- 1 <u>Title</u>
- 2 Bacterial contact induces polar plug disintegration to mediate whipworm egg hatching
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- 4 Short title
- 5 Whipworm egg structure during bacteria-mediated hatching
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30 Abstract

31 The bacterial microbiota promotes the life cycle of the intestine-dwelling whipworm *Trichuris* by 32 mediating hatching of parasite eggs ingested by the mammalian host. Despite the enormous 33 disease burden associated with Trichuris colonization, the mechanisms underlying this 34 transkingdom interaction have been obscure. Here, we used a multiscale microscopy approach 35 to define the structural events associated with bacteria-mediated hatching of eggs for the murine 36 model parasite *Trichuris muris*. Through the combination of scanning electron microscopy (SEM) 37 and serial block face SEM (SBFSEM), we visualized the outer surface morphology of the shell 38 and generated 3D structures of the egg and larva during the hatching process. These images 39 revealed that exposure to hatching-inducing bacteria catalyzed asymmetric degradation of the 40 polar plugs prior to exit by the larva. Although unrelated bacteria induced similar loss of electron 41 density and dissolution of the structural integrity of the plugs, egg hatching was most efficient in 42 the presence of bacteria that bound poles with high density such as *Staphylococcus aureus*. 43 Consistent with the ability of taxonomically distant bacteria to induce hatching, additional results 44 suggest chitinase released from larva within the eggs degrade the plugs from the inside instead 45 of enzymes produced by bacteria in the external environment. These findings define at 46 ultrastructure resolution the evolutionary adaptation of a parasite for the microbe-rich environment 47 of the mammalian gut.

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53 Introduction

54 Soil transmitted helminths (STH) are parasitic worms that affect nearly 1.5 billion people 55 worldwide (1). The developmental maturation of certain STHs occurs in the gastrointestinal (GI) 56 tract where the parasites encounter trillions of bacteria that are part of the gut microbiota, as 57 exemplified by the whipworm Trichuris. Infection with Trichuris species initiates when 58 embryonated eggs in the environment are ingested, which then hatch in the cecum and large 59 intestine where the bacterial community is most diverse and dense (2-4). In this environment, the 60 hatched larvae then embed themselves in the intestinal epithelium where they remain anchored 61 as they undergo several molts to become sexually reproductive adult worms (3). Heavy worm 62 burden in individuals infected by the human parasite *Trichuris trichiura* is associated with colitis, 63 anaemia, and dysentery (5-8). Single doses of the available anthelmintic drugs for T. trichiura 64 infections display poor efficacy with an average cure rate of less than 50% (9). A better 65 understanding of how *Trichuris* species have adapted to their host may reveal vulnerabilities that 66 can be targeted for intervention.

67 Recent studies using the murine parasite Trichuris muris have revealed a remarkable 68 degree of co-adaptations with the mammalian host and the gut microbiota. Bacteria, which trigger 69 egg hatching in vitro, are necessary for T. muris to establish infection (10-12). In turn, T. muris 70 colonization alters the microbiota composition of mice (13), which we and others have shown is 71 consequential for the host. In a model of inflammatory bowel disease (IBD), we found the type 2 72 immune response to T. muris colonization protects against intestinal inflammation by increasing 73 Clostridiales and reducing Bacteroidales species within the gut microbiota (14). Another murine 74 intestinal parasite Heligmosomoides bakeri also participates in three-way interactions with the 75 host and microbiota including inducing an outgrowth of Clostridiales that attenuates allergic airway 76 inflammation (15, 16). Colonization of indigenous people in Malaysia with T. trichiura is associated 77 with a similar expansion of Clostridiales and reduction in Bacteroidales in the microbiota,

indicating that these relationships are conserved in humans (14). Further, we found that Clostridiales species induce superior egg hatching of *T. muris* and *T. trichiura* compared with Bacteroidales species (17). These observations suggest that *Trichuris* colonization alters the microbiota composition in a manner that may be mutually beneficial for the parasite and mammalian host in certain contexts.

83 Given this dependence on the microbiota, egg hatching may be a point of vulnerability in 84 the parasite lifecycle. Trichuris eggs are ovoid shaped with a multi-layered shell. A membrane-85 like outer vitelline layer covers the entire surface of the egg, and underneath is a middle layer 86 consisting of mainly chitin and a lower lipid layer (18, 19). These eggs have two openings, one at 87 each end, that are blocked by chitinous polar plugs (4). Although these plugs are also 88 multilayered, they have a higher proportion of chitin in its middle layer than the rest of the shell 89 (4). During the hatching process (eclosion), the oral spear in the anterior end of the larva pierces 90 the outer vitelline membrane and exits through a polar plug on one side of the egg (20, 21). The 91 role of bacteria in this process remains obscure. For the Gram-negative bacterial species 92 Escherichia coli, type 1 fimbriae mediate binding to the polar ends of eggs and are necessary for 93 optimal hatching (10). However, Gram-positive species that do not have fimbriae such as 94 Staphylococcus aureus can also induce efficient hatching in vitro (10, 11, 17). It is unclear whether 95 the structural events associated with hatching in the presence of Gram-negative and -positive 96 species are similar.

In this study, we show that although many Gram-positive bacterial species are strong hatching inducers, *Enterococci* fail to trigger hatching. We also show that the ability of *S. aureus* to form high density clusters is associated with superior hatching rates and that binding of this bacterium to the egg is essential. High resolution 3D volume electron microscopy imaging revealed the ultrastructural organization of the eggshell and larva, and identified striking asymmetrical morphological changes to the polar plug regions that occur upon exposure to *E. coli* and *S. aureus*, representative Gram-negative and -positive bacteria that induce hatching. We

104 further showed that the degree of bacterial binding to eggs is directly proportional to the hatching 105 rate yet hatching required bacteria to be metabolically active. Finally, we show that eggs from 106 both *T. muris* and the human pathogen *T. trichiura* harbor chitinase activity and propose that this 107 activity plays a role in hatching by degrading the chitinous polar plug of the egg from within.

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110 **Results**

111 Bacteria display species-dependent effects on *T. muris* egg hatching.

112 To establish conditions for investigating structural changes that occur during T. muris egg 113 hatching, we first investigated the degree to which bacterial taxa differ in their ability to induce 114 hatching. Although previous studies have shown that both the Gram-negative species E. coli and 115 the Gram-positive species S. aureus can induce hatching (10, 11), only a limited number of 116 bacterial taxa have been examined under the same conditions and it was unclear whether the 117 time course differs between bacterial species. We used previously optimized conditions in which 118 bacteria cultures grown to their maximal density are added to embryonated T. muris eggs in a 119 culture well and monitored over time for hatching by light microscopy under aerobic conditions 120 (12) (Fig 1A, S1 Video). We found that E. coli and S. aureus induced comparable levels of T. 121 *muris* egg hatching by four hours post-incubation, while untreated eggs remained unhatched (Fig 122 1B). However, a higher proportion of eggs were hatched in the presence of S. aureus at earlier 123 time points compared with E. coli under these conditions (Figs 1B and E). Pseudomonas 124 aeruginosa and Salmonella enterica Typhimurium – Gram-negative Proteobacteria related to E. 125 coli – were previously shown to induce T. muris egg hatching under similar conditions (10, 17). 126 To increase the number of different Gram-positive bacterial species investigated in this assay, we 127 examined Staphylococcus epidermidis as a member of the same genus as S. aureus, and Bacillus 128 subtilis and Enterococcus faecalis as unrelated taxa. S. epidermidis induced hatching at a similar

rate as *S. aureus*, while *B. subtilis* also induced efficient hatching, albeit at a modestly slower rate. In contrast, hatching failed to occur in the presence of *E. faecalis*. To determine whether this was specific to *E. faecalis*, we tested another member of the genus, *Enterococcus faecium*, and found that it also did not induce hatching (Fig 1C). These differences in hatching rates cannot be explained by the culture media used to grow each bacterial species because *S. aureus* grown in each of the three media types (BHI, TSB, and LB) induced similar levels of hatching across time points (Fig 1D).





138 Fig 1. Bacteria display species-dependent effects on *T. muris* egg hatching.

- (A) Representative light microscopy image of *T. muris* eggs induced to hatch after incubation with
- 140 S. aureus at 37°C for 45 mins. White arrowhead denotes unhatched egg and black arrowhead
- 141 denotes hatched egg.
- (B) Percent of *T. muris* eggs hatched after incubation in aerobic conditions with overnight cultures
- 143 of *E. coli* and *S. aureus*, compared with their respective broth controls determined by light
- 144 microscopy at indicated time points. Colony forming units (CFUs) of each bacterial species added
- to the eggs are indicated on the graphs.
- 146 (C) Percent of *T. muris* eggs hatched after incubation in aerobic conditions with overnight cultures
- 147 of S. epidermidis, B. subtilis, E. faecalis and E. faecium compared with their respective broth
- 148 controls determined by light microscopy at indicated time points. Colony forming units (CFUs) of
- each bacteria added to the eggs are indicated on the graphs.
- 150 (D) Percent of *T. muris* eggs hatched after incubation with *S. aureus* grown overnight in tryptic
- soy broth (TSB), Luria Bertani (LB) broth and Brain Heart Infusion (BHI) broth.
- (E) Percent of *T. muris* eggs hatched at 1 hour after 37°C incubation with a maximum number of
- 153 E. coli, S. aureus and E. faecalis.
- **(F)** Percent of *T. muris* eggs hatched at 1 hour after 37° C incubation with an equal number (~ 10^{8}
- 155 CFU) of *E. coli* and *E. faecalis*.
- 156 (G) Percent of *T. muris* eggs hatched after incubation with *S. aureus* and *E. faecalis* alone or
- 157 together compared with broth controls. For the *S. aureus* + *E. faecalis* condition, bacterial cultures
- 158 were grown separately overnight, and then were mixed the day of the experiment.
- (H) Percent of *T. muris* eggs hatched after incubation in anaerobic conditions with overnight
- 160 cultures of *E. coli* and *S. aureus* compared with their respective broth controls determined by light
- 161 microscopy at indicated time points.
- 162 (I) Number of worms harvested from the caecum per mouse (n = 9-12 mice per group).

(J) Proportion of germ-free and *S. aureus* monocolonized mice that had harbored adult *T. muris*worms after double-dose infection.

165 Data points and error bars represent mean and SEM from 3 independent repeats for (B), (C), (D), 166 (G), and (H). Dots represent a single well and bars show means and SEM from 3 independent 167 repeats for (E) and (F). Dots represent a single mouse and bars show means and SEM from 3 168 independent repeats for (I). Welch's t test was used to compare area under the curve between 169 each condition and its respective media control for (B), (C), and (H). Ordinary one-way analysis 170 of variance (ANOVA) test followed by a Turkey's multiple comparisons test was used to compare 171 AUC of hatching induced by different conditions to each other for (D) and (G). Kruskal-Wallis test 172 followed by a Dunn's multiple comparisons test was used for (E). Mann Whitney test was used 173 for (F) and (I). Fisher's exact test was used to determine whether there was a significant 174 association between gut microbial composition of mice and the presence of adult worms in the 175 cecum for (J).

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177 Consistent with the ability of staphylococci to grow as tightly packed clusters, a higher 178 number of S. aureus (~10⁸ colony forming units; CFUs) were added to eqgs compared with E. *coli* (\sim 5x10⁷ CFUs) in the above experiments in which we normalized based on maximal growth 179 180 in media. Therefore, bacterial density may explain differential *T. muris* egg hatching rates. Indeed, 181 the difference in early hatching was eliminated when we concentrated E. coli to normalize the number of bacteria in the inoculum to match the maximal growth of S. aureus (~10⁸ CFUs) (Fig 182 183 1F). However, increasing the number of *E. faecalis* led to negligible amounts of hatching (Fig 1F). 184 This lack of hatching was not due to the production of an inhibitory or toxic factor by this bacterium 185 because adding *E. faecalis* did not negatively impact *S. aureus*-mediated hatching (Fig 1G). 186 These results show that taxonomically distant Gram-positive and -negative bacteria induce 187 hatching with similar efficiency when the number of bacteria added to the culture are normalized. 188 although there are Gram-positive taxa such as enterococci that are unable to induce hatching.

189 Given that hatching occurs predominantly in the anaerobic environment of the cecum and large intestine (2), we confirmed that E. coli and S. aureus were able to induce hatching in 190 191 anaerobic conditions (Fig 1H). We previously demonstrated that monocolonization of germ-free 192 mice with E. coli is sufficient to support T. muris development following inoculation with 193 embryonated eggs, although the number of worms recovered is less than conventional mice (12). 194 To determine whether a Gram-positive species can support parasite colonization, we 195 monocolonized mice with S. aureus and inoculated them with two doses of 100 T. muris eggs. 196 We recovered adult worms from the cecum from all S. aureus monocolonized mice 42 days after 197 T. muris inoculation, whereas most germ-free control mice did not harbor worms (Figs 1I and J). 198 Together with our prior findings, these results show that E. coli and S. aureus, despite their 199 taxonomic and structural dissimilarities, are individually sufficient to induce egg hatching and 200 promote T. muris colonization.

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202 Physical contact with egg is essential for *S. aureus*-mediated hatching.

203 T. muris egg hatching is impaired in the presence of E. coli when direct contact is inhibited 204 (10, 11). Given the structural and growth differences between E. coli and S. aureus, we revisited 205 contact-dependence of hatching using the above conditions. We initially tested whether S. aureus 206 secretes a soluble factor that is sufficient to induce hatching. Similar to our previous finding with 207 E. coli (12), the addition of filtered S. aureus culture supernatant to T. muris eggs did not result in 208 hatching (Fig 2A). It is possible that the presence of T. muris eggs is required for S. aureus to 209 produce a pro-hatching factor, which we would miss in a simple supernatant transfer experiment. 210 To address this possibility, we incubated S. aureus with eggs, collected and filtered the 211 supernatant from the co-culture, and then added this egg-primed supernatant to a new batch of 212 eggs (Fig 2B). The co-culture supernatant was also unable to induce hatching of eggs (Fig 2C). 213 To further determine whether contact is necessary, we separated eggs by placing them inside a 214 0.4µm transwell insert and added bacteria to the outside well. Although moderate amounts of

- 215 hatching occurred when eggs were segregated from *E. coli* in this manner, no hatching was
- observed when eggs were placed in transwells that were incubated with S. aureus (Fig 2D). These
- findings indicate that direct contact with eggs is required for *S. aureus* to induce hatching.
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Fig 2. Physical contact between the bacterial cell and egg is essential for *S. aureus* mediated hatching.

(A) Percent of *T. muris* eggs hatched after incubation with total overnight *S. aureus* culture, culture
 pellet resuspended in fresh media, culture supernatant filtered with a 0.22μm filter, or media
 control.

(B) Experimental approach for determining whether a soluble hatching inducing factor is produced
by *S. aureus* in response to exposure to eggs. Filtered supernatant from *S. aureus* grown with or
without *T. muris* eggs for 4 h were transferred to a dish containing *T. muris* eggs. The ability of
the supernatant to mediate egg hatching was evaluated over 4 hours.
(C) Percent of *T. muris* eggs hatched after incubation with total overnight *S. aureus* culture or
filtered supernatant obtained from *S. aureus* incubated with or without eggs as in (B) compared

- 250 Intered supernatant obtained norm 0. adreas incubated with or without eggs as in (b) com
- with media controls.

(D) Percent of *T. muris* eggs hatched when placed in a 0.4µm transwell separated from bacteria
or control media in the outer well compared with eggs incubated bacteria without transwell
separation.

Data points and error bars represent the mean and SEM from 3 independent experiments for (A) and (C). Dots represent a single well and bars show means and SEM from 3 independent experiments for (D). Ordinary one-way ANOVA test followed by a Turkey's multiple comparisons test was used to compare resuspended pellet and supernatant conditions with total O/N culture for (A) and (C). Two-way ANOVA followed by a Turkey's multiple comparisons test was used for (D).

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242 Collapse of the polar plug precedes hatching mediated by bacteria.

243 Bacteria-induced T. muris egg hatching has not been examined with ultrastructural 244 resolution. We examined eggs exposed to S. aureus and E. coli by scanning electron microscopy 245 (SEM) and were able to capture eggs at different stages in the hatching process as evidenced by 246 larvae in mid-ejection and morphological changes to the plug not observed in untreated eggs and 247 eggs exposed to *E. faecalis*, a poor hatching-inducing species (Figs 3A-E). For both untreated 248 and *E. faecalis*-treated eggs, the plugs appeared as crater-like structures approximately 5 µm in 249 diameter with an inner surface displaying a slightly rounded wrinkled morphology (Figs 3C and 250 D). Diplococci characteristic of Enterococci were present at the plugs of eggs incubated with E. 251 faecalis (Fig 3C). Untreated eggs, despite lack of hatching, had a high concentration of bacterial 252 cells on the plugs (Fig 3D). Because eggs were harvested from adult worms isolated from the 253 cecum of mice, these bacteria were likely derived from the mouse gut microbiota (13).

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256 Fig 3. Collapse of the polar plug precedes hatching mediated by bacteria.

257 (A, B, C, D) Representative low (left) and high magnification (right) SEM images of eggs (clear 258 arrowhead) that were exposed to S. aureus for 1 hour (A), E. coli for 1.5 hours (B) and E. faecalis 259 for 1 hour (C) or untreated (D). White arrowheads correspond to bacteria on polar plug regions of 260 the eggs denoted by black arrowheads. Yellow arrow in right panel of (A) and (B) indicates woolly 261 substance present among bacteria. Debris on untreated eggs is denoted by the white arrow (D). 262 (E) Representative low (left) and high magnification (right) SEM images of hatching eggs (clear 263 arrowhead) that were exposed to S. aureus for 1 hour (top) and E. coli for 1.5 hours (bottom). 264 White arrowheads correspond to bacteria on polar plug regions of the eggs. Emerging larvae are 265 denoted by white diamonds.

- 266 (F) Number of bacterial cells visible on polar plugs of eggs incubated with *E. coli*, *S. aureus*, or *E.*
- 267 *faecalis*. Bars showing mean from 2 eggs per condition.
- 268 (G) Width of collar openings on eggs that were treated with either *E. coli* or *S. aureus* and were
- either unhatched or in the process of hatching.

For low magnification images, scale bar represents 2μ m for unhatched egg in (A), (B), (C) and (D) and 10μ m for hatched egg in (A) and (B). For high magnification images, scale bar represents 1 μ m for (A), (B) and (D) and 2μ m for (C). Dots represent a single plug and bars show mean and SEM of collar sizes from 4-7 eggs per condition for (G). Two-way ANOVA followed by a Turkey's multiple comparisons test was used for (G).

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276 Eggs incubated with S. aureus and E. coli were enriched for the presence of bacteria at 277 their plugs (Figs 3A and B), which displayed a collapsed morphology distinct from untreated eggs 278 (Fig 3D) and eggs incubated with E. faecalis (Fig 3C). S. aureus and E. coli were densely 279 clustered within or near depressions. Additionally, there was a wooly substance that was closely 280 associated with the bacteria on the plug region (Figs 3A and B, right). The vitelline membrane of 281 the plug was also visible and resembled a deflated balloon (Fig 3B, right). Bacteria, while present 282 sporadically across the eggshell, were more densely packed on the plug region than other 283 regions. Examination of eggs incubated with S. aureus and E. coli in which worms were captured 284 mid-ejection showed that, in both instances, larvae exited the egg through the plug, rupturing it 285 along with the outer vitelline layer in the process (Fig 3E). In these eggs in which larvae were 286 partially out of the shell, bacteria were found near the polar plug region (Fig 3E, top and bottom). 287 We observed more individual bacterial cells per disrupted plug when eggs were incubated

with *S. aureus* than other conditions, likely reflecting their capacity for growing in clusters. However, the number of visible *E. faecalis* cells on the polar plug region was comparable to the number of *E. coli* cells on the structurally distinct polar plug, suggesting that differences in the density of bacterial species at the plug are not sufficient to explain why Enterococci are poor hatching inducers (Fig 3F). Additionally, the diameter of the collar region on eggs in the process of hatching was generally wider than on eggs exposed to bacteria, but that had not yet begun to hatch, consistent with the previously described observation that plugs swell during the hatching

process (21) (Fig 3G). These results show that exposure of eggs to strong hatching-inducing
bacteria is associated with a collapse and loss of structural integrity of the polar plugs.

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298 Hatching-inducing bacteria trigger disintegration of the polar plug

299 Advances in the volume electron microscopy technique serial block face-scanning 300 electron microscopy (SBFSEM) present an opportunity to gain additional insight into the above 301 structural changes that we observed in eggs exposed to S. aureus and E. coli. Preparation of 302 samples for ultrastructural electron microscopy imaging normally involves chemical fixation. 303 staining, dehydration and then finally embedding of the specimen in resin (22). However, the 304 impermeability of the eggshell made preservation of the egg contents using routine chemical 305 fixation and high pressure freezing and freeze substitution methods difficult (23). To overcome 306 this challenge, we used a microwave assisted sample preparation method to increase the 307 penetration of fixative and stains into the egg, which enabled identification of structures 308 corresponding to the eggshell, polar plugs, and the larvae (Figs 4A-C, S2 Video). The eggshell 309 thickness was comparable between both bacteria-treated eqgs, 2.17µm and 2.18µm for E. coli 310 and S. aureus-exposed eggs, respectively (Figs 4A-C). Cells and other structures within the 311 larvae and the cuticle confining the larvae were visible. Characteristic granules of the larval 312 intestinal tract were observed, and we propose that those granules contain lipids based on the 313 low electron density of the contents (Fig 4A, S2 Video) (20). The plug on one end (top) displayed 314 a more granular and less electron dense morphology compared with the contralateral plug 315 (bottom) for both S. aureus- and E. coli-exposed eggs. The anterior end of the larva was closer 316 to the top plug that was granular and pointed towards the bottom plug in both cases (Figs 4A-C, 317 S2 Video). 3D renderings generated from the data collected showed that both S. aureus and E. 318 coli were sporadically present over the entire surface of the egg and enriched at the plugs (Figs 319 4B and C respectively, right; S2 Video). In the case of S. aureus exposed eggs, a large aggregate

- 320 of a characteristic grape-like cluster of cocci was visible on one of the poles, versus the shell
- 321 where most bacteria were present as either single cells or in pairs (Figs 4B, far right).

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(A) Representative electron micrograph of a longitudinal cross-section of a *T. muris* egg exposed
to *S. aureus* for 1 hour. Image shows granules containing lipids (LG) within the larva, eggshell
(ES) and larval cuticle (C). Distances measured are also indicated. The width of the collar opening
(i), width of the widest part of the plug (ii), the height of the plug (distance from the top of the collar
to the top of the plug) (iii) and the eggshell thickness (iv) were measured for all conditions. Insets
show the regions boxed in black dotted line.

(B, C) Representative electron micrographs and 3D reconstructions from SBFSEM data of a *T. muris* egg exposed to *S. aureus* for 1 hour (B) and *E. coli* for 1.5 hours (C). For the longitudinal section, the original micrograph (left), the micrograph with color overlays indicating segmented egg components and bacteria (middle) and 3D reconstruction of the egg (right) are shown. For the complete structure, two different angles are shown (left and right). *S. aureus* (red), *E. coli* (blue), polar plugs (green), eggshell (yellow) and larvae (purple) are all shown. Scale bars represent 10μ m.

(D) High magnification images of polar plugs from (B) and (C) on eggs exposed to *S. aureus* and *E. coli* compared with equivalent regions from an egg untreated with bacteria. Outer vitelline layer
is denoted by white arrowheads and eggshell is denoted by black arrowheads.

341 (E) 3D reconstruction of polar plugs on eggs exposed to *E. coli*. Multiple angles are shown (left to342 right).

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344 For Trichuris eggs not treated with bacteria, we found that the fixative did not penetrate 345 through the intact shell even with the microwave assisted fixation method, precluding structural 346 analysis of the larva. However, we were able to obtain high resolution images of the eggshell and 347 pole regions of untreated eggs for comparison with bacteria-treated eggs. Eggshells were 2.31µm 348 thick, similar to bacteria-treated eggs (Figs 4A-D). Consistent with previous reports analyzing 349 eggs in non-hatching conditions (18, 21), we found that the polar plugs appeared as uniformly 350 electron dense structures rounded towards the outer surface for egg samples that were not 351 exposed to hatching-inducing bacteria (Fig 4D, right). While being rounded, the height of the plug 352 of the untreated egg (Fig 4A) did not extend far beyond the top of the collar on this untreated egg 353 (3.29µm top, 3.49µm bottom. These intact plugs also contained an electron dense outer vitelline 354 layer that was continuous with the rest of the shell and overlayed a similarly electron dense 355 chitinous layer enclosed by an electron dense, ridged collar (Fig 4D, right).

356 In contrast, eggs incubated with S. aureus or E. coli displayed plugs that were 357 morphologically distinct from untreated egg plugs. First, they were much less electron dense and 358 non-uniform in electron density. As noted above, these changes were asymmetric. For both 359 bacteria, one plug completely lost structural integrity and appeared more granular and 360 disintegrated than the contralateral plug on the same egg (Fig 4D). Instead of a smooth rounded 361 surface, these plugs displayed indentations on the surface, which was more pronounced for the 362 E. coli-treated egg in which the plug had a large depression (Figs 4D and E; S2 Video). The 363 contralateral pole was more intact compared with the top pole but was less electron dense 364 compared with plugs on the egg not exposed to bacteria. There were two distinct regions of 365 electron density for these contralateral poles exposed to S. aureus or E. coli, with an outer region 366 of moderate density encompassing an inner less electron dense area (bottom) (Fig 4D, left and 367 middle). In the S. aureus condition, the plug with the disintegrated morphology corresponded with 368 the side associated with the bacterial clusters. Although different in electron density, both plugs 369 on eggs incubated with bacteria were swollen and extended beyond the width of the collar region. 370 Specifically, plugs on *E. coli* exposed eggs were 9.51µm and 10.43µm in width while their 371 respective collars were $8.87\mu m$ and $8.68\mu m$ in diameter for the top and bottom, respectively (Fig. 372 4D). Similarly, plugs on S. aureus exposed eggs were 9.46µm and 10.39µm wide while their 373 respective collars were 8.62µm and 9.46µm wide (Fig 4D). Moreover, the heights of the plugs 374 were larger on eggs exposed to E. coli and S. aureus (4.47um top and 7.40um bottom, and 375 $5.63\mu m$ top and $6.51\mu m$ bottom, respectively) than on the untreated egg (Fig 4D).

We repeated the SBFSEM experiment with new bacteria-exposed egg samples to determine whether we can capture additional stages of plug disintegration. In this second run of SBFSEM on *S. aureus*-exposed eggs, one of the plugs (pole 1, S1 Fig A) displayed similar morphology as the contralateral (bottom) plug on the *S. aureus*-exposed egg in Fig 4D, while the other plug (pole 2, S1Fig A) had a similar electron density to the outer region of the pole 1 plug

but was uniformly electron dense. Additionally, the second run of SBFSEM on *E. coli*-exposed eggs showed plugs with a uniform and high electron density that were not swollen, similar to the plugs seen on eggs that were not treated with bacteria (S1 Fig B). Lastly, in the second run on *S. aureus* exposed eggs, we observed that the larva appeared to be directly interacting with the inner surface of pole 1 (S1 Fig C). This result could be representative of additional and earlier stages of plug degradation.

Together, these results indicate that exposure to hatching inducing bacteria is associated with substantial morphological changes of the polar plugs, including swelling and a decrease in structural integrity.

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391 Degree of bacterial binding is proportional to the rate of *T. muris* egg hatching.

392 The strict requirement of physical binding for S. aureus-induced hatching and the striking 393 images showing that this bacterial species was densely bound to the plug motivated us to 394 quantitatively define the nature of this interaction. Towards this end, we used spinning-disk 395 confocal microscopy to quantify the location and number of bacteria by incubating eggs with a S. 396 aureus strain that expressed GFP (Fig 5A). At 30 minutes post-incubation, we found that the pole region contained a substantially higher proportion of GFP+ structures compared with the sides of 397 398 the eggs (Fig 5B). Our cross-species comparisons (Fig 1) suggest that hatching efficiency is 399 dependent on the number of bacteria incubated with the eggs. Consistent with this observation, 400 when we added serial dilutions of *S. aureus* to eggs, we observed that the rate of hatching was 401 proportional to the concentration of bacteria (Fig 5C). To test whether this relationship between 402 bacterial concentration and hatching rates reflected binding events, we quantified the number and 403 location of bacteria associated with eggs incubated with dilutions of S. aureus-GFP. The 404 percentage of eggs associated with GFP+ puncta and the number of GFP+ puncta per egg after 405 a 30-minute incubation were proportional to the concentration of S. aureus added to the culture 406 (Figs 5D-F). In contrast to the higher concentration conditions in which almost all GFP+ structures

407 were associated with poles, the proportion of *S. aureus* bound to the poles versus other parts of 408 the eggshell was reduced at lower concentrations (Figure 5G). These results are consistent with 409 the ultrastructural analyses, suggesting that the degree of *S. aureus* binding to the polar plug 410 region determines the rate of hatching.

411



Fig 5. Degree of bacterial binding is directly proportional to the rate of *T. muris* egg hatching.

- 415 (A) Representative confocal microscopy image of eggs incubated with S. aureus GFP for 30
- 416 minutes. Regions defined as poles and sides are indicated. Scale bar represents $34\mu m$.
- 417 **(B)** Percent of GFP+ puncta present on the poles versus the sides of the eggs.
- 418 (C) Percent of *T. muris* eggs hatched after incubation with 10-fold dilutions of overnight *S. aureus*
- 419 culture ranging from approximately $10^5 10^8$ CFU (n = 3).
- 420 (D) Percent of eggs associated with GFP+ puncta after incubation with 10-fold dilutions of
- 421 overnight S. aureus-GFP culture ranging from approximately $10^5 10^8$ CFU.
- 422 (E) Correlation analysis comparing log₁₀ CFUs of bacteria used and percent of eggs with GFP+
- 423 puncta present.
- 424 **(F)** Number of GFP+ puncta bound per egg from (D).
- 425 (G) Percent of GFP+ puncta present on the poles of the eggs versus the sides of the eggs from426 (D).
- 427 Bars and error bars show means and SEM from 3 independent repeats for (B), (D), (F) and (G).

Data points and error bars represent mean and SEM from 3 independent repeats for (C). Dots represent percentage of 40 eggs that were GFP+ from a single experiment for (D) and (E) and number of GFP+ puncta found on a single egg for (F). Mann-Whitney test was used for (B). Kruskal-Wallis test followed by a Dunn's multiple comparisons test was used for (D) and (F). Simple linear regression was performed for (E). Two-way ANOVA followed by a Sidak's multiple comparisons test was used for (G).

434

435 Bacterial metabolic activity is essential for *S. aureus*-mediated hatching.

Egg hatching was low or completely absent within the first 30 minutes post-incubation with *S. aureus* (Figs 5C and 6A). Given our results showing a direct relationship between bacterial binding and hatching rate, it is possible this delay in hatching reflects the time required for bacteria to bind eggs. However, even at 0 minutes post-incubation with *S. aureus*-GFP, the majority of eggs were bound by bacteria based on their association with GFP+ puncta (Figure 6B). By 10

441 minutes post-incubation, the proportion of eggs bound to bacteria and the number of bacterial 442 cells bound to each egg reached the maximum value (Figs 6B and C). *S. aureus* was consistently 443 enriched at plugs, and the ratio of bacteria bound to plugs versus other regions of the shell was 444 similar across time points (Fig 6D). These findings indicate that bacterial binding is not the rate 445 limiting step for hatching induction.



(A) Percent of *T. muris* eggs hatched after incubation with 10⁸ CFU overnight *S. aureus*culture at time points indicated.

(B) Percent of eggs that had GFP+ puncta bound after incubation with 10⁸ CFU of overnight S.

452 *aureus* culture for different lengths of time.

453 **(C)** Number of GFP+ puncta bound per egg after incubation with 10⁸ CFU of overnight *S. aureus*

454 culture for different lengths of time.

(D) Percent of GFP+ puncta present on the poles of the eggs versus the sides of the eggs after
 incubation with 10⁸ CFU of overnight *S. aureus* culture for different lengths of time.

457 (E) Percent of *T. muris* eggs hatched after incubation with UV killed overnight *S. aureus* culture.

458 (F) Percent of *T. muris* eggs hatched after incubation with untreated *S. aureus*, chloramphenicol

459 (CAM)-treated *S. aureus* (100μg/ml), and CAM-treated *S. aureus* together with CAM-resistant *S.*

460 *aureus* GFP for 4hrs. *S. aureus* GFP was spiked-in at the 2hr time point.

461 Bars and error bars show means and SEM from 3 independent repeats for (B), (C), and (D). Data

462 points and error bars represent mean and SEM from 3 independent repeats for (A), (E) and (F).

463 Dots represent mean percentage of 40 eggs that were GFP+ from a single experiment for (B) and

464 number of GFP+ puncta found on a single egg for (C). Kruskal-Wallis test followed by a Dunn's

466 multiple comparisons test was used for (D). Ordinary one-way ANOVA followed by Turkey's

multiple comparisons test was used for (B) and (C). Two-way ANOVA followed by a Sidak's

467 multiple comparisons test was used to compare AUC of different conditions tested for (F).

468

465

We previously showed that hatching required metabolically active *E. coli*, specifically arginine biosynthesis (12). Consistent with a requirement for viability, UV-killed *S. aureus* cells failed to induce egg hatching (Fig 6E). Also, the bacteriostatic antibiotic chloramphenicol (CAM), which inhibits protein translation in *S. aureus* (24, 25), reduced bacteria-mediated hatching (Fig 6F). This impaired hatching was not due to an effect of CAM on the egg itself because spiking-in

474 *S. aureus*-GFP that is CAM-resistant at 2 hours after the beginning of the assay immediately 475 restored hatching in the presence of CAM (Fig 6F). In conclusion, egg binding and metabolic 476 activity are both required for *S. aureus* to mediate hatching.

477

478 *Trichuris* eggs harbor chitinase activity.

479 The biochemical process involved in *Trichuris* egg hatching is unknown. Hatching of eggs 480 from the STH Ascaris lumbricoides is mediated by a parasite-derived chitinase enzyme (26). 481 However, A. lumbricoides hatching occurs independently of bacteria. Thus, we explored the 482 possibility that bacteria provided the chitinase for *T. muris* eggs to hatch. *S. aureus*, for example, 483 produces degradative enzymes that have chitinase activity (27). To test this hypothesis, we used 484 a S. aureus mutant with a deletion in a gene encoding a chitinase-related protein, which we refer 485 to as CRP. Hatching occurred at a similar rate in the presence of S. aureus Δcrp and wild-type, 486 indicating that CRP is not required (Fig 7A). In fact, S. aureus mutants deficient in all the major 487 secreted proteases (S. aureus \(\triangle aur\(\triangle sspAB\)\(\triangle scpAspl::erm\) (28) as well as a S. aureus mutant 488 lacking the three major lipases (S. aureus $\Delta gehA \Delta gehB \Delta gehE$) (29) retained the ability to induce 489 hatching with similar efficiency as wild-type bacteria (Figs 7B and C). These findings raise the possibility that the enzymatic activity responsible for the disintegration of the poles is derived from 490 491 the parasite rather than bacteria.



494 Fig 7. Trichuris eggs harbor chitinase activity.

- 495 (A-C) Percent of *T. muris* eggs hatched after incubation with overnight *S. aureus* WT, *S. aureus*
- 496 \triangle crp (A), S. aureus \triangle aur \triangle sspAB \triangle scpAspl::erm (B) and S. aureus \triangle gehA \triangle gehB \triangle gehE (C) culture.
- 497 (D, E) Amount of fluorescence detected in wells containing eggs hatched in response to S. aureus
- 498 (D) and *E. coli* (E). AU = arbitrary unit. 5mU of stock chitinase were used as a positive control.
- 499 (F) Amount of fluorescence detected in wells containing crushed eggs that were exposed to S.
- 500 *aureus* for 20 minutes. 0.02mU of stock chitinase were used as a positive control.
- 501 **(G)** Amount of fluorescence detected in wells containing crushed *T. muris* and *T. trichiura* eggs.
- 502 0.02mU of stock chitinase were used as a positive control.
- (H) Percent of *T. muris* eggs hatched after incubation with fluid from hatched eggs (purple open circles) and bacteria resuspended in fluid from hatched eggs (purple filled circles). Fluid from unhatched eggs was used as a control (yellow filled and open circles).
- 506 (I) Graphical depiction of proposed mechanism.
- 507 Data points and error bars represent means and SEM from 3 independent repeats for (A), (B),
- 508 (C) and (H). Dots represent fluorescence detected in a single well and bars and error bars show
- 509 means and SEM from 2-3 independent repeats for (D), (E), (F) and (G). Welch's t test was used
- 510 to compare area under the curve between mutant S. aureus strains and WT S. aureus for (A), (B)
- and (C). Kruskal-Wallis test followed by a Dunn's multiple comparisons test was used for (D), (E),
- 512 (F) and (G). Ordinary one-way ANOVA followed by Turkey's multiple comparisons test was used
- 513 to compare AUC of different conditions tested for (H).
- 514

515 Chitinase is released into the media during *A. lumbricoides* egg hatching (26). If *T. muris* 516 also uses chitinase to degrade the polar plug from the inside, we should be able to detect chitinase 517 activity in the post-hatching fluid, similar to *A. lumbricoides*. Using a previously described assay 518 in which a substrate produces a fluorescent product when cleaved by chitinase (30), we detected 519 chitinase activity in media from eggs incubated with *S. aureus*, but not in media containing 520 untreated eggs or *S. aureus* alone (Fig 7D). Release of chitinase was not specific to *S. aureus*-

521 mediated hatching as chitinase activity was also detected in media from eggs incubated with E. 522 coli (Fig 7E). To determine whether chitinase within T. muris eggs is induced by exposure to 523 bacteria, we measured chitinase activity in crushed eggs with or without pre-incubation with S. 524 aureus. Chitinase activity was detected in the media containing crushed eggs that were exposed 525 to bacteria as well as in eggs that were not exposed to bacteria (Fig 7F). Thus, eggs harbor 526 chitinase activity independent of bacteria treatment. This experiment also rules out the possibility 527 that bacteria were the source of the chitinase activity. We detected chitinase from crushed T. 528 trichiura eggs (Fig 7G), indicating that the human parasite also produces this enzyme.

Lastly, we tested whether media containing chitinase released from *T. muris* eggs induce hatching of intact eggs from the external environment ("outside-in"). Fluid from hatched eggs was obtained by incubating eggs with bacteria for 4hrs and then filtering the supernatant. Fluid obtained from unhatched eggs was included as a negative control. We observed that fluid from hatched eggs did not induce hatching and was also unable to enhance *S. aureus*-mediated hatching (Fig 7H). These results suggest that enzymes likely mediate hatching from the inside rather than from the outside (Fig 7I).

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538 **Discussion**

In this study we illuminated structural events that occur during bacteria-mediated hatching of *T. muris* eggs by using multiscale microscopy techniques. Notably, we overcame technical challenges to apply SBFSEM to generate high resolution 3D images of eggs in the process of hatching. This technique along with SEM showed that, despite being taxonomically distant species with significant structural and biological differences, *E. coli* and *S. aureus* induce similar morphological changes in the polar plugs of eggs. These structural changes include swelling of

the plugs, and disintegration of the plug material, resulting in loss of structural integrity of the plugsthat appear as dips and depressions on the plug surface.

547 A previous study describing the egg hatching cascade showed that prior to hatching, the 548 head of the larva inserts into the plug space and then uses its stylet to tear through the vitelline 549 membrane of the plug and exit the egg (21). Therefore, it is possible that bacteria-induced plug 550 disintegration plays an important role in hatching by either hollowing out the space in the collar 551 region to accommodate the larval head or by reducing the density of the plug to facilitate 552 movement into the plug space. The difference in electron density we observed between 553 contralateral plugs on the same egg might be indicative of plugs at different stages of degradation, 554 with the less dense plugs being further along in the disintegration process. This model is 555 supported by our comparison of plugs from different eggs, which highlighted the heterogenous 556 degrees of collapse and granularity. The fact that a loss of electron density is observed during 557 hatching induction by two different bacterial species increases our confidence that dissolution of 558 the integrity of the polar plugs is an essential step during hatching. A mechanism in which 559 disintegration of the poles is rate-limiting and occurs in an asynchronous manner would explain 560 why bacteria-induced egg hatching is also asynchronous.

561 The strict requirement for physical binding between S. aureus and T. muris eggs, and the 562 general absence of information regarding how Gram-positive bacteria mediate hatching, 563 motivated us to further investigate hatching of eggs exposed to this bacterium by confocal 564 microscopy. Although the number of bacterial cells bound to the poles was a critical determinant 565 of hatching efficiency, physical association was not sufficient. Also, although we observed E. 566 faecium bound to egg poles, the two Enterococcus species we tested failed to induce significant 567 amounts of hatching. Similar to our previous findings that bacterial byproducts were necessary 568 for E. coli-mediated hatching (12), S. aureus metabolic activity contributed to hatching. These 569 results support a requirement for both bacterial binding and bacterial products.

570 Given that the polar plug predominantly consists of chitin, we initially considered the 571 possibility that binding led to local production of chitinase derived from bacteria, thereby degrading 572 the pole from the outside. However, hatching remained efficient in the presence of S. aureus 573 mutants lacking candidate enzymes including a homolog of chitinase. Additionally, fluid collected 574 from hatched eggs was unable to induce hatching of unexposed eggs, suggesting that polar plug 575 disintegration is occurring via an "inside-out" mechanism where the larva itself produces the 576 enzymes that facilitate eclosion (Fig 7I). Consistent with this possibility, we detected chitinase 577 within the eggs. A mechanism by which the larva within the egg is the producer of degradative 578 enzymes would explain why unrelated bacterial taxa can induce hatching – it would be unlikely 579 that all the bacterial species shown to induce hatching possess the same enzyme necessary to 580 dissolve the chitinous plug. This model would be further supported if a conserved bacterial product 581 were shown to induce hatching. Such a factor has not been identified. The lack of a genetic 582 system to interrogate T. muris and the impermeable nature of the eggshell also pose challenges 583 for elucidating detailed mechanism. Therefore, technical advances may be required to confirm 584 the role for worm-derived chitinase and examine how products released from bacteria relate to 585 this chitinase activity. Although bacteria exposure is not required for the presence of chitinase 586 activity, it may influence the location of the enzymatic activity (Fig 7I). It is possible, for instance, 587 that bacterial exposure triggers the release of chitinase from vesicles within the worm and into 588 the perivitelline space in the egg, as suggested for *H. bakeri* (31).

Gram-positive species are a prominent component of the gut microbiota and certain taxa are enriched in *Trichuris*-colonized individuals (17). *S. aureus* and other staphylococci colonize the gut of humans, especially early in life as the microbiota develops (32, 33). It is possible that microbiota composition contributes to the susceptibility of young children to high worm burden and disease. Differences in microbiota composition may also partially explain why helminth infections tend to be aggregated in the host population, with a minority of the population harboring a majority of the worms (34-36). Our findings indicate that, although not all bacteria induce

596 hatching of *T. muris* eggs, those that mediate hatching cause similar structural events to occur at 597 the polar plug. We suggest that targeting enzymatic activity involved in plug disintegration would 598 be difficult if it initiates from within an impermeable eggshell that evolved to be resilient to the 599 external environment. Instead, understanding the chemical and physical interaction between 600 bacteria and eggs may uncover a vulnerability in the life cycle of the parasite that is suitable for 601 intervention.

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- 603

604 Materials and methods

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606 Experimental model details

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608 Parasite maintenance

Stock eggs of *Trichuris muris* E strain (14) were propagated and maintained in the NOD.Cg-*Prkdc*^{scid}/J (Jax) mouse strain as previously described (37). Each egg batch was confirmed to hatch at \geq 90% *in vitro* using method below and WT *S. aureus* before use in subsequent experiments.

Stock eggs of Trichuris trichiura were provided by the Trichuris trichiura egg Production 613 614 Unit (TTPU) located at the Clinical Immunology Laboratory at the George Washington University. 615 Whipworm eggs were isolated from the feces of a chronically infected human volunteer following 616 a qualified standard procedure that includes a modified Simulated Gastric Fluid (SGF) method. 617 After isolation, the eggs were stored for two months in flasks containing sulfuric acid (H_2SO_4) 618 maintained at 25-30°C in a monitored incubator. Once embryonation was achieved, the eggs 619 were transferred to a locked and monitored refrigerator at 2-8°C until further use. Controls for the 620 manufacturing process involved: (i) tests for viability (hatching), which confirmed that more than

621 80% of the eggs were viable; (ii) species confirmation by polymerase chain reaction (PCR); and
622 (iii) evaluation of the microbiological burden determined by bioburden testing by an outside623 certified laboratory.

624

625 Bacterial Strains

626 Escherichia coli strain used was a kanamycin resistant transformant of strain BW25113 627 kindly provided by E. Jane Hubbard (NYU Grossman School of Medicine) (12). Staphylococcus 628 aureus strain USA300 LAC clone AH1263 (38), S. epidermidis (ATCC12228) and Enterococcus 629 faecalis OG1RF were provided by V. J. Torres (NYU Grossman School of Medicine) and were 630 originally sourced from Alex Horswill (University of Colorado Anschutz Medical Campus), Eric 631 Skaar (Vanderbilt University) and Lynn Hancock (University of Kansas) respectively. AH1263 was 632 used as wild-type (WT) S. aureus unless otherwise specified. Bacillus subtilis was sourced from 633 the ATCC (ATCC 6633). E. faecium Com15 was kindly provided by Howard Hang (Scripps 634 Research). Frozen glycerol stocks (30% glycerol) of all bacteria were prepared. Glycerol stocks 635 of E. coli was streaked onto Luria Bertani (LB) kanamycin 25 mg/ml plates (NYU Reagent 636 Preparation Core), S. aureus, S. epidermidis and B. subtilis were streaked onto Tryptic Soy Agar 637 (TSA) plates (NYU Reagent Preparation Core) and Enterococcus species were streaked onto bile 638 esculin azide (BEA) plates (Millipore) and incubated overnight at 37°C. Single colonies of E. coli, 639 Staphylococcus species and Enterococcus species were then spiked into 5mls of LB broth, 640 Tryptic Soy Broth (TSB) and Bacto Brain Heart Infusion (BHI) broth (BD) respectively unless 641 otherwise specified and grown overnight at 37°C with shaking at 225 rpm. B. subtilis was also 642 spiked into TSB and was grown overnight under the same conditions. To quantify colony forming 643 units, we performed serial dilutions of liquid culture in sterile PBS and plated on LB (Kan) agar for 644 E. coli, TSA for Staphylococcus species and B. subtilis and BEA for Enterococcus species.

645

646 S. aureus mutants

647 The S. aureus $\Delta aur\Delta sspAB\Delta scpAspl::erm$ mutant. S. aureus $\Delta gehA\Delta gehB\Delta gehE$ mutant. 648 S. aureus Acrp mutant (and its parental strain LAC (ErmS)) and S. aureus GFP strain were all 649 provided by V. J. Torres (NYU Grossman School of Medicine) and were originally sourced from 650 A. Horswill (University of Iowa) (28), Francis Alonzo (University of Illinois at Chicago), Lindsey 651 Shaw (University of South Florida), and the V. Torres lab respectively. S. aureus GFP strain used 652 was a chloramphenicol resistant transformant of strain AH-LAC USA300 transformed with a pOS1 653 vector containing superfolder gfp with a sarA promoter (39). Frozen glycerol stocks (30% 654 alvcerol) of all bacteria were prepared. Glycerol stocks of S. aureus \(\Delta aur\(\Delta sspAB\)\) scpAspl::erm, 655 S. aureus \triangle geh $A \triangle$ geh $B \triangle$ gehE mutant, S. aureus \triangle crp mutant and S. aureus GFP were streaked 656 onto TSA plates with 5 µg/ml of erythromycin, 25 µg/ml Kan/ 25 µg/ml neomycin (Neo), 50 µg/ml 657 Kan/ 50 µg/ml Neo and 10 mg/ml of chloramphenicol respectively (NYU Reagent Preparation 658 Core and V. Torres lab). Single colonies of all species were spiked into 5mls of TSB and grown 659 overnight at 37°C with shaking at 225 rpm. To quantify colony forming units, we performed serial 660 dilutions of liquid culture in sterile PBS and plated on TSA plates.

661

662 Germ-free Mice

663 Germ-free (GF) C57BL/6J mice were bred and maintained in flexible-film isolators at the 664 New York University Grossman School of Medicine Gnotobiotics Animal Facility. Absence of fecal 665 bacteria was confirmed monthly by evaluating the presence of 16S DNA in stool samples by 666 gPCR as previously described (40). Mice were transferred into individually ventilated Tecniplast 667 ISOcages for infections to maintain sterility under positive air pressure. Female mice 6-11 weeks 668 of age were used for all experiments in this study. All animal studies were performed according 669 to protocols approved by the NYU Grossman School of Medicine Institutional Animal Care and 670 Use Committee (IACUC) and Institutional Review Board.

671

672 Method details

673

674 In vitro hatching of T. muris eggs

675 T. muris eggs were hatched in vitro by mixing 25 µL of embryonated eggs at a 676 concentration of 1 egg/1 µL suspended in sterile water with 10 µL of overnight bacterial culture 677 and 15 µL sterile media in individual wells of a 48 well plate. Media control wells contained an 678 additional 10 μ l of sterile media instead of bacterial culture. Plates were incubated at 37°C and observed every 10 mins, 30 mins and/or hour for 4-5 hrs on the Zeiss Primovert microscope to 679 680 enumerate the percentage of hatched eggs by counting hatched and embryonated unhatched 681 eggs in each well. Unembryonated eggs, which lack visible larvae and have disordered contents, 682 were excluded due to their inability to hatch. For anaerobic hatching specifically, plates were 683 incubated at 37°C in an anaerobic chamber and a separate 48-well plate was used per timepoint 684 as the plate needed to be removed from the anaerobic chamber to count hatching each timepoint. 685 Experiments utilizing transwell inserts (Millicell) were performed as previously described (11). For 686 experiments where cell-free supernatant was used, supernatant and cells were isolated by 687 centrifugation and filtration through a 0.22 µm syringe filter or after wash with autoclaved PBS, 688 respectively. Incubation with embryonated eggs was performed by adding 30 µL T. muris eggs to 689 5 mL of bacterial culture and incubating at 37°C for four hours. E. coli and E. faecalis were 690 concentrated by centrifuging 4 ml of each and then resuspending the pellets in 400 μ l of their 691 respective media. For the co-incubation experiment, 2 ml of S. aureus overnight culture was 692 added to 2 ml of *E. faecalis* overnight culture and then the mixture was added to eggs.

693

694 *T. muris in vivo* infection in germ-free mice

Female germ-free C57BL/6J mice were monocolonized at 6–11 weeks of age by oral gavage with $\sim 1 \times 10^9$ colony forming units (CFU) of *S. aureus*. A subculture of *S. aureus* was made by diluting overnight culture 1:100 in TSB and then incubating at 37°C while shaking at 225 rpm for ~ 4 hrs, until 1×10^9 CFU/mL was reached. 5ml of subculture was pelleted by

699 centrifugation at 3480 rpm for 10 min and washed once with sterile 1x PBS. Pellets were then 700 resuspended in 500 μ L sterile 1x PBS and mice were inoculated by oral gavage with 1x10⁹ CFU in 701 a volume of 100 μ L. Inoculum was verified using dilution plating of alignots.

7 and 28 days later, mice were infected by oral gavage with ~100
embryonated *T. muris* eggs. 14 days after the second gavage of *T. muris* eggs, mice were
euthanized, and individual worms were collected and enumerated from the cecal contents of mice.

705

706 Electron microscopy

707 Scanning electron microscopy

708 T. muris eggs were incubated with S. aureus or E. faecalis for 1hr, E. coli for 1.5 hr or 709 bacteria free media for 1 hr. All eggs were fixed with 2% paraformaldehyde (PFA), 2.5% 710 glutaraldehyde in 0.1 M sodium cacodylate buffer (CB, pH 7.2) at 4°C for a week. ~50 eggs were 711 then loaded into 12 mm, 0.44 mm transwells (#PICM01250, Millipore Sigma) to avoid losing eggs 712 during sample processing. The eggs were washed 3 times with PBS, post fixed with 1% osmium 713 tetroxide (OsO₄) in aqueous solution for 1 hour, then dehydrated in a series of ethanol solutions 714 (30%, 50%, 70%, 85%, 95%) for 15 mins each at room temperature. Eggs were then dehydrated 715 with 100% ethanol 3 times for 20 mins each. The eggs were critical point dried using the Tousimis 716 Autosamdri®-931 critical point dryer (Tousimis, Rockville, MD), put on the SEM stabs covered 717 with double sided electron conducted tape, coated with gold/palladium by the Safematic CCU-718 010 SEM coating system (Rave Scientific, Somerset, NJ), and imaged with the Zeiss Gemini300 719 FESEM (Carl Zeiss Microscopy, Oberkochen, Germany) using secondary electron detector (SE₂) 720 at 5 kV with working distance (WD) 18.3 mm.

721

722 Serial block face-scanning electron microscopy

723 *T. muris* eggs were incubated with *S. aureus*, *E. coli* or bacteria-free media and fixed as 724 described above. For further sample processing, we adopted a previously described protocol

(41) and made modifications based on personal communications with Rick Webb (Queensland
University) using a PELCO Biowave (Ted Pella Inc., Redding, CA) for microwave processing. The
detailed sample processing steps are listed in table 1. Samples were embedded with Spurr resin
(Eletron Microscopy Sciences, Hatfield, PA) between ACLARE using a sandwich method.

729 For SBF-SEM imaging, the sample block was mounted on an aluminum specimen pin 730 (Gatan, Pleasanton, CA) using silver conductive epoxy (Ted Pella Inc.) to electrically ground the 731 block. The specimen was trimmed again and sputter coated with 10 nm of gold (Rave Scientific, 732 Somerset, NJ). Serial block face imaging was performed using a Gatan OnPoint BSE detector in 733 a Zeiss Gemini 300 FESEM equipped with a Gatan 3View automatic microtome unit. The system 734 was set to cut sections with 100 nm thickness, imaged with a pixel size of 12 nm and a dwell time 735 of 3.0 µs/pixel, with each frame sized at 50 x 90 µm. SEM beam acceleration was set at 1.5 keV 736 and Focus Charge Compensation gas injection set at 12% (7.9E-04mBar) to reduce charging 737 artifacts. Images of the block face were recorded after each sectioning cycle with a working 738 distance of 6.6 nm. Data acquisition and sectioning were automatically controlled using Gatan 739 Digital Micrograph software to manage imaging parameters. A stack of 250 slices was aligned 740 and assembled using ImageJ. Semi-automated segmentation and video renders were generated with ORS Dragonfly 4.1 (Object Research Systems, Montréal, QC). 741

742

743 Optical microscopy

744 Live Imaging

20 µL of overnight *S. aureus*-GFP culture were added to 50 µL of embryonated eggs (1
egg/1 µL) and 30 µL sterile media in a 35 mm Petri dishes with No. 1.5 coverglass at the bottom.
Mixture was then imaged using a Nikon 40x N.A. 1.3 oil immersion objective lens on a Nikon
Eclipse Ti microscope. An environmental chamber set at 37°C and a lens heater set at 45°C were
both used to observe hatching for approximately 45 minutes.

751 Binding Assay

An *in vitro* hatching assay was prepared as described above using *S. aureus*-GFP. After incubating the plate at 37°C, eggs with bacteria were fixed using 4% PFA, 0.5% glutaraldehyde in PBS at room temperature for 1 hr. Plate contents were then transferred to 1.5 ml Eppendorf tubes. Eggs in this mixture were pelleted by centrifugation at 1000 rpm for 5 mins with a slow brake speed (4). Samples were then washed twice with PBS and then resuspended in PBS for imaging.

Fixed samples were imaged on 35 mm Petri dishes with No. 1.5 coverglass at the bottom using a Nikon 60x N.A. 1.4 oil immersion objective lens on a Nikon Eclipse Ti microscope. Number of puncta on the sides and on the poles were enumerated by eye.

761

762 UV killing of S. aureus

2 ml overnight of *S. aureus* culture were added to 8 ml of sterile PBS in a 150mm Petri dish. The dish was then placed uncovered in a UV Stratalinker 2400 (Stratagene) for 1 hr. Killed bacteria were then transferred to a 15 ml conical tube and then centrifuged at 3480 rpm and 4°C for 10 mins. Pellets were then resuspended in different amounts of sterile TSB to obtain bacterial suspensions of different concentration that were then used in an *in vitro* hatching assay as described above.

769

770 Chloramphenicol treatment of S. aureus

Overnight cultures of *S. aureus* were centrifuged at 3480 rpm and 4°C for 10 mins, resuspended in fresh, sterile TSB and then incubated with 100 μ g/ml of chloramphenicol or 4% ethanol (vehicle control) on ice for 45 mins as previously described (25). Treated bacteria was then used in an *in vitro* hatching assay described above. After 2 hrs of incubation, 10 μ l of untreated chloramphenicol resistant *S. aureus* (*S. aureus*-GFP) was added to the wells of the hatching assay and hatching was monitored for 2 more hours.

777

778 Chitinase detection assay

779 In vitro hatching assay was performed in a 96-well black-walled, clear, flat-bottom plate by 780 adding 12.5 µl of eggs to 7.5 µl of sterile media and 5 µl of overnight bacterial culture and 781 incubating the plate at 37°C for 4 hrs. 6.25 μ l of 4-Methylumbelliferyl β - D-N,N',N'-782 triacetylchitotrioside hydrate (4MeUmb, Millipore Sigma) were then added to hatching assay wells 783 as well as an additional well containing 25 µl of chitinase from Streptomyces griseus (positive 784 control, Millipore Sigma). After incubating at 37°C for 1 hr, reaction was stopped by adding 6.25 785 µl of 1M glycine/NaOH (NYU Reagent Preparation Core) to all wells. Plate was then read at 786 355/450nm ex/em by using a SpectraMax M3 plate reader (Molecular Devices) (30).

787 Chitinase activity within eggs exposed to bacteria was measured by first preparing an *in vitro* hatching assay in a 48-well plate as described above and incubating it at 37°C for 20 mins.
789 16 wells per condition were prepared. Well contents for each condition were then pooled together
790 into 1.5 ml Safe-lock tubes (Eppendorf) with 1.0 mm diameter glass beads (BioSpec Products).
791 Tubes were then homogenized by performing 4, 20 s homogenization cycles at 4.5 m/s using a
792 bead beater. 25 µl of homogenate was then added to a 96-well black-walled, clear, flat-bottom
793 plate and chitinase detection assay was performed as described above.

Chitinase activity within *Trichuris* eggs that were not exposed to bacteria was measured by first centrifuging equal numbers of eggs (500g, 3 mins, brake=3 for *T. trichiura* and 4000rpm, 5 mins, brake=4 for *T. muris*) and then resuspending them in sterile TSB. Eggs were then homogenized and chitinase detection assay performed as described above. In both homogenization experiments, chitinase from *Streptomyces griseus* (Millipore Sigma) was also homogenized as a positive control.

800

801 Determining effect of hatching fluid on eggs

An *in vitro* hatching assay was prepared in a 48-well plate as described above and incubated at 37°C for 4 hrs. 36 wells per condition were prepared. Well contents for each condition were pooled together, centrifuged at 3480 rpm for 10 mins, and then supernatant was filtered as described above and added to new eggs in another *in vitro* hatching assay. Hatching was measured over time.

Additional *S. aureus* overnight cultures were also centrifuged at 3480 rpm for 10 mins and the remaining pellet was resuspended in the same filtered hatching assay fluid obtained above. Resuspended *S. aureus* was then added to new eggs in another *in vitro* hatching assay and hatching was measured over time.

811

812

813 Quantification and statistical analysis

The number of repeats per group is annotated in corresponding figure legends. Significance for all experiments was assessed using GraphPad Prism software (GraphPad). Specific tests are annotated in corresponding figure legends. p values are also annotated on the figures themselves. A p value of ns or no symbol = not significant.

818

819 Table 1. Sample preparation process for SBF-SEM imaging.

Step	Reagent Used	Time	# of	Power	Vacuum
			Repeats	(Watts)	
fixation	2.5% glutaraldehyde in 0.1M CB	2min on/2min off	4 x	100 W	on
washing	0.1M CB	10 min	2 x	on bench	N/A
washing	0.1M CB	40 secs	3 x	100 W	off

post-fix	2% OsO ₄ and 1.5% potassium	2min on/2min off	4 x	100 W	on
	ferrocyanide in 0.1M CB				
washing	ddH ₂ O	10 min	2 x	on bench	N/A
washing	ddH ₂ O	40 secs	3 x	100 W	off
staining	1% aqueous thiocarbohydrazide (TCH)	20 min		on bench	N/A
washing	ddH ₂ O	10 min	2 x	on bench	N/A
washing	ddH ₂ O	40 secs	3 x	100 W	off
staining	2% aqueous OsO ₄	2min on/2min off	4 x	100 W	on
washing	ddH ₂ O	10 min	2 x	on bench	N/A
washing	ddH ₂ O	40 secs	3 x	100 W	off
staining	1% aqueous uranyl acetate	2min on/2min off	4 x	150 W	on
washing	ddH ₂ O	10 min	2 x	on bench	N/A
washing	ddH ₂ O	40 secs	3 x	100 W	off
staining	lead aspartate solution	20 min		60°C	N/A
washing	ddH ₂ O	10 min	2 x	on bench	N/A
washing	ddH ₂ O	40 secs	3 x	100 W	off
dehydration	30%, 50%, 70% ethanol	1min on/1min off/1min	1x each	150 W	off
dehydration	85%, 95%, 100%, 100% acetone	1min on/1min off/1min	1x each	150 W	off
infiltration	acetone : spurr = 1 : 1	3 mins	2x	150 W	off
infiltration	acetone : spurr = 1 : 2	3 mins	2x	150 W	off
infiltration	acetone : spurr = 1 : 2	overnight		on bench	N/A
infiltration	pure spurr w/o NMA	3 mins	2x	150 W	off
infiltration	pure spurr w/o NMA	overnight	2x	on bench	N/A
infiltration	pure spurr w/ NMA	3 hrs	2x	37°C	N/A
embedding	pure spurr w/ NMA	overnight		37°C	N/A
polymerization	pure spurr w/ NMA	72 hrs		60°C	N/A

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946 Supporting information



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948 S1 Fig. Plugs on eggs from bacteria exposed samples show different stages of

949 disintegration, related to Fig 4.

950 (A, B) Representative electron micrograph of a section of polar plugs (black asterisk) on eggs

- 951 exposed to *S. aureus* (left) or *E. coli* (right). Images of Pole 1 (top) and Pole 2 (bottom) were
- collected from the same egg. Outer vitelline layer is denoted by white arrowheads and eggshell
- 953 is denoted by black arrowheads.
- 954 (C) Representative electron micrograph of a section of polar plug (black asterisk) on an egg
 955 exposed to *S. aureus*. Outer vitelline layer is denoted by the white arrowhead and point of contact
- 956 between inner surface of the plug and the larva is denoted by the yellow arrowhead.
- 957
- S1 Video. Live imaging of *T. muris* egg hatching. *T. muris* eggs were incubated with 2 x 10¹⁰
 cfu/ml of *S. aureus* and imaged continuously at 37°C for 45 mins. White arrows indicate eggs
 that hatch. Eggs were imaged using a 40x N.A. 1.3 objective.
- 961

962 S2 Video: 3D reconstruction of a *T. muris* egg incubated with *E. coli* for 1.5 hours at 37°C
 963 from SBFSEM

- 964 Video begins with slices through the egg and bacteria being shown. The 3D reconstruction of
- 965 the larva (purple) and the polar plugs (green) is then revealed. One plug curves inward and the
- 966 other plug appears rounded. Lastly, the reconstructed shell (yellow) and bacteria (blue) are
- 967 shown.
- 968
- 969
- 970