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# The magmatic and eruptive evolution of the 1883 caldera-forming eruption of Krakatau

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1	The magmatic and eruptive evolution of the 1883 caldera-forming eruption
2	of Krakatau: integrating field- to crystal-scale observations
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#### 22 Abstract

Explosive, caldera-forming eruptions are exceptional and hazardous volcanic phenomena. The 1883 23 24 eruption of Krakatau is the largest such event for which there are detailed contemporary written 25 accounts, allowing information on the eruptive progression to be integrated with the stratigraphy and 26 geochemistry of its products. Freshly exposed sequences of the 1883 eruptive deposits of Krakatau, 27 stripped of vegetation by a tsunami generated by the flank collapse of Anak Krakatau in 2018, shed 28 new light on the eruptive sequence. Matrix glass from the base of the stratigraphy is chemically 29 distinct and more evolved than the overlying sequence indicating the presence of a shallow, silicic 30 melt-rich region that was evacuated during the early eruptive activity from May 1883 onwards. 31 Disruption of the shallow, silicic magma may have led to the coalescence and mixing of chemically 32 similar melts representative of a range of magmatic conditions, as evidenced by complex and varied 33 plagioclase phenocryst zoning profiles. This mixing, over a period of two to three months, culminated in the onset of the climactic phase of the eruption on 26<sup>th</sup> August 1883. Pyroclastic density currents 34 35 (PDCs) emplaced during this phase of the eruption show a change in transport direction from north 36 east to south west, coinciding with the deposition of a lithic lag breccia unit. This may be attributed to 37 partial collapse of an elevated portion of the island, resulting in the removal of a topographic barrier. 38 Edifice destruction potentially further reduced the overburden on the underlying magmatic system, 39 leading to the most explosive and energetic phase of the eruption in the morning of 27<sup>th</sup> August 1883. 40 This phase of the eruption culminated in a final period of caldera collapse, which is recorded in the 41 stratigraphy as a second lithic lag breccia. The massive PDC deposits emplaced during this final phase 42 contain glassy blocks up to 8 m in size, observed for the first time in 2019, which are chemically similar 43 to the pyroclastic sequence. These blocks are interpreted as representing stagnant, shallow portions 44 of the magma reservoir disrupted during the final stages of caldera formation. This study provides new 45 evidence for the role that precursory eruptions and amalgamation of shallow crustal magma bodies 46 potentially play in the months leading up to caldera-forming eruptions.

47 **Keywords:** Caldera, stratigraphy, petrology, geochemistry, fieldwork

48

# 49 **1. Introduction**

50 The 1883 eruption of Krakatau (or Krakatoa) is one of the most infamous volcanic disasters, and 51 the first caldera-forming eruption in history to make headline news around the world (Symons et 52 al., 1888). The eruption demonstrated that violent, caldera-forming eruptions can have not only devastating local effects, but also global impacts (Verbeek, 1884; Symons et al., 1888; Simkin and 53 54 Fiske, 1983). Subsequent work on other caldera-forming systems shows that these high-55 magnitude, low-frequency events may have very long build-up phases, characterised by both 56 effusive and weakly explosive activity (e.g., Forni et al., 2018; Druitt et al., 2019). Long-term shifts 57 in eruptive style are poorly understood in general, but this is particularly the case for caldera 58 systems, as there has not yet been the opportunity to monitor any volcano during the long run-59 up through to a caldera-forming eruption; this poses challenges for the interpretation of 60 contemporary geophysical data in volcano monitoring (e.g., Newhall and Dzurisin, 1988; 61 Gottsmann and Marti, 2008; Druitt et al., 2012). Petrological tools offer a unique insight into the 62 architecture of pre-eruptive magma plumbing systems, and the conditions leading up to eruption. 63 The 1883 eruption of Krakatau is particularly useful in this regard, as it is the largest known eruption for which there are multiple documented observations in contemporary accounts, which 64 65 allow the eruption progression to be integrated with other datasets, analogous to studies of the 66 79 AD eruption of Vesuvius (e.g. Sigurdsson et al., 1982; Cioni et al., 2000).

67 Physical and chemical properties of a magma influence its ascent rate and ability to outgas, and 68 are in turn strongly linked with eruptive style (Cassidy et al., 2018). Petrological data allow pre-69 eruptive magma storage conditions, such as pressure, temperature and volatile content, to be 70 constrained. Pressure is an important constraint, as it controls water solubility, and thus viscosity, 71 and pressure estimates can also be used to infer magma storage depths. Temperature is a key parameter when estimating timescales of magmatic processes based on diffusion models (e.g.,
Costa et al., 2020). In addition, petrological data can provide information on magmatic processes
like mafic recharge, magma mixing, assimilation and fractional crystallisation (e.g., Knesel et al.,
1999; Ruprecht et al., 2012; Cassidy et al., 2015; 2016).

76 Past studies place some constraints on the structure of the upper-crustal magma storage system at Krakatau. Mandeville et al. (1996a) proposed that the 1883 eruption was fed from a chemically 77 78 and thermally zoned magma reservoir at 5 to 8 km depth. From the analysis of co-existing iron-79 titanium oxides, Mandeville et al. (1996a) inferred that a rhyodacite magma (880-890°C) overlay progressively hotter dacites (890-913°C), and andesites (980-1000°C). Fugacity of oxygen was 80 81 estimated by the same method for rhyodacitic pumice at -10.79 to  $-11.07 \log fO_2$  (or approximately 0.96 to 1.43 above nickel-nickel-oxide (NNO) buffer; assuming pressure at 100 82 83 MPa). Dahren et al. (2012) used petrological and seismic techniques to analyse the structure of the plumbing system beneath Anak Krakatau, the post-caldera volcano that in 1927 emerged 84 85 above sea-level within the 1883 Krakatau caldera. They concluded that the contemporary and 86 1883 plumbing systems were likely to have been subject to comparable structural controls, based 87 on compositionally similar phenocrysts. From mineral data, they inferred that magmas stalled in 88 three lithologically-controlled zones: 23 – 28 km (plagioclase cores), 7 – 12 km (clinopyroxene) 89 and 3 - 7 km (plagioclase rims). The only constraints on pre-eruptive magmatic water contents 90 come from analysis of volatiles by difference of glass inclusions, at 4 +/- 0.5 wt% (Mandeville et 91 al., 1996a). In this study, we provide further constraints on critical magmatic storage conditions 92 (temperature, pressure,  $H_2O$  content,  $fO_2$ ) prior to the 1883 eruption.

There is still debate regarding the main triggers involved at various stages of the 1883 eruption.
Potentially important processes include (i) fractional crystallisation, potentially leading to "second
boiling" (Camus et al., 1987; Mandeville et al., 1996a), (ii) magma mixing (Francis and Self, 1983;
Self and Wohletz, 1983) and (iii) phreatomagmatism, which has been proposed as a trigger for the

97 main explosions on the morning of 27<sup>th</sup> August (Verbeek, 1884). Self and Rampino (1981) ruled 98 out phreatomagmatism as they found no textural evidence for interaction of the magma with 99 water during fragmentation. Verbeek (1884) reported two distinct ash compositions that were 100 erupted during May 1883: dacite and a high-alumina basalt (Stehn 1929). This led Francis and Self 101 (1983) and Self and Wohletz (1983) to suggest that magma mixing triggered the initial stage of the 102 Krakatau 1883 eruption. Several studies have noted the presence of rare, banded pumice clasts 103 from the main phase of the eruption (e.g., Self and Rampino 1981), which is often used as an 104 indicator for magma mixing (Sparks et al., 1977; Andrews and Manga, 2014; Rossi et al., 2019). 105 However, the two visually distinct glasses are of very similar chemical compositions (Self, 1992). 106 Camus et al. (1987) and Mandeville et al. (1996a) suggested that fractional crystallisation was the 107 most important process prior to the 1883 eruption, increasing the SiO<sub>2</sub> content of the residual 108 melt and enriching it in volatiles. Both factors make an eruption more likely and potentially more 109 explosive (Blake, 1984). A final process recognised for the 1883 and contemporary Krakatau 110 system is assimilation of crustal material (Gardner et al., 2012). Using mineral and whole rock Sr isotope data, Gardner et al. (2013) showed that evolving basaltic andesite (Anak Krakatau) to 111 rhyodacite (1883 compositions), required 45% crystallisation, accompanied by assimilation of 5-112 113 23% carbonate or quartzo-feldspathic siltstone.

114 This study aims to integrate the known eruptive progression at Krakatau in 1883 – based on 115 historical accounts – with new studies of the stratigraphy, crystal zoning and glass geochemistry. 116 The new exposure of pyroclastic sequences from the 1883 eruption by the tsunami generated by 117 the flank collapse of Anak Krakatau in December 2018 (Grilli et al., 2019; Novellino et al., 2020), 118 means that it is possible to build significantly on prior work (e.g., Self, 1992; Mandeville et al., 119 1996a). Whole-rock and matrix glass data were collected and analysed in the context of this 120 sequence and help to constrain the chemical structure of the plumbing system, allowing the 121 magma reservoir zonation hypothesis to be tested. Thermodynamic modelling using Rhyolite-122 MELTS (Gualda et al., 2012), provides further insight into the role of fractional crystallisation prior

to the 1883 eruptions. Chemical analyses of both plagioclase and pyroxene phenocrysts at higher
 spatial resolution than previous studies (e.g., Mandeville et al., 1996a) allow the crystal growth
 history to be constrained in more detail. Furthermore, thermobarometric and hygrometric models
 provide improved constraints on magmatic conditions. These field observations, geochemical and
 petrological data shed new light on this highly active caldera system, and provide new context for
 the monitoring of the present-day activity of Anak Krakatau, as well as providing broader lessons
 applicable to other similar systems globally.

130

131 2. Geological Context

132 2.1 Tectonic setting

133 The Krakatau complex comprises four islands: Panjang, Sertung, Rakata, and Anak Krakatau 134 (Figure 1). Panjang and Sertung are remnant islands left behind after a caldera-forming eruption prior to that in 1883; Rakata is the southern remnant of a pre-existing larger island 135 136 that lay between Panjang and Sertung, the northern two thirds of which was destroyed in 137 1883 (dashed line, Figure 1). Prior to the 1883 caldera collapse, this main island of Krakatau 138 consisted of three volcanic centres aligned NNW: Perboewatan, in the north, Danan, in the 139 centre, and Rakata forming a higher peak to the south (Figure 1). Anak Krakatau is the current subaerial volcanic cone, which first emerged above sea-level in 1927 on the same alignment 140 141 as the 1883 vents, and lies between the positions of Perboewatan and Danan.

The Krakatau archipelago is part of the Sunda Arc; volcanism in this region is caused by subduction of the Indo-Australian Plate beneath the Eurasian Plate (Figure 1). Krakatau lies in the Sunda Strait, between Java and Sumatra, at the intersection of a NNE trending lineament of Quaternary volcanic edifices roughly perpendicular to the Java trench (Nishimura et al., 1992) and a fault connecting Krakatau with the Sunda Graben (e.g. Harjono et al., 1989). The Sunda Strait is extending, as Sumatra rotates relative to Java (Ninkovich, 1976; Hall and Spakman, 2002; Hall, 2012). Therefore, magmatism in the Sunda Strait is not only a function
of subduction, but also of rifting and extension (Harjono et al., 1989) associated with slabthinning and mantle upwelling beneath Krakatau (Abdurrachman et al., 2018).



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152 Figure 1: Tectonic map showing the Sunda Straits, with insert (a) showing the Java trench and (b) showing the Krakatau islands and field sites visited. The green stars in insert (b) are new field sites, 153 presented here for the first time, whereas localities matching, or very close to, the orange circles were 154 155 also visited by Self and Rampino (1986) and/or Mandeville et al. (1996b). Localities with two names 156 were visited in both field campaigns (2017 and 2019). All sites visited in 2019 had considerably more 157 exposure than observed and/or presented previously, due to erosion by the 2018 tsunami associated with a flank collapse on Anak Krakatau. The black star labelled approx. caldera centre marks the 158 159 deepest part of the caldera structure, estimated from bathymetric data from Deplus et al. (1995). The 160 dotted line represents the island prior to collapse in 1883, with Perboewatan and Danan, the 1883 161 active cones, marked as black circles. Figure based on Mandeville et al. (1996b), Schlüter et al. (2002), 162 Lunt et al. (2009), Susilohadi et al. (2009) and Dahren et al. (2012).

#### 163 2.2 Pre-1883 eruptive history

164 The ages of pre-1883 eruptions are uncertain. Drill core data suggests an eruption in the Sunda Straits at ca. 60 ka, however this cannot be definitively attributed to Krakatau (Ninkovich, 165 166 1979). The Javanese chronicle Pararaton, or the Book of Kings, describes a very large eruption, with "heavy rains of stone" in 416 AD originating from the straits of Sunda (Symons et al., 167 168 1888). However, no geological evidence presented to date substantiates this eruption. In May 169 1681, observations of earthquakes and pumice were made in the diaries of Johann Wilhelm 170 Vogel and Elias Hesse, likely pertaining to an eruption of Krakatau (Vogel, 1690; Hesse, 1690; Hesse, 1694; Verbeek, 1884). 171

172

# 173 **2.3 1883 Eruption**

The eruption of Krakatau volcano on 26<sup>th</sup> and 27<sup>th</sup> August 1883 was the culmination of at least 174 175 four months of unrest (Figure 2). The climactic eruption ejected  $18 - 21 \text{ km}^3$  of dominantly rhyodacite ejecta (9 – 10 km<sup>3</sup> dense rock equivalent; Self and Rampino, 1981) in a sequence 176 177 of pyroclastic density currents (PDCs) that swept across the straits of Sunda, causing volcanogenic tsunamis. In total there were approximately 36,000 fatalities (Self, 1992). This 178 eruption resulted in the destruction of two-thirds of the main island of Krakatau, forming a 179 submarine caldera, which manifests today as a 250 m deep depression in the seafloor (Deplus 180 181 et al., 1995). The soundwaves produced from the eruption were the greatest ever recorded in the audible range (Gorshkov, 1959), and the atmospheric effects were seen around the world, 182 with vivid sunsets observed up to a year after the eruption (Symons et al., 1888). 183

The first record of the 1883 eruption of Krakatau is for 20<sup>th</sup> May, with contemporary descriptions suggesting Vulcanian to Sub-Plinian activity from Perboewatan (Verbeek, 1885), which declined after 22<sup>nd</sup> May (Symons et al., 1888). Self (1992) suggested that this precursory 187 eruption column reached 20 km, with ash fall up to ~375 km away. There are no records of
188 activity between 23<sup>rd</sup> and 26<sup>th</sup> May (Symons et al., 1888). On 27<sup>th</sup> May a party visited the island
189 and observed explosions every 10 minutes (Verbeek, 1885). While there are no specific
190 records from 28<sup>th</sup> May to 19<sup>th</sup> June, Krakatau was reported to have continuously expelled
191 "smoke" throughout June according to the newspaper *Javasche Courant*, and Symons et al.
192 (1888) report no interruption in activity "according to reports from lighthouses... and vessels".
193 Krakatau began to "smoke heavily" again on 19<sup>th</sup> June (Simkin and Fiske, 1983) (Figure 2).

On 24<sup>th</sup> June, a second column of "smoke" was observed from Java for the first time, likely 194 195 emanating from Danan. This coincided with the reported disappearance of the summit of 196 Perboewatan (Symons et al., 1888). However, Ferzenaar, who was the last person to set foot on the island on 11<sup>th</sup> August prior to the climactic phase of the eruption, instead reported that 197 198 Danan had partially collapsed (Verbeek, 1885). The few records that exist suggest that activity continued to fluctuate. Verbeek (1885) observed "no ash" but "a hazy red glimmer" on 3rd 199 200 July, interpreted to be lava extrusion, and Symons et al. (1888) reported "continued eruptions, 201 earthquakes and occasional violent explosions" throughout July.

Plinian activity began on 26<sup>th</sup> August (Figure 2). At 2pm local time, a black eruption column
rose ~ 26 km into the atmosphere, with explosions every 10 minutes (Sturdy, 1884). By 3pm,
explosions were heard ~ 670 km away (Symons et al., 1888), and the first abnormal sea wave
was recorded in Batavia (modern-day Jakarta; Latter, 1981). By 5pm explosions were heard all
over Java (Symons et al., 1888). There was intense volcanic lightning through the night and a
strong sulphurous smell was reported on nearby ships, such as the Charles Bal (Sturdy, 1884).



209 The most powerful explosions, based on pressure deviations recorded on the Batavia 210 gasometer record (Latter, 1981), occurred at 5.30 am, 6.44 am, ~10:00 am and 10.45 am on 211 27<sup>th</sup> August (Symons et al., 1888); the third of these was the most violent, and was heard 850 km away in Singapore (Strachey 1888). Multiple tsunami waves traversed the Sunda Straits 212 from the 26<sup>th</sup> to 27<sup>th</sup> August and caused the majority of casualties (Symons et al., 1888). The 213 largest of these waves originated at approximately 10 am on 27<sup>th</sup> August (Verbeek, 1885; 214 215 Latter, 1981), broadly coinciding with the most powerful recorded explosion. The whole of the 216 northern portion of the island disappeared into the sea during caldera formation (Lindemann, 217 1884). However, the precise timing of caldera collapse has not yet been determined (Self, 218 1992). Details of historical sources are summarised in Supplementary Material 1.

219

# 220 2.4 Anak Krakatau

In 1927 Anak Krakatau, or "child of Krakatau", emerged above the sea surface, forming a new
subaerial volcanic cone (Stehn, 1929). This soon became a permanent island, which has grown
rapidly. Since the 1960s, when the active vent stopped interacting with seawater, Anak
Krakatau has predominantly erupted effusively, punctuated with Vulcanian and Strombolian
explosions (Abdurrachman et al., 2018).

A period of elevated activity from July to October 2018 culminated in the collapse of the southwestern portion of Anak Krakatau on December 22<sup>nd</sup> 2018. This created a volcanogenic tsunami which greatly impacted the coast along the straits of Sunda, killing 437 people (Grilli et al., 2019; Novellino et al., 2020).

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#### 233 **3.** Methods and material

#### 234

## 3.1 1883 Stratigraphy and samples

Field campaigns to the Krakatau islands were undertaken in September 2017 and August 2019. Some field locations, originally described by Self and Rampino (1981) and Mandeville et al. (1996b), were revisited (Figure 1b). The 2019 field campaign provided new constraints on the stratigraphic sequence, as erosion associated with the December 2018 tsunami increased exposure of the 1883 sequence considerably at all localities. Stratigraphic logs were created from field observations, and cross correlated using lithological and stratigraphic characteristics, as well as glass chemistry.

In addition to samples collected on both field campaigns, three archived samples from the British Geological Survey (BGS) were analysed: two from the mail steamer Norham Castle (08 and 72), which was 92 km from Krakatau in the Sunda Straits on 26<sup>th</sup> and 27<sup>th</sup> August 1883, and one sample of ash from "Districts of java opposite Krakatau and on volcanic island itself" (07). Field locations and sample details are presented in Analytical Data 1 in the published dataset Madden-Nadeau (2020).

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# 249 3.2 X-ray fluorescence (XRF)

A selection of samples collected through the 1883 stratigraphy were analysed for whole-rock major and minor element chemistry by X-Ray Fluorescence (XRF) at the Department of Geology, University of Leicester on a PANalytical Axios Advanced XRF spectrometer. Most of these samples were powdered pumice clasts, except for one sample comprising fine ash aggregates (U22.2), and four samples of bulk tephra, containing both ash and pumice. For the pumice analyses, multiple clasts were powdered from a single sample and analysed in a single aliquot.

257	Major and minor elements (SiO <sub>2</sub> , TiO <sub>2</sub> , Al <sub>2</sub> O <sub>3</sub> , Fe <sub>2</sub> O <sub>3</sub> , MnO, MgO, CaO, NaO <sub>2</sub> , K <sub>2</sub> O, P <sub>2</sub> O <sub>5</sub> and SO <sub>2</sub> )
258	and Loss on Ignition (LOI) were determined on fused glass beads prepared from ignited
259	powders, while trace elements (As, Ba, Ce, Co, Cr, Cs, Cu, Ga, La, Mo, Nb, Nd, Ni, Pb, Rb, Sb,
260	Sc, Se, Sn, Sr, Th, U, V, W, Y, Zn, Zr) were determined on pressed pellets.

261

262 **3.3 Scanning electron microscope (SEM)** 

Back-scatter electron (BSE) images of plagioclase and pyroxene phenocrysts and matrix glass
picked from crushed pumice samples and mounted in resin blocks, were obtained with an FEI
Quanta 650 field emission gun (FEG) scanning electron microscope (SEM) in the Department
of Earth Sciences, University of Oxford, and a Zeiss Merlin Compact FEG-SEM at the Sir William
Dunn School of Pathology, University of Oxford. Operating conditions were 20 KeV with a 15micron aperture.

269

# 270 **3.4 Electron Probe Microanalysis (EPMA)**

Phenocryst phases were analysed on a FEG CAMECA SX-5 electron microprobe at the
Department of Earth Sciences, University of Oxford. Sodium was always analysed first with a
10 s peak count time, to prevent Na migration.

Compositional profiles (n = 56) for Al, Si, Na, Ca, K, Fe, Ti, Mn and Mg were collected by Electron Probe Microanalysis (EPMA) for plagioclase phenocrysts at 15 kV acceleration voltage and 20 nA beam current, with 5-micron defocussed beam size. Point spacings in line analyses were approximately 10 microns. Points were also analysed for BSE image calibration for anorthite content with the same operating conditions. Plagioclase phenocrysts were picked from archived ash collected at the time of the eruption (BGS samples), as well as samples collected in the field. Phenocrysts were picked from samples of multiple crushed, cm-scale 281 pumices, one ash sample, one sample of ash aggregates and two bulk tephra samples282 containing both pumice and ash.

Compositional profiles for pyroxene phenocrysts picked from archived ash collected at the time of the eruption (BGS samples; n = 46) were obtained at 15 kV, with a focused beam of 20 nA for Al, Si, Na, Ca, Fe, Ti, Mn, Cr and Mg. Fe/Ti oxides (n = 419) partially included into the rim of pyroxene phenocrysts and in contact with the melt from both archive and field samples, were also analysed as points under the same conditions.

Point analyses of matrix glass, mounted in resin, were analysed on a Jeol JXA-8200 electron 288 289 Microprobe in the School of Archaeology, University of Oxford. Glass analyses were conducted 290 at 15 kV with a 5-micron defocussed beam of 6 nA for Al, Si, Na, Ca, K, Fe, Ti, Mn, Mg, P and 291 Cl. Secondary standards of a similar composition to the target glass were analysed to check 292 the accuracy of the calibration (see Supplementary Material 2). Most matrix glass analyses 293 were obtained from grains picked from multiple crushed, cm-scale pumices, however two 294 samples contained only ash, one ash-aggregates, and one was bulk tephra sample, containing 295 both pumice and ash. Multiple pumices were crushed per sample, and glass clasts were picked 296 and mounted from crushed material. We also analysed a sample of crushed obsidian, and 297 three samples taken from large glassy blocks at D2S2/U23. For each sample, analyses are based on at least two clasts, except for D1S1.2 and U23.7 where the only viable analyses came 298 299 from the same clast.

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### 3.5 Vesicularity and crystallinity

Thin sections of pumice were used to estimate phenocryst content of the 1883 samples through the stratigraphy. Five images per sample were photographed through a transmitted light microscope, and crystals were traced by hand using image processing software to provide an average estimate of crystallinity reported on a vesicle-free (VF) basis. An example of a
pictomicrograph used is in Supplementary Material 3. Average vesicularity of the 1883
samples was estimated from BSE images (five to nine grains imaged per sample), picked from
crushed samples, and thresholded using image processing software (e.g., Burgisser et. al.,
2010). Estimates for vesicularity are likely to be slightly underestimated, as a result of plane
of section effects.

311 Error will also be incurred as a result of the degree of user-defined thresholding chosen for 312 each BSE image, and differences in how phenocrysts are traced by hand. Repeat 313 measurements of thresholding and crystal tracing was carried out 10 times on a single image, 314 averaged over 5 images, to give a 1  $\sigma$  error in crystallinity estimates of +/- 0.1 %, and in 315 vesicularity of +/- 2.5 %.

316

# 317 **3.6 BSE image calibration for Plagioclase**

Back scatter electron (BSE) intensity profiles of plagioclase phenocrysts were calibrated for anorthite content using quantitative point analyses obtained by EPMA following the approach outlined by Ginibre et al. (2002). Most phenocrysts were calibrated individually where enough EMPA data was available, whilst a global calibration was used for other crystals where brightness and contrast settings on the BSE images made the images comparable. Global calibrations were only used where R<sup>2</sup> > 0.8 for the correlation between anorthite and grey scale, determined using imageJ.

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#### 329 4 Results

330

# 4.1 Fieldwork and stratigraphy

331 The stratigraphic sequence of the 1883 eruptive deposits has previously been established by 332 Self and Rampino (1981) and Mandeville et al. (1996b), however, the significantly increased 333 exposure of deposits in coastal cliffs following the 2018 tsunami made this worth revisiting. 334 The stratigraphy as presented by Self (1992) (after Self and Rampino, 1981) and Mandeville et 335 al. (1996b), is shown in Figure 3. Terminology for deposits pertaining to pyroclastic density 336 currents (PDCs), and particularly dilute-PDCs, has been subject to some ambiguity. Here, we use the terms PDC deposit to refer to any unit with characteristics consistent with flow-driven 337 338 transport and sedimentation; and we use the term dilute-PDC deposit specifically for those PDC units which show cross stratification and are generally finer-grained and better sorted, 339 following Branney and Kokelaar (2002). Previous authors have used the terms ignimbrite to 340 refer to PDC deposits with more massive or poorly sorted characteristics, and surge deposits 341 342 for those with dilute-PDC deposit characteristics. However, when discussing stratigraphic 343 descriptions of previous authors we use their original terminology.

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345

4.1.1. Previous work

346 Self and Rampino (1981) reported that the proximal stratigraphy comprised sub-347 Plinian fall deposits interbedded with surge (i.e., dilute-PDC) deposits up to 20 m thick, 348 overlain by up to 55 m of massive ignimbrite (i.e., PDC deposit). Mandeville et al. (1996b) reported a layer of olive- to bluish-grey, fine-ash fall deposit up to 4 cm thick 349 350 at the base of the 1883 deposit, which overlies a soil horizon on West Rakata and West 351 Panjang (equivalent to our localities U24/D1S1 and NP3/D3S1, respectively). They 352 attributed this to phreatomagmatic activity in May to August 1883. Next in the 353sequence, Mandeville et al., (1996b) reported 5 - 20 cm of coarse, light-grey pumice354fall deposit, followed by 4 to 6 m of fall deposits interbedded with surge deposits. At355the top of the stratigraphy, Mandeville et al. (1996b) report thick accumulations of356massive pyroclastic flow (i.e., PDC) deposits. The thickness of the fall deposit layer is357therefore disputed, with Self and Rampino (1981) observing fall deposits interbedded358with surges up to 20 m thick, and Mandeville et al. (1996b) observing up to 6.2 m of359fall deposit interbedded with surge deposits.



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# **4.1.2. This study**

365 We present our new composite stratigraphy in Figure 3; field logs, stratigraphic 366 correlations and field photographs for each locality are shown in Supplementary Material 4 – 11. Like Mandeville et al. (1996b), we found a distinctive olive-green ash fall deposit at the base of the sequence (Figure 3), which we term Unit A (Figure 3; 4a). Unit A is between 2 and 5 cm thick and was only found exposed on West Rakata and South Panjang. This layer overlies a red paleosol, and is composed of fine-ash aggregates. A back-scatter electron image of the ash aggregates can be found in Supplementary Material 12.



Figure 4: Panel (a) shows Unit A, as observed from locality MS-Swim1 on South Panjang, which is a thin layer of olive-green ash aggregates, overlying a red paleosol. Panel (b) shows the base of Unit B at locality D4S1/U20 on North Panjang, which is composed of clast-supported (C-S) fallout layers, interbedded with dilute-PDC deposits that show cross-stratification (X-strat). Panel (c) shows the units overlying those in panel (b), at the same locality (D4S1/U20). These are also part of Unit B, and show matrix-supported (M-S) dilute-PDC deposits interbedded with pumice-supported lenses. All locations are shown in Figure 1. People (b) or measuring tape (a and c) for scale; measuring tape is 1 m in (c).

381

Unit B is characterised by well-sorted beds of angular lapilli, interbedded with poorly 382 383 sorted layers which are matrix-supported (Figure 4b; 4c; 5a). Clasts within the well-384 sorted layers are predominantly juvenile pumice (>80%), the remainder being dense, 385 angular, and visibly altered volcanic lithics; Mandeville et al. (1996b) determined that 386 the majority of these lithics are basalt and basaltic andesite. Of the juvenile clasts, 387  $\sim$ 90% are white pumice, although pink, grey and yellow pumice are also observed. 388 Clasts within the poorly sorted units show similar proportions. Some of the poorly sorted beds are cross-bedded, and interpreted as dilute-PDC deposits (after Branney 389 390 and Kokelaar, 2002). These dilute-PDC deposits also contain laterally restricted, 391 discontinuous lenses of well-sorted, sub-rounded, pumiceous lapilli. Unit B is 392 interpreted to comprise fall deposits interbedded with PDC and dilute-PDC deposits. 393 We find that Unit B is up to 20 m thick, in agreement with observations made by Self 394 and Rampino (1981) (Figure 3). Charcoal and tree moulds were found towards the 395 base of Unit B at two localities. Carbonised logs were also reported by Mandeville et 396 al. (1996b), but not attributed to a specific unit within the stratigraphy.

397 Next in the sequence is Unit C, which is characterised by lithic blocks (up to 50 cm) in 398 a poorly sorted, juvenile matrix (Figure 5a; 5b). This section of the sequence is 399 interpreted to be a lithic lag breccia (Druitt and Sparks, 1982; Branney and Kokelaar, 400 2002). Both Self and Rampino (1981) and Mandeville et al. (1996b) identify lithic lag 401 breccias in the sequence (Figure 3), however only Self and Rampino (1981) used them 402 as a correlating horizon. The lag breccia is variable in stratigraphic thickness (0.3 to 4 403 m), and bifurcates in some outcrops (Figure 4a). The proportion of lithic blocks within 404 this unit also varies between localities.

405 Unit D is a massive, poorly sorted, matrix-supported unit containing predominantly 406 pumice clasts (80-90% of clasts) in an ash-rich matrix (Figure 5b). Both Self and Rampino (1981) and Mandeville et al. (1996b) identify a similar unit towards the top 407 of the sequence (Figure 3). The structureless nature of Unit D suggests it was likely 408 409 deposited by large volume, high-concentration PDCs. Another characteristic feature of Unit D is the presence of obsidian clasts. Frothy, glassy, and banded obsidian clasts 410 411 are present (e.g., Shields et al., 2016), making it likely that the obsidian is juvenile (Self 412 and Rampino, 1981). Rare black and white banded pumices, as reported by Self and Rampino (1981), were also observed. 413



Figure 5: Panel (a) shows Units B, C and part of Unit D, as observed from locality D2S3a/U13, North
Sertung. Unit B comprises clast-supported (C-S) pumice layers interbedded with matrix-supported (MS) layers. Panel (b) shows Units C and D at locality D3S2/NP4, North Panjang. Unit C in both panels (a)
and (b) show lithic-rich (L-R) layers. Panel (c) shows the top of Unit D at locality D2S2/U23, South

Sertung. Unit D is comprised of massive PDC deposits in both panels (b) and (c). In panel (b), there is
a second lithic-rich layer at the very top of the sequence. In panel (c) there are large, glassy blocks up
to 8 m in size. All locations are shown in Figure 1. People for scale.

422

423 A second lithic lag breccia horizon is observed at the top of Unit D on North Panjang 424 (Figure 4b), which concurs with an updated stratigraphic log presented by Self (1992) 425 (Figure 3). New exposure on South Sertung (D2S2/U23, Figure 1), observed for the 426 first time in the August 2019 field campaign, contains large glassy blocks up to 8 m in length within the massive PDC unit (Figure 5c). Although the blocks are intact, they 427 428 are intensely fractured and have sub-rounded irregular shapes, aligned broadly 429 horizontally, but not confined to a single horizon within the deposit (Figure 4c). Some 430 of these fractured blocks are black in colour and vitreous, and look similar to the 431 smaller obsidian clasts already identified by Self and Rampino (1981), with a low 432 phenocryst content, whereas other blocks are dull grey in colour, with a higher 433 phenocryst content (comparable to pitchstone). Their geochemistry will be discussed 434 further in section 4.5. This section of Unit D also contains clasts of mudstone. We also 435 noted crude horizontal stratification of the PDCs delineated by subtle colour changes; this was also observed in massive PDC units described by Mandeville et al. (1996b). 436

Locality U22 (Figure 5) is the only outcrop where the entire sequence (Units A to D) can be observed. The sequence appears to be condensed (2.8 m), and we use this as a type locality. Key marker beds in the 1883 stratigraphic sequence include: a thin, green, ash-aggregate layer overlying a red paleosol at the very base of the sequence, delineating Unit A (Mandeville et al., 1996b; Figure 3; 6a); Pumice fallout units interbedded with PDC and dilute-PDC deposits (Self and Rampino, 1981; Figure 3), some of which contain charcoal aligned east-west (Unit B; Figure 6b); the lithic lag breccia overlying Unit B (Unit C; Figure 6c); and Unit D, consisting of massive PDC
deposits containing obsidian (Self and Rampino, 1981; Mandeville et al., 1996b; Figure
3) (Figure 6c). Figure 7 shows the logs from each locality cross-correlated by unit.



447

Figure 6: Photographs and log of deposit at locality U22 to show the entire sequence (E.
Rakata; location shown in Figure 1). Panel (a) shows Unit A at the base of the sequence, (b)
shows Unit B, (c) shows Units C and D, and (d) shows the cross correlated stratigraphic log,
with sample numbers down the right-hand side. C-S stands for clast-supported, X-strat for

452 cross-stratification, M-S for matrix-supported and L-R for lithic-rich. Measuring tape for scale

453 (a-c).



**Figure 7:** Cross correlated stratigraphic logs from all localities visited. It should be noted that the scale varies between logs. Insert map (a) shows the location of each locality, with the locations visited by Mandeville et al. (1996) and Self and Rampino (1986) shown in orange, and fieldsites new to this study indicated by a green star. Letters represent the discussed units, with A corresponding to blue, B to orange, C to yellow and D to green. Bars below logs represent the relative thicknesses of each unit. Symbols by sample numbers represent thin sections and chemical analyses. Distance above log is from the assumed caldera centre, taken from Deplus et al. (1995).

The spatial distribution and thicknesses of PDC (including dilute-PDC) deposits change 455 456 through the stratigraphic sequence (Figure 8). Unit B is thickest to the north east, with 457 approximately 20 m of vertical exposure (Figure 8a). Unit C appears to be more evenly distributed around the main island (Figure 8b), whereas Unit D is thickest in the south 458 459 west (14 m; Figure 8c), rather than north as reported by Self and Rampino (1981). 460 These thicknesses are based on only limited exposures, and there are only two localities where it was possible to observe the base of the sequence (MS-Swim1; south 461 462 Panjang and U22; east Rakata); this will lead to underestimates of unit thickness. Self 463 and Rampino (1981) noted a lack of fall deposits in the south west, which we confirmed. 464



466

#### 4. 2 Vesicularity and crystallinity

467 Crystallinity of juvenile clasts increase up the stratigraphic sequence. Unit A (n = 1) is 468 comprised of poorly vesicular ash aggregates, with crystallinity (on a vesicle-free 469 (VF)/porosity-free basis) at 5 %. In Unit B (n = 7), crystallinity (VF) ranges from 10 - 20 % and 470 Unit D (n = 3) has a crystallinity (VF) of 30 %. Vesciularity in both Units B and D ranges from 70 471 - 80 %.

472

#### 473 4.3 XRF whole-rock chemistry

474 XRF whole-rock data are presented in Analytical Data 2 in the dataset Madden-Nadeau (2020). 475 The 1883 whole-rock samples have anomalous major element compositions compared with 476 previously reported values (Oba et al., 1982; Self 1992; Mandeville et al., 1996a; Turner and Foden 2001; and Gardner et al., 2013). These samples also have high Loss on Ignition (LOI; 3.1 477 478 - 8 wt%). We suspect these samples have experienced seawater alteration, based on the 479 positive correlation between LOI and Na<sub>2</sub>O (Figure 9a), and negative correlation with SiO<sub>2</sub> 480 (Figure 9b). The values for major elements are anomalous, even after correcting for LOI, in all 481 but two of the samples, and thus cannot be used to assess the chemostratigraphy.

482 Trace element compositions show no evidence for alteration, when plotted against Zr, which 483 is a high field-strength element, assumed to be immobile (Figure 9c-f). Correlations between 484 Zr and other incompatible elements provide no evidence for loss or gain of these elements 485 during alteration, as the ratios between them remain constant. Zr vs Y (Figure 9e) has the 486 strongest positive correlation, with an R<sup>2</sup> value of 0.87. Once normalised to LOI to correct for 487 the dilution effect (Figure 9d and f), there is no discernible systematic correlation with 488 stratigraphic height for Ba and Y with Zr.



Figure 9: (a) a plot showing a positive correlation between LOI vs Na<sub>2</sub>O and (b) the negative correlation between LOI and SiO<sub>2</sub>, indicating that the samples are likely to have undergone seawater alteration.
(c) and (d) show Ba plotted against Zr, which is assumed to be immobile, with (d) showing the data corrected for LOI (transparent symbols). (e) and (f) show Y against Zr, with (f) being corrected for LOI (transparent symbols). (e) and (f) show Y against Zr, with (f) being corrected for LOI (transparent symbols). The dashed lines are the least squares linear regression, and in (c) and (e) represent the constant ratio between the two trace elements, and thus no losses or gains during alteration. The higher the R<sup>2</sup> value, the less those elements have been lost or gained during alteration.

- (e) Y vs Zr has the highest R<sup>2</sup> value of 0.87, thus being the elements least affected by alteration, and
  (f) shows that there is no correlation in chemistry of Y and Zr with stratigraphic height.
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# 1 4.4 Phenocryst textures and chemistry

#### 502 **4.4.1 Plagioclase**

503 Plagioclase is the dominant phenocryst phase in pumice clasts throughout the 1883 504 eruption stratigraphy and, as is typical of subduction settings, most plagioclase crystals 505 are zoned. When discussed here, cores and rims refer to the centre and edge of euhedral, 506 unbroken crystals. Zoning profiles vary between crystals from very complex to unzoned 507 (Figure 10). Anorthite (An = molar Ca/(Ca+Na)) compositions present across all sampled 508 plagioclase phenocryst cores, ranges from An<sub>29</sub> to An<sub>90</sub> (mean An<sub>54</sub>; 95 analysed crystals), 509 bearing in mind that the 'core' may be subject to section effects. Phenocrysts also have a 510 wide range in rim compositions:  $An_{26}$  to  $An_{65}$  (mean  $An_{48}$ ; 77 analysed crystals) (Figure 11). 511 There is no discernible inter-unit variation in crystal textures, or the core and rim anorthite 512 contents. There is also no correlation between anorthite content of the rims and cores. Resorption textures are common (e.g., Figure 10a), with some crystals having patchy cores 513 and/or zones (~ 30 %; e.g., Figure 10c). Both normal (~ 35 % of rims) and reverse zoning 514 515 (~ 60 % of rims) are also common at various stages in plagioclase crystallisation histories, 516 with  $\sim$  5 % of crystals showing no zoning at all (Figure 10e). The anorthite content of crystals does not converge on a single value at the rims. A high anorthite ( $>An_{70}$ ) core is 517 observed in ~ 20 % of phenocrysts (e.g., Figure 10a). Due to the complexity in the 518 plagioclase phenocryst zoning profiles, it is not possible to identify discrete typologies, or 519 common phases of growth, as has been achieved in previous studies (e.g., Druitt et al., 520 521 2012). All whole plagioclase anorthite traverses, from euhedral rim to core, are shown on 522 a single plot in Supplementary Material 13.

523 Analytical data for plagioclase phenocrysts, with accompanying BSE images, from Units B 524 to D and archived BGS samples, can be found in Analytical Data 3 – 11 in the dataset 525 Madden-Nadeau (2020). Analytical data for Unit A, and one Unit B sample (U22.3) can be 526 found in Supplementary Material 14.



527

Figure 10: BSE images of a selection of plagioclase phenocrysts, with anorthite traverses shown in
red. Note that the BSE grey scale differs between each image. Plagioclase (a) has a high anorthite
core, followed by oscillatory zoning, resorption textures and a reverse zone at the rim; (b) has a
single reverse zone between core and mantle, with normal zoning at the rim; (c) has a high An core,

followed by a patchy, reabsorbed zone, then oscillatory zoning, with normal zoning at the rim; (d) has an  $An_{46}$  core with resorption textures, followed by oscillatory zoning, with the rim being reverse zoned; (e) appears largely unzoned with consistent composition of ~ $An_{50}$  and (f) has an  $An_{47}$  core with resorption textures, followed by two reverse zones.

536



**Figure 11:** Histogram showing the frequency of core and rim compositions in anorthite for plagioclase phenocrysts; 97 individual crystals were used to create this histogram, taken from the 3 archive samples, as well as 4 samples from Units B, 3 samples from Unit D, and one sample from Unit A. The three inserted histograms show distributions of anorthite compositions for Unit A, B and D individually. It should be noted that there will be some error in crystal core compositions as a result

of plane of section effects. Not all crystals had both viable core and rim analyses, as rim analyses wereonly included for euhedral, unbroken phenocrysts.

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546

# 4.4.2 Pyroxene and Fe-Ti oxides

547	Pyroxene phenocrysts were picked from archived BGS ash samples, collected at the time
548	of the eruption. When discussed here, cores and rims refer to the centre and edge of
549	euhedral, unbroken crystals, and cores may be subject to plane of section effects.
550	Orthopyroxene rims and cores range in Mg# (Mg# = molar Mg/(Mg+Fe)) from 0.67 to 0.72
551	(n = 22, $\sigma$ = 0.1), whilst the range for clinopyroxene is 0.73 to 0.77 (n = 23, $\sigma$ = 0.1).
552	Pyroxene phenocrysts are largely unzoned in major elements. All data for pyroxene
553	phenocrysts can be found in Analytical Data 12 – 14 in the dataset Madden-Nadeau
554	(2020), and a plot showing all whole pyroxene traverses from euhedral rim to core, is
555	shown in Supplementary Material 15.

Fe-Ti oxides (n = 419) partially included in pyroxene rims, and in contact with the melt, were analysed from Units B to D in the stratigraphy; they are magnetite and ilmenite. All chemical data for Fe-Ti oxides can be found in Analytical Data 15 in the dataset Madden-Nadeau (2020).

560

# 561 4.5 Matrix glass

562 Matrix glass data for the 1883 pyroclastic sequence normalised to 100 % anhydrous 563 compositions are plotted in Figure 12, with full data table in Analytical Data 16 in the dataset 564 Madden-Nadeau (2020). Totals range from 96.0 to 101.2 wt % prior to normalisation. Matrix 565 glasses from Unit A are distinct, and more evolved than the rest of the proximal 1883 566 sequence, with SiO<sub>2</sub> ranging from 72.6 to 74.7 wt%. Data for Units B, C and D overlap, with a broad trend towards marginally less evolved compositions, showing a slight decrease in both
SiO<sub>2</sub> and total alkalis, moving up the sequence (Figure 12). SiO<sub>2</sub> for Unit B glasses range from
71.4 to 72.7 wt%, Unit C from 70.9 to 72.3 wt%, with Unit D ranging from 70.6 to 72.3 wt%.

570 Matrix glass of an obsidian clast and samples from three of the large glassy blocks on the south 571 of Sertung (U23/D2S2.2) within Unit D were also analysed. Matrix glass from the blocks and 572 smaller obsidian fragments are broadly similar to the matrix glass of pumice and ash ejected 573 in the 1883 eruption, extending to more evolved compositions, with SiO<sub>2</sub> ranging from 71.8 to 574 77.6 wt% (Figure 12). Full chemical data for the matrix glass of the blocks can be found in 575 Supplementary Material 16.



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578

**Figure 12:** (a) Matrix glass data on a total alkali and silica diagram (Le Bas et al., 1986). Colours of symbols and transparent areas correspond to stratigraphic units: A = blue, B = orange, C =

yellow and D = green, as in Figure 6. Each symbol represents a separate sample, with the key
in stratigraphic order. Unit A is more evolved than, and distinct from, Units B to D. Average
error is given to 1 σ. (b) shows the same data, overlain with data from the glassy blocks and
obsidian found within Unit D at U23/D2S2 on south of Sertung.

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# 4.6.1 Fe-Ti oxides

4.6 Thermobarometry

586Pre-eruptive temperatures and oxygen fugacities of the 1883 magmas were estimated587from the compositions of titanomagnetite and hemoilmenite crystals, using the Fe2+-588Mg exchange model of Ghiorso and Evans (2008). Fe-Ti oxide pairs were tested for589equilibrium using the Mg-Mn equilibrium line (Bacon and Hirchsmann, 1988). Fe-Ti590oxides are likely to yield the last equilibration temperature before quenching591(Rutherford and Devine, 1988; Geschwind and Rutherford, 1992; Lindsley and Frost,5921992).

593 The temperature range for all Fe-Ti oxide pairs (n = 64; 24 touching, 47 partially 594 included within the same pyroxene phenocryst and in contact with the melt; chemical 595 data in Analytical Data 15; dataset Madden-Nadeau 2020) was 890 to 935 °C, with a mean value of 914 +/- 50 °C. The range of temperatures generated for the 24 touching 596 597 Fe/Ti oxide pairs is between 891 and 935 °C, with an average of 913 °C. The oxygen fugacity (fO<sub>2</sub>) range for all Fe-Ti oxide pairs is 0.6 to 0.85 log units above the nickel-598 599 nickel-oxide (NNO) buffer, with the mean being NNO + 0.76. The same range of values 600 for  $fO_2$  is seen within the data set for the 24 touching pairs (average of NNO + 0.74). 601 The temperatures and  $fO_2$  of Units B, C and D all show considerable overlap. A 602 temperature fO<sub>2</sub> plot can be found in Supplementary Material 17. These temperatures 603 are interpreted as the final pre-eruption equilibration temperature.

604

#### 4.6.2 Plagioclase hygrometer

605 Pre-eruptive dissolved water content of the melt for samples from Units A, B and D 606 are estimated using a plagioclase-melt hygrometer (Waters and Lange, 2015) which is 607 based on the crystal-liquid exchange reaction between the anorthite and albite 608 components. From the anorthite contents of plagioclase rims and matrix glass data (n = 59), and the mean Fe/Ti oxide temperature (914  $^{\circ}$ C), we infer a mean H<sub>2</sub>O content 609 610 of 3.4 wt% for Unit A, 3.6 wt % for Unit B, and 3.5 wt% for Unit D. Allowing for 611 uncertainty in Fe/Ti oxide temperatures (890 to 935 °C), gives a range of water contents from 2.6 to 4.3 wt% for all three units. These results are consistent over 612 613 pressure inputs of 100 to 250 MPa. It should be noted that these are estimates of the 614 final pre-eruptive H<sub>2</sub>O contents, and will not reflect the entire range of conditions 615 experienced by plagioclase phenocrysts throughout their crystallisation histories.

616Equilibrium in plagioclase is difficult to test, owing to composition being a function of617both temperature and H2O content, however only anorthite compositions for618plagioclase with paired liquid anorthite compositions that overlap with the619compositional data for which the hygrometer was calibrated has been used to620generate these estimates of water content (Supplementary Material 18).

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622

#### 4.7 Rhyolite-MELTS modelling

623 Melt evolution was modelled using Rhyolite-MELTs (Gualda et al., 2012) (Figure 13). The 624 starting composition is the most primitive composition (4.63 wt% MgO) from Anak 625 Krakatau analysed by Dahren et al. (2012). Initial H<sub>2</sub>O contents of 1 - 3.5 wt% (steps of 0.5 626 wt%; varied between runs) were modelled isobarically. Pressure was varied between runs 627 from 50 - 300 MPa (steps of 50 MPa). In all runs, temperature was dropped from 1200 to 628 700 °C in increments of 2 °C. The lines of descent that fit the best with whole-rock and

629	matrix glass compositions for the 1883 pyroclastic sequence were modelled at an initial
630	$H_2O$ content of 1.5 wt%, and lie between 125 and 250 MPa (Figure 13; dashed lines). The
631	evolution of $H_2O$ within the melt indicates that it is water-undersaturated prior to
632	eruption under these conditions, as otherwise the $H_2O$ content would have plateaued on
633	the $H_2O$ vs SiO <sub>2</sub> plot in Figure 13(f). Varying the initial water content has a much larger
634	effect on the liquid line of descent than varying the pressure.



635

Figure 13: (a) shows a total alkali and silica diagram (Le Bas et al., 1986), and (b) to (e) show Harker
diagrams containing matrix glass analyses from 1883 samples analysed in this study (MG, coloured
squares), as well as matrix glass and whole rock analyses from previous studies (Colourless squares
and diamonds, respectively; Oba et al., 1982; Self 1992; Mandeville et al., 1996a; Turner and Foden
2001; and Gardner et al., 2013). Colours denote the different stratigraphic units, as presented in Figure
7. Matrix glass data are all normalised to 100 % anhydrous. Liquid lines of descent are shown, modelled

642 isobarically in Rhyolite-MELTS (Gualda et al., 2012), with (f) showing how H<sub>2</sub>O content varies with SiO<sub>2</sub> 643 in these models. Red lines were modelled with an initial  $H_2O$  content of 1 wt%, orange 1.5 wt%, yellow 644 2 wt%, green 2.5 wt%, blue 3 wt% and purple 3.5 wt%. Arrows indicate direction of increasing pressure 645 (steps of 50 MPa from 50 to 300 MPa) between liquid lines of descent modelled at the same initial 646 water content (same colour). The effect of varying the initial H<sub>2</sub>O content in these models is much 647 greater than that of pressure. The dashed lines were modelled at 125 MPa and 250 MPa, with an initial 648 H<sub>2</sub>O content of 1.5 wt%, and liquid lines of descent that fall between these lines provide best estimates 649 of the conditions under which the 1883 melt evolved. The input composition is indicated by the star 650 and is from Anak Krakatau (Darhen et al., 2012). The triangles in (f) indicate  $H_2O$  contents of the 651 average melt composition for Units A, B and D, as modelled using plagioclase hygrometry (Waters and 652 Lange, 2015) using the average temperature generated by Fe/Ti oxide thermometry (914 °C; Ghiorso and Evans, 2008). Standard error of 0.35 wt% is indicated by the error bars, along with 2  $\sigma$  of the 653 654 average  $SiO_2$  content. Where the lines flatten in (f) represents water saturation of the melt at the 655 corresponding pressure.

656

#### 657 5 Discussion

658 5.1 Structure of plumbing system

The structure of the pre-eruption magma plumbing system remains a key unknown, and yet an understanding of the reservoir architecture is critical to assessing long-term volcanic hazards (Edmonds, 2008). The lack of systematic change in trace element chemistry with stratigraphic height (Figure 9) is not consistent with the postulated existence of a single chemically zoned, and sequentially tapped, magma reservoir at shallow depths, as suggested previously for the 1883 system (Mandeville et al., 1996a; Gardner et al., 2013), and contrasts with the prominent chemical zonations recognised in the deposits of some other large 666 magnitude explosive eruptions, such as the Green Tuff, Pantelleria and the Bishop Tuff, 667 California (Williams et al., 2014; Hildreth and Wilson, 2007).

668 Fe-Ti oxide temperatures reported by Mandeville et al., (1996a), based on the Anderson and Lindsley (1988) model, were interpreted as evidence for a stratified magma reservoir, with 669 homogenous rhyodacite (880-890 °C) overlying dacite (890-913 °C), and andesite (980-1000 670 °C). However, the more mafic components form only a minor constituent of the erupted 671 volume (~ 6 %), and there is no evidence that their frequency varies with stratigraphic height. 672 673 We also found no evidence for these lower silica compositions as a juvenile melt component in the samples analysed in this study. Our Fe-Ti oxide temperature estimates, outlined in 674 675 section 4.6.1 and based on the Ghiorso and Evans (2008) model, show no variation with 676 stratigraphic height from Units B to D, with an average final equilibration temperature of 914 677 °C and a narrower range than previously reported (890 to 935 °C). The lack of systematic variation in temperature with stratigraphic height is not consistent with the existence of a 678 simply zoned magma reservoir. 679

The diverse and heterogeneous populations of plagioclase phenocrysts, which display a large 680 681 range of crystallisation histories over the entire stratigraphic sequence (Figure 10 and 11), is 682 not easy to reconcile with a zoned magma reservoir. Patchy cores/zones are observed in 30% of crystals, however some have higher anorthite compositions and show resorption (e.g., 683 Figure 10c), while others show the reverse (e.g., Figure 10a). Plagioclase rim compositions 684 685 vary between An<sub>26</sub> to An<sub>65</sub>, with no clear modality, indicating a spectrum of different 686 crystallisation conditions, and requiring crystals to have been mixed only shortly prior to 687 eruption to prevent their rims from equilibrating (Figure 11). Normal zoning of rims accounts 688 for 35 % of crystals, with 60 % being reverse zoned; there is no convergence towards a single 689 anorthite content, as might be expected from simple homogenisation of two or more discrete 690 magma compositions. Unzoned phenocrysts account for 5% of the population (e.g., Figure 691 10e). This variability in the phenocryst population is inconsistent with a simple zoned magma
692 reservoir, and instead suggests that these phenocrysts grew under a range of different
693 conditions until shortly before eruption.

The matrix glass in Unit A is chemically distinct and more evolved than the rest of the sequence 694 (Figure 12; with the exception of the Unit D glassy blocks, discussed below), indicating the 695 presence of a more silica-rich, and likely shallower melt-rich region that was erupted first. The 696 697 matrix glass SiO<sub>2</sub> contents for Units B to D (Figure 12) show subtle compositional differences 698 with stratigraphic height, with B presenting slightly more evolved compositions and spanning a wider range of total alkali contents than Units C and D. The broad compositional consistency 699 700 means that the melt present throughout the shallow system was likely of similar bulk 701 compositions, with the variations in total alkali content (Figure 14a) within the 2 sigma error. 702 The subtle trend towards a slightly more homogenous and less evolved melt is more 703 consistent with syn-eruptive mixing of chemically similar melts, than a large-scale zoned 704 reservoir.

705 Our rhyolite-MELTS modelling suggest that the shallow magma system lies at 125 – 250 MPa 706 (Figure 13), equivalent to approximately 5 to 10 km within the crust, consistent with the 5-8707 km depths estimated by Mandeville et al. (1996a) from plagioclase melt barometry (Housh 708 and Luhr, 1991). At present-day Anak Krakatau, Dahren et al. (2012) inferred that 709 crystallisation of plagioclase and two pyroxenes takes place at 4 to 8 km, suggesting that the 710 depth range of the pre-2018 magma plumbing system is little different from that of 1883, due 711 to lithological controls on reservoir formation; these depths are corroborated by seismic 712 tomography data (Jaxybulatov et al., 2011; Harajono et al., 1989).

In contrast to the complex and varied zoning profiles observed in the plagioclase phenocryst
population (Figures 10 and 11), which suggests that they grew under a variety of conditions,
pyroxene phenocrysts lack strong zoning. Muted zoning in pyroxene phenocrysts would

therefore either suggest consistent magmatic conditions for growth, or that the crystals were
 resident in hot magma long enough for diffusion to smooth out their zoning profiles.

To determine whether the zoning in pyroxene phenocrysts was smoothed out via timedependant diffusion, we can estimate diffusion timescales for pyroxene and plagioclase from
the 1-dimensional diffusion equation:

721 
$$x = \sqrt{Dt}$$

where x is the mean diffusion length, D is the diffusion coefficient and t is time. We have 722 chosen to investigate Mg diffusion, as Mg zoning is present in plagioclase but is considerably 723 muted in pyroxene. Given a diffusion coefficient for Mg in clinopyroxene (5x10<sup>-21</sup> m<sup>2</sup>s<sup>-1</sup>; 724 725 Schwandt et al., 1998), it would take centuries for diffusion to homogenise a length scale of 726 order 10 microns. When the above diffusion equation is applied to plagioclase, for the same 727 temperature, over the same distance, using the Mg diffusion coefficient given by LaTourrette and Wasserburg (1998) of  $7.19 \times 10^{-18} \text{ m}^2/\text{s}$ , the timescales are in the order of months to years. 728 729 Therefore, both plagioclase and pyroxene phenocrysts could not have been resident in a hot 730 magma for sufficient time to smooth the Mg profiles in the pyroxene population, as significant 731 Mg variations in plagioclase are retained. This implies that diffusion is unlikely to be the cause 732 of the discrepancy in zoning complexity between the two crystal populations.

733 The compositional consistency in matrix glass (Figure 12) and pyroxene phenocryst chemistry 734 means that the complex crystallisation histories recorded by plagioclase are therefore likely to reflect variations in magmatic conditions e.g., P<sub>H2O</sub>, temperature, fO<sub>2</sub>, rather than melt 735 composition. Plagioclase crystallisation is more sensitive to these changes in magmatic 736 737 conditions than other phenocryst phases (e.g., Mollo et al., 2011), meaning that, in a system where the melt composition was similar but other conditions varied, plagioclase could record 738 complex crystallisation histories, whilst pyroxene remain largely unzoned. One hypothesis for 739 740 generating varied magmatic conditions would be the existence of multiple, discrete melt-rich

regions of similar composition. To investigate the crystallisation histories in more detail, and
test this hypothesis, automated processing of a much larger sample of phenocrysts could be
used to identify common growth zones, and events within plagioclase zoning profiles (e.g.,
Cheng et al., 2017; Probst et al., 2018).

745 The presence of anorthite-rich (An<sub>70</sub> to An<sub>92</sub>) plagioclase cores in 20 % of plagioclase phenocrysts (e.g., Figure 10a) likely indicates the existence of a deeper, more mafic reservoir 746 747 feeding the shallow system, as suggested by Mandeville et al. (1996a) for the 1883 eruption, 748 and Dahren et al. (2012) for the Anak Krakatau system. Highly anorthitic plagioclase may reflect growth in a deeper, more mafic melt at earlier stages in the development of the 749 750 reservoir. The relatively homogeneous glass chemistry suggests these more mafic 751 compositions were largely erased by the time the reservoir had matured to its pre-eruptive 752 state, as the alternative would require very efficient mixing of the melt phase. The range of 753 high-anorthite plagioclase core compositions (An<sub>70</sub> to An<sub>91</sub>; Figure 11) indicates that they grew under differing conditions (e.g., fO<sub>2</sub>, T, P, H<sub>2</sub>O content) within the deeper system. Sieve 754 755 textures in high anorthite cores, as well as strong resorption textures, followed by lower 756 anorthite mantles (e.g., Figure 11a), are most consistent with decompression (e.g., Nelson and 757 Montana, 1992; Viccaro et al., 2010), potentially during extraction from this deeper reservoir, 758 and subsequent mixing with magmas in the shallow system. The presence of a large range of 759 different crystal textures in significant proportions is also consistent with accumulation of the 760 magma body through many different episodes of mixing (e.g., El Chichón; Andrews et al., 761 2008). The lack of significant variation in matrix glass data (Figure 12) can be explained if the 762 1883 melt is sourced from a similar host composition.

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#### 5.2 Role of pre-eruptive fractional crystallisation

The importance of fractional crystallisation processes in both the compositional evolution of the magma, and in concentrating volatiles, varies greatly between volcanoes, with other processes such as magma mixing often cited as eruption triggers (e.g., Sparks et al., 1977). It is thus important to discuss the role of fractional crystallisation in the magmatic system prior to the 1883 eruption of Krakatau.

Mandeville et al. (1996a) conclude that fractional crystallisation is likely to have been the dominant process in the plumbing system beneath Krakatau prior to its 1883 eruption. They found limited evidence for magma mixing, and an abundance of normal zoning in phenocrysts. Normal zoning is a common feature in plagioclase within this dataset, and accounts for ~35 % of all rims, supporting an interpretation that a proportion of phenocrysts were growing under stable conditions not long before eruption.

778Our Rhyolite-MELTS modelling (Figure 13) shows that it is possible to evolve from an Anak779Krakatau composition (basaltic andesite) to the 1883 melt composition by simple fractional780crystallisation, with a starting H2O content of 1.5 wt%, and final water content of 3.4 - 3.6 wt%.781Thus, no other internal processes need to be invoked to reach the composition of the 1883782magmas from an Anak Krakatau basaltic andesite starting composition. Small discrepancies783between the model and natural samples may be as a result of the system not being entirely784isobaric.

Eruptions of silica-rich magma bodies at < 300 MPa can be internally triggered by volatile saturation resulting from fractional crystallisation, without efficient degassing (Tramontano et al., 2017). The modelled evolution of H<sub>2</sub>O using Rhyolite MELTS (Gualda et al., 2014; Figure 13f) suggests that it is likely that the pre-1883 reservoir did not reach water saturation prior to eruption, because the water content does not plateau under the best-fit pressure and temperature conditions. Using VolatileCalc (Newman and Lowenstern, 2002) we model H<sub>2</sub>O 791 saturation at a shallower crustal pressure of ~95 MPa (compared with best-estimates of 125-792 250 MPa for the 1883 reservoir), for a rhyodacite magma with 3.6 wt%  $H_2O$  at a temperature 793 of 914 °C. However, over the pressure range of 125 – 250 MPa, an estimated, additional 175 794 -925 ppm of CO<sub>2</sub> would be sufficient for the system to reach volatile saturation. These values 795 are certainly within the range expected for arc magmas (e.g., Blundy et al., 2010), lending 796 credence to the idea that the shallow magma system was volatile saturated prior to eruption, 797 and that fractional crystallisation is likely to have primed the magmatic system for eruption, 798 and is a potential trigger, if degassing was inefficient.

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#### 5.3 Role of pre-eruptive magma mixing

Recharge of magma reservoirs and magma mixing are often cited to be eruptive triggers (Sparks et al., 1977). Mandeville et al., (1996a) rule out magma mixing as a trigger for the 1883 eruption, based on a lack of reverse zoning in phenocrysts, and a low abundance of mixed pumices. However, ~60% of plagioclase phenocrysts in this study show reverse zoning at the rim, a result likely obtained by investigating crystal zoning transects at a spatial resolution of unicrons, rather than just core and rim point analyses. It is important therefore to discuss the role that magma mixing had prior to the 1883 eruption of Krakatau.

808 As discussed in section 5.1, it is likely that the compositional variations in plagioclase crystals 809 (Figures 10 and 11) represent different P, T and/or  $H_2O$  conditions, rather than differing melt 810 composition, because the matrix glass and pyroxene phenocryst chemistry are broadly 811 consistent. Therefore, the high proportion of reverse zoning at plagioclase rims is likely to 812 indicate changes in magmatic conditions, rather than mixing of compositionally distinct magmas. The volume of visually mingled pumice is also small (Mandeville et al., 1996a), and 813 814 not observed in Units A – C of the stratigraphy. Furthermore, the visually mingled pumices do 815 not show compositional differences between the dark and light matrix glasses (Self 1992).

Mafic enclaves are also entirely absent within the stratigraphy. Stehn (1929) reports some mafic ash collected during the precursory eruptive phase, however evidence of this ash was not observed on either field campaign, suggesting it may not have been preserved, possibly because it is volumetrically minor component similar to the andesite glass (~1 %) reported by Mandeville et al., (1996a).

821 However, syn-eruptive mixing of chemically similar magmas (rather than pre- or syn-eruptive 822 mixing of chemically distinct magmas) crystalising under different magmatic conditions, could 823 account for the wide range in plagioclase rim compositions (Figure 10 and 11), as they would not have had time to equilibrate with the new host melt. The matrix glass compositions also 824 825 support this, as they are broadly consistent, but become slightly more homogenous as the 826 eruption progresses through Units B to D (Figure 12), which could result from mixing of 827 chemically similar melts syn-eruption. One hypothesis might be that different melt-rich 828 regions within the shallow system coalesced and mixed syn-eruptively as a result of magma 829 withdrawal and system restructuring. In this case, magma mixing would be considered a 830 consequence of magma body destabilisation during eruption, rather than an eruptive trigger 831 (e.g., Christopher et al., 2015); this has been invoked as an explanation for homogeneity in 832 pyroclasts coexisting with complex phenocryst zoning for similar crystal-poor caldera systems 833 (Cashman and Giordano, 2014).

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# 835 5.4 Eruptive progression

A schematic diagram illustrating the proposed evolution of the magmatic system and how this links with the eruptive progression is shown in Figure 14. Unit A (Figure 4a; 6a; 7) is comprised of a green ash aggregate fall deposit at the base of stratigraphy, which is distinct in its chemistry, being more evolved than the eruptive material that follows (Figure 12). Unit A is also volumetrically minor when compared with the overlying sequence. We concur with Mandeville et al. (1996b) that Unit A likely represents the sub-Plinian May phase of the 1883 eruption of Krakatau. This is supported by reported observations of a maximum of 50 cm of green ash at the coast in June 1883 (Symons et al., 1888). We also suggest that Unit A erupted from a more-evolved, shallower melt-rich region.

845 The observation that Unit A is composed of aggregates of fine ash could indicate that the May 1883 activity was phreatomagmatic (Mandeville et al., 1996b). This interpretation is 846 consistent with historical accounts made by inhabitants of the nearby island of Sebesi, who 847 848 visited the main island of the Krakatau complex on 21<sup>st</sup> of May: "the earth burst open at their feet" on the beach. European officials who travelled from Anjer the following day also 849 reported: "near the beach, the earth was belching fire and smoke" (Furneaux 1964). According 850 851 to these accounts, Krakatau was erupting close to the coast in May, making magmatic 852 interaction with seawater possible. This is corroborated by reports from the ship Prins Hendrick, which passed close to Krakatau on 12<sup>th</sup> August: "I passed the island on the north 853 854 side... the new opening of the crater... appeared to be a small hole, maybe 100 ft in diameter, 855 only a few meters above sea level" (Macleod 1884). However, Ferzenaar, a visitor to the island 856 the day before, noted only subaerial vents. Brown et al., (2010) also note that ash aggregates 857 can also form in a moist atmosphere, so interaction of the magma with seawater during 858 fragmentation is not required to account for the Unit A deposits. Therefore, there is ambiguity 859 in both the evidence from deposits and the historical accounts regarding the degree of 860 phreatomagmatic character of the 1883 May phase.

There are periods in both early June and late July 1883 which have sparse historical records (Figure 2), and it is not known whether the activity was continuous. The volcano was attracting much less attention, and therefore it is at least likely that the eruptions were smaller, if not less frequent. It is thus difficult to know whether any activity after 27<sup>th</sup> May contributed significantly to the deposition of Unit A or not. 866 As discussed in section 5.3, the wide range in plagioclase rim compositions (Figures 10 and 867 11), and subtle changes in matrix glass chemistry throughout the stratigraphy towards a more 868 homogenous melt (Figure 12), suggest that after the initiation of the eruption in May 1883 869 there was a continuing process of coalescence and syn-eruptive mixing of shallow-stored 870 melt-rich regions. The emptying of the shallow silicic reservoir from May onwards may have 871 promoted the restructuring and mixing of the shallow system. Partial collapse of the summits 872 of Perboewatan or Danan from June to August (Verbeek 1885; Symons et al. 1888; Figure 2) 873 may also have led to downward propagating stress changes in the magmatic system, aiding 874 reorganisation of the magmatic system (e.g., Tarasewicz et al., 2012).

875 Our hypothesis is that magma body destabilisation occurred over the ~ 2-3 months leading up 876 to the onset of the paroxysmal phase of the eruption. Unit B represents the onset of this 877 climactic eruptive phase, comprising interbedded PDC and fall units (Figure 4b; 5a; 7), and tree remains (Figure 6b). The deposition of Unit B may have started as early as 22<sup>nd</sup> August (Figure 878 879 2). The increase in deposit thickness from Unit A (<5 cm) to B (up to 20 m) is likely as a result 880 of increased eruptive flux, assuming no significant topographic changes. The increase in 881 eruptive flux is likely to be due to vent widening, however stress changes related to the 882 reduction in overburden on the shallow magmatic system, may have contributed (e.g., Watt, 883 2019).

The PDC deposits of Unit B appear to be thicker in the north east (Figure 8a), suggesting that this was the dominant direction of travel. Accounts from 23<sup>rd</sup> August corroborate this, with ash reported in the north east in the straits of Sunda by ships such as the Princess Wihelmina (Macleod 1884), and heavy rains of pumice in the north in Lampong Bay on 26<sup>th</sup> August. The north east direction of travel may indicate that the summit of Danan (> 400 m) to the south, acted as a topographic barrier to PDC transport. The massive PDC deposits of Unit D (Figures 5b; 5c; 7) emanate from the inferred caldera centre, with a dominant direction of travel towards the south west (Figure 8c). The change in PDC transport direction could indicate that the topographic barrier formed by Danan had been diminished by this stage, perhaps due to collapse associated with the emplacement of the lithic lag breccias comprising Unit C (Figures 4c; 5a; 5b).

The thick, structureless PDC deposits emplaced as part of Unit D may correspond to the 895 896 paroxysmal explosions in the morning of 27<sup>th</sup> August 1883. At the top of Unit D, a lithic lag 897 breccia is observed at locality D3S2/NP4 (Figure 5b), and at D2S2/U23 there are intensely fractured metre-scale glassy blocks (Figure 5c). These blocks are reminiscent of the large lava 898 899 blocks found in phase 3 of 4 of the Late Bronze Age (Minoan) eruptive sequence in Santorini 900 (Druitt and Francaviglia 1992; Sparks and Wilson 1990). By analogy, lava blocks originating 901 from the volcanic island may have been entrained by PDCs. However, matrix glass analyses of 902 the blocks, and smaller obsidian clasts, in Unit D of the 1883 stratigraphy, show that their 903 chemistry is similar to the 1883 pyroclastic sequence, but extend to more evolved 904 compositions (Figure 12). This chemistry is distinct from the range of 1883 lithic compositions 905 noted by Mandeville et al. (1996b), which are predominantly basalt and basaltic andesite. The 906 broadly consistent matrix glass chemistry of these block means it is likely that they are derived 907 from the same source. Given the broad compositional similarities, they may be directly related 908 to the magma that fed the 1883 eruption. In addition, the rounded, irregular "tear-drop" 909 shapes of these blocks (Figure 5c) suggest ductile deformation during hot emplacement, 910 making it likely that they erupted concurrently with the pyroclastic material. Given that the 911 blocks are found in Unit D, at the top of the stratigraphic sequence (Figure 7), and the fact 912 that they have not undergone fragmentation, one hypothesis for their formation is that they 913 were small, stagnant, pockets of highly evolved melt in the shallow crust that were excavated 914 and incorporated with the pyroclastic material during the final stages of caldera formation.

915 This section of the stratigraphy potentially corresponds to the largest explosion, and most 916 devastating tsunami, at 10 am on 27<sup>th</sup> August 1883.

917 No textural evidence for phreatomagmatic activity was found in Unit D. This is contrary to what might be expected if the caldera collapse promoted significant magma-water 918 interaction. However, there are records of ash falling as "rounded accretions" in Java after the 919 main paroxysm (equated to Unit D; Figure 2), which may provide evidence for some magma 920 921 interacting with water during or after caldera collapse. However, the lack of fine-grained, 922 highly fragmented ash in Unit D is quite striking, particularly when compared with a "wet" ignimbrite such as Oruanui (Allan et al., 2012), making it unlikely that water entered the vent 923 924 in significant amounts.

925 The deposits from the 1883 Krakatau eruption are consistent with those observed in many 926 caldera-forming eruptions, e.g., Bishop Tuff (Hildreth and Wilson, 2007), Crater Lake (Bacon, 927 1983; Kamata et al., 1993) and Santorini (Druitt et al., 2019). These eruptions commonly start with a Plinian plume, with the single vent widening through time and PDCs contributing to an 928 increasing proportion of the erupted products. Caldera collapse then occurs when a critical 929 930 volume of magma has been removed from the plumbing system beneath. The tapping of 931 multiple melt batches has been documented for many crystal-poor, caldera-forming 932 eruptions, particularly in systems undergoing active extension, similar to Krakatau, e.g., the Snake River Plain (Ellis et al., 2010; Ellis and Wolff, 2012) and the Taupo Volcanic Zone (Brown 933 934 et al., 1998; Charlier et al., 2003; Gravley et al., 2007; Wilson and Charlier, 2009; Bégué et al., 935 2014). In these cases, melt was stored in laterally (rather than vertically) extensive systems, 936 with a consistent bulk chemistry between melt lenses (Cashman and Giordano 2014). Many 937 caldera-forming eruptions are preceded by some form of precursory eruption, however the 938 period of time between this and the climactic eruption is often poorly constrained (e.g., Allan 939 et al., 2012; Cashman and Giordano 2014; Druitt et al., 2019). The 1883 eruption of Krakatau

940 is unique in the sense that we know when this precursory activity began (Figure 2; 20<sup>th</sup> May 1883) from direct observations. This may be invaluable for monitoring volcanoes with a 941 history of producing explosive, caldera-forming eruptions, because it highlights the potential 942 for a large event to follow a relatively moderate explosive eruption, on a timescale of months. 943 944 However, the identification of such events as precursory to an incipient larger eruption 945 remains challenging. Although a precursory Plinian eruption might increase the risk of a larger eruption, one does not always follow on from the other. Top-down factors, such as the 946 947 removal of mass from the volcano edifice (discussed in section 5.4), potentially have a role in triggering these devastating paroxysms. Therefore, it would also be useful to carefully monitor 948 949 surface deformation and any significant losses of mass at Anak Krakatau, as well as other similar volcanic systems. 950



Figure 14: Series of schematic diagrams coupled with plan view maps illustrating one possible
model for the magmatic and eruptive evolution of the 1883 eruption of Krakatau (lithological
structure and crystallisation depths from Darhen et al., 2012). On maps orange stars indicate

955 an active edifice, and black arrows represent the dominant direction of PDCs. Dashed lines on 956 schematic cross sections indicate ephemeral transport pathways of magma. Red arrows 957 indicate melt extraction. Colours of melt represent melt composition: Purple for the least 958 evolved, followed by brown, bright red, with pink representing the most silicic melt. Panel (a) 959 shows the deposition of Unit A (Figure 7), with evidence for a more silicic reservoir derived 960 from matrix glass analyses (Figure 12), and the observed eruption dynamics (Figure 2). Panel (b) shows observed changes to the eruption dynamics (Figure 2), and proposed system 961 962 restructuring. Panel (c) shows the deposition of Unit B (Figure 7), and the start of the 963 paroxysmal phases of the eruption, as determined by observations (Figure 2), as well as syn-964 eruptive mixing of chemically similar melt batches hypothesised as a result of varied and 965 complex plagioclase textures and chemistry (Figure 10; 11), and only minor variations in 966 matrix glass chemistry (Figure 12). Panel (d) shows the deposition of Unit C and Unit D in a 967 piecemeal caldera collapse, as there are two lithic lag breccias (Figure 7), and a substantial change in PDC and dilute-PDC directionality (Figure 8). 968

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# 970 6 Conclusions

971 This study presents field observations from new exposures of the 1883 eruptive deposits of Krakatau, revealed as a result of the removal of vegetation by the 2018 tsunami generated by 972 973 flank collapse of Anak Krakatau. This has allowed for the stratigraphy of the eruption to be 974 considerably better constrained (Figure 3; Figure 7). Examination of the geochemistry in the 975 context of this refined eruptive stratigraphy does not support previous studies (e.g., Mandeville 976 et al., 1996a) that have proposed that the eruption emanated from a chemically zoned magma 977 reservoir. An updated model for the magmatic system is proposed, accounting for the chemical 978 variations in context with the stratigraphic sequence.

979 The existence of a distinct, green, basal ashfall (Figure 4a) is consistent with written accounts of 980 precursory activity in May (e.g., Symons et al., 1888; Figure 2), with the matrix glass chemistry 981 indicating a shallow, more silicic melt-rich region was tapped during this preliminary eruptive phase (Figure 12). It is likely that restructuring of the magmatic system and syn-eruptive mixing of 982 983 multiple chemically similar melt batches then occurred, to account for the chemical homogeneity 984 in pyroclasts in Units B to D (Figure 12), and complex plagioclase phenocryst zoning profiles and 985 textures (Figures 10 and 11). This restructuring may have occurred simply as a result of gradual 986 emptying of the initial silicic reservoir, however the loss of mass reported from the summits of 987 Perboewatan and/or Danan (Figure 1) may also have played a role. The stress changes and 988 reservoir reconstruction may have contributed to the onset of the climactic phase of the eruption on 26<sup>th</sup> August. 989

990 There is a substantial change in the directionality of the PDCs throughout the climactic phase of 991 the eruption (Figure 8), which coincides with the deposition of a lithic lag breccia occupying a distinct horizon within the stratigraphy (Figure 4c; 5a; 5b). The lithic lag breccia is thus attributed 992 993 to partial collapse of the island, and the removal of the edifice Danan as a topographic barrier. 994 The 1883 eruption culminated in total caldera collapse, which, together with the PDC production 995 at this stage, was a potential cause of the largest tsunami at 10 am on 27<sup>th</sup> August. This final 996 caldera collapse is recorded in the stratigraphy as a second lithic lag breccia (Figure 5b), and at 997 one locality (U23/D2S2) glassy blocks up to 8 m in size are present (Figure 5c), which are reported 998 within this sequence for the first time here. The matrix glass chemistry of these blocks (Figure 12) 999 suggests that they are derived from the same melt as the 1883 pyroclastic material, and their 1000 shapes suggest that they underwent ductile deformation during hot emplacement.

1001 The identification of at least two lag breccias may indicate piecemeal caldera formation, where 1002 the first stage of collapse is the driving force behind the most energetic and explosive, climactic 1003 part of the eruption. Precursory Plinian eruptions are therefore very useful phenomena to be aware of for the future monitoring of volcanoes with a history of producing silicic caldera-forming
 eruptions. The 1883 eruption of Krakatau provides an example of an event where relatively
 moderate explosive eruptions may potentially have run-away effects culminating in cataclysmic
 caldera-collapse several months later.

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# 1023 8 Data availability

1024Datasetsrelatedtothisarticlecanbefoundat1025<a href="https://www.bgs.ac.uk/services/ngdc/accessions/index.html#item137445">https://www.bgs.ac.uk/services/ngdc/accessions/index.html#item137445</a>, hosted at National1026Geological Data Centre by the British Geological Survey (Madden-Nadeau, 2020).

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