



 **Opin vísindi**

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# Deep-sea eruptions boosted by induced fuel-coolant explosions

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**The majority of Earth's volcanic eruptions occur beneath the sea, but few direct observations and samples limit our understanding of these unseen events. Subaerial eruptions lend some insights, but direct extrapolation from subaerial to deep-sea is precluded by the great differences in pressure, thermal conditions, density, rheology, and the interplay among them. Here we present laboratory fragmentation experiments that mimic deep-sea explosive eruptions and compare our laboratory observations with those from the kilometre-deep submarine eruption of Havre volcano, Kermadec arc, New Zealand in 2012. We find that the Havre eruption involved explosive fragmentation of magma by a pressure-insensitive interaction between cool water and**

23 **hot magma, termed induced fuel-coolant interaction. The laboratory experiments show**  
24 **that this water-magma interaction is initiated by the formation of cracks in cooling**  
25 **magma into which the water coolant can infiltrate, driving explosive fragmentation.**  
26 **Explosive submarine eruptions have previously been considered unlikely because the**  
27 **stabilisation of a vapour film at the magma-water contact was thought to be a key**  
28 **requirement and is suppressed at depths exceeding a hundred metres. However, here we**  
29 **demonstrate that these induced fuel-coolant interactions between magma and water can**  
30 **occur in a range of wet environments regardless of pressure, from subaerial to the deep**  
31 **sea, and may operate on different planets, as well as apply to materials other than**  
32 **magma and water.**

33 The 2012 eruption of Havre volcano, Kermadec arc, New Zealand, produced in a day a  
34 ~400 km<sup>2</sup> pumice raft<sup>1-3</sup>, and on the seafloor an abundance of fine ash<sup>4</sup>. Over 35 km<sup>2</sup> on the  
35 volcano the ash, most widely distributed from the main eruptive phase, shows no thinning  
36 trend, so the total volume of erupted ash must substantially exceed the 100 million m<sup>3</sup> in the  
37 mapped area<sup>4</sup>. We focus on ash (S1, S2) from this main phase<sup>4</sup>; it signals expenditure of a  
38 large amount of energy to fragment magma<sup>5</sup>, which is surprising at this depth<sup>6-8</sup>. The ash,  
39 erupted from 900-1100 m below sea level (bsl), has abundant blocky or curvi-planar shaped  
40 particles <125 μm with low vesicularity and stepped fracture surfaces. Abundant fine to very  
41 fine ash suggests explosive fragmentation<sup>9</sup>, but particle shapes are not those of pumiceous  
42 ash produced by expansion of magmatic gases. Instead we find fingerprints like those from  
43 fuel-coolant (thermohydraulic) ash-forming explosions. Havre is too deep, and rhyolite too  
44 viscous, for 'normal' fuel-coolant interactions<sup>6,7,10-18</sup>, and we infer fragmentation by "Induced  
45 Fuel Coolant Interaction" (IFCI). Experimental evidence is presented for its role at Havre.  
46 We suggest that IFCI aids ash production and increases explosivity during most submarine

47 eruptions, and for deep-water volcanic eruptions it is likely to be the dominant ash-forming  
48 process.

49

### 50 **Kindred explosive processes - MFCI and IFCI**

51 In volcanology highly explosive energy release from interaction of magma with water  
52 (“Molten-Fuel Coolant Interaction” (MFCI)) is known as the driving mechanism for  
53 subaerial, mostly basaltic, phreatomagmatic eruptions<sup>10,11,19–22</sup>. The key MFCI process is a  
54 non-equilibrium thermohydraulic feedback cycle (Fig. 1). Heat transferred from hot fuel into  
55 entrapped, expandable, coolant induces strong hydraulic pressure on the fuel-coolant  
56 interface, creating brittle fractures that penetrate the fuel. Expanding liquid coolant pushes  
57 into these cracks, driving within fractions of milliseconds their further propagation which  
58 increases the interface area, accelerates heat transfer, and releases strong shock-waves. The  
59 feedback persists until the system is opened and the superheated entrapped coolant is released  
60 as expanding vapour, carrying away with high kinetic energy the crack-bounded fragments.  
61 In a study on Tepexitl<sup>23</sup>, a desert volcano, it was experimentally verified that explosive fuel-  
62 coolant interaction processes are possible even for a highly viscous magma, and without  
63 water entrapment. Here we distinguish two conceptual models (Fig.1) for fuel-coolant  
64 interaction and introduce the term “Induced Fuel-Coolant Interaction” (IFCI) as separate  
65 from the well-explored ‘regular’ MFCI mechanism. The differences between these two are  
66 particularly relevant for submarine eruptive settings below the depth of stable film boiling  
67 (>1 MPa; deeper than 100 m bsl<sup>8,16</sup>), and we show that IFCI produced most Havre 2012 ash  
68 using experiments, particle morphology, and thermodynamic analysis. This changes our view  
69 of deep-submarine eruptions, for which it is a deeply embedded concept that magma-water  
70 explosivity is unimportant.

71 Induced fuel coolant interactions function through "shortcutting" initiation of the  
72 thermohydraulic feedback loop, which is the underappreciated core of MFCI explosions.  
73 IFCI occurs when coolant enters cracks that open in a fuel being fragmented by other  
74 processes. While resembling the MFCI process in how heat is thermohydraulically converted  
75 in a feedback loop (see Fig. 1) to release fragmentation energy, IFCI does not require vapour  
76 films and occurs under less restrictive initial and boundary conditions. Based on our analysis  
77 of initial threshold conditions, experiments and heat transfer simulations we show that IFCI,  
78 here first diagrammed and established as a separate fuel-coolant process, can convert heat to  
79 produce ash in submarine eruptions at any depth.

80 We investigated IFCI characteristics and explored its role in the 2012 Havre eruption, by  
81 conducting two series of fragmentation experiments with crushed and remelted Havre rock  
82 together with statistical ash sampling strategies (Methods). In "dry" runs, melt was deformed  
83 and fragmented by injecting pressurized gas, whereas in "IFCI" runs a layer of water was  
84 added prior to gas injection (see Fig. 2). In IFCI runs, fragments were produced by (1) dry  
85 gas-driven cracking, and (2) thermohydraulic processes during IFCI ("IFCI particles").

86 Thermohydraulic IFCI processes advanced downward in our setup, tracing the opening  
87 tensional cracks from the top of the melt (Methods). IFCI particles were thus much more  
88 abundant in the leading part of the ejected cloud of fragments than in the following ejecta. In  
89 both dry and IFCI runs ejecta was collected from the ground ("DG" and "IG" for dry and  
90 IFCI runs, respectively). IFCI products were also captured in a water bowl positioned  
91 alongside the open conduit ("IB"), and as deposits inside water droplets ("IW") adhering to  
92 the walls and ceiling around the experimental area (see Extended Data Fig. 1). Furthermore,  
93 in a novel subseries of IFCI experiments one end of a U-shaped tube was mounted above the  
94 crucible with its other end leading to a water bowl. In these "U-tube" runs, the tube detached

95 ~30 ms after particle ejection began (Methods), dynamically separating the leading front of  
96 fine particles (“IU”, Fig. 2).

97 We compared the shape of the ash-sized experimental products with natural ash retrieved at  
98 six locations close to Havre volcano, labelled “Nat1”-“Nat6” (Extended Data Fig. 1).

99

### 100 **IFCI vs dry-fragmentation experiments**

101 The effect of IFCI on fragmentation is evident if maximum recoil force  $F_{max}$  (i.e., the  
102 repulsion force exerted on the crucible) is normalized relative to maximum pressure  $P_{max}$  and  
103 plotted over itself (Fig. 3). Dry runs plot below the dashed line; most IFCI runs plot above.

104 For dry runs  $F_{max}$  did not exceed ~2.9 kN (for many, <1 kN), whereas IFCI run peak values  
105 reached ~5.8 kN (Extended Data Fig. 2). In dry runs  $F_{max}$  correlates with  $P_{max}$  because driving

106 pressure is their only energy source for deformation and stress-induced fragmentation. In

107 contrast, most of the IFCI runs reveal the anticipated thermohydraulic “boost”, which added  
108 significantly more energy (and thus also entropy) into the process of fragmentation,

109 producing also a considerably larger deviation of data points from the diagonal in the upper

110 IFCI field compared to those in the dry regime (Fig. 3a). We infer that the three outlier IFCI

111 runs (IFCI01, IFCI03 and IFCI09), which plot among the dry runs, did not experience

112 significant interaction of magma with coolant, despite having equivalent initial conditions.

113 This suggests that IFCI energy yield is sensitive to subtle dynamic controls at laboratory

114 (decimeter) scale. We tested different melt masses (thickness of melt plug), and they had no

115 significant influence on explosivity, nor on the likelihood of achieving efficient IFCI runs.

116 The increase in fragmentation efficiency with IFCI is also reflected in particle grainsizes.

117 Sieve data for particles  $\leq 2$  mm (“ash”) from seven dry runs and five IFCI runs show

118 increased IFCI fragmentation. On average IFCI runs generated 2.25 times the mass of ash

119 particles compared to dry runs (Extended Data Fig. 3). Particularly interesting is the increase

120 in fine ash (<125  $\mu\text{m}$ ) – this matches the bulk of seafloor ash discovered at Havre<sup>4</sup>, and is not  
121 a particle population that is predicted as significant for submarine volcanism<sup>7</sup>. The proportion  
122 of fine ash with IFCI is ~240% of that produced in dry runs (14.1% vs 5.8%, see Fig. 3c).

123

#### 124 **Identifying IFCI's morphometric fingerprint at Havre**

125 Curvi-planar Havre ash grains do not share the morphology of ground-sampled particles from  
126 dry runs (DG) or open IFCI runs (without U-tube, IG) (Fig. 4). We infer that ground-sampled  
127 particles from IFCI runs are a mixture of dry-formed and thermohydraulically fragmented  
128 grains, with population differences in shape that place them apart from the other samples.

129 U-tube and wall-sampled particles from IFCI runs show, unlike ground-sampled ones, clear  
130 similarities with natural ash samples (Nat1 - Nat6), verified by t-tests and e(quivalence)-tests  
131 for all 23 shape parameters. SEM imagery and high-resolution micro X-ray tomography show  
132 that these grains share surface features such as steps (Fig. 4). IU contain predominantly grains  
133 from the leading ejecta front generated by IFCI, and IW samples show no significant  
134 differences to IU ones in any tested shape parameter. Twelve parameters are verified as  
135 significantly similar according to e-tests (other shape parameters had large variance  
136 differences that precluded meaningful e-tests), indicating high similarity. This suggests that  
137 IW samples originated, like IU samples, from the leading ejecta front, and that in our  
138 experiments IFCI processes generated particles with a characteristic morphometric  
139 “fingerprint”. This fingerprint is shared by Havre's curvi-planar natural grains, which are  
140 dominant in Havre's seafloor deposits of fine to extremely fine, 3-8 phi, ash<sup>4</sup>. We therefore  
141 infer that IFCI played the major role in generating Havre's ash.

142

#### 143 **Deep-sea conditions favour IFCI**

144 The ratio of thermohydraulic to dry-generated grains in open IFCI experiments is measurably  
145 lower than in the main Havre ash deposits, in which curvi-planar particles are dominant<sup>4</sup>.  
146 IFCI processes at Havre were thus apparently more efficient than in the lab.  
147 One of the biggest differences between lab conditions vs. Havre's ones is Havre's much  
148 higher ambient pressure (0.1 vs ~10 MPa). A mechanical consequence of higher confining  
149 pressure is stronger driving of water into opening cracks, and this water ingress, as well as an  
150 initial magmatic expansion that opens them, are necessary to initiate IFCI. Two  
151 thermodynamic factors most strongly control the efficiency of IFCI: heat transfer rates from  
152 melt to water (controlling energy available for thermohydraulic conversion) and expansion  
153 rates of water that acts as a “wedge” inside the crack (controlling how effectively this energy  
154 is converted into mechanical work).

155

156 Our laboratory heat-transfer rates were mediated by vapour almost instantaneously generated  
157 when water contacted melt (Leidenfrost effect<sup>21,24</sup>). Unlike for MFCI<sup>18,21,25</sup>, no stable vapour  
158 film is required for IFCI; instead, vapour films inhibit IFCI because they prevent efficient  
159 heat transfer from melt to water. Shock waves are generated the moment pressurized gas hits  
160 the ceramic barrier and overlying melt, and cause the vapour film to collapse.<sup>21,23,26</sup> We  
161 cannot be sure, however, that there is no re-formation, even locally, of a vapour film by the  
162 time of crack opening.

163 Under lab conditions 30% of IFCI runs with water flooding did not produce recognizable  
164 IFCI fragmentation according to Fig. 3a. We infer that these intended IFCI runs “failed”  
165 because the required pre-condition – synchronous crack opening and water invasion (Fig. 1) –  
166 was not met, possibly because local film boiling inhibited interaction. This finding suggests  
167 that IFCI requires a critical density of simultaneous cracking with direct water-melt contact.

168 With increasing water depth and pressure, vapour ceases to impede IFCI, and this happens at  
169 pressures much less than critical for seawater (~30 MPa)<sup>16</sup>. The key process, stable film  
170 boiling, is strongly suppressed from 1 MPa, and at 10 MPa (~1 km bsl) meta-stable film  
171 boiling becomes impossible<sup>16</sup>. Consequently, IFCI is *favoured* in a deep-sea environment,  
172 compared to shallower settings with lower ambient pressure.

173 With no vapour film, water directly contacts melt, and heat transfer rates are controlled by  
174 conduction and convection. Experiments have found that the efficiency of heat transfer  
175 between hot rock and water flowing into cracks is increased under increased pressures.<sup>27</sup>

176

177 Higher pressures also enhance the rate of water expansion. Fig.5 compares the evolution of  
178 the thermal expansion coefficient  $\beta$  with temperature, computed for water under lab and  
179 seafloor pressure conditions. In addition, values for sea water at 1 km depth are plotted, based  
180 on measurements under oceanic temperatures<sup>28</sup>. At low water temperatures,  $\beta$  is slightly  
181 larger for pure water under high pressure conditions, (e.g.,  $1.8^{\circ}\text{C}^{-1}$  vs  $1.6^{\circ}\text{C}^{-1}$  at  $30^{\circ}\text{C}$ ). This  
182 difference is more pronounced for the measurements of seawater at high pressure (e.g.,  $2.4^{\circ}\text{C}^{-1}$   
183 at  $30^{\circ}\text{C}$  which is 48% larger than  $\beta$  for pure water at 0.1 MPa). The dominant controlling  
184 factor on  $\beta$  is, however, the water temperature (see also Extended Data Fig. 4). At a water  
185 temperature of  $100^{\circ}\text{C}$ ,  $\beta$  is  $4.5^{\circ}\text{C}^{-1}$ . At  $310^{\circ}\text{C}$ ,  $\beta$  is more than three times as large ( $14.8^{\circ}\text{C}^{-1}$ ),  
186 so the same amount of transferred heat would therefore result in significantly larger  
187 expansion rates, and considerably enhance the efficiency of IFCI. In contrast to water at  
188 atmospheric pressure (boiling point  $\sim 100^{\circ}\text{C}$ ), water at 10 MPa vaporizes not under  $\sim 311^{\circ}\text{C}$   
189 and can therefore exploit the full range of IFCI intensification. Thermodynamically, IFCI  
190 processes should thus be more efficient under deep submarine conditions than in the lab.

191

192 **“Explosive” ash generation under unexpected conditions**

193 IFCI is an unusual “explosive” process. This thermohydraulic non-equilibrium mechanism  
194 generates fine ash at extreme rates by converting heat into intense mechanical work. It is  
195 based on an inherently accelerating feedback mechanism fed by increases in both contact  
196 surface and heat-transfer rates, leading to rapid and accelerating expansion at microsecond  
197 timescales, an “explosive” rate. However, unlike MFCI, at deep seafloor conditions (or  
198 similar ones beneath thick glaciers) no dramatic steam expansion would follow the hydraulic  
199 stage. Although considerable kinetic energy release can be expected (based on the observed  
200 surplus of  $F_{max}$  in the IFCI experiments), the higher mass density and viscosity of water  
201 compared to air in subaerial volcanic settings would cause significantly reduced ejection  
202 speeds for ash particles. Instead, newly generated fine ash particles would be primarily  
203 transported by convective movements of the heated ocean water. Their small settling  
204 velocities enable widespread dispersion, but explosive expansion has been effectively  
205 suppressed.

206  
207 A study of Havre pumice concluded that the pumice-raft-generating phase was not driven by  
208 magmatic explosions<sup>3</sup>. However, a ~70 km-long subaerial vapour plume was observed above  
209 the eruptive centre of Havre<sup>1</sup>, which is evidence for high rates of heat and energy transfer  
210 during one observed stage of the Havre eruption. Furthermore, a concomitant bluish semi-  
211 opaque plume at the ocean surface, offset from the pumice raft<sup>1</sup>, indicates that fine particles  
212 (ash) were scattering light down-current from the thermal source of the vapour plume. Fine  
213 ash of the main event comprises mostly curvi-planar particles<sup>4</sup>, which morphologically match  
214 our experimentally generated IFCI particles. We infer that at some point(s) during the main  
215 phase of the eruption, ash production and heat-transfer rates at Havre were increased by IFCI  
216 processes, and that IFCI was the mechanism which fuelled ash transport towards the surface  
217 in a strengthened convective plume.

218 The presence of fluidal ash particles in the same deposits implies that in Havre's main phase  
219 inhomogeneous magma was erupted and fragmented by multiple processes<sup>4</sup>. This is  
220 consistent with our findings that suggest that the thermohydraulic boost of IFCI was  
221 superimposed on rapid and extensive, magmatically driven, fragmentation, which might have  
222 either occurred simultaneously or in alternation with other processes during the pumice-raft-  
223 generating phase.

224

225 We conclude that IFCI can operate in all deep-sea eruptive settings. Because its major  
226 requirement is initial magmatic cracking, we suggest it can operate with all magmatic  
227 compositions. The primary effects of IFCI at depth are intensified fragmentation and heat  
228 transfer, but without necessary vapour-driven particle transport. Deep-sea eruptions in which  
229 it is a major process are far more energetic than classic (subaerial) "effusive" eruptions,  
230 resulting in an increased production of fine ash, yet may lack the jets and plumes driven by  
231 gas expansion typifying "explosive" ones. Any process cracking a hot fuel into which a  
232 coolant can be driven invites IFCI – its fingerprint can be detected through morphometric  
233 analysis of the resulting small particles. Beyond submarine volcanoes, IFCI is insensitive to  
234 many controls thought to limit explosive magma-water interaction, and it may be expected  
235 with hot dry rocks or other hot brittle materials in the deep ocean, as well as with magmatism  
236 beneath thick ice on glacier-bearing planets.

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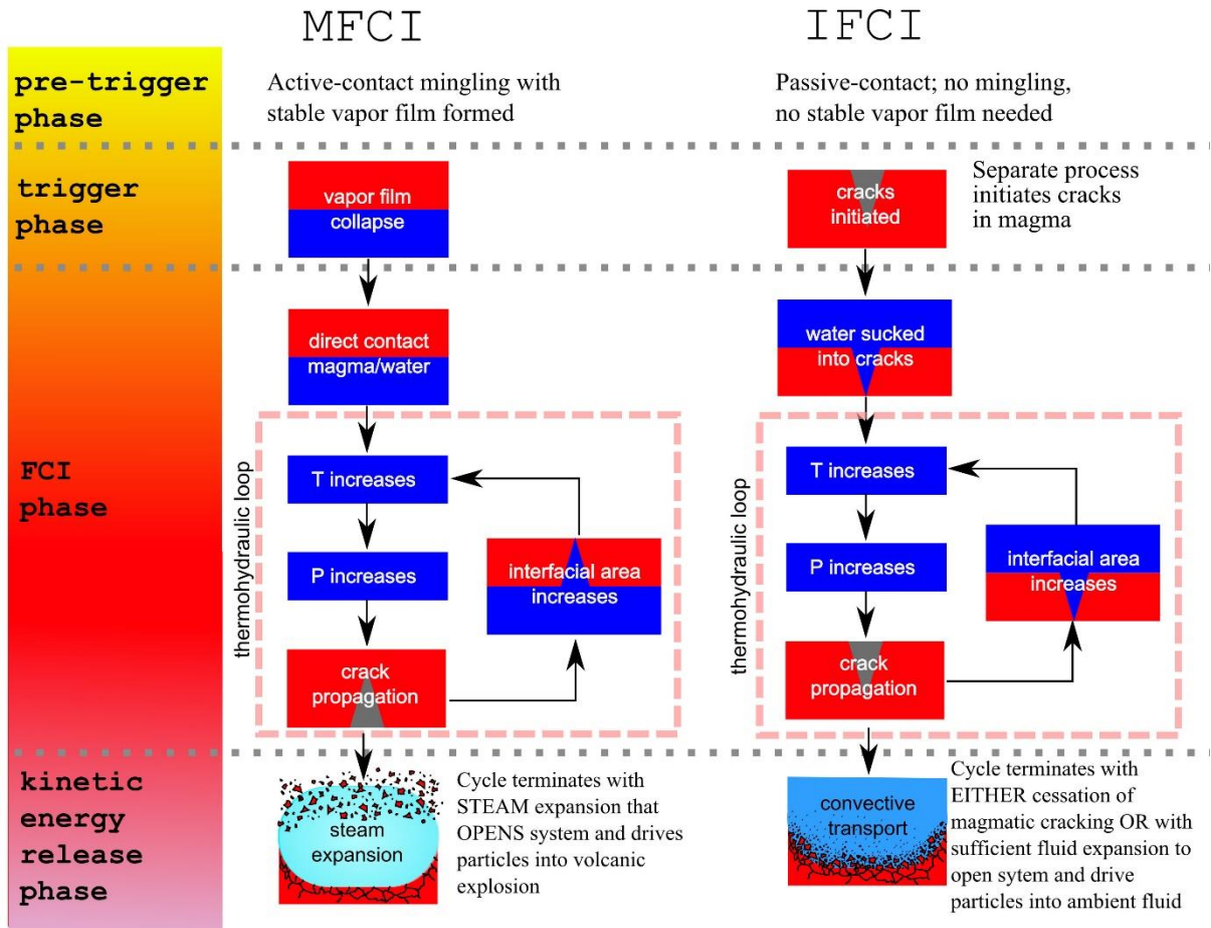
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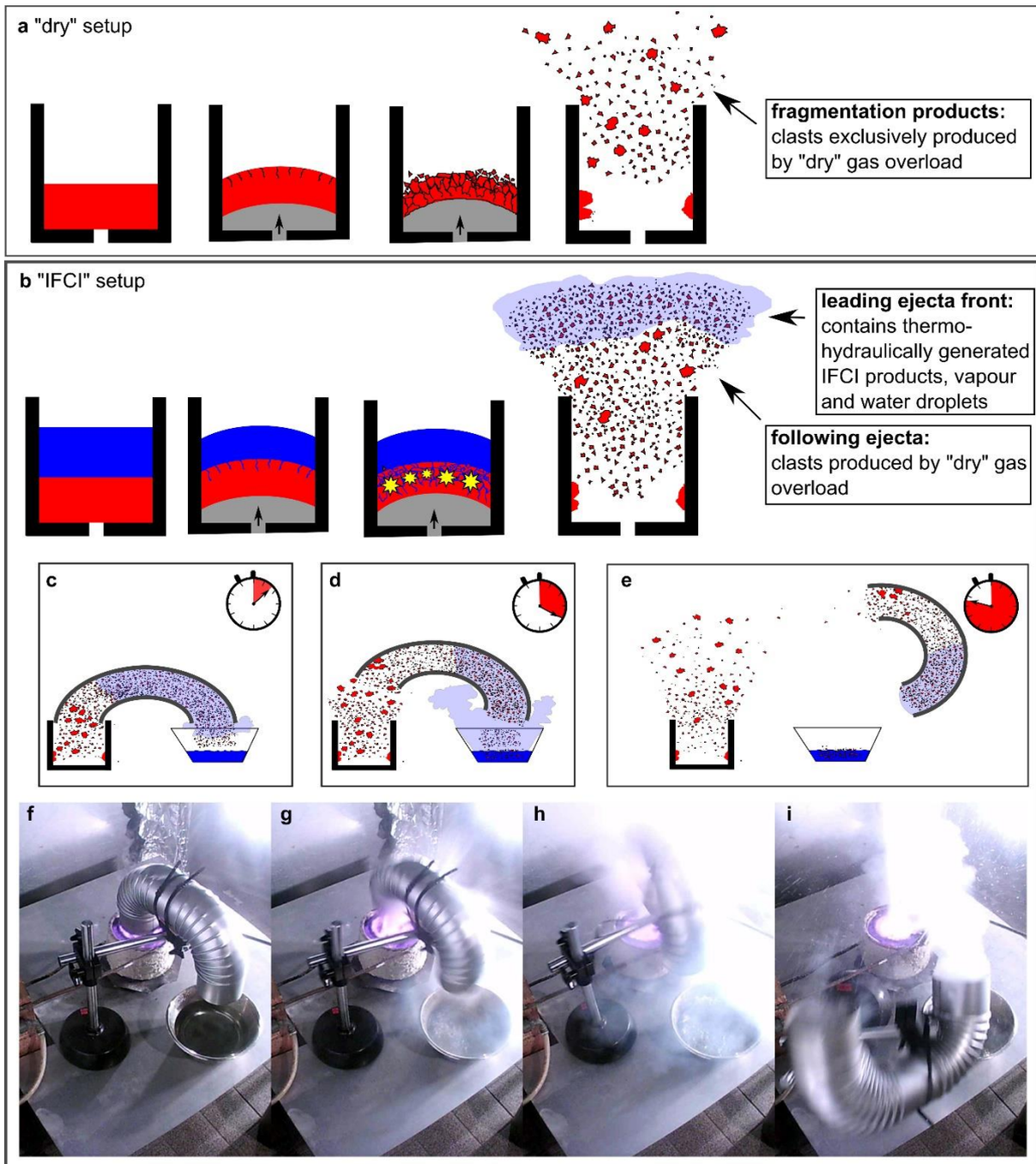
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323 Figure 1



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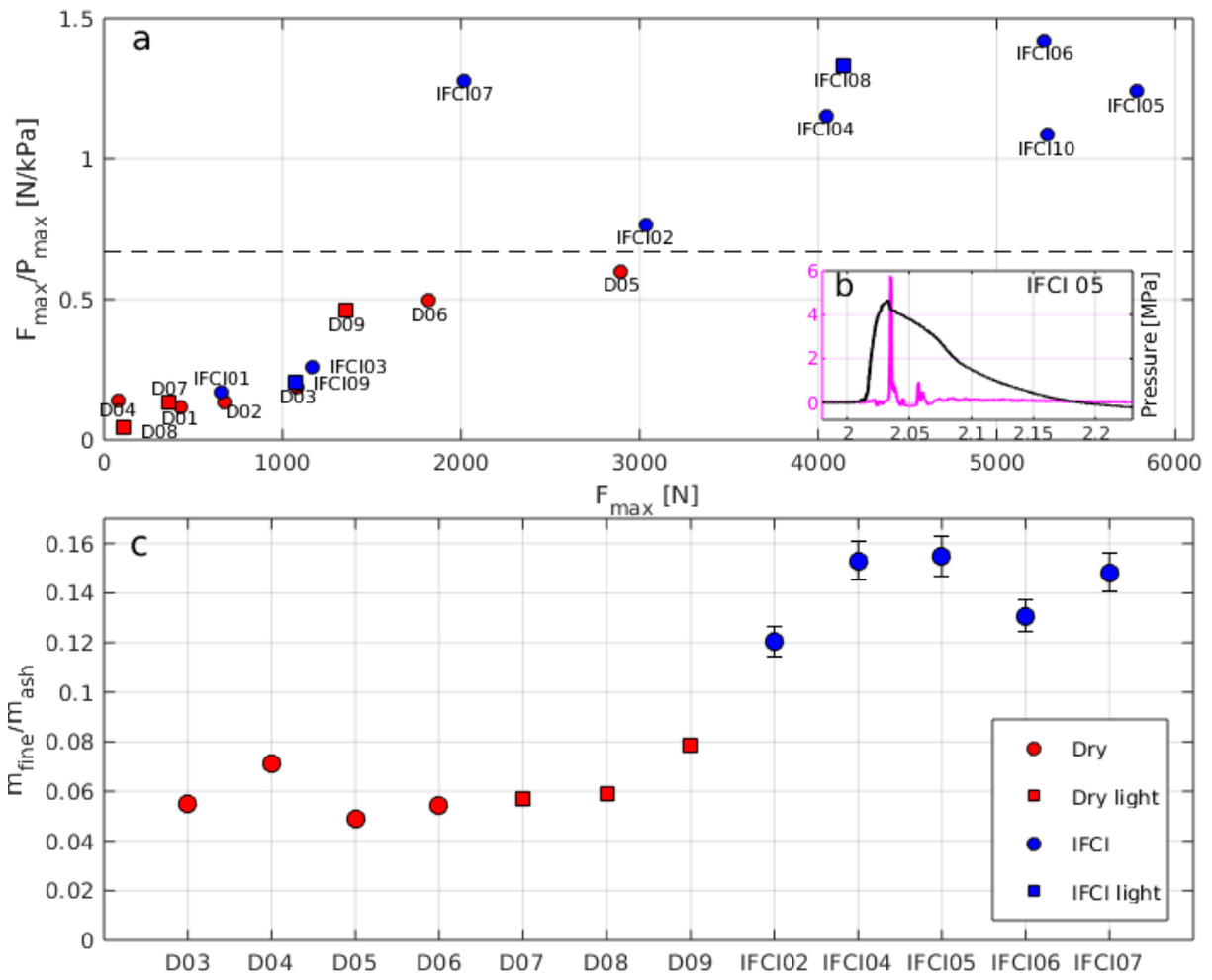
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329 Figure 3



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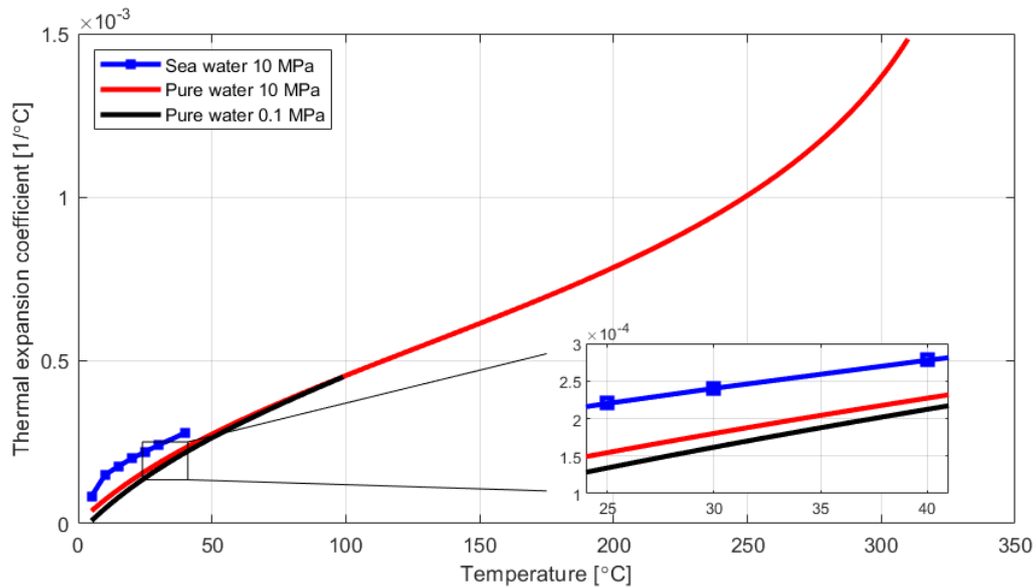
332 Figure 4

	Nat	IU	IW	IG	DG
	curvi-planar Havre ash	IFCI (U-tube)	IFCI (wall)	IFCI (ground)	dry (ground)
SEM					
μCT					
					50μm
Nat		✓	✓	✗	✗
IU	✓		✓	✗	✗
IW	✓	✓		✗	✗
IG	✗	✗	✗		✗
DG	✗	✗	✗	✗	
legend:	✓ similarity verified		✓ similarity likely		✗ significant differences

333

334

335 Figure 5



336

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345

346 **Author contributions:**

347 J.D.L.W. designed and supervised the Marsden research project. T.D., J.D.L.W., B.Z., R.B.,  
348 and A.P.M. conducted the lab experiments at Physikalisch Vulkanologisches Labor,  
349 University of Würzburg. T.D. and N.S. sampled the experimental particles. T.D. conducted  
350 2D morphometry and statistical analysis of both experimental and natural ash with support  
351 from D.M. and P.D.. D.M., P.D. and T.D. conducted X-ray microtomography and image

352 reconstruction. L.S.S. computed thermal expansion coefficients. R.J.C was leader of the NSF  
353 cruise and helped formulate the Marsden project. J.D.L.W., R.J.C. and A.P.M. took part at  
354 the NSF cruise and collected the dome rock used as starting material for melt fragmentation  
355 experiments. R.J.C. and A.P.M. provided Havre ash samples and related meta data, including  
356 the bathymetric map. Figures and plots were produced by T.D. and L.S.S.. T.D. and J.D.L.W.  
357 wrote the manuscript, with contributions from all the authors.

358

### 359 **Financial and non-financial competing interests**

360 The authors declare no competing interests.

361

### 362 **Figure legends**

#### 363 **Figure 1| Comparison of molten (MFCI) with induced fuel-coolant interaction (IFCI).**

364 Magma (red) is volcanic fuel, and water (blue) the coolant. MFCI needs a pre-trigger vapor  
365 film, limiting it to shallow water depths. IFCI initiation requires a sudden increase of the  
366 fuel's surface by cracking, *e.g.* by brittle failure of magma highly strained during eruption.

367

#### 368 **Figure 2| Melt fragmentation experiments. a**, dry runs: the melt plug (red) was deformed

369 by injecting gas, causing stress-induced brittle fragmentation. **b**, IFCI runs: seconds before

370 gas injection, water was added atop the plug, subsequently intruding cracks opened when gas

371 deformed the plug, initiating downward-advancing IFCI that thermohydraulically “boosted”

372 fragmentation. IFCI particles are most abundant in the leading ejecta front. **c**, "U-tube"

373 experiments: the fine particles from the leading ejecta front were collected in a water bowl

374 via a U-shaped tube. **d**, following coarser ejecta particles enter the tube shortly after. Impact

375 momentum causes the tube to detach and **e**, move away from the crucible, with very few

376 particles deposited into the bowl. Bottom-row images: **f**, before the run, and at **g**, 33 ms, **h**,

377 67 ms and **i**, 167 ms after initiation of fragmentation (see also Supplementary Video 1).

378

379 **Figure 3| Increased explosivity in IFCI fragmentation experiments. a,** recoil-force peaks  
380  $F_{max}$  normalized over maximum driving pressure  $P_{max}$  and plotted over themselves for dry  
381 and IFCI runs. Squares indicate experiments with reduced melt mass. Dashed line indicates  
382 the empirical boundary between two regimes. Dry runs plot below the line, whereas most  
383 IFCI runs plot above it, reflecting the additional thermohydraulic fragmentation “boost”. IFCI  
384 runs of low efficiency plot together with dry runs. **b,** example showing pressure (black) and  
385 force (magenta) signals for IFCI run plotted over time (range: 250 ms). **c,** mass ratio of fine  
386 ash (<125  $\mu\text{m}$ ) over total ash mass. Error bars indicate measurement uncertainties and are  
387 displayed where larger than marker.

388

389 **Figure 4| Morphometric analysis results by t-tests and e-tests.** Typical particles imaged  
390 with SEM and X-ray microtomography (image pixel size 0.56  $\mu\text{m}$ ). Natural ash (“Nat”) is  
391 significantly similar to particles from the leading ejecta front of IFCI experiments (IU and  
392 IW), which are highly similar to one another.

393

394 **Figure 5| Thermal expansion coefficient at lab and seafloor pressures.** The volumetric  
395 thermal expansion coefficient  $\beta$  was computed for water at atmospheric pressure (0.1 MPa;  
396 black) and 10 MPa (red). In addition, measured data for saline water (within 0.1%  
397 accuracy)<sup>28</sup> are plotted for oceanic temperatures (solid blue lines). Water boils much hotter on  
398 the seafloor, allowing higher thermal expansion coefficient values that increase IFCI  
399 efficiency under deep-sea conditions.

400 **Methods**

401

402 **Fragmentation experiments**

403 For each run crushed rock of mass  $m_{melt}$  was inductively heated under non-equilibrium  
404 conditions in a 10 cm diameter cylindrical steel crucible. Standard runs used 250 g of rock;  
405 “light” experiments used reduced  $m_{melt}$  of 100 g. Within an 80 minute period, the crushed  
406 dome rock material was heated up to a temperature of 1573 K then equilibrated for 30  
407 minutes. It was then cooled over a 30-minute period to the experimental temperature of 1423  
408 K. Throughout this process the crucible was covered by a steel lid, which did not contact the  
409 melt and which was removed only seconds before the experiment. The heating procedure was  
410 found to provide homogeneous melt temperatures (within a range of 2 K) in calibration  
411 experiments in which this parameter was measured at different times, depths and locations  
412 using a type S thermocouple.

413 “Dry run” routines were based on standard experiments for determining material-specific  
414 fragmentation and kinetic energy release rates and used that standard setup<sup>26,29</sup> (see Extended  
415 Data Fig. 5).

416 In preparation for each run, a steel-enforced rubber hose 8 m long with 8 mm (inner)  
417 diameter, leading from an argon gas reservoir to a closed high-speed solenoid valve, was  
418 pressurized. At a pressure of 8.5 MPa the valve of the gas reservoir was closed. The inner  
419 volume of the hose hence represents the driving volume of the pressurized gas. A pipe  
420 connected the solenoid with the gas inlet into the crucible, which was covered by a ceramic  
421 barrier (diaphragm) placed, without bonding, to block rock fragments or melt from entering  
422 the gas inlet. With initiation of the experiment the high-speed solenoid was opened, and the  
423 pressurized argon was injected from below into the cylindrical plug of melt.

424 When rapidly deformed by the expanding injected gas, a silicate melt behaves  
425 brittlely.<sup>23,26,29,30</sup> Like a glass pane hit by a football, the plug deformed until the critical shear

426 stress was reached, at which point it broke, with elastically stored energy converted into  
427 fragmentation and kinetic energy.<sup>26,29,30</sup> In the setup geometry used, melt fragmentation is  
428 initiated by cracks opening at the top and subsequently progressing downwards.<sup>23,26</sup>  
429 The pressure of the driving gas was measured by a Kistler® 603B pressure transducer (see  
430 Extended Data Fig. 5). A Kistler® 9031A force transducer measured the vertical component  
431 of the repulsion force exerted to the crucible, hereby termed “recoil force”. Force and  
432 pressure signals were measured with a sampling rate of 100 kHz.  
433 Heating procedure, geometry and sensor setup for “IFCI runs” were the same as for dry runs,  
434 but we added a hosepipe leading to the top of the crucible. Two seconds before the gas valve  
435 opened, 240 ml of deionized water flooded the crucible, forming a water layer atop the melt.  
436 This produced no force signal, audible cracking, or fragmentation visible on highspeed video.  
437 Only when injected gas initiated deformation and fragmentation did water intrude into the  
438 opening cracks and produce IFCI.  
439 To separately sample leading ejecta, mostly from IFCI, we modified some runs with a U-  
440 shaped 10 cm-diameter steel tube mounted so that one opening was a few centimetres above  
441 the crucible orifice (see Fig. 2). The other end led to a 600 ml bowl of deionized water. In  
442 these “U-tube” runs, small particles (plus water and steam) of the leading ejecta front were  
443 guided into the water bowl. The tube remained fixed until larger fragments of the following  
444 ejecta entered it (~30 ms after fragmentation began), pushing the U-tube upward and  
445 removing it from the particle-ejection path (see Fig. 2c-e; Supplementary Video 1).  
446 Fragments ejected after U-tube separation followed free ballistic trajectories and were  
447 deposited across the whole experimental area (“ground”). U-Tube separation ensured most  
448 fragments in the water bowl were from the leading ejecta front.

449

#### 450 **Analysis of force and pressure signals**

451 Force and pressure signals, their temporal development and the relation between them carry  
452 information about how the melt plug is accelerated and fragmented.<sup>23,26,29</sup>  
453 Technically, for all fragmentation experiments the same initial gas expansion energy was  
454 provided as input. However, due to the stochastic nature of material failure processes and  
455 crack paths<sup>31</sup>, the times when fragmentation is completed, and the gas jet breaks through  
456 differed between each run. Thus, the actual gas-driven energy input before breakthrough  
457 varied. We used the maximum driving pressure  $P_{max}$  as indicator for this effective expansion  
458 work. This energy causes plastic deformation and is (mainly) stored as elastic energy, which  
459 then drives the fragmentation and is released as kinetic energy of the ejecta.<sup>26,29</sup>  
460 In this study we used the maximum recoil force  $F_{max}$  exerted on the crucible as a measure for  
461 the mechanical response of the plug towards the gas expansion with pressure  $P_{max}$ .  
462 While for dry runs we would expect  $F_{max}$  to be correlated with  $P_{max}$ , in case of IFCI runs  
463 rapid thermohydraulic expansion has to be considered as an additional source of mechanical  
464 energy.

465

#### 466 **Particle sampling**

467 Before each run the experimental area was thoroughly cleaned to avoid any contamination.  
468 Particles generated by the fragmentation experiments were collected after each run, using  
469 three different methods:

- 470 • Ballistically transported particles deposited on table and floor (DG and IG, for  
471 notation see also Extended Data Fig. 1a) were retrieved using a vacuum cleaner with micro-  
472 porous paper bags
- 473 • Particles generated in open IFCI runs, which were deposited in water droplets on the  
474 ceiling and walls (IW, see also Extended Data Fig. 1b), were collected using paper tissues,  
475 which were subsequently dried.

476 • Particles collected in “U-tube” runs in the water bowl (IU) were retrieved by using  
477 paper filters and were subsequently dried.

478 We note that it was not possible to avoid particle loss in any of the configurations. In  
479 particular, it is expected that not all of the fine IU and IW particles collected could be  
480 retrieved from the paper tissues and filters. Therefore, the sieving curves of IFCI particles are  
481 biased toward larger grains. The bias is probably not large, given the very low per-particle  
482 mass of the fine ash-grade particles.

483 Glassy vesicular Havre ash grains fit three sub-classes: curvi-planar, angular, and fluidal<sup>4</sup>.

484 The experimental grains of the 4 phi (>64 $\mu$ m and <125 $\mu$ m) sieve fraction were compared  
485 with ash particles of the same size from six Havre seafloor sites, “Nat1” to “Nat6” (Extended  
486 Data Fig. 1c). We exclusively focused on the curvi-planar grains dominant in Havre ash  
487 samples, which can be attributed to the main eruptive phase (layers S1 and S2)<sup>4</sup>.

488

#### 489 **Grain size analysis**

490 All particles collected were sieved at 1 phi intervals down to a grain size of 4phi (64-125 $\mu$ m),  
491 with the smallest particles accumulating in the pan. One of the characteristics of fuel coolant  
492 interaction processes is the increased production of fine ash grains. In order to check if this  
493 effect is also measurable for IFCI on Havre material, the two finest fractions, 4phi (125 $\mu$ m-  
494 64 $\mu$ m) and smaller than 4phi (<64 $\mu$ m) were binned, and their mass  $m_{fine}$  relative to the total  
495 amount of “ash”  $m_{ash}$ , i.e., mass of particles smaller than -1phi (<2mm), was determined as a  
496 ratio. We estimate the accuracies for  $m_{ash}$  and  $m_{fine}$  to be within 1% and 5%, respectively.

497 According to propagation of error, the ratio  $m_{fine}/m_{ash}$  has therefore an estimated uncertainty  
498 of 5.1%.

499

#### 500 **Morphometric analysis**

501 For morphometric analysis, a population of 4phi particles was randomly selected and  
502 mounted on carbon-coated tape, ensuring that the grains did not touch one another.  
503 Backscatter electron scans were produced with a resolution of 2048 x 1536 pixels, using a  
504 Zeiss Sigma® VP FEG scanning electron microscope (SEM). The image of each particle was  
505 next isolated and binarized, resulting in a black and white image showing its silhouette, i.e  
506 the projection of the particle onto an underlying plane. These binarized images were used as  
507 input data for the particle shape analyzer software PARTISAN©<sup>32</sup>. This program was  
508 developed to quantify shapes of particle outlines, based on 5 morphometric systems<sup>33-37</sup>, and  
509 computes 23 dimensionless shape descriptors (of which several are redundantly used in  
510 different systems, sometimes with different labels/names). The shape parameters and  
511 references to their computation are listed in Extended Data Fig. 6.

512 Each data set was tabulated, then tested pairwise, e.g. A-B, A-C, A-D, B-C, B-D, C-D, by  
513 applying a sequence of statistical tests, following a test procedure presented in an earlier  
514 study<sup>38</sup> (see also Extended Data Fig. 7). All tests were performed with the software SPSS©,  
515 selecting a level of significance  $\alpha$  of 5%.

516 First a Levene test<sup>39</sup> was used to determine whether the variances between compared data sets  
517 were homogeneous or not. As a second step, two-tailed pooled variance t-tests<sup>38,39</sup> or  
518 separated variance t-tests<sup>40</sup> were used. We identified parameters of indicated significant  
519 differences, without corrections<sup>41</sup>, and thereby determined types of experiments that  
520 reproduced grains with features *most similar* to the natural Havre ash (for sample sizes and  
521 results of each test see can be downloaded from the PANGAEA depository<sup>42</sup>). This approach  
522 minimises the number of type I errors (indications of differences where there are none).

523

524 In order to verify, for any two data sets that did not show significant differences from one  
525 another in t-tests, a high degree of similarity, we applied equivalence tests (“e-tests”). This

526 method was introduced for image particle analysis with volcanic ash<sup>38</sup> and tests whether the  
527 confidence interval  $C$  of the tested data set lies within a given range  $D$ , specified by the  
528 threshold  $D_{max}$ . For  $C$ , a range of 5% was used. For mathematical details on this method, the  
529 reader is referred to the 2012 study by Dürig et al.<sup>38</sup>. E-tests are based on the pooled Student's  
530 t-function and thus only provide reliable results for data sets with homogeneous variances. E-  
531 test results based on data sets with unequal variances have therefore been omitted and are  
532 displayed as “(...)”, indicating that no statement about possible equivalence can be made (see  
533 Extended Data Fig. 8-10).

534 Ranges of  $D$  were calibrated to specific shape parameters of Havre ash by applying e-tests  
535 pairwise to the natural curvi-planar ash samples, i.e., the samples of Nat1 to Nat6.

536 In these 15 benchmark tests, the threshold values for  $D$  were increased stepwise, starting from  
537 0.01 by a step size of 0.01, until the test indicated a “significant equivalence” in the  
538 respective shape parameter. Variances must be homogeneous for this test, so not every e-test  
539 yielded results for each of the tested shape factors. For all shape parameters, however, at least  
540 one valid threshold value for  $D$  was found. Based on the results of these benchmark tests, the  
541 largest of the up to 15  $D$  values was used as the shape parameter-specific threshold  $D_{max}$  for  
542 testing the experimental grains (Extended Data Fig. 6).  $D_{max}$  can thus be seen as a quantity  
543 which specifies the “natural variation” of the respective parameter. We note, that in this  
544 approach it is implicitly assumed that these data sets originate from the same particle  
545 population, which implies that the curvi-planar ash particles in Nat1 – Nat6 were generated in  
546 the same event.

547

#### 548 **X-ray microtomography ( $\mu$ -CT)**

549 For each sample suite, the 3D external morphology of about 30 particles of the grain fraction  
550 between 125 and 64  $\mu\text{m}$  was determined by X-ray microtomography using a Bruker  
551 Skyscan® 1172 high-resolution  $\mu$ -CT scanner. Particles were cleaned in an ultrasonic bath

552 and mounted on a graphite rod holder using vinyl glue. Particles were scanned with a pixel  
553 size of 0.54  $\mu\text{m}$ , an X-ray voltage of 34 kV, an X-ray current of 210  $\mu\text{A}$ , a rotation step of  
554  $0.46^\circ$ , with no filter and a frame averaging over 5 scans. Bruker's software NRecon<sup>®</sup> was  
555 used to reconstruct the  $\mu\text{-CT}$  projection images into two-dimensional cross sections (slices),  
556 using a smoothing parameter of 1, a ring artefact correction index of 4 and a beam hardening  
557 correction of 36%. The programs CTAn<sup>®</sup> and CTVox<sup>®</sup> (both by Bruker) were used for  
558 particle image segmentation, and for rendering and displaying the 3D objects.

559

### 560 **Volumetric temperature expansion**

561 The change in the volume of a unit under changing temperatures can be expressed as

$$562 \quad \beta = \frac{dV}{V_0 dT}$$

563 where  $\beta$  is the volumetric thermal expansion coefficient,  $dV$  is the change in volume,  $V_0$  is the  
564 initial volume, and  $dT$  is the temperature change. The volumes were computed for pressures  
565 between 0.1 and 10 MPa and temperatures from  $4^\circ\text{C}$  to the boiling point, using the Matlab  
566 toolbox X Steam<sup>43</sup>. The thermal expansion coefficient  $\beta$  was calculated assuming an initial  
567 temperature of  $4^\circ\text{C}$ . A plot showing the resulting values for  $\beta$  is provided in Extended Data  
568 Figure 4. The volume changes for sea water with temperature at 10 MPa presented in Fig. 5  
569 are based on experimental measurements for oceanic temperatures.<sup>28</sup>

### 570 **Data availability**

571 Raw data and results of t-tests, including p-values, t-values and degrees of freedom, are  
572 available on the open access data archive PANGAEA<sup>42</sup> and can be retrieved under  
573 <https://doi.org/10.1594/PANGAEA.908865>. Additional experimental data, including videos  
574 are available from the corresponding author on request.

575

576 **Code availability**

577 MATLAB routines used for data processing and modelling are available from the  
578 corresponding author on request.

579

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