (Fig. 4a, b). Therefore, it is likely that the composition of the film is not the original one. Chips of crystals resulting from fracturing along preexisting inclusions (see Scolamacchia et al. 2005) are preserved by the film close to the locus of rupture (Fig. 4b). These textural features indicate that the accretion occurred during the early stages of particle ejection, before they were cooled in the atmosphere.

**Evidence of other aggregation regimes: sulfur as a binder**

The common occurrence of individual components larger than 1 mm in aggregates of El Chichón and Imlatepec volcanoes contrasts with the size of individual particles inside aggregates found in different types of pyroclastic deposits worldwide, where clasts >500 μm are only found as nuclei for the accretion of smaller ones (Schumacher and Schoenke 1995).

Nevertheless, individual components exceeding 850 μm, in many cases 1.4 mm, have been reported in aggregates of different shapes (i.e., irregular, elongated, or spherical) found on sulfur cones formed inside the crater of Poás volcano in 1989 (Oppenheimer 1992). Such aggregates had an overall poor internal sorting. Spherical aggregates up to 5 cm displayed, in some cases, a weak internal layering. Such aggregates could have formed within the conduits of sulfur cones, during eruptions consisting of “geyser-like” bursts of mud and liquid sulfur up to heights of 5 m, at rates 1/s, attributed to the venting of fumarolic gases toward the surface through shallow layers of molten sulfur, whereby the individual components were cemented by liquid sulfur derived from the remelting of S-grains (Oppenheimer 1992). Oppenheimer and Stevenson (1989) attributed the accumulation of layers of elemental sulfur (Sₐ) to the reactions occurred between ascending SO₂ and H₂S (either SO₂ disproportionation or, oxidation of H₂S by SO₂, O₂, H₂O, H₂SO₄, or Fe³⁺ contained in the lake waters) from a degassing andesitic magma at shallow depths and the lake sediments. Sulfur melting would have occurred after that the lake waters, which were acting as a temperature buffer, disappeared, allowing the sediment interface temperature to rise above the sulfur liquidus. The low viscosity of S at temperatures near its melting temperature would have facilitated its migration and coalescence with sediments (Oppenheimer 1992).
The similarities (i.e., internal grain-size distribution, sorting, morphology, and composition of the cementing film) between the sulfur aggregates described at Poás (Oppenheimer 1992) and those reported at El Chichón and Ilimatepec indicate that, also in these cases, sulfur acted as a binder among particles. Nevertheless, in contrast to Poás, the pristine composition of the binding substance of aggregates described here cannot be determined and reflects the sum of different contamination processes, which are difficult to separate due to the peculiar behavior of sulfur (see Discussion section).

The abundance of such aggregates in deposits emplaced during different phases of the 1982 eruption of El Chichón (right up to the final ones) indicates that liquid sulfur was explosively ejected and that its supply was sustained. Moreover, the poor internal sorting (Figs. 2a and 3a), and their textural features (Fig. 4a, b), suggests a fast process of accretion, completed before pumice cooling and before the fragments from fractured crystals were separated, in contrast with respect to those described elsewhere (Fig. 5).

Evidence of S-layers

The explosive ejection of liquid sulfur is not unusual and has been reported at other andesitic volcanoes hosting hydrothermal systems, beside Poás. At Kusatsu-Shirane (Japan), the ejection of liquid sulfur occurred during several phreatic eruptions, through the acidic lake inside the crater (Takano et al. 1994). At Ruapehu, the most active volcano of New Zealand, the accumulation of a pool of elemental sulfur (>50 m in diameter and 6 m deep), with temperatures between 173 and 177 °C, was discovered in 1991 at depths of 134 m beneath a hot, acidic lake hosting sulfate-chloride waters (Crater Lake) inside the crater (Christenson and Wood 1993). Historical activity consisted mostly of phreatic and phreatomagmatic eruptions, with the occurrence of few Plinian eruptions, and the effusive emplacement of viscous andesitic lava domes (Johnston et al. 2000). According to the model of hydrothermal system envisaged at Ruapehu (Hurst et al. 1991; Christenson and Wood 1993; Christenson 2000), the condensation of more soluble gases (SO2 and H2S) among those exsolved from an andesitic magma reacts with water-saturated sediments, within a convective hydrothermal system, precipitating native sulfur (S0) through the reaction 2H2S + SO2 = 3S0 + H2O (Christenson 2000). Such process would have caused the deposition of shrouds of elemental S around the conduits and an S-pool at the lake bottom, in a time lapse between 20 and 50 years following major magmatic eruptions, in 1945 and 1995–1996, despite the occurrence of phreatic and phreatomagmatic eruptions (Christenson 2000; Christenson et al. 2010). Modeling by Christenson and Woods (1993) of hydrothermal circulation before the 1995–1996 eruption indicated that the condensation of 1 kg of magmatic gases at 250 °C and 0.85 MPa would have determined the precipitation of 10 g of elemental S followed by the precipitation of anhydrite and alunite. Christenson (2000) estimated that 0.75 Tg of elemental S was introduced in the hydrothermal system between 1967 and 1995; assuming an average input of 30 Gg per year after 1945, a total amount of 1.5 Tg was stored in the hydrothermal system in 1995. Apparently, this amount was expelled during this eruption.

Similarly, Rowe et al. (1992) estimated deposition of 1.15×107 kg S/year in the hydrothermal system of Poás volcano.

Even if the storage of layers of elemental sulfur was not reported at El Chichón volcano, several lines of evidence support this hypothesis.

The 1982 eruption breached through a sealed hydrothermal system (Taran et al. 2008), uncovering its upper 500 m, forming a 1-km wide and 300-m deep crater (Taran et al. 1998), and releasing 5–9 Tg of SO2 in the stratosphere (Krueger et al. 1995), 7.5 Tg of SO2 according to more recent estimates (Krueger et al. 2008).

Sulfur content in fresh pumices was estimated in ≈1.25 wt% SO3, almost entirely in the form of ≈1 vol% anhydrite crystals precipitated directly from a trachyandesitic magma at temperatures of 810±40 °C, as attested by both 34Sδ34S and δ18O isotopic data (Rye et al. 1984) and textural evidences (Luhr et al. 1984; Luhr and Logan 2002). The total content of sulfur in magma, estimated in 2.6 wt% SO3, was nearly an order of magnitude greater than contents typical for subduction-related magmas (Luhr et al. 1984; Varekamp et al. 1984).

The 1982 pumice consisted of a stable equilibrium assemblage of hornblende, augite, pyrrhotite, anhydrite, and
titanomagnetite (Luhr et al. 1984). Anhydrite microcrystals account for 1.3 wt% of the total S emitted. Nevertheless, the low S content (<0.1 wt%) in the matrix glass of the 1982 pumices and evidence from trapped glass inclusions both suggest that the remaining 1.2 wt% was present in a separate gas phase, in which the fugacity of H₂S was greater than SO₂ (Luhr et al. 1984). This hypothesis is consistent with the high δ³⁴S values (5.8‰) estimated for the bulk magma (i.e., melt, crystals, and gas), suggesting a loss of H₂S depleted in δ³⁴S, into a liquid or a gas phase at depth (Rye et al. 1984). Beside primary anhydrite in juvenile material, lithic clasts ejected during the eruption showed intense sulfate mineralization and anhydrite veins, with isotopic values of δ³⁴S=13.3‰. These isotopic values were explained with the presence of heavy hydrothermal components beneath El Chichón (Rye et al. 1984).

Varekamp et al. (1984) and Luhr et al. (1984) have suggested that almost all sulfur not present as microphenocrysts of anhydrite was released as a H₂S gas inside the eruptive clouds and then either absorbed onto ash after its oxidation to sulfate or alternatively condensed as sulfuric acid droplets, leading to a clustering of fines (<63 μm) and producing a polymodal distribution (with modes at 4 and 1 mm) of fallout at distances of 17, 22, and 110 km NE from the vent. Both δ¹⁸O and δ³⁴S isotopic data indicate that the rapid oxidation of H₂S into SO₄²⁻ into eruptive clouds was kinetically prohibited and that a more likely source of sulfate adsorbed on ash consisted of primary oxidized sulfur (Rye et al. 1984). The lower isotopic values of oxygen and sulfur of the bulk ash leachates (δ³⁴S=+7.8 to 8.4‰), compared with those of anhydrites crystals in pumices, suggested that the excess S did not derive from the dissolution of primary anhydrite (Rye et al. 1984). Varekamp et al. (1984) indicated that using only the main phases occurring in pumices to obtain the modal composition of ashes always yielded negative quantities of augite by including excess Na and S as plagioclase and anhydrite. Nevertheless, the fits improved when additional Na-K-rich phases (accounting for vapor phase condensates and altered lithic fragments rich in alkalis), and elemental S (accounting for sulfur not present as anhydrite), were included. Pyrite and anhydrite, found in accidental blocks ejected during the 1982 eruption, were also common minerals precipitated from hydrothermal solutions at Ruapehu.

Therefore, it is likely that elemental sulfur was slowly accumulating also in the hydrothermal system at El Chichón, by precipitation from condensing SO₂ and H₂S, emitted from the degassing trachyandesitic magma, similar to what reported at other andesitic volcanoes hosting active hydrothermal systems (e.g., Oppenheimer and Stevenson 1989; Christenson and Wood 1993; Takano et al. 1994; Christenson 2000; Christenson et al. 2010).

In fact, Mülliered (1932) had described the presence of stalactites of native sulfur near fumaroles discharging H₂S in 1932, 50 years before the 1982 eruption, similar to the timescale invoked by Christenson et al. (2010) for the formation of a pool of elemental sulfur below Crater Lake at Ruapehu.

The thermal activity before 1982 consisted of fumaroles, hot springs, H₂S emissions, and acid-sulfate alteration concentrated around and on the flanks of a central dome plugging a Somma crater (Mülliered 1932; Molina Berbeyer 1974; Templos et al. 1981), formed after an eruption occurred approximately 500 years BP (Macías et al. 2003). These manifestations were considered consistent with the existence of a hot-water hydrothermal system overlain by a relatively small volume, vapor-dominated cap (Casadevall et al. 1984; Rye et al. 1984). Most geothermal manifestations are still present today. According to Taran et al. (1998), the springs on the southern slopes of the volcano were only minor affected by the 1982 eruption. The steaming vents and hot springs described at the base of the dome plugging the Somma crater before the 1982 eruption eventually represented the top of the steam cap of the hydrothermal system (Taran et al. 2008; Taran and Peiffer 2009).

As observed at other volcanoes hosting active hydrothermal systems, El Chichón experienced a very fast transition from magmatic to hydrothermal stage following the 1982 eruption (Casadevall et al. 1984; Peiffer et al. 2011). The formation of a lake inside the crater occurred shortly after the end of the eruption, in late April 1982, and was attributed to a refilling by groundwaters (Casadevall et al. 1984), with a minor contribution by rainfall. The salinity and volume of the lake has changed considerably with time due to a group of geyser-like boiling springs, inside the crater (Soap Pools of Taran et al. 1998). The springs discharge clear highly saline (TDS>20,000 mg/kg) boiling Na-Ca-Cl waters, with a periodic behavior (Taran et al. 2008; Rouwet et al. 2008) in which vapor exhalations, with periods of few months to 1 year, alternate with periods of discharge of near-neutral CI-rich waters that directly feed the lake at rates between 10 and 30 l/s (Taran and Rouwet 2008). The composition of these Soap Pool springs (currently ~3,000±1,000 mg/l of CI) has been linked to a group of springs located on the SE and SW flanks, and Sr isotopes and Ca/Sr ratios suggest that all of them are fed by a shallow volcanic aquifer located at depths of 280 m beneath the crater, with a maximum horizontal extension of 3.5 km (Peiffer et al. 2011).
Nevertheless, considering the mechanisms that lead to the deposition of elemental S at Ruapehu, the low S contents of the SP springs, and the lake waters, could be also explained with the precipitation of elemental S at depth, as envisaged at Ruapehu and Poás volcanoes. This process may have occurred also before the 1982, leading to the formation of layer(s) of elemental S.

The occurrence of a similar cementing film in aggregates found in PDC deposits emplaced during the 2005 eruption of Ilamatepec volcano (El Salvador) (Scolamacchia et al. 2010) suggests that also during this eruption, liquid sulfur was explosively ejected. In fact, a vapor-dominated hydrothermal system is inferred to exist at low depths beneath the acid lake (Bernard et al. 2004).

**Discussion**

According to all experimental results to date (Gilbert and Lane 1994; Schumacher and Schmincke 1995; James et al. 2002; Van Eaton et al. 2012a), aggregation processes are strongly particle size dependent. Yet, the validity of such a conclusion may only apply when considering water or electrostatic attraction as binding agents between particles. Different experimental results agree in the existence of a lower limit around 15 wt% liquid water for the formation of stable spherical aggregates and an upper limit of 25 wt%, after which the aggregates began to clot. Under these conditions, the scavenging of particles of different sizes is controlled by liquid contents, but particles >1 mm are rarely accreted, unless as nucleation sites (i.e., cores) for sticking smaller particles; in addition, capillary forces are ineffective for high amounts of liquid water (>25 wt%; >50 vol%), for which the formation of stable aggregates is prevented for any particle size (Schumacher and Schmincke 1995; Van Eaton et al. 2012a, b).

Particles larger than 1 mm are instead common inside aggregates found at Poás, El Chichón, and Ilamatepec volcanoes; at El Chichón and Ilamatepec, they occur also in the most external rims, which are instead characteristically made of particles only tens of microns in size in pellets formed by the binding action of water.

Such evidence indicates that liquid sulfur is able to remove coarser particles (i.e., size of lapilli) from the eruptive clouds, with respect to those (tens to hundreds μm) removed by either water bridges or electrostatic attraction, as suggested by observations and experimental studies (see Background section).

Such a binding property would depend on sulfur viscosity. Sulfur viscosity is highly temperature dependent, varying from 0.01 Pa s around its melting point (116.3 °C, for its triclinic modification according to Cernosek et al. 2009) to values up to 3 orders of magnitude greater at 160 °C due to polymerization (Meyer 1976; Sagan 1979). At Poás, the temperature of ejected sulfur during S-eruptions in 1989 was 96.7 °C near the vents of sulfur cones and 55 °C on their flanks. Direct measurements of sulfur lakes established within the craters of sulfur cones after the eruptions ceased yielded values of 116 °C, but temperatures up to 159 °C were measured in one of them in 1990 (Oppenheimer 1992).

Nevertheless, sulfur properties depend on its thermal history and its potentially metastable state (Meyer 1964, 1976; Sagan 1979; Cernosek et al. 2009). Meyer (1976) reported at least 19 different melting points for elemental sulfur (i.e., 94 to 133 °C), which were explained by the slow kinetics of the melting process and the different reaction paths followed on cooling (resulting in various metastable mixtures of different allotropes). S-allotropes can form, or transform, under the influence of minor disturbances, such as sunlight, recrystallization, and time alone. Moreover, experimental determination of the S-phase composition evidenced that during sample preparation, a variety of allotropes formed and co-precipitate with sulfides, oxyacids, and solvents. The viscosity of sulfur around its (variable) melting point is ten times the viscosity of water at 20 °C (Sagan 1979). This much higher viscosity than water can explain both its capacity to accrete coarser particles and to cement them. Sulfur is therefore able to provide stronger bonds with respect to water, considered to date the strongest binding substance among particles of aggregates in pyroclastic deposits (Brown et al. 2012 and references therein).

The formation of sulfur layers of different spatial extent has been reported at several volcanoes with active hydrothermal systems, and this work infers a similar process at El Chichón and Ilamatepec volcanoes. All volcanoes show a common eruptive pattern, in which periods of frequent phreatic and phreatomagmatic events through a crater lake are followed by rare periods of effusive dome growth or explosive magmatic activity.

All these cases suggest that the storage of sulfur layers in andesitic volcanoes hosting active hydrothermal systems may be more common than previously assumed.

**Concluding remarks and suggestions for further work**

The poor internal sorting of particles inside aggregates described at El Chichón and Ilamatepec volcanoes suggests a faster process of accretion with respect to those described elsewhere. Such a hypothesis is confirmed by the textural features observed inside cemented aggregates, which attest that some particles were still molten at the moment of accretion, and by the distribution of the aggregates, limited to distances of few kilometers from the eruptive vents, in contrast to aggregates described elsewhere (tens to hundreds of km). Such distribution suggests that when explosively ejected, sulfur is able to cause the settling of coarser particles (>2 mm) from eruptive clouds, already at distances of few kilometers from the eruptive vents.
kilometers from eruptive vents and eventually at lower atmospheric levels, compared to what was commonly thought. In contrast to what observed in sulfur eruptions at Poás, the aggregates described here were likely transported at greater distances with respect to their locus of formation, by turbulent PDCs, in which aggregates are found.

Such evidence has important implications on the volumetric estimates of different grain sizes inside eruptive clouds and should be considered in models of particle dispersion and sedimentation to improve hazard assessment.

The maximum dimensions, the shape, and the internal structure of aggregates found at El Chichón and Ilimaatepec are similar to those of aggregates formed by the binding action of water. The only differences are the maximum grain size of individual particles accreted and the presence of a film cementing the grains. Such characteristics are not visible at the outcrop scale, but only apparent after the ultrasonic cleaning of samples (Scolamacchia et al. 2005). Considering that the accumulation of sulfur layers has been described at several volcanoes hosting hydrothermal systems, it is probable that aggregates similar to those described here also occurred in other PDC deposits but were not properly recognized because of the lack of proper analyses. Future work should concentrate on their identification, characterization, and spatial distribution in order to provide an adequate basis for modeling.

Due to the intrinsic complexity of the state of sulfur, the minimum amount necessary to stick particles of different sizes is difficult to estimate and likely represent the sum of different processes, especially when its ejection is contemporaneous with that of other fluids (e.g., hydrothermal). As suggested by textural features of aggregates showed here, eventually liquid S continued to react with nearby crystals and glass accreted. Previous studies indicate that the cooling of molten S involves different reaction paths, producing a range of metastable allotropes, and due to the high heat capacity of S-melts, the isolation of impurities is difficult (Meyer 1976). Experimental efforts should concentrate in finding solutions to determine the optimum amount of S necessary to accrete particles of different sizes.

The settling velocity of aggregates is considered to be lower than that of single particles, due to the presence of lighter fluids (air or water). Notwithstanding the fact that the bulk densities of sulfur aggregates reported by Oppenheimer (1992) at Poás range between 900 and 1,500 kg/m³, similar to those reported for aggregates elsewhere (1,200–1,600 kg/m³, Brown et al. 2012), the variations in density and porosity of aggregates with respect to single particles in the presence of S need experimental investigation. In fact, the density of solid elemental sulfur is higher than water (i.e., in a range between 1.92 and 2.07 g/cm³ depending on the allotrope), and the presence of other elements, as impurities (e.g., Fe), could contribute to an increase in the density of the binding material.

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